

*Thomas Lengauer (Ed.)*  
**Bioinformatics –**  
**From Genomes to Drugs**  
**Vol. I: Basic Technologies**  
**Vol. II: Applications**

# **Methods and Principles in Medicinal Chemistry**

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**Bioinformatics – From Genomes  
to Drugs**

**Volume I: Basic Technologies**

**Volume II: Applications**

*Edited by Thomas Lengauer*

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## Preface

The present volume of our series “Methods and Principles in Medicinal Chemistry” focuses on a timely topic: Bioinformatics. Bioinformatics is a multidisciplinary field, which encompasses molecular biology, biochemistry and genetics on the one hand, and computer science on the other. Bioinformatics uses methods from various areas of computer science, such as algorithms, combinatorial optimization, integer linear programming, constraint programming, formal language theory, neural nets, machine learning, motif recognition, inductive logic programming, database systems, knowledge discovery and database mining. The exponential growth in biological data, generated from national and international genome projects, offers a remarkable opportunity for the application of modern computer science. The fusion of biomedicine and computer technology offers substantial benefits to all scientists involved in biomedical research in support of their general mission of improving the quality of health by increasing biological knowledge. In this context, we felt that it was time to initiate a volume on bioinformatics with a particular emphasis on aspects of designing new drugs.

The completion of the human genome sequence, published in February 2001, marks a historic event, not only in genomics, but also in biology and medicine in general. We are now able to read the text; but we understand only minor parts of it. “Making sense of the sequence” is the task of the coming years. Bioinformatics will play the leading role in this field, in understanding the regulation of gene expression, in the functional description of the gene products, the metabolic processes, disease, genetic variation and comparative biology. Correspondingly, the publication of this book is “just in time” to jump into the post-genomic era.

Basically, there are two ways of structuring the field of bioinformatics. One is intrinsically by the type of problem that is under consideration. Here, the natural way of structuring is by layers of information that are compiled, starting from the genomic data. The second is extrinsically, by the application scenario in which bioinformatics operates and by the type of molecular biology experiment that it supports. This new contribution to bioinformatics is roughly structured according to this view. The wealth of

information bundled in this volume necessitated a subdivision into two parts.

The intrinsic view is the subject of Part 1: it structures bioinformatics in methodical layers. Lower layers operate directly on the genomic text that is the result of sequencing projects. Higher layers operate on higher-level information derived from this text. Accordingly, Part 1 discusses sub-problems of bioinformatics that provide components in a global bioinformatics solution. Each chapter is devoted to one relevant component: after an introductory overview, Chapters follow that are devoted to Sequence Analysis (*written by Martin Vingron*), Structure, Properties and Computer Identification of Eukaryotic Genes (*by Victor Solovyev*), Analyzing Regulatory Regions in Genomes (*by Thomas Werner*), Homology-Based Protein Modeling in Biology and Medicine (*by Roland Dunbrack*), Protein Structure Prediction and Applications in Structural Genomics, Protein Function Assignment and Drug Target Finding (*by Ralf Zimmer and Thomas Lengauer*), Protein–Ligand Docking and Drug Design (*by Matthias Rarey*) and Protein–Protein and Protein–DNA Docking (*by Mike Sternberg and Gidon Moont*). An appendix *by Thomas Lengauer*, sketching the algorithmic methods that are used in bioinformatics, concludes this first Part.

The extrinsic view is the focus of the second Part: Chapters concentrate on several important application scenarios that can only be supported effectively by combining components discussed in Part 1. These Chapters cover Integrating and Accessing Molecular Biology Resources (*by David Hansen and Thure Etzold*), Bioinformatics Support of Genome Sequencing Projects (*by Xiaoqiu Huang*), Analysis of Sequence Variations (*by Christopher Carlson et al.*), Proteome Analysis (*by Pierre-Alan Binz et al.*), Target Finding in Genomes and Proteomes (*by Stefanie Fuhrman et al.*) as well as Screening of Drug Databases (*by Martin Stahl et al.*). In a concluding Chapter, *Thomas Lengauer* highlights the Future Trends in the field of bioinformatics.

The series editors are grateful to Thomas Lengauer, who accepted the challenging task to organize this volume on bioinformatics, to convince authors to participate in the project and to finish their chapters in time, despite the fact that research runs hot these days. We are sure that the result of his coordinative work constitutes another highlight in our series on Methods and Principles in Medicinal Chemistry. In addition, we want to thank Gudrun Walter and Frank Weinreich, Wiley-VCH, for their effective collaboration.

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## ***For my son Nico***

### **Foreword**

*Computational biology* and *bioinformatics* are terms for an interdisciplinary field joining information technology and biology that has skyrocketed in recent years. The field is located at the interface between the two scientific and technological disciplines that can be argued to drive a significant if not the dominating part of contemporary scientific innovation. In the English language, computational biology refers mostly to the scientific part of the field, whereas bioinformatics addresses mainly the infrastructure part. In some other languages (e.g. German) bioinformatics covers both aspects of the field.

The goal of this field is to provide computer-based methods for coping with and interpreting the genomic data that are being uncovered in large volumes through the diverse genome sequencing projects and other new experimental technology in molecular biology. The field presents one of the grand challenges of our times. It has a large basic research aspect, since we cannot claim to be close to understanding biological systems on an organism or even cellular level. At the same time, the field is faced with a strong demand for immediate solutions, because the genomic data that are being uncovered encode many biological insights whose deciphering can be the basis for dramatic scientific and economical success. At the end of the pre-genomic era that was characterized by the effort to sequence the human genome we are entering the postgenomic era that concentrates on harvesting the fruits hidden in the genomic text. In contrast to the pregenomic era which, from the announcement of the quest to sequence the human genome to its completion, has lasted less than 15 years, the postgenomic era can be expected to last much longer, probably extending over several generations.

While it will encompass many basic and general aspects of the field, the specific aim of the book is to point towards perspectives that bioinformatics can open towards the design of new drugs. It is this pharmaceutically oriented side of the field that provides the strongest fuel for the current widespread interest in bioinformatics.

Before this background, the book is intended as an introduction into the field of bioinformatics and is targeted to readers with a variety of backgrounds. Biologists, biochemists, pharmacologists, pharmacists and medical doctors can get from it an introduction into basic and practical issues of the computer-based part of handling and interpreting genomic data. In particular, many chapters of the book point to bioinformatics software and data resources that are available on the internet (often at no cost) and make an attempt at classifying and comparing those resources. For computer scientists and mathematicians the book contains an introduction into the biological backgrounds and the necessary information to begin to appreciate the difficulties and wonders of modeling complex biochemical and biomolecular issues in the computer.

Bioinformatics is a quickly progressing field. Both the experimental technologies and the computer-based method are in a dynamic phase of development. This book presents a snapshot of where the field stands today.

I am grateful to many people that helped make this book possible. Hugo Kubinyi first approached me with the idea for this book and since then accompanied the project with a finely balanced mixture of pressure and encouragement. Raimund Mannhold and Henk Timmerman supported the project as the other editors of this series. Above all, I thank the authors of the various chapters of the two volumes who found the time to write well thought-out chapters in a period of dramatic growth of the field that presents every participating scientist with an especially high workload. Gudrun Walter and Frank Weinreich handled the production process of the book very well. Finally I would like to express my deep gratitude to my wife Sybille and my children Sara and Nico who had to cope with my physical or mental absence while the project was ongoing.

**Thomas Lengauer**

Sankt Augustin, July 2001

# Part I: Basic Technologies

# 1

## From Genomes to Drugs with Bioinformatics

*Thomas Lengauer*

In order to set the stage for this two-volume book, this Chapter provides an introduction into the molecular basis of disease. We then continue to discuss modern biological techniques with which we have recently been empowered to screen for molecular drugs targets as well as for the drugs themselves. The Chapter finishes with an overview over the field of bioinformatics as it is covered in the two volumes of this book.

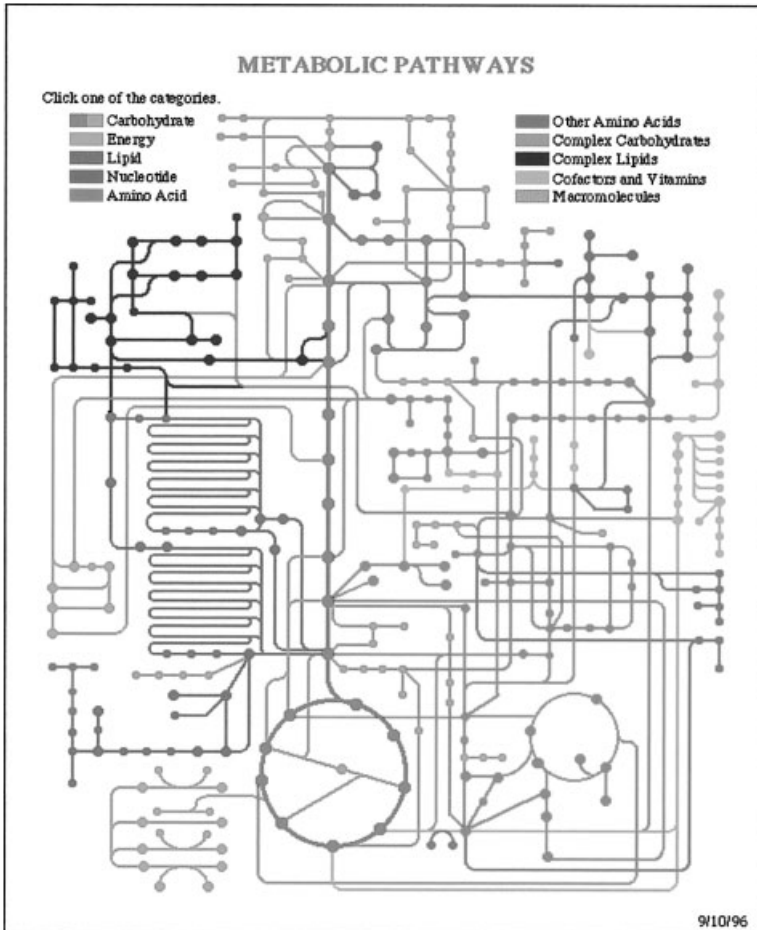
### 1.1

#### The molecular basis of disease

Diagnosing and curing diseases has always been and will continue to be an art. The reason is that man is a complex being with many facets much of which we do not and probably will never understand. Diagnosing and curing diseases has many aspects include biochemical, physiological, psychological, sociological and spiritual ones.

Molecular medicine reduces this variety to the molecular aspect. Living organisms, in general, and humans in particular, are regarded as complex networks of molecular interactions that fuel the processes of life. This “molecular circuitry” has intended modes of operation that correspond to healthy states of the organism and aberrant modes of operation that correspond to diseased states. In molecular medicine, the goal of diagnosing a disease is to identify its molecular basis, i.e. to answer the question what goes wrong in the molecular circuitry. The goal of therapy is to guide the biochemical circuitry back to a healthy state.

As we pointed out, the molecular basis of life is formed by complex biochemical processes that constantly produce and recycle molecules and do so in a highly coordinated and balanced fashion. The underlying basic principles are quite alike throughout all kingdoms of life, even though the processes are much more complex in highly developed animals and the human than in bacteria, for instance. Figure 1.1 gives an abstract view of such an



**Fig. 1.1**  
 Abstract view of part of the metabolic network of the bacterium *E. coli*. From <http://www.genome.ad.jp/kegg/kegg.html>

underlying biochemical network, the so-called *metabolic network* of a bacterial cell (the intestinal bacterium *E. coli*). The figure affords an incomplete and highly simplified account of the actual molecular interactions, but it nicely visualizes the view of a living cell as a biochemical circuit. The figure has the mathematical structure of a *graph*. Each dot (*node*) stands of a small organic molecule that is metabolized within the cell. Alcohol, glucose, and ATP are examples for such molecules. Each line (*edge*) stands for chemical reaction. The two nodes connected by the edge represent the substrate and the product of the reaction. The colors in Figure 1.1 represent the role that the respective reaction plays in metabolism. These roles include the con-



struction of molecular components that are essential for life – nucleotides (red), amino acids (orange), carbohydrates (blue), lipids (light blue), etc. – or the breakdown of molecules that are not helpful or even harmful to the cell. Other tasks of chemical reactions in a metabolic network pertain to the storage and conversion of energy. (The blue cycle in the center of the Figure is the citric acid cycle.) A third class of reactions facilitates the exchange of information in the cell or between cells. This include the control of when and in what way genes are expressed (*gene regulation*), but also such tasks as the opening and closing of molecular channels on the cell surface, and the activation or deactivation of cell processes such as replication or apoptosis (induced cell death). The reactions that facilitate communication within the cell or between cells are often collectively referred to as the *regulatory network*. Figure 1.1 only includes metabolic and no regulatory reactions. Of course the metabolic and the regulatory network of a cell are closely intertwined, and many reactions can have both metabolic and regulatory aspects. In general, much more is known on metabolic than on regulatory networks, even though many relevant diseases involve regulatory rather than metabolic dysfunction.

The metabolic and regulatory network can be considered as composed of partial networks that we call *pathways*. Pathways can fold in on themselves, in which case we call them *cycles*. A metabolic pathway is a group of reactions that turns a substrate into a product over several steps (pathway) or recycle a molecule by reproducing in several steps (cycle). The *glycolysis pathway* (the sequence of blue vertical lines in the center of Figure 1.1), is an example of a pathway that decomposes glucose into pyruvate. The *citric acid cycle* (the blue cycle directly below the glycolysis pathway in Figure 1.1) is an example of a cycle that produces ATP, the universal molecule for energy transport. Metabolic cycles are essential, in order that the processes of life not accumulate waste or deplete resources. (Nature is much better at recycling than man.)

There are several ways in which Figure 1.1 hides important detail of the actual metabolic pathway. In order to discuss this issue, we have extracted a metabolic cycle from Figure 1.1 (see Figure 1.2). This cycle is a component of cell replication, more precisely; it is one of the motors that drive the synthesis of thymine, a molecular component of DNA. In Figure 1.2, the

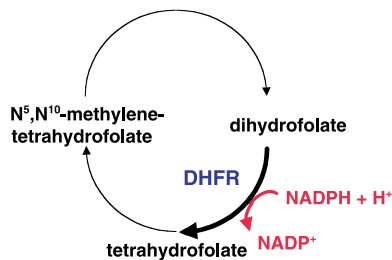
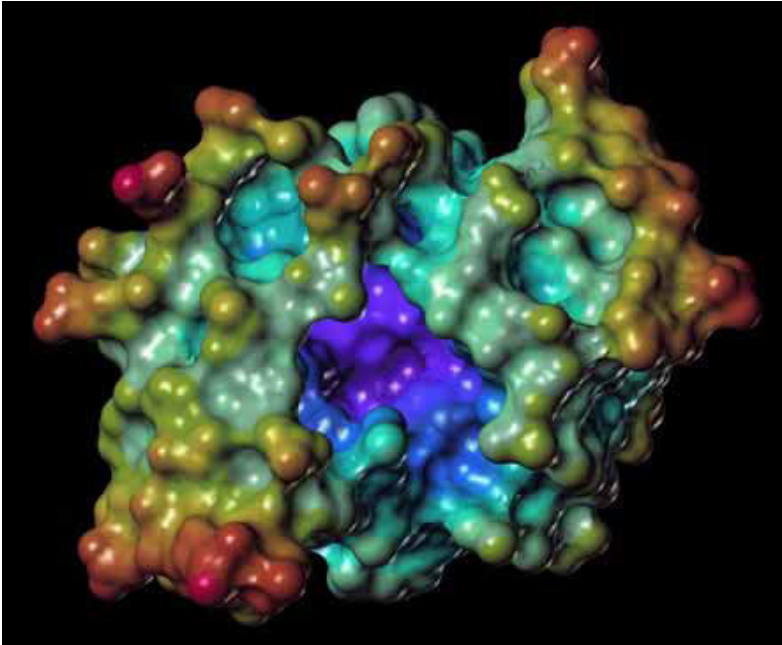


Fig. 1.2  
A specific metabolic cycle



**Fig. 1.3**  
The three dimensional structure of dihydrofolate reductase colored by its surface potential. Positive values are depicted in red, negative values in blue.

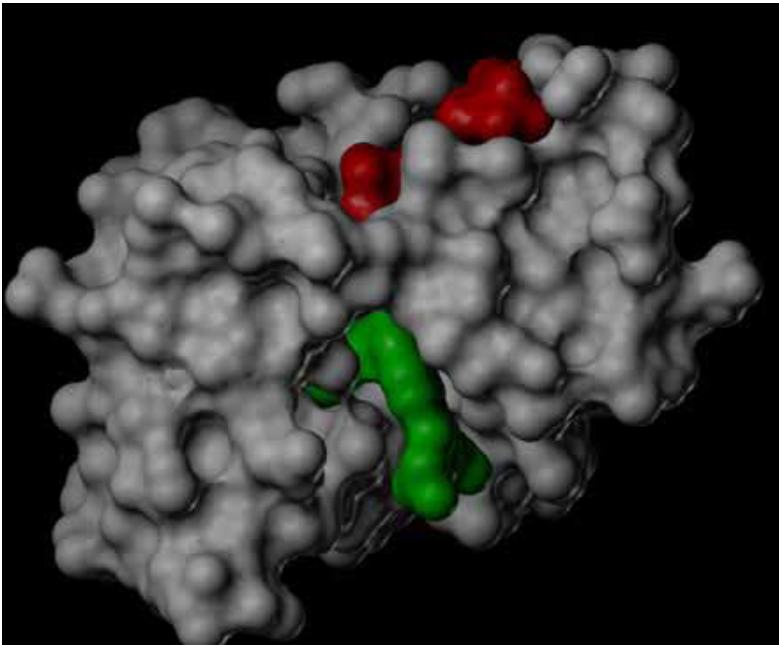
nodes of the metabolic cycle are labeled with the respective organic molecules, and the edges point in the direction from the substrate of the reaction to the product. Metabolic reactions can take place spontaneously under physiological conditions (in aqueous solution, under room temperature and neutral pH). However, nature has equipped each reaction (each line in Figure 1.1) with a specific molecule that catalyzes that reaction. This molecule is called an *enzyme* and, most often, it is a protein. An enzyme is a tailor-made binding site for the transition state of the catalyzed chemical reaction. Thus the enzyme speeds up the rate of that reaction tremendously, by rates of as much as  $10^7$ . Furthermore, the rate of a reaction that is catalyzed by an enzyme can be regulated by controlling the effectivity of the enzyme or the number of enzyme molecules that are available. Even the direction of a reaction can effectively be controlled with the help of several enzymes.

How does the enzyme do its formidable task? For an example, consider the reaction in Figure 1.2 that turns dihydrofolate into tetrahydrofolate. This reaction is catalyzed by an enzyme called *dihydrofolate reductase (DHFR)*. The surface of this protein is depicted in Figure 1.3. One immediately recognizes a large and deep pocket that is colored blue (representing its negative charge). This pocket is a *binding pocket* or *binding site* of the enzyme,

and it is ideally formed in terms of geometry and chemistry, such as to bind to the substrate molecule dihydrofolate and present it in a conformation that is conducive for the desired chemical reaction to take place. In this case, this pocket is also where the reaction is catalyzed. We call this place the *active site*. (There can be other binding pockets in a protein that are far removed from the active site.)

There is another aspect of metabolic reactions that is not depicted in Figure 1.1: Many reactions involve *co-factors*. A co-factor is an organic molecule, a metal ion, or – in some cases – a protein or peptide that has to be present in order for the reaction to take place. If the co-factor is itself modified during the reaction, we call it a *co-substrate*. In the case of our example reaction, we need the co-substrate NADPH for the reaction to happen. The reaction modifies dihydrofolate to tetrahydrofolate and NADPH to  $\text{NADP}^+$ . Figure 1.4 shows the molecular complex of DHFR, dihydrofolate (DHF) and NADPH before the reaction happens. After the reaction has been completed, both organic molecules dissociate from DHFR and the original state of the enzyme is recovered.

Now that we have discussed some of the details of metabolic reactions let us move back to the global view of Figure 1.1. We have seen that each of the edges in that figure represents a reaction that is catalyzed by a specific pro-



**Fig. 1.4**  
DHFR (gray) complexed with DHF (green) and NADPH red)

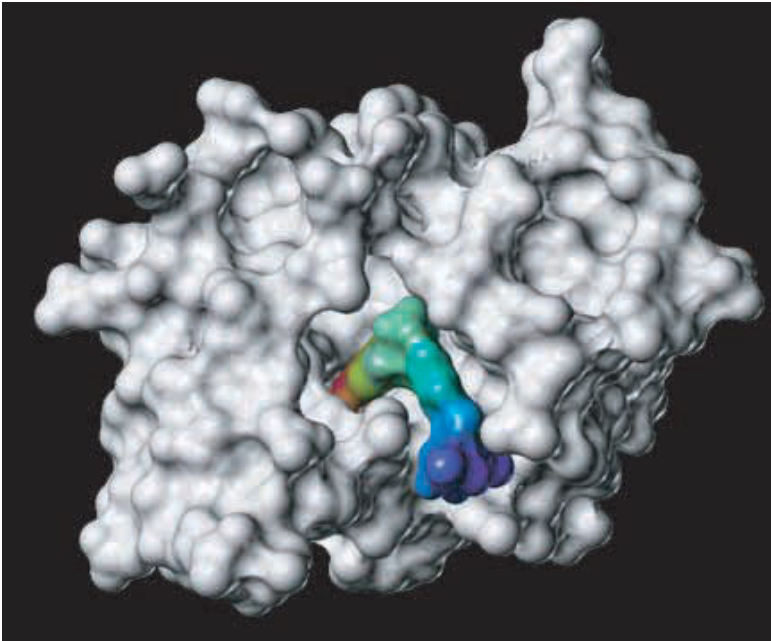
tein. (However, the same protein can catalyze several reactions.) In *E. coli* there are an estimated number of 3000 enzymes, in the human there are at least twice as many. The molecular basis of a disease lies in modifications of the action of these biochemical pathways. Some reactions do not happen at their intended rate (sickle cell anemia, diabetes), resources that are needed are not present in sufficient amounts (vitamin deficiencies) or waste products accumulate in the body (Alzheimer's disease). In general imbalances induced in one part of the network spread to other parts. The aim of therapy is to replace the aberrant processes with those that restore a healthy state. The most desirable fashion, in which this could be done, would be to control the effectiveness of a whole set of enzymes, in order to regain the metabolic balance. This set probably involves many, many proteins, as we can expect many proteins to be involved in manifesting the disease. Also, each of these proteins would have to be regulated in quite a specific manner. The effectiveness of some proteins would possibly have to be increased dramatically whereas other proteins would have to be blocked entirely etc. It is obvious that this kind of therapy involves a kind of global knowledge of the workings of the cell and a refined pharmaceutical technology that is far beyond what man can do today and for some time to come.

## 1.2

### The molecular approach to curing diseases

For this reason, the approach of today's pharmaceutical research is far more simplistic. The aim is to regulate a single protein. In some cases we aim at completely blocking an enzyme. To this end, we can provide a drug molecule that effectively competes with the natural substrate of the enzyme. The drug molecule, the so-called *inhibitor*, has to be made up such that it binds more strongly to the protein than the substrate. Then, the binding pockets of most enzyme molecules will contain drug molecules and cannot catalyze the desired reaction in the substrate. In some cases, the drug molecule even binds very tightly (covalently) to the enzyme (suicide inhibitor). This bond persists for the remaining lifetime of the protein molecule. Eventually, the deactivated protein molecule is broken down by the cell and a new identical enzyme molecule takes its place. Aspirin is an example of a suicide inhibitor. The effect of the drug persists until the drug molecules themselves are removed from the cell by its metabolic processes, and no new drug molecules are administered to replace them. Thus, one can control the effect of the drug by the time and dose it is administered.

There are several potent inhibitors of DHFR. One of them is *methotrexate*. Figure 1.5 shows methotrexate (color) both unbound (left) and bound (right) to DHFR (black and white). Methotrexate has been administered as an effective cytostatic cancer drug for over two decades.



**Fig. 1.5**  
Methotrexate (colored by surface potential, see Figure 3)  
and bound DHFR (gray)

There are many other ways of influencing the activity of a protein by providing a drug that binds to it. Drugs interact with all kinds of proteins:

- with receptor molecules that are located in the cell membrane and fulfill regulatory tasks
- with ion channels and transporter systems (again proteins residing in the cell membrane) that monitor the flux of molecules into and out of the cell

The mode of interaction between drug and protein does not always have the effect to block the protein, but we are generally looking for drugs that bind tightly to the protein.

Most drugs that are on the market today modify the enzymatic or regulatory action of a protein by strongly binding to it as described above. Among these drugs are long-standing, widespread and highly popular medications and more modern drugs against diseases such as AIDS, depression, or cancer. Even the life-style drugs that came into use in the past few years, such as Viagra and Xenical, belong to the class of protein inhibitors.

In this view, the quest for a molecular therapy of a disease decomposes into two parts:

- *Question 1: Which protein should we target?* As we have seen, there are many thousands candidate proteins in the human. We are looking for one of them that, by binding the drug molecule to it, provides the most effective remedy of the disease. This protein is called the *target protein*.
- *Question 2: Which drug molecules should be used to bind to the target protein?* Here, the molecular variety is even larger. Large pharmaceutical companies have compound archives with millions of compounds at their disposal. Every new target protein raises the questions, which of all of these compounds would be the best drug candidate. Nature is using billions of molecules. With the new technology of combinatorial chemistry, where compounds can be synthesized systematically from a limited set of building blocks, this number of *potential* drug candidates is also becoming accessible to the laboratory.

We will now give a short summary of the history of research on both of these questions.

### 1.3 Finding protein targets

Question 1 could not really be asked realistically until a few years ago. Historically, few target proteins were known at the time that the respective drug has been discovered. The reason is that new drugs were developed by modifying known drugs, based on some intuitive notion of molecular similarity. Each modification was immediately tested in the laboratory either *in vitro* or *in vivo*. Thus, the effectiveness of the drug could be assessed without even considering the target protein. To this day, all drugs that are on the marketplace world-wide target to an estimated set of 500 proteins [1]. Thus the search for target proteins is definitely the dominant bottleneck of today's pharmaceutical research.

Today, new experimental methods of molecular biology, the first versions of which have been developed just a few years ago, afford us with a fundamentally new way – the first systematic way – of looking for protein targets. We exemplify this progress at a specific DNA chip technology [2]. However, the general picture extends to many other experimental methods under development.

Figure 1.6 shows a DNA-chip that provides us with a differential census of the proteins manufactured by a yeast cell in two different cell states, one governed by the presence of glucose (green) and one by the absence of glucose (red). In effect the red picture is that of a starving yeast cell, whereas the green picture shows the “healthy” state. Each bright green dot stands for a protein that is manufactured (expressed) in high numbers in the “healthy” state. Each bright red dot stands for a protein that is expressed



**Fig. 1.6**  
A DNA chip (from <http://cmgm.stanford.edu/pbrown/explore/>)

highly in the starving cell. If the protein occurs frequently in both the healthy and the starving state, the corresponding dot is bright yellow (resulting from an additive mixture of the colors green and red). Dark dots stand for proteins that are not frequent, the tint of the color again signifies whether the protein occurs more often in the healthy cell (green), equally often in both cells (yellow) or more often in the diseased cell (red).

At this point it is of secondary importance, exactly which experimental procedures generate the picture of Figure 1.6. What is important is, how much information is attached to colored dots in the picture. Here, we can make the following general statements.

1. The identity of the protein is determined by the coordinates of the colored dot. We will assume, for simplicity, that dots at different locations also represent different proteins. (In reality, multiple dots that represent

the same protein are introduced, on purpose, for the sake of calibration.) The exact arrangement of the dots is determined before the chip is manufactured. This involves identifying a number of proteins to be represented on the chip and laying them out on the chip surface. This layout is governed by boundary conditions and preferences of the experimental procedures and is not important for the interpretation of the information.

2. Only rudimentary information is attached to each dot. At best, the experiment reveals the complete sequence of the gene or protein. Sometimes, only short segments of the relevant sequence are available.

Given this general picture, the new technologies of molecular biology can be classified according to two criteria, as shown by the next subsections.

### 1.3.1

#### **Genomics vs proteomics**

In genomics, not the proteins themselves are monitored but rather we screen the expressed genes whose translation ultimately yields the respective proteins. In proteomics, the synthesized proteins themselves are monitored. The chip in Figure 1.6 is a DNA-chip, i.e., it contains information on the expressed genes and, thus, only indirectly on the final protein products. The advantage of the genomics approach is that genes are more accessible experimentally and easier to handle than proteins. For this reason, genomics is ahead of proteomics, today. However, there also are disadvantages to genomics. First the expression level of a gene need not be closely correlated with the concentration of the respective protein in the cell. But the latter figure may be more important to us if we want to elicit a causal connection between protein expression and disease processes. Even more important, proteins are modified post-translationally (i.e. after they are manufactured). These modifications involve glycosylation (attaching complex sugar molecules to the protein surface), and phosphorylation (attaching phosphates to the protein surface), for instance, and they lead to many versions of protein molecules with the same amino-acid sequence. Genomics cannot monitor these modifications, which are essential for many diseases. Therefore, it can be expected that, as the experimental technology matures, proteomics will gain importance over genomics.

### 1.3.2

#### **Extent of information available on the genes/proteins**

Technologies vary widely in this respect. The chip in Figure 1.6 is generated by a technology that identifies the gene sequence. We are missing information on the structure and the function of the protein, its molecular interac-



tion partners and its location inside the metabolic or regulatory network of the organism. All of this information is missing the majority of the genes on the chip.

There are many variations on the DNA-chip theme. There are technologies based on so-called *expressed sequence tags (ESTs)* that tend to provide more inaccurate information on expression levels, and various sorts of microarray techniques (see Chapter 5 of Volume 2). Proteomics uses different kinds of gels, and experiments involving mass spectrometry (see Chapter 4 of Volume 2). As is the case with genomics, proteomics technologies tend to generate information on the sequences of the involved proteins and on their molecular weight, and possibly information on post-translational modifications such as glycosylation and phosphorylation. Again, all higher-order information is missing. It is infeasible to generate this information exclusively in the wet lab – we need bioinformatics to make educated guesses here. Furthermore basically all facets of bioinformatics that start with an assembled sequence can be of help. This includes the comparative analysis of genes and proteins, protein structure prediction, the analysis and prediction of molecular interactions involving proteins as well as bioinformatics for analyzing metabolic and regulatory networks. This is why all of bioinformatics is relevant for the purpose of this book. Thus, part 1 of the book summarizes the state of the art of the relevant basic problems in bioinformatics.

If, with the help of bioinformatics, we can retrieve this information then we have a chance of composing a detailed picture of the disease process that can guide us to the identification of possible target proteins for the development of an effective drug. Note that the experimental technology described above is universally applicable. The chip in Figure 1.6 contains all genes of the (fully sequenced) organism *S. cerevisiae* (yeast). The cell transition analyzed here is the diauxic shift – the change of metabolism upon removal of glucose. But we could exchange this with almost any other cell condition of any tissue of any conceivable organism. The number of spots that can be put on a single chip goes into the hundred thousands. This is enough for putting all human genes on a single chip. Also, we do not have to restrict ourselves to disease conditions; all kinds of environmental conditions (heat, cold, low or high pH, chemical stress, drug treatment, diverse stimuli etc.) or intrinsic conditions (presence or absence of certain genes, mutations etc.) can be the subject of study.

This paradigm of searching for target proteins in genomics data is considered to be so promising that pharmaceutical industry has been investing heavily in it in over the past years. Contracts between pharmaceutical companies and new biotech industry that is providing the data and clues to new targets have involved hundreds of million dollars. In turn, each new target protein can afford a completely new approach to disease therapy and a potentially highly lucrative worldwide market share. Yet, the paradigm still has to prove its value. These commercial aspects of the research have also

strongly affected the scientific discipline of bioinformatics. We will comment more on this in the second volume of this book.

Providing adequate bioinformatics for finding new target proteins is a formidable challenge that is the focus of much of this book. But once we have a target protein our job is not done.

## 1.4

### Developing drugs

If the target protein has been selected, we are looking for a molecule that binds tightly to a binding site of the protein. Nature often uses macromolecules, such as proteins or peptides to inhibit other proteins. However, proteins do not make good drugs: They are easily broken down the digestive system, they can elicit immune reaction, and they cannot be stored for a long time. Thus, after a short excursion into drug design based on proteins, pharmaceutical research has gone back to looking for small drug molecules. Here, one idea is to use a peptide as the template for an appropriate drug (peptidomimetics).

Due to the lack of fundamental knowledge of the biological processes involved, the search for drugs was governed by chance, until recently. However, as long as chemists have thought in terms of chemical formulas, pharmaceutical research has attempted to optimize drug molecules based on chemical intuition and on the concept of molecular similarity. The basis for this approach is the lock-and-key principle formulated by Emil Fischer [3] about a hundred years ago. Figure 1.4 nicely illustrates that principle: In order to bind tightly the two binding molecules have to be complementary to each other both sterically and chemically (colors). The drug molecule fits into the binding pocket of the protein like a key inside a lock. The lock-and-key principle has been the dominating paradigm for drug research ever since its invention. It has been refined to include the phenomenon of induced fit, by which the binding pocket of the protein undergoes subtle steric changes in order to adapt to the geometry of the drug molecule.

For most of the century, the structure of protein binding pockets has not been available to the medicinal chemist, so drug design was based on the idea that molecules whose surface is similar in shape and chemical features should bind to the target protein with comparable strength. Thus drug design was based on comparing drug molecules, either intuitively or, more recently, systematically with the computer. As 3D protein structures became available, the so-called *rational* or *structure-based* approach to drug development was invented, which exploited this information to develop effective drugs. Rational drug design is a highly interactive process with the computer originally mostly visualizing the protein structure and allowing queries on its chemical features. The medicinal chemist modified drug mole-

cules inside the binding pocket of the protein at the computer screen. As rational drug design began to involve more systematic optimization procedures interest rose in *molecular docking*, i.e. the prediction of the structure and binding affinity of the molecular complex involving a structurally resolved protein and its binding partner, in the computer. Synthesizing and testing a drug in the laboratory used to be comparatively expensive. Thus it was of interest to have the computer suggest a small set of highly promising drug candidates. However, to this day rational drug design has been hampered by the fact that the structure of the target protein will not be known for many pharmaceutical projects for some time to come. For instance, many diseases involve target proteins that reside in the cell membrane, and we cannot expect the 3D structure of such proteins to become known soon.

With the advent of *high-throughput screening*, the binding affinity of as many as several hundred thousands drug candidates to the target protein can now be assayed within a day. Furthermore, *combinatorial chemistry* allows for the systematic synthesis of molecules that are composed of preselected molecular groups that are linked with preselected chemical reactions. The number of molecules that is accessible in such a combinatorial library, in principle, can exceed many billions. Thus we need the computer to suggest promising sublibraries that contain a large number of compounds that bind tightly to the protein.

As in the case of target finding, the new experimental technologies in drug design require new computer methods for screening and interpreting the voluminous data assembled by the experiment. These methods are seldom considered part of bioinformatics, since the biological object, namely the target protein is not the focus of the investigation. Rather, people speak of *cheminformatics*, the computer aspect of medicinal chemistry. Whatever the name, it is our conviction that both aspects of the process that guides us from the genome to the drug have to be considered together, and we will do so in this book.

## 1.5

### A bioinformatics landscape

In the Sections above, we have described the application scenario that is the viewpoint from which we are interested in bioinformatics. In this Section, we attempt to chart out the field of bioinformatics in terms of its scientific subproblems and the application challenges that it faces.

Basically, there are two ways of structuring the field of bioinformatics. One is *intrinsically*, by the type of problem that is under consideration. Here, the natural way of structuring is by layers of information that are compiled starting from the genomic data and working our way towards various levels of the phenotype. The second is *extrinsically*, by the medical or pharmaceu-

tical application scenario to which bioinformatics contributes and by the type of biological experimentation that it supports. This book is roughly structured according to this distinction. The intrinsic view is the subject of Volume 1. This volume discusses subproblems of bioinformatics that provide components in a global bioinformatics solution. Each Chapter is devoted to one relevant component. The extrinsic view is the focus of Volume 2. This volume has Chapters which each concentrate on some important application scenarios that can only be supported effectively by combining components discussed in Volume 1.

Both views are summarized in an integrated fashion in Figure 1.7, where the bioinformatics circle is depicted reaching from the genome sequence to the macroscopic phenotype. We will be guided by this figure as we now chart out the bioinformatics landscape in more detail.

### 1.5.1

#### The intrinsic view

The intrinsic view may be the more basic research-oriented one. This view on bioinformatics tends to produce reasonably well-defined and timeless scientific problems that often have the character of grand challenges. As described above, we will structure this view of bioinformatics in layers of derived information that we add on top of the primary information provided by the genomic sequence.

##### 1.5.1.1 Layer –1: Sequencing support

This part of bioinformatics addresses problems that occur even before the genomic sequence is available. The object is to interpret experimental data that are generated by sequencing efforts. Problems to be addressed here are *base calling*, i.e. the interpretation of the signals output by sequencers in terms of nucleic acid sequences, *physical mapping*, i.e. providing a rough map of relevant marker loci along the genome, and *fragment assembly*, i.e. the process of piecing together short segments of sequenced DNA to form a contiguous sequence of the genome or chromosome. Layer –1 plays a special role, since it is both a subproblem and an application scenario. The application-oriented character of this problem originates from its close relationship with the applied sequencing procedure. The two competing approaches to sequencing the human genomes, namely whole-genome-shotgun sequencing, as performed by Celera Genomics [4] and BAC-assembly as performed by the publically funded human genome project [5], which lead to quite different approaches for assembly, have illustrated this point very visibly. We chose to discuss this layer in Volume 2 (Chapter 2 by Xiaoqiu Huang).

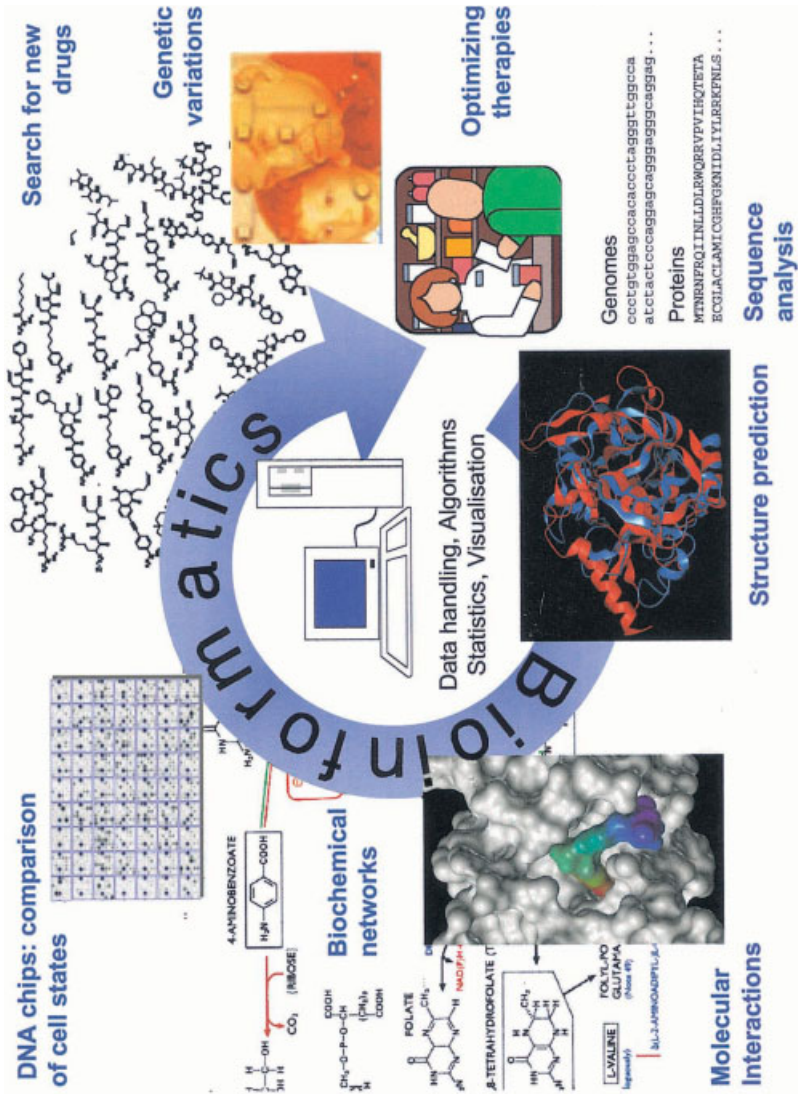


Fig. 1.7  
 A schematic overview of bioinformatics

After exercising layer –1 of bioinformatics successfully the raw genomic sequence is available.

#### 1.5.1.2 Layer 1: Analysis of nucleic acid sequences (5 o'clock position in Figure 1.7)

This layer is concerned with annotating the raw genomic sequence with information that can be derived directly from the sequence. Problems addressed in this layer are *gene finding* (Chapter 3 of Volume 1, Victor Solovyev), i.e. the identification of those – as it happens, rare – stretches of genomic DNA that code for proteins, and the analysis of the much more prevalent and quite a bit more mysterious *non-coding regions* in the genome (Chapter 4 of Volume 1, Thomas Werner). The former problem is our entry to understanding the proteome. It is so hard, that today experimental procedures that go directly after protein sequences are developed to help along the way. The latter problem currently concentrates on the regions upstream of sequences that encode proteins. These regions are believed to contain patterns that govern the regulation of the expression of the genes, i.e. their translation to proteins. Both problems can be considered basic research problems and scientific grand challenges. Numerous other annotations are desirable, such as the annotation of genomic sequences with homologous sequences in the same or different species, the distinction between orthologous<sup>1)</sup> and paralogous<sup>2)</sup> sequences. A comparison between many sequences at a time can yield a *multiple alignment* or, if we are interested in the lineages of the participating sequences, an *evolutionary tree*.

The processes of layer 1 provide the data for layer 2.

#### 1.5.1.3 Layer 2: Analysis of protein sequences (5 o'clock position in Figure 1.7)

This layer annotates protein sequences. Here, the most voluminous source of information is the comparison with other homologous sequences, either on the protein or on the nucleic acid level. The results desired are relationships between different proteins that allow for making conclusions about

- |   |   |
|---|---|
| <p>1) Orthologous sequences are homologous sequences in <i>different</i> species that have common origin and whose distinctions come about by gradual evolutionary modifications from the common ancestor. Orthologous sequences tend to have the same function in different species.</p> | <p>2) Paralogous sequences are homologous sequences that exist within a species. Paralogous sequences have a common origin but involve gene duplication events to arise. Often the gene duplication has the purpose of using the sequence to implement a new function. Thus paralogous sequences often perform different functions.</p> |
|---|---|

protein function, cellular localization and the like. The methods in layer 2 are basically the same ones as in layer 1. Chapter 2 of Volume 1 (Martin Vingron) concentrates on the bioinformatics methods for layers 1 and 2.

#### 1.5.1.4 Layer 3: Analysis and prediction of molecular structure (6 o'clock position in Figure 1.7)

This layer generates three-dimensional models of the biomolecules under consideration. In the introductory part of the Chapter we have seen that this structure is the key to the function of a molecule. Thus, modeling molecular structure is a very important problem of bioinformatics. There are several kinds of molecules to be considered.

##### 1.5.1.4.1 DNA

The structure of DNA is quite well preserved throughout nature. It is the double helix. There are small differences, the so-called fine structure of DNA, which are modeled in the computer with methods like energy minimization and molecular dynamics. These methods are also used for modeling other biomolecules, notably, proteins. Therefore modeling the structure of DNA is not covered in this book.

##### 1.5.1.4.2 RNA

This molecule is structurally more flexible than DNA. The database of resolved three-dimensional RNA structures is much smaller than that for proteins; it contains only a few hundred molecules. Modeling three-dimensional structures of RNA is basically an unsolved problem, however, there are effective methods for modeling two-dimensional RNA structures. We do not discuss modeling RNA structures in this book and refer the reader to [6].

##### 1.5.1.4.3 Proteins

Proteins display a wide variety of structures. The protein structure prediction problem has been a challenge for a long time. Many successes have been accomplished but the general problem is still far from being solved. What can be achieved is to model (target) proteins given a structurally resolved protein that acts as a template. However this process is only successful if the target and the template protein exhibit a sufficient amount of similarity in sequence. Chapter 5 of Volume 1 (Roland Dunbrack, Jr) describes the methods used in this approach to protein modeling. If the sequence simi-

larity is low, then one cannot generate an accurate full-atom model of the protein, in general. But often one can still find out a lot of significant aspects of the structure of the protein, such as the overall architecture, the coordinates of the protein backbone, or even more accurate models of relevant binding pockets. Chapter 6 of Volume 1 (Ralf Zimmer, Thomas Lengauer) reports on the state of the art in this field.

#### 1.5.1.4.4 Lipids, carbohydrates

Lipids form membranes inside and around the cell. Carbohydrates form complex tree-like molecules that become attached to the surface of proteins and cellular membranes. In both cases, the three-dimensional molecular structure is not unique, but the molecular assemblies are highly flexible. Thus, analyzing the molecular structure involves the inspection of a process in time. Molecular dynamics is the only available computer-based method for doing so. Compared with protein structures there are relatively few results on lipids and carbohydrates. The book does not detail this topic.

#### 1.5.1.5 Layer 4: Molecular interactions (7 o'clock position in Figure 1.7)

If the molecular structure is known we can attempt to analyze the interactions between molecules. Among the many possible pairs of interaction partners two are both relevant and approachable by computer.

##### 1.5.1.5.1 Protein–ligand interactions

Here, one molecular partner is a protein; and the other is a small, often flexible, organic molecule. This is the problem discussed in detail in the introductory parts of this Chapter. The problem is a basic research problem and, at the same time, is of prime importance for drug design. The problem has two aspects. One is to determine the correct geometry of the molecular complex. The second is to provide an accurate estimate of the differential free energy of binding. Whereas much progress has been made on the first aspect, the second aspect remains a grand challenge. Chapter 7 of Volume 1 (Matthias Rarey) discusses this problem.

##### 1.5.1.5.2 Protein–protein interactions

Here two proteins bind to each other. This problem is important if we want to understand molecular interaction inside living organisms; it plays less of a role for drug design. The problem is different from protein–ligand docking in several respects. In protein–ligand docking, the binding mode is mostly determined by strong enthalpic forces between the protein and the



ligand. In addition, the contributions of desolvation (replacing the water molecules inside the pocket by the ligand) are essential. The notion of molecular surface is not as much relevant, especially, since the ligand can be highly flexible and does not have a unique surface. In contrast, geometric complementarity of both proteins is a dominating issue in protein–protein docking, where both binding partners meet over a much larger contact surface area. Issues of desolvation can be essential here. Induced fit, i.e. subtle structural changes on the protein surface to accommodate binding are important in both problems, but seem to be more essential in protein–protein docking. Chapter 8 of Volume 1 (Michael Sternberg) discusses this problem. Estimating the free energy of binding is a major bottleneck in all docking problems.

#### 1.5.1.5.3 Other interactions

Of course, all kinds of molecular interactions are important, be it protein–DNA (essential for understanding gene regulation) or reactions involving RNA, lipids or carbohydrates. Protein–DNA docking seems even harder than protein–protein docking. Because of the rigidity of DNA stronger conformational changes seem to be necessary on the protein side. Also, water plays an important role in protein–DNA docking. Chapter 8 discusses protein–DNA docking. The other problems are not discussed in this book.

#### 1.5.1.6 Layer 5: Metabolic and regulatory networks (9 o'clock position in Figure 1.7)

This layer takes the step from analyzing single interactions to analyzing networks of interactions. It has a database aspect, which just collects the voluminous data and makes them generally accessible, and an algorithmic aspect that performs simulations on the resulting networks. Both aspects are in a preliminary stage. The most development has taken place in metabolic databases. Here, we have quite a detailed picture of parts of the metabolism for several organisms. This book does not detail the simulation of metabolic networks.

## 1.6

### The extrinsic view

This view structures bioinformatics according to different application scenarios and types of experimental data that are under scrutiny.

## 1.6.1

**Basic contributions: molecular biology databases and genome comparison**

The first completely sequenced bacterial genome, that of the bacterium, *H. influenzae*, was available in 1994. Since then, a few dozen bacterial genomes have been sequenced. Sequencing a bacterium has become state of the art, and we can expect many more genomes to appear in the near future. The first eucaryote (*S. cerevisiae*, yeast [7]) has been sequenced in 1996, and the genome of the first multicellular organism (*C. elegans*, nematode, [8]) became available in 1998. The genome of *D. melanogaster* (fruitfly [9]) became available in February 2000, and two drafts of the human genome sequence have been published in February 2001 [4, 5]. As the number and variety of completely sequenced genomes rises, their comparison can yield important scientific insights. What is it that makes the bacterium *E. coli* a symbiotic beneficial parasite, whereas the bacterium *H. influenzae* that generates ear infections basically has toxic effects in the relevant parts of the human body? Huynen et al. [10] have carried out a differential analysis that looks for homologies among proteins in both bacteria and concentrates on those (about 200) proteins that occur only in *H. influenzae*. Using homology searches over the database of available protein sequences from all kinds of organisms, Huynen et al. [10] found that many of these 200 proteins are similar to proteins that act toxic in other organisms. Thus, in some sense, the set of candidate proteins that are responsible for the adverse effects of *H. influenzae* has been narrowed down from the about 1800 protein in the bacterium to 200. This is an example of the kinds of results genome comparison can achieve.

In order to carry out such tasks, we need to combine a wide variety of methods and data. Chapter 1 of Volume 2 (Thure Etzold et al.) describes a successful and widely used approach for linking the vastly growing variety of biomolecular databases.

## 1.6.2

**Scenario 1: Gene and protein expression data (10 o'clock position in Figure 1.7)**

The measurement and interpretation of expression levels of genes and proteins is at the center of today's attention in genomics, proteomics, and bioinformatics. The goal is to create a detailed differential functional picture of the cell's inner workings, as it is determined by the cell state at hand. Cell states can be influenced and determined by external environmental factors (temperature, pH, chemical stress factors), and by internal factors such as the cell belonging to a special tissue or be in a certain disease state.

Historically the first version of expression of data is *expressed sequence tags* (ESTs). These are short stretches of cDNA (complementary DNA), i.e. DNA that has been translated back from messenger RNA (mRNA) expressed in the cell. The messenger RNA is in a mature state, i.e. introns have been spliced out before the cDNA was taken from the mRNA. ESTs provide a census of the genes expressed in a cell that is subject to certain internal and external conditions. The level of expression of a certain gene is roughly correlated with the frequency with which this gene is hit by sequenced ESTs. Today, there are large collections of ESTs available for many organisms and cells subject to many different conditions.

The DNA chips described above (see Figure 1.6) provide a different variant of the kind of data provided by ESTs. The advantage of DNA chips is that they can provide more accurate expression data on expression levels. Whereas, EST counts can be low and generally provide little resolution. One way of reading out expressions levels from DNA chips is via fluorescent marking. The high correlation between the light intensity and the expression level provides more resolution than EST counts.

Expression levels on DNA or protein gels are generally deduced from the size of spots on the gel. 2D gels and mass spectrometry can be used as fast tools for identifying expressed proteins in various states of the cell. These experimental data and their interpretation with bioinformatics are subject to Chapter 4 (Roland Appel et al., proteomics) and Chapter 5 (Roland Somogyi, genomics) of Volume 2.

### 1.6.3

#### **Scenario 2: Drug screening (1 o'clock position in Figure 1.7)**

Once the target protein is identified, the search for the drug starts. That search is again composed of a mixture of experimental and computer-based procedures. In a first phase, one is looking for a *lead compound*. The only requirement on the lead compound is that it binds tightly to the binding site of the target protein. In a second step of *lead optimization*, the lead compound is modified to be non-toxic, have few side effects and be bio-accessible, i.e. be easily delivered to the site in the body, at which its effect is desired.

Basically, there are two approaches to developing a new drug. One is to create one from scratch. This has been attempted on the basis of the knowledge of the three-dimensional structure of the binding site of the protein. In effect, the drug molecule can be designed to be complementary to the binding site in steric and physicochemical terms. This approach is called *ab initio* drug design (not to be confused with *ab initio* protein structure prediction or *ab initio* methods in theoretical chemistry). *Ab initio* drug design has been hampered by the fact that often the developed molecules

were hard to synthesize. Furthermore lead optimization turned out to be hard for many lead compounds developed with *ab initio* design techniques.

The second approach is to screen through a large database of known molecules and check their binding affinity to the target protein. The advantage of this approach is that, mostly, the compounds that have been accumulated in a database have been investigated before – though mostly in a different context. Their bioaccessibility and toxicity may have been studied and they probably have been synthesized. Screening compounds from a library can be done in the laboratory or in the computer. In the latter case, the process is called *virtual screening*. Compound databases become large, so a mixture of the two processes may be advisable. Here, a set of compounds is preselected from a large library with virtual screening. These compounds are then tested in the laboratory. Chapter 6 of volume 2 by Matthias Rarey, Martin Stahl, and Gerhard Klebe gives more detail on virtual drug screening.

#### 1.6.4

##### **Scenario 3: Genetic variability (2 o'clock position in Figure 1.7)**

Recently, new experimental methods have been developed that uncover variations of genomic text for organisms of the same species. The sequencing data discriminate between different external or internal states of the organism or provide insight into intra-species genetic variability. Genetic variability is just starting to be investigated on a genomic scale both experimentally and with bioinformatics. Chapter 3 of Volume 2 (Deborah Nickerson) introduces this new and exciting field of molecular biology and bioinformatics.

Genetic variability does not only occur in the host (patient) but also in the guest (infectious agent). For many antibacterial and several antiviral therapies (e.g. AIDS), the guest population reacts to the administering of drugs, and resistant strains develop. In this context, drug selection is a difficult problem. Given appropriate genotypic and phenotypic data, bioinformatics can eventually help in optimizing therapies in this context (3 o'clock position in Figure 1.7).

The following Chapters of both volumes provide the necessary detail for an in-depth introduction into bioinformatics. Chapter 7 of volume 2 will make an attempt at wrapping up where we stand today and balance hopes and expectations with limitations of this new and rapidly developing field.

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## 2

# Sequence Analysis

*Martin Vingron*

### 2.1

#### Introduction

Although a theoretical method to predict the three-dimensional structure of a protein from its sequence alone is not in sight yet, researchers have uncovered a multitude of connections between the primary sequence on one hand and various functional features of proteins on the other hand. Among the success stories are the recognition of transmembrane proteins or the classification of proteins into classes of similar function based on sequence similarity. The latter achievement uses the observation that proteins that are similar in sequence are likely to share similar functional features. This is also giving rise to the enormous utility of similarity searches in sequence data bases. This Chapter will deal with the kinds of analysis and predictions based on primary sequence alone and the algorithms used in this field.

Roughly speaking, methods fall into two classes, based on whether they analyze the individual protein sequence in isolation or whether implications are drawn from a comparison with or among many sequences. Individual sequences are analyzed primarily based on characteristics of amino acids as derived either from physical chemistry or statistically. Such characteristics are hydrophobicity scales or statistical preferences for a particular secondary structure. Section 2.2 will cover these methods.

Naturally the type of information that can be extracted from a sequence in isolation is very much restricted compared to what can be gained from analyzing many sequences. The key observation allowing us to exploit the information that is collected in our databases is the close relationship between homology, i.e., common ancestry and sequence similarity. In searching for sequence similarity we strive for the recognition of homology because homology lets us suggest common functions. The basis for the study of sequence similarity is the comparison of two sequences which will be dealt with in Section 2.3.

Sequence comparisons are effected in large numbers when searching sequence databases for sequences that are similar to a query sequence. Al-

gorithms for this purpose need to be fast even at the expense of sensitivity. Section 2.4 discusses the widely used heuristic approaches to database searching. However, the algorithms we are designing for the purpose of quantifying sequence similarity can only be as good as our understanding of evolutionary processes and thus they are far from perfect. Therefore, results of algorithms need to be subjected to a critical test using statistics. Methods for the assessment of the statistical significance of a finding are introduced in Section 2.5.

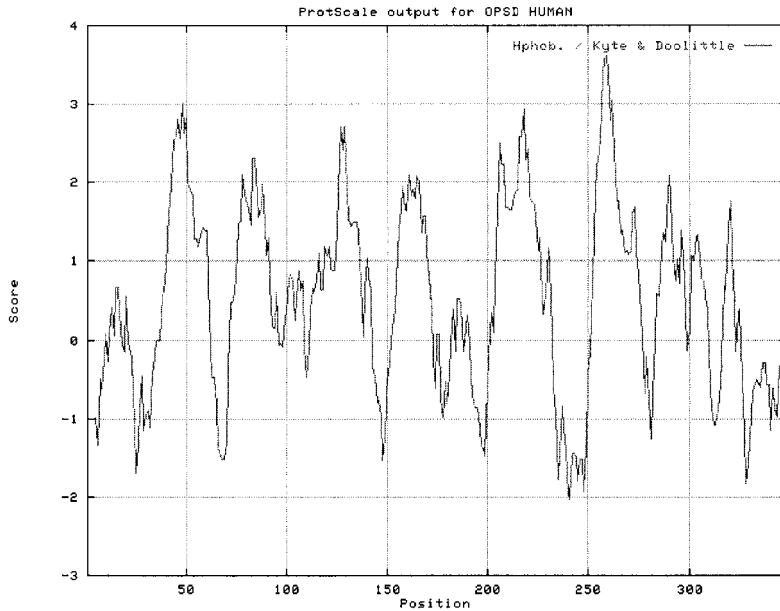
Genes do not come in pairs but rather in large families. Consequently, the need arises to align more than two sequences at a time which is done by multiple alignment programs. Computationally a very hard problem, it has found considerable attention from the side of algorithms development. Section 2.6 presents the basic approaches to multiple sequence alignment. Section 2.7 will build on the knowledge of a multiple alignment and introduce how to exploit the information contained in several related sequences for the purpose of identifying additional related sequences in a database. The last section will cover methods for identifying domains in one or among several sequences. For the annotation of sequences and for prediction of function this aspect has become increasingly important during recent years.

## 2.2

### Analysis of individual sequences

This Chapter focuses on analyses that can be performed based solely on the primary sequence of a protein. Several rationales can be applied. Physico-chemical characteristics of individual amino acids are one basis for predictions of gross structural features. For example, particular repetitive patterns may suggest a coiled-coil structure while in general secondary structure can be predicted based on an a statistical analysis of the primary sequence. The definition of signals recognized by the cellular transport machinery allow the prediction of subcellular location. Although somewhat unsystematic such observations can provide valuable hints as to the structure and/or function of a protein.

Amino acids side chains differ in their physico-chemical features. For example, some like to be exposed to water, i.e., they are hydrophilic, while the hydrophobic amino acids tend to avoid exposure to water. Charge, size, or flexibility in the backbone are only some of the other examples of amino acid parameters. These parameters are usually measured on a numerical scale such that for every parameter there exists a table assigning a number to each amino acid. For the case of hydrophobicity two such scales have become famous. The first is due to Hopp and Woods [1] while the other is due to Kyte and Doolittle [2]. A large collection of amino acid parameters



**Fig. 1**  
Hydrophobicity plot for the transmembrane protein opsin.  
Note the peaks in hydrophobicity which roughly denote  
the putative membrane spanning regions.

have been collected by Argos [3] who found that subgroups of them are correlated with each other such that the real information content of this large number of parameters is in fact lower than it seems. He selected the following parameters as a non-redundant set: hydrophobicity, turn preference [4], residue bulk [5], refractivity index [6], and antiparallel strand preference [7].

Standard sequence analysis software today offers programs that plot various parameters for a given protein (Figure 2.1). Serious software packages tend to provide the user with a selection of informative and non-redundant parameters similar to the one given. Some other packages pretend to offer new insights by plotting large numbers of parameters. In practice, the various parameters are used to plot a curve along the amino acid chain. Values are averaged within a sliding window to smoothen the curve. The selection of the window width is, of course, arbitrary but values between 7 and 15 would generally seem appropriate. Within a window a weighting scheme may be applied which would typically assign more importance to residues in the middle of the window than to the ones on the edges.

Features of the individual amino acids also play a key role in protein secondary structure formation. Based on experimental data, scales have been



assigned to the amino acids describing their preference for assuming, e.g., a helical structure [8]. Consequently, early secondary structure prediction methods have assigned preferences to the amino acids according to which secondary structure they tend to assume. For example, Glutamate is frequently found in alpha helices while Valine has a preference for beta strands and Proline is known to be strongly avoided in helices. Modern secondary structure prediction methods are more involved though (see below).

The functional features of proteins that are grasped by such parameters are manifold. Hydrophobic amino acids tend to occur in the interior of globular proteins, while at the surface of a protein one will preferentially find hydrophilic residues. One application of the latter is the prediction of antigenic epitopes because these are assumed to coincide with patches of hydrophilic residues. In transmembrane proteins, the regions of the chain that span the membrane tend to be strongly hydrophobic and the recognition of several such regions is an integral part of methods aiming at the recognition of transmembrane proteins based on their sequence alone [9, 10]. Recognition of transmembrane regions has been found to be remarkably successful, also leading to the acceptance of the output of these programs as annotation in sequence databases.

Certain periodicities in the occurrence of hydrophilic and hydrophobic residues may indicate particular secondary structures. Exposed helices tend to display a hydrophilic residue every 3–4 positions, while the other residues are rather hydrophobic. This is well visible in a “helical wheel representation” where the amino acid sequence is printed on a circle with the appropriate 3.6 amino acids per  $360^\circ$ . Likewise, a strand whose one side is exposed may display a hydrophilic residue at every other position. Based on these observations, Eisenberg devised the method of hydrophobic moments [11]. Many structural proteins interact with each other through intertwined helices forming a so-called coiled-coil structure. The physical constraints on the interface between these helices is reflected in a certain periodic arrangement of hydrophilic and hydrophobic amino acids. Generally, the sequence of an alpha helix that participates in a coiled-coil region will display a periodicity with a repeated unit of length 7 amino acids. Denote those 7 positions by  $a$  through  $g$ , then position  $e$  and  $g$  tend to be charged or polar while  $a$  and  $d$  are hydrophobic [12]. Prediction methods for coiled-coil regions are making use of these preferences [13] and even attempt to distinguish between two-stranded and three-stranded coiled-coils [14].

The primary sequence of a protein also contains the information whether a protein is secreted or which cellular compartment it is destined for. In particular, secretory proteins contain a N-terminal signal peptide for the recognition of which programs have been developed [15]. Other signals like, e.g., nuclear localization signals are notoriously hard to describe. Many posttranslational modifications, on the other hand, are linked to particular amino acid patterns as described in the Prosite database.

## 2.2.1

**Secondary structure prediction**

Linus Pauling already suggested that amino acid chains could assume regular local structures, namely alpha helices and beta strands [16]. In between these secondary structure elements there are turns or loops [17]. There is a long tradition of attempts to predict local secondary structure based on sequence. Secondary structure prediction generally aims at correlating the frequencies of occurrences of short amino acid words with particular secondary structures. The data set for this statistic must be derived from known protein structures with the secondary structures assigned to the primary sequence which is in itself not a trivial task [18]. Based on such a learning set, statistical methods are applied that range from information theory (the GOR method [19]) to neural networks [20] and linear discriminant analysis [21].

The basic approach essentially due to Chou and Fasman [22] uses a log-odds ratio comparing the frequency with which a particular amino acid assumes, say, a helical structure over the overall frequency of an arbitrary amino acid to occur in an alpha helix. The logarithm of this ratio is a measure indicating when this residue preferentially assumes a helical structure (positive value) as opposed to avoiding a helix (negative value). These preference values can be learnt from a large data set. For a prediction the preference is used like one of the parameters from above. Values are averaged within a sliding window and the resulting curves for helix, strand and loop prediction are compared, looking for the highest one in every part of the sequence.

The GOR method phrases a very similar approach in an information theoretic framework, computing not only preferences for individual residues but aiming at the delineation of preferences for short stretches of amino acids. Since the given data set will generally not supply sufficient data for estimation of the log-odds for every  $k$ -tuple certain approximations have to be made [23]. At the same time it has become clear that even this approach is unlikely to give perfect predictions because, in known crystal structures, one and the same 5-mer of residues will be found in different secondary structures [24]. Other approaches like the one due to Solov'yev and Salamov [25] assemble different characteristics for a short stretch (singlet and doublet secondary structure preferences, hydrophobic moment) of amino acids and apply linear discriminant analysis in order to derive a predictor for the secondary structure of a region.

While the information contained in one primary sequence alone seems to be insufficient to predict a protein's secondary structure, multiply aligned sequences (see below) offer a means to push the limits of the prediction. Originally due to Zvebil et al. [26] this line of attack on the problem is the basis of the PHD method by Rost and Sander [27]. The PHD method takes as input a multiple alignment of a set of homologous protein sequences and

uses several neural networks to effect the prediction. The first neural network maps the distribution vectors of amino acid occurrences in a window of adjacent alignment columns onto one of the states helix, strand, and loop. Outputs of this neural net are then reconciled by a second neural net which computes a consensus prediction. A last level of consensus-taking combines several predictions from neural nets that were trained on different data or with different techniques. The learning set for this procedure is taken from the HSSP database [28] of multiply aligned sequences that are annotated with their common secondary structure.

Development of prediction methods can, of course, only become fruitful when there are standards available to test and compare methods. It has been the achievement of Kabsch and Sander [29] to put forward a validation scheme for secondary structure prediction. This is based on an automatic assignment of secondary structures for a given crystal structure followed by an assessment of the degree of accuracy of helix, sheet and loop predictions. As long as one restricts oneself to the problem of secondary structure prediction for a single sequence the limit in predicting accuracy today seems to lie around 63% [30]. PHD claims an accuracy up to 72% due to the inclusion of multiply aligned sequences. In practical applications, however, the concrete data set will decide about the success of secondary structure prediction. Even when a substantial number of protein sequences can be aligned prior to predicting secondary structure, the information that they contribute depends on the degree of divergence between them. Some methods (like GOR and PHD, among others) supply the user with an estimate of how reliable a prediction in a particular area is which is of course helpful in practice. The success of overlaying the output from several secondary structure prediction programs is hard to predict because it is not clear whether the individual methods are sufficiently different to actually produce new information through such an approach. Recent methods by Cuff et al. [31] and by Selbig et al. [32] have automated and evaluated this approach.

The expasy server [33] offers references to many Web sites for protein sequence analysis and secondary structure prediction.

### 2.3 Pairwise sequence comparison

The methods of the last chapter analyzed a sequence by its own virtues. We now turn to the comparison of two sequences. The rationales behind the comparison of sequences may be manifold. Above all, the theory of evolution tells us that gene sequences may have derived from common ancestral sequences. Thus it is of interest to trace the evolutionary history of mutations and other evolutionary changes. Comparison of biological sequences in this context is understood as comparison based on the criteria of evolu-

tion. For example, the number of mutations, insertions, and deletions of bases necessary to transform one DNA sequence into another one is a measure reflecting evolutionary relatedness. On the other hand, a comparison may be more pragmatic in that it is not aimed at a detailed reconstruction of the evolutionary course of events but rather aims at pinpointing regions of common origin which may in turn coincide with regions of similar structure or similar function. Physical characteristics of amino acids play a more important role in this viewpoint than they do when studying evolution.

### 2.3.1

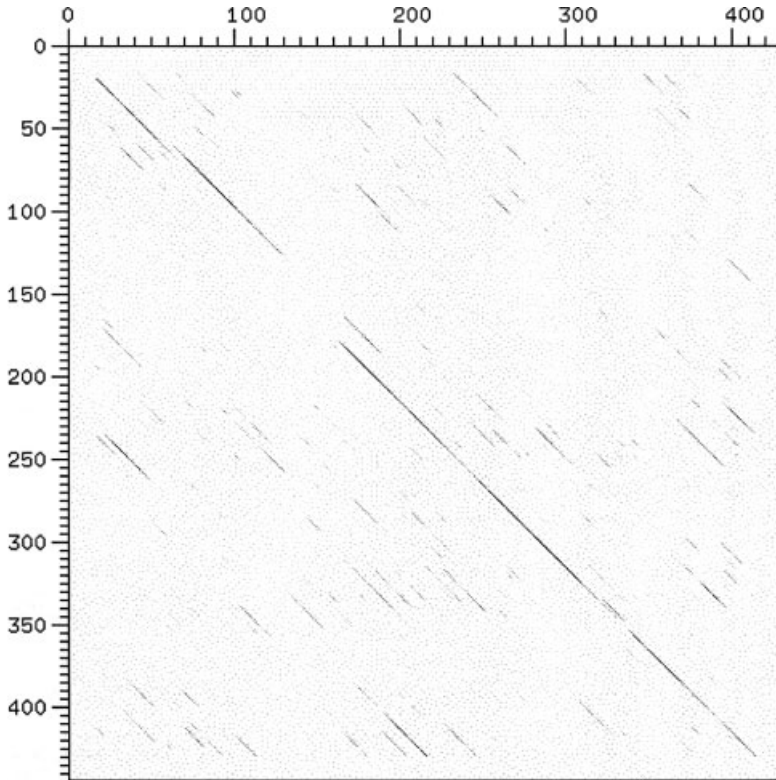
#### Dot plots

Dot plots are probably the oldest way of comparing sequences [34]. A dot plot is a visual representation of the similarities between two sequences. Each axis of a rectangular array represents one of the two sequences to be compared. A window length is fixed, together with a criterion when two sequence windows are deemed to be similar. Whenever one window in one sequence resembles another window in the other sequence, a dot or short diagonal is drawn at the corresponding position of the array. Thus, when two sequences share similarity over their entire length a diagonal line will extend from one corner of the dot plot to the diagonally opposite corner. If two sequences only share patches of similarity this will be revealed by diagonal stretches.

Figure 2.2 shows an example of a dot plot. There, the coding DNA sequences of the alpha chain of human hemoglobin and of the beta chain of human hemoglobin are compared to each other. For this computation the window length was set to 31. The program adds up the matches within a window and the grayscale value of the diagonal is set according to the quality of the match at that position. One can clearly discern a diagonal trace along the entire length of the two sequences. Note the jumps where this trace changes to another diagonal of the array. These jumps correspond to position where one sequence has more (or fewer) letters than the other one. The figure was produced using the program “dotter” [35].

Dot plots are a powerful method of comparing two sequences. They do not predispose the analysis in any way such that they constitute the ideal first-pass analysis method. Based on the dot plot the user can decide whether he deals with a case of global, i.e., beginning-to-end similarity, or local similarity. “Local similarity” denotes the existence of similar regions between two sequences that are embedded in the overall sequences which lack similarity. Sequences may contain regions of self-similarity which are frequently termed internal repeats. A dot plot comparison of the sequence itself will reveal internal repeats by displaying several parallel diagonals.

Instead of simply deciding when two windows are similar a quality function may be defined. In the simplest case, this could be the number of



**Fig. 2**  
Dot plot comparing two hemoglobin sequences.

matches in the window. For amino acid sequences the physical relatedness between amino acids may give rise to a quantification of the similarity of two windows. For example, when a similarity matrix on the amino acids (like the Dayhoff matrix, see below) is used one might sum up these values along the window. However, when this similarity matrix contains unequal values for exact matches this leads to exactly matching windows of different quality. The dot plot method of Argos [36] is an intricate design that reflects the physical relatedness of amino acids. The program *dotter* [37] is an X-windows based program that allows to display dot plots for DNA, for proteins, and for comparison of DNA to protein.

### 2.3.2

#### **Sequence alignment**

A sequence alignment [38] is a scheme of writing one sequence on top of another where the residues in one position are deemed to have a common

evolutionary origin. If the same letter occurs in both sequences then this position has been conserved in evolution (or, coincidentally, mutations from another ancestral residue has given rise to the same letter twice). If the letters differ it is assumed that the two derive from an ancestral letter, which could be one of the two or neither. Homologous sequences may have different length, though, which is generally explained through insertions or deletions in sequences. Thus, a letter or a stretch of letters may be paired up with dashes in the other sequence to signify such an insertion or deletion. Since an insertion in one sequence can always be seen as a deletion in the other one sometimes uses the term 'indel', or simply 'gap'.

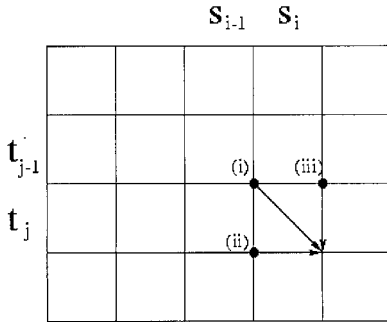
In such a simple evolutionarily motivated scheme, an alignment mediates the definition of a distance for two sequences. One generally assigns 0 to a match, some positive number to a mismatch and a larger positive number to an indel. By adding these values along an alignment one obtains a score for this alignment. A distance function for two sequences can be defined by looking for the alignment which yields the minimum score.

Naively, the alignment that realizes the minimal distance between two sequences could be identified by testing all possible alignments. This number, however, is prohibitively large but luckily, using dynamic programming, the minimization can be effected without explicitly enumerating all possible alignments of two sequences. To describe this algorithm [39], denote the two sequences by  $s = s_1, \dots, s_n$  and  $t = t_1, \dots, t_m$ . The key to the dynamic programming algorithm is the realization that for the construction of an optimal alignment between two stretches of sequence  $s_1, \dots, s_i$  and  $t_1, \dots, t_j$  it suffices to inspect the following three alternatives:

- (i) the optimal alignment of  $s_1, \dots, s_{i-1}$  with  $t_1, \dots, t_{j-1}$ , extended by the match between  $s_i$  and  $t_j$ ;
- (ii) the optimal alignment of  $s_1, \dots, s_{i-1}$  with  $t_1, \dots, t_j$ , extended by matching  $s_i$  with a gap character '-';
- (iii) the optimal alignment of  $s_1, \dots, s_i$  with  $t_1, \dots, t_{j-1}$ , extended by matching a gap character '-' with  $t_j$ .

Each of these cases also defines a score for the resulting alignment. This score is made up of the score of the alignment of the so far unaligned sequences that is used plus the cost of extending this alignment. In case: (i) this cost is determined by whether or not the two letters are identical and in cases (ii) and (iii) the cost of extension is the penalty assigned to a gap. The winning alternative will be the one with the best score (Figure 2.3).

To implement this computation one fills in a matrix the axes of which are annotated with the two sequences  $s$  and  $t$ . It is helpful to use north, south, west, and east to denote the sides of the matrix. Let the first sequence extend from west to east on the north side of the matrix. The second sequence extends from north to south on the west side of the matrix. We want to fill the matrix starting in the north-western corner, working our way southward



**Fig. 3** Schematic representation of the edit matrix comparing two sequences. The arrows indicate how an alignment may end according to the three cases described in the text.

row by row, filling each row from west to east. To start one initializes the northern and western margin of the matrix, typically with gap penalty values. After this initialization the above rules can be applied. A cell  $(i, j)$  that is already filled contains the score of the optimal alignment of the sequence  $s_1, \dots, s_i$  with  $t_1, \dots, t_j$ . The score of each such cell can be determined by inspecting the cell immediately north-west of it (case (i)), the cell west (case (ii)), and the one north (case (iii)) of it and deciding for the best scoring option. When the procedure reaches the south-eastern corner that last cell contains the score of the best alignment. The alignment itself can be recovered as one backtracks from this cell to the beginning, each time selecting the path that had given rise to the best option.

The idea of assigning a score to an alignment and then minimizing or maximizing over all alignments is at the heart of all biological sequence alignment. However, many more considerations have influenced the definition of the scores and made sequence alignment applicable to a wide range of biological settings. Firstly, note that one may either define a distance or a similarity function to an alignment. The difference lies in the interpretation of the values. A distance function will define positive values for mismatches or gaps and then aim at minimizing this distance. A similarity function will give high values to matches and low values to gaps and then maximize the resulting score. The basic structure of the algorithm is the same for both cases. In 1981, Smith and Waterman [40] showed that for global alignment, i.e., when a score is computed over the entire length of both sequences, the two concepts are in fact equivalent. Thus, it is now customary to choose the setting that gives more freedom in appropriately modeling the biological question that one is interested in.

In the similarity framework one can easily distinguish among the different possible mismatches and also among different kinds of matches. For example, a match between two Tryptophanes is usually seen to be more important than a match between two Alanines. Likewise, the pairing of two hydrophobic amino acids like Leucine and Isoleucine is preferable to the pairing of a hydrophobic with a hydrophilic residue. Scores are used to de-

scribe these similarities and are usually represented in the form of a symmetric 20 by 20 matrix, assigning a similarity score to each pair of amino acids. Although easy to understand from the physical characteristics of the amino acids, the values in such a matrix are usually derived based on an evolutionary model that allows one to estimate whether particular substitutions are preferred or avoided. This approach has been pioneered by M. Dayhoff [41] who computed a series of amino acid similarity matrices. Each matrix in this series corresponds to a particular evolutionary distance among sequences. This distance is measured in an arbitrary unit called 1 PAM, for 1 Accepted Point Mutation (in 100 positions). The matrices carry names like PAM120 or PAM250 and are supposed to be characteristic for evolutionary distances of 120 or 250 PAM, respectively. Other more recent series of matrices are the BLOSUM matrices [42] or Gonnet's [43] matrices. For every matrix one needs to find appropriate penalties for gaps.

The treatment of gaps deserves special care. The famous algorithm by Needleman and Wunsch [44] did not impose any restrictions on the penalty assigned to a gap of a certain length. For reasons of computational speed this was later specified to assigning a cost function linear in the number of deleted (inserted) residues [45]. This amounts to penalizing every single indel. However, since a single indel tends to be penalized such that it is considerably inferior to a mismatch, this choice resulted in longer gaps being quite expensive and thus unrealistically rare. As a remedy, one mostly uses a gap penalty function which charges a *gap open penalty* for every gap that is introduced and penalizes the length with a *gap extension penalty* which is charged for every inserted or deleted letter in that gap. Clearly, this results in an affine linear function in the gap length, frequently written as  $g(k) = a + b * k$  [46].

With the variant of the dynamic programming algorithm first published by Gotoh [47] it became possible to compute optimal alignments with affine linear gap penalties in time proportional to the product of the lengths of the two sequences to be aligned. This afforded a speed-up by one order of magnitude compared to a naive algorithm using the more general gap function. A further breakthrough in alignment algorithms development was provided by an algorithm that could compute an optimal alignment using computer memory only proportional to the length of one sequence instead of their product. This algorithm by Myers and Miller [48] is based on work by Hirschberg [49].

Depending on the biological setting several kinds of alignment are in use. When sequences are expected to share similarity extending from the beginning of the sequences to their ends they are aligned globally. This means that each residue of either sequence is part either of a residue pair or a gap. In particular it implies that gaps at the ends are charged like any other gap. This, however, is a particularly unrealistic feature of a global alignment. While sequences may very well share similarity over their entire length (see the example dot plot of two hemoglobin chains in Figure 2.2), their spec-



tive N- and C-termini usually are difficult to match up and differences in length at the ends are more of a rule than an exception. Consequently, one prefers to leave gaps at the ends of the sequences un-penalized. This variant is easy to implement in the dynamic programming algorithm. Two modifications are required. Firstly the initialization of the matrix needs to reflect the gap cost of 0 in the margin of the matrix. Secondly, upon backtracking, one does not necessarily start in the corner of the matrix but much rather searches the margins for the maximum from which to start. Variants of this algorithm that penalize only particular end-gaps are easy to derive and can be used, e.g., to fit one sequence into another or to overlap the end of one sequence with the start of another.

In many cases, however, sequences share only a limited region of similarity. This may be a common domain or simply a short region of recognizable similarity. This case is dealt with by so-called local alignment in an algorithm due to Smith and Waterman [50]. Local alignment aims at identifying the best pair of regions, one from each sequence, such that the optimal (global) alignment of these two regions is the best possible. This relies on a scoring scheme that maximizes a similarity score because otherwise an empty alignment would always yield the smallest distance. Naively, the algorithm to compute a local alignment would need to inspect every pair of regions and apply a global alignment algorithm to it. The decisive idea of Smith and Waterman was to offer the maximization in each cell of the matrix a fourth alternative: a zero to signify the beginning of a new alignment. After filling the dynamic programming matrix according to this scheme, backtracking starts from the cell in the matrix that contains the largest value.

Upon comparing a dot plot and a local alignment one might notice regions of similarity visible in the dot plot but missing in the alignment. While in many cases there exist gap penalty settings that would include all interesting matching regions in the alignment, generally it requires the comparison with the dot plot to notice possible misses. This problem is remedied by an algorithm due to Waterman and Eggert [51] which computes suboptimal, local, and non-overlapping alignments. It starts with the application of the Smith-Waterman algorithm, i.e., a dynamic programming matrix is filled and backtracking from the matrix cell with the largest entry yields the best local alignment. Then the algorithm proceeds to delineate a second-best local alignment. Note that this cannot be obtained by backtracking from the second-best matrix cell. Such an approach would yield an alignment largely overlapping the first one and thus containing little new information. Instead, those cells in the dynamic programming matrix are set to zero from where backtracking would lead into the prior alignment. This can be seen as “resetting” the dynamic programming matrix after having deleted the first alignment. Then the second best alignment is identified by looking for the maximal cell in the new matrix and starting backtracking from there. Iteration of this procedure yields one alternative,

non-overlapping alignment after the other in order of descending quality. Application of this algorithm avoids possibly missing matching regions because even under strong gap penalties the procedure will eventually show all matching regions.

There is an interesting interplay between parameters, in particular the gap penalty, and the algorithmic variant used. Consider a pair of sequences whose similar regions can in principle be strung together into an alignment (as opposed to sequences containing repeats which cannot all be seen in a single alignment). Under a weak gap penalty the Smith–Waterman algorithm has a chance to identify this entire alignment. On the other hand, not knowing about the similarity between the sequences ahead of time, a weak gap penalty might also yield all kinds of spurious aligned regions. The Waterman–Eggert algorithm is a valid alternative. The gap penalty can be chosen fairly stringently. The first (i.e., the Smith–Waterman) alignment will then identify only the best matching region out of all the similar regions. By iterating the procedure, though, this algorithm will successively identify the other similar regions as well. For a detailed discussion of these issues see Vingron and Waterman [52].

## 2.4

### **Database searching I: single sequence heuristic algorithms**

This section takes a first look at the problem of identifying those sequences in a sequence database that are similar to a given sequence. This task arises, e.g., when a gene has been newly sequenced and one wants to determine whether a related sequence already exists in a database. Generally, two settings can be distinguished. The starting point for the search may either be a single sequence with the goal of identifying its relatives, or a family of sequences with the goal of identifying further members of that family. Searching a data base needs to be fast and sensitive but the two objectives contradict each other. Fast methods have been developed primarily for searching with a single sequence and this shall be the topic of this section.

When searching a database with a newly determined DNA or amino acid sequence – the so-called query sequence – the user will typically lack knowledge of whether an expected similarity might span the entire query or just part of it. Likewise, he will be ignorant of whether the match will extend along the full length of some database sequence or only part of it. Therefore, one needs to look for a local alignment between the query and any sequence in the database. This immediately suggests the application of the Smith–Waterman algorithm to each database sequence. One should take care, though, to apply a fairly stringent gap penalty such that the algorithm focuses on the regions that really match. After sorting the resulting scores the top scoring database sequences are the candidates one is interested in.

Several implementations of this procedure are available, most prominently the SSEARCH program from the FASTA package [53]. There exist implementations of the Smith–Waterman algorithm that are tuned for speed like one using special processor instructions [54] and, among others, one by Barton [55]. Depending on implementation, computer, and database size, a search with such a program will take on the order of one minute.

The motivation behind the development of other database search programs has been to emulate the Smith–Waterman algorithm’s ability to discern related sequences while at the same time performing the job in much less time. To this end, one usually makes the assumption that any good alignment as one wishes to identify contains, in particular, some stretch of ungapped similarity. Furthermore, this stretch will tend to contain a certain number of identically matching residues and not only conservative replacements. Based on these assumptions most heuristic programs rely on identifying a well-matching core and then extending it or combining several of these. With hindsight, the different developments in this area can further be classified according to a traditional distinction in computer science according to which one either preprocesses the query or the text (i.e., the database). Preprocessing means that the string is represented in different form that allows for faster answer to particular questions, e.g., whether it contains a certain subword.

The FASTA program (part of a package [56] that usually goes by the same name) sets a size  $k$  for  $k$ -tuple subwords. The program then looks for diagonals in the comparison matrix between query and search sequence along which many  $k$ -tuples match. This can be done very quickly based on a preprocessed list of  $k$ -tuples contained in the query sequence. The set of  $k$ -tuples can be identified with an array whose length corresponds to the number of possible tuples of size  $k$ . This array is linked to the indices where the particular  $k$ -tuples occur in the query sequence. Note that a matching  $k$ -tuple at index  $i$  in the query and at index  $j$  in the database sequence can be attributed to a diagonal by subtracting the one index from the other. Therefore, when inspecting a new sequence for similarity, one walks along this sequence inspecting each  $k$ -tuple. For each of them one looks up the indices where it occurs in the query, computes the index-difference to identify the diagonal and increases a counter for this diagonal. After inspecting the search sequence in this way a diagonal with a high count is likely to contain a well-matching region. In terms of the execution time, this procedure is only linear in the length of the database sequence and can easily be iterated for a whole database. Of course this rough outline needs to be adapted to focus on regions where the match density is high and link nearby, good diagonals into alignments.

The other widely used program to search a database is called BLAST [57, 58]. BLAST follows a similar scheme in that it relies on a core similarity, although with less emphasis on the occurrence of exact matches. This program also aims at identifying core similarities for later extension. The core

similarity is defined by a window with a certain match density on DNA or with an amino acid similarity score above some threshold for proteins. Independent of the exact definition of the core similarity, BLAST rests on the precomputation of all strings which are in the given sense similar to any position in the query. The resulting list may be on the order of thousand or more words long, each of which if detected in a database gives rise to a core similarity. In Blast nomenclature this set of strings is called the neighborhood of the query. The code to generate this neighborhood is in fact exceedingly fast.

Given the neighborhood, a finite automaton is used to detect occurrences in the database of any string from the neighborhood. This automaton is a program, constructed on the fly and specifically for the particular word neighborhood that has been computed for a query. Upon reading through a database of sequences, the automaton is given an additional letter at a time and decides whether the string that ends in this letter is part of the neighborhood. If so, BLAST attempts to extend the similarity around the neighborhood and if this is successful reports a match.

Like FASTA, BLAST has also been adapted to connect good diagonals and report local alignments with gaps. BLAST converts the database file into its own format to allow for faster reading. This makes it somewhat unwieldy to use in a local installation unless someone takes care of the installation. FASTA, on the other hand, is slower but easier to use. There exist excellent web servers that offer these programs, in particular at the National Center for Biotechnology Information (NCBI [59]) and at the European Bioinformatics Institute (EBI [60]) where BLAST or FASTA can be used on up-to-date DNA and protein databases.

According to the above mentioned distinction among search methods into those that preprocess the pattern and those that preprocess the text, there also is the option of transforming a DNA or amino acid database such that it becomes easier to search. This route was taken, e.g., by a group from IBM developing the FLASH [61] program. They devised an intricate though supposedly very space consuming technique of transforming the database into an index for storing the offsets of gapped  $k$ -tuples. The work of Heumann and Mewes [62] has focused on further developing the well known data structure of a suffix tree. A suffix tree allows for quick lookup of where subwords of arbitrary length occur in a database. The problem with a suffix tree is that its size is several times the magnitude of the original data which for a sequence database means that it is unlikely to fit into main memory. Heumann and Mewes developed a data structure that allows for efficient mapping to disk such that the size problems can be circumvented. Likewise, the QUASAR program by Burkhard et al. [63] preprocesses the database into a so-called suffix array, similar to a suffix tree yet simple to keep on disk.

Progress in computational speed using either specially designed or massively parallel hardware has led to the availability of extremely fast versions

of the Smith–Waterman algorithm. The EBI, among other institutions, offers a service where this algorithm is executed on a massively parallel computer resulting in search times of a few seconds. Companies like Compugen [64] or Paracel [65] have developed special hardware to do this job.

With the availability of expressed sequence tags (ESTs) it has become very important to match DNA sequence with protein sequence in such a way that a possible translation can be maintained throughout the alignment. Both the FASTA and the BLAST package contain programs for this and related tasks. When coding DNA is compared to proteins, gaps are inserted in such a way as to maintain a reading frame. Likewise, a protein sequence can be searched versus a DNA sequence database. The search of DNA vs. DNA with an emphasis on matching regions that allow for a contiguous translation is not so well supported. Although a dynamic programming algorithm for this task is feasible the existing implementation in BLAST compares all reading frames.

## 2.5

### Alignment and search statistics

Alignment score is the product of an optimization, mostly a maximization procedure. As such it tends to be a large number sometimes suggesting biological relatedness where there is none. In pairwise comparisons the user still has a chance to study an alignment by eye in order to judge its validity but upon searching an entire database automatic methods are necessary to attribute a statistical significance to an alignment score.

In the early days of sequence alignment, the statistical significance of the score of a given pairwise alignment was assessed using the following procedure. The letters of the sequences are permuted randomly and a new alignment score is calculated. This is repeated roughly 100 times and mean and standard deviation of this sample are calculated. The significance of the given alignment score is reported in ‘number of standard deviations above the mean’, also called the Z-score. Studying large numbers of random alignments is in principle correct. However, the significance of the alignment should then be reported as the fraction of random alignments that score better than the given alignment. The procedure described assumes that these scores were normally distributed. Since the random variable under study – the score of an optimal alignment – is the maximum over a large number of values this is not a reasonable assumption. In fact, when trying to fit a normal distribution to the data the lack of fit quickly becomes obvious. The second argument against this way of calculating significance is a pragmatic one. The procedure needs to be repeated for every alignment

under study because the effect of the sequence length cannot be accounted for.

Based on the work of several researchers [66, 67] it has meanwhile become apparent that alignment score as well as scores from database searches obey a so-called extreme value distribution. This is not surprising given that extreme value distributions typically describe random variables which are the result of a maximization. In sequence alignment, there are analytical results confirming the asymptotic convergence to an extreme value distribution for the case of local alignment without gaps, i.e., the score of the best-matching contiguous diagonal in a comparison [68]. This is also a valid approximation to the type of matching effected in the database search program BLAST. Thus this approach has become widely used and in fact has contributed significantly to the popularity of database search programs because significance measures have made the results of the search much easier to interpret.

The statistical significance of an event like observing a sequence alignment of a certain quality is the probability to observe a better value as a result of chance alone. Thus one needs to model chance alignments, which is precisely what the statistician means by deriving the distribution of a random variable. The probability that a chance result would exceed an actually obtained threshold  $S$  is 1 minus the value of the cumulative distribution function evaluated at that threshold. (If  $S$  is the score of a given sequence segment this probability is the famed P-value of the segment. It is to be contrasted with the E-value which gives the expected number of segments in the sequence that exceed the score. For interesting outcomes both of these values are very small and almost identical.) In sequence alignment, the respective cumulative distribution function is generally expressed as [69]:

$$\exp(-mnKe^{-\lambda S}).$$

Here,  $m$  and  $n$  are the lengths of the sequences compared and  $K$  and  $\lambda$  are parameters which need to be computed (where possible) or derived by simulation.  $K$  and  $\lambda$  depend on the scoring matrix used (e.g., the PAM120 matrix) and the distribution of residues. Hence, for any scoring system these parameters are computed beforehand and the statistical significance of an alignment score  $S$  is then computed by evaluating the formula with the length of the two sequences compared.

The most prominent case where the parameters  $K$  and  $\lambda$  can be defined analytically is local alignment without gaps. Algorithmically this amounts to computing a Smith–Waterman alignment under very high gap penalties such that the resulting alignment will simply not contain any gaps. Since this notion of alignment also guides the heuristic used by the BLAST database search program the resulting statistical estimates are primarily used in

database searching. In this application, one of the lengths is the length of the input sequence and the other length is on the order of the length of the concatenated sequences from the database that is being searched.

When gaps are allowed the situation is more complex because an approximation of the distribution function of alignment score by an extreme value distribution as above is not always valid. Generally speaking, it is allowed only for sufficiently strong gap penalties where alignments remain compact as opposed to spanning the entire sequences. Under sufficiently strong gap penalties, though, it has been demonstrated that the approximation is indeed valid just like for infinite gap penalties [70]. However, it is not possible any more to compute analytically the values of the parameters  $K$  and  $\lambda$ . As a remedy, one applies simulations where many alignments of randomly generated sequences are computed and the parameters are determined based on fitting the empirical distribution function with an extreme value distribution [71]. Like in the case above, this procedure allows to determine parameters beforehand and compute significance by putting the lengths of the sequences into the formula.

The question remains how to determine whether approximation by an extreme value distribution is admissible for a certain scoring scheme and gap penalty setting one is using. This can be tested on randomly generated (or, simply, unrelated) sequences by computing a global alignment between sequences under that particular parameter setting. If the result has negative sign (averaged over many trials or on very long random sequences) then the approximation is admissible. This is based on a theorem due to Arratia and Waterman [72] and subsequent simulation results reported by Waterman and Vingron [73]. In particular, a gap open penalty of 12 with extension penalty of 2 or 3 for the case of the PAM250 matrix, as well as any stronger combination allow for approximation by the extreme value distribution.

In database searching the fitting need not be done on randomly generated sequences. Under the assumption that the large majority of sequences in a database are not related to the query, the bulk of the scores generated upon searching can be used for fitting. This approach is taken by W. Pearson in the FASTA package. It has the advantage that the implicit random model is more realistic since it is taken directly from the data actually searched. Along a similar line of thought, Spang and Vingron [74] tested significance calculations in data base searching by evaluating a large number of search results. Their study showed that one should not simply use the sum of the lengths of all the sequences in the database as the length-parameter in the formula for the extreme value distribution. This would overestimate the length that actually governs the statistics. Instead, a considerably shorter effective length can be determined for a particular database using simulations. This effect is probably due to the fact that alignments cannot start in one sequence and end in the next one which makes the number of feasible starting points for random alignments smaller than the actual length of the database.

## 2.6

### Multiple sequence alignment

For many genes a database search will reveal a whole number of homologous sequences. One then wishes to learn about the evolution and the sequence conservation in such a group. This question surpasses what can reasonably be achieved by the sequence comparison methods described in Section 2.3. Pairwise comparisons do not readily show positions that are conserved among a whole set of sequences and tend to miss subtle similarities that become visible when observed simultaneously among many sequences. Thus, one wants to simultaneously compare several sequences.

A multiple alignment arranges a set of sequences in a scheme where positions believed to be homologous are written in a common column. Like in a pairwise alignment, when a sequence does not possess an amino acid in a particular position this is denoted by a dash. There also are conventions similar to the ones for pairwise alignment regarding the scoring of a multiple alignment. The so-called sum-of-pairs (SOP) [75] score adds the scores of all the induced pairwise alignments contained in a multiple alignment. For a linear gap penalty this amounts to scoring each column of the alignment by the sum of the amino acid pair scores or gap penalties in this column. Although it would be biologically meaningful, the distinctions between global, local, and other forms of alignment are rarely made in a multiple alignment. The reason for this will become apparent below, when we describe the computational difficulties in computing multiple alignments.

Note that the full set of optimal pairwise alignments among a given set of sequences will generally overdetermine the multiple alignment. If one wishes to assemble a multiple alignment from pairwise alignments one has to avoid “closing loops”, i.e., one can put together pairwise alignments as long as no new pairwise alignment is included to a sequence which is already part of the multiple alignment. In particular, pairwise alignments can be merged when they align one sequence to all others, when a linear order of the given sequences is maintained, or when the sequence pairs for which pairwise alignments are given form a tree. While all these schemes allow for the ready definition of algorithms that output multiply aligned sequences, they do not include any information stemming from the simultaneous analysis of several sequences.

An alternative approach is to generalize the dynamic programming optimization procedure applied for pairwise alignment to the delineation of a multiple alignment that maximizes, e.g., the SOP score. The algorithm used [76] is a straight-forward generalization of the global alignment algorithm. This is easy to see, in particular, for the case of the column-oriented SOP scoring function avoiding affine gap penalty in favor of the simpler linear one. With this scoring, the arrangement of gaps and letters in a column can be represented by a Boolean vector indicating which sequences contain a gap in a particular column. Given the letters that are being com-



pared, one needs to evaluate the scores for all these arrangements. However conceptually simple this algorithm may be, its computational complexity is rather forbidding. For  $n$  sequences it is proportional to  $2^n$  times the product of the lengths of all sequences.

In practice this algorithm can be run only for a modest number of sequences being compared. There exists software to compare three sequences with this algorithm that additionally implements a space-saving technique [77]. For more than three sequences, algorithms have been developed that aim at reducing the search space while still optimizing the given scoring function. The most prominent program of this kind is MSA2 [78, 79]. An alternative approach is used by DCA [80, 81] which implements a divide-and-conquer philosophy. The search space is repeatedly subdivided by identifying anchor points through which the alignment is highly likely to pass.

None of these approaches, however, would work independent of the number of sequences to be aligned. The most common remedy is reducing the multiple alignment problem to an iterated application of the pairwise alignment algorithm. However, in doing so, one also aims at drawing on the increased amount of information contained in a set of sequences. Instead of simply merging pairwise alignments of sequences, the notion of a profile [82] has been introduced in order to grasp the conservation patterns within subgroups of sequences. A profile is essentially a representation of an already computed multiple alignment of a subgroup. This alignment is “frozen” for the remaining computation. Other sequences or other profiles can be compared to a given profile based on a generalized scoring scheme defined for this purpose. The advantage of scoring a sequence versus a profile over scoring individual sequences lies in the fact that the scoring schemes for profile matching reflect the conservation patterns among the already aligned sequences.

Given a profile and a single sequence, the two can be aligned using the basic dynamic programming algorithm together with the accompanying scoring scheme. The result will be an alignment between sequence and profile that can readily be converted into a multiple alignment now comprising the sequences underlying the profile plus the new one. Likewise, two profiles can be aligned with each other resulting in a multiple alignment containing all sequences from both profiles. With these tools various multiple alignment strategies can be implemented. Most commonly, a hierarchical tree is generated for the given sequences which is then used as a guide for iterative profile construction and alignment. This alignment strategy was introduced in papers by Taylor [83], Corpet [84], and Higgins [85]. Higgins’ program Clustal [86] has meanwhile become the de facto standard for multiple sequence alignment [87]. Other programs in practical use are the MSA2 program, DCA, and Dialign. Dialign [88, 89] is different in that it aims at the delineation of regions of similarity among the given

sequences. Chapter 8 will discuss approaches to multiple sequence comparison where common patterns are sought rather than overall alignments.

Since iterative profile alignment tends to be guided by a hierarchical tree, this step of the computation is also influencing the final result. Usually the hierarchical tree is computed based on pairwise comparisons and their resulting alignment scores. Subsequently this score matrix is used as input to a clustering procedure like single linkage clustering or UPGMA [90]. However, it is well understood that in an evolutionary sense such a hierarchical clustering does not necessarily result in a biologically valid tree. Thus, when allowing this tree to determine the multiple alignment there is the danger of pointing further evolutionary analysis of this alignment into the wrong direction. Consequently, the question has arisen of a common formulation of evolutionary reconstruction and multiple sequence alignment. The cleanest although biologically somewhat simplistic model attempts to reconstruct ancestral sequences to attribute to the inner nodes of a tree [91]. Such reconstructed sequences at the same time determine the multiple alignment among the sequences. In this 'generalized tree alignment' one aims at minimizing the sum of the edge-lengths of this tree, where the length of an edge is determined by the alignment distance between the sequences at its incident nodes. As to be expected, the computational complexity of this problem again makes its solution unpractical. The practical efforts in this direction go back to the work of Sankoff [92]. Hein [93] and Schwikowski and Vingron [94] produced software [95, 96] relying on these ideas.

## 2.7

### **Multiple alignments and database searching**

Information about which residues are conserved and thus important for a particular family is crucial not only for the purpose of multiply aligning a set of sequences. Also in the context of identifying related sequences in a database this information is very valuable. A multitude of methods has been developed that aim at identifying sequences in a database which are related to a given family. The first one was the notion of a profile that was described above and was actually introduced in the context of data base searching. Like in multiple alignment, profiles help in emphasizing conserved regions in a database search. Thus, a sequence that matches the query profile in a conserved region will receive a higher score than a database sequence matching only in a divergent part of an alignment. This feature is of enormous help in distinguishing truly related sequences.

Algorithmically, profile searching simply uses the dynamic programming alignment algorithm for aligning a sequence to a profile on each sequence

in the database. Of course, this is computationally quite demanding and much slower than the heuristic database search algorithms like BLAST or FASTA. Typically, the multiple alignment underlying the profile will describe a conserved domain which one expects to find within a database sequence. Therefore, in this context it is important that end gaps should not be penalized. Furthermore, gap penalties for profile matching frequently vary along the profile in order to reflect the existence of gaps within the underlying multiple alignment. Through this mechanism, one attempts to allow new gaps preferentially in regions where gaps have been observed already. However, different suggestions exist as to the choice and derivation method for these gap penalties [97].

In 1994, Haussler and co-workers [98] introduced Hidden Markov Models (HMMs) for the purpose of identifying family members in a database. HMMs are a class of probabilistic models well suited for describing the relevant parameters in matching a given multiple alignment against a database. For HMMs there exist automatic learning algorithms that adapt the parameters of the HMM for best identification of family members. Thus, they offer in particular a solution to the question of gap penalty settings along a profile. Sequences are matched to HMMs in much the same way as they are aligned to a profile although the interpretation of the procedure is different. The HMM is thought of as producing sequences by going through different states. Aligning a sequence to an HMM amounts to delineating the series of model-states that is most likely to have produced the sequence.

Based on this interpretation one can make an interesting distinction between the optimal alignment of a sequence with a HMM and the computation of the probability that a model has produced a particular sequence. The optimal alignment is computed with an algorithm exactly analogous to the dynamic programming algorithm and maximizing the probability that a series of states has given rise to a particular sequence. In contrast hereto, in absence of knowledge of the correct path of states, the probability that a model has given rise to a particular sequence should rather be computed as the sum over the different sets of states that could have produced the sequence. This interpretation leads to a summation over all paths instead of the choice of the best one. These issues are discussed by Durbin et al. [99] (Section 2.5.4) although, practically, there is little known about the difference in performance between the two approaches. Bucher and Karplus [100] introduced generalized profiles and showed that the two concepts are equally powerful in their abilities to model sequence families and detect related sequences.

The fact that a profile or HMM can pick out new sequences also related to the given family suggests that these should be used to update the profile or HMM used as search pattern. This idea leads to iterative search algorithms where the database is searched repeatedly, each time updating the query pattern with some or all of the newly identified sequences. Psi-Blast [101] is a very successful implementation of this idea. It starts with a single

sequence and after the first search builds profiles from conserved regions among the query and newly identified sequences. Without allowing for gaps (to speed up the search) these new profiles are used to repeat the search. Generally, Psi-Blast quickly converges after updating these profiles again and generally is very successful in delineating all the conserved regions a sequence may share with other sequences in a database. In the realm of HMMs, SAM is a very careful implementation of the idea of iterated searches [102, 103].

It is the generally held view that searching a database with a profile or HMM will produce extreme-value distributed random scores, just like single-sequence database searching. The quality of the fit to the extreme-value distribution may depend on the particular given alignment, though. This has been substantiated with mathematical arguments only for the case of ungapped profile matching. Nevertheless, this basic understanding of the statistical behavior of database matching methods is a crucial element in the iterative search programs. Without clear and reliable cutoff values one could not decide which sequences to integrate into the next search pattern and would run the danger of including false positives, thus blurring the information in the pattern.

Both single sequence search methods and profile/HMM-based ones have been thoroughly validated during recent years [104]. Databases of structurally derived families like, e.g., SCOP [105, 106] have made it possible to search a sequence database with a query and exactly determine the number of false positives and false negatives. For every search one determines how many sequences one misses (false negatives) in dependence of the number of false positive matches. If the sequence statistics is accurate the number of false positives correlates well with the E-value which is the number of false positives expected by chance. Although not always quite decided, this way of validating search methods allows to make objective comparisons and to determine how much quality one actually gains with slower methods over faster, less accurate one.

## 2.8

### Protein families and protein domains

The dual question to the one that assigns related sequences from a database to a given query sequence or family is the question that tries to assign to a query sequence the family that it is a member of or the domains that it contains. One simple yet very effective resource for this purpose is the Prosite database [107, 108] which contains amino acid patterns that are descriptive for particular domains, families, or functions. These patterns allow to specify alternative residues in particular positions or variable length spacers between positions. Matching a sequence against a Prosite entry

amounts simply to looking for the particular pattern of characters in the given sequence. For many sequences that are contained in a protein sequence database this is not even necessary any more because they have already been annotated with the Prosite patterns they contain. The expasy server [109] offers the possibility to screen a sequence versus the entire collection of Prosite patterns.

For certain protein sequence domains character patterns may be rather poor descriptors, i.e., they may cause many false positive and false negative matches. The Pfam database remedies this situation by supplying pre-computed Hidden Markov Models for protein domains. A query sequence can be matched against this library of HMMs in order to identify known domains in the query sequence. Here, too, match statistics plays a crucial role in order to determine the significantly matching domains. A server that allows to scan a sequence versus all Pfam domains can be found at the Sanger center [110]. Software has also been developed to recognize the Pfam HMMs in either coding DNA or in genomic DNA. In the latter case the program combines the HMM matching with the distinction between coding and non-coding DNA. Recently, a database of domains, InterPro [111], has emerged that unifies sources like Prosite and Pfam.

Many attempts have been made at developing algorithms that will automatically determine domains shared by several input sequences. In a way this is a generalization of the multiple alignment problem in the direction of local alignment, however in this context it seems to be considerably more difficult than in the pairwise case. There exist two programs that identify Prosite-like patterns contained in many sequences. These are Pratt [112, 113] and TEIRESIAS [114, 115]. The program Blockmaker [116, 117] identifies non-gapped sequence blocks that are characteristic of a given set of sequences. This program has also been used to construct a large collection [118] of Blocks that can be searched like Prosite or Pfam. Krause and Vingron [119] used an iterated search procedure SYSTEMS [120] to delineate protein families and supply consensus sequences of these families to be searched with a DNA or protein query sequence. The most sophisticated programs for the detection of common sequence motifs in a given set use probabilistic modeling and/or machine learning approaches. In particular, the mathematical technique of the Gibbs sampler has lent its name also to a motif-finding program, the Gibbs Motif Sampler [121, 122]. Bailey and Elkan [123] designed the MEME [124] program which relies on an expectation maximization algorithm.

## 2.9

### Conclusion

The problems and methods introduced above have been instrumental in the advance in our understanding of genome function, organization, and

structure. While some years ago human experts would check every program output, nowadays sequence analysis routines are being applied in an automatic fashion creating annotation that is included in various database. This holds true for similarity relationships among sequences and extends all the way to the prediction of genomic structure or to function prediction based on similarity. Although the quality of the tools has increased dramatically, the possibility of error and in particular its perpetuation by further automatic methods exists. Thus, it is apparent that the availability of these high-throughput computational analysis tools is a blessing and a problem at the same time.

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### 3

## Structure, Properties and Computer Identification of Eukaryotic Genes

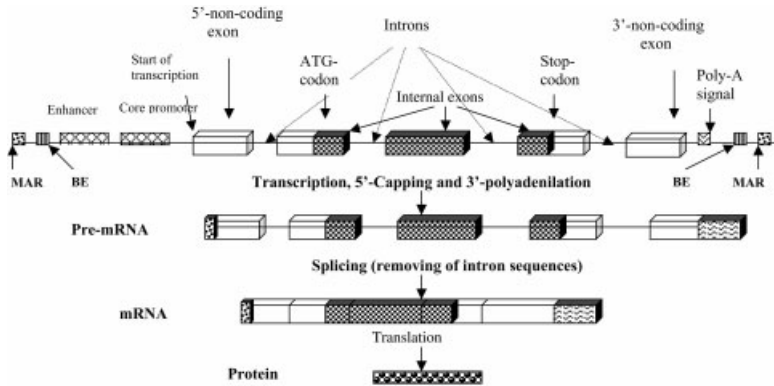
*Victor Solovyev*

### 3.1

#### Structural characteristics of eukaryotic genes

The gene is a fragment of nucleic sequence that carries the information representing a particular polypeptide or RNA molecule. In eukaryotes, genes lie in a linear array on chromosomes, which consist of a long molecule of duplex DNA and chromatin proteins (mostly histones that form a structure called a nucleosome). The complex of DNA and proteins (chromatin) can maintain genes in an inactive state by restricting access to RNA polymerase and its accessory factors. To activate a gene, the chromatin encompassing that gene and its control region must be modified to permit transcription [1]. The principal steps in gene expression of protein-coding genes are transcription and post-transcriptional processing of messenger RNA precursors including 5'-capping, 3'-polyadenylation and splicing. The order in which these events occur is not entirely clear: some splicing events could take place during transcription [2]. The processing events produce the mature mRNA in the nucleus and then it is transported to the cytoplasm for translation. The mature mRNA includes sequences called exons that encode the protein product according to the rules of the genetic code. However, the gene sequence often includes non-coding regions: introns that are removed from the primary transcript during RNA splicing and 5'- and 3'-untranslated regions. Model and stages of gene expression of a typical protein-coding gene are presented in Figure 3.1.

Knowledge about structural gene characteristics is accumulated in GenBank and EMBL nucleotide sequence databases, where one gene can be described in dozens of entries assigned to partially sequenced gene regions, alternative splicing forms or mRNAs. Processing GenBank [3] data, a gene-centered database InfoGene was created [4, 5], which contains description of known genes and their basic functional signals and structural components. All major organisms are presented in the separate divisions. The Human InfoGene division contains about 21 000 genes (including 16 000 partially sequenced genes), 53 435 coding regions, 83 488 exons and about



**Fig. 3.1**

Model of eukaryotic gene structure and gene expression stages. BEs are boundary elements that bind proteins to prevent the enhancer effects on outside genes. MARs are matrix attachment regions.

58 000 donor and acceptor splice sites. Table 3.1 shows the major structural characteristics of Human, Mouse, *D. melanogaster*, *C. elegans*, *S. cerevisiae* and *A. thaliana* genes deposited in GenBank, release 119.

We do not observe any significant difference in the size of protein coding mRNAs in different types of organisms, but the gene sizes are often larger in vertebrates and especially in primates. We can notice that the human coding exons are significantly shorter than the sizes of the respective genes. The average size of an exon is about 200 bp, that is close to the DNA length associated with the nucleosome particle. Coding and non-coding exons can be as short as several bases and as long as dozens or thousands of bases. Usually protein coding exons occupy just a few percent of the gene size. Different kind of repeats cover 41% of sequenced human DNA, and coding exons account only for 2–3% of the genomic sequence.

The structural characteristics of eukaryotic genes considered above create two major problems in computational gene identification.

1. Low quality recognizers will generate a lot of false positive predictions, the number of which is comparable with the true exon number (Figure 3.2);
2. Recognition of small exons (1–20 bp) can not be done using any composition-based coding measure that is often successful for prokaryotes.

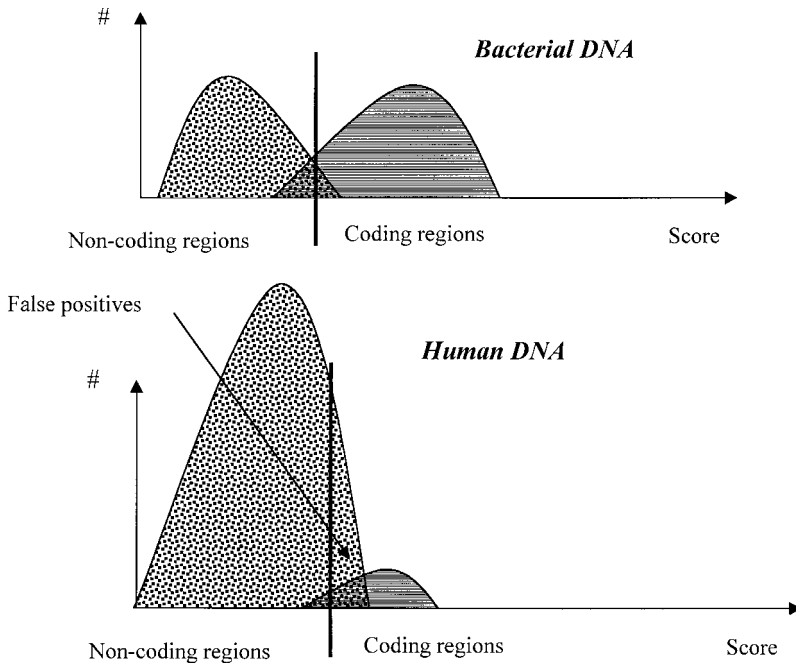
We need to develop gene prediction approaches that are based significantly on the recognition of functional signals encoded in the gene sequence.

The main information about exon location is encoded in splice site sequences. In the next section we will consider essential characteristics of splice sites. The other functional signals as promoter, poly-A, start and stop

Tab. 3.1

Structural characteristics of genes in eukaryotic model organisms. The numbers reflect gene characteristics described in GenBank (Release 119, 2000), which might deviate from the average parameters for organisms. Gene numbers were calculated for DNA loci only. Many long genes have partially sequence introns, therefore the actual average gene size is bigger. The range (or maximal) and average values are shown. For donor and acceptor sites the percentage of annotated canonical (GT-AG) splice pairs is provided.

	<i>Homo sapiens</i>	<i>Mus musculus</i>	<i>Drosophila melanogaster</i>	<i>C. elegans</i>	<i>S. cerevisiae</i>	<i>Arabidopsis thaliana</i>
CDS/partial	53435/29404	24527/13060	20314/1510	20634/526	12635/1016	31194/1461
Exons/partial	83488/21342	24508/7913	66960/19343	122951/38293	13572/13127	145942/42844
Genes/partial	20791/16141	7428/5573	17435/1154	19658/1263	12513/1098	28346/1023
Alternative splicing	2167, 10.4%	749, 10%	1785, 10%	1194, 6.1%	598, 4.7%	227, 0.1%
No introns genes	1552, 7.4%	748, 10%	3583, 20%	669, 3%	11070, 88.5%	5776, 19.7%
Number of exons	117, 5.7	64, 4.72	50, 3.88	52, 6.1	3, 1.03	78, 5.1
Exon length	1-1088, 201.6	1-6642, 207.1	6-10785, 419.5	1-14975, 22125	1-7471, 1500.0	3-75916, 192.0
Intron length	259776, 2203.5	42573, 818.3	205244, 613.7	19397, 244.0	7317, 300	118637, 174.4
Gene length	401910, 9033	150523, 3963	155515, 2854	45315, 2624	14733, 1462	170191, 2027
Donor sites	58707, 98.0%	6225, 96.9%	49592, 98.0%	102872, 99.5%	471, 93.0%	117658, 99.2%
Acceptor sites	58112, 98.53%	5627, 97.5%	49602, 97.9%	102933, 99.7%	475, 95.6%	121917, 96.9%



**Fig. 3.2**  
The huge amount of non-coding DNA in the human genome can significantly increase the number of false positive coding exon predictions. The score x-axis

reflects the value of a given classifying function to assign a sequence region to coding or non-coding classes. The y-axis presents the number of sequence fragments with a particular score.

of translation will be discussed later with description of their recognition functions used in many gene prediction programs.

### 3.2 Classification of splice sites in mammalian genomes

Since the discovery of split genes it was observed that practically all introns contain two very conservative dinucleotides. The donor site has GT exactly at the intron's 5'-boundary and the acceptor site has AG exactly at its 3'-boundary [6, 7]. We call splice sites of this type canonical. Introns flanked by the standard GT-AG pairs excised from pre-mRNA by the spliceosome including U1, U2, U4/U6 and U5 snRNPs [7]. Recently, a few examples of a new type of splice pair, a AT-AC, has been discovered. It is processed by a related, but different splicing machinery [8, 9]. AT-AC introns are excised by a novel type of spliceosome composed of snRNPs U11, U12, U4atac/U6atac, and U5 [10, 11, 12]. Several other cases of non-canonical splice sites with



GC-AG, GG-AG, GT-TG, GT-CG or CT-AG dinucleotides at the splice junctions have been reported [8, 13].

The most comprehensive investigation of different types of annotated splice sites has been performed by Bursset, Seledtsov and Solovyev [14]. They have extracted 43 337 pairs of exon-intron boundaries and their sequences from the InfoGene database [15] covering all annotated genes in mammalian genomic sequences. Annotation errors present a real problem in obtaining accurate information on eukaryotic gene functional signals from nucleotide sequence databases, such as GenBank or EMBL [3, 16]. This is especially crucial for the analysis of non-standard splice sites. For instance, carefully checking 50 such examples revealed 21 cases of clear EMBL annotation errors [17]. To verify the annotated splice sites Bursset et al. [14] used the alignment of spliced exons with known mammalian ESTs and high throughput genomic sequences.

Of 43 337 pairs of donor and acceptor splice sites (splice pairs) 22 489 were supported by EST sequences. 98.71% of those contain canonical dinucleotides GT and AG for donor and acceptor sites. 0.56% hold non-canonical GC-AG splice site pairs. The remainder 0.73% occurs in a lot of small groups (with maximum size of 0.05%). 53.6% of canonical and just 27.3% of non-canonical splice pairs were supported by ESTs. Based on these figures it was supposed that at least half of annotated non-canonical sites presents annotation errors, as was shown in some previous works [8, 17].

In addition to the conserved dinucleotides AG-GT, canonical splice sites demonstrate well defined, conserved donor site consensus: AG|GTRAGT<sup>1)</sup>, and acceptor site consensus: YTTTYYYYYNC|AGG [18]. For the much smaller AT-AC group different conserved positions have been noticed: |ATATCCTTT for donor site and YAC| for acceptor site [12, 19, 20]. These differences reflect some specific interactions with the components of splicing machineries and they can be used to judge if a particular splice site group belongs to the GT-AG or AT-AC splice system. Weight matrices for the GT-AG, GC-AG pairs constructed based on EST-supported splice pairs are presented in Table 3.2 and consensus sequences for the AT-AC pair are depicted in Figure 3.3.

The occurrence of canonical dinucleotides upstream or downstream in each EST-supported non-canonical splice site group was carefully analyzed. For example, the telethonin gene has only one intron annotated in positions 639–885. This junction is completely supported by ESTs and the annotated splice sites are GG-CA. Analyzing the telethonin sequence uncovered an occurrence of a canonical splice pair GT-AG just one position downstream from the annotated site (Figure 3.4). Taking into account that the non-canonical splice junctions occur very rarely, we can suspect that the canon-

1) Here, we use the common IUPAC ambiguity codes: e.g., R = A, G, Y = C, T, S = C, G, N = A, C, G, T

**Tab. 3.2**

Sequence composition of major group of splice sites.

**GT-AG group (98.70%).**

Frequency of nucleotides in donor splice site. 1 and 2 are positions of conserved GT dinucleotide.

%	-3	-2	-1	1	2	3	4	5	6
A	34.0	60.4	9.2	0.0	0.0	52.6	71.3	7.1	16.0
C	36.3	12.9	3.3	0.0	0.0	2.8	7.6	5.5	16.5
G	18.3	12.5	80.3	100	0.0	41.9	11.8	81.4	20.9
U	11.4	14.2	7.3	0.0	100	2.5	9.3	5.9	46.2

Frequency of nucleotides in acceptor splice site. -1 and -2 are positions of conserved AG dinucleotide.

%	-14	-13	-12	-11	-10	-9	-8	-7	-6	-5	-4	-3	-2	-1	1
A	9.0	8.4	7.5	6.8	7.6	8.0	9.7	9.2	7.6	7.8	23.7	4.2	100	0.0	23.9
C	31.0	31.0	30.7	29.3	32.6	33.0	37.3	38.5	41.0	35.2	30.9	70.8	0.0	0.0	13.8
G	12.5	11.5	10.6	10.4	11.0	11.3	11.3	8.5	6.6	6.4	21.2	0.3	0.0	100	52.0
U	42.3	44.0	47.0	49.4	47.1	46.3	40.8	42.9	44.5	50.4	24.0	24.6	0.0	0.0	10.4

**GC-AG group (0.56%).**

Frequency of nucleotides in donor splice site. 1 and 2 are positions of conserved GT dinucleotide.

%	-3	-2	-1	1	2	3	4	5	6
A	40.5	88.9	1.6	0.0	0.0	87.3	84.1	1.6	7.9
C	42.1	0.8	0.8	0.0	100	0.0	3.2	0.8	11.9
G	15.9	1.6	97.6	100	0.0	12.7	6.3	96.8	9.5
U	1.6	8.7	0.0	0.0	0.0	0.0	6.3	0.8	70.6

Frequency of nucleotides in acceptor splice site. -1 and -2 are positions of conserved AG dinucleotide.

%	-14	-13	-12	-11	-10	-9	-8	-7	-6	-5	-4	-3	-2	-1	1
A	11.1	12.7	3.2	4.8	12.7	8.7	16.7	16.7	12.7	9.5	26.2	6.3	100	0.0	21.4
C	36.5	30.9	19.1	23.0	34.9	39.7	34.9	40.5	40.5	36.5	33.3	68.2	0.0	0.0	7.9
G	9.5	10.3	15.1	12.7	8.7	9.5	16.7	4.8	2.4	6.3	13.5	0.0	0.0	100	62.7
U	38.9	41.3	58.7	55.6	42.1	40.5	30.9	37.3	44.4	47.6	27.0	25.4	0.0	0.0	7.9

Donor: S<sub>90</sub>|ATA<sub>100</sub>T<sub>100</sub>C<sub>100</sub>C<sub>100</sub>T<sub>100</sub>T<sub>90</sub>T<sub>70</sub>Acceptor: T<sub>70</sub>G<sub>50</sub>C<sub>70</sub>NC<sub>60</sub>AC|A<sub>60</sub>T<sub>60</sub>**Fig. 3.3**

Consensus sequences for the AT-AC pair of the alternative splicing machinery.

	Donor	Acceptor
<b>Annotated:</b>	... <b>CCCGAGGAGG</b>  ggtgagtgtg.....cctctcccca  <b>GCTGCTCCCT</b> ...	
<b>Possible:</b>	... <b>CCCGAGGAGGG</b>  gtgagtgtg.....cctctcccccag  <b>CTGCTCCCT</b> ...	
<b>Annotated junction:</b>	... <b>CCCGAGGAGG</b>   <b>GCTGCTCCCT</b> ...	
(EST=AI802984)		
<b>Possible junction:</b>	... <b>CCCGAGGAGGG</b>   <b>CTGCTCCCT</b> ...	

**Fig. 3.4**  
Possible errors in EST-supported splice pairs. Example of an annotated non-canonical junction from the *H. sapiens* telethonin gene, intron 1 (Genbank accession #: AJ010063). This junction is

supported by ESTs, which also supports a canonical splice junction. The annotated non-canonical junction and the putative canonical junction produce the same final spliced sequence.

ical splice sites are very likely the real ones. This suggested to explain these observations of shifted canonical dinucleotides by annotation errors involving inserting/deleting one nucleotide, which is actually absent/present in the real genomic sequence. This hypothesis was tested by comparing human gene sequences deposited to GenBank earlier with the sequence of the same region obtained in high throughput genomic sequencing projects (HTGs). Several examples of clear annotation and sequencing errors identified by the comparison are presented in Figure 3.5. 156 out of 171 human non-canonical and EST-supported splice site sequences had a clear match in the human HTG. They can be classified after corrections as: 79 GC-AG pairs (of which 1 was an error that corrected to GC-AG), 61 errors that were corrected to GT-AG canonical pairs, 6 AT-AC pairs (of which 2 were errors that corrected to AT-AC), 1 case that was produced from non-existent intron, 7 cases that were found in HTG that were deposited to GenBank, and finally there were only 2 cases left of supported non-canonical splice sites.

It was concluded that 99.24% of the splice site pairs should be GT-AG, 0.69% GC-AG, 0.05% AT-AC and finally only 0.02% could consist of other types of non-canonical splice sites (Table 3.3). Therefore, gene finding approaches using just standard GT-AG splice sites can potentially predict accurately 97% genes (assuming 4–5 exons per gene on average). If the

#### Sequences of homeodomain protein, HOXA9EC (AF010258)

	Donor	Acceptor
<b>Genbank:</b>	<b>CGATCCCAAT</b>  aa-tgtctcct	cccgcagaat  <b>AACCCAGCAG</b>
<b>High throughput:</b>	<b>CGATCCCA</b>  gtaagtgtctcct	cccgcag  <b>AT-AACCCAGCAG</b>

#### Sequences of poly(A) binding protein II, PABP2 (AF026029)

	Donor	Acceptor
<b>Genbank:</b>	<b>TCCAGGCAAT</b>  gctgagtaac	tttctgata  <b>GCTGGCCCGG</b>
<b>High throughput:</b>	<b>TCCAGGCAATG</b>  gtgagtaac	tttctgatag  <b>CTGGCCCGG</b>

**Fig. 3.5**  
Errors found by comparing GenBank and human high throughput sequences for several annotated non-canonical splice sites.

**Tab. 3.3**

Frequencies of canonical and non-canonical splice sites in human genes.

GT-AG	99.24%
GC-AG	0.69%
AT-AC	0.05%
Other non-canonical	0.02%

method takes into consideration GC-AG splice pairs, it can increase the recognition quality to 99%. The GC-AG splice group has several interesting features. The first is its relatively high frequency (0.56% of all EST-supported splicing pairs belong to this type and 0.69% is the final estimated frequency of this group). The frequency matrix of this site shows significantly higher degree of conservation in relation to the canonical donor matrix. It provides a possibility to implement this information in gene prediction programs without generating many false-positive predictions. The Fgenesh HMM-based gene prediction program [21] has been modified to predict genes with canonical as well as with GC-donor splice sites. This version of Fgenesh (<http://www.softberry.com/gf/gf.html>) can identify non-standard GC-exons with approximately the same level of false-positive predictions as the original program.

22 199 verified examples of canonical splice pairs are presented in the SpliceDB database, which is publicly available through the WWW (<http://genomic.sanger.ac.uk/SpliceDB.html>) [14]. It also includes 1615 annotated and 292 EST-supported and shift-verified non-canonical pairs.

### 3.3

#### Methods for the recognition of functional signals

This Section is intended to describe several approaches for the recognition of functional signals in genes and some features of these signals used in gene identification. A traditional way to find functional sites is based on using consensus sequences or weight matrices reflecting conservative nucleotides of a signal.

#### 3.3.1

##### Search for nonrandom similarity with consensus sequences

A statistical method for estimating the significance of the similarity between a consensus of a functional signal and a similar sequence fragment is briefly described below [22, 23].

Let us assume that we are searching for a site in a sequence of length  $N$  with a random arrangement of nucleotides A, T(U), G and C, where the frequencies of these nucleotides are  $P_A, P_{T(U)}, P_G$  and  $P_C$ , respectively. If we accept that  $P_1 = P_A, P_2 = P_G, P_3 = P_T, P_4 = P_C$ , then the frequencies of the nucleotides of the other classes  $P_j$  ( $j = 5, \dots, 15$ ) representing the non-empty subsets of the set {A, C, G, T} of nucleotides are determined as sums of frequencies of nucleotides of all the types of the  $j$ -th class.  $P_{15}$  represents the set of all 4 nucleotides, depicted by the IUPAC code N.

### 3.3.1.1 Single block site

Let us consider a functional site of length  $L$ . Let  $N_l$  be the number of positions in the consensus sequence whose bases belong to the  $l$ -th class and let  $N_1 + N_2 + \dots + N_{15} = L$ . Assume that the consensus has  $M$  conserved positions  $M_l$  ( $l = 1, \dots, 14$ ), where  $M$  is the number of conserved nucleotides of the  $l$ -th class ( $M_1 + M_2 + \dots + M_{14} = M$ ). Then  $k$  ( $k = 0, 1, \dots$ ) mismatches between the site and the segment of length  $L$  belonging to the sequence under consideration are allowed only at the  $L-M$  variable positions.  $R_l$  ( $l = 1, \dots, 14$ ) are the numbers of mismatches between the consensus and the DNA segment of the  $l$ -th class of nucleotides. The number of mismatches should meet the following conditions:  $0 \leq R_1 \leq \min(k, N_1 - M_1), 0 \leq R_2 < \min(k - R_1, N_{15} - M_{15}), \dots, 0 \leq R_{15} < \min(k - R_1 - R_2 - \dots - R_{13}, N_{14} - M_{14})$ .

Therefore, the probability  $P(L, k)$  of detecting the segment  $(L, k)$  of length  $L$  with mismatches in  $k$  variable positions, what we will call an  $(L, k)$  site, is:

$$P(L, k) = \sum_{R_1=0}^{\min(k, N_1 - M_1)} \dots \sum_{R_{15}=0}^{\min(k - R_1 - R_2 - \dots - R_{14}, N_{15} - M_{15})} C_{N_1 - M_1}^{R_1} P_1^{N_1 - P_1} (1 - P_1)^{R_1} \dots C_{N_{15} - M_{15}}^{R_{15}} P_{15}^{N_{15} - R_{15}} (1 - P_{15})^{R_{15}} \quad (1)$$

In this case the expected number  $\bar{T}(L, k)$  of structures  $(L, k)$  in a random sequence of length  $N$  is:

$$\bar{T}(L, k) = P(L, k) \times F_L$$

Here  $F_L$  is the number of possible site positions in the sequence:  $F_L = N - L + 1$ .

The probability of having precisely  $T$  structures  $(L, k)$  in the sequence may be estimated using the binomial distribution:

$$P(T) = C_{F_L}^T P^T(L, k) [1 - P(L, k)]^{F_L - T}. \quad (2)$$

The probability of detecting  $T$  structures with  $k$  or fewer mismatches is:

$$P(T) = \sum_{z=0}^k C_{F_L}^T P^T(L, z) [1 - P(L, z)]^{F_L - T}. \quad (3)$$

Now we can derive the upper boundary of the confidence interval  $T_0$  (with the significance level  $q$ ) for the expected number of structures in a random sequence:

$$\sum_{t=0}^{T_0-1} P(t) < q \quad \text{and} \quad \sum_{t=0}^{T_0} P(t) \geq q. \quad (4)$$

If the number of  $(L, k)$  sites detected in the sequence meets the condition  $T \geq T_0$ , they can be considered as potential functional signals with the significance level  $q$ .

### 3.3.1.2 Composite (two-blocks) site

Let us consider a composite consensus from two blocks of lengths  $L_1$  and  $L_2$  at a distance  $D$  ( $D_1 \leq D \leq D_2$ , i.e.,  $D_1$  and  $D_2$  are, respectively, the minimum and maximum allowed distances between the blocks). Let  $N_{1l}$  and  $N_{2l}$  be the number of nucleotides of the  $l$ -th class in the first and second blocks, respectively ( $l = 1, \dots, 15$ ). It is clear that  $N_{j1} + N_{j2} + \dots + N_{j15} = L_j$  ( $j = 1, 2$ ).

Let the first and second blocks have  $M_{1l}$  and  $M_{2l}$  conserved positions of the nucleotides of  $l$ -th class. Then the probability  $P(L_j, k_j)$  of finding in a random sequence the segment  $(L_j, k_j)$  of size  $L_j$  differing in  $k_j$  non-conserved positions from the  $j$ -th block of the site is calculated using Eq. (2) with the substitutions  $L, k, N_l$  and  $M_l$  by  $L_j, k_j, N_{jl}$  and  $M_{jl}$  ( $l = 1, \dots, 14$ ;  $j = 1, 2$ ), respectively. The probability of simultaneous and independent occurrence of  $(L_1, k_1)$  and  $(L_2, k_2)$  in the random sequence is:

$$P(L_1, k_1, ; L_2, k_2) = P(L_1, k_1) \times P(L_2, k_2). \quad (5)$$

The number of possible ways of arranging the segments  $(L_1, k_1)$  and  $(L_2, k_2)$  in the random sequence of length  $N$  is:

$$F(L_1, L_2, D_1, D_2) = (D_2 - D_1 + 1) \left[ N - L_1 - L_2 - \frac{D_1 + D_2}{2} + 1 \right]. \quad (6)$$

Then, the expected number of sequences  $(L_1, k_1, ; L_2, k_2, D_1, D_2)$  is:

$$(L_1, k_1, L_2, k_2, D_1, D_2) = F(L_1, L_2, D_1, D_2) \times P(L_1, k_1, ; L_2, k_2) \quad (7)$$

The probability  $P(T)$  of detecting  $T$  sequences  $(L_1, k_1, ; L_2, k_2, D_1, D_2)$  and the upper boundary of the confidence interval  $T_0$  can be computed analogously to Eq. (2-4).

This statistics has been implemented in the program NSITE (<http://www.softberry.com/gf.html>) [24], that identifies nonrandom similarity between fragments of a given sequence and consensuses of regulatory motifs from the TRANSFAC database [25].

### 3.3.2

#### Position-specific sensors

Weight matrices are often applied for functional signal description [26, 27, 28]. They are usually more accurate than the consensus technique and often incorporated as a signal scoring method in gene finding algorithms. A weight matrix can be considered as a simple model based on a set of position-specific probability distributions  $\{p_s^i\}$ , that provide probabilities of observing a particular type of nucleotide  $s$  in a particular position  $i$  of the functional signal sequence  $S$ . The probability of generating a sequence  $X(x_1, \dots, x_k)$  under this model is:

$$P(X/S) = \prod_{i=1}^k p_{x_i}^i, \quad (8)$$

where nucleotides of the signal are generated independently. A corresponding model can also be constructed for non-site ( $N$ ) sequences:  $\{\pi_s^i\}$ . A good discriminative score based on these models is the log-likelihood ratio:

$$\text{LLR}(X) = \log P(X/S)/P(X/N) \quad (9)$$

This score can be computed as an average sum of weights of observed nucleotides in a given sequence fragment using a corresponding weight matrix  $w(i, s) = \{\log(p_s^i/\pi_s^i)\}$ :

$$\text{LLR}(X) = \frac{1}{k} \sum_{i=1}^k w(i, x_i). \quad (10)$$

The more the  $\text{LLR}(X)$  (log likelihood ratio) exceeds 0, the better chances this sequence has to represent a real functional signal.

There are some other weight functions that are used to search for functional signals, for example, weights can be received by optimization procedures such as perceptrons or neural networks [29, 30]. Also, different position-specific probability distributions  $\{p_s^i\}$  can be considered. One typical generalization is to use position-specific probability distributions  $\{p_s^i\}$  of  $k$ -base oligonucleotides (instead of mononucleotides), another one is to exploit Markov chain models, where the probability to generate a particular nucleotide  $x_i$  of the signal sequence depends on  $k_0 - 1$  previous bases (i.e.

depends on the  $k_0 - 1$  base oligonucleotide ending at the position  $i - 1$ . Then the probability of generating the signal sequence  $X$  is:

$$P(X/S) = p_0 \prod_{i=k_0}^k p_{s_{i-1}, x_i}^{i-1, i} \quad (11)$$

where  $p_{s_{i-1}, x_i}^{i-1, i}$  is the conditional probability of generating nucleotide  $x_i$  in position  $i$  given oligonucleotide  $s_{i-1}$  ending at position  $i - 1$ ,  $p_0$  is the probability of generating oligonucleotide  $x_1 \dots x_{k_0-1}$ . For example, a simple weight matrix represents the independent mononucleotide model (or 0-order Markov chain), where  $k_0 = 1$ ,  $p_0 = 1$  and  $p_{x_{i-1}, x_i}^{i-1, i} = p_{x_i}^i$ . When we use dinucleotides (1<sup>st</sup> order Markov chain)  $k_0 = 2$ ,  $p_0 = p_{x_1}^1$ , and  $p_{x_{i-1}, x_i}^{i-1, i}$  is the conditional probability of generating nucleotide  $x_i$  in position  $i$  given nucleotide  $x_{i-1}$  at position  $i - 1$ . The conditional probability can be estimated from the ratio of the observed frequency of the  $k_0$ -base oligonucleotide ( $k_0 > 1$ ) ending at position  $i$  divided by the frequency of the  $k_0 - 1$ -base oligonucleotide ending at position  $i - 1$  in a set of aligned sequences of some functional signal.

$$p_{s_{i-1}, x_i}^{i-1, i} = f(s_{i-1}, x_i) / f(s_{i-1})$$

By the same method we can construct a model for non-site sequences for computing  $P(X/N)$ , where often the 0-order Markov chain with genomic base frequencies (or even equal frequencies (0.25)) is used. A log likelihood ratio (10) with Markov chains was used to select CpG island regions [31] as well in as a description of functional signals in gene finding programs such as Genscan [32], Fgenesh [21, 33] and GeneFinder [34].

Useful discriminative measure taking into account some *a priori* knowledge can be based on computing Bayesian probabilities as components of position specific distributions  $\{p_s^i\}$ :

$$P(S/o_s^i) = P(o_s^i/S)P(S) / (P(o_s^i/S)P(S) + P(o_s^i/N)P(N)), \quad (12)$$

where  $P(o_s^i/S)$  and  $P(o_s^i/N)$  can be estimated as position-specific frequencies of oligonucleotides  $o_s^i$  in the set of aligned sites and non-sites;  $P(S)$  and  $P(N)$  are the *a priori* probabilities of site and non-site sequences, respectively.  $s$  is the type of oligonucleotide starting (or ending) in the  $i$ -th position [35]. If one assumes independence of oligonucleotides in different positions the probability of a sequence  $X$  belonging to the signal is given by:

$$P(S/X) = \prod_{i=1}^k P(S/o_m^i).$$

Another empirical discriminator called ‘‘Preference’’ uses the average posi-



tional probability of belong to a signal:

$$\Pr(S/X) = 1/k \sum_{i=1}^k P(S/o_m^i). \quad (13)$$

This measure which is used in constructing discriminant functions for the Fgenes gene finding program [36] can be more stable than the above one on short sequences and has a simple interpretation: if  $\Pr(S/X) > 0.5$ , then our sequence is more likely belong to signal than to non-signal sequences.

### 3.3.3

#### Content-specific measures

Some functional signal sequences have distinctive general oligonucleotide composition. For example, many eukaryotic promoters are found in GC-rich chromosome fragments. We can characterize these regions applying scoring functions that are similar to the ones discussed above, but using probability distributions and their estimations by oligonucleotide frequencies, which are computed on the whole sequence of functional signal and are not position specific. For example, the Markov chain based probability (11) of generating the signal sequence  $X$  will be:

$$P(X/S) = p_0 \prod_{i=k_0}^k p_{s_{i-1}, x_i} \quad (14)$$

### 3.3.4

#### Frame-specific measures for recognition of protein coding regions

An important problem is to compute the probability of generating a protein coding sequence  $X$ . A coding sequence is a sequence of triplets (codons) read continuously from a fixed starting point. Three different reading frames with different codons are possible for any nucleotide sequence (or six if the complementary chain is also considered). It was noted that nucleotides are distributed quite unevenly among different codon positions, therefore the probability of observing a specific oligonucleotide in coding sequences depends on its position relative to the reading frame (three possible variants) as well as on adjacent nucleotides [37, 38, 39, 40]. Asymmetry in base composition between codon positions arises due to uneven usage of amino acids and synonymous codons in addition to the particular structure of the genetic code [41]. Comprehensive assessment of various protein coding measures was done by Fickett and Tung [42]. They estimate the quality of more than 20 measures and showed that the most powerful

measures, such as the ‘in phase hexanucleotide composition’, codon or amino acids usage give about 81% accuracy as coding region recognition functions on 54 base windows. Some powerful recognisers of coding gene regions based on neural network approaches have been constructed [43, 44]. Their sensitive coding region sensor, based on codon and dicodon statistics in six frames and a neural network output function, reaches 99% accuracy on predicting 180 nt ORFs (open reading frame), but it was analyzed only for genes expressed in liver and tested on 1000 examples. The prediction accuracy over 54 bp coding windows was about 85% [44]. Using a neural network to combine the information from 7 sensors that describe large amounts and diverse types of information, Uberbacher and Mural created the “coding recognition module”, which identifies 90% of the coding exons of length 100 nt or greater and about 50% of those less than 100 nt long [43]. Solovyev and Lawrence [45] used as the recognition function a modification of a Bayesian prediction scheme Eq. (12) that demonstrated good results on prokaryotic coding regions prediction [40]. The accuracy of classification using oligonucleotides of 8 bp length and function Eq. (13) is 90% for 54 nt and 95% for sequences 108 nt long [45], which is better than the accuracy for the combined six most powerful measures by LDA function in the Fickett and Tung [42] investigation.

In Markov chain approaches the frame-dependent probabilities  $p_{s_{i-1}, x_i}^f$  ( $f = \{1, 2, 3\}$ ) can be used to model coding regions:

$$P(X/S) = p_0 \prod_{i=k_0}^k p_{s_{i-1}, x_i}^f \quad (15)$$

where  $f$  is equal 1, 2 or 3 for oligonucleotides ending at codon position 1, 2 or 3, respectively.

### 3.3.5

#### Accuracy measures

For estimation of the performance of an algorithm or a recognition function we will use several quality measures [42, 46, 47]. True positives ( $T_P$ ) is the number of correctly predicted and false positives ( $F_P$ ) is the number of falsely predicted authentic sites. True negatives ( $T_N$ ) is the number of correctly predicted and false negative ( $F_N$ ) is the number of falsely predicted non-sites. Sensitivity ( $S_n$ ) measures the fraction of the true examples that are correctly predicted:  $S_n = T_P / T_P + F_N$ . Specificity ( $S_p$ ) measures the fraction of the predicted examples that are correct:  $S_p = T_P / T_P + F_P$ . Only consideration of both  $S_n$  and  $S_p$  values makes sense when we aim at providing accuracy information. If we want to concentrate on a single value for accuracy estimation the average of the correctly predicted number of sites and non-sites:  $AC = 0.5 (T_P + T_N)$  is a suitable measure. However, this

latter measure does not take into account possible difference in sizes of site and non-sites sets. A better single measure (correlation coefficient) takes into account the relation between correctly predictive positives and negatives as well as false positives and negatives [48]:

$$CC = (T_p T_n - F_p F_n) / \sqrt{(T_p + F_p)(T_n + F_n)(T_p + F_n)(T_n + F_p)}.$$

### 3.3.6

#### Application of linear discriminant analysis

Many eukaryotic functional signals have very short conservative regions and we should use their additional specific features (other than oligonucleotide frequencies) to increase the accuracy of signal identification. Different features of a functional signal may have different significance for the recognition and might not be independent. Classical linear discriminant analysis provides a method to combine such features in a discriminant function. In general, a discriminant function, when applied to a pattern, yields an output that is an estimate of the class membership of this pattern. The discriminative technique provides a minimization of the error rate of classification [49]. Let assume that each given sequence can be described by vector  $X$  of  $p$  characteristics  $(x_1, x_2, \dots, x_p)$ , which could be measured. The procedure of linear discriminant analysis is to find a linear combination of the measures (called the linear discriminant function or LDF), that provides maximum discrimination between sites sequences (class 1) and non-site examples (class 2).

The LDF:

$$Z = \sum_{i=1}^p a_i x_i$$

classifies ( $X$ ) into class 1 if  $Z > c$  and into class 2 if  $Z < c$  with a few misclassification as possible. The vector of coefficients  $(\vec{a}_1, \vec{a}_2, \dots, \vec{a}_p)$  and threshold constant  $c$  are derived from the training set by maximizing the ratio of the between-class variation  $z$  to within-class variation and are equal to [49]:

$$\vec{a} = s^{-1}(\vec{m}_1 - \vec{m}_2)$$

and

$$c = \vec{a}(\vec{m}_1 + \vec{m}_2)/2,$$

where  $\vec{m}_i$  are the sample mean vectors of characteristics for class 1 and class

2, respectively;  $s$  is the pooled covariance matrix of characteristics:

$$s = \frac{1}{n_1 + n_2 - 2} (s_1 + s_2)$$

$s_i$  is the covariation matrix, and  $n_i$  is the sample size of class  $i$ . Thus, based on these equations we can calculate the coefficients of the LDF and the threshold constant  $c$  using the values of characteristics of site and non-site sequences from the training sets and then test the accuracy of the LDF on the test set data. The significance of a given characteristic or set of characteristics can be estimated by the generalized distance between two classes (called the Mahalanobis distance or  $D^2$ ):

$$\vec{D}^2 = (\vec{m}_1 - \vec{m}_2) s^{-1} (\vec{m}_1 - \vec{m}_2)$$

which is computed based on values of the characteristics in the training sequences of classes 1 and 2. To find sequence features a lot of possible characteristics are generated, such as score of weight matrices, distances, oligonucleotide preferences within different sub-regions etc. Selection of the subset of significant characteristics  $q$  (among the tested  $p$ ) is performed by a step-wise discriminant procedure including only characteristics, which significantly increase the Mahalanobis distance. The procedure to test this significance uses the fact that the quantity:

$$F = \frac{n_1 + n_2 - p - 1}{p - q} \frac{n_1 n_2 (D_p^2 - D_q^2)}{(n_1 + n_2)(n_1 + n_2 - 2) + n_1 n_2 D_q^2}$$

has an  $F(p - q, n_1 + n_2 - p - 1)$  distribution [49] when testing hypothesis  $H_0: \Delta_p^2 = \Delta_q^2$ , where  $\Delta_m^2$  is the population Mahalanobis distance based on  $m$  variables. If the observations come from multivariate normal populations, the posterior probability that the example belongs to class 1 may be computed as [49]

$$\Pr(\text{class1}/\vec{x}) = \frac{1}{1 + \frac{n_2}{n_1} \exp\{-z + c\}}.$$

The posterior probability  $\Pr(\text{class2}/\vec{x}) = 1 - \Pr(\text{class1}/\vec{x})$ .

### 3.3.7

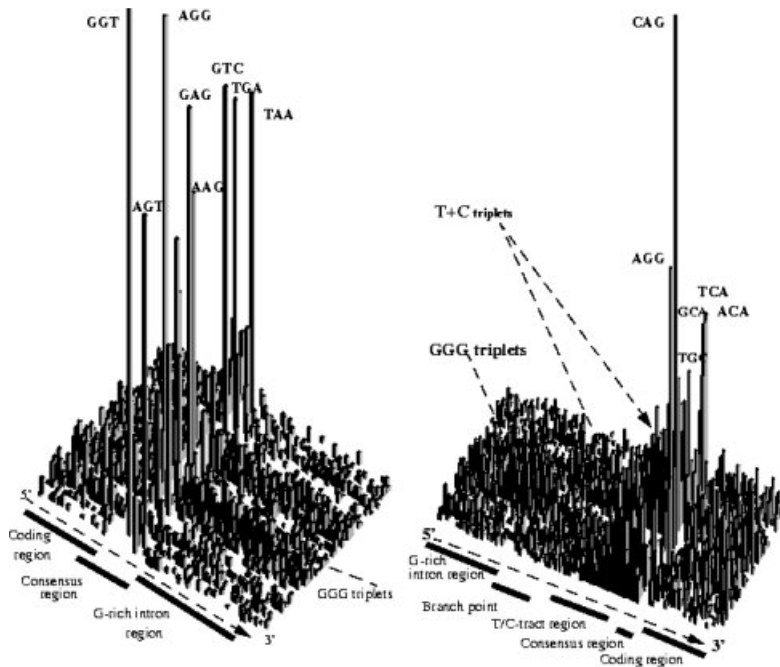
#### Prediction of donor and acceptor splice junctions

Recognition of RNA splice sites by the spliceosome is very precise [50–52] indicating the presence of specific signals for their function. Splice site patterns are mainly defined by nucleotides at the ends of introns, because

deletions of large intron parts often turn out not to effect their selection [7, 53]. A sequence of 8 nucleotides is highly conserved at the boundary between an exon and an intron (donor or 5'-splice site) and a sequence of 4 nucleotides, preceded by a pyrimidine-rich region, is also highly conserved between an exon and an intron (acceptor or 3'-splice site). The third less conserved sequence of about 5–8 nucleotides and containing an adenosine residue, lies within the intron, usually between 10 and 50 nucleotides upstream of the acceptor splice site (branch site). These sequences provide specific molecular signals by which the RNA splicing machinery can select the precise splice sites [18]. There are many efforts to analyze the sequences around these conserved regions [7, 18, 54]. It was shown that their consensus differ slightly between different classes of organisms [18, 55] and certain important information may be provided by the sequences outside the short conserved regions. Scoring schemes based on consensus sequences or weight matrices, which take into account information on open reading frames, free energy of base-pairing of RNA with snRNA and other peculiarities, yield an accuracy of about 80% for the prediction splice site positions [56, 57]. More accurate prediction is shown by neural network algorithms [58, 59]. The optimal network architecture (15 nucleotides window and 40 hidden units for donor sites and 41 nucleotides window and 20 hidden units) has an accuracy about of 94% (111/118) for predicting donor and 85% (100/118) for predicting acceptor splice sites in a test set of 118 examples [59]. Using joint coding/noncoding and splice site classification the prediction accuracy of about 95% with a low level of false positive sites was obtained. This work provides a good benchmark for testing new predictive algorithms, because the authors analysed large learning (331 examples) and testing (118 examples) sets. Because practically all donor sites contain the conserved dinucleotide GT and all acceptor sites contain AG, Mural et al. [60] used for splice sites selection the preferences of tabulated triplets in and about authentic splice junctions and also in and about pseudo-junctions which contain either a GT or an AG base pair. The authors obtained a good 91% accuracy for donor and 94% accuracy for acceptor splice sites prediction for primate genes. However, this data was based on small learning (about 150) and test (about 50) sets of splice junctions.

The overall view on the difference of triplet composition in splice and pseudosplice sequences is shown in Figure 3.6 [61]. This figure clearly demonstrates the various functional parts of splice sites. We see that only short regions around splice junctions have a great difference in triplet composition. Their sequences usually are used as the only determinants of donor or acceptor splice site positions. However, dissimilarity in many other regions can also be seen: for the donor site coding region, the G-rich intron region may be distinguished; for acceptor sites – intron G-rich region, branch point region, poly(*T/C*)-tract, and coding.

Splice site prediction methods using a linear function that combines triplet preferences around splice junction and preferences to be coding and intron of adjacent regions have been developed [45, 61]. 692 sequences with



**Fig. 3.6** Difference of the triplet composition within donor and GT-containing non-donor sequences (left); around acceptor and AG-containing non-acceptor sequences (right) in 692 human genes. Each column

presents the difference of specific triplet numbers between sites and pseudosites in a specific position. For comparison the numbers were calculated for equal quantities of sites and pseudosites.

2037 donor splice sites and 2054 acceptor splice sites having the GT and AG conserved dinucleotide in flanking intron positions were extracted from GenBank [62]. Also, 89 417 pseudodonor and 134 150 pseudoacceptor sites that contain either a GT or an AG base pair (and are not annotated as splice sites) were extracted from these sequences. The characteristics of sequences around splice sites and pseudosites were used for developing and testing a human splice site recognition function to distinguish them. The training set includes 2/3 of all sequences, and the test set contains the remaining ones. The data set for computing octanucleotide preferences in coding and intron regions included 4 074 593 bp of coding regions and 1 797 572 bp of intron sequences.

### 3.3.7.1 Donor splice site recognition

The characteristics used for classifying donor site are: the triplet preferences (Eq. 13) in the potential coding region ( $-30$  to  $-5$ ); conserved consensus region ( $-4$  to  $+6$ ) and G-rich region ( $+7$  to  $+50$ ); the number of significant

**Tab. 3.4**  
Significance of selected characteristics of donor splice sites.

	<b>Characteristics</b>	<b>1</b>	<b>2</b>	<b>3</b>	<b>4</b>	<b>5</b>	<b>6</b>	<b>7</b>
a	Individual $D^2$	9.3	2.6	2.5	0.01	1.5	0.01	0.4
b	Combined $D^2$	9.3	11.8	13.6	14.9	15.5	16.6	16.8

1, 2, 3 are the triplet preferences of consensus, intron G-rich and coding regions, respectively; 4 is the number of significant triplets in the consensus region, 5 and 6 are the octanucleotide preferences for being the coding 54 bp region on the left and for being the intron 54 bp region on the right of donor splice site junction; 7 is the number of G bases, GG-doublets and GGG-triplets in intron G-rich region.

triplets in the conserved consensus region; octanucleotide preferences (Eq. 15) for being coding in the ( $-60$  to  $-1$ ) region and being intron in the ( $+1$  to  $+54$ ) region; the number of G-bases, GG-doublets and GGG-triplets in the ( $+6$  to  $+50$ ) region. The values of these 6 characteristics of donor site were calculated for 1375 authentic donor site and for 60 532 pseudosite sequences from the learning set. The Mahalanobis distances showing the significance of each characteristic are given in Table 3.4a. We can see that the strongest characteristic for donor sites is the triplet composition in the consensus region ( $D^2 = 9.3$ ) and then the adjacent intron region ( $D^2 = 2.6$ ) and coding region ( $D^2 = 2.5$ ). Other significant characteristics are: the number of significant triplets in the conserved consensus region; the number of G-bases, GG-doublets and GGG-triplets; the quality of the coding and intron regions. The accuracy of the discriminant function based on these characteristics was tested on the recognition of 662 donor sites and 28 855 pseudosite sequences. The general accuracy of donor site prediction was 97%. This accuracy is better than in the neural network-based method, which has  $CC = 0.61$  at 95% accuracy [59], comparing to  $CC = 0.63$ .

### 3.3.7.2 Acceptor splice site recognition

The characteristics used for acceptor splice sites are: the triplet preferences (Eq. 13) in the branch point region ( $-48$  to  $-34$ ); poly( $T/C$ )-tract region ( $-33$  to  $-7$ ); conserved consensus region ( $-6$  to  $+5$ ); coding region ( $+6$  to  $+30$ ); and octanucleotide preferences (Eq. 15) of being coding in the ( $+1$  to  $+54$ ) region and in the ( $-1$  to  $-54$ ) region; and the number of  $T$  and  $C$  in poly( $T/C$ )-tract region. The values of 7 characteristics of acceptor sites were calculated for 1386 authentic acceptor site and 89 791 pseudosite sequences from the learning set. The Mahalanobis distances showing individual significance for each characteristic are given in Table 3.5a. We can see that strongest characteristics for acceptor sites are: the triplet composition in poly( $T/C$ )-tract region ( $D^2 = 5.1$ ); consensus region ( $D^2 = 2.7$ ); adjacent

Tab. 3.5

Significance of various characteristics of acceptor splice sites.

	<b>Characteristics</b>	<b>1</b>	<b>2</b>	<b>3</b>	<b>4</b>	<b>5</b>	<b>6</b>	<b>7</b>
a	Individual $D^2$	5.1	2.6	2.7	2.3	0.01	1.05	2.4
b	Combined $D^2$	5.1	8.1	10.0	11.3	12.5	12.8	13.6

1, 3, 4, 6 are the triplet preferences of poly(*T/C*)-tract, consensus, coding and branch point regions, respectively; 7 is the number of *T* and *C* in intron poly(*T/C*)-tract region, 2 and 5 are the octanucleotide preferences for being coding 54 bp region on the left and 54 bp region for being intron on the right of donor splice site junction.

coding region ( $D^2 = 2.3$ ); and branch point region ( $D^2 = 1.0$ ). Some significance is found using the number of *T* and *C* in the adjacent intron region ( $D^2 = 2.4$ ); and the quality of the coding region ( $D^2 = 2.6$ ).

The general accuracy of acceptor site prediction was 96% ( $C = 0.47$ ). Table 3.6 illustrates the performance for donor and acceptor sites and compares the results with the work of Mural et al. [60] who used triplet composition for splice site prediction, and the work of Brunak et al. [59] who used a complex neural network for site discrimination.

### 3.3.8

#### Recognition of promoter regions in human DNA

Eukaryotic polymerase II promoter sequences are the main regulatory elements of eukaryotic genes. Their recognition by computer algorithms will increase the quality of gene structure identification as well as provide the possibility to study gene regulation. The development of computer algorithms to recognize Pol II promoter sequences in genomic DNA is an extremely difficult problem in computational molecular biology. The 5'-flanking region of a promoter is very poorly described in general. It may contain dozens short motifs (5–10 bases) that serve as recognition sites for

Tab. 3.6

Splice site prediction accuracy.

<b>Method</b>	<b>Solovyev et al. [61]</b>	<b>Mural et al. [60]</b>	<b>Brunak et al. [59]</b>
Donor Sn	0.97	0.91	0.95
Donor CC	0.63	0.41	0.61
Acceptor Sn	0.96	0.93	0.96
Acceptor CC	0.47	0.36	0.40
Traning sites	2037	135	449
Test sites	662	50	118



proteins providing initiation of transcription as well as specific regulation of gene expression. These motifs differ among various groups of genes and even such a well known promoter element as the TATA-box is often absent in 5'-regions of many house-keeping genes. Each promoter has a unique selection and arrangement of these elements providing a unique program of gene expression. A comprehensive description of promoter recognition problems is considered in Chapter 4 of this book. Here we will consider some general features of PolII promoters that can be taken into account in gene prediction programs.

A review of the prediction accuracy of many general purpose promoter prediction program was presented by Fickett and Hatzigeorgiou [63]. This paper surveys oligonucleotide content based [64, 65], neural network [66, 67] and linear discriminant approaches [68] among others. Although the test set is relatively small (18 sequences) and has several problems that were noticed by [69], the results demonstrated that the programs can recognize no more than 50% of the promoters with a false positive rate of about 1 in 700–1000 bp. With the average size of a human gene being more than 5000 bases and many genes occupying hundreds of kilobases, we expect significantly more false positive predictions than real promoters. However, these programs can be used to find the promoter position (the start of transcription and the TATA-box) in a given 5'-region or to help select the correct 5'-exon in gene prediction approaches.

We will describe an improved version of the promoter recognition program TSSW (Transcription Start Site, W stands for using functional motifs from the Wingender [25] database) [68] to demonstrate sequence features that can be used to identify eukaryotic promoter regions. In this version it was suggested that TATA+ and TATA- promoters have very different sequence features and these groups were analyzed separately. Potential-TATA+ promoter sequences were selected by the value of score computed using the Bucher TATA box weight matrix [70] with the threshold closed to the minimal score value for the TATA+ promoters in the learning set. This choice of the threshold divides the learning set of known promoters into two approximately equal-size parts. Significant characteristics of both groups found by discriminant analysis are presented in Table 3.7. Values of Mahalanobis distances ( $D^2$ ) of individual characteristics reflect the power of the feature to separate the signal from non-signal sequences. This analysis demonstrated that TATA- promoters have much weaker general features than TATA+ promoters. Probably the TATA- promoter possesses more gene specific structure and will be extremely difficult to predict by any general-purpose methods.

The recognition quality of the program was tested on 200 promoters, which were not included in the learning set. We provide the accuracy values for different levels of correctly predicted promoters in Table 3.8. The data demonstrate a poor quality of TATA- promoter recognition on long sequences and show that their recognition function can provide relatively

Tab. 3.7

Selected characteristics of promoter sequences used by TSSW programs for TATA+ and TATA- promoters.

<i>Characteristics</i>	<i>D<sup>2</sup> for TATA+ promoters</i>	<i>D<sup>2</sup> for TATA- promoters</i>
• Hexaplets -200 to -45	2.6	1.4 (-100 to -1)
• TATA box score	3.4	0.9
• Triplets around TSS	4.1	0.7
• Hexaplets +1 to +40		0.9
• Sp1-motif content		0.9
• TATA fixed location	0.7	
• CpG content	1.4	0.7
• Similarity -200 to -100	0.3	0.7
• Motif Density(MD) -200 to +1	4.5	3.2
• Direct/Inverted MD -100 to +1	4.0	3.3 (-100 to -1)
Total Maxalonobis distance	11.2	4.3
Number promoters/non-promoters	203/4000	193/74 000

unambiguous predictions within regions less than 500 bp long. In contrast, 90% of the TATA+ promoters can be identified in the range about 2000 bp, which makes valid their incorporation into gene finding programs.

Recently several improvements of promoter prediction approaches have been published. Ohler et al. [69] used interpolated Markov chains in their approach and slightly improved the previous results. They identify 50% of the promoters in the Fickett and Hatzigeorgiou [63] promoter set, while having one false positive prediction every 849 bp. The old version of TSSW had an accuracy of 42% with a false positive rate of 1/789 bp. Another new program (Promoter 2.0) was designed by Knudsen [71] applying a combination of neural networks and genetic algorithms. Promoter 2.0 was tested on promoters in the complete Adenovirus genome, which is 35 937 bases long. The program predicted all 5 known promoter sites on the plus strand and 30 false positive promoters. The average distance of real and closest predicted promoter is about 115 bp. The TSSW program with the threshold

Tab. 3.8

Accuracy of promoter identification by TSSW program.

<i>Type of promoter</i>	<i>Number of test sites</i>	<i>Correctly predicted</i>	<i>1 false positive per bp</i>
TATA +	101	98%	1000
		90%	2200
		75%	3400
		52%	6100
TATA -	96	52%	500
		40%	1000

to predict all 5 promoters produces 35 false positives, but an average distance between the predicted TSS (transcription start site) and the real promoter is just 4 bp (2 predicted exactly, 1 with 1 bp shift, 1 with 5 bp shift and the weakest promoter was predicted with 15 bp shift).

### 3.3.9

#### Prediction of poly-A sites

The 3'-untranslated region (3'UTR) has several cytoplasmic functions affecting the localization, stability and translation of mRNAs [71]. Almost all eukaryotic mRNAs undergo 3'-end processing, which involves the endonucleotic cleavage followed by the polyadenylation of the upstream cleavage product [72, 73]. Recognition of essential sequences involves the formation of several large RNA-protein complexes [74]. RNA sequences directing binding of specific proteins are frequently poorly conserved and often recognized in a cooperative fashion [72]. Numerous experiments have revealed three types of RNA sequences defining a 3'-processing site [72, 75] (Figure 3.7). The most conserved is the hexamer signal AAUAAA (poly-A signal), situated 10–30 nucleotides upstream of the 3'-cleavage site. About 90% of the sequenced mRNAs have a perfect copy of this signal. Two other types, the upstream and the downstream elements are degenerate and have not been properly characterized. Downstream elements are frequently located within approximately 50 nucleotides 3' of the cleavage site [76]. These elements are often GU or U rich, although may have various base compositions and locations. On the basis of sequence comparisons McLachlan et al. [77] have suggested a consensus of the downstream element: YGUGUUY.

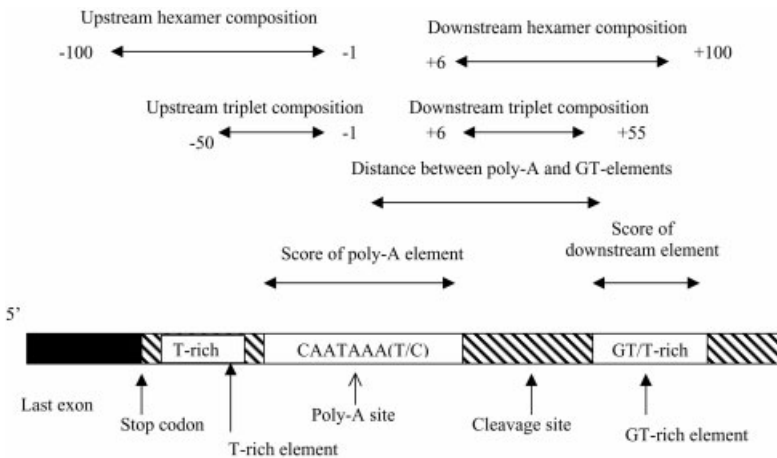


Fig. 3.7

Sequence characteristics selected to describe poly-A signal sequences.

The efficiency of polyadenylation in a number of genes can be also increased by sequences upstream of AAUAAA, which are generally U-rich [72]. All these RNA sequences serve as nucleation sites for the multi-component protein complex catalyzing the polyadenylation reaction. Yada et al. [78] have conducted a statistical analysis of human DNA sequences in the vicinity of the poly-A signal in order to distinguish them from AATAAA sequences occurring in human DNA (pseudo polyA-signals) that are not involved in polyadenylation. They found that a base C frequently appears on the upstream side of the AATAAA signal and a base T or C often appears on the downstream side, implying that CAATAAA(T/C) can be regarded as a consensus of the poly-A signal. Kondrakhin et al. [79] constructed a generalized consensus matrix using 63 sequences of cleavage/polyadenylation sites in vertebrate pre-mRNA. The elements of the matrix were absolute frequencies of triplets at each site position. Using this matrix they have provided a multiplicative measure for recognition of polyadenylation regions. However this method has a very high false positive rate.

Salamov and Solovyev developed a LDF recognition function for the poly-A signal. The datasets for 3'-processing sites and "pseudo" poly-A signals were extracted from GenBank (Version 82). 3'-processing sites were taken from the human DNA entries, containing a description of the poly-A signal in the feature table. Pseudo sites were taken out of human genes as the sequences comprising (-100, +200) around the patterns revealed by the poly-A weight matrix (see below), but not assigned to poly-A sites in the feature table. Sequences submitted to GenBank before 1994 were included to the training set and those after 1994 to the test set. As a result there were 248 positive and 5702 pseudosites in the training set and respectively 131 and 1466 in the test set.

### 3.3.9.1 A model for recognition of 3'-processing sites

As the hexamer AATAAA is the most conservative element of 3'-processing sites it was considered as the main block in our complex recognition function. Although the hexamer is highly conserved, variants of this signal are observed. For example, in the training set 43 of 248 poly-A sites have hexamer variants of AATAAA with one mismatch. To consider such variants the position weight matrix for recognizing this signal has been used. The other characteristics such as content statistics of hexanucleotides, positional triplets in the upstream and downstream regions were defined relative to the position of the conservative hexamer sequence (Figure 3.7).

The Mahalanobis distances for each characteristic calculated on the training set are given in Table 3.9. The most significant characteristic is the score of the AATAAA pattern (estimated by the position weight matrix), which indicates the importance of occurrences of an almost perfect poly-A signal (AATAAA). The second most valuable characteristic is the hexanucleotide preferences of the downstream (+6 to +100) region. Although the discrim-

**Tab. 3.9**  
Significance of selected characteristics of poly-A signal.

Characteristics	1	4	2	5	3	6	8	7
Individual $D^2$	7.61	3.46	0.01	2.27	0.44	1.61	0.16	0.17
Combined $D^2$	7.61	10.78	11.67	12.36	12.68	12.97	13.09	13.1

1 is the score of position weight matrix of poly-A signal, 2 is the score of the position weight matrix of the downstream GT/T-rich element, 3 is the distance between the poly-A signal and the predicted downstream GT/T-rich element, 4 is the hexanucleotide composition of the downstream (+6, +100) region, 5 is the hexanucleotide composition of the upstream (-100, -1) region, 6 is the positional triplet composition of the downstream (+6, +55) region, 7 is the positional triplet composition of the upstream (-50, -1) region, 8 is the positional triplet composition of the GT/T-rich downstream element.

inating ability of the GT-rich downstream element itself (characteristic 2) is very weak, combining it with the other characteristics significantly increases the total Mahalanobis distance.

According to the observation of Yada et al. [78], we take into account the hexamer AATAAA and its flanking nucleotides. Only sequences with a score that exceeds some threshold were considered as candidates of poly-A sites. Around the consensus we calculated the other characteristics (Figure 3.8) and the value of the linear discriminant function. The threshold was chosen as the minimal score observed for authentic poly-A signals in the training set. The poly-A signal weight matrix provides a higher score for typical poly-A signals (like CAATAAAT) and a smaller score for minor variants with other sequences. Such variants are often observed much more frequently among pseudosites and therefore the weight matrix is a good discriminant itself.

In general, characteristics from the downstream region of the AATAAA sequence are more informative than those from the upstream region. Such selected characteristics as positional triplet composition of the upstream and downstream regions as well as the distance between AATAAA pattern and GT-rich element might indicate the importance of definite location of sequence elements in the 3'-processing region. Altogether, these elements can create the bedding for cooperative interactions between proteins of multicomponent complex catalyzing the polyadenylation reaction. It is worth mentioning that we tested and selected only the most general characteristics of poly-A site regions.

The algorithm for 3'-processing site identification is realized in the POLYAH program (<http://genomic.sanger.ac.uk>). First, it searches for the AATAAA pattern using the position weight matrix and, after the pattern is found, it computes the value of the linear discriminant function defined by the characteristics around this signal. The poly-A site is predicted if the

value of this function is greater than some empirically selected threshold. An accuracy of the method has been estimated on a set of 131 positive and 1466 negative sites, which were not used in training. When 86% of polyA-regions are predicted correctly, the fraction of correctly predicted sites among all predicted (specificity) is 51% and the correlation coefficient is 0.62.

Kondrakhin et al. [79] reported the error rates of their method at different thresholds for poly-A signal selection. If the threshold is set to predict 8 of 9 real sites their function also predicts 968 additional false sites. We have tested the POLYAH program with the same sequence of Ad2 genome and for 8 correctly predicted sites it gives only 4 false sites. A certain improvement of Poly-A recognition was reached in using a pair of quadratic discriminant function in Polyadq program [80]. This program outperforms the POLYAH detection method and is the first that can detect significant numbers of ATTAAA-type signals.

### 3.4

#### Gene identification approaches

Most gene prediction systems combine information about functional signals and the regularities of coding and intron regions. Initially several algorithms predicting internal exons were constructed. The program SORFIND [81] was designed to predict internal exons based on codon usage and Berg & von Hippel [82] discrimination energy for intron-exon boundaries recognition. The accuracy of exact internal exons prediction (at both 5'- and 3'-splice junctions and in the correct reading frame) by the SORFIND program reaches 59% with a specificity of 20%. A dynamic programming approach (alternative to the rule-based approach) was applied by Snyder and Stormo (1993) to internal exon prediction in the GeneParser algorithm. It recognized 76% of the internal exons, but the structure of only 46% exons was exactly predicted when tested on entire GenBank entry sequences. HEXON (Human EXON) program [61] based on linear discriminant analysis was the most accurate in exact exon prediction that time.

Later a number of single gene prediction programs has been developed to assemble potential eukaryotic coding regions into translatable mRNA sequences selecting optimal combinations of compatible exons [47, 66, 83, 84]. Dynamic programming was suggested as a fast method to find an optimal combination of preselected exons [45, 85, 86], that is different from the approach suggested by Snyder and Stormo [46] in GeneParser algorithm to search for exon-intron boundary positions. The Genie algorithm uses a Generalized Hidden Markov Model. Genie is similar in design to GeneParser, but is based on a rigorous probabilistic framework [87]. The FGENEH (Find GENE in Human) algorithm includes 5'-, internal and 3'-exon identification linear discriminant functions and dynamic program-

ming approach [61, 88]. A comprehensive test of FGENEH and the other gene finding algorithms has been carried out by Burset and Guigo [89]. The FGENEH program was one of the best in the tested group having the exact exon prediction accuracy 10% higher than for the others and the best level of accuracy on the protein level. The same best performance of FGENEH was shown by the developers of Genie [87].

### 3.5

#### Discriminative and probabilistic approaches for multiple gene prediction

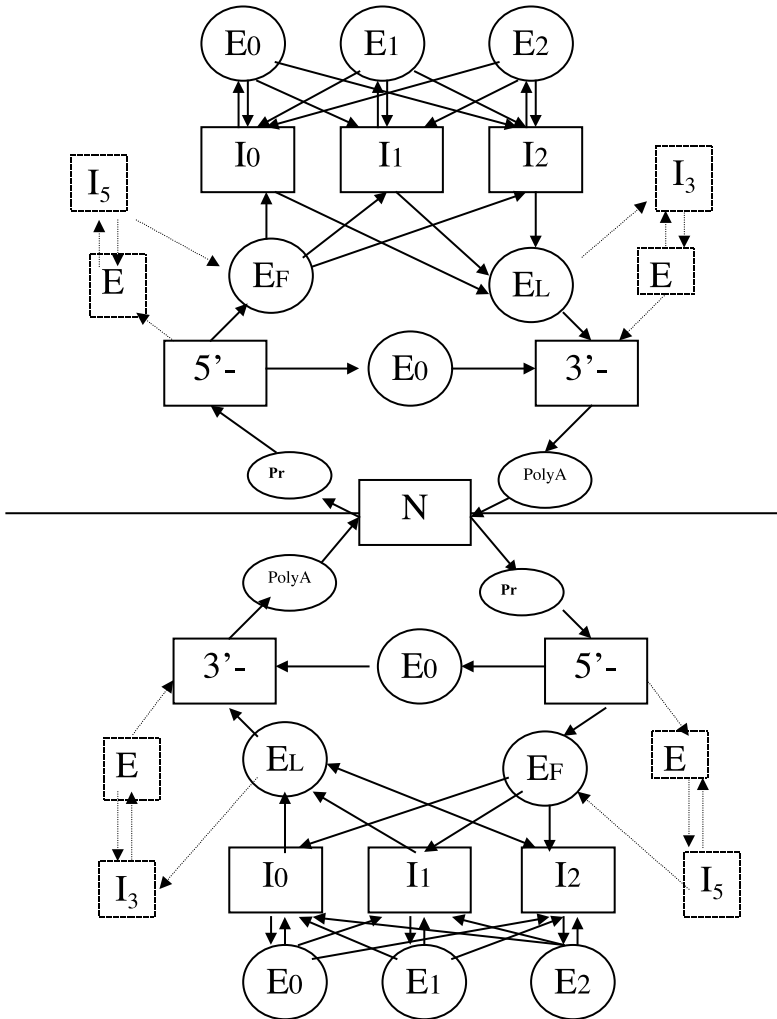
International genome sequencing projects generate hundreds of megabases each year and require gene-finding approaches able to identify many genes encoded in the produced sequences. The value of sequence information for the biomedical community will strongly depend on availability of candidate genes computationally predicted in these sequences. Moreover, the initiative that created a 'rough draft' of the human genome can allow other scientists to proceed more rapidly with discovering disease genes [90]. The best multiple gene prediction programs include Genscan [32] and Fgenesh [21] (HMM based, probabilistic approach) and Fgenes (discriminative approach) [36]. Initially we will describe a general scheme of HMM based gene prediction (first realized by the Haussler group [87, 91]) as the most general description of a gene model. The pattern-based approach can be considered as a particular case where transition probabilities are not taken into account.

#### 3.5.1

##### Multiple gene prediction using the HMM approach

A gene sequence can be considered as a succession of segments  $x_i$  representing exons, introns, 5' and 3'-untranslated regions and the like. These segments can be considered as different sequence states. There are 35 states describing the eukaryotic gene model considering direct and reverse chains as possible gene locations (Figure 3.8). In the current gene prediction approaches non-coding 5'- and 3'-exons (and introns) are not considered, because the absence of protein coding characteristics accounts for lower accuracy of their prediction. Also they do not code any protein sequences. The remaining 27 states include 6 exon states (first, last single and 3 types of internal exons in 3 possible reading frame) and 7 noncoding states (3 intron, noncoding 5'- and 3'-promoter and poly-A) in each chain plus noncoding intergenic region. The latter 27 states are connected with solid arrows in Figure 3.8, the eight unused states are connected with broken arrows.

The predicted gene structure can be considered as an ordered set of states/sequence pairs,  $\phi = \{(q_1, x_1), (q_2, x_2), \dots, (q_k, x_k)\}$ , called the parse, such that probability  $P(X, \phi)$  of generating  $X$  according to  $\phi$  is maximal over



**Fig. 3.8** Different sequence states and transitions in eukaryotic gene model.  $E_i$  and  $I_i$  are different exon and intron states, respectively ( $i = 0, 1, 2$  reflect 3 possible different ORF).  $E$  marks non-coding exons and  $I_5/I_3$  are 5'- and 3'-introns adjacent to non-coding exons.

all possible parses (or some score is optimal in some meaningful sense, i.e., best explains the observations [93]):

$$P(X, \varphi) = P(q_1) \left( \prod_{i=1}^{k-1} P(x_i | l(x_i), q_i) P(l(x_i) | q_i) (P(q_{i+1}, q_i)) \right) \times P(x_k | l(x_k), q_k) P(l(x_k) | q_k)$$

where  $P(q_1)$  is denote the initial state probabilities;  $P(x_i | l(x_i), q_i) P(l(x_i) | q_i)$



and  $P(q_{i+1}, q_i)$  are the independent joint probabilities of generating the subsequence  $x_i$  of length  $l$  in state  $q_i$  and transiting to state  $i + 1$ , respectively.

Successive states of this HMM model are generating output according to the Markov process with inclusion of explicit state duration density. A simple technique based on the dynamic programming method for finding the optimal parse (or the single best state sequence) is called the Viterbi algorithm [92]. The algorithm requires on the order of  $N^2 D^2 L$  calculations, where  $N$  is the number of states,  $D$  is the longest duration and  $L$  is the sequence length [93]. A helpful technique was introduced by Burge [28] to reduce the number of states and simplify computations by modeling non-coding state length with a geometric distribution. We consider shortly the algorithm of gene finding using this technique, which was initially implemented in the Genscan program [32] and used later in the Fgenesh program [21]. Since any valid parse will consist of only alternating series of Non-coding and Coding states:  $NCNCNC, \dots, NCN$ , we need only 11 variables, corresponding to the different types of  $N$  states. At each step corresponding to some sequence position we select the maximum joint probability to continue in the current state or to move to another non-coding state defined by a coding state (from a precomputed list of possible coding states), which is ending in analyzed sequence position.

Let us define the best score (highest joint probability)  $\gamma_i(j)$  of the optimal parse of the subsequence  $s_{1,j}$ , which ends in state  $q_i$  at position  $j$ . Let we have a set  $A_j$  of coding states  $\{-c_k\}$  of lengths  $\{d_k\}$ , starting at positions  $\{m_k\}$  and ending at position  $j$ , which have the previous states  $\{b_k\}$ . The length distribution of state  $c_k$  is denoted by  $f_{ck}(d)$ . The searching procedure can be stated as follows:

*Initialization:*

$$\gamma_i(1) = \pi_i P_i(S_1) p_i, \quad i = 1, \dots, 11.$$

*Recursion:*

$$\gamma_i(j+1) = \max \left\{ \gamma_i(j) p_i P_i(S_{j+1}), \right. \\ \left. \max_{c_k \in A_j} \{ \gamma_i(m_k - 1) (1 - p_{bk}) t_{bk, ck} f_{ck}(d_k) P(S_{m_k, j}) t_{ck, i} p_i P_i(S_{j+1}) \} \right\} \\ i = 1, \dots, 11, \quad j = 1, \dots, L - 1.$$

*Termination:*

$$\gamma_i(L+1) = \max \left\{ \gamma_i(L), \max_{c_k \in A_j} \{ \gamma_i(m_k - 1) (1 - p_{bk}) t_{bk, ck} f_{ck}(d_k) P(S_{m_k, j}) t_{ck, i} \} \right\} \\ i = 1, \dots, 11.$$

For each step we record the location and type of transition maximizing the functional to restore the optimal set of states (gene structure) by a back-tracking procedure. Most parameters of these equations can be calculated on the learning set of known gene structures. Instead of scores of coding states  $P(S_{mk,j})$  it is better to use log-likelihood ratios which do not produce scores below the limits of computer precision.

Genscan [32] was the first algorithm to predict multiple eukaryotic genes. Several similar HMM-based gene prediction programs were developed later, among them Veil [94], HMMgene [95], Fgenesh [21, 33], a variant of Genie [87] and GeneMark [96]. Fgenesh (Find GENES Hmm) is currently the most accurate program. It is different from Genscan in that in the model of gene structure signal terms (such as splice site or start site score) have some advantage over content terms (such as coding potentials). In log-likelihood terms this means that the splice sites and other exon functional signals have an additional score, depending on the environment of the sites. Also in computing the coding scores of potential exons, a priori probabilities of exons were taken into account according to Bayes' theorem. As a result, the coding scores of potential exons are generally lower than in Genscan. The donor recognition function is based on a second order position specific HMM (13) instead of 8 matrix, weight describing different groups of splice sites used in Genescan. Fgenesh works with separately trained parameters for each distinct model organism such as human, drosophila, nematode, yeast, arabidopsis monocot plants etc. Coding potentials were calculated separately for 4 isochores<sup>2)</sup> (human) and for 2 isochores (other species). The run time of Fgenesh is practically linear and the current version has no practical limit on the length of analyzed sequence. The prediction about 800 genes in 34 MB of Chromosome 22 sequence takes about 1.5 minutes on a Dec-alpha processor EV6 for the latest Fgenesh version.

### 3.5.2

#### **Pattern based multiple gene prediction approach**

FGENES (Find GENES) is the multiple gene prediction program based on dynamic programming. It uses discriminant classifiers to generate a set of exon candidates. Similar discriminant functions were developed initially in Fexh (Find Exon), Fgeneh (Find GENE) ( $h$  stands for version to analyze human genes) and described in detail [35, 68, 88].

The following major steps describe analysis of genomic sequences by the Fgenes algorithm:

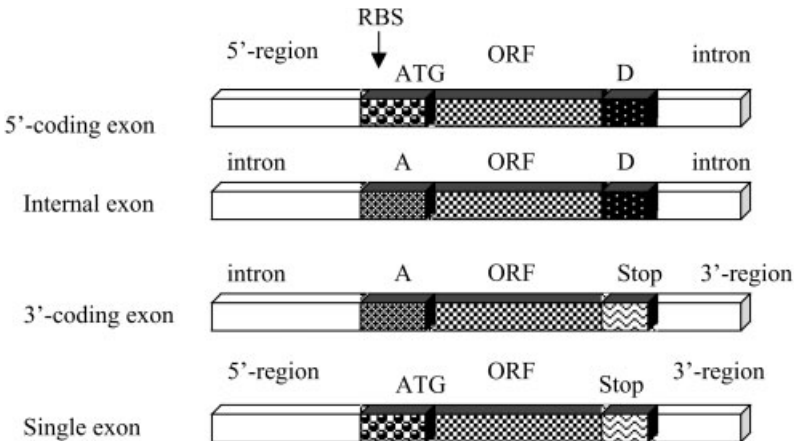
- 2) An isochore is a very long stretch (> 300 kb) of DNA that is homogeneous in base composition and compositionally correlated with the coding sequence that it embeds

- 1) Create a list of potential exons, selecting all ORFs: ATG ... GT, AG ... GT, AG ... Stop with exons scores higher than the specific thresholds depending on GC content (4 groups);
- 2) Find the set of compatible exons with maximal total score. Guigo [41] described an effective algorithm of finding such a set. Fgenes uses a simpler variant of the algorithm: Order all exon candidates according to their 3'-end positions. Going from the first to the last exon select for each exon the maximally scoring path (compatible exons combination) terminated by this exon using the dynamic programming approach. Include to optimal gene structure either this exon or exon with the same 3'-splicing pattern ending at the same position or earlier (which has higher the maximal score path).
- 3) Take into account promoter or poly-A scores (if predicted) in terminal exons scores.

The run time of the algorithm grows approximately linearly with the sequence length. Fgenes is based on the linear discriminant functions developed for identifying splice sites, exons, promoter and poly-A sites [61, 68]. We consider these functions in the following sections to see which sequence features are important in exon prediction.

### 3.5.2.1 Internal exon recognition

For internal intron prediction we consider all open reading frames in a given sequence that flanked the AG (on the left) and GT (on the right) base pairs as potential internal exons. The structure of such exons is presented in Figure 3.9. The components of the recognition function for internal exon



**Fig. 3.9**

Different functional regions of the first (a), internal (b), last (c) and single exons corresponding to components of recognition functions.

**Tab. 3.10**

Significance of selected internal exon characteristics. Characteristics 1 and 2 are the values of donor and acceptor site recognition functions; 3 is the octanucleotide preferences for being coding of potential exon region; 4 are the octanucleotide preferences for being intron 70 bp region on the left and 70 bp region on the right of potential exon region.

	<b>Characteristics</b>	<b>1</b>	<b>2</b>	<b>3</b>	<b>4</b>	<b>5</b>
a	Individual $D^2$	15.0	12.1	0.4	0.2	1.5
b	Combined $D^2$	15.0	25.3	25.8	25.8	25.9

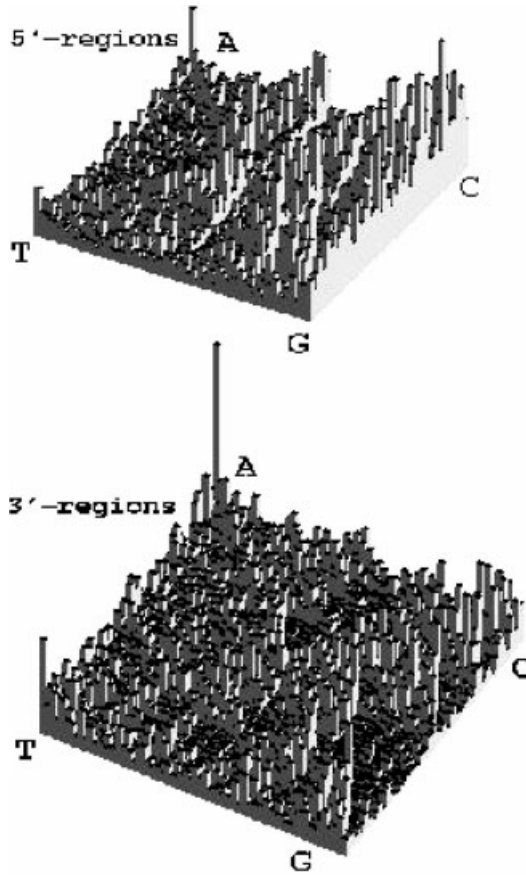
prediction consist of the octanucleotide preferences for an intron 70 bp to the left of the potential intron region; the value of the acceptor splice site recognition function, the octanucleotide preferences for the coding ORF, the value of the donor splice site recognition function and the octanucleotide preferences for intron 70 bp to the right of potential intron region. The values of 5 characteristics were calculated for 952 authentic exons and for 690 714 pseudo-exon training sequences from the set. The Mahalanobis distances showing significance of each characteristic are given in Table 3.10. We can see that the strongest characteristics for exons are the values of recognition functions of flanking donor and acceptor splice sites ( $D^2 = 15.04$  and  $D^2 = 12.06$ , respectively). The preference of ORF being a coding region has  $D^2 = 1.47$  and adjacent left intron region has  $D^2 = 0.41$  and right intron region has  $D^2 = 0.18$ .

The accuracy of the discriminant function based on these characteristics was calculated on the recognition of 451 exon and 246 693 pseudo-exon sequences from the test set. The general accuracy of exact internal exon prediction is 77% with a specificity of 79%. At the level of individual nucleotides, the sensitivity of exon prediction is 89% with a specificity of 89%; and the sensitivity of the intron prediction positions is 98% with a specificity of 98%. This accuracy is better than in the dynamic programming and neural network-based method [46], which has 75% accuracy of the exact internal exons prediction with a specificity of 67%. The method has 12% fewer false exon assignments with the better level of correct exon prediction.

### 3.5.2.2 Recognition of flanking exons

Figure 3.10 shows the three-dimensional histograms reflecting oligonucleotide composition of gene flanking regions created based on the graphical fractal representation of nucleotide sequences [97, 98, 99]. The clear differences in composition were exploited in the development of recognisers of these regions.

**Fig. 3.10**  
Fractal graphical representation of the number of different hexanucleotides in the 5'- (top) and 3'- (bottom) gene regions. Each colon represents the number of a particular hexanucleotide in the set of sequences.



#### 3.5.2.2.1 5'-terminal exon coding region recognition

For 5'-exon prediction, all open reading frames in a given sequence starting with an ATG codon and ending with a GT dinucleotide were considered as potential first exons. The structure of such exons is presented in Figure 3.9. The components of the 5'-exon recognition function included hexanucleotide preferences in the  $(-150, -101)$ ,  $(-100, -51)$  and  $(-50, 1)$  regions to the left of the potential coding region; the average value of the positional triplet preferences in the  $(-15, +10)$  region around the ATG codon; octanucleotide preferences of the coding ORF, the value of the donor splice site recognition function and the octanucleotide preferences to be intron 70 bp to the right of the potential coding region. The Mahalanobis distances showing the significance of each characteristic are given in Table 3.11. The accuracy of the discriminant function based on these characteristics was computed on the recognition of 312 the first exon and 246 693 pseudo-exon sequences. The gene sequences was scanned the 5'-exon with maximal weight were selected for each of them. The accuracy of the exact first coding

**Tab. 3.11**

Significance of selected 5'-exon characteristics. Characteristic 1 is the value of the donor site recognition function; 2 is the average value of positional triplet preferences in the (-15, 10) region around the ATG codon; 4 is the octanucleotide preferences for being intron in the 70 bp region to the right of the potential exon; 3, 5 and 7 are the hexanucleotide preferences in the (-150, -101), (-100, -51) and (-50, -1) regions to the left of the potential exon, respectively; 6 is the octanucleotide preferences for being coding in the exon region.

	<b>Characteristics</b>	<b>1</b>	<b>2</b>	<b>3</b>	<b>4</b>	<b>5</b>	<b>6</b>	<b>7</b>
a	Individual $D^2$	5.1	2.6	2.7	2.3	0.01	1.05	2.4
b	Combined $D^2$	5.1	8.1	10.0	11.3	12.5	12.8	13.6

exon prediction is 59%. It must be noted that the competition with the internal exons was not considered in this test.

#### 3.5.2.2.2 3'-exon coding region recognition

All ORF regions that were flanked by GT (on the left) and finished with a stop codon were considered as potential last exons. The structure of such exons is presented in Figure 3.9. The characteristics of their discriminant function and their Mahalanobis distances are presented in Table 3.12. The accuracy of the discriminant function was tested on the recognition of the last 322 exon and 247 644 pseudo-exon sequences. The gene sequences were scanned and the 3'-exon with the maximal weight was selected for each of them. The function can identify exactly 60% of the annotated last exons.

The recognition function of single exons combines the corresponding characteristics of 5'- and 3'-exons.

#### 3.5.2.2.3 Combined prediction of the first, internal and last exons in human genes

The program FEX (FindEXon) predicts coding regions in a given sequence. The program initially predicts the internal exons based on the internal exon

**Tab. 3.12**

Significance of selected 3'-exon characteristics. Characteristic 1 is the value of the acceptor site recognition function; 2 is the octanucleotide preferences for being coding of ORF region; 3, 5 and 7 are the hexanucleotide preferences (+100, 150), (+50, +100) and (+1, +50) regions to the left of the coding region, respectively; 4 is the average value of the positional triplet preferences in the (-10, +30) region around the stop codon; 6 is the octanucleotide preferences for being intron in the 70 bp region to the left of the exon sequence.

	<b>Characteristics</b>	<b>1</b>	<b>2</b>	<b>3</b>	<b>4</b>	<b>5</b>	<b>6</b>	<b>7</b>
a	Individual $D^2$	10.0	3.2	0.8	2.2	1.2	0.2	1.6
b	Combined $D^2$	10.0	11.4	12.0	13.8	14.3	14.5	14.6

discriminant function. Then we search for the 5'-coding region starting from the beginning of the sequence until the end of the first predicted internal exon. In this region the 5'-coding exon with the maximal weight of the discriminant function for the first exon is selected. After that we search for the 3'-coding region starting from the beginning of the last predicted internal exon until the end of the sequence. In this region the 3'-coding exon with the maximal weight of the last exon discriminant function is selected. This program can be useful in producing a list of potential exons, which are not effected by their assembling, because exon assembling in some cases might significantly corrupt actual gene structure due to underprediction of some small exons. However, exon assembling reduces significantly the number of false positive exons and this strategy is used in the most accurate gene prediction approaches.

### 3.5.3

#### Accuracy of gene identification programs

Most gene recognition programs were tested on a specially selected set of 570 one-gene sequences [89] of mammalian genes (Table 3.13). We can see that, on the average, the best programs predict accurately 93% of the exon nucleotides ( $S_n = 0.93$ ) with just 7% false positive predictions. Because the most difficult task is to predict small exons and to exactly identify the 5'-

**Tab. 3.13**

Accuracy of gene prediction programs for single gene sequences from the [89] data set.  $S_n$  (sensitivity) is the number of exactly predicted exons/number of true exons (or nucleotides);  $S_p$  (specificity) is the number of exactly predicted exons/number of all predicted exons. Accuracy data for the programs developed before 1996 were estimated by Burset and Guigo [89]. The other data were received by the authors of the respective programs.

<i>Algorithm</i>	<i>S<sub>n</sub> (exons)</i>	<i>S<sub>p</sub> (exons)</i>	<i>S<sub>n</sub> nucleotides</i>	<i>S<sub>p</sub> nucleotides</i>	<i>Reference</i>
Fgenesh	0.84	0.86	0.94	0.95	[33]
Fgenes	0.83	0.82	0.93	0.93	[36]
GenScan	0.78	0.81	0.93	0.93	[32]
Fgeneh	0.61	0.64	0.77	0.88	[88]
Morgan	0.58	0.51	0.83	0.79	[108]
Veil	0.53	0.49	0.83	0.79	[94]
Genie	0.55	0.48	0.76	0.77	[87]
GenLang	0.51	0.52	0.72	0.79	[47]
Sorfind	0.42	0.47	0.71	0.85	[81]
GeneID	0.44	0.46	0.63	0.81	[66]
Grail2	0.36	0.43	0.72	0.87	[86]
GeneParser2	0.35	0.40	0.66	0.79	[46]
Xpound	0.15	0.18	0.61	0.87	[109]

and 3'-ends of an exon, the accuracy at the exon level is usually lower than at the nucleotide level.

The Table demonstrates that the modern multiple gene prediction programs as Fgenesh, Fgenes and Genescan significantly outperform the older approaches (some of them may have been improved since the initial publication). These data show definite progress in accuracy of gene recognition. The exon identification rate is actually even higher than the presented data since overlapping exons were not counted in exact exon predictions. However there is a lot of room for future improvements. Thus, the accuracy at the level of exact gene prediction is only 59% for Fgenesh, 56% for Fgenes and 45% for Genescan programs even on this relatively simple test.

The real challenge for *ab initio* gene identification is presented by long genomic sequences containing many genes in both DNA strands. There is not much information about real genes in such sequences. One example, which was experimentally studied in the Sanger Centre (UK), is the human BRCA2 region (1.4 MB) that contains 8 genes and 169 experimentally verified exons. This region is one of the most difficult cases in genome annotation, because it has genes with many exons and almost no genes encode amino acid sequences having similarity with known proteins. Moreover the region contains 4 pseudogenes and at least 2 genes have alternative splicing variants. The results of gene prediction initially provided by Hubbard and Bruskiewich (The Sanger Centre Genome Annotation Group) are shown in Table 3.14.

Fgenesh predicts 20% fewer false positive exons in this region than Gen-

**Tab. 3.14**

Accuracy of gene prediction programs for the BRCA2 1.4 MB human genomic sequence. When repeats have been defined by the RepeatMasker [100] program in the analyzed sequence these regions were masked for the prediction and excluded from the potential exon locations during prediction. The region consisted of 20 sequences with 8 verified genes, 4 pseudogenes and 169 exons. Later one sequence was constructed and 3 additional exons were identified. The results of the predictions on this sequence is marked in bold. CC is the correlation coefficient reflecting the accuracy of prediction at the nucleotide level. Snb, Spb – sensitivity and specificity at the base level (in %), Pe – number of predicted exons, Ce – number of correctly predicted exons, Sne, Spe – sensitivity and specificity at the exon level, Snep – exon sensitivity, including partially correct predicted exons (in %). Ov results by including as correctly predicted not only predicted exons which exactly coincide with the real exons, but count also predicted exons that overlapped with the real exons.

	CC	Snb	Spb	Pe	Ce/Ov	Sne	Sn_ov	Spe/Spe_ov
Genescan	0.68	90	53	271/271	109/131	65	80/76	40/49
Fgenesh	0.80	89	73	188/195	115/131	69	80/76	61/67
Fgenes	0.69	79	62	298/281	110/136	66	86/78	37/48
Genescan masked	0.76	90	66	217	109	65	80	50
Fgenesh masked	0.84	89	82	172/168	114/131	68	79/73	66/76
Fgenes masked	0.73	80	68	257/228	107/133	64	85/75	42/58



Tab. 3.15

Performance of multiple gene-finding for 38 genomic sequences. Me is missing exons and W is wrong exons. M\_r lines provide predictions on sequences with masked repeats. Sn\_o is exon prediction accuracy including overlapping exons.

Program	Sequences/ Genes	Accuracy per nucleotide			Accuracy per exon		Me	We	Genes/ Entries
		Sn	Sp	CC	Sn/Sn_o	Sp			
Fgenesh	38/77 M_r	0.94	0.87	0.90	0.85/0.93	0.80	0.08	0.14	0.36/0.11
		0.94	0.78	0.85	0.84/0.92	0.75	0.08	0.21	0.34/0.08
Genscan	38/77 M_r	0.93	0.82	0.87	0.80/0.90	0.74	0.10	0.18	0.29/0.03
		0.92	0.70	0.79	0.79/0.90	0.66	0.11	0.30	0.29/0.03
Fgenes	38/77 M_r	0.91	0.80	0.84	0.84/0.92	0.72	0.08	0.21	0.36/0.18
		0.92	0.76	0.83	0.84/0.93	0.68	0.07	0.30	0.39/0.21

scan with practically the same number of correctly predicted exons. Also, we can observe that even for such a difficult region, about 80% of the exons were identified by *ab initio* approaches. It is interesting to note that, when we take the subset of exons predicted by both programs (Fgenes+Fgenesh), then the predicted exons are correct in 93% cases. It is very important to know such reliable exons to start verification of a gene structure by experimental techniques.

For another test we selected a set of 19 long genomic sequences with between 26 000 and -240 000 bp and 19 multigene sequences with between 2 and 6 genes from GenBank. Table 3.15 demonstrates the results of gene prediction on these data. This results show that the accuracy is still pretty good on the nucleotide and exon level, but exact gene prediction is lower than for the test with short single gene sequences. Sensitivity for exact internal exon prediction is 85–90%, but 5'-, 3'- and single exons have a prediction sensitivity of ~50–75%, which can partially explain the relatively low level of exact gene prediction. As a result we observe splitting up of some actual genes and/or joining some other multiple genes into a single one. A significant limitation of the current gene-finding programs is that they could not detect nested genes, i.e., genes located inside introns of another genes. While this is probably a rare event for the human genome, for organisms like *Drosophila* it presents a real problem. For example, annotators identified 17 examples of nested genes in the Adh region. [110]. Masking repeats seems important to increase (~10%) the specificity of prediction.

### 3.5.4

#### Using protein or EST similarity information to improve gene prediction

Manual annotation experience shows that it is often advantageous to take into account all available information to improve gene identification. Auto-

matic gene prediction approaches also take into account similarity information with known proteins or ESTs [101, 102]. Fgenesh+ [21] is a version of Fgenesh, which uses additional information from the available protein homolog. When exons initially predicted by Fgenesh show high similarity to some protein from the database, it is often advantageous to use this information to improve the prediction accuracy. Fgenesh+ requires an additional file with a protein homolog, and aligns all predicted potential exons with that protein using the SCAN 2 iterative local alignment algorithm [103]. To decrease the computation time, all overlapping exons in the same reading frame are combined into one sequence and aligned only once.

The main additions to the algorithm, relative to Fgenesh, include:

- 1) augmentation of the scores of exons with detected similarity by an additional term proportional to the alignment score.
- 2) additional penalty included for the adjacent exons in dynamic programming (Viterbi algorithm), if their corresponding aligned protein segments are not close in the corresponding similar protein.

Fgenesh+ was tested on the selected set of 61 GenBank human sequences, for which the Fgenesh predictions were not accurate (correlation coefficient  $0.0 \leq CC < 0.90$ ) and which have protein homologs from other organisms. The percent sequence identity between encoded proteins and homologs was varied from 99% to 40%. The prediction accuracy on this set is presented in Table 3.16. The results show that if the alignment covers the whole length of both proteins, then Fgenesh+ usually increases the accuracy relative to Fgenesh. This phenomenon does not depend significantly on the level of sequence identity, as long as level exceed 40%. This property makes knowledge of proteins from even distant organisms valuable for improving the accuracy of gene identification. A similar approach of exploiting known EST/cDNA information was realized in Fgenesh\_c program [21].

**Tab. 3.16**

Comparison of the accuracy of Fgenesh and Fgenesh+ on the set of human genes that are poorly predicted by *ab initio* methods, with known protein homologs from other organisms. The set contains 61 genes and 370 exons. CG – percent of correctly predicted genes; Sne, Spe are the sensitivity and specificity at the exon level (in %); Snb, Spb are the sensitivity and specificity at the base level (in %), respectively; CC – correlation coefficient.

	CG	Sne	Spe	Snb	Spb	CC
Fgenesh	0	63	68	86	83	0.74
Fgenesh+	46	82	85	96	98	0.95

## 3.6

## Annotation of sequences from genome sequencing projects

The first task in analyzing these sequences is finding the genes. Knowledge of genes opens a new way of performing biological studies called ‘functional genomics’. The other problem is to find out what all these new genes do, how they interact and are regulated [104]. Comparisons between genes of different genomes can provide additional insights into the details of the structure and function of genes.

We cannot predict exactly all gene components due to the limitation of our knowledge of the complex biological processes and signals regulating gene expression. In this respect, computer analysis of the genetically well studied Adh region of *D. melanogaster* by several gene-finding approaches [105] gives us a unique opportunity to define the reliability and limitations of our predictions and provide the strategy of right interpretation of prediction results in analysis of new genomic sequences. The predictions were evaluated by using two standards, one based on previously unreleased high quality full-length cDNA sequences and a second based on the set of annotations generated as part of an in-depth study of the region by a group of *Drosophila* experts. The performance of several accurate annotations is presented in Table 3.17. The CGG1 annotation comprised the non-

Tab. 3.17

Performance of several programs on the Adh region of *Drosophila*. The std3 contains 222 genes and 909 exons, the std1 set contains 43 genes and 123 exons. The annotated exons are taken from the set presented by organizers of the Genome Annotation Assessment Project (GASP1) at the time of the initial data analysis. Pe – number of predicted exons, Ce – number of correctly predicted exons. Pg – number of predicted genes, Pe – number of correctly predicted genes. Sn – sensitivity (in %), Sp – specificity (in %). At the exon level the second figure shows sensitivity taking into account exactly predicted and overlapped exons.

		CGG1	Fgenesh	Fgenesh pruned	Genie	Genie EST
Base Level	Sn std1	89	98	98	96	97
	Sn std3	87	92	88	79	79
	Sp std3	77	71	86	92	91
Exon Level	Pe	1115	1671	979	786	849
	Ce std1	80	100	100	86	95
	Sn std1	65/89	81/97	81/97	70	77
	Ce std3	544	601	565	447	470
	Sn std3	60/82	66/89	62/82	49	52
	Sp std3	49	36	58	57	52
	Gene Level	Pg	288	530	262	241
	Cg std1	22	31	31	24	28
	Sn std1	51	72	72	56	65
	Cg std3	102	108	106	86	92
	Sn std3	46	49	48	39	41
	Sp std3	36	20	39	37	38

ambiguous gene set. The genes were included by the following rules (descending in priority):

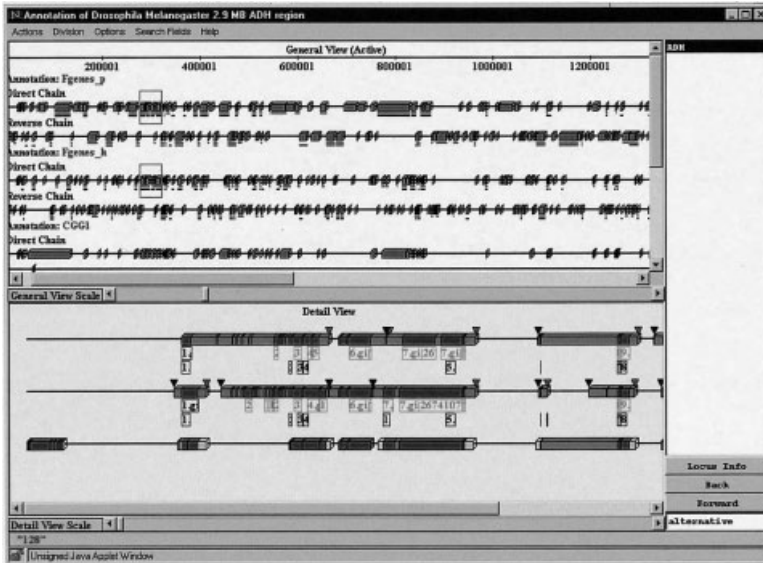
- a) all genes that were predicted by Fgenesh+;
- b) genes predicted identically by both Fgenes (human parameters) and Fgenesh (drosophila parameters) programs;
- c) in the regions of overlapped (but not exactly coincide) predictions, only one predicted gene was included with the priority given to the genes producing the longer protein.

Fgenesh+ was used to improve the accuracy of prediction for 49 genes. 37 of them were predicted using *D. melanogaster's* own proteins, already deposited in protein databases. Analysis of these predictions demonstrates that even for such cases prediction of accurate gene structure may not be trivial, although in most cases Fgenesh+ improved the prediction accuracy relative to *ab initio* methods.

The annotation CGG1 predicts about 87% of the real coding nucleotides and has just about 23% false positives (some of them might happen to be coding due to absence of experimental data in many regions). 89% of the exons are predicted exactly or with overlapping. These data show that *ab initio* predictions can provide information about practically all of protein coding genes (just 13% of coding region was not predicted) and can serve as a reasonable base for further experimental analysis.

It is interesting to note that the usage of two programs provided stable prediction accuracy on both (std1 and std3) sets. The Genie program, for example, demonstrated 20% decrease of sensitivity (Table 3.17). Because there is no version of Fgenes with all parameters computed on *Drosophila* genes, the optimal variant of automatic annotation was performed by using only the Fgenesh program.

It was found that the Fgenesh pruned predictions provided the best accuracy of annotation of the 2.9 MB Adh sequence. In this simple variant from the set of predicted genes all low-scoring genes (with average gene score less than 15) were discarded. This yields 98% of the coding nucleotide prediction on the set of 43 verified genes and 88% accurate coding nucleotide predictions with only 14% of false positives on a 222 gene set (where the significant parts of the genes were derived from Genscan predictions). The results demonstrate that most of the annotated genes in std3 are at least partially covered by predictions. For example, just 5 genes from the std3 set do not overlap with Fgenesh predictions (two of them are also included in the std1 set). From these 5 genes, 4 are located inside introns of other genes and 4 are single-exon genes. So one of the limitations of current gene-finding programs is that they cannot detect nested genes, i.e. genes located inside introns of other genes and this is one of the future directions for improvement of gene-finding software. Another drawback of the current gene-finding programs is that predictions of terminal exons are generally



**Fig. 3.11** InfoGene viewer [15] visualization of genes and fgenes predictions in Adh region. In the lower panel fragments of annotations are presented that are marked in the upper panel. The last level presents

the std3 manual annotation. Coding exons are marked by red color and introns by grey color. Green triangles show the start of transcription and red ones mark the poly-A signal.

much worse than the internal ones. This results in splitting up of some actual genes and/or joining some other multiple genes into a single one. Several examples of such a situation can be clear seen with the InfoGene Java viewer [15] (Figure 3.11) developed to present information about gene structures described in Genbank (collecting information about a gene from many entries) or annotated using gene prediction programs. On the std1 set Fgenesh predicts all internal exons correctly (100%), while only 72% of initial exons and 77% of terminal exons were predicted correctly. Thus better predicting the terminal exons and the related problem of better recognizing the beginnings (transcription start sites) and endings (polyA sites) of genes are the other areas of possible future improvements of gene-finding programs.

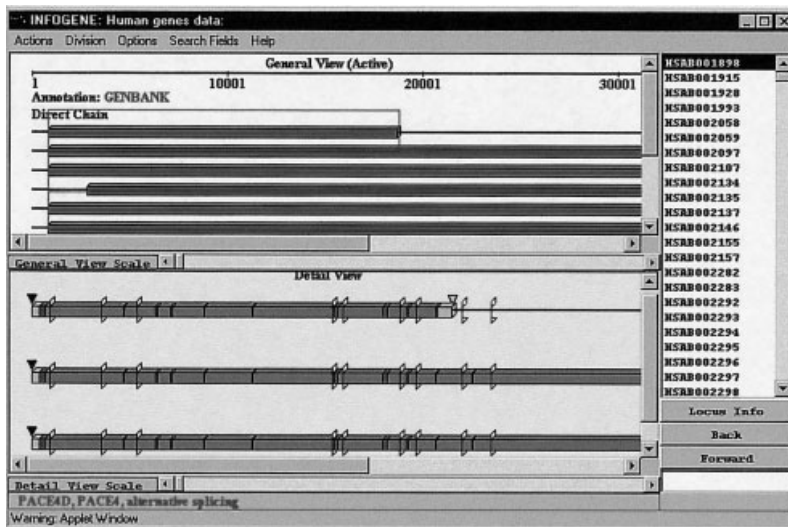
### 3.7 InfoGene: database of known and predicted genes

Genomic information is growing faster every day, but unfortunately the proportion of experimentally confirmed data is decreasing and the com-

plexity of extraction of useful information is increasing. The InfoGene database [4, 5] provides one of the most complete gene-centered genomic databases practically with all coherent information that can be obtained from the GenBank feature tables. We can obtain the necessary information without looking into many GenBank entries, where the information about a particular gene might be stored.

The InfoGene database includes known and predicted gene structures with a description of their basic functional signals and gene components. All major organisms are presented in separate divisions. The information about a gene structure might be collected from dozens GenBank entries. This information can be applied to create different sets of functional gene component for extraction their significant characteristics used in gene prediction systems. InfoGene is realized under a JAVA interactive environment system [15] that provides visual analysis of known information about complex gene structure (Figure 3.12) and searches different gene components and signals. The database is available through WWW server of Computational Genomics Group at <http://genomic.sanger.ac.uk/infodb.shtml>.

The value of sequence information for the biomedical community will strongly depend on the availability of candidate genes that are computationally predicted in these sequences. Currently information about predicted genes is absent in sequence databases if the gene has no similarity on the protein level with a known protein. Using gene prediction the scientific community can start experimental work with most human genes, because



**Fig. 3.12**  
InfoGene [15] presentation of human PACE4 (AB001898) gene. This gene has several alternative forms and described in

17 entries of GenBank. Continuous sequences regions corresponding different GenBank entries are separated by the vertical bars.

gene-finding programs usually predict accurately at least the major part of the exons in a gene sequence. InfoGene includes all predicted genes for Human and *Drosophila* genome drafts and several chromosomes of the *Arabidopsis* genome. The database is available currently through WWW server at <http://www.softberry.com/infodb.html>. Recently, the similar project Ensembl was started as a collaboration between the Sanger Center and European Bioinformatics Institute (<http://www.ensembl.org/>).

### 3.7.1

#### Annotation of human genome draft

The nucleotide sequence of nearly 90% of the human genome (3 GB) has been determined by international sequencing effort. Assembly of the current draft of the human genome was performed by the Human Genome Project Team at UC Santa Cruz. On this sequence (with masked repeats) the FGENESH program was used to predict exons and assemble predicted genes. Annotation of similarity of each exon with the PfamA protein domain database [106] was produced by the Blast program [107]. Totally 49 171 genes and 282 378 coding exons were predicted. On average one gene was found per about 68 623 bp and one exon per 11 949 bp. A complete summary of this analysis including the gene and exon numbers in different human chromosomes is presented at [http://www.softberry.com/inf/humd\\_an.html](http://www.softberry.com/inf/humd_an.html) and can be viewed in the InfoGene database. Sequences of predicted exons and gene annotation data can be copied from this site also. 1154 types of PfamA different domains were found in the predicted proteins. The top part of the domain list is presented in Table 3.18.

## 3.8

### Functional analysis and verification of predicted genes

Large scale functional analysis of predicted and known genes might be done using expression micro-array technology (see Chapter 5 of Volume 2 of this book). Often genes are presented on the chips by unique oligonucleotides close to the 3'-end of the mRNA. But there are many predicted new genes that have no known corresponding EST sequences. We can study the expression of such genes in a large number of human tissues using predicted exon sequences represented on one or several Affymetrix type DNA chips. As a result we will know not only expression properties of genes, but we can identify what exons are real. Observing coordinated expression of neighboring exons in different tissues it will often be possible to define gene boundaries, which is very difficult using *ab initio* gene prediction. Moreover such experiments might have additional value in defining disease

Tab. 3.18

PfamA domains identified in the predicted human genes. Domain of the same type localized in neighboring exons were counted only once.

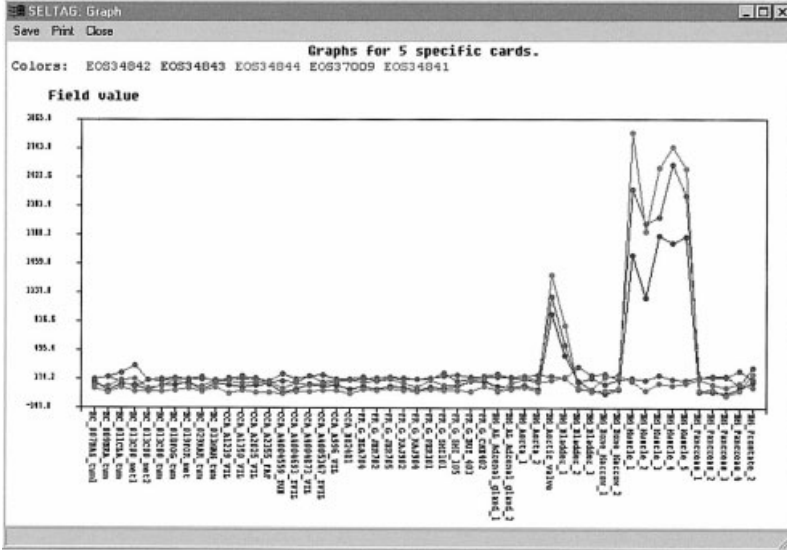
<b>Number</b>	<b>PfamA short name</b>	<b>Name</b>
467	Pkinase	Eukaryotic protein kinase domain
372	7tm_1	7 transmembrane receptor (rhodopsin family)
308	Myc_N_term	Myc amino-terminal region
256	Topoisomerase_I	Eukaryotic DNA topoisomerase I
224	Ig	Immunoglobulin domain
183	Rrm	RNA recognition motif
182	PH	PH domain
180	Myosin_tail	Myosin tail
166	EGF	EGF-like domain
159	Filament	Intermediate filament proteins
154	Syndecan	Syndecan domain
143	Ras	Ras family
138	RNA_pol_A	RNA polymerase A/beta'/A" subunit
123	BTB	BTB/POZ domain
119	Granin	Granin (chromogranin or secretogranin)
119	Troponin	Troponin
113	Herpes_glycop_D	Herpesvirus glycoprotein D
111	Homeobox	Homeobox domain
110	SH3	SH3 domain
102	Trypsin	Trypsin
102	helicase_C	Helicases conserved C-terminal domain
100	KRAB	KRAB box
98	dehydrin	Dehydrins
96	ABC_tran	ABC transporter
95	ERM	Ezrin/radixin/moesin family
89	Collagen	Collagen triple helix repeat
87	Tryp_mucin	Mucin-like glycoprotein
84	Fn3	Fibronectin type III domain
81	pro_isomerase	Cyclophilin type peptidyl-prolyl cis-trans isomerase
81	HMG_box	HMG (high mobility group) box
79	SH2	Src homology domain 2

tissue specific genes, which can be used for the development of potential therapeutics.

The chip designed by EOS Biotechnology included all exons from Chromosome 22 predicted by Fgenesh and Genescan as well as exons from human genomic sequences of Phase 2 and 3 predicted by Fgenesh. It was found that the predicted exon sequences present a good alternative to EST sequences that open a possibility to work with predicted genes on a large scale.

In Figure 3.13 we have an example of expression behavior of five sequential exons along the Chromosome 22 sequence (expression data were received in EOS Biotechnology Inc.). Exons 2, 3 and 4 are Myoglobin gene exons. Tissue specific expression of them is clear seen with the major peaks located





**Fig. 3.13**  
Coordinative expression of three exons (EOS34842, EOS34842, EOS34842) of the Human Myoglobin gene from Chromosome 22 (exons were predicted by the Fgenesh program and used to design the EOS Biotechnology Human genome chip) in 50 different tissues. The high level of expression is observed only in several

specific tissues. The two exons of Myoglobin predicted on the left side (EOS34841) and on the right side (EOS37009) respectively show completely different patterns of expression. The visualization is presented by the SELTAG program for analysis of expression data developed by Softberry Inc.

in skeletal muscle, heart and diaphragm tissues. The level of expression in these tissues is 10–100 times higher than the level of signals for other tissues as well as the average level of expression for randomly chosen exons. We found that three Myoglobin exons have expression level correlation coefficient 0.99, when for random exons it is about 0.06. These exons were predicted correctly by the Fgenesh program and were used for selection of oligonucleotide probes. From this result we can conclude that the predicted exons can be used as gene representatives. At the same time two flanking exons (1 and 5) from different genes show no correlation with the Myoglobin exons. This clearly demonstrates how expression data can be used to define gene boundaries. Another application of expression data is functional analysis and identification of alternatively spliced genes (exons), if in particular tissues some exons (or their parts) have very different expression intensities comparing with the other exons from the same gene. If 5'-alternative exons define different functional forms of genes (normal and disease-specific, for example), then the 3'-EST generated probes cannot be used for identification of disease specific gene variants.

## 3.9

## Internet sites for gene finding and functional site prediction

Gene identification, and finding of ORFs, promoters and splice sites by the methods described above is available via World Wide Web. Table 3.19 presents a list of some useful programs but does not provide a comprehensive collection.

Tab. 3.19

Public web servers for eukaryotic gene and functional signal prediction.

<b>Program/function</b>	<b>World Wide Web address</b>
<b>Fgenesh</b> /HMM-based gene prediction (Human, Drosophila, Dicots, Monocots, C.elegans, S.pombe)	<a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a> <a href="http://searchlauncher.bcm.tmc.edu:9331/seq-search/gene-search.html">http://searchlauncher.bcm.tmc.edu:9331/seq-search/gene-search.html</a> <a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>
<b>Genscan</b> /HMM-based gene prediction (Human, Arabidopsis, Maize)	<a href="http://genes.mit.edu/GENSCAN.html">http://genes.mit.edu/GENSCAN.html</a>
<b>HMM-gene</b> /HMM-based gene prediction (Human, C.elegans)	<a href="http://www.cbs.dtu.dk/services/HMMgene/">http://www.cbs.dtu.dk/services/HMMgene/</a>
<b>Fgenes</b> /Disciminative gene prediction (Human)	<a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a> <a href="http://searchlauncher.bcm.tmc.edu:9331/seq-search/gene-search.html">http://searchlauncher.bcm.tmc.edu:9331/seq-search/gene-search.html</a>
<b>Fgenes-M</b> /Prediction of alternative gene structures (Human)	<a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a> <a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>
<b>Fgenesh + /Fgenesh_c</b> /gene prediction with the help of similar protein/EST	<a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a> <a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>
<b>Fgenesh-2</b> /gene prediction using 2 sequences of close species	<a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>
<b>BESTORF</b> /Finding best CDS/ORF in EST (Human, Plants, Drosophila)	<a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a> <a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>
<b>Mzef</b> /internal exon prediction (Human, Mouse, Arabidopsis, Yeast)	<a href="http://argon.cshl.org/genefinder/">http://argon.cshl.org/genefinder/</a>
<b>TSSW/TSSG</b> /eukaryotic promoter prediction	<a href="http://searchlauncher.bcm.tmc.edu:9331/seq-search/gene-search.html">http://searchlauncher.bcm.tmc.edu:9331/seq-search/gene-search.html</a> <a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a>
<b>Promoter 2.0</b> /promoter prediction	<a href="http://www.cbs.dtu.dk/services/Promoter/">http://www.cbs.dtu.dk/services/Promoter/</a>
<b>PromoterScan</b> /eukaryotic promoter prediction	<a href="http://cbs.umn.edu/software/proscan/promoterscan.htm">http://cbs.umn.edu/software/proscan/promoterscan.htm</a>
<b>CorePromoter</b> /promoter prediction	<a href="http://argon.cshl.org/genefinder/CPROMOTER/index.htm">http://argon.cshl.org/genefinder/CPROMOTER/index.htm</a>
<b>SPL</b> /splice site prediction (Human, Drosophila, Plants, Yeast)	<a href="http://genomic.sanger.ac.uk/gf/gf.shtml">http://genomic.sanger.ac.uk/gf/gf.shtml</a> <a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>
<b>NetGene2/NetPGene</b> /splice site prediction (Human, C.elegans, Plants)	<a href="http://www.cbs.dtu.dk/services/NetPGene/">http://www.cbs.dtu.dk/services/NetPGene/</a>
<b>Dbscan</b> /searching for similarity in genomic sequences and its visualization altogether with known gene structure	<a href="http://www.softberry.com/nucleo.html">http://www.softberry.com/nucleo.html</a>

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## 4

# Analyzing Regulatory Regions in Genomes

*Thomas Werner*

### 4.1

#### General features of regulatory regions in eukaryotic genomes

Regulatory regions share several common features despite their obvious divergence in sequence. Most of these common features are not evident directly from the nucleotide sequence but result from the restraints imposed by functional requirements. Therefore, understanding of the major components and events during the formation of regulatory DNA–protein complexes is crucial for the design and evaluation of algorithms for the analysis of regulatory regions. Transcription initiation from polymerase II (pol II) is the best understood example so far and will be a major focus of this chapter. However, the mechanisms and principles revealed from promoters are mostly valid for other regulatory regions as well.

Algorithms for the analysis and recognition of regulatory regions draw from the underlying biological principles, to some extent, in order to generate suitable computational models. Therefore, a brief overview over the biological requirements and mechanisms is necessary to understand what are the strengths and weaknesses of the individual algorithms. The choice of parameters and implementation of the algorithms largely control sensitivity and speed of a program. The specificity of software recognizing regulatory regions in DNA is determined, to a large extent, by how closely the algorithm follows what will be called the biological model from hereon. A detailed overview of this topic was recently published [1].

### 4.2

#### General functions of regulatory regions

The biological functionality of regulatory regions is generally not a property evenly spread over the regulatory region in total. Functional units usually are defined by a combination of defined stretches that can be delimited

and possess an intrinsic functional property (e.g., binding of a protein or a curved DNA structure). Several functionally similar types of these stretches of DNA are already known and will be referred to as *elements*. Those elements are neither restricted to regulatory regions nor individually sufficient for the regulatory function of a promoter or enhancer. The function of the complete regulatory region is composed from the functions of the individual elements either in an additive manner (independent elements) or by synergistic effects (modules).

#### 4.2.1

##### **Transcription factor binding sites (TF-sites)**

Binding sites for specific proteins are most important among these elements. They consist of about 10–30 nucleotides, not all of which are equally important for protein binding. As a consequence individual protein binding sites vary in sequence, even if they bind to the same protein. There are nucleotides contacted by the protein in a sequence-specific manner which are usually the best conserved parts of a binding site. Different nucleotides are involved in DNA backbone contacts, i.e., contacting the sugar-phosphate framework (not sequence specific as they do not involve the bases A, G, C, or T), and there are internal “spacers” not contacted by the protein at all. In general, protein-binding sites exhibit enough sequence conservation to allow for the detection of candidates by a variety of sequence similarity-based approaches. Potential binding sites can be found almost all over the genome and are not restricted to regulatory regions. Quite a number of binding sites outside regulatory regions are also known to bind their respective binding proteins [2]. Therefore the abundance of predicted binding sites is not just a shortcoming of the detection algorithms but reflects biological reality.

#### 4.2.2

##### **Sequence features**

Regulatory DNA also contains several features not directly resulting in recognizable sequence conservation. For example, the sequences of two copies of a direct repeat (approximate or exact) are conserved in sequence with respect to each other but different direct repeats are not similar in sequence at all. Nevertheless, direct repeats are quite common within regulatory DNA regions. They consist either of short sequences which are repeated twice or more frequently within a short region or they can be complex repeats which repeat a pattern of two or more elements. Repeat structures are often associated with enhancers. Enhancers are DNA structures that enhance transcription over a distance without being promoters themselves. One example

of a highly structured enhancer is the interleukin-2 enhancer [3]. Other sequence features that are hard to detect by computer methods include the relatively weak nucleosomal positioning signals [4], DNA stretches with intrinsic 3-D structures (like curved DNA [5]), methylation signals (if there are definite signals for methylation at all), and other structural elements.

#### 4.2.3

##### **Structural elements**

Secondary structures are currently the most useful structural elements with respect to computer analysis. Secondary structures are mostly known for RNAs and proteins but they also play important roles in DNA. Potential secondary structures can be easily determined and even scored via the negative enthalpy that should be associated with the actual formation of the hairpin (single strand) or cruciform (double strand) structure. Secondary structures are also not necessarily conserved in primary nucleotide sequence but are subject to strong positional correlation within the structure. Three-dimensional aspects of DNA sequences are without any doubt very important for the functionality of such regions. However, existing attempts to calculate such structures in reasonable time met with mixed success and cannot be used for a routine sequence analysis at present.

#### 4.2.4

##### **Organizational principles of regulatory regions**

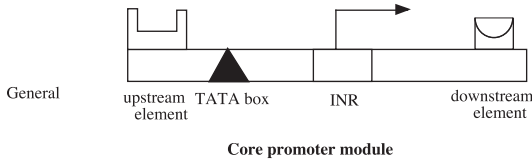
This paragraph will mainly concentrate on eukaryotic polymerase II promoters as they are currently the best-studied regulatory regions. The TF-sites within promoters (and likewise most other regulatory sequences) do not show any general patterns with respect to location and orientation within the promoter sequences. Even functionally important binding sites for a specific transcription factor may occur almost anywhere within a promoter. For example, functional AP-1 (Activating protein 1, a complex of two transcription factors, one from the fos and one from the Jun family) binding sites can be located far upstream, as in the rat bone sialoprotein gene where an AP-1 site located about 900 nucleotides upstream of the transcription start site (TSS) inhibits expression [6]. An AP-1 site located close to the TSS is important for the expression of Moloney Murine Leukemia Virus [7]. Moreover, functional AP-1 sites have also been found inside exon 1 (downstream of the TSS) of the proopiomelanocortin gene [8] as well as within the first intron of the fra-1 gene [9], both locations outside the promoter. Similar examples can be found for several other TF-sites, illustrating why no general correlation of TF-sites within specific promoter regions can be defined. TF binding sites can be found virtually everywhere in promoters

but in individual promoters possible locations are much more restricted. A closer look reveals that the function of an AP-1 binding site often depends on the relative location and especially on the sequence context of the binding site. The AP-1 site in the above mentioned rat bone sialoprotein gene overlaps with a set of glucocorticoid responsive element (GRE, the DNA sequence that is bound by the glucocorticoid receptor which is a transcription factor) half sites which are crucial for the suppressive function.

The context of a TF-site is one of the major determinants of its role in transcription control. As a consequence of context requirements, TF sites are often grouped together and such functional groups have been described in many cases. A systematic attempt of collecting synergistic or antagonistic pairs of TF binding sites has been made with the COMPEL database [10, 11]. In many cases, a specific promoter function (e.g., a tissue-specific silencer) will require more than two sites. Promoter subunits consisting of groups of TF binding sites that carry a specific function independent of the promoter will be referred to as *promoter modules*. A more detailed definition of promoter modules has been given recently by Arnone and Davidson [12]. In summary, promoter modules contain several transcription factor binding sites which act together to convey a common function like tissue-specific expression. The organization of binding sites (and probably also of other elements) of a promoter module appears to be much more restricted than the apparent variety of TF sites and their distribution in the whole promoter suggests. Within a promoter module both sequential order and distance can be crucial for function indicating that these modules may be the critical determinants of a promoter rather than individual binding sites. Promoter modules are always constituted by more than one binding site. Since promoters can contain several modules that may use overlapping sets of binding sites, the conserved context of a particular binding site cannot be determined from the primary sequence. The corresponding modules must be detectable separately before the functional modular structure of a promoter or any other regulatory DNA region can be revealed by computer analysis. One well known general promoter module is the core promoter, which will be discussed in more detail below. However, the basic principles of modular organization are also true for most if not all other regulatory regions and are neither peculiar nor restricted to promoters.

#### 4.2.4.1 Module properties of the core promoter

The core promoter module can be defined functionally by its capability to assemble the transcription initiation complex (see Figure 4.3) and orient it specifically towards the TSS of the promoter [13], defining the exact location of the TSS. Various combinations of about four distinguishable core promoter elements that constitute a general core promoter can achieve this. This module includes the TATA box, the initiator region (INR), an upstream activating element, and a downstream element. (The TATA box is a basic



**Fig. 4.1** General structure of a polymerase II core promoter. Simultaneous presence of all four elements is not always essential. INR = initiator region. The shapes above the bar symbolize additional protein binding sites and the arrow indicates the transcription start site.

transcription element, which is located about 20–30 nucleotides upstream of the actual transcription start site and is known to bind to the TATA box binding protein TBP.) However, this is also where the straightforward definition of a core promoter module ends because not all four elements are always required, or some elements can be too variable to be recognizable by current computer tools.

#### 4.2.4.2 First group: TATA box containing promoters without known initiator

Successful positioning of the initiation complex can start at the TATA box containing promoters by the TFIID complex, which contains the TATA box binding protein as well as several other factors. Together with another complex of general transcription factors, termed TFIIB, this leads to the assembly of an initiation complex [14]. If an appropriate upstream TF binding site cooperates with the TATA box, no special initiator or downstream sequences might be required, which allows for the assembly of a functional core promoter module from just two of the four elements (Figure 4.1). This represents one type of a distinct core promoter that contains a TATA box, common among cellular genes in general.

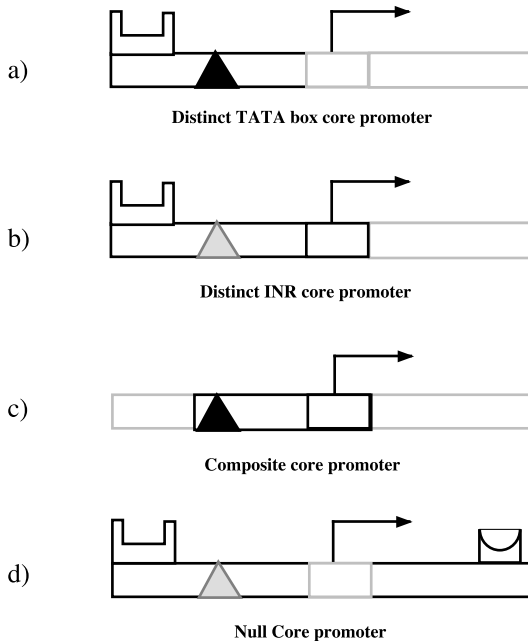
#### 4.2.4.3 Second group: TATA-less promoters with functional initiator

However, as it is known from a host of TATA-less promoters, the TATA box is by no means an essential element of a functional core promoter. An INR combined with a single upstream element has also been shown to be capable of specifically initiating transcription [15], although initiators cannot be clearly defined on the sequence level so far. Generally, a region of 10–20 nucleotides around the transcription start site is thought to represent the initiator. A remarkable array of four different upstream TF sites (SP1, AP-1, ATF, or TEF1) was shown to confer inducibility by T-antigen to this very simple promoter. T-antigen is a potent activating protein from a virus called SV40. This is an example of a TATA-less distinct promoter that can be found in several genes from the hematopoietic lineage (generating blood cells).

The third combination is called a composite promoter and consists of both a TATA box and an initiator. This combination can be found in several viral promoters and it has been shown that an additional upstream TF binding site can influence whether the TATA box or the initiator element will be determining the promoter properties [16]. The authors showed that upstream elements can significantly increase the efficiency of the INR in this combination while especially SP-1 sites made the TATA box almost obsolete in their example. The combination of TATA box with an INR had the general effect to induce resistance against the detrimental effects of a TFIIB mutant, which interfered with expression from TATA-only promoters. This is also an example for the more indirect effects of specific arrangements in promoters that may not be apparent unless special conditions occur.

The last group consists of so-called null-promoters which have neither a TATA box nor an initiator and rely exclusively on upstream and downstream elements [17].

Basically, at least the four different core promoter types detailed above have been identified so far, which all represent valid combinations of core promoter sites (reviewed in [17]). If the combinations involving upstream and downstream elements are also considered, a total of 7 possible core promoter modules are possible (most of which can be actually found in genes and consist of the four variants in Figure 4.2 adding upstream or downstream elements or both).



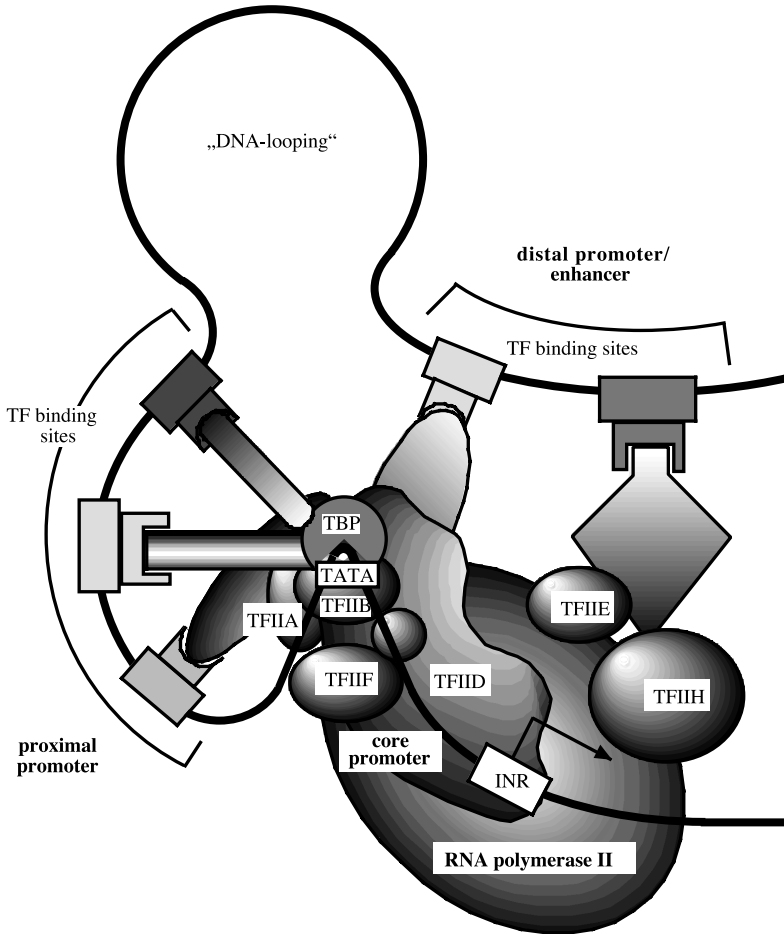
**Fig. 4.2**  
Four different setups of a polymerase II core promoter. Graphical symbols as in Figure 4.1.

The only apparent common denominator of transcription initiation within a promoter would be that there must be at least one (or more) core promoter element within a certain region. This assumption is wrong. Both spacing and/or sequential order of elements within the core promoter module are of utmost importance regardless of the presence or absence of individual elements (as a rule, there appear to be some exceptions). Moreover, many distinct promoters have requirements for specific upstream or downstream elements and will function with their specific TF. Moving around the initiator, the TATA box and, to some extent, also upstream elements can have profound effects on promoter functions. For example, insertion of just a few nucleotides between the TATA box and an upstream TF binding site (MyoD) in the desmin gene promoter cuts the expression levels by more than half [18]. Moreover, the promoter structure can affect later stages of gene expression like splicing [19]. It was also shown for the rat beta-actin promoter that a few mutations around the transcription start site (i.e. within the initiator) can render that gene subject to translational control [20].

As a final note, the mere concept of one general TATA box and one general INR is an oversimplification. There are several clearly distinguishable TATA boxes in different promoter classes [21, 22, 23, 24] and the same is true for the INR region which also has several functionally distinct implementations as the glucocorticoid-responsive INR in the murine thymidine kinase gene [25], the C/EBP binding INR in the hepatic growth factor gene promoter [26], or the YY-1 binding INR [27].

Most of the principles of variability and restrictions detailed above for the core promoter modules are also true for other promoter modules that modify transcriptional efficiency rather than determining the start point of transcription as the core promoter does. The bottom line is that the vast majority of alternative arrangements of the elements that can be seen in a particular promoter might not contribute to the function of the promoter. Module-induced restrictions are not necessarily obvious from the primary sequences. Figure 4.3 shows a schematic pol II promoter with the initiation complex assembled which illustrates that it matters where a specific protein is bound to the DNA in order to allow for proper assembly of the molecular jigsaw puzzle of the initiation complex. This is not immediately obvious from inspection of promoter sequences because there exist several (but a strictly limited set of) alternative solutions to the assembly problem. As complicated as Figure 4.3 may appear, it still ignores all aspects of chromatin rearrangements and nucleosomal positions which also play an important role in transcription regulation. Stein et al. [28] has detailed an example of the profound influence of these effects on promoter-protein complex assembly and function for the osteocalcin promoter in a study. However, chromatin-related effects are not yet considered in any of the promoter prediction methods. Therefore, we do not go into any more details here.





**Fig. 4.3**  
Transcription initiation complex bound to a schematic promoter

#### 4.2.5

#### Bioinformatics models for the analysis and detection of regulatory regions

Algorithms used to analyze and detect regulatory regions are necessarily based on some kind of usually simplified model of what a regulatory region should look like. All of these models are inevitably compromising between accuracy with respect to the biological model (the standard of truth) and computational feasibility of the model. For example, a computational model based on *a priori* three-dimensional structure prediction derived from

molecular dynamics using sophisticated force fields may be the most accurate model for a region but cannot be used for the analysis of real data due to excessive demand on computational resources. On the other hand a model based on simple sequence similarities detected by IUPAC consensus (see Chapter 3 for detailed definition) sequences can be easily used on a PC but results will usually not match the biological truth in an acceptable manner.

#### 4.2.5.1 Statistical models

It has been noted several years ago that promoters and most likely also other regulatory regions like enhancers contain more transcription factor binding sites than non-regulatory sequences. An analysis of the relative frequencies of such sites within a sliding window therefore can yield some information about the potential regulatory character of a stretch of DNA, which is the prototype of simple statistical models. Several programs exist that rely to some extent on this type of statistics.

#### 4.2.5.2 Mixed models

Of course, it is quite clear from section 4.1 that a pure statistical model is an oversimplification that will adversely affect the accuracy of prediction despite its attractive ease of implementation. Therefore, mixed models are also used that take at least some regional information into consideration and can be seen as statistical models split into compartments. Within the compartments solely statistical features are considered, but promoter organization is somewhat reflected by the arrangement of the compartments which represent different promoter regions.

#### 4.2.5.3 Organizational models

The last category consists of models that try to closely follow the organizational principles of real regulatory regions. In order to accomplish that, individual promoter elements like transcription factor binding sites as well as their relative order and distances are encoded in a formal model which reflects the setup of a single promoter or a small group of functionally similar promoters. Although they are matching the biological situation best they are not yet suitable for widespread application. High quality sets of training sequences are required to generate such models because many parameters concerning nature, order and distance of elements have to be determined. This is a process far from being an easy task and limits application of such models to relatively few examples that require elaborate analysis.

### 4.3

#### Methods for element detection

##### 4.3.1

#### Detection of transcription factor binding sites

Transcription factor binding sites are the most important elements within regulatory DNA regions like promoters or enhancers. The majority of the known transcription factors recognize short DNA stretches of about 10–15 nucleotides in length that show different degrees of internal variation. Successful detection of protein binding sites in DNA sequences always relies on precompiled descriptions of individual binding sites. Such descriptions are usually derived from a training set of four or more authentic binding sites. However, the criteria applied for the decision whether a site is authentic or not vary considerably among authors of different publications. One of the first approaches to define protein binding sites used IUPAC consensus sequences, which indicate the predominant nucleotide or nucleotide combination at each position in a set of example sequences. The IUPAC string TGASTCA indicates that the first three positions are most frequently T, G, and A while the fourth position may be C or G followed by T, C, and A in most cases. IUPAC consensus sequences became very popular as they are extremely easy to define from even a small set of sequences and their definition does not require more than a pencil and a sheet of paper.

However, IUPAC consensus sequences strongly depend on the sequence set used for definition. The final IUPAC consensus sequence remains arbitrary depending on the rules used to determine the consensus. Cavener defined some rules that we have used at GSF for several years now and, in our experience, IUPAC consensus sequences defined that way can be useful [29]. However, IUPAC consensus sequences may reject biologically functional binding sites due to a single mismatch (or an ill-defined IUPAC sequence).

The concept of nucleotide weight matrix (NWM) descriptions has been developed in the 1980s as an alternative to IUPAC strings [30, 31]. However, although weight matrices proved to be generally superior to IUPAC strings their biggest disadvantage is the absolute requirement for predefined matrices, which are more complicated to construct than IUPAC strings and require specific software. This delayed widespread use of weight matrices for almost a decade although the methods were principally available. They remained mostly unused because only a few special matrices had been defined [32]. The situation changed when in 1995 two (overlapping) matrix libraries for TF sites were compiled and became widely available almost simultaneously [33, 34]. Matrix Search [34] transformed the TRANSFAC database as complete as possible (starting at two binding sites for one factor) into matrices using a log-odds scoring approach. The MatInspector library is largely based on a stringent selection from the matrix table

of the TRANSFAC database, including the matrices derived from the ConInspector library [35, 36, 37]. and several genuine matrices. The Information Matrix Database was compiled from the TRANSFAC matrix table and the TFD. The MatInspector library is updated with each new TRANSFAC release while IMD has not been updated recently. Available matrix detection programs have been reviewed recently [36] and a comparison of these methods by application to a test set of sequences was published [37]. For convenience Table 4.1 summarizes which methods for the detection of transcription factor binding sites are available in the internet with emphasis on programs featuring a WWW-interface.

#### 4.3.2

##### **Detection of structural elements**

Regulatory sequences are associated with a couple of other individual elements or sequence properties in addition to the factor binding sites. Among these are secondary structure elements like the HIV-1 TAR region (Trans-ActivatingRegion, which constitutes an RNA enhancer [38], cruciform DNA structures [39], or simple direct repeats [40]. Three-dimensional structures like curved DNA [41] also influence promoter function. Most of these elements can be detected by computer-assisted sequence analysis [42, 43, 44, 45] but none of them is really promoter specific and all such elements can be found frequently outside of promoters. The promoter or enhancer function arises from the combination of several elements that need to cooperate to exert transcription control which none of them can achieve alone. This also illustrates the main problem of promoter recognition. It is necessary to compile several individually weak signals into a composite signal, which then indicates a potential promoter without being overwhelmed by the combinatorial complexity of potential element combinations.

#### 4.3.3

##### **Assessment of other elements**

Several methods employ statistical measures of sequence composition to include features of regulatory sequences, which cannot be described by the three types discussed above. This includes frequencies of oligonucleotides (dinucleotides, trinucleotides, and hexamers are used most frequently), CpG islands (CG dinucleotides are usually underrepresented in mammalian genomes except in part of coding and regulatory sequences. CpG islands are regions where the dinucleotide is NOT underrepresented [46], and periodicity of weak sequence patterns (AA, TT etc). Definitions of such elements are usually too weak to allow any significant contribution to current prediction programs. However, this situation might well change due to

**Tab. 4.1**  
Internet accessible methods to detect promoter elements (transcription factor binding sites)

<b>Program</b>	<b>Availability</b>	<b>Comments</b>
MatInspector	<a href="http://www.gsf.de/cgi-bin/matsearch.pl">http://www.gsf.de/cgi-bin/matsearch.pl</a>	MatInspector matrix library (includes TRANSFAC matrices)
SIGNAL SCAN	<a href="http://genomatix.gsf.de/cgi-bin/matinspector/matinspector.pl">http://genomatix.gsf.de/cgi-bin/matinspector/matinspector.pl</a>	IUPAC consensus library
MATRIX SEARCH	<a href="http://bimas.dcr.t.nih.gov/molbio/signal/">http://bimas.dcr.t.nih.gov/molbio/signal/</a>	IMD matrix library (TRANSFAC + TFD)
TFSearch	<a href="http://bimas.dcr.t.nih.gov/molbio/matrix/">http://bimas.dcr.t.nih.gov/molbio/matrix/</a>	TRANSFAC matrices
TESS	<a href="http://pdap1.trc.rwcp.or.jp/research/db/TFSEARCH.html">http://pdap1.trc.rwcp.or.jp/research/db/TFSEARCH.html</a>	TRANSFAC matrices
	<a href="http://agave.humgen.upenn.edu/utess/tess/">http://agave.humgen.upenn.edu/utess/tess/</a>	TRANSFAC matrices

the unprecedented amounts of continuous genomic sequences that become available in the course of the current genome sequencing projects.

#### 4.4

#### **Analysis of regulatory regions**

Basically two different tasks can be distinguished in the analysis of regulatory regions. The first task is analysis aiming at the definition of common features based on sets of known regulatory sequences. This is a prerequisite for the definition of descriptions suitable for large-scale application for prediction of potential regulatory regions within new anonymous sequences which can be regarded as the second task.

##### 4.4.1

#### **Training set selection**

One of the most important steps in comparative sequence analysis is the selection of suitable training sets of sequences. If a training set of promoters consists only of constitutively expressed sequences (constant level of expression, no or little regulation) little can be learned about any kind of tissue-specific expression regardless of the methods applied. Also inclusion of too many wrong sequences (e.g., that are no promoters at all) may prevent any meaningful analysis. Although this appears a bit trivial at first, it is a real issue when data are scarce and less well-characterized sequences have to be used.

Control sets known not to be functionally similar to the training sets are about as important as the training sets themselves. However, true negative regions are even scarcer than known regulatory regions. Negative often means just “no positive functions found” which can also be due to failures or simply means that the sequences have not been tested at all. Therefore, statistical negative control sequences are often required. Random sequences can be easily generated but often are of limited use, as they do not represent several important features of natural DNA correctly. This includes under-represented features (e.g., CpG islands), asymmetric features (e.g., strand specificity), or repetitive DNA elements. Selection of appropriate control sequences can be a major effort, but is also crucial for the validity of the evaluation of any method. Common problems with controls are either known or unknown biases in the control set or circularity problems, i.e. the training and the test sets of sequences are related or overlap. The availability of large continuous stretches of genomic DNA from the genome sequencing projects constantly improves this situation. Genomic sequences should always be the first choice for controls as they reflect the natural situation.

## 4.4.2

**Statistical and biological significance**

The quality of sequence pattern recognition is often optimized to improve the correlation of the methods with the data (positive and negative training sets). However, in most cases it is not possible to collect sufficient data to perform a rigorous correlation analysis. Therefore, bioinformatics methods often rely on statistical analysis of their training sequences and optimize for statistically most significant features. Unfortunately, this kind of optimization does not always reflect the evolutionary optimization of regulatory sequences that is always optimizing several features at once.

The dynamics of biological function often necessitates suboptimal solutions. For example, real sequences usually do not contain binding sites with the highest affinity for their cognate protein because binding AND dissociation of the protein are required for proper function. The perfect binding site would interfere with the dissociation and is therefore strongly selected against.

## 4.4.3

**Context dependency**

The biological significance of any sequence element is defined by the regulatory function it can elicit. This is usually dependent on a functional context rather than being a property of individual elements. Therefore, statistical significance of the features or scores of individual elements is neither necessary nor sufficient to indicate biological significance. Recognition of the functional context in an essentially linear molecule like DNA can be achieved by correlation analysis of individual elements, which became an important part of all semi-statistical or specific modeling approaches discussed below.

## 4.5

**Methods for detection of regulatory regions**

There are several methods available for the prediction of regulatory DNA regions in new sequence data. Table 4.2 lists methods available with a special focus on programs that provide a WWW-interface. Unfortunately, there is no “one-does-it-all” method and all methods have their individual strong and weak points. A program doing an excellent job in one case might be a complete failure in another case where other methods are successful. Therefore, we will describe a whole lot of methods without intending any rank by order of discussion. We will rather follow the functional hierarchy that appears to apply to the different regulatory regions.

Tab. 4.2  
Available promoter/promoter regions prediction tools

<b>Program</b>	<b>Availability</b>	<b>Comments</b>
<b>Promoter prediction</b>		<b>WWW-accessible</b>
FunSiteP	<a href="http://transfac.gbf.de/dbsearch/funsitep/fsp.html">http://transfac.gbf.de/dbsearch/funsitep/fsp.html</a>	includes proximal promoter
NNPP	<a href="http://www-hgc.lbl.gov/projects/promoter.html">http://www-hgc.lbl.gov/projects/promoter.html</a>	core promoter
PromFD	<a href="http://beagle.colorado.edu/~cheng/Hypertexts/PromFD.html">http://beagle.colorado.edu/~cheng/Hypertexts/PromFD.html</a> <a href="mailto:chenq@beagle.colorado.edu">chenq@beagle.colorado.edu</a>	includes proximal promoter for further information
Promoter Scan	<a href="http://biosci.umn.edu/software/proscan/promoterscan.htm">http://biosci.umn.edu/software/proscan/promoterscan.htm</a>	includes proximal promoter
TSSG/TSSW	<a href="http://dot.imgen.bcm.tmc.edu:9331/gene-finder/gf.html">http://dot.imgen.bcm.tmc.edu:9331/gene-finder/gf.html</a>	includes proximal promoter triplet & hexamer frequencies discrimination analysis
Core Promoter Finder	<a href="http://sciclio.cshl.org/genefinder/CPROMOTER/human.htm">http://sciclio.cshl.org/genefinder/CPROMOTER/human.htm</a>	
	<b>Download/inquire</b>	
Audic/Claverie	<a href="mailto:audic@newton.cnrs-mrs.fr">audic@newton.cnrs-mrs.fr</a>	Markov models
PromFind	<a href="ftp://iubio.bio.indiana.edu/molbio/ibmpc">ftp://iubio.bio.indiana.edu/molbio/ibmpc</a> <a href="http://www.rabbithutch.com">http://www.rabbithutch.com</a>	hexamer frequencies for further information
XGRAIL	<a href="ftp://arthur.epm.ornl.gov/pub/xgrail">ftp://arthur.epm.ornl.gov/pub/xgrail</a>	core promoter in gene context
<b>Promoter module/region recognition</b>		<b>WWW-accessible</b>
FastM library	<a href="http://www.gsf.de/cgi-bin/fastm.pl">http://www.gsf.de/cgi-bin/fastm.pl</a> <a href="http://genomatix.gsf.de/cgi-bin/fastm2/fastm.pl">http://genomatix.gsf.de/cgi-bin/fastm2/fastm.pl</a> <a href="http://gcg.tigem.it/TargetFinder.html">http://gcg.tigem.it/TargetFinder.html</a>	module of 2 TF sites uses MatInspector
TargetFinder	<b>Download/inquire</b> <a href="http://www.gsf.de/biodv/genomeinspector.html">http://www.gsf.de/biodv/genomeinspector.html</a> <a href="ftp://ariane.gsf.de/pub/unix/genomeinspector/ficketjw@molbio.sbphtd.com">ftp://ariane.gsf.de/pub/unix/genomeinspector/ficketjw@molbio.sbphtd.com</a> <a href="ftp://beagle.colorado.edu/pub/Landscape/xland.v1.tar.Z">ftp://beagle.colorado.edu/pub/Landscape/xland.v1.tar.Z</a>	module of 1 TF site combined with 1 annotated feature correlation analysis, e.g 2 TF binding sites
GenomeInspector		
Muscle-specific regions		contact for download
Xlandscape		word frequencies, not promoter specific



## 4.5.1

**Types of regulatory regions****4.5.1.1 Matrix attachment regions**

A chromatin loop is the region of chromosomal DNA located between two contact points of the DNA with the nuclear matrix marked by so-called Scaffold/Matrix Attachment Regions (S/MARs). The nuclear matrix is a mesh of proteins lining the inner surface of the nuclear envelope. Transcriptional regulation requires the association of DNA with this nuclear matrix, which retains a variety of regulatory proteins. S/MARs are composed of several elements including transcription factor binding sites, AT-rich stretches, potential cruciform DNA, and DNA-unwinding regions to name a few of the most important S/MAR elements. There is an excellent review on chromatin domains and prediction of MAR sequences by Boulikas [47], explaining S/MARs and their elements in detail. Kramer et al. [48] published a method to detect potential S/MAR elements in sequences and made the method available via WWW (<http://www.ncgr.org/MAR-search/>) [49]. Their method is based on a statistical compilation of the occurrence of a variety of S/MAR features (called rules). Accumulation of sufficient matches to these rules will be predicted as potential S/MAR regions. The specificity of the method depends critically on the sequence context of the potential S/MAR sequences. Therefore, results are difficult to evaluate by comparisons. However, so far this is the only method available to predict S/MAR regions. We developed another approach to define especially AT-rich MARs called SMARTest which is available on the web at <http://genomatix.gsf.de>. SMARTest is based on a library of MAR-associated nucleotide weight matrices and determines S/MARs independent of any larger sequence context. Therefore, the method is suitable to test isolated S/MAR fragments. MARFinder and SMARTest are complementary and should be seen in combination rather than as alternatives.

**4.5.1.2 Enhancers/silencers**

Enhancers are regulatory regions that can significantly boost the level of transcription from a responsive promoter regardless of their orientation and distance with respect to the promoter as long as they are located within the same chromatin loop. Silencers are basically identical to enhancers and follow the same requirements but exert a negative effect on promoter activities. At present there are no specific programs to detect enhancers and silencers. However, programs designed to detect the internal organization of promoters are probably also suitable to detect enhancers and silencers since these elements often also show a similar internal organization as promoters.

### 4.5.1.3 Promoters

Promoters are DNA regions capable of specific initiation of transcription (start of RNA synthesis) and consist of three basic regions (See Figure 4.2). The part determining the exact nucleotide for transcriptional initiation is called the core promoter and is the stretch of DNA sequence where the RNA polymerase and its cofactors assemble on the promoter.

The region immediately upstream of the core promoter is called the proximal promoter and usually contains a number of transcription factor binding sites responsible for the assembly of an activation complex. This complex in turn recruits the polymerase complex. It is generally accepted that most proximal promoter elements are located within a stretch of about 250–500 nucleotides upstream of the actual transcription start site (TSS).

The third part of the promoter is located even further upstream and is called the distal promoter. This region usually regulates the activity of the core and the proximal promoter and also contains transcription factor binding sites. However, distal promoter regions and enhancers exhibit no principal differences. If a distal promoter region acts position and orientation independent it is called an enhancer.

## 4.5.2

### Programs for recognition of regulatory sequences

There are several ways promoter recognition tools can be categorized. We will focus on the main principles and intended usage of the programs rather than technical details, as this will also be the main interest of experimentally working scientists. Two generally distinct approaches have been used so far in order to achieve *in silico* promoter recognition [50]. The majority of programs focused on *general promoter recognition*, which represents the first category. One group of programs in this category (see below) concentrates on recognition of core promoter properties and infers promoter location solely on that basis while the other group consists of programs that take into account also the proximal promoter region of about 250–300 nucleotides upstream of the TSS. General recognition models are usually based on training sets derived from the Eukaryotic Promoter Database (EPD [51]) and various sets of sequences without known promoter activities. The EPD is an excellent collection of DNA sequences that fulfil two conditions: They have been shown experimentally to function as promoters and the transcription start site is known. The beauty of these approaches is their generality which does not require any specific knowledge about a particular promoter in order to make a prediction. This appears ideal for the analysis of anonymous sequences for which no *a priori* knowledge is available. The bad news is that the specificity of all such general approaches implemented so far is limited. The inevitable huge burden of false-positive predictions in

longer sequences precludes large scale application of such approaches to genomic sequences.

The second category of tools aims at *specific promoter recognition* relying on more detailed features of promoter subsets like combinations of individual elements. The beauty of this approach is excellent specificity, which is extremely helpful if only promoters of a certain class are of interest or megabases of sequences have to be analyzed. The bad news here is limited applicability, i.e., each promoter group or class requires a specifically pre-defined model before sequences can be analyzed for these promoters. This may result in a huge number of false negatives in large-scale analysis.

We will briefly discuss individual methods in these two categories with emphasis on the implementation of the biological principles of promoter features. Recently, a practical comparison of the majority of available tools based on general promoter models has been carried out, which has shown that none of these methods is clearly superior to its peers [50]. Therefore, I will not go into details about performance of the methods here (see Chapter 3 for details).

#### 4.5.2.1 Programs based on statistical models (general promoter prediction)

These programs aim at the detection of pol II promoters by a precompiled general promoter model that is part of the method. Despite the complicated modular structure of promoters outlined above there is a solid rational basis for this general model. All promoters must have a functional core promoter module often containing a TATA box which is the prime target of the majority of the general promoter prediction tools. This is also one of the reasons why some programs confine their analysis to the core promoter region which avoids problems with the much more diverse proximal regions. General models that include the proximal region consequently treat this part of the promoter as a purely statistical problem of TF binding site accumulations, sometimes fine tuned by some sort of weighting based on occurrence frequencies of TF binding sites in promoters as compared to a negative sequence set.

However, the cost of generality without exception is a huge number of false positive predictions (typically about one match in 10 000–30 000 nucleotides). Sacrifice of a considerable percentage of true promoters (30% or more) is also a necessity to maintain at least some specificity. There also is inevitably no clue what kind of promoter was detected in case of a match.

##### 4.5.2.1.1 PromoterScan

Several of the general promoter prediction programs followed the basic design of Prestridge who used the Eukaryotic Promoter Database (EPD) by Bucher [51] to train his software for promoter recognition. His program *PromoterScan* was the first published method to tackle this problem [52].

He utilized primate non-promoter sequences from GenBank as a negative training set and included the proximal promoter region in the prediction. The program uses individual profiles for the TF binding sites indicative of their relative frequency in promoters to accumulate scores for DNA sequences analyzed. PromoterScan employs the SignalScan IUPAC library of TF binding sites [53], introducing a good deal of biological knowledge into the method, although modular organization of the proximal region is necessarily ignored. Results of the first version were combined with the Bucher NWM for the TATA box, which served as a representation of the core promoter module [32].

There is now a new version of *PromoterScan* available, PromoterScan 2.0. This new version is supposed to provide more information from inside the “black box” which a promoter used to be for version 1.0 and is also able to compare a predicted promoter to EPD promoter sequences on the basis of the pattern of TF sites. Although this moves promoter comparisons with PromoterScan 2.0 effectively closer to the specific recognition of individual elements this approach is not used for the initial promoter prediction. PromoterScan is available via WWW, which is a definitive advantage for occasional promoter testing.

#### 4.5.2.1.2 PromFD

This program [54], extends the model behind *PromoterScan* in two ways. One major difference is use of the IMD matrix library [34] instead of the SIGNAL SCAN IUPAC strings. The other feature is the inclusion of patterns of strings of 5–10 nucleotides in length that were found to be overrepresented in the training set of promoters. Basically, the intrinsic model is the same as in *PromoterScan* but the overrepresented strings may account for some so far unknown binding sites that are missing in the libraries. PromFD requires local installation and is not available via a WWW-interface. There have been no updates so far.

#### 4.5.2.1.3 TSSG/TSSW

Two other methods (*TSSG* and *TSSW*) also including proximal promoter regions are available via WWW and share the basic algorithm. They center on detection of a TATA box as most prominent part of the core promoter. Promoter prediction is then based on the score of the TATA box, and nucleotide triplet distributions around the putative TSS. In analogy to PromFind (see below) hexamer frequencies in three 100 nucleotide wide windows upstream of the putative TSS are also considered. These data are combined with potential TF binding sites, which are predicted either based on a TFD derived compilation by Prestridge (in TSSG), or on the TRANSFAC database [11] in TSSW. Fickett and Hatzigeorgiou did not report significant improvements in the predictive capabilities by inclusion of triplets and

hexamers [50]. However, the general aim of these programs is not independent promoter prediction but to assist in finding exon-intron structures of complete genes.

#### 4.5.2.1.4 **XLandscape**

The group of Stormo developed another method, which is called *XLandscape* and essentially determines nucleotide strings of various lengths called words which are specifically associated with promoters, exons or introns [55]. Then a score for a sequence is determined indicating by the different promoter, exon, and intron scores whether a particular region is likely to belong to one of these groups. Although this method was not developed for promoter prediction it fares about as well as all other general promoter prediction programs although it completely ignores even the core promoter module. The program requires local installation.

#### 4.5.2.1.5 **PromFind**

This program [56] is similar to *XLandscape*. It relies on the difference in hexamer frequencies between promoters and regions outside of promoters which is a more restricted view as compared to the landscaping approach. The advantage is less computational complexity. The program was also trained on sequences from the EPD and corresponding coding and non-coding regions outside the promoters taken from GenBank. Any region in which the ratio of promoter to non-promoter hexamer-frequencies reaches a threshold is considered and only the region where this measure is maximal is defined as the promoter. This again relies on pure statistics disregarding any organizational features of promoters. I am not aware of any updates since the initial publication.

#### 4.5.2.1.6 **NNPP**

This program [57] utilizes time delay neural networks to locate a TATA box combined with an initiator region and thus is a representative of the second subgroup which focuses on the minimum promoter region. Although the program does allow for variable spacing between the elements, especially distinct and null promoters will pose principal problems for this method as it includes some modeling of a TATA box. An improved version of the program was published recently and the program features a lower number of false negatives (about 1 match/kb). This is good news if a short region of DNA is to be analyzed (length should be less than 2 kb). However, long DNA regions pose a problem, as the number of false positives becomes overwhelming. The program is available via a WWW-interface (<http://www-hgc.lbl.gov/projects/promoter.html>).

#### 4.5.2.1.7 CorePromoter Search

Michael Zhang [58] published a new method to detect TATA-box containing core promoters by discrimination analysis. This method is available via a WWW-interface, which already requires restriction of the sequence length to 1 kb. Core Promoter Search and NNPP are alternative implementations of a similar general promoter model and can be applied in parallel.

#### 4.5.2.2 Programs utilizing mixed models

These programs also rely on statistical promoter models but include directly or indirectly some organizational features of promoters placing them in between the pure statistical models and attempts to approximate the biologically important structured organization of promoters.

##### 4.5.2.2.1 FunSiteP

This program [59] also takes into account proximal promoter regions and utilizes a collection of TF binding sites [60] with which a promoter set taken from EMBL (472 promoters) was analyzed. From this analysis a weight matrix of TF binding site localization was derived representing regions in promoters with lower or higher concentrations of TF binding sites. FunSiteP not only reports potential promoter matches but also assigns them to one of seven promoter classes. These were taken from Bucher's definition (from EPD) and consist of small nuclear RNAs, structural proteins, storage and transport proteins, enzymes, hormones, growth factors, and regulatory proteins, stress or defense related proteins, and unclassified proteins. Although these classes are very broad as defined by biological function, they represent an attempt towards more specific promoter recognition. FunSiteP is also available via a WWW-interface.

##### 4.5.2.2.2 Audic/Claverie

A program designed by Audic and Claverie [61] uses Markov models of vertebrate promoters generated again by training on the EPD and non-promoter sequences outside of the promoters. Markov models principally allow for the inclusion of organizational features and the nature of the training set determines whether this becomes part of the model or not. We describe the program in this section because the many different promoters in EPD most likely cause these Markov models to be more general than specific.

#### 4.5.2.3 Programs relying on organizational models

This category of methods introduces the functional context in form of heuristic rules or tries to learn the context from comparative sequence analysis.

These methods emphasize specific modeling of promoters or promoter substructures rather than general recognition. Therefore, it is not possible to directly assess the promoter prediction capabilities of these methods. However, in many cases recognizing a common substructure between promoters can be very helpful especially for experimental design.

#### 4.5.2.3.1 **FastM**

This method was derived from the program ModelGenerator [22] and takes advantage of the existence of NWM libraries. It can be accessed via a WWW-interface (<http://www.genomatix.de> “free services”) and allows for a straightforward definition of any modules of two TF binding sites by simple selection from the MatInspector Library [33]. This now enables definition and detection of wide variety of synergistic TF binding site pairs. These pairs often are functional promoter modules conferring a specific transcriptional function to a promoter [10, 62]. *FastM* models of two binding sites can successfully identify promoters sharing such composite elements but are not promoter specific. Composite elements can also be located in enhancers or similar structures. A commercial version of FastM is available that enables definition of complete, highly specific promoter class models including up to 10 individual elements.

#### 4.5.2.3.2 **TargetFinder**

Another approach aiming at modeling of promoter substructures consisting of two distinct elements is TargetFinder [63]. This method combines TF binding sites with features extracted from the annotation of a database sequence to allow selective identification of sequences containing both features within a defined length. The advantage is that TargetFinder basically also follows the module-based philosophy but allows inclusion of features that have been annotated by experimental work for which no search algorithm exists. Naturally, this excludes analysis of new anonymous sequences. The program is accessible via a WWW-interface (<http://gcg.tigem.it/TargetFinder.html>).

It should be mentioned here that Fickett also employed the idea of a two TF binding site module to successfully detect a subclass of muscle-specific regulatory sequences governed by a combination of MEF2 and MyoD [64]. However, this was also a very specific approach and no general tool resulted out of that work. The MEF2/MyoD model can be used to define a corresponding module with FastM. Wasserman and Fickett [65] recently published a modeling approach based on clustering of a preselected set of NMW (defined in this study) correlated with muscle-specific gene expression. They were able to detect about 25% of the muscle-specific regulatory regions in sequences outside their training set and more than 60% in their

training set. They classify their method as regulatory module detection. However, their results suggest that they probably detect a collection of different more specific modules with respect to the definition given above. Although the method is not promoter-specific and the specificity is moderate, it is a very interesting approach, which has potential for further development. The authors will make the non-commercial software used in their approach available on request (contact J. Fickett).

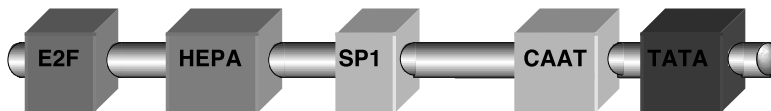
Generally, this group of methods achieves much higher specificity than programs following general models. However, the price for this increase in specificity is usually restriction of the promoter models to a small subset (class) of promoters.

#### 4.5.2.4 The organizational model of histone H1 promoters

The specific modeling of a promoter class can be demonstrated on a well-known example from the cell-cycle regulation. Histone genes are required during the DNA replication and they show up as a group of coregulated genes in array analyses [66]. This suggests that they might have a common promoter structure. Histone H1 gene promoters for example can be found in the Eukaryotic Promoter Database (EPD, [51]).

The model for Histone H1 genes (Figure 4.4) was based on only 9 training sequences taken from homology group 17 of the EPD. They share a 100 bp fragment around the transcription start site but not much similarity elsewhere in the 600 nucleotides (standard length of EPD promoter sequences). A model containing 5 different transcription factor binding sites was derived from the set of sequences shown in Figure 4.4 using GEMS Launcher (Genomatix Software, Munich).

Notably, the model generated contains one factor, E2F that is known to be involved in cell cycle regulation of genes [67]. The model appears to be very selective and only one match per 12 million base pairs of the mammalian sections of the EMBL database was found. Almost all matches (total of 59) are known Histone H1 genes, except in the human section where the majority of matches (25) are within anonymous sequences. However, given the extraordinary specificity throughout the mammalian sections, it seems safe to assume that most of the unknown matches within the human database section actually identified new so far unannotated genes, that are subject to



**Fig. 4.4**  
Organizational model of histone H1 promoters containing six transcription factor binding sites.



a similar regulation as histones (There is no direct evidence that these are histone genes). However, this is a working hypothesis which needs to be verified.

#### 4.6

#### Annotation of large genomic sequences

Almost all of the methods discussed above were developed before the databases started to be filled with sequence contigs exceeding 100 000 nucleotides in length. The complete human genome draft now contains more than 3 billion nucleotides. This changes the paradigm for sequence annotation. While complete annotation remains an important goal, specific annotation becomes mandatory when even individual sequences exceed the capabilities of researchers for manual inspection. Annotation of genomic sequences has to be fully automatic in order to keep pace with the rate of generation of new sequences. Simultaneously, annotations are embedded into a large natural context rather than residing within relatively short isolated stretches of DNA. This has several quite important consequences.

##### 4.6.1

##### The balance between sensitivity and specificity

I will confine the discussion here to regulatory regions but the problems are general. A very sensitive approach will minimize the amount of false negative predictions and thus is oriented towards a complete annotation. However, this inevitably requires accepting large numbers of false positive hits, which easily outnumber the true positive predictions by one order of magnitude.

In order to avoid this methods can be designed to yield the utmost specificity (e.g., specific promoter modeling as discussed above). Here, the catch is that inevitably a high number of false-negative results, which also may obscure 70% to 90% of the true positive regions.

A little thought experiment demonstrates the dilemma of current methods for annotation of sequences. Assume we are analyzing a region of 3 billion bp of contiguous human genomic DNA (the total genome). The human genome is estimated to contain about  $60\,000 \pm 30\,000$  promoters. On average general promoter prediction programs detect one promoter per 1000 to 10 000 nucleotides according to the respective authors. Assuming an optimistic value (1/10 000) this would result in about 300 000 predictions. Given the true match rate determined by Fickett and Hatzigeorgiou [50] of less than 20% on average the result would be 20 000 true predictions versus 280 000 or more than 90% false positives.

On the other hand specific methods were shown to produce more than 50% true positive matches in their total output [21] but recognize just a small fraction of all promoters. A single specific model like the actin class model [21] matches about once every 2.5 million bps and thus would yield a total of 1200 matches, 600 of which can be expected to be true. This is great in terms of specificity, but losing more than 90% of the true promoters present is certainly far from what sequence annotation is aiming at. The existence of 100 or, most likely, even more promoter models of the specificity of the actin model would be required to achieve specific recognition of most of the promoters present within the genome. Definition of the required number of specific models based on current technologies is not a feasible task. Therefore, new developments have already been initiated to overcome the current obstacles.

The numbers mentioned above are necessarily very rough estimates. However, two- or three-fold variations would not change the general results. It is quite evident that functional promoter analysis in laboratories is capable of dealing with several hundred or even thousand predicted regions while predicting several hundred thousand or even millions of regions remains out of reach. It is safe to assume that further improvement of laboratory high throughput technologies and enhancement of the specificity of promoter recognition *in silico* will meet somewhere in the future to close the gap in our knowledge about the functional regulation of the genome. It is also quite clear from the past and present developments that bioinformatics will probably cover significantly more than half of that path.

#### 4.6.2

##### **The larger context**

There will be help on the way towards more specific modeling of functional regions in genomic sequences. The almost unlimited natural context of regions in genomic sequences will allow for completely new approaches to comparative sequences analysis, which has already proven to be the most powerful approach in bioinformatics of genomic sequences (e.g., in detection of new sequence elements). Comparative analysis will be instrumental in determining the anatomy of regulatory networks including MARs, LCRs, enhancers, and silencers in addition to the promoter sequences. It can be safely assumed that context-sensitive sequence analysis will prevail in the long run over any methods dealing with short isolated sequences.

#### 4.6.3

##### **Aspects of comparative genomics**

The context information is by no means restricted to other regions within the same genome. Approaches based on comparative genomics employing

sequence information from two or more genomes proved to be very powerful as they introduce phylogenetic aspects into the analysis. Phylogenetic footprints have already shown this [67]. This way, discrimination between noise and functionally conserved regions can be improved resulting in easier detection of regions useful for predictive efforts.

#### 4.6.4

#### **Analysis of data sets from high throughput methods**

Another field to which the bioinformatics of regulatory DNA regions can be expected to contribute significantly is the analysis of results from high throughput experiments in expression analysis (e.g., all forms of expression arrays). Due to the discontinuous nature of regulatory regions there is no way of deducing common regulatory features from the expression data directly that are usually based on coding regions. However, the general availability of the corresponding genomic regulatory regions for many (and very soon all) of the genes analyzed in an expression array experiment enables attempts to elucidate the genomic structures underlying common expression patterns of genes. Expression arrays (described in detail in Chapter 6, Volume 2) directly deliver information, *which* genes are *where* expressed under the conditions tested. However, they cannot provide any clue to *why* this happens or how the same genes would behave under yet untested conditions. Identification of functional features by comparative sequence analysis (e.g., promoter modules) can reveal different functional subgroups of promoters despite common regulation under specific conditions. Consequently, the detection of known functional modules can suggest expression patterns under yet untested conditions. Moreover, the organizational structures of promoters can also be used to identify additional potential target genes either within the same organism in other genomes or via comparative genomics. Given the exponential number of possibilities for combinations of conditions, bioinformatics of regulatory sequences will also become instrumental for the rational design of expression arrays as well as for selection of experimental conditions.

#### 4.7

#### **Conclusions**

The experimental dissection of functional mechanisms of transcription control has gained an enormous momentum during recent years. The ever rising number of publications on this topic bears witness to this development which found one hallmark manifestation in the introduction of a new section in the *Journal of Molecular and Cellular Biology* entirely devoted to analysis of transcription control. The complex interleaved networks of tran-

scription control certainly represent one of the cornerstones on which to build our understanding of how life functions, in terms of embryonic development, tissue differentiation, and maintenance of the shape and fitness of adult organisms throughout life. This is also the reason why both the experimental analysis as well as the bioinformatics of transcription control will move more and more into the focus of medical/pharmaceutical research. A considerable number of diseases are directly or indirectly connected to alterations in cellular transcription programs (e.g., most forms of cancer). Furthermore, many drugs influence transcription control via signaling pathways (triggering transcription factors) which could also be connected to certain side effects of drugs. The human genome sequencing project will provide us with a complete catalog of the components of a human probably within a year. This will constitute a blueprint of the material basis of a human. However, only the analysis of the regulatory part of the genome and the corresponding expression patterns and the complex metabolic networks will provide deeper insight into how the complex machinery called life is working. Definition and detection of regulatory regions by bioinformatics will contribute to this part of the task.

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## 5 Homology Modeling in Biology and Medicine

Roland L. Dunbrack, Jr.

### 5.1 Introduction

#### 5.1.1 The concept of homology modeling

To understand basic biological processes such as cell division, cellular communication, metabolism, and organismal development and function, knowledge of the three-dimensional structure of the active components is crucial. Proteins form the key players in all of these processes, and study of their diverse and elegant designs is a mainstay of modern biology. The Protein Databank (PDB) of experimentally determined protein structures [1, 2] now contains some 18 000 entries, which can be grouped into between 300 and 700 related families based either on structure or on sequence similarity [3–5]. The fact that proteins that share very little or no sequence similarity can have quite similar structures has led to the hypothesis that there are in fact on the order of 1000–7000 different families [6, 7] which have been adapted by a process of duplication, mutation, and natural selection to perform all the biological functions that proteins perform.

Since it was first recognized that proteins can share similar structures [8], computational methods have been developed to build models of proteins of unknown structure based on related proteins of known structure [9]. Most such modeling efforts, referred to as homology modeling or comparative modeling, follow a basic protocol laid out by Greer [10, 11]: 1) identify a *template* or *parent* structure related to the *target* sequence of unknown structure, and align the target sequence to the template sequence and structure; 2) for core secondary structures and all well-conserved parts of the alignment, borrow the backbone coordinates of the template according to the sequence alignment of the target and template; 3) for segments of the target sequence for which coordinates cannot be borrowed from the template because of insertions and deletions in the alignment (usually in loop regions of the protein), build these segments using some construction

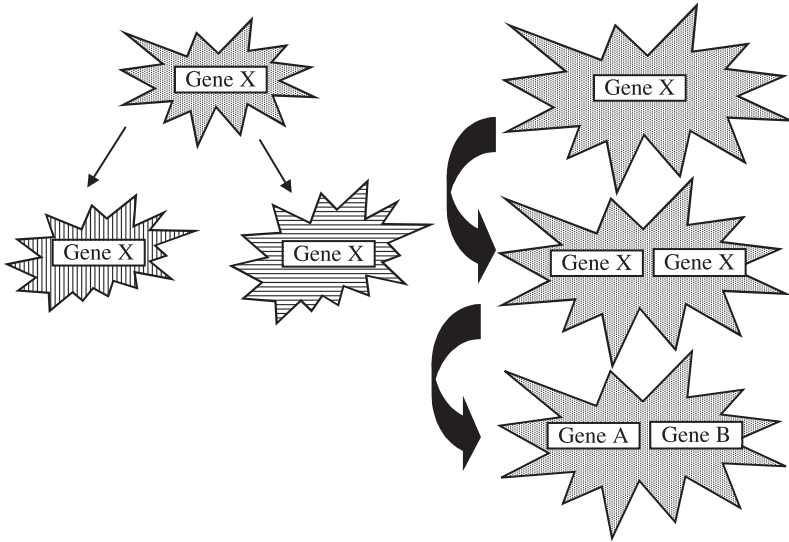
method based on our knowledge of the determinants of protein structure; 4) build side chains determined by the target sequence on to the backbone model built from the template structure and loop construction. In practice, identifying a structural homologue requires aligning the sequences, and so steps 1 and 2 are performed together. Step 2 is often a manual adjustment of the alignment, but automated sequence alignment methods different from those used to make the identification may be used. Also, Steps 3 and 4, backbone and side-chain modeling, may be coupled, since certain backbone conformations may be unable to accommodate the required side chains in any low-energy conformation. An alternative strategy has been developed by Blundell and colleagues, based on averaging a number of template structures, if these exist, rather than using a single structure [12–14]. More complex procedures based on reconstructing structures (rather than perturbing a starting structure) by satisfying spatial restraints using distance geometry [15] or molecular dynamics and energy minimization [16–19] have also been developed.

Many methods have been proposed to perform each of the steps in the homology modeling process. There are also a number of research groups that have developed complete packages that take as input a sequence alignment or even just a sequence and develop a complete model. In this Chapter, we describe some of the basic ideas that drive loop and side-chain modeling individually and the programs publicly available that implement them. We also discuss some of the programs that perform all of the steps in modeling. The identification and alignment steps will be covered in the next Chapter of this volume.

### 5.1.2

#### **How do homologous proteins arise?**

By definition, homologous proteins arise by evolution from a common ancestor. But there are several different mechanisms in play, and these are illustrated in Figure 5.1. The first is random mutation of individual nucleotides that change protein sequence, including missense mutations (changing the identity of a single amino acid) as well as insertions and deletions of a number of nucleotides that result in insertion and deletion of amino acids. As a single species diverges into two species, a gene in the parent species will continue to exist in the divergent species and over time will gather mutations that change protein sequence. In this case, the genes in the different organisms will usually maintain the same function. These genes are referred to as orthologues of one another. A second mechanism is duplication of a gene or of a gene segment within a single organism or germ line cell. As time goes by, the two copies of the gene may begin to gather mutations. If the parent gene performed more than one function, for example similar catalytic activity on two different substrates, one of the



**Fig. 5.1**  
Orthologues vs. paralogues. Schematic of evolutionary process that gives rise to homologous proteins. Left: A single gene X in one species is retained as the species diverges into two separate species. The genes in these two species are *orthologous*. Right: A single gene X in one species is duplicated. As each gene gathers mutations, it may begin to perform new

functions, or the two genes may specialize in carrying out two or more functions of the ancestral gene, thus improving the fitness of the organism. These genes in one species are *paralogous*. If the species diverges, each daughter species may maintain the duplicated genes, and therefore each species contains an orthologue and a paralogue to each gene in the other species.

duplicated genes may gain specificity for one of the reactions, while the other gene gains specific activity for the other. If this divergence of specificity in the two proteins is advantageous, the duplication will become fixed in the population. These two genes are paralogues of one another. If the species with the pair of paralogues diverges into two species, each species will contain the two paralogues. Each gene in each species will now have an orthologue and a paralogue in the other species.

### 5.1.3

#### The purposes of homology modeling

Homology modeling of proteins has been of great value in interpreting the relationships of sequence, structure, and function. In particular, orthologous proteins usually show a pattern of conserved residues that can be interpreted in terms of three-dimensional models of the proteins. Conserved

residues often form a contiguous active site or interaction surface of the protein, even if they are distant from each other in the sequence. With a structural model, a multiple alignment of orthologous proteins can be interpreted in terms of the constraints of natural selection and the requirements for protein folding, stability, dynamics, and function.

For paralogous proteins, three-dimensional models can be used to interpret the similarities and differences in the sequences in terms of the related structure but different functions of the proteins concerned [20]. In many cases, there are significant insertions and deletions and amino acid changes in the active or binding site between paralogues. But by grouping a set of related proteins into individual families, orthologous within each group, the evolutionary process that changed the function of the ancestral sequences can be observed. Indeed, homology models can serve to help us identify which protein belongs to which functional group by the conservation of important residues in the active or binding site [21]. A number of recent papers have been published that use comparative modeling to predict or establish protein function [22–26].

Another important use of homology modeling is to interpret point mutations in protein sequences that arise either by natural processes or by experimental manipulation. Now that the human genome project has produced a rough draft of the complete human genome sequence, there are starting to be collected voluminous data concerning polymorphisms and other mutations related to differences in susceptibility, prognosis, and treatment of human disease. There are now many such examples, including the Factor V/Leiden R506Q mutation [27] that causes increased occurrence of thrombosis, mutations in cystathionine beta synthase that cause increased levels of homocysteine in the blood, a risk factor for heart disease [28], and BRCA1 for which many sequence differences are known, some of which may lead to breast cancer [29]. At the same time, there are many polymorphisms in important genes that have no discernible effect on those who carry them. At least for some of these, there may be some effect that has yet to be measured in a large enough population of patients, and the risk of cancer, heart disease, or other illness to these patients is unknown. This is yet another important application of homology modeling, since a good model may indicate readily which mutations pose a likely risk and which do not.

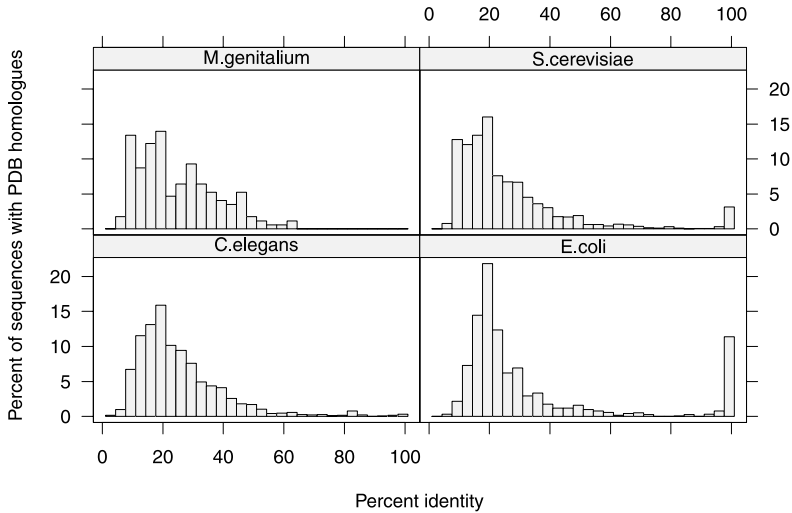
Homology models may also be used in computer-aided drug design, especially when a good template structure is available for the target sequence. For enzymes which maintain the same catalytic activity, the active site may be sufficiently conserved that a model of the protein provides a reasonable target for computer programs which can suggest the most likely compounds that will bind to the active site (see also Chapter 7 of this volume). This has been used successfully in the early development of HIV protease inhibitors [30, 31] and in the development of anti-malarial compounds that target the cysteine protease of *P. falciparum* [32].

## 5.1.4

**The effect of the genome projects**

The many genome projects now completed or underway have greatly affected the practice of homology modeling of protein structures. First, the many new sequences have provided a large number of targets for modeling. Second, the large amount of sequence data makes it easier to establish remote sequence relationships between proteins of unknown structure and those of known structure on which a model can be built. The most commonly used methods for establishing sequence relationships such as PSI-BLAST [33] are dependent on aligning many related sequences to compile a pattern or profile of sequence variation and conservation for a sequence family. This profile can be used to search among the sequences in the PDB for a relative of the target sequence. The more numerous and more varied sequences are in the family the more remote are the homologous relationships that can be determined, and the more likely it is that a homologue of known structure for a target sequence can be found. Third, it is likely that the accuracy of sequence alignments between the sequence of unknown structure that we are interested in (referred to as the target sequence) and the protein sequence of known structure used for model building (referred to as the template structure) are also greatly improved with profiles established from many family members of the target sequence [34]. Fourth, the completion of a number of microbial genomes has prodded a similar effort among structural biologists to determine the structures of representatives of all common protein sequence families, or all proteins in a prototypical genome, such as *E. coli* or yeast [35]. Protein structures determined by X-ray crystallography or NMR spectroscopy are being solved at a much faster pace than was possible even 10 years ago. The great increase in the number of solved protein structures has a great impact on the field of homology modeling, since it becomes ever more likely that there will be a template structure in the PDB for any target sequence of interest.

Given the current sequence and structure databases, it is of interest to determine what fraction of sequences might be modeled and the range of sequence identities between target sequences and sequences of known structure. In Figure 5.2, we show histograms of sequence identities of the sequences in several genomes and their nearest relatives of known structure in the Protein Data Bank. These relationships were determined with PSI-BLAST as described in the legend. PSI-BLAST is fairly sensitive in determining distant homology relationships [34, 36, 37]. The results indicate that on average 30–40% of genomic protein sequences are related to proteins of known structure, which presents a large number of potential targets for homology modeling. But it should also be pointed out that the average sequence identity between target sequences and template structures in the PDB is less than 25%. It is likely that as sequence comparison methods improve, our ability to identify increasingly remote homologies will in-



**Fig. 5.2**

Distribution of sequence identities between proteins in four genomes and their closest homologues in the Protein Databank for those sequences in genomes with homologues in the PDB. PSI-BLAST was used to search the non-redundant protein sequence database with a representative set of PDB sequences as queries. The program was run for four iterations, with a maximum expectation value of 0.0001 (see Chapter 2 for an

explanation of this value) used to determine sequences which are included in the position-specific similarity matrix. After four iterations, each matrix was used to search each of the four genomes. Coiled-coil and low-sequence complexity sequences were removed from each genome and the non-redundant sequence database. All hits in the genomes with E-values less than 0.001 were saved, and the histograms were built from the PSI-BLAST derived sequence identities.

crease, and we will be able to determine relationships at even lower sequence identity with confidence [38].

The low sequence identity between target and template sequences in Figure 5.2 presents a major challenge for homology modeling practitioners, since a major determinant in the accuracy of homology modeling is the sequence identity between the target sequence and the sequence of the template structure. At levels below 30% sequence identity, related protein structures diverge significantly and there may be many insertions and deletions in the sequence [39]. At 20% sequence identity, the average RMSD of core backbone atoms is 2.4 Å [39]. But as demonstrated in Figure 5.2, it is likely that we will most often face a situation where the target and template sequences are remotely related. Most widely used homology modeling methods have been predicated on much higher sequence identities between template and target, usually well above 30% [40–42]. What methods should be used at sequence identities in the 10–30% range is of crucial importance in this so-called post-genomic era.

## 5.2

### Input data

To produce a protein model that will be useful and informative requires more than placing a new sequence onto an existing structure. A large amount of sequence data and other kinds of experimental data can often be gathered on the target sequence and on its homologue of known structure to be used for model building. This information can be used to build a better model *and* as the data to be interpreted in light of the model. The goal is to forge an integrated model of the protein sequence, structure, and function, not merely to build a structure. In Table 5.1, we list the kinds of information which might be available for a target protein and how these data might be processed. With the large amount of sequence information available, it is almost always possible to produce a multiple alignment of

**Tab. 5.1**  
Input information for homology model building

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#### Target sequence

- Target orthologous relatives (from PSI-BLAST)
- Target paralogous relatives (from PSI-BLAST)
- Multiple sequence alignment of orthologues and paralogues (either BLAST multiple alignment or (preferably) other multiple alignment program)
- Sequence profile of ortho/paralogues

#### Template sequence

- Homologue(s) of known structure (template(s)) determined by database search methods (BLAST, PSI-BLAST, intermediate-sequence-search methods, HMM's, fold recognition methods)
- Template orthologous sequences
- Template paralogous sequences
- Multiple sequence alignment of parent orthologues and paralogues

#### Alignment of target sequence to template sequence and structure

- Pairwise alignment
- Profile alignment
- Multiple sequence alignment of target and template sequence relatives
- Profile-profile alignment
- Fold recognition alignment
- Visual examination of proposed alignments and manual adjustment
- Assessment of confidence in alignment by residue (some regions will be more conserved than others)

#### Structure alignment of multiple templates, if available

- Align by structure (fssp, VAST, CE, etc.)
- Compare sequence alignments from structure to sequence alignments from multiple sequence alignments (see above)

#### Experimental information

- Mutation data (site-directed, random, naturally occurring)
  - Functional data – e.g. DNA binding, ligands, metals, catalysis, etc.
-

sequences related to the target protein. The first step in modeling therefore is to use a database search program such as PSI-BLAST [33] against a non-redundant protein sequence database such as NCBI's *nr* database [43] or the curated SwissProt database [44]. With some care, a list of relatives to the target sequence can be gathered and aligned. PSI-BLAST provides reasonable multiple alignments, but it may be desirable to take the sequences identified by the database search and realign them with a multiple sequence alignment program such as CLUSTAL W [45], prrp [46], and multalin [47]. Another source of protein sequences related to the target is the database of expressed sequence tags (ESTs) available at NCBI. ESTs are derived from sequencing DNA transcriptions of mRNAs derived from cellular samples from various species or tissue types. While these sequences contain some non-coding nucleotides, the bulk of the sequence represents transcribed codons. This DNA sequence database can be searched with protein sequence queries using the TBLASTN program (<http://www.ncbi.nlm.nih.gov/blast>). NCBI also makes available the sequences of unfinished microbial genomes for TBLASTN searches ([http://www.ncbi.nlm.nih.gov/Microb\\_blast/unfinishedgenome.html](http://www.ncbi.nlm.nih.gov/Microb_blast/unfinishedgenome.html)), and a search of these may provide additional sequences related to the target.

It may be that a database search consisting of several rounds of PSI-BLAST will provide one or more sequences of known three-dimensional structure. If this is not the case then more sensitive methods based on fold recognition [48–79] (see Chapter 6 of this volume) or hidden Markov models (see Appendix to this volume) [80–87] of protein superfamilies may identify a suitable template structure. Once a template structure is identified, a sequence database search will provide a list of relatives of the template, analogous to searches for relatives of the target. At this stage it is useful to divide the sequences related to the target into orthologues and paralogues of either the target or the template (or both). The sequence variation within the set of proteins that are orthologous to the target provides information as to what parts of the sequence are most conserved and therefore likely to be most important in the model. Similarly variation in the set of proteins that are orthologous to the template provide a view of the template protein family that can be used to identify features in common or distinct in the template and target families. These features can be used to evaluate and adjust a joint multiple alignment of both families.

If there are multiple structures in the Protein Data Bank that are homologous to the target sequence, then it is necessary to evaluate them to determine which PDB entry will provide the best template structure and whether it will be useful to use more than one structure in the modeling process. In the case of a single sequence that occurs in multiple PDB entries, it is usually a matter of selecting the entry with the highest resolution or the most appropriate ligands (DNA, enzyme inhibitors, metal ions). In other cases, there may be more than one homologue related to the target sequence, and the task is to select the one more closely related to the target



or to combine information from more than one template structure to build the model. To do this, a structure alignment of the potential templates can be performed by one of a number of computer programs available on the Internet (Dali [88], CE [89], etc., see Table 5.6). From alignments of the target to the available templates, the location of insertions and deletions can be observed and often it will be clear that one template is better than others. This may not be uniform, however, such that some regions of the target may have no insertions or deletions with respect to one template, but other regions are more easily aligned with the other template. In this case, a hybrid structure may be constructed [13].

Finally, any other experimental information available on the target or template proteins may be very helpful in producing and interpreting a structural model. This can include inhibitor studies, DNA binding and sequence motifs, proteolysis sites, metal binding, mutagenesis data, and so forth. A number of databases are available on the web that summarize information on particular genes, or that collect information on mutations and polymorphisms linked to disease including: the Cancer Genome Anatomy Project [90]; the Online Mendelian Inheritance in Man (OMIM) [91]; YDB (Yeast database), WormDB (*C. elegans* database), and PombeDB (*Schizosaccharomyces pombe* database), all available from <http://www.proteome.com/databases/>; and the Human Gene Mutation Database [92].

Finally, we note that many methods in homology modeling rely on statistical data on protein conformations. We present a primer on some aspects of protein structure in the Appendix to this Chapter.

## 5.3 Methods

### 5.3.1 Modeling at different levels of complexity

Once an alignment is obtained between the target and a protein of known structure (as described in “Input Data” or in Chapter 6 of this volume), it is possible to build a series of models of increasing sophistication.

#### 5.3.1.1 Simple model

Keep backbone and conserved side chains by renaming and renumbering coordinates in the template structure with the new sequence using the alignment of target and template; rebuild other side chains using a side-chain modeling program (e.g., SCWRL [93, 94]); do not model insertions or deletions; that is, do not build new loops and do not close up gaps. For this purpose, we have made available a Perl program called *blast2model* that

uses BLAST alignments as input and the SCWRL program to build model structures (<http://www.fccc.edu/research/labs/dunbrack/software.html>).

#### 5.3.1.2 Stepwise model

Borrow core backbone from template structure, minus coil regions with insertions or deletions in the sequence alignment; rebuild core side chains; rebuild coil regions with loop prediction method in conjunction with side-chain prediction method. Core backbone and side chains may or may not be held fixed during loop prediction (e.g., CONGEN [95]).

#### 5.3.1.3 Global model

Build entire protein from spatial restraints drawn from known structure(s) and sequence alignment (e.g., MODELLER [16, 17]).

#### 5.3.1.4 Choosing a model

It is not always the case that more sophisticated models are better than simpler, less complete ones. If elements of secondary structures are allowed to move away from their positions in the template and large changes are made to accommodate insertions and deletions, it may be the case that the model is further away from the target structure (if it were known) than the template structure was to begin with. This is the “added value” problem discussed by John Moult at the CASP meetings [96–98]. We would like methods that change the template structure closer to the target structure, such that they “add value” to a simple model based on an unaltered template structure, perhaps with side chains replaced. Extensive energy minimization or molecular dynamics simulations often bring a model further away from the correct structure than toward it [99].

The simple model is sometimes justified when there are no insertions and deletions between the template and target or when these sequence length changes are far from the active site or binding site of the protein to be modeled. This often occurs in orthologous enzymes that are under strong selective pressure to maintain the geometry of the active site. Even in non-orthologous enzymes, sometimes we are most interested in an accurate prediction of the active site geometry and not in regions of the protein distant from the active site. Unless loop modeling is accurate, it is possible that modeled loops in more sophisticated models may get in the way of side-chain modeling. If uncertain parts of the chain are not modeled, then side chains may find their correct positions and the model may be superior to a more complete version.

A stepwise model is probably the most common method used in homology modeling, since it is conceptually simpler than the more complex

models and since each piece can be constructed and examined in turn. Some programs therefore proceed by taking the sequence–structure alignment, removing all regions where there are insertions and deletions, and reconstructing loops and side chains against the fixed template of the remaining atoms. Some methods may also allow all parts of the template structure to adjust to the changes in sequence and insertions and deletions. This usually takes the form of a Monte Carlo or molecular dynamics simulation [18]. A global model, as described above, rebuilds a structure according to constraints derived from the known template structure or structures. This is in contrast to stepwise models that proceed essentially by replacing parts of the template structure and perhaps perturbing the structure.

Many computer programs for homology modeling are developed to solve a single problem – such as loop or side-chain building, and may not be set up to allow all atoms of the protein to adjust or to model many components simultaneously. In many cases these methods have been tested by using simplified modeling situations with the newly developed software. Such examples include experiments with removing and rebuilding loops onto single protein structures, and stripping and rebuilding all side chains. Therefore in the next sections we review some of the work in these two areas.

### 5.3.2

#### **Loop modeling**

##### 5.3.2.1 **Input information**

In stepwise construction methods, backbone segments which differ in length between the template and target (according to the sequence alignment) need to be rebuilt. In some situations, even when the sequence length of a coil segment is maintained, it may be necessary to consider alternative conformations to accommodate larger side chains or residues with differing backbone conformational requirements, Gly  $\leftrightarrow$  non-Gly, or Pro  $\leftrightarrow$  non-Pro mutations (see Appendix). Most such loop construction methods have been tested only on native structures from which the loop to be built has been removed. But the reality in homology modeling is more complicated, requiring several choices to be made in building the complete structure. These include how much of the template structure to remove before loop building; whether to model all side-chains of the core before rebuilding the loops; and whether to rebuild multiple loops simultaneously or serially.

Deciding how much of the template structure to remove before loop building depends on examination of the sequence alignment and the template structure. Sequence alignments with insertions and deletions are usually not unambiguous. Most sequence alignment methods ignorant of

structure will not juxtapose a gap in one sequence immediately adjacent to a gap in another sequence. That is, they will produce an alignment that looks like this alignment:

```
AGVEPMENYKLS
SG---LDDFKLT
```

rather than like this one:

```
AGVEPMEN---YKLS
SGL-----LDDFKLT
```

However, the latter alignment is probably more realistic [100], indicating that a 5 amino acid loop in the first sequence and structure is to be replaced with a 3 amino acid loop in the second sequence. The customary practice is to remove the whole segment between two conserved secondary structures units. Even with this practice, ambiguity remains, since the ends of secondary structures, especially  $\alpha$ -helices, are not well determined. If loop building methods were accurate, then removing more of the segment would be a good idea. But long loops (longer than 6 amino acids) are difficult to rebuild accurately, and hence there is cause to preserve as much of the starting structure as possible. Once the backbone has been borrowed from the template in stepwise modeling, one has to decide the order of building the core side chains, the backbone of loops to be built, and their side chains. They may be built sequentially, or allowed to vary simultaneously. Side chains from the core may guide the building of the loop, but at the same time may hinder correct placement. It is certainly the case that in the final structure there must be a reasonably low-energy conformation that can accommodate all loops and side chains simultaneously. Different authors have made different choices, and there has been little attempt to try vary the procedure while keeping the search algorithm and potential energy function used [74, 101–105] fixed.

### 5.3.2.2 Loop conformational analysis

Loop structure prediction is always based in one way or another on an understanding of loop conformations in experimentally determined structures. Loop conformational analysis has been performed on a number of levels, ranging from classification of loops into a number of distinct types to statistical analysis of backbone dihedral angles. Loop classification schemes have usually been restricted to loops of a particular size range: short loops of 1–4 residues, medium loops of 5–8 residues, and long loops of 9 residues or longer.

Thornton and coworkers have classified  $\beta$ -turns, which are short loops of 2–5 amino acids that connect two anti-parallel  $\beta$ -sheet strands [106–109]. These loops occur in a limited number of conformations that depend on the sequence of the loop, especially on the presence of glycine and proline residues at specific positions. The backbone conformation can be characterized by the conformations of each amino acid in terms of regions of the Ramachandran map occupied (usually defined as  $\alpha_R$ ,  $\beta_P$ ,  $\beta_E$ ,  $\gamma_R$ ,  $\alpha_L$ , and  $\gamma_L$ ; see Appendix) [108]. Usually one or more positions in the loops requires an  $\alpha_L$  conformation and therefore a glycine, asparagine, or aspartic acid residue. One useful aspect of this analysis is that if a residue varies at certain positions or there are short insertions at certain positions, the effect on the loop can be predicted [109] since the number of possibilities for each length class is small. The program BTPRED is available (see Table 5.6) to predict the locations of specific types of  $\beta$ -turns from protein sequences and secondary structure predictions derived from other programs [110]. Single amino acid changes tend to maintain the loop conformations, except when Pro residues substitute for residues with  $\phi > 0^\circ$  (see Appendix to this Chapter), while insertions change the class of the loop.

In recent years with a larger number of structures available, medium length loops have also been classified [111–117] by their patterns of backbone conformation residue by residue ( $\alpha_R$ ,  $\beta_P$ , etc.). A number of regularly occurring classes have been found, depending on length, type of secondary structure being connected, and sequence. These classes cover many but by no means all of the loops seen in non- $\beta$  turn contexts. The work of Oliva et al. [114] is probably the most thorough classification to date.

Longer loops (>8 amino acids) have been investigated by Martin et al. [118] and Ring et al. [119]. Martin et al. found that long loops fall into 2 classes, those that connect spatially adjacent secondary structures and those that connect secondary structures separated by some distance. Ring et al. provided a useful classification of longer loops as either strap (long extended loops),  $\Omega$  loops (similar to those described by Leszczynski [113]) which resemble the Greek letter, and  $\zeta$  loops, which are non-planar and have a zigzag appearance. The different loop types were found to have different distributions of virtual  $C\alpha$ - $C\alpha$ - $C\alpha$ - $C\alpha$  dihedrals to accommodate their shapes.

Swindells et al. [120] have calculated the intrinsic  $\phi$ ,  $\psi$  propensities of the 20 amino acids from the coil regions of 85 protein structures. The distribution for coil regions is quite different than for the regular secondary structure regions, with a large increase in  $\beta_P$  and  $\alpha_L$  conformations and much more diverse conformations in the  $\beta_E$  and  $\alpha_R$  regions. Their results also indicate that the 18 non-Gly, Pro amino acid type are in fact quite different from each other in terms of their Ramachandran distributions, despite the fact that they are usually treated as identically distributed in prediction methods [95, 121]. Their analysis was divided into the main broad

regions of the Ramachandran map, ignoring the  $\alpha_L$  region. The results are intriguing, in that the probability distributions are distinct enough even when calculated from a relatively small protein dataset. See the Appendix to this Chapter for examples.

### 5.3.2.3 Loop prediction methods

Loop prediction methods can be analyzed for a number of important factors in determining their usefulness: 1) method of backbone construction; 2) what range of lengths are possible; 3) how widely is the conformational space searched; 4) how are side chains added; 5) how are the conformations scored (i.e., the potential energy function); and 6) how much has the method been tested (length, number, self/non-self). We summarize a large number of published methods in Table 5.2. Only a subset of these methods are available as programs that can be downloaded or as web-servers that can perform calculations on input structures. This is indicated in Table 5.2, and the addresses for available programs are listed in Table 5.6.

#### 5.3.2.3.1 Database methods

The most common approach to loop modeling involves using “spare parts” from other (unrelated) protein structures [11, 13, 119, 122–133]. These database methods begin by measuring the orientation and separation of the backbone segments flanking the region to be modeled, and then search the PDB for segments of the same length which span a region of similar size and orientation. This work was pioneered by Jones and Thirup [122]. They defined a procedure in which  $C\alpha$ - $C\alpha$  distances were measured among six residues, three on either side of a backbone segment to be constructed. These 15  $C\alpha$ - $C\alpha$  distances were used to search structures in the Protein Databank for segments with similar  $C\alpha$ - $C\alpha$  distances and the appropriate number of intervening residues. Other authors have used the same method for locating potential database candidates for the loop to be constructed [124, 127, 128, 130].

In recent years, as the size of the PDB has increased, database methods have continued to attract attention. With a larger database, recurring structural motifs have been classified for loop structures [111, 114, 116, 129, 134], including their sequence dependence. Database methods have been applied only for loops of up to 8 residues. Fidelis et al. [126] found that for loops of length greater than 7 there is not likely to be a segment in the PDB that corresponds to the correct loop conformations for the 58 protein they looked at. Some authors report that when the database contains a loop of similar structure to the target to be modeled, their methods perform well, with RMSD values around 1 Å or better. Otherwise they tend to fail [124, 127, 130].

Although many methods have been published, they have usually only been tested on a small number of loops, and then usually in the context of rebuilding loops onto their own backbones, rather than in the process of homology modeling. These numbers are listed in Table 5.2. One exception to the small numbers of tests is the work of Fechteler et al. [128], who developed a database method (implemented in the program BRAGI) and tested it on 71 insertion and 74 deletion regions of 1–3 amino acids in length. Another is the work of Rufino et al., who used their loop classification scheme predictively on 1785 loops [129]. Unfortunately, they found their database not particularly successful in making predictions. Their database could only be used to make a prediction in 63% of cases, and only 54% of these predictions were considered correct.

#### 5.3.2.3.2 Construction methods

The main alternative to database methods is construction of loops by random or exhaustive search mechanisms. Moult and James [135] early on used a systematic search to predict loop conformations up to 6 residues long. They pioneered several useful concepts in loop modeling by construction: the use of a limited number of  $\phi, \psi$  pairs for construction; construction from each end of the loop simultaneously; discarding conformations of partial loops that cannot span the remaining distance with those residues left to be modeled; using side-chain clashes to reject partial loop conformations; and the use of electrostatic and hydrophobic free energy terms in evaluating predicted loops. Their method successfully predicted the structures of two loops in trypsin. The CONGEN program of Brucoleri and Karplus [18, 95, 136, 137] is also based on an exhaustive search algorithm, this time based on an evenly spaced grid of backbone conformations (either 15° or 30°) in accessible regions of the Ramachandran map. Gly and Pro are treated as separate classes, while the remaining 18 amino acids are treated identically in terms of their backbone energies and allowed range. Amino acids are built in turn, with partial conformations discarded if it will be impossible to close the loop with the remaining amino acids to be built. Energies are evaluated with the CHARMM polar-hydrogen force field [138]. Side chains are built by exhaustive search on 30° increments of side-chain dihedrals, as each residue is built. Since the method has been tested only on a small number of immunoglobulin CDR loops, it is difficult to tell how generally successful the algorithm is likely to be.

A particularly interesting loop construction algorithm is the scaling-relaxation method of Zheng et al. [139–141]. In this method, a full segment is sampled and its end-to-end distance is measured. If this distance is longer than the segment needs to be, then the segment is scaled in size so that it fits the end-to-end distance of the protein anchors. This results in very short bond distances, and unphysical connections to the anchors. From

Tab. 5.2  
Loop methods

<b>Authors</b>	<b>Availability</b>	<b>Side chains</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Abagyan [147, 149, 292]	ICM (\$3250 per year)	Searched with Monte Carlo probability function	Molecular mechanics + solvation terms	Construction: Internal coordinate Monte Carlo from biased probability distribution function for dihedrals	5 CASP2 deletion loops of 4–5 aa and 2 CASP2 insertion loops (10,13 aa)	Deletion loops 1.19–2.55 Å RMSD; Insertion loops 8.1, 11.4 Å RMSD
Brucoleri, Karplus [95, 136, 201]	CONGEN	Generated on grid of $\chi$ angles	CHARMM	Construction: Grid-based search + closure with Go-Scheraga + minimization with CHARMM; unproductive branches are truncated	5 self-proteins, 12 loop, 5–9 aa [95] 12 Ig loops [136]	Ig loops: 0.6–2.6 Å for backbone atoms
Carlaacci, Chou, Englander [293, 294]		Part of simulation	ECEPP/2; vdW + electrostatic, hydrogen bonds, torsional potential; $\epsilon = 2$ ; united-atom + polar hydrogens; harmonic force on N, C $\alpha$ , C, O atoms of terminal residues on x-ray positions	Construction: MC/ simulated annealing; fixed template; fixed bond lengths and angles; random translations and rotations of modeled segment + random changes in dihedrals in simulated annealing	helix, sheet, loops 5–9 aa of BPTI only	5 res loop = 1.0 Å backbone rms; 7 aa loop = 1.6 Å backbone rms; 9 aa loop = 1.87 Å backbone rms



Collura, Higo, Robson, Garnier [295–297]	Part of simulation	Robson-Platt potential [298]; vdW + electrostatic; torsions on side chains; $r$ -dependent dielectric; united-atom + polar hydrogens	Construction: MC/ simulated annealing; increasing weight of different energy terms non-uniformly; anchors fixed; initial conformation is extended peptide with harmonic potential centered on x-ray positions applied to 4 backbone atoms at each end; rigid geometry except dihedrals	8 self-protein loops in Ig, BPTI, trypsin, 6–8 aa	RMS 1 Å for backbone; more complicated solvent models not necessarily helpful
Dudek, Palmer, Scheraga [299–301]	Added after backbone deformation + minimization	ECEPP + hydration function	Construction: Random deformation (in acceptable $\phi/\psi$ regions spec. for each aa) + minimization + add side chains + minimization	5 BPTI self-loops 1 BT self-loop, all length 7	Depends on knowing initial structure. Low RMS 0.16, 0.27, 0.91, 1.15, 0.32, 0.89. Deformations are random $\phi/\psi$ pairs for 3 residues and analytic calculation of remaining $\phi/\psi$ to close loop

Tab. 5.2 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Side chains</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Janardhan, Vajda [102]	Can be implemented in CONGEN	Built with CONGEN	CHARMM + surface-solvation term + side-chain entropy term	Construction: CONGEN: systematic search	6 CASP1 loops	Evaluating free energies for CASP1 predictor loops + CONGEN side chains; importance of internal energy terms; side-chain entropy in loop predictions
Moult, James, Fidelis [126, 135]		Rotamer library, complete search	Electrostatic term, image-charge solvation term, van der Waals, hydrophobic term	Construction: by 11 $\phi\psi$ pairs; truncate unproductive branches; build many copies from each end and link; allow closer than normal contacts	2SGT loops based on bovine trypsin, 4, 5 residues; 11 homology loops in 4 proteins; 4–6 aa in length	Hydrophobic term very useful; database methods are not adequate even for short loops
Rao, Teeter [148]	Can be implemented in XPLOR	Part of MD simulation	AMBER $\epsilon = 4f$ ; counterions present	Construction: AMBER 50 ps simulation	One 7-residue loop in purothionin built from the corresponding 7-residue loop in crambin	Successfully reproduced purothionin loop with MD and simulated and annealing

Rapp, Friesner [104]	Can be implemented in MacroModel, (\$995)	Added with RSA of Shenkin, rotamer based, after initial generated loops	AMBER, AMBER94 + Born solvation implemented in MacroModel	Construction: Initial loops generated randomly with $\phi/\psi$ library. Loops that fit are saved. 10 000 step substructure minimization; then MD and simulated annealing; minimization	2 self-loops, 8 and 12 aa from ribonuclease A	r-dependent dielectric should fail badly; 12 aa loop at 0.81 Å; AMBER94 needed, not AMBER united atom model
Ring, Cohen [119, 302]	Drawbridge	Isoteric placement + Ponder & Richards rotamers	Simple steric clash check	Construction: genetic algorithm used to search among statistical distribution of tetrapeptide conformers; TWEAK algorithm used to refine model [143, 144]	4 CDR loops in antibody HY-HEL5	Better results than other methods on this antibody
Shenkin, Fine, Wang, Yarmush, Levinthal [143, 144]	Implemented in InsightII as "TWEAK"	None in early steps; added for low energy loops and searched on 10° grid, then minimized with PAKGGRAF	DISCOVER, CHARMM	Construction: Random-structure is constrained by Lagrange multipliers to fit the Jones-Thirup base distances; max change in dihedrals = 10° at each step, until convergence; followed by MD/min or min.	CDR's of MCP603	Success on short loops; multiple minima for longer CDR; need to add side chains to restrict some of the minima

Tab. 5.2 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Side chains</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Sowdhamini, Ramakrishnan, Balaram [145]		Added after search and disulfide bond formation, minimized; Janin rotamers	MODIP force field	Construction: Uniform random $\phi/\psi$ (C, P, non-GP) in allowed regions; if can form disulfide, it is made, followed by energy minimization	Disulfide bonded structures, 15 aa	Disulfide loop predictions: conotoxin, endothelin, 3–5 A rms; generate 350 000 conformations to find loops for 15 aa
Sudarsanam, Srinivasan [303, 304]	None		Simple steric checks.	Construction: Random draws from statistical distribution of $\psi_i-\phi_{i+1}$ for all 400 possible pairs of aa types, further subdivided by neighboring aa types (resulting in tetramer distributions). 10 000 conformers generated and clustered and those that fit anchors used as prediction	7 peptides, 6–20 aa in length	Various conformations can be generated, including helix and sheet.

Zheng, Delisi, Kyle [139–141]	Can be implemented in CHARMM and CONGEN	Included in simulation	CHARMM, no solvation; $\epsilon = 4\epsilon$	Construction: Scaling-relaxation method in CONGEN; reduce random loops to fit anchors and then scale to size gradually with ABNR minimization; random loops from $\phi/\psi$ distribution Database: [114] like Jones-Thirup based on C $\alpha$ positions; followed by annealing in Quanta	8 self-loops from 6 proteins, 7 aa each	Lowest E is lowest RMS 7/8 times; Zheng93 0.83–1.44 Å RMS bb Zheng96 has 0.38 Å
Bates, Oliva, Sternberg [114, 131, 305]	Summers procedure to construct similar side chains; otherwise McGregor ss-dep		CHARMM		9 CASP2 loops, 2–9 aa in length	CASP2: best loop may be most probable, not one with lowest energy because of secondary structure shifts
Fechteler, Lessel, Schomburg [128, 132, 134]	BRAGI	Not predicted	Contact distances only	Database: clustered fragments to reduce library size [134]; fit by comparison of internal distance C $\alpha$ matrices	71 insertions, 74 deletions, 1–8 aa	Good results on short indel regions
Greer [11]	Maximal overlap with old side chain		Hagler potential	Database: Alignment of multiple structures related to target to define variable regions (VR's); database fragments aligned to C $\alpha$ steric check, build side chains with overlap, restrained minimization	No validation	No comparison to known structures

Tab. 5.2 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Side chains</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Koehl, Delarue [127]	Homology program	handled with Tuffery library and SCMF search	van der Waals only	Database: Database of 81 proteins scanned for Jones-Thirup distances; minimized first; mean field theory	8 gaps in BPTI, each 4 aa; same 8 based on homology; 6 other non-self loops in homo.	Loop closure: 0.28–1.8 Å RMS; 67% $\chi_1$ (as in sc SCMF); 0.19–0.90 for homology loops
Pellequer, Chen [306]	Can be implemented in CHARMM	Tuffery rotamer library [165]	CHARMM + finite difference Poisson Boltzmann electrostatics	Database: Placement from database followed by energy minimization; selection based on free energy cycle calculation with CHARMM + Poisson-Boltzmann electrostatics	6 CDR loops from 1 antibody	Gas phase energy most discriminative; solvation free energy helped in some cases.
Rooman, Wodak, Thornton [115]		None	None	Database: Sequence pattern to predict loops	1464 loops	Sequence alone in loop prediction is not very powerful.

Rufino, Donate, Canard, Blundell [111, 129]	Sloop	Not included	None. Loops chosen by sequence and anchor positions in comparison to database loops.	Database: by class of loop, 1–8 aa long, based on length, secondary structure, vectors, sequence compared to class [111]	1785 loops	Predictions for loops that can be matched to a class in database: under lax criteria, predictions can be made for 63% of loops, of which 54% are correct.
Samudrala, Moulton [198, 199]	RAMP	Side-chain rotamer sampled and evaluated with database potential	Interatomic potential from database	Database: Graph theory and clique finding among backbone segments derived from PDB	22 CASP2 loops in 3 proteins; 4 CDR loops in antibody	Database method remains limited; improvement over CASP1
Shepherd, Gorse, Thornton [110]	BTPRED	None	None	Database: Prediction of $\beta$ -turn loop type by sequence in comparison to conformational analysis of $\beta$ turns in structures with a neural network	3359 $\beta$ turns	Correct prediction of turn type in 47/59 cases (80%).

Tab. 5.2 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Side chains</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Summers, Karplus [124]		Isosteric replacements	CHARMM, $r$ -dependent dielectric	Database: uses Jones-Thrup 15 C $\alpha$ -C $\alpha$ distances of 3 residues on either side of loop; 31 protein database; constrained energy minimization of fragments Database: Jones-Thrup distances + C $\beta$ atoms	18 non-self loops, 0–4 aa in length	2/3 of loops were well-modeled
van Vlijmen, Karplus [130]		Reduced side-chain model of Levitt; CB-only in minimization of initial loops; later bbdep rotamer library	CHARMM		18 self-loops	8/18 up to 9 aa < 1.07 Å; 17/18 had correct loop in top 3 conformations
Wilnot and Thomson [107, 108]		None	None	Database: Prediction of $\beta$ -turn loop type by sequence in comparison to conformational analysis of $\beta$ turns in structures	59 loops	Correct prediction of turn type in 47/59 cases (80%).



Wojcik, Mornon, Chomilier [116, 133]	No side chains	CHARMM (X-PLOR)	Database: Classification database of 3–8 aa loops [116]; choice determined by sequence + distance of ends; let anchors rotate $\phi\psi$ minimize to close loops; minimization with X-PLOR	3–8 aa in length; 13 CASP3 loops	0.5–2.8 Å backbone RMS for CASP3 loops
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Methods for loop prediction are listed. When publicly available programs can be identified, they have been listed by name. Some methods have been implemented in publicly available programs, but no specific software or scripts are available.

Abbreviations used in table: MC – Monte Carlo; aa – amino acid; vdW – van der Waals potential; Ig – immunoglobulin or antibody; CDR – complementarity-determining regions in antibodies; RMS – root-mean-square deviation;  $r$ -dependent dielectric – distance-dependent dielectric constant;  $\epsilon$  – dielectric constant; MD – molecular dynamics simulation; self-loops – prediction of loops performed by removing loops from template structure and predicting their conformation with template sequence; bbdep – backbone-dependent rotamer library; SCMF – self-consistent mean field; PDB – Protein Data Bank; Jones-Thirup distances – interatomic distances of 3 C $\alpha$  atoms on either side of loop to be modeled.

there, energy minimization is performed on the loop, slowly relaxing the scaling constant, until the loop is scaled back to full size. The method is efficient, since many loops can be generated which span the anchors, closing the loop with reasonable dihedral angles. Again, this method has been tested on only a few proteins, mostly immunoglobulins. Using the same methodology, Rosenbach and Rosenfeld have addressed the important issue of simultaneously modeling two or more loops in proteins [142]. In one test case, they found it advantageous to model two loops simultaneously, so that each felt the presence of the other in energy calculations

Other methods have built chains by sampling Ramachandran conformations randomly, keeping partial segments as long as they can complete the loop with the remaining residues to be built [143–145]. For longer loops, these methods seem to be much more promising, since they spend less time in unlikely conformations searched in the grid method. Many of these methods are based on Monte Carlo or molecular dynamics simulations with simulated annealing to generate many conformations which can then be energy minimized and tested with some energy function to choose the lowest energy conformation for prediction [143, 144, 146–148]. Several authors have developed Monte Carlo methods that draw  $\phi, \psi$  values from probability distributions derived from the PDB [149, 150] for other purposes, such as NMR structure refinement and *ab initio* peptide folding. These methods are promising because they are faster than exhaustive calculations, but have not been designed or tested on loop generation.

One important aspect in the development of a prediction method based on random (or exhaustive) construction of backbone conformations is the free energy function used to discriminate among those conformations which successfully bridge the anchors (see Table 5.2). Janardhan and Vajda [102] have found that a free energy function including a molecular mechanics term for the conformational energy, and a hydrophobic surface free energy term for the solvation is able to discriminate between decoy loops and real loops and to locate the correct conformation. Rapp and Friesner [104] reported recently that the AMBER94 force field and a generalized Born solvation model coupled with molecular dynamics and Monte Carlo simulations was able to regenerate conformations close to the crystal structure for one 8 and one 12 amino acid loop segment of ribonuclease. Their energy function was able to discriminate between good and poor conformations.

#### 5.3.2.4 Available programs

Apart from complete modeling packages (discussed below), only a small number of programs are freely and publicly available from the many methods that have been published (see Table 5.6). Among database methods, this includes only Swiss-PDBViewer [151], and BRAGI which implements the method of Fichteler et al. [128]. Several loop databases are available. Among

construction methods, CONGEN is currently available [95]. CODA runs both a database and an *ab initio* construction method [152].

### 5.3.3

#### Side-chain modeling

##### 5.3.3.1 Input information

Side-chain modeling is a crucial step in predicting protein structure by homology, since side-chain identities and conformations determine the specificity differences in enzyme active sites and protein binding sites. The problem has been described as “solved” [153], although new methods [154] or improvements on older ones [94] continue to be published. Some side-chain prediction methods stand on their own, and are meant to be used with a fixed backbone conformation and sequence to be modeled given as input. Other methods have been developed in the context of general homology modeling methods, including the prediction of insertion-deletion regions. Even when using general modeling procedures, such as MODELLER, it may be worthwhile subsequently to apply a side-chain modeling step with other programs optimized for this purpose. This is especially the case when side-chain conformations may be of great importance to interpretation of the model. It is also often the case that insertion-deletion regions are far away from the site of interest, and loop modeling may be dispensed with. Indeed, significant alterations of the backbone of the template, if they are not closer to the target to be modeled (if it were known) than the template, may in fact result in poorer side-chain modeling than if no loop modeling were performed. As described above, the choice of the template may depend not only on sequence identity but also on the absence of insertions and deletions near the site of interest. If this is successful, side-chain modeling rises in importance in relation to loop prediction.

Side-chain prediction methods described in detail in the literature have a long history, although only a small number of programs are currently publicly available (see Tables 5.3 and 5.6). Nearly all assume a fixed backbone, which may be from a homologous protein of the structure to be modeled, or may be the actual X-ray backbone coordinates of the protein to be modeled. Many methods have in fact only been tested by replacing side chains onto backbones taken from the actual three-dimensional coordinates of the proteins being modeled (“self-backbone predictions”). Nevertheless, these methods can be used for homology modeling by substituting the target sequence onto the template backbone. When a protein is modeled from a known structure, information on the conformation of some side chains may be taken from the template. This is most frequently the case when the template and target residue are identical, in which case the template residue’s Cartesian coordinates may be used. These may be kept fixed as the other side chains are placed and optimized, or they may be used only as a

starting conformation and optimized with all other side chains. Only a small number of methods use information about non-identical side chains borrowed from the template. For instance, Phe  $\leftrightarrow$  Tyr substitutions only require the building or removal of a hydroxyl group while Asn  $\leftrightarrow$  Asp substitutions require changing one of  $\delta$  atoms from NH2 to O or vice versa. Summers and Karplus [155, 156] used a more detailed substitution scheme, where for instance the  $\chi_1$  angle of very different side-chain types (e.g., Lys  $\leftrightarrow$  Phe) might be used in building side chains (see Appendix for definition of side-chain dihedral angles). In the long run, this is probably not advantageous, since the conformational preferences of non-similar side-chain types may be quite different from each other.

### 5.3.3.2 Rotamers and rotamer libraries

Nearly all side-chain prediction methods depend on the concept of side-chain *rotamers*. From conformational analysis of organic molecules, it was predicted long ago [157, 158] that protein side chains should attain a limited number of conformations because of steric and dihedral strain within each side chain and between the side chain and the backbone (dihedral strain occurs because of Pauli exclusion between bonding molecular orbitals in eclipsed positions) [159]. For  $sp^3$ - $sp^3$  hybridized bonds, the energy minima for the dihedral are at the staggered positions that minimize dihedral strain at approximately  $60^\circ$ ,  $180^\circ$ , and  $-60^\circ$ . For  $sp^3$ - $sp^2$  bonds, the minima are usually narrowly distributed around  $+90^\circ$  or  $-90^\circ$  for aromatics and widely distributed around  $0^\circ$  or  $180^\circ$  for carboxylates and amides (e.g., Asn/Asp  $\chi_2$  and Glu/Gln  $\chi_3$ ).

As crystal structures of proteins have been solved in increasing numbers, a variety of rotamer libraries have been compiled with increasing amounts of detail and greater statistical soundness; that is, with more structures at higher resolution [160–169]. The earliest rotamer libraries were based on a small number of structures [160–163]. Even the widely used Ponder & Richards library was based on only 19 structures, including only 16 methionines [163]. The most recent libraries are based on over 600 structures with resolution of 1.8 Å or better and mutual sequence identity less than 50% between any two chains used.

Most rotamer libraries are backbone-conformation-independent. In these libraries, the dihedral angles for side chains are averaged over all side chains of a given type and rotamer class, regardless of the local backbone conformation or secondary structure. These libraries include two in common use in side-chain conformation prediction methods, that of Ponder & Richards and that of Tuffery et al. [165]. It should be noted that the Ponder-Richards library is based on a very small sample of proteins and should *not* be used for conformation prediction (which was not its intended use anyway). The Tuffery library is based on 53 structures, which is also a very small sample compared to the PDB now available. Kono and Doi also published a rotamer

library based on a cluster analysis of 103 structures [170]. Richardson et al. have compiled rotamers recently from proteins with resolution better than 1.7 Å (compared to the more common 2.0 Å resolution cutoff), while excluding residues whose conformation is likely to be poorly determined, for example, partially disordered residues (high X-ray temperature factors) and those with unfavorable steric overlaps. This winnowing process leaves a smaller sample but one with more tightly clustered dihedral values [171].

Several libraries have been proposed that are dependent on the conformation of the local backbone [164, 166–169]. McGregor et al. [164] and Schrauber et al. [167] compiled rotamer probabilities and dihedral angle averages in different secondary structures. To my knowledge these libraries are not used in any available side-chain conformation prediction programs. We have used Bayesian statistical methods to compile a backbone-dependent rotamer library with rotamer probabilities and average angles and standard deviations at all values of the backbone dihedral angles  $\phi$  and  $\psi$  in 10° increments [166, 168, 169]. The current version of this library is based on 699 chains with resolution better than 1.8 Å and less than 50% mutual sequence identity. The library is described in greater detail in the appendix.

### 5.3.3.3 Side-chain prediction methods

Side-chain prediction methods can be classified in terms of how they treat side-chain dihedral angles (rotamer library, grid, or continuous dihedral angle distribution), potential energy function used to evaluate proposed conformations, and search strategy. These factors are summarized in Table 5.3 for nearly all side-chain prediction methods published to date. It is also useful to know how well each method has been tested, and this information is also given in the table.

As demonstrated in Table 5.3, the potential energy functions in side-chain prediction methods have varied tremendously from simple steric exclusion terms to full molecular mechanics potentials. In most cases, the potential energy function is a standard Lennard-Jones potential:

$$E(r) = 4\epsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - B \left( \frac{\sigma}{r} \right)^6 \right] \quad (5.1)$$

In this equation,  $r$  is the distance between two non-bonded atoms and  $\epsilon$  and  $\sigma$  are parameters that determine the shape of the potential. This potential has a minimum at the distance  $r = 2^{1/6}\sigma$  and a well depth of  $\epsilon$ . Different values of  $\sigma$  and  $\epsilon$  may be chosen for different pairs of atom types. Some potential energy functions for side chains may also include a hydrogen bond term. Depending on the potential parameters, these potentials may not accurately model the relative energies of rotamers for each side-chain type that are determined from local interactions within each side chain and between the side chain and the local backbone. For instance, in molecular

Tab. 5.3  
Side-chain methods

Authors	Availability	Rotamers & Flexibility	Potential energy	Search method	Test set + criteria	Claims
Bower, Cohen, Dunbrack [93, 94]	SCWRL	Backbone-dependent rotamer library [169]	Log(rotamer prob) + linear-truncated steric term	Combinatorial search of clashing clusters	300 high-resolution self-bb proteins; 1490 non-self proteins; whole PDB as self-bb predictions	80% $\chi_1$ correct
Chinea, Vriend [194, 284]	WHAT IF	Position specific rotamers extracted from PDB, 5–7 aa	Packing + HB + vdw	Ordered placement by entropy	15 non-self proteins	No statistical results given
Desmet, Lasters, DeMaeyer, Hazes [181, 182, 184, 307]		Backbone-independent rotamer library: 437 rotamers; DeMaeyer = Ponder & Richards [163] + Schrauber [167] + $2 \times 10^\circ$ steps for Aro $\chi_1 \times 2 \times 20^\circ$ Aro $\chi_2$	Force field based on CHARMM implemented in BRUGEL program	Dead-end elimination	2 self-bb proteins [181]; 19 self-bb proteins [307]	72% $\chi_1$ correct ~70% in [307]
Dunbrack & Karplus [166]	Can be implemented in CHARMM	Backbone-dependent rotamer library	CHARMM force field with no electrostatics	Placement with most probable backbone-dependent rotamer, followed by energy minimization of clashing pairs of residues	6 self-bb proteins	82% $\chi_1$ correct

Eisenmenger, Abagyan, Argos [308]	ICM (\$3250 per year)	Canonical dihedral angles, $-60^\circ$ , $180^\circ$ , $+60^\circ$	vdW + HB + torsions from ECEPP/2	Local energy search, followed by global energy minimization; ICM program	6 self:bb proteins, 2 non-self proteins	80% $\chi_1$ , 67% $\chi_{12}$ ; 60% $\chi_1$ for homology models
Holm, Sander [176]	TORSO	Tuffery library [165]	vdW	MC; side-chain/backbone energy > 10 kcal/mol removes rotamer	33 self:bb proteins, 5 non-self proteins	72% aver $\chi_1$ ; 81% buried $\chi_1$ ; non-self 54–73% correct $\chi_1$ with fixed conserved
Hwang, Liao [179]		Neural network on $20^\circ \chi$ grid	Levitt vdW truncated at 7 kcal/mol	Simulated annealing, Markov chain Monte Carlo	12 self:bb proteins	74% $\chi_1$ , 60% $\chi_{12}$
Keller et al. [309]		Tuffery library	6–12 potential of Lee and Subbiah [190]	Dead-end elimination	15 self:bb proteins	73.4% $\chi_1$ , 60% $\chi_{12}$
Koehl, Delarue [173]	Confmat	Tuffery library + modifications	vdW; disulfides, truncated to 10 kcal/mol (Levitt, 1983)	Mean field theory	30 self:bb proteins, 6 non-self proteins	72% $\chi_1$ , 62% $\chi_{12}$ , buried 82% $\chi_1$ , 72% $\chi_{12}$ buried non-self; 55–72% correct $\chi_1$ , 64–81% on correct backbone for these proteins
Kono, Doi [170, 192]	SIDEMOD	Tuffery library + additions	vdW + HB	Neural network	21 self:bb proteins + 2 non-self proteins	87.7% $\chi_1$ for buried
Laughton [310]		Database search for local environment similarity	Repulsive only vdW as $(r-r_0)^{-*2}$ , $r < r_0$	Monte Carlo search on rotamers	8 self:bb proteins, 1 non-self protein	72.8% $\chi_1$ , 79.6% $\chi_1$ buried; 64.1% on non-self test

Tab. 5.3 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Rotamers &amp; Flexibility</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Leach, Lemon [311]		Desmet/Lavery backbone-independent rotamer library [181]	AMBER force field	DEE + A* Algorithm	8 self-bb proteins	65–73% of all dihedral angles correct; Provides global minimum energy + states of system within specified energy range of global minimum
Lee, Subbiah [190, 312]	GeneMine/ Look	10° increments on $\chi$	vdW + 3-fold alkane potential	Simulated annealing	9 self-bb proteins	80–90% of interior residues; 70% of all $\chi_1$
Levitt [177]	Segmod module in Look	Segment matching to PDB	ENCAD force field, including vdW + Hbonds	Monte Carlo simulation to choose segments from PDB to model each side chain, followed by averaging; energy minimization with ENCAD	8 proteins rebuilt from C $\alpha$ positions	80% correct
Mendes et al. [154]		Tuffery214 + Gaussian distribution for flexibility around rotamer angles	GROMOS, distance-dependent dielectric	Mean-field approximation	20 self-bb proteins	87.7% $\chi_1$ , 85.7% $\chi_{12}$ ; core better than SCWRL, surface not



Ogata & Umeyama [313, 314]	FAMS	Side chains from 72 proteins used as spare parts for side chains with similar arrangements of backbone atoms in local environment, as determined from principal component analysis	None	Starting from center of mass, each residue is placed in most likely conformation in turn; a steric conflict restarts the procedure beginning with the conflicting residue; in homology modeling, $\chi$ angles are borrowed from template for geometrically similar amino acids.	15 self-bb proteins	65% $\chi_1$ on 15 self proteins
Samudrala & Moult [193]	RAMP	Canonical dihedral angles, $60^\circ$ , $180^\circ$ , $-60^\circ$	Database atom-atom distances	Top 5 interactions with mainchain saved; then by best interactions in pairs	15 self-bb proteins	75% $\chi_1$
Schiffer et al. [174]		Ponder & Richards	AMBER + surface-area a solvation term	Molten zone around each mutated side chain is minimized starting from Ponder & Richards rotamers	1 non-self bb protein	solvation important for external residues

Tab. 5.3 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Rotamers &amp; Flexibility</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Shenkin et al. [315]		Ponder & Richards, extended through $\chi_4$	$E = B - \epsilon \log P$ where B is the total number of bad contacts, and log P is log P&R rotamer probabilities	Monte Carlo simu- lated annealing	49 self:bb proteins	74% $\chi_1$
Snow, Amzel [316]		15° increments in dihedral angles	Molecular mechanics energy	“Coupled perturba- tion procedure”; Potential energy surface of side chain determined first with nearby side chains trun- cated to alanine; then model is energy minimized with all atoms present	6 side-chains in 1 non-self antibody	Side chains cor- rectly modeled
Summers, Karplus [155, 156]		Rigid rotation by 10°	CHARMM	Copy template dihedrals by maximal atomic occupancy, search by 10° for best + many checks	1 non-self protein	92% of C-terminal lobe of rhizopus- pepsin from penicillopepsin

Sutcliffe, Blundell [13]	COMPOSER	Secondary-structure dependent rotamers	None	Build from structurally aligned positions in set of homologous proteins	1 non-self protein	No results given
Tanimura, Kidera, Nakamura [317]		Backbone-independent rotamer libraries from 49 proteins	AMBER, $\epsilon = 2\tau$	Dead-end elimination	11 self-bb proteins	68% correct $\chi_1$ , 80% buried $\chi_1$ sc-mc and sc-sc are consistent with each other to favor low energy rotamers; principle of "minimal frustration" and Go's (1983) consistency principle
Tuffery, Etchebest, Hazout, Lavery [165, 191]	SMD	Tuffery library	"Flex" force field: vdW + elec ( $\epsilon = 3$ ), torsions	Sparse-matrix-driving interacting rotamers + genetic algorithm	15 self-bb proteins	percents not published
Vasquez [189]		Backbone-independent rotamer library calculated at modes of distributions (not means), similar to Tuffery; minimization in each step	vdW 12-6 and 9-6 + torsional PE for continuum searches	Mean-field, heat bath	60 self-bb proteins	80% $\chi_1$ , 70% $\chi_{12}$ ; only slight improvement for flexible rotamers

Tab. 5.3 (continued)

<b>Authors</b>	<b>Availability</b>	<b>Rotamers &amp; Flexibility</b>	<b>Potential energy</b>	<b>Search method</b>	<b>Test set + criteria</b>	<b>Claims</b>
Wendoloski, Salemmme [318]		Borrowed from similar backbone segments of 5–9 aa in PDB	Simple bump check	Placed in order of highest probability first according to segment matching + steric checks	3 self:bb proteins	59, 53, 70% $\zeta_1$ 54, 39, 59% $\zeta_2$
Wilson et al. [175]		Ponder & Richards + more Met, Asn	AMBER all atom + Eisenbeg-McLachlan solvation term	Exhaustive search for small number of residues in active site		Designed enzyme with altered specificity

Methods for side-chain prediction are listed. When publicly available programs can be identified, they have been listed by name. Some methods have been implemented in publicly available programs, but no specific software or scripts are available.

Abbreviations used in table (see also caption to Table 5.2): HB – hydrogen bonding potential energy term; self:bb – prediction of side-chain conformation on to template backbone of modeled sequence; non-self – prediction of side-chain conformation onto backbone borrowed from a homologous protein structure.

mechanics potentials, interactions between atoms connected by 3 covalent bonds (atoms  $i$  and  $i + 3$  in a chain) are not usually treated by van der Waals terms, but rather in torsion terms of the form [172],

$$E(\tau) = \sum_m K_m \cos(m\tau + a_m) \quad (5.2)$$

where the sum over  $m$  may include one-fold, two-fold, three-fold, four-fold, and six-fold cosine terms. The  $K_m$  and  $a_m$  are constants specific for each dihedral angle and each term in the sum. These torsion terms are included in some side-chain prediction methods, but ignored in others [173].

Electrostatic interactions in the form of a Coulomb potential have been included in methods that rely on full molecular mechanics potentials, usually with a distance-dependent dielectric,  $\epsilon(r) = r$ :

$$E = \frac{q_i q_j}{\epsilon(r)r} \quad (5.3)$$

Solvent interactions are also usually ignored, since these can be difficult or expensive to model properly (for exceptions see [174, 175]).

SCWRL uses an alternative strategy, based on a probabilistic potential based on the backbone-dependent rotamer library. There are two terms: the internal side-chain energy and the local side-chain-backbone interaction are modeled with an energy term proportional to  $-\ln p_{rot}$  where  $p_{rot}$  is the probability of the rotamer for the particular side-chain type and backbone conformation and a simple truncated linear steric term that models the repulsive interactions between atoms [93, 94].

Side-chain conformation prediction is a combinatorial problem, since there are on the order of  $n_{rot}^N$  possible conformations, where  $n_{rot}$  is the average number of rotamers per side chain and  $N$  is the number of side chains. But in fact the space of conformations is much smaller than that, since side chains can only interact with a small number of neighbors, and in most cases clusters of interacting side chains can be isolated and each cluster can be solved separately [93, 165]. Also, many rotamers have prohibitively large interactions with the backbone and are at the outset unlikely to be part of the final predicted conformation. These can be eliminated from the search early on.

Many standard search methods have been used in side-chain conformation prediction, including Monte Carlo simulation [176–178], simulated annealing [179], self-consistent mean field calculations [154, 173, 180], and neural networks [170]. Self-consistent mean field calculations represent each side chain as a set of conformations, each with its own probability. Each rotamer of each side chain has a certain probability,  $p(r_i)$ . The total energy is a weighted sum of the interactions with the backbone and interactions of side chains with each other:

$$E_{\text{tot}} = \sum_{i=1}^N \sum_{r_i=1}^{n_{\text{rot}}(i)} p(r_i) E_{bb}(r_i) + \sum_{i=1}^{N-1} \sum_{r_i=1}^{n_{\text{rot}}(i)} \sum_{j=i+1}^N \sum_{r_j=1}^{n_{\text{rot}}(j)} p(r_i) p(r_j) E_{sc}(r_i, r_j) \quad (5.4)$$

In this equation,  $p(r_i)$  is the density or probability of rotamer  $r_i$  of residue  $i$ ,  $E_{bb}(r_i)$  is the energy of interaction of this rotamer with the backbone, and  $E_{sc}(r_i, r_j)$  is the interaction energy (van der Waals, electrostatic) of rotamer  $r_i$  of residue  $i$  with rotamer  $r_j$  of residue  $j$ . Some initial probabilities are chosen for the  $p$ 's in Eq. (5.4), and the energies calculated. New probabilities  $p'(r_i)$  can then be calculated with a Boltzmann distribution based on the energies of each side chain and the probabilities of the previous step:

$$E(r_i) = E_{bb}(r_i) + \sum_{j=1, j \neq i}^N \sum_{r_j=1}^{n_{\text{rot}}(j)} p(r_j) E_{sc}(r_i, r_j)$$

$$p'(r_i) = \frac{\exp(-E(r_i)/kT)}{\sum_{r_i=1}^{n_{\text{rot}}(i)} \exp(-E(r_i)/kT)} \quad (5.5)$$

Alternating steps of new energies and new probabilities can be calculated from the expressions in Eq. (5.5) until the changes in probabilities and energies in each step become smaller than some tolerance.

The dead-end elimination algorithm is a method for pruning the number of rotamers used in a combinatorial search by removing rotamers that can not be part of the global minimum energy conformation [181–185]. This method can be used for any search problem that can be expressed as a sum of single-side-chain terms and pairwise interactions. Goldstein's improvement on the original DEE can be expressed as follows [183]. If the total energy for all side chains is expressed as the sum of singlet and pairwise energies,

$$E = \sum_{i=1}^N E_{bb}(r_i) + \sum_{i=1}^{N-1} \sum_{j>i}^N E_{sc}(r_i, r_j) \quad (5.6)$$

then a rotamer  $r_i$  can be eliminated from the search if there is another rotamer  $s_i$  for the same side chain that satisfies the following equation:

$$E_{bb}(r_i) - E_{bb}(s_i) + \sum_{j=1, j \neq i}^N \min_{r_j} \{E_{sc}(r_i, r_j) - E_{sc}(s_i, r_j)\} > 0 \quad (5.7)$$

In words, rotamer  $r_i$  of residue  $i$  can be eliminated from the search if another rotamer of residue  $i$ ,  $s_i$ , always has a lower interaction energy with all other side chains regardless of which rotamer is chosen for the other side chains. More powerful versions have been developed that eliminate certain

pairs of rotamers from the search [183, 185]. DEE-based methods have also proved very useful in protein design, where there is variation of residue type as well as conformation at each position of the protein [186–188].

Our side-chain prediction program SCWRL [93, 94] begins by determining the  $\phi$  and  $\psi$  angles of the input backbone conformation, and then placing side chains initially in the most favored position according to the backbone-dependent rotamer library [166, 168, 169]. The  $\chi_1$  side-chain dihedrals vary as a function of the backbone conformation for each rotamer type, in contrast to the use of backbone-independent rotamer libraries. Once the side chains are placed, steric clashes are calculated and side chains are grouped into clusters of clashing residues. If any rotamer for any side chain in one of the original clusters can produce steric overlaps with other side chains not in the clusters, then these side chains are added to the clusters. The rotamers of these side chains are searched for interactions with other side chains, and so on. Once the clusters are determined, they are solved using a branch-and-bound algorithm. It should also be noted that SCWRL uses what is essentially a dead-end elimination step, since side chains which never get added to the clusters are not searched, since they are already in their optimum rotamer with respect to the backbone. Any other rotamer choice would increase the energy, consistent with Eq. (5.7) above.

If the clusters are too large to be solved quickly, the clusters are divided by finding a residue which when removed will divide the cluster into two parts with the smallest number of combinations for solving each subcluster. For instance if a cluster of  $N$  residues can be broken into two non-interacting parts by removing one residue (called the keystone residue), then the number of combinations required to solve each group separately once for each rotamer of the keystone residue, is approximately  $n_{rot}(n_{rot}^a + n_{rot}^b)$  where  $N = a + b$  and  $n_{rot}$  is the average number of rotamers for each side chain. If  $a \approx b$ , then the number of combinations is approximately the square root of the number of combinations of the undivided cluster.

In most methods, the search is over a well-defined set of rotamers for each residue. As described above, these represent local minima on the side-chain conformational potential energy map. In several methods, however, non-rotamer positions are sampled. Summers and Karplus used CHARMM to calculate potential energy maps for side chains based on  $10^\circ$  grids [155, 156]. Dunbrack and Karplus used CHARMM to minimize the energy of rotamers from canonical starting conformations ( $-60^\circ$ ,  $180^\circ$ , and  $+60^\circ$ ) [166]. Vasquez also used energy minimization [189], Lee and Subbiah used a search over  $10^\circ$  increments in dihedral angles with a simple van der Waals term and a threefold alkane potential on side-chain dihedrals [190]. Very recently Mendes et al. [154, 180] used a mean-field method to sample from Gaussian distributions about the conformations in the rotamer library of Tuffery et al. [165]. On a test-set of 20 proteins, they claim a higher rate of prediction of core residues than SCWRL but a lower rate for surface-exposed residues.

#### 5.3.3.4 Available programs for side-chain prediction

While many methods for side-chain prediction have been presented over the years, only a small number of programs are publicly available at this time. Information on obtaining these programs is given in Table 5.6. The available programs include SMD [165, 191], Sidemod [192], Conformat [173], Torso [176], GeneMine [190], RAMP [193], and SCWRL [93, 94]. We found most of these programs easy to obtain, compile, and execute. One complication for using Torso and Conformat in modeling situations is that the programs do not maintain the residue numbering of the input file, but instead renumber the residues from 1 to  $N$ , the number of residues in the input file. This complicates the use of these programs for homology modeling. Also, only SCWRL provides a method for constraining the conformations of certain side chains, usually those that are preserved in amino acid type between the target sequence and template structure used for modeling.

Modeling programs which include backbone modeling also produce side-chain conformation predictions, although in most cases their side-chain prediction rates have not been studied. These programs include WhatIf, which uses a segment library (5–7 amino acids long) to determine a population of side-chain conformations for the central residue with a similar backbone conformation over the segment [194]. WhatIf's side-chain modeling is available on the web (see Table 5.6). MODELLER [16, 17] uses constraints based on the known side-chain conformations in the template structure. As such, it is not appropriate for modeling side chains on a backbone with no prior information on side-chain conformation. SwissModel [151] uses a backbone-independent rotamer library followed by energy minimization. ICM [147] uses a statistical potential energy function to sample side-chain conformations.

#### 5.3.4

#### Methods for complete modeling

Homology modeling is a complex process. Automated protocols that begin with a sequence and produce a complete model are few, and the resulting models should be examined with great care (as of course should all models). But these methods usually allow for (and indeed recommend) some manual intervention in the choice of template structure or structures, and in the sequence alignment. In these steps, manual intervention is likely to have important consequences. Later stages of modeling (actual building of the structure) are more easily automated, and there are not usually obvious manual adjustments to make.

There are several publicly available programs available for homology modeling that are intended to make complete models from input sequences. These are MODELLER, developed by Andrej Šali and colleagues [16, 17, 41, 195], SwissModel, developed by Manuel Peitsch and colleagues [151, 196, 197], RAMP developed by Ram Samudrala and John Moult [193,



198, 199], and COMPOSER by Blundell and colleagues [13, 14]. Some details are summarized in Table 5.4. Program availability is given in Table 5.6. We described each program in turn.

#### 5.3.5.1 Modeller

Modeller takes as input a protein sequence and a sequence alignment to the sequence(s) of known structure(s), and produces a comparative model. The program uses the input structure(s) to construct constraints on atomic distances, dihedral angles, and so forth, that when combined with statistical distributions derived from many homologous structure pairs in the Protein Data Bank, form a conditional probability distribution function for the degrees of freedom of the protein. For instance, a probability function for the backbone dihedrals of a particular residue to be built in the model can be derived by combining information in the known structure (given the alignment) and information about the amino acid type's Ramachandran distribution in the PDB. The number of restraints is very large; for a protein of 100 residues there may be as many as 20 000 restraints. The restraints are combined with the CHARMM force field to form a function to be optimized. This function is optimized using conjugate gradient minimization and molecular dynamics with simulated annealing.

Modeller provides some help with homologue identification and sequence alignment, but the full PDB should be searched outside of Modeller unless a very recent update of the program and database is obtained. Additional sequences of unknown structure can also be added to the information used by Modeller, although these must be obtained with other programs such as BLAST and FASTA. Modeller can combine the sequences and structures into a complete alignment which can then be examined using molecular graphics programs and edited manually.

#### 5.3.5.2 SwissModel

SwissModel is intended to be a complete modeling procedure accessible via a web server that accepts the sequence to be modeled, and then delivers the model by electronic mail [151, 196]. In contrast to Modeller, SwissModel follows the standard protocol of homologue identification, sequence alignment, determining the core backbone, and modeling loops and side chains. SwissModel will search a sequence database of proteins in the PDB with BLAST, and will attempt to build a model for any PDB hits with  $p$ -values less than  $10^{-5}$  and at least 30% sequence identity to the target (for a description of  $p$ -values and BLAST, see Chapter 2 of this volume). SwissModel allows for user intervention by specifying the template(s) and alignments to be used. If more than one structure is found, they will be superimposed on the template structure closest in sequence identity to the target. The structural superposition is accomplished by aligning the structures according to the initial sequence alignment, and then minimizing  $C\alpha$ - $C\alpha$  distances be-

**Tab. 5.4**  
 Programs for complete homology modeling (alignment, loops, side chains)

<b>Authors</b>	<b>Availability</b>	<b>Potential energy</b>	<b>Search method</b>
Sutcliffe, Blundell [13, 14]	COMPOSER		Averaging several structures; picking fragments from each most similar to average
Sali, Blundell, Sanchez [16, 17, 195]	MODELLER	CHARMM + constraints from statis- tical analysis	Satisfaction of spatial restraints (prob. distribution functions) by minimization and simu- lated annealing
Yang, Honig [204]	PrISM		Multiple structures are aligned, and the most appropriate template is used for each segment of the target to be built. Loops are built <i>ab</i> <i>initio</i> and side chains are built using the template or based on mainchain torsions and a neural network algorithm.
Li, Tejero, Brucoleri, Montelione [18, 19]	CONGEN + 2 set-up programs	CHARMM; $\epsilon = r$	Distance constraints derived from known structure and alignment; random subset is used as input to CONGEN simulated annealing/ restrained molecular dynamics simulations
Peitsch, Guex [151, 196]	SwissModel	GROMOS	Alignment determines core backbone, while loops are built with database method. Side chains are built using the template side chain where possible, and GROMOS minimization otherwise.
Havel, Snow [15, 210]	DISGEO/Co- nsensus (MSI, Inc.)	None	Distance geometry; constraints based on alignment and multiple structures
Samudrala, Moult [193, 198, 199, 209]	RAMP	Atom distance energy func- tion from statistical analysis	Graph theory method that selects compatible confor- mations from multiple templates, multiple database loops, and side-chain rotamers

tween the proteins. This is in contrast to methods that compare *internal* distances in one structure with internal distances in the other, such as DALI [88] and CE [89].

SwissModel determines the core backbone from the alignment of the target sequence to the template sequence(s) by averaging the structures according to their local degree of sequence identity with the target sequence. The program builds new segments of backbone for loop regions by a database scan of the PDB using anchors of 4 C $\alpha$  atoms on each end. This method is used to build only the C $\alpha$  atoms, and the backbone is completed with a search of pentapeptide segments in the PDB that fit the C $\alpha$  trace of the loop. Side chains are now built for those residues without information in the template structure by using the most common (backbone-independent) rotamer for that residue type. If a side chain can not be placed without steric overlaps, another rotamer is used. Some additional refinement is performed with energy minimization with the GROMOS [200] program.

SwissModel has been extensively tested by building over 1200 models of proteins of known structure, with template-target sequence identities ranging from 25 to 95%. It was found that 30% of models with sequence identity less than 30% could be built with backbone RMSD less than 3 Å. This rose rapidly with sequence identity, such that over 80% of models with sequence identity better than 50% could be built with RMSD's better than 3 Å. Assessments of side chain and loop modeling quality however have not been published.

#### 5.3.5.3 CONGEN with Homology-derived constraints

Montelione et al. have used CONGEN [95, 201] to produce comparative models [18, 19, 202, 203]. Their method derives distance restraints from the template structure and the sequence alignment of the target and template for atoms they consider “homologous.” These atoms include all backbone atoms and side-chain atoms of the same chemical type and hybridization state (i.e.,  $sp^3$ ,  $sp^2$ ). In the vicinity of an insertion or deletion in the alignment ( $\pm 3$  amino acids), no homologous atoms are defined so that loop regions are free to move. From the very large number of constraints generated by this procedure, a small fraction ( $< 2\%$ ) are chosen randomly and used with restrained molecular dynamics and simulated annealing in the CONGEN program. CONGEN's side-chain loop and search routines can also be used. The method has only been tested on 4 proteins [18, 19], and so it is difficult to assess the accuracy.

#### 5.3.5.4 PrISM

Yang and Honig have developed the PrISM package of programs that performs structure alignment, PDB sequence search, fold recognition, se-

quence alignment, secondary structure prediction, and homology modeling [204–208]. Its methods for homology modeling [204], in contrast to structure alignment [206–208], have not been published in detail. If multiple structures are available, PrISM aligns them and builds a composite template by selecting each secondary structure from the most appropriate template. Loop modeling is performed *ab initio* and side-chain dihedrals are taken either from the template or predicted based on mainchain torsion angles and a neural network algorithm.

#### 5.3.5.5 COMPOSER

COMPOSER by Blundell and colleagues [13, 14] exploits the use of multiple template structures for building homology models. If a target sequence is related to more than one template (of different sequence) then all templates are used to provide an “average” framework for building the structure. Modeling of loops and side-chains is accomplished by borrowing from one of the template structure where possible, and by database methods in other cases.

#### 5.3.5.6 RAMP

Samudrala and Moult described a method for “handling context sensitivity” of protein structure prediction, that is, simultaneous loop and side-chain modeling, using a graph theory method [198, 209] and an all-atom distance-dependent statistical potential energy function [199]. Their program RAMP is listed in Table 5.6.

#### 5.3.5.7 DISGEO/Consensus

Havel and coworkers [15, 210] have described a method of producing comparative models by distance geometry based on restricted ranges for certain C $\alpha$ -C $\alpha$  distances, dihedrals, and mainchain–side-chain distances. This method has become part of the InsightII package from Molecular Simulations, Inc.

## 5.4 Results

### 5.4.1 Range of targets

A very large number of homology models have been built over the years by many authors. Targets have included antibodies [211–216] and many proteins involved in human biology and medicine [19, 31, 217–256].

```

* *
SecStr : EEEE EEE EEEEEEE EEEEEEE EEEE
lps0   1:VDRQPLENYLDMYF+FGTIGTTPAQDFTV+FDTGSSNLWVSPVYCSSLACTNHNRPED: 60
hbace1 62:EMVDNLRGKSGQGYVEMTVGSPQTLN+ILVDTGSSNFAVGAAPH---PFLHRY+YQRQL:117
mbace1 62:EMVDNLRGKSGQGYVEMTVGSPQTLN+ILVDTGSSNFAVGAAPH---PFLHRY+YQRQL:117
rbace1 62:EMVDNLRGKSGQGYVEMTVGSPQTLN+ILVDTGSSNFAVGAAPH---PFLHRY+YQRQL:117
bbace1 :
sbace  :
zbace  :DMINNLKGD+SGR+GYM+QMIIGT+PGQTLN+ILVDTGSSNFAVAAAAH---PYITHY+FNRL:
hbace2 79:AMVDNLQD+SGR+GYL+EMLIGT+PPQK+QLVDTGSSNFAVAGT+PH---SYIDTY+FDTER:134
mbace2 75:AMVDNLQD+SGR+GYL+EMLIGT+PPQ+KVQ+ILVDTGSSNFAVAG+APH---SYIDTY+FDSE:130

**
SecStr : EEEE EEEEE EEEEEEEEEEE EEEEEEEEEEE HHHHH
lps0   61:SSTYQSTSE-TVSI+TYGTGSM+TGILGYD+TVQV+G---GISDTN+QIFGLSETEPGSFL+YAP:116
hbace1 118:SSTY+RDLRK-GVYV+PY+TQ+GK+WEGELG+TDLV+SI+PH-GPNV+TV-RANIAA+ITES+DKFF+INGSN:175
mbace1 118:SSTY+RDLRK-GVYV+PY+TQ+GK+WEGELG+TDLV+SI+PH-GPNV+TV-RANIAA+ITES+DKFF+INGSN:175
rbace1 118:SSTY+RDLRK-GVYV+PY+TQ+GK+WEGELG+TDLV+SI+PH-GPNV+TV-RANIAA+ITES+DKFF+INGSN:175
bbace1 :SSTY+RDLRK-GVYV+PY+TQ+GK+WEGELG+TDLV+SI+PH-GPNV+TV-RANIAA+ITES+DKF-----:
sbace  :SSSV+INS+GP+VD+IEY+TE+GF+WK+G+PLV+TDL+V+SI+PEA+GLTE+QV-RV+IV+KIT+SS+KK+FF+INGSG:
zbace  :SSTY+Q+STER-AVAV+KY+TQ+EWE+GELG+TDL+ITIP
hbace2 135:SSTY+R+SKGF-DVT+V+KY+TQ+GS+WT+GF+V+GED+L+V+TIP+K-GFN+T+SF-LV+NI+ATI+FES+EN+FF+LP+G+IK:192
mbace2 131:SSTY+H+SKGF-DVT+V+KY+TQ+GS+WT+GF+V+GED+L+V+TIP+K-GFN+SS+F-LV+NI+ATI+FES+EN+FF+LP+G+IK:188

*
SecStr : EEEE HHHHHHHH EEEEE
lps0   117:FDGILGLAY+PS+ISS--S-GATP+V+FD+N+W+QGLV+SQD+L+FS+VY+LS+ADD-----QSGS:163
hbace1 176:WEGILGLAY+AE+IAR+PDD-SLE+PF+F+DS+L+V+K+Q+TH+VP-N+IF+SL+QL+CGA+GF+PL+NQ+SE+VL+AS+VGG:233
mbace1 176:WEGILGLAY+AE+IAR+PDD-SLE+PF+F+DS+L+V+K+Q+TH+IP-N+IF+SL+QL+CGA+GF+PL+NQ+TE+AL+AS+VGG:233
rbace1 176:WEGILGLAY+AE+IAR+PDD-SLE+PF+F+DS+L+V+K+Q+TH+IP-N+IF+SL+QL+CGA+GF+PL+NQ+TE+AL+AS+VGG:233
sbace  :WQGI+IGL+G+YDEL+V+R+P+N+N+K+VF+MT+S+V+IEN+T+SV+R-NV+FS+IOL+CAA---NTM+FS+V+T+TG:
hbace2 193:WNGILGLAY+ATL+AK+PSS-SLE+T+F+DS+L+V+TQ+AN+IP-NV+FS+M+Q+CGA+GL+PV+AGS---GT+NGG:247
mbace2 189:WNGILGLAY+AA+L+AK+PSS-SLE+T+F+DS+L+V+AQ+AK+IP-D+IF+SM+Q+CGA+GL+PV+AGS---GT+NGG:243

*
SecStr : EEEE EEEE EEEEEEEEE EEE EEEEEEE
lps0   164:V+IFGG+ID+SS+Y+YT+G+SL+N+W+V+F+V+TE+GY+Q+IT+V+DS+IT+M+NGE+AI+AC---AEG+CQ+AI+VD+TG+TS:219
hbace1 234:SMI+IG+ID+HS+LY+TG+SL+W+Y+TP+IR+RE+WY+EV+II+VR+VE+ING+Q+DL+K+MD+CKE+YN+Y+DK+S+IV+DS+GT+:293
mbace1 234:SMI+IG+ID+HS+LY+TG+SL+W+Y+TP+IR+RE+WY+EV+II+VR+VE+ING+Q+DL+K+MD+CKE+YN+Y+DK+S+IV+DS+GT+:293
rbace1 234:SMI+IG+ID+HS+LY+TG+SL+W+Y+TP+IR+RE+WY+EV+II+VR+VE+ING+Q+DL+K+MD+CKE+YN+Y+DK+S+IV+DS+GT+:293
bbace1 143:--IG+ID+HS+LY+M+GS+L+W+Y+TP+IR+RE+WY+EV+II+VR+VE+ING+Q+DL+K+MD+CKE+YN+Y+DK+S+IV+DS+GT+:200
sbace  :SLV+FG+DY+DR+T--DG+T+IF+R+TR+IV+H+WY+EV+IV+LG+M+KVCRE+FN+ND+KS+IV+DS+GT+:
hbace2 248:SLV+GGI+E+PS+LY+K+GD+I+WY+TP+KE+WY+Q+IE+LL+K+LE+IG+GS+N+L+DC+RE+YN+AD+KAI+VD+SG+TT:307
mbace2 244:SLV+GGI+E+PS+LY+K+GD+I+WY+TP+KE+WY+Q+IE+LL+K+LE+IG+GS+N+L+DC+RE+YN+AD+KAI+VD+SG+TT:303
rbace2 :
IL+K+LE+IG+GS+N+L+DC+RE+YN+AD+KAI+VD+SG+TT:

+
SecStr : EEE HHHHHHHHHH EE EE EEEEE EEEE
lps0   220:LLTG+PTS+PI+ANI+QSD+I+GAS+NSD+GD-----MV+SC+SA+ISS+LP+DIV+TIN-----:260
hbace1 294:N+LR+L+P+K+K+VF+E+AA+V+KS+I+KAA+S+TE+K+FP+DG+FW+L+GE+QL+VC+W+QAG+TP+PN+IF+FP+VI+SL+YL+M+GE+V:352
mbace1 294:N+LR+L+P+K+K+VF+E+AA+V+KS+I+KAA+S+TE+K+FP+DG+FW+L+GE+QL+VC+W+QAG+TP+PN+IF+FP+VI+SL+YL+M+GE+V:352
rbace1 294:N+LR+L+P+K+K+VF+E+AA+V+KS+I+KAA+S+TE+K+FP+DG+FW+L+GE+QL+VC+W+QAG+TP+PN+IF+FP+VI+SL+YL+M+GE+V:352
bbace1 201:N+LR+L+P+K+K+VF+E+AA+V+KS+I+KAA+S+TE+K+FP+DG+FW+L+GE+QL+VC+W+QAG+TP+PN+IF+FP+VI+SL+YL+M+GE+V:259
sbace  :NL+HL+PE+K+VEN
hbace2 308:LL+RL+P+Q+K+VD+AV+VEA+V+ARS+SL+IP+EF+SD+GF+WT+GS+QL+AC+WT+NS+ET+PW+Y+FP+K+ISI+YL+RD+EN:366
mbace2 304:LL+RL+P+Q+K+VD+AV+VEA+V+ARS+SL+IP+EF+SD+GF+WT+GA+QL+AC+WT+NS+ET+PW+Y+FP+K+ISI+YL+RD+EN:362
L+RL+P+Q+K+VD+AV+VEA+V+ARS+SL+IP+EF+SD+GF+WT+GA+QL+AC+WT+NS+ET+PW+Y+FP+K+ISI+YL+RD+EN:362

```

Fig. 5.3

Multiple sequence alignment of BACE family members and human pepsin (Protein Data Bank entry 1ps0, used as the basis of homology modeling). Identical residues are shaded gray, highly conserved residues in yellow, and active-site residues in the model in contact with the peptide substrate are indicated by an asterisk (\*). BACE residue Arg 296, which is most often serine or threonine at this position in

aspartyl proteases, is shown with a plus sign (+). The secondary structure of the human pepsin template structure is given to show that gaps in the alignment between elements of secondary structure (H is  $\alpha$ -helix; E is  $\beta$ -strand). In sequence names, h = human; m = mouse; r = rat; b = bovine; z = zebrafish; s = sea squirt (*Halocynthia roretzi*). Fragment sequences of bbace1, rbace2, bbace2, and sbace were derived from EST sequences.

Two databases of homology models are available on the Internet – ModBase and SwissModel. ModBase [257], produced by Andrej Šali and colleagues at Rockefeller University, contains homology models built with the program MODELLER [16] for as many sequences in several complete genomes that can be modeled with structural homologues of better than 30% sequence identity. The models include evaluations for predicting their accuracy.

#### 5.4.2

##### **Example: amyloid precursor protein $\beta$ -secretase**

Alzheimer's disease is characterized by plaques in the brain consisting primarily of the 40–42 amino acid amyloid  $\beta$ -peptide ( $A\beta$ ) [258].  $A\beta$  derives from proteolysis of the amyloid precursor protein (APP) by the  $\beta$  and  $\gamma$  secretases to create the N and C-termini of the peptide respectively [259]. The  $\beta$ -secretase has recently been identified as a 501 amino-acid transmembrane protein by several research groups [260–263]. The enzyme, variously named BACE, memapsin2, and Asp2, is an aspartic protease related to pepsin, cathepsin D, and renin, with all the properties expected of the  $\beta$ -secretase.

Because BACE is a potential target for drug design to treat or prevent Alzheimer's disease, we recently built models of the BACE extracellular domain based on crystal structures of two aspartyl proteases. As described above, perhaps the most important step in homology modeling is to gather available experimental data on the protein's function. Even before the identification of BACE, the specificity of the  $\beta$ -secretase had already been studied extensively [264]. The sequence of APP at the protease cleavage site is EVKM-DAEF, where the protease breaks the peptide bond between M and D. These residues are labeled P4-P3-P2-P1-P1'-P2'-P3'-P4' in standard protease nomenclature. The protease also cleaves at another position within the  $A\beta$  peptide, at Glu 11, with sequence DSGY-EVHH. It is clear that there is at least some preference for a negatively charged residue at the P1' position, an unusual feature in aspartic proteases. This led us to hypothesize that the substrate binding site was very likely to contain a positively charged residue.

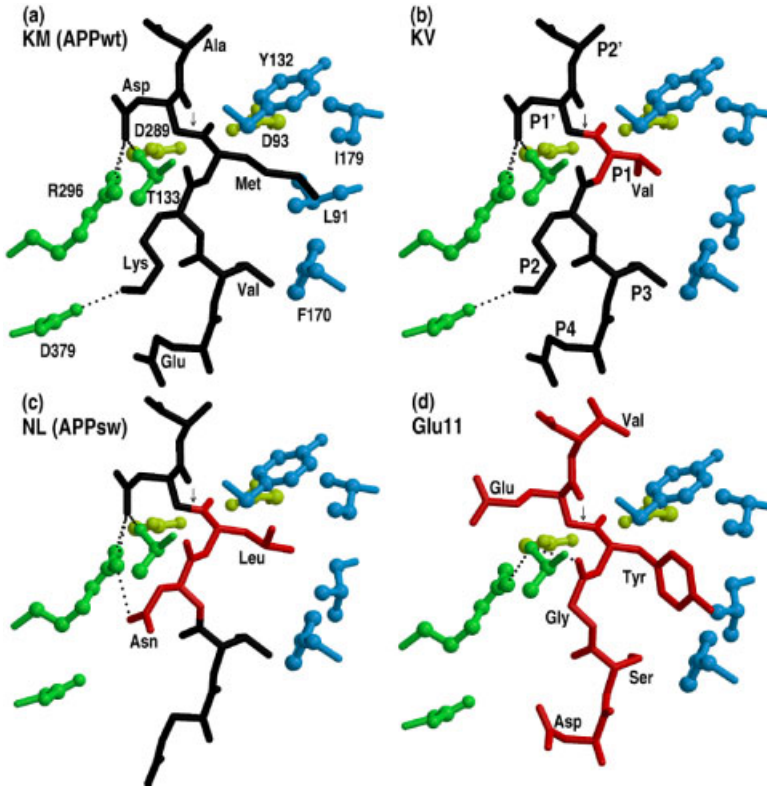
BACE also cleaves at the same location of APP with the so-called "Swedish mutation", KM  $\rightarrow$  NL at P2-P1, found in families with early-onset Alzheimer's disease. This indicates that a hydrophobic residue seems to be preferred at the P1 position and a hydrophilic residue at P2. One other piece of data was a mutant of the APP substrate of M  $\rightarrow$  V at position P1, which greatly reduced activity of BACE for the substrate [264, 265].

The second step was to gather *orthologous* sequences of BACE to define the family of sequences to which BACE belongs. We searched both the non-redundant protein sequence database at NCBI as well as the EST databases. The resulting alignment of BACE orthologues is shown in Figure 5.3

(s. p. 189). In mammals, there are two closely related sequences, now called BACE1 and BACE2, that appear to have arisen from a recent duplication. In zebrafish and seq squirrels, there is a single BACE sequence in EST databases that shares feature of both BACE1 and BACE2. We include BACE1 and BACE2 in the alignment of orthologues, even though these sequences exist in the same species, because they are very closely related and their specific functions have not yet been determined.

To build the model we used PSI-BLAST to search the non-redundant protein sequence database and to produce a position-specific scoring matrix for BACE. This matrix was used to search the Protein Data Bank to identify homologues of known structure that could be used as templates to build a model of BACE. This process identified many aspartyl proteases. After looking at the alignments, we decided to use a 2.0 Å resolution crystal structure of human pepsin in a complex with pepstatin (Protein Databank entry 1PSO) [266]. This structure had no insertions and deletions in close proximity to the enzyme active site, while most other aspartyl proteases had insertions in the loops covering the active site. We were also interested in building in a model of the substrate sequence from APP. The crystal structures of most aspartyl proteases in the PDB contain a peptide-like inhibitor in the active site. To make these small molecules inhibitors rather than substrates, the peptide bond at the cleavage site is replaced with non-hydrolysable linkages. This is almost always accomplished with an extra atom or two in the backbone of the inhibitor. The only exception to this is a reduced peptide inhibitor in a 1.8 Å crystal structure of rhizopuspepsin (PDB entry 3APR) [267]. In this inhibitor, the carbonyl group of the scissile bond has been replaced with a methylene group ( $-\text{CH}_2-$ ). We chose this substrate to model the APP substrate since the backbone of the substrate to be modeled could be borrowed directly from the substrate in the rhizopuspepsin structure. To model the interaction of the substrate with the enzyme, we superimposed the structure of the reduced peptide inhibitor onto the peptide portion of the pepstatin inhibitor in 1PSO. The backbone atoms of the peptide inhibitor before the scissile bond superimpose with the pepstatin backbone with an RMSD of 0.16 Å.

In Figure 5.4 we show the major proposed interactions of the APP substrate with the enzyme. For the wild-type peptide in Figure 5.4A with sequence EVKMDA, the P1' aspartic acid makes a salt-bridge with Arg296 of the enzyme. This is the hypothesized positive charge in the BACE active site. It is conserved in the closely related human transmembrane protein BACE2 [268], which indicates that this enzyme is likely to have similar substrate specificity to BACE. In other aspartic proteases, this residue is an Ala, Ser, Thr, Val, Leu, or Ile residue. In the model, the P2 lysine also makes a salt-bridge with the enzyme at Asp379. There are several hydrophobic contacts between the P1 methionine and enzyme residues Leu91, Tyr132, and Ile179, while the P3 valine interacts with Phe170.

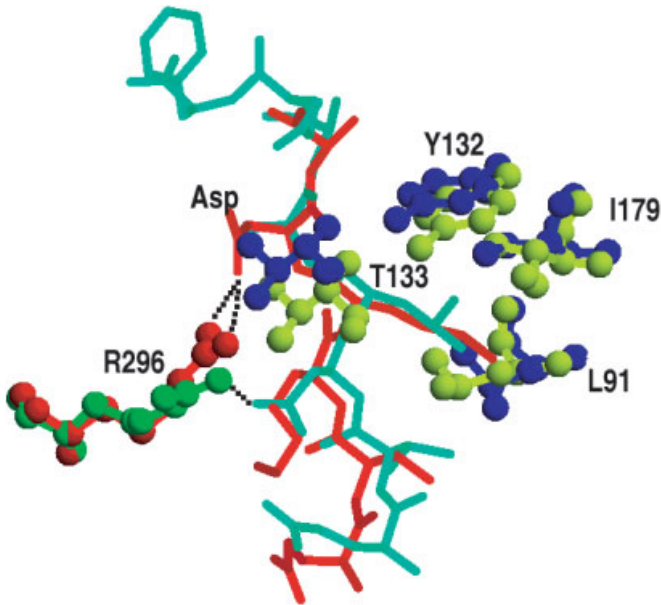


**Fig. 5.4**  
Models of  $\beta$ -secretase with APP-derived substrates. **(A)** Wild-type substrate residues 592–597 of APP (GenBank accession CAA312830.1) with sequence EVKM\*DA (P4-P3-P2-P1-P1'-P2') with cleavage site marked with \*. **(B)** Mutant substrate with Val (red) substitute for P1

Met (sequence EVKV\*DA). **(C)** Swedish mutation substrate with P2-P1 sequence of Asn-Leu (red) in place of wild-type Lys-Met (sequence EVNL\*DA). **(D)** Alternative  $\beta$ -secretase cleavage site at APP residue Glu607 (P1') (sequence DSGY\*EV). The figure was generated with MOLSCRIPT [287], Raster3D [288], and the GIMP [289].

We also modeled the mutant substrate EVKVDA shown in Figure 5.4B. The differences in interaction between the side chain and the enzyme explain why this substrate is cleaved less well by BACE [264]. While the  $C_{\gamma 1}$  of Val and  $C_{\gamma}$  of Met are in the same location ( $\chi_1 = -65^\circ$ ), the  $C_{\gamma 2}$  atom of Val contacts the catalytic Asp93  $C_{\gamma}$ ,  $O_{\delta 1}$ , and  $O_{\delta 2}$  atoms. This is likely to interfere with the catalytic activity. Also, many of the hydrophobic contacts between P1 Met and the enzyme are missing in P1 Val. In Figure 5.4C a model of the “Swedish mutation” KM  $\rightarrow$  NL at P2, P1, which is associated with early-onset Alzheimer’s disease [269], shows that the Asn can hydrogen bond to





**Fig. 5.5**  
Superposition of BACE crystal structure with homology model of BACE. The crystal structure of BACE (PDB entry 1FKN) was superimposed on the model using the MIDAS program [290]. The model peptide is shown in green stick figure and the crystal structure peptide is shown in red.

The crystal structure side-chains in contact with the peptide are shown in blue ball-and-stick figures, while the model side-chains are shown in yellow. Crystal structure Arg296 is shown in red and model structure Arg296 is shown in green, with a hydrogen bond depicted by dotted lines to Asp at P1'.

Arg296, which also hydrogen bonds to the P1' Asp. The leucine also makes good hydrophobic contacts with Leu91, Tyr132, and Ile179. In Figure 5.4D, the minor cleavage site for  $\beta$ -secretase, at Glu11 of the  $A\beta$  peptide, is modeled as the substrate. The P1' Glu is within hydrogen bonding distance of Arg296, although the hydrogen bond geometry is poor. This may explain why the Glu11 site is cleaved by BACE, but only as a minor component of BACE activity on APP.

The crystal structure of BACE was recently determined by Tang and colleagues [270]. In Figure 5.5, we show the substrate binding site from a structure alignment of the model and the crystal structure. A number of the substrate-specificity-determining residues were correctly determined both in location and conformation. In this crystal structure, an inhibitor is bound in the active site. This inhibitor has sequence EVNL-AAEF, where the L-A sequence has a non-peptide backbone. The P1' position is occupied by an

alanine rather than a negatively charged residue, as in both APP substrate positions. In our model, Arg296 forms a salt-bridge to Asp at P1'. This arginine is in nearly the exact same position in the crystal structure, but in a different rotamer and making a hydrogen bond to substrate Asn at P2. In the natural substrate, P2 is Lys and therefore very unlikely to make a hydrogen bond to Arg296. We believe that with a conformational change, Arg296 is easily able to form a salt bridge with P1' negatively charged residues and is thus one of the prime determinants of specificity of BACE for APP. It is notable that because the crystal structure did not contain the native substrate sequence this interaction of Arg296 with P1' was not discussed by the crystallographers, even though it is the only evident positively charged residue able to interact with P1' side chains.

It is clear from this example that choosing templates and modeling goals is a process that takes some care and understanding of biological function. Automated procedures are less than satisfactory for the purposes described here.

## 5.5 Strengths and limitations

The strengths of homology modeling are based on the insights provided for protein function, structure, and evolution which would not be available in the absence of an experimental structure. In many situations, a model built by homology is sufficient for interpreting a great deal of experimental information and will provide enough information for designing new experiments. Homology modeling may also provide functional information beyond the identification of homologous sequences to the target. That is, a model may serve to distinguish orthologous and paralogous relationships.

The limitations are due to decreasing accuracy as the evolutionary distance between target and template increases. Alignment becomes more uncertain, insertions and deletions more frequent, and even secondary structural units may be of different lengths, numbers, and positions in very remote homologues. Predicting the locations of secondary structure units that are not present in the template structure is a difficult problem and there has been little attention paid to it.

The limitations of homology modeling also arise when we have insufficient information to build a model for an entire protein. For instance, we may be able to model one or more domains of a multi-domain protein or a multisubunit complex, but it may not be possible to predict the relative organization of the domains or subunits within the full protein. This remains a challenge for further research. And we are of course limited by

structures present in the PDB, which are almost exclusively soluble proteins. Up to 30% of some genomes are membrane proteins, which are at present difficult to model because of the small number of membrane proteins of known structure. The recent structure of the G-protein-coupled receptor rhodopsin [271] creates an opportunity to model many of these membrane proteins more accurately than the previously available structure of bacteriorhodopsin [272, 273].

Another problem is the quality of data in sequencing and structure determination. There are substantial errors in determining protein sequences from genome sequences, either because of errors in the DNA sequence or in locating exons in eukaryotic DNA [274]. Over 50% of X-ray structures are solved at relatively low resolution, levels of greater than 2.0 Å. Despite progress in determining protein structures by NMR, these structures are of lower resolution than high quality X-ray structures. While high throughput structure determination will be of great value to modeling by homology, one concern is the quality of structure determination when the function of the proteins being determined is unknown.

## 5.6

### Validation

Validation for homology modeling is available in two distinct ways: 1) the prediction rates for each method based on the prediction of known structures given information from other structures; 2) criteria used to judge each model individually. As shown in Tables 5.2–5.4, most structure prediction method papers have included predictions of known structures, serving as test sets of their accuracy. However, in many cases the number of test cases is inadequate. It is also very easy to select test structures that behave particularly well for a given method, and many methods do not stand up to scrutiny of large test sets performed by other researchers. Test sets vary in number of test cases as well as whether predictions of loops or side chains are performed by building replacements on the template structure scaffold, or in real homology modeling situations where the loops/side-chains are built on non-self scaffolds. The realistic case is more difficult to perform in a comprehensive way, since it requires many sequence-structure alignments to provide the input information on which models are to be built. Another problem is that each method is judged using widely varying criteria, and so no head-to-head comparison is possible from the published papers. The problem of biased test sets and subsequent development of larger benchmarks has a long history in the secondary structure prediction field [275, 276], but testing of loop and side-chain methods in this way has been much more limited.

## 5.6.1

**Side-chain prediction accuracy**

To remedy this situation for protein side-chain prediction, we developed a benchmark and tested several publicly available side-chain prediction programs. In the original publications, these methods were tested on very different test sets and using varying criteria for a “correct” prediction. In some cases, the test sets are rather small. To see why, recall that the standard deviation of the mean of a sample is the population standard deviation divided by the square root of the number of data points. If we treat the fraction of correct  $\chi_1$  rotamers for a single protein as a data point, we observe that the standard deviation of the data is approximately 0.06 (6%) on a large test set. If a test set consists of 10 proteins, then the standard deviation of the mean is  $0.06/\sqrt{10} = 0.02$ . The 95% confidence interval is therefore the mean  $\pm 0.04$ , or an eight percentage point spread. Larger test sets are therefore called for.

We performed a test on 180 monomeric (at least within each asymmetric unit of the crystal structure) proteins with resolution better than 1.8 Å and less than 30% sequence identity between any two proteins in the list. The same criteria were used to assess each of the methods: a  $\chi$  angle prediction within 40° of the crystal structure conformation. Pro residues were judged correct if the  $\chi_1$  angle was within 20° of the crystal structure conformation. All residues other than Gly and Ala were included. The results for the 18 amino acid types and for all residues together are shown in Table 5.5. In this table, “bbdep” means the prediction that would be made simply by choosing the most common rotamer in the backbone-dependent rotamer library, that is, the rotamer determined by  $\phi, \psi$  for each residue. “bbind” indicates a prediction based on simply choosing the most common rotamer for each side chain in the backbone-independent rotamer library of Dunbrack. In this prediction, all non-Pro side chains are given a  $\chi_1$  dihedral of approximately  $-60^\circ$ , except for Val which is most commonly near  $180^\circ$  (trans), and Ser and Thr which are most commonly near  $+60^\circ$ . The most interesting result is that some of the methods currently available provide *worse* predictions than simply choosing the most common rotamer in the backbone-dependent rotamer library, a prediction that proceeds with simply looking up a conformation in a table. The bbdep library on its own can predict 73% of  $\chi_1$  rotamers correctly. SCWRL, Conformat, and Torso perform better than this prediction by optimizing rotamers based on interactions between side chains and between side chains and the backbone.

## 5.6.2

**The CASP meetings**

Another forum for testing homology modeling methods has been the ongoing series of CASP meetings (for “Critical Assessment of Protein Struc-

Tab. 5.5  
Side chain program comparison

Residue	Number	BBIND	BBDEP	CONFMAT	RAMP	SCWRL	SIDEMOD	SMD	TORSO
ILE	2127	75.9	87.6	91.1	54.1	91.3	89.7	66.3	90.7
VAL	2779	70.7	85.9	88.7	72.3	88.8	87.6	65.3	88.4
THR	2439	48.0	84.5	78.5	80.5	87.1	69.0	65.1	80.6
PRO	1996	60.4	76.8	51.1	*	74.8	39.2	*	52.9
PHE	1644	53.5	74.9	90.4	70.3	91.2	87.7	68.1	91.8
TYR	1626	53.0	74.2	88.0	67.3	90.0	81.6	65.0	89.8
LEU	3396	66.1	73.3	86.7	68.5	87.1	78.3	65.2	87.9
ASP	2483	50.2	71.3	79.8	62.4	81.1	58.2	57.9	70.0
HIS	895	54.3	70.8	82.8	63.4	83.1	74.4	59.5	84.1
CYS	678	57.8	70.5	85.3	60.5	83.0	78.0	81.5	78.5
TRP	668	46.8	69.9	83.9	60.5	86.1	76.1	57.9	84.0
ASN	2027	52.6	67.7	74.3	55.9	77.6	62.2	53.2	74.7
LYS	2311	56.4	67.4	71.4	58.9	73.2	57.1	57.0	73.1
GLN	1562	60.0	65.9	73.8	56.8	73.6	63.9	54.2	73.3
ARG	1885	57.1	65.4	75.4	59.5	74.7	64.0	54.7	69.0
MET	788	59.5	65.0	74.5	59.3	75.5	70.8	53.0	77.0
SER	2661	44.8	62.4	57.4	56.8	64.2	38.6	41.4	60.1
GLU	2255	55.0	62.0	67.3	52.6	68.7	56.8	51.3	66.8
ALL	34220	57.4	72.6	77.2	63.0	80.5	66.6	61.5	76.8

Percentage  $\chi_1$  dihedral angle prediction within 40° of crystal structure (20° for Pro) for predictions on 180 proteins of known structure, with resolution better than or equal to 1.8 Å. Residue types are sorted in increasing difficulty of prediction, according to backbone-dependent rotamer library prediction (“BBDEP”). “BBIND” column lists

backbone-independent rotamer library prediction. The programs tested were CONFMAT [173], RAMP [193], SCWRL [93, 94], Sidemod [192], SMD [165, 191], and TORSO [319].

\* RAMP and SMD do not predict proline conformations.

ture Prediction'') organized by John Moult and colleagues [96–98, 277–280]. In the spring and summer before each meeting held in December 1994, 1996, 1998, and 2000, sequences of proteins whose structure was under active experimental determination by NMR or X-ray crystallography were distributed via the Internet. Anyone could submit structure predictions at various levels of detail (secondary structure predictions, sequence alignments to structures, and full 3-dimensional coordinates) before specific expiration dates for each target sequence. The models were evaluated via a number of computer programs written for the purpose, and then assessed by experts in each field, including comparative modeling, fold recognition, and *ab initio* structure prediction. The organizers invited predictors whose predictions were outstanding to present their methods and results at the meeting, and to describe their work in a special issue of the journal *Proteins*, published in the following year.

Ordinarily when protein structure prediction methods are developed, they are tested on sets of protein structures where the answer is known. Unfortunately, it is easy to select targets, even subconsciously, for which a particular method under development may work well. Also, it is easy to optimize parameters for a small test set that do not work as well for larger test sets. While the number of prediction targets in CASP is limited to numbers on the order of 10–20 per category, these numbers are still higher than many of the test sets used in testing new methods under development, as shown in Tables 5.2 and 5.3.

### 5.6.3

#### Protein health

A number of programs have been developed to ascertain the quality of experimentally determined structures and these can be used to determine whether a protein model obeys appropriate stereochemical rules. The two most popular programs are ProCheck [281–283] and WhatCheck [284]. These programs check bond lengths and angles, dihedral angles, planarity of  $sp^2$  groups, non-bonded atomic distances, disulfide bonds, and other characteristics of protein structures. One of the more useful checks is to see whether backbone geometries are in acceptable regions of the Ramachandran map. Backbone conformations in the forbidden regions are very likely to be incorrect. We have also developed a program, *bbdep*, that assesses whether side-chain rotamers in a structure are low-energy or high-energy conformations. The program reports unusual rotamers for the  $\phi, \psi$  positions of each residue, as well as unusual  $\chi_1, \chi_2, \chi_2, \chi_4$  rotamer combinations, and unusual dihedral angles.

It should be noted once again that correct geometry is no guarantee of correct structure prediction. In some cases, it may be better to tolerate a few steric conflicts or bad dihedral angles, rather than to minimize the struc-

ture's energy. While the geometry may look better, the final structure may be further away from the true structure (if it were known) than the unminimized structure.

The availability of these programs is listed in Table 5.6.

## 5.7

### Availability

Many programs are publicly available for the various steps in homology modeling and for evaluating and comparing structures, and for sequence alignment. I define "publicly available" as those programs which can be downloaded or used on a webserver for free or for a nominal charge to academic and non-profit research groups. There are commercial programs that cost on the order of \$2000–\$3000 per year or more (such as InsightII and ICM), and these are not included in this list. I have included as many programs as I could find by searching the Internet and contacting authors of papers that present methods for loop, side-chain, or full comparative modeling. The websites or electronic mail addresses for publicly available programs are listed in Table 5.6.

## 5.8

### Appendix

#### 5.8.1

##### Backbone conformations

In this Appendix, we describe some of the basic structural properties of proteins. Proteins are heteropolymers of 20 amino acid types with a backbone consisting of  $-\text{NH}-\text{C}(\text{H})\text{R}-\text{CO}-$  (except proline) where R is some functional group defined for each amino acid type. A fragment of 3 amino acids is shown in Figure 5.6. As with all organic molecules, there are characteristic bond lengths and angles associated with the types of chemical bonds formed, and proteins have been found to conform to the properties of small organic molecules as determined by very high resolution structures of small peptides [285]. The structures of proteins can therefore be defined approximately by a set of *dihedral angles* that determine the orientation of chemical groups in the peptide chain. As shown in Figure 5.7, a dihedral angle is defined for a set of 4 bonded atoms, say A-B-C-D, as the angle between the planes A-B-C and B-C-D. Looking down the B-C bond (atom B nearest the viewer, C further away), a positive dihedral angle is determined by rotation of bond A-B (or rather plane A-B-C) in a counter-clockwise

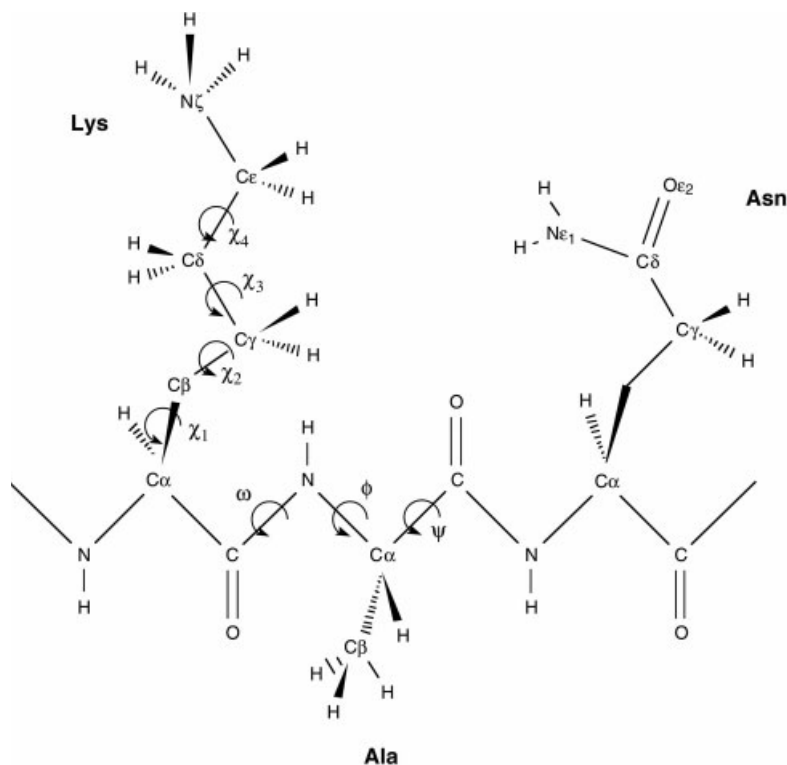
**Tab. 5.6**  
Web-sites and publicly available programs

	<b>Down- load</b>	<b>Web- server</b>	<b>Internet address</b>
<b>Structure alignment</b>			
CE	X	X	<a href="http://cl.sdsc.edu/ce.html">http://cl.sdsc.edu/ce.html</a>
Dali	X	X	<a href="http://www2.ebi.ac.uk/dali/">http://www2.ebi.ac.uk/dali/</a>
VAST		X	<a href="http://www.ncbi.nlm.nih.gov/80/Structure/VAST/vast.shtml">http://www.ncbi.nlm.nih.gov/80/Structure/VAST/vast.shtml</a>
MINAREA	X		<a href="http://www.cmpharm.ucsf.edu/cohen/pub/minarea.html">http://www.cmpharm.ucsf.edu/cohen/pub/minarea.html</a>
LOCK	X	X	<a href="http://gene.stanford.edu/lock/">http://gene.stanford.edu/lock/</a>
SARF2		X	<a href="http://genomic.sanger.ac.uk/123D/sarf2.html">http://genomic.sanger.ac.uk/123D/sarf2.html</a>
SSAP	X		<a href="http://www.biochem.ucl.ac.uk/~orengo/ssap.html">http://www.biochem.ucl.ac.uk/~orengo/ssap.html</a>
CATH			<a href="http://www.biochem.ucl.ac.uk/bsm/cath/">http://www.biochem.ucl.ac.uk/bsm/cath/</a>
STAMP		X	<a href="http://www.hgmp.mrc.ac.uk/Registered/Option/stamp.html">http://www.hgmp.mrc.ac.uk/Registered/Option/stamp.html</a>
COMPARER	X	X	<a href="http://www-cryst.bioc.cam.ac.uk/~robert/cpgs/COMPARER/comparer.html">http://www-cryst.bioc.cam.ac.uk/~robert/cpgs/COMPARER/comparer.html</a>
<b>Loop libraries</b>			
Protein Loop Classification (Oliva and Sternberg)		X	<a href="http://www.bmm.icnet.uk/loop/index.html">http://www.bmm.icnet.uk/loop/index.html</a>
Sloop (Burke, Deane, Blundell)		X	<a href="http://www-cryst.bioc.cam.ac.uk/~sloop/">http://www-cryst.bioc.cam.ac.uk/~sloop/</a>
Lessel & Schomburg	X		<a href="ftp://ftp.uni-koeln.de/institute/biochemie/pub/loop_db">ftp://ftp.uni-koeln.de/institute/biochemie/pub/loop_db</a>
<b>Loop prediction methods</b>			
BRAGI	X		<a href="ftp://ftp.gbf.de/pub/Bragi/">ftp://ftp.gbf.de/pub/Bragi/</a>
BTPRED		X	<a href="http://www.biochem.ucl.ac.uk/bsm/btpred">http://www.biochem.ucl.ac.uk/bsm/btpred</a>
CODA		X	<a href="http://www-cryst.bioc.cam.ac.uk/~charlotte/Coda/coda.html">http://www-cryst.bioc.cam.ac.uk/~charlotte/Coda/coda.html</a>
RAMP	X		<a href="http://www.ram.org/computing/ramp/ramp.html">http://www.ram.org/computing/ramp/ramp.html</a>
CONGEN	X		<a href="http://www.congenomics.com/congen/congen_toc.html">http://www.congenomics.com/congen/congen_toc.html</a>
Drawbridge	X		<a href="http://www.cmpharm.ucsf.edu/cohen/pub/">http://www.cmpharm.ucsf.edu/cohen/pub/</a>
Confmat	X		Contact: <a href="mailto:koehl@allegro.stanford.edu">koehl@allegro.stanford.edu</a>
Protein Loop Classification (Oliva et al.)		X	<a href="http://www.bmm.icnet.uk/loop/">http://www.bmm.icnet.uk/loop/</a>
Swiss-PdbViewer	X	X	<a href="http://www.expasy.ch/spdbv">http://www.expasy.ch/spdbv</a>
<b>Rotamer libraries</b>			
Backbone-dependent rotamer library (Dunbrack)	X		<a href="http://www.fccc.edu/research/labs/dunbrack/sidechain.html">http://www.fccc.edu/research/labs/dunbrack/sidechain.html</a>



Tab. 5.6 (continued)

	<i>Down- load</i>	<i>Web- server</i>	<i>Internet address</i>
Lovell	X		<a href="http://kinemage.biochem.duke.edu/website/rotamer.htm">http://kinemage.biochem.duke.edu/ website/rotamer.htm</a>
DeMaeyer			<a href="http://www.fccc.edu/research/labs/dunbrack/sidechain/demaeyer.rot">http://www.fccc.edu/research/labs/ dunbrack/sidechain/demaeyer.rot</a>
Tuffery			<a href="http://condor.urbb.jussieu.fr/rotamer.html">http://condor.urbb.jussieu.fr/ rotamer.html</a>
Ponder & Richards	X		<a href="http://www.fccc.edu/research/labs/dunbrack/sidechain/ponder_richards.rot">http://www.fccc.edu/research/labs/ dunbrack/sidechain/ ponder_richards.rot</a>
<b>Side-chain prediction methods</b>			
Confmat	X		Contact: Patrice Koehl, <a href="mailto:koehl@allegro.stanford.edu">koehl@allegro.stanford.edu</a>
FAMS		X	<a href="http://physchem.pharm.kitasato-u.ac.jp/FAMS/fams.html">http://physchem.pharm.kitasato-u.ac.jp/ FAMS/fams.html</a>
RAMP	X		<a href="http://www.ram.org/computing/ramp/ramp.html">http://www.ram.org/computing/ramp/ ramp.html</a>
SCWRL	X		<a href="http://www.fccc.edu/research/labs/dunbrack/scwrl">http://www.fccc.edu/research/labs/ dunbrack/scwrl</a>
Segmod/CARA (Genemine/Look3)	X		<a href="http://www.bioinformatics.ucla.edu/~genemine">http://www.bioinformatics.ucla.edu/ ~genemine</a>
Sidemod	X		<a href="http://www.rtc.riken.go.jp/~hkono/SideChain">http://www.rtc.riken.go.jp/~hkono/ SideChain</a>
SMD	X		<a href="http://condor.urbb.jussieu.fr/Smd.html">http://condor.urbb.jussieu.fr/Smd.html</a>
torso	X	X	<a href="http://www2.ebi.ac.uk/dali/maxsprout/maxsprout.html">http://www2.ebi.ac.uk/dali/maxsprout/ maxsprout.html</a>
WhatIf	X	X	<a href="http://www.cmbi.kun.nl/whatif">http://www.cmbi.kun.nl/whatif</a>
<b>General Modeling Programs</b>			
COMPOSER	X		<a href="http://www-cryst.bioc.cam.ac.uk/">http://www-cryst.bioc.cam.ac.uk/</a>
MODELLER	X		<a href="http://guitar.rockefeller.edu/modeller/modeller.html">http://guitar.rockefeller.edu/modeller/ modeller.html</a>
PrISM	X		<a href="http://www.columbia.edu/~ay1/">http://www.columbia.edu/~ay1/</a>
CONGEN (Montelione)	X		<a href="http://www-nmr.cabm.rutgers.edu/software/html/nmr_software.html">http://www-nmr.cabm.rutgers.edu/ software/html/nmr_software.html</a>
RAMP	X		<a href="http://www.ram.org/computing/ramp/ramp.html">http://www.ram.org/computing/ramp/ ramp.html</a>
SwissModel		X	<a href="http://www.expasy.ch/swissmod/SWISSMODEL.html">http://www.expasy.ch/swissmod/ SWISSMODEL.html</a>
WhatIf	X	X	<a href="http://www.cmbi.kun.nl/whatif">http://www.cmbi.kun.nl/whatif</a>
<b>Protein health</b>			
ProCheck	X		<a href="http://www.biochem.ucl.ac.uk/~roman/procheck/procheck.html">http://www.biochem.ucl.ac.uk/~roman/ procheck/procheck.html</a>
WhatCheck	X		<a href="http://www.cmbi.kun.nl/whatif/whatcheck/">http://www.cmbi.kun.nl/whatif/ whatcheck/</a>
Promotif	X		<a href="http://www.biochem.ucl.ac.uk/bsm/promotif/promotif.html">http://www.biochem.ucl.ac.uk/bsm/ promotif/promotif.html</a>
BBDEP	X		<a href="http://www.fccc.edu/research/labs/dunbrack/sidechain.html">http://www.fccc.edu/research/labs/ dunbrack/sidechain.html</a>



**Fig. 5.6**  
Diagram of tripeptide with sequence Lys-Ala-Asn. Backbone dihedrals  $\phi$ ,  $\psi$ ,  $\omega$  of the central Ala residue are indicated. Side-chain dihedrals  $\chi_1$ ,  $\chi_2$ ,  $\chi_3$ , and  $\chi_4$  of Lys are also indicated.

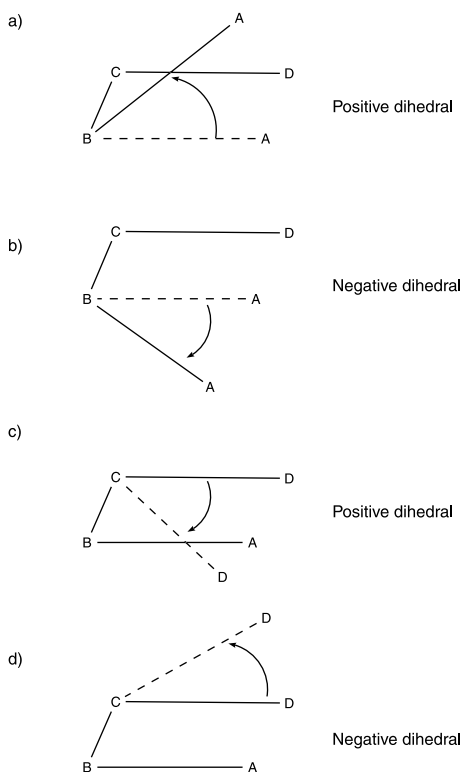
direction about the B-C axis. Alternatively, a positive dihedral is determined by rotation of bond C-D in a clockwise direction about the B-C axis.

Along the protein backbone, the dihedral angles  $\omega$ ,  $\phi$ , and  $\psi$  are used to describe the conformation, as shown in Figure 5.6. These are defined as  $\omega_i = \text{C}\alpha_{i-1}-\text{C}_{i-1}-\text{N}_i-\text{C}\alpha_i$ ,  $\phi_i = \text{C}_{i-1}-\text{N}_i-\text{C}\alpha_i-\text{C}_i$ ,  $\psi_i = \text{N}_i-\text{C}\alpha_i-\text{C}_i-\text{N}_{i+1}$ . Because the peptide bond between C and N has partial double-bond character, the amide group is nearly flat. That is, the atoms N,  $\text{C}\alpha$ , H,  $\text{C}_{i-1}$ ,  $\text{O}_{i-1}$ , and  $\text{C}\alpha_{i-1}$  all lie in a plane.  $\omega$  is therefore always close to  $180^\circ$  (*trans*) or  $0^\circ$  (*cis*). For all residues except proline,  $\omega$  is nearly always *trans* (less than 0.1% are *cis*). Approximately 5% of proline peptide bonds are *cis*.

Only certain  $\phi$ ,  $\psi$  conformation pairs are observed. This is usually described in terms of the Ramachandran diagram, as shown in Figure 5.8, with  $\phi$  as the horizontal axis and  $\psi$  as the vertical axis. In Figure 5.8, the experimentally determined backbone dihedral angles for the alanine, glycine, and proline residues of 699 proteins are depicted. There are three basic

**Fig. 5.7**

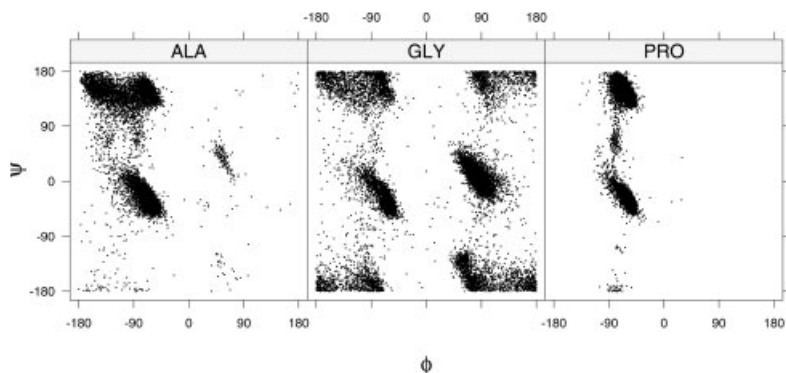
Dihedral angle definition. The dihedral angle of atoms A,B,C,D is shown as (A) a positive angle by counter-clockwise rotation of the B-A bond about the B-C bond; (B) a negative angle by clockwise rotation of B-A; (C) a positive angle by clockwise rotation of the C-D bond; (D) a negative angle by counterclockwise rotation of C-D.



factors that determine the broad features of backbone conformations observed in proteins as shown in Figure 5.8: (1) steric interactions along the protein backbone that prevent the existence of some conformations; (2) electrostatic interactions in the formation of secondary structures by hydrogen bonding of backbone NH and CO groups; and (3) backbone-side-chain interactions, both steric and electrostatic (and covalent in the case of proline), that account for the variation in Ramachandran distributions among the 20 amino acids. We describe each of these in turn.

#### 5.8.1.1 Steric interactions

Both the  $\phi$  and  $\psi$  dihedrals are composed of 4 atoms with hybridization  $sp^2$ - $sp^2$ - $sp^3$ - $sp^2$ . The energy of a conformation with either  $\phi$  or  $\psi$  approaching  $0^\circ$  is up to 5 kcal/mol higher than the energy minimum because of steric interactions of the terminal atoms. In the case of  $\phi$  these are the carbonyl carbon of succeeding residues. In the case of  $\psi$ , these are the nitrogen atoms of succeeding residues. The Ramachandran distribution for alanine clearly demonstrates that these conformations ( $\phi \sim 0^\circ$  or  $\psi \sim 0^\circ$ ) are rare in

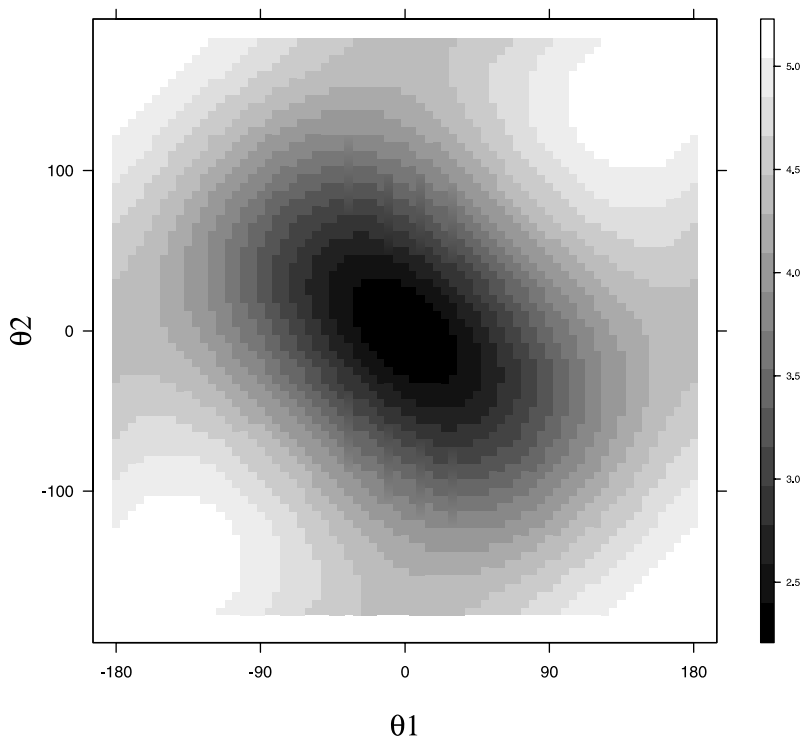


**Fig. 5.8**  
Ramachandran plot. The density of  $\phi$ ,  $\psi$  values for alanines, glycines, and prolines in 699 proteins of known structure. Proteins were selected from the PDB with

resolution better than or equal to 1.8 Å and less than 50% sequence identity among all pairs in the list using the *culledpdb* algorithm (<http://www.fccc.edu/research/labs/dunbrack/culledpdb.html>).

the database. We must also consider the presence of the  $\beta$  carbon in all non-glycine amino acids. The dihedral  $C_{i-1}-N_i-C\alpha_i-C\beta_i$ , is equal to  $\phi_i - 120^\circ$ . This dihedral is near  $0^\circ$  when  $\phi = +120^\circ$ . There are very few alanines (or other residues with  $\beta$  carbons) in the database with this conformation. The dihedrals  $C\beta_i-C\alpha_i-C_i-N_{i+1}$  and  $C\beta_i-C\alpha_i-C_i-O_i$  are equal to  $\psi_i + 120^\circ$  and  $\psi_i - 60^\circ$  respectively. These dihedrals are  $0^\circ$  when  $\psi_i$  is  $-120^\circ$  and  $+60^\circ$ . Conformations with  $\psi_i$  near these values are also clearly relatively rare for residues with  $C\beta$  atoms. Because glycine lacks  $C\beta$  (its side chain is simply a hydrogen atom), glycine can not incur these steric penalties. In Figure 5.8B, the Ramachandran conformations of glycine residues in the same 699 proteins are plotted.

Another important effect is the steric interactions of non-hydrogen atoms separated by 4 covalent bonds. Let us label them A-B-C-D-E. There are two dihedrals that determine the separation in space of atoms A and E in this chain:  $\theta_1 = A-B-C-D$  and  $\theta_2 = B-C-D-E$ . When both  $\theta_1$  and  $\theta_2$  are  $0^\circ$ , atoms A and E are very close together. The distance between A and E falls off quickly when both  $\theta_1$  and  $\theta_2$  increase, or when they both decrease. The distance falls off much more slowly when  $\theta_1$  increases while  $\theta_2$  decreases or *vice versa*. In Figure 5.9, the distance between atoms A and E is shown as a function of the two dihedrals  $\theta_1$  and  $\theta_2$ , assuming the bond lengths are 1.55 Å and the bond angles are  $115^\circ$  (the average of  $sp^2$  and  $sp^3$  angles). To determine the effect of this interaction on Ramachandran distributions we have to consider all chains of 5 atoms that vary with  $\phi$  and/or  $\psi$  (along with the dihedrals A-B-C-D and B-C-D-E connecting them). These are shown in Table 5.7. The reader can confirm that these interactions reduce the



**Fig. 5.9**

Distance between atoms A and E in a five atom chain, A-B-C-D-E as a function of the A-B-C-D ( $\theta_1$ ) and B-C-D-E ( $\theta_2$ ) dihedrals.

The A–E distance was calculated by assuming the bond lengths are 1.55 Å and the bond angles are 115° (the average of  $sp^2$  and  $sp^3$  angles).

probability of certain Ramachandran conformations as seen in Figure 5.8A. Because glycine lacks  $C\beta$  it can attain backbone conformations that other residue types can not – those listed in Table 5.7 that include  $C\beta$ . Because proline contains a five-membered ring consisting of N-C $\alpha$ -C $\beta$ -C $\gamma$ -C $\delta$  atoms, the  $\phi$  dihedral is highly constrained to be near  $\phi = -60 \pm 30^\circ$ . The Ramachandran distribution for proline is shown in Figure 5.8C.

### 5.8.1.2 Electrostatic interactions

Because the carbonyl and amide NH groups are highly polar, they form hydrogen bonds easily with water molecules. To bury these groups away from solvent (on the interior of a protein structure), requires that they are almost always involved in hydrogen bonds with other portions of the backbone or with side-chain groups. An efficient way of fulfilling this obligation occurs in the regular secondary structures of  $\alpha$ -helices and  $\beta$ -sheets. In the

**Tab. 5.7**  
Steric interactions along backbone that determine Ramachandran distributions

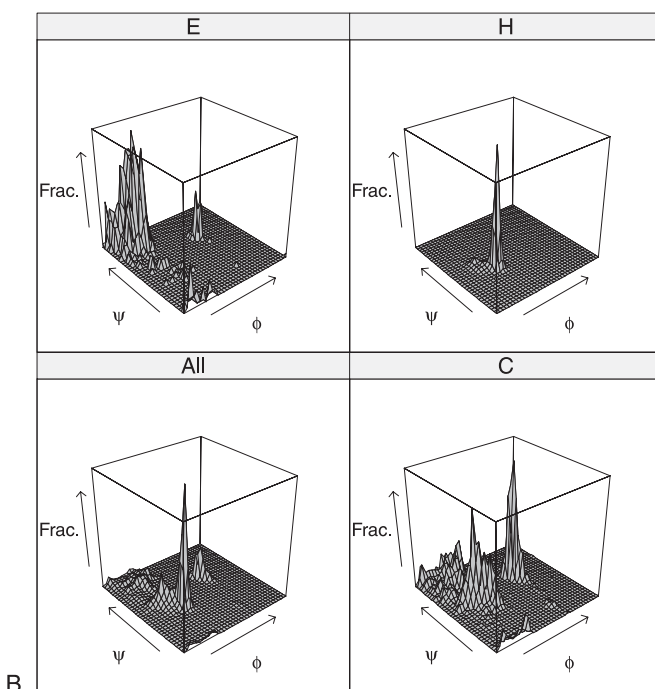
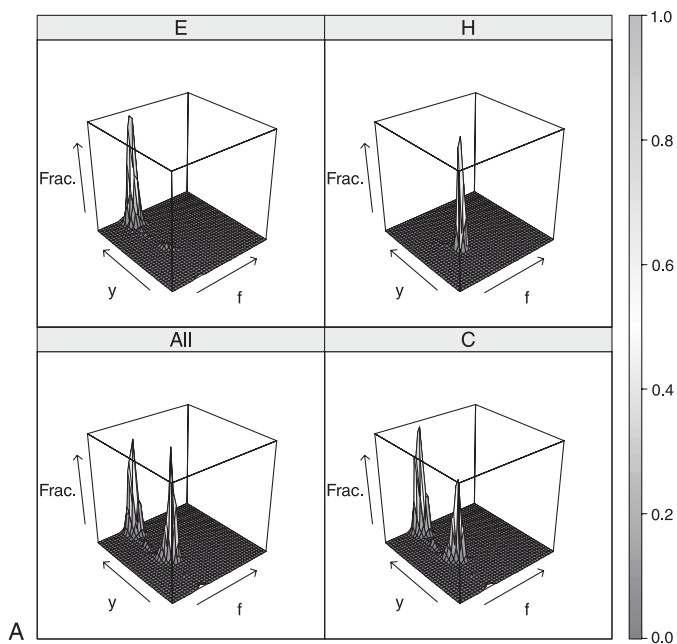
A	B	C	D	E	$\theta_1$	$\theta_2$	$\omega$	$\phi$	$\psi$
$C\alpha_{i-1}$	$C_{i-1}$	$N_i$	$C\alpha_i$	$C_i$	$\omega$	$\phi$	0	0	
$C\alpha_{i-1}$	$C_{i-1}$	$N_i$	$C\alpha_i$	$C\beta_i$	$\omega$	$\phi - 120^\circ$	0	120	
$O_{i-1}$	$C_{i-1}$	$N_i$	$C\alpha_i$	$C_i$	$\omega - 180^\circ$	$\phi$	180	0	
$O_{i-1}$	$C_{i-1}$	$N_i$	$C\alpha_i$	$C\beta_i$	$\omega - 180^\circ$	$\phi - 120^\circ$	180	120	
$C_{i-1}$	$N_i$	$C\alpha_i$	$C_i$	$N_{i+1}$	$\phi$	$\psi$	0	0	
$C_{i-1}$	$N_i$	$C\alpha_i$	$C_i$	$O_i$	$\gamma$	$\psi - 180^\circ$	0	180	
$N_i$	$C\alpha_i$	$C_i$	$N_{i+1}$	$C\alpha_{i+1}$	$\psi$	$\omega$	0		0
$C\beta_i$	$C\alpha_i$	$C_i$	$N_{i+1}$	$C\alpha_{i+1}$	$\psi + 120^\circ$	$\omega$	0		-120

Possible steric interactions of 2 atoms (A and E) on the ends of a 5-atom chain (A-B-C-D-E). The dihedral A-B-C-D is denoted  $\theta_1$  and the dihedral B-C-D-E is denoted  $\theta_2$ . These dihedrals for each set of atoms are expressed as functions of backbone dihedrals  $\phi$ ,  $\psi$ , and  $\omega$ . The values of  $\phi$ ,  $\psi$ , and  $\omega$  when  $\theta_1$  and  $\theta_2$  are zero are given in the last three columns. Large steric interactions will occur when  $\theta_1$  and  $\theta_2$  are both  $0^\circ$ . This interaction will fall off quickly as both dihedrals decrease or both dihedrals increase, but much more slowly as one increases from  $0^\circ$  and the other decreases (see Figure 5.7).

case of  $\alpha$ -helices, hydrogen bonds are formed between the C=O of amino acid  $i$  and the amide HN group of amino acid  $i + 4$  when the backbone dihedral angles are close to  $(-45^\circ, -55^\circ)$ . Most  $\alpha$ -helices are at least 2 turns long, and often longer.  $\beta$ -sheets are secondary structures consisting of two or more parallel or anti-parallel backbone segments that form hydrogen bonds between them with their backbone NH and C=O groups. For several residue types, we have drawn Ramachandran diagrams for residues in  $\alpha$ -helices,  $\beta$ -sheets, and coil regions in separate plots in Figure 5.10 to demonstrate that regular secondary structures populate only certain portions of the map, and that coil residues occur in many areas.

### 5.8.1.3 Backbone-side-chain interactions

We describe the effect of backbone conformations on side-chain conformations below, but first we show that non-Gly, non-Pro amino acids do not have identical Ramachandran distributions, due to interactions between the side chain and the backbone. In Figure 5.10B and 5.10C we show the Asn and Val Ramachandran distributions in each secondary structure to demonstrate this effect. Asn in particular has high content of residues with  $\phi > 0^\circ$  in coil because of backbone-side-chain hydrogen bonds. Val has very few residues with  $\phi > 0^\circ$  because of steric clashes of the two  $C\gamma$  atoms with the backbone.



**Fig. 5.10**  
 Ramachandran distributions of A) Alanine; B) Asparagine; C) Valine separated by secondary structure type. Data was taken from the same 699 proteins of known

structure as described in the caption to Figure 5.8. Secondary structure was determined with the STRIDE program [291]. H denotes Helix, E denotes Sheet, and C denotes Coil.

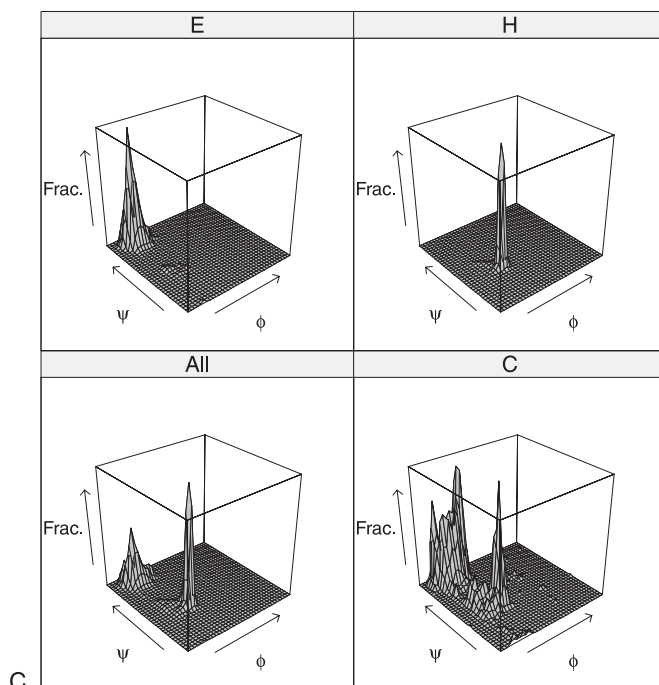


Fig. 5.10 (continued)

## 5.8.2

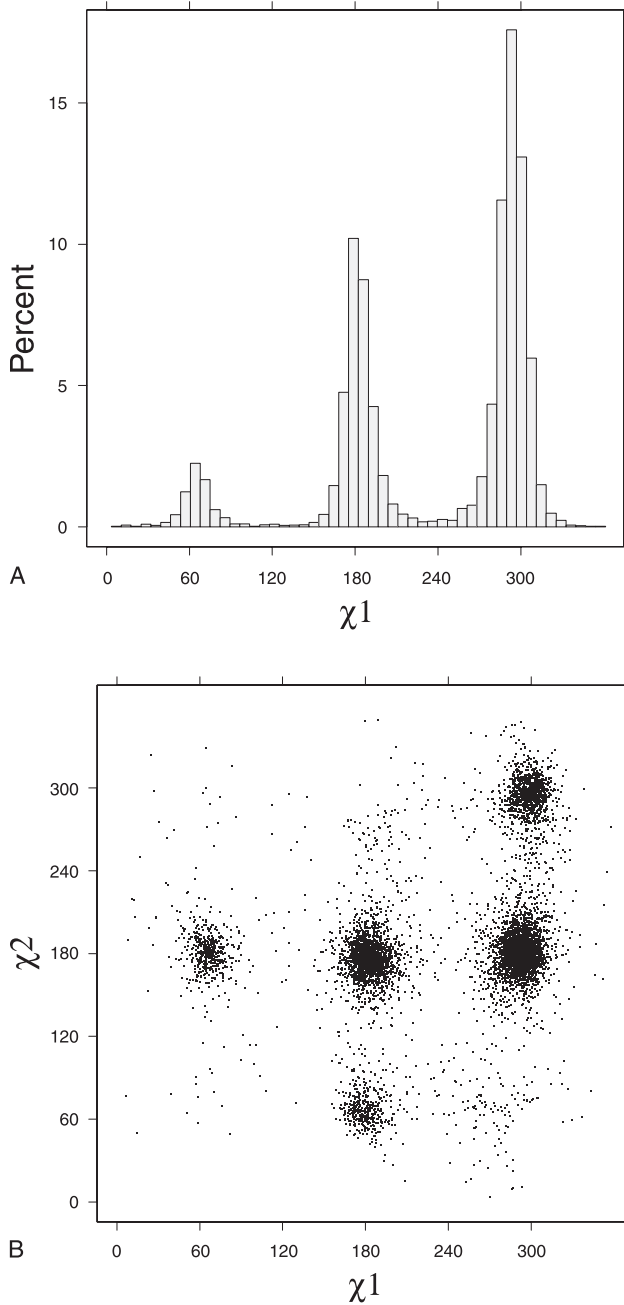
### Side-chain conformational analysis

#### 5.8.2.1 Backbone-conformation-independent interactions

Side chains can also be described by their dihedral angles, and these are denoted  $\chi_1, \chi_2, \chi_3$ , and  $\chi_4$ . Except for glycine, whose side chain is only a single hydrogen atom bonded to the  $\alpha$ -carbon of the backbone, and alanine, whose side chain is only a single methyl group bonded to the  $\alpha$ -carbon of the backbone, all other side chains have a  $\chi_1$  dihedral angle defined as N-C $\alpha$ -C $\beta$ -X where X is either C $\gamma$ , or C $\gamma$ 1 (Val, Ile), O $\gamma$  (Ser), O $\gamma$ 1 (Thr), or S $\gamma$  (Cys). Longer side chains have additional  $\chi$  dihedrals, as shown in Figure 5.6. In Figure 5.11 we show the distribution of  $\chi_1$  dihedrals and  $\chi_1, \chi_2$  for Lys. As we expect from organic chemistry, the dihedrals are clustered near  $+60^\circ$ ,  $180^\circ$ , and  $-60^\circ$ . These conformations are denoted as  $g^+$ ,  $t$ , and  $g^-$  respectively, where  $g$  represents “gauche” and  $t$  represents trans. It should be noted that some authors reverse the  $g^+$  and  $g^-$  definitions.

The rotamers are not at all evenly distributed among the possible staggered dihedrals. The reason for this can be seen in the Newman diagram in





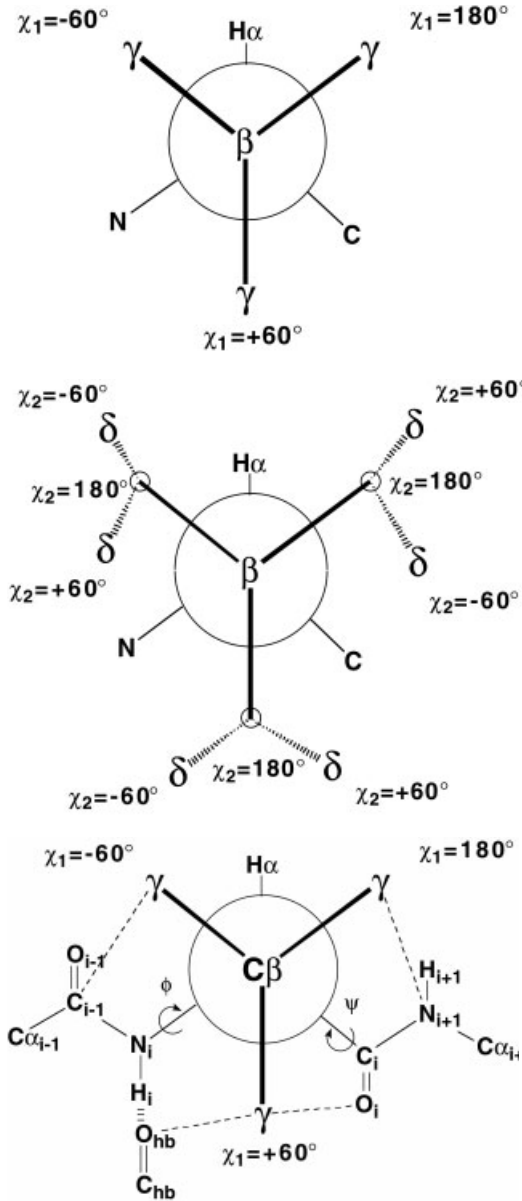
**Fig. 5.11**  
Distribution of A)  $\chi_1$  and B)  $\chi_1$ - $\chi_2$  side-chain dihedrals for Lysine. Lysines in the

same 699 proteins as described in the caption to Figure 5.8 were used to produce the plot.

Figure 5.12. First, the gauche interaction (dihedral =  $+60^\circ$  or  $-60^\circ$ ) of the  $\chi_1$  dihedral, N-C $\alpha$ -C $\beta$ -C $\gamma$ , increases the energy by about 0.9 kcal/mol per gauche interaction [286]. The  $t$  and  $g^-$  rotamers each have one gauche interaction with the backbone (backbone C and N respectively), while the  $g^+$  has gauche interactions with *both* the backbone C and N atoms. Its energy is therefore about 0.9 kcal/mol higher than the other two rotamers. This is a backbone-conformation independent interaction; that is, it does not depend on backbone dihedrals  $\phi$  and  $\psi$ . Second, both gauche and 1–5 interactions (as shown in Figure 5.9) affect the distribution of  $\chi_1$ – $\chi_2$  pairs of dihedrals. As discussed earlier for interactions along the backbone, the distance between atoms A and E of a 5-atom chain remain close to one another if the dihedrals are of opposite sign. For  $\chi_1$ – $\chi_2$  dihedral pairs, this means that when  $\chi_1$  is  $+60^\circ$  and  $\chi_2$  is  $-60^\circ$ , or vice versa, the energy rises because of the steric interaction of the nitrogen and C $\delta$  atoms in the 5-atom chain N-C $\alpha$ -C $\beta$ -C $\gamma$ -C $\delta$ . But this kind of interaction also occurs between C $\delta$  and the carbonyl carbon. This occurs when  $\chi_1$  is  $+60^\circ$  and  $\chi_2$  is  $+60^\circ$  and when  $\chi_1$  is  $180^\circ$  and  $\chi_2$  is  $-60^\circ$ . The effects of these interactions can be observed in the  $\chi_1$ – $\chi_2$  distribution for lysine in Figure 5.11B.

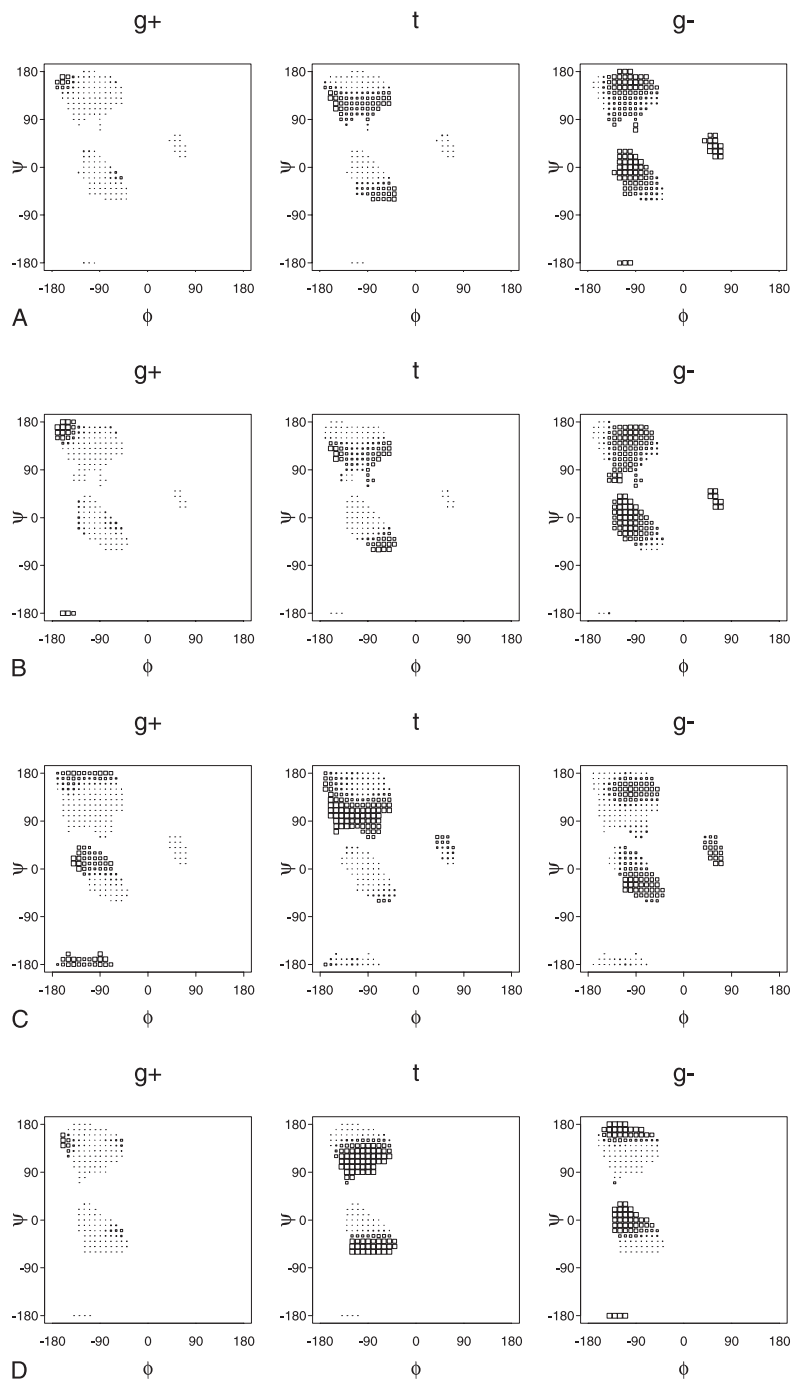
### 5.8.2.2 Backbone-conformation-dependent interactions

One of the prime determinants of  $\chi_1$  rotamer choice is steric interaction of the  $\gamma$  heavy atom with the backbone in a manner that is dependent on the conformation of the backbone, that is, the dihedrals  $\phi$  and  $\psi$ . In Figure 5.13, the proportions of the  $g^+$ ,  $t$ , and  $g^-$  rotamers are shown on the Ramachandran map for Lys, Phe, Asp, and Val. The patterns of low energy and high energy rotamers are immediately apparent, with some regularities (and some differences) between the different amino acid types. The reason for strong backbone dependence can be observed in Figure 5.12B. The possible interactions are listed in Table 5.8. Steric interactions can occur between heavy atoms at positions  $i$  and  $i + 4$  when the two dihedrals connecting them take on values in the gray region of Figure 5.9. We count from the  $\gamma$  carbon to atoms C $_{i-1}$ , O $_i$ , and N $_{i+1}$  of the backbone, and determine the disallowed combinations of dihedrals in these 5-atom chains, C $_{i-1}$ -N $_i$ -C $\alpha$ -C $\beta$ -C $\gamma$ , O $_i$ -C $_i$ -C $\alpha$ -C $\beta$ -C $\gamma$ , and N $_{i+1}$ -C $_i$ -C $\alpha$ -C $\beta$ -C $\gamma$  respectively. The second dihedral in each of these sequences is either  $\chi_1$  or  $\chi_1 - 120^\circ$  (see Figure 5.6). Since the  $\chi_1$  dihedral is likely to be in a staggered position ( $60^\circ$ ,  $180^\circ$ ,  $-60^\circ$ ) relative to backbone N and C, we need to determine the range of the dihedrals C $_{i-1}$ -N $_i$ -C $\alpha$ -C $\beta$ , O $_i$ -C $_i$ -C $\alpha$ -C $\beta$ , and N $_{i+1}$ -C $_i$ -C $\alpha$ -C $\beta$  allowed. For instance, if  $\chi_1$  is  $-60^\circ$ , Figure 5.9 indicates that C $_{i-1}$ -N $_i$ -C $\alpha$ -C $\beta$  is disallowed when it takes on values of  $+30^\circ \pm 50^\circ$ . Since this dihedral is equal to  $\phi - 120^\circ$ , the  $g^+$  rotamer is disallowed when  $\phi = +150^\circ \pm 50^\circ$ . This is evident in the plots of Figure 5.13, where the  $g^-$  rotamer is uncommon on the far left of each plot ( $\phi \sim -180^\circ$ ). By the same reasoning, the  $g^+$  rotamer is not allowed when  $\phi = +90^\circ \pm 50^\circ$ .



**Fig. 5.12** Newman diagrams for protein backbone-side-chain interactions. **(A)** Backbone-independent interactions, looking down the  $C\alpha$ - $C\beta$  bond from  $C\beta$ ; certain combinations of  $\chi_1$  and  $\chi_2$  are forbidden because of close interactions between  $C\delta$  and backbone N and C of the same residue. In the lower diagram, the  $\chi_2 = 180^\circ$  rotamer is pointing towards the viewer. The four  $\chi_1, \chi_2$  combinations which

are forbidden are those with  $C\delta$  (denoted “ $\delta$ ”) are closest to the backbone N and C atoms. **(B)** Backbone-dependent interactions, looking down the  $C\alpha$ - $C\beta$  bond from  $C\beta$ ; certain combinations of  $\chi_1$  and  $\phi$  and certain combinations of  $\chi_1$  and  $\psi$  are essentially forbidden because of interactions between  $C\gamma$  and backbone atoms  $C_{i-1}$ ,  $N_{i+1}$ ,  $O_i$ , and a hydrogen bond acceptor to  $HN_i$ .



**Fig. 5.13**

Distribution of  $\chi_1$  side-chain rotamers as a function of backbone dihedrals  $\phi$  and  $\psi$  for (A) Lys; (B) Phe; (C) Asp; (D) Val. Each square represents the fraction of each

rotamer type in  $10^\circ$  ranges of  $\phi$  or  $\psi$  in a sample of 699 proteins as described in Figure 5.8. Only ranges of  $\phi$  or  $\psi$  for which there were at least 20 residues available in this sample of proteins are plotted.

Tab. 5.8  
Steric interactions along backbone that determine Ramachandran distributions

A	B	C	D	E	$\theta_1$	$\theta_1$ value	$\theta_2$	$\theta_2$ value	$\chi_1$	$\phi$	$\psi$
C $\gamma$	C $\beta$	C $\alpha$	N $_i$	C $_{i-1}$	$\chi_1$	+60°	$\phi$	-30°	+60°	-30°	+30°
C $\gamma$	C $\beta$	C $\alpha$	N $_i$	O..HN $_i$	$\chi_1$	+60°	$\phi - 180^\circ$	-30°	+60°	150°	-150°
C $\gamma$	C $\beta$	C $\alpha$	N $_i$	C $_{i-1}$	$\chi_1$	-60°	$\phi$	+30°	-60°	+30°	+90°
C $\gamma$	C $\beta$	C $\alpha$	N $_i$	O..HN $_i$	$\chi_1$	-60°	$\phi - 180^\circ$	+30°	-60°	-150°	-90°
C $\gamma$	C $\beta$	C $\alpha$	C $_i$	O $_i$	$\chi_1 - 120^\circ$	+60°	$\psi - 60^\circ$	-30°	180°		+30°
C $\gamma$	C $\beta$	C $\alpha$	C $_i$	N $_{i+1}$	$\chi_1 - 120^\circ$	+60°	$\psi + 120^\circ$	-30°	180°		-150°
C $\gamma$	C $\beta$	C $\alpha$	C $_i$	O $_i$	$\chi_1 - 120^\circ$	-60°	$\psi - 60^\circ$	+30°	+60°		+90°
C $\gamma$	C $\beta$	C $\alpha$	C $_i$	N $_{i+1}$	$\chi_1 - 120^\circ$	-60°	$\psi + 120^\circ$	+30°	+60°		-90°
C $\gamma$ 2	C $\beta$	C $\alpha$	N $_i$	C $_{i-1}$	$\chi_1 + 120^\circ$	+60°	$\phi$	-30°	-60°	-30°	
C $\gamma$ 2	C $\beta$	C $\alpha$	N $_i$	O..HN $_i$	$\chi_1 + 120^\circ$	+60°	$\phi - 180^\circ$	-30°	-60°	150°	
C $\gamma$ 2	C $\beta$	C $\alpha$	N $_i$	C $_{i-1}$	$\chi_1 + 120^\circ$	-60°	$\phi$	+30°	180°	+30°	
C $\gamma$ 2	C $\beta$	C $\alpha$	N $_i$	O..HN $_i$	$\chi_1 + 120^\circ$	-60°	$\phi - 180^\circ$	+30°	180°	-150°	
C $\gamma$ 2	C $\beta$	C $\alpha$	C $_i$	O $_i$	$\chi_1$	+60°	$\psi - 60^\circ$	-30°	+60°		+30°
C $\gamma$ 2	C $\beta$	C $\alpha$	C $_i$	N $_{i+1}$	$\chi_1$	+60°	$\psi + 120^\circ$	-30°	+60°		-150°
C $\gamma$ 2	C $\beta$	C $\alpha$	C $_i$	O $_i$	$\chi_1$	-60°	$\psi - 60^\circ$	+30°	-60°		+90°
C $\gamma$ 2	C $\beta$	C $\alpha$	C $_i$	N $_{i+1}$	$\chi_1$	-60°	$\psi + 120^\circ$	+30°	-60°		-90°

Possible steric interactions of 2 atoms (A and E) on the ends of a 5-atom chain (A-B-C-D-E). The dihedral A-B-C-D is denoted  $\theta_1$  and the dihedral B-C-D-E is denoted  $\theta_2$ . These dihedrals for each set of atoms are expressed as functions of side-chain dihedral  $\chi_1$  and the backbone dihedrals  $\phi$  and  $\psi$ . Because  $\chi_1$  must be near one of the canonical staggered dihedral values (+60°, 180°, and -60°), steric interactions can occur when the value of  $\theta_1$  is +60° or -60°, and  $\theta_2$  is -30°  $\pm$  50°

or +30°  $\pm$  50° respectively (see Figure 5.7). For the side-chain types with one  $\gamma$  heavy atom, the entries listed for C $\gamma$  apply. For side-chains with two  $\gamma$  heavy atoms (Val, Thr, Ile), interactions listed for C $\gamma$  and C $\gamma$ 2 apply. Because of the definition of  $\chi_1$  for Ile and Thr,  $\chi_1$  for these residues is equal to  $\chi_1 + 120^\circ$  as listed in the Table. So for instance the -60° rotamer in the Table is equivalent to the +60° rotamer for Ile and Thr.

The dihedral  $O_i-C_i-C\alpha-C\beta$  is equal to  $\psi - 60^\circ$ .  $\chi_1 - 120^\circ$  is  $-60^\circ$  for the  $g^+$  rotamer, and therefore the  $g^+$  rotamer is forbidden when  $\psi - 60^\circ$  is  $+30^\circ \pm 50^\circ$ , or when  $\psi = +90^\circ \pm 50^\circ$ , because of a steric interaction between  $C\gamma$  and  $O_i$ . The interaction with backbone atom  $N_{i+1}$  occurs  $180^\circ$  away from this, or when  $\psi$  is  $-90^\circ \pm 50^\circ$ . Both of these interactions are evident in the low proportions of the  $g^+$  rotamers in these regions in Figure 5.13. Finally, the same interactions occur for the  $t$  rotamer ( $\chi_1 - 120^\circ = +60^\circ$ ) when  $\psi - 60^\circ$  is  $-30^\circ \pm 50^\circ$  or when  $\psi = +30^\circ \pm 50^\circ$  and again  $180^\circ$  away when  $\psi = -150^\circ \pm 50^\circ$ . Because valine, isoleucine, and threonine have two  $\gamma$  heavy atoms, interactions with both atoms must be considered in this analysis. It is clear in Figure 5.13C that valine is highly restricted by the backbone conformation, since in most regions of the Ramachandran map two out of three  $\chi_1$  rotamers are disallowed, leaving only one likely conformation.

## 5.9

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## 6

### Protein Structure Prediction

*Ralf Zimmer and Thomas Lengauer*

Proteins fold into nice-looking, three-dimensional structures [1, 2]. These structures are an essential factor of the protein's function. Therefore, extensive experimental efforts have been invested to solve the structures for the most interesting, abundant, or medically, biologically, or pharmaceutically important proteins. Classically, X-ray crystallography has been the dominant technique for resolving protein structures. Here the protein has to be purified and crystallized, before a diffraction diagram can be measured by exposing the crystal to X-ray radiation. The diffraction diagram then has to be Fourier transformed to obtain the electron density map of the protein. The latter problem is aggravated by the fact that the diffraction diagram only reveals the intensity but not the phase of the interfering X-rays. In order to obtain the electron density from the diffraction diagram computer-based modeling methods have been established. More recently, NMR techniques have been developed and refined to solve larger and larger proteins structures in solution. These methods yield 2D data that have to be interpreted by a laborious computer-based process, in order to reveal the electron density [1]. Thus, for both approaches, not only are the experimental costs enormous but also the computer methods involved. The motivation to go through the time-consuming and expensive process of determining a detailed atomic structure of a bio-molecule ranges from the fundamental research interest to "see" the molecule under investigation and to structurally understand how it works, to being able to exploit the protein structure for rationally designing molecular interaction partners, which influence the protein by either inhibiting or activating it (see also Chapters 1, 7 and 8).

As part of the protein folding problem, determining the 3D structure of a protein from its 1D sequence is a grand challenge in molecular biology ("the second half of the genetic code"). Since Anfinsen's [3] refolding experiments it seems that the primary sequence often completely determines the tertiary structure according to the principles of thermodynamics. But also often protein folding appears to be dependent on a number of additional factors: post-translational editing, covalent modifications, interactions

with other molecules and membranes, and on cofactors or chaperones necessary for correct folding.

Bioinformatics approaches towards solving protein structures are called protein structure prediction methods. They can roughly be categorized into three types according to the input data used and the available knowledge to be exploited: 2D prediction, *ab initio* structure prediction, and homology-based modeling. 2D or secondary structure prediction methods just assign secondary structures to the residues of a protein, i.e. they decide whether the residue occurs in a  $\alpha$ -helix, a  $\beta$ -strand or neither. *Ab initio* methods predict structure from sequence without any additional information on the protein. They usually employ energy minimization or molecular dynamics methods (see Chapter 5 and Appendix). Homology-based prediction as a first step needs to construct a rough model for a protein sequence after the known structure of another protein (the so-called template) via a sequence-structure alignment, a so-called threading, as a basis for further homology-based modeling (see Chapter 5). Threading methods only predict the course of the protein backbone, whereas homology-based modeling methods compute a full-atom model of the protein 3D structure. Different structure prediction methods have different applications. E.g., protein threading can extend the current capabilities of sequence analysis methods (see Chapter 2) with respect to detecting remote homology and similarities in order to derive functional and structural features. Homology-based modeling aims at computing detailed 3D structural information to be used either in the understanding of the protein function or as a basis for rational drug design.

## 6.1

### Overview

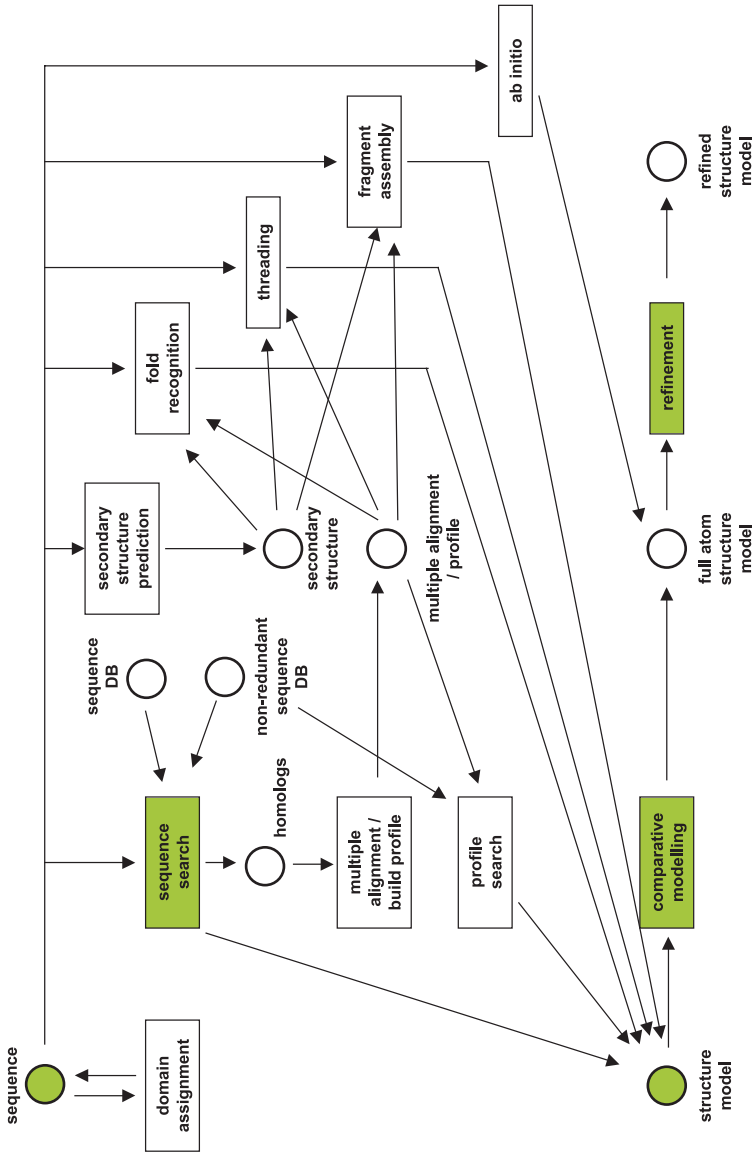
This Chapter concentrates on secondary structure prediction and on protein threading methods since, given current data and state-of-the-art of the field, these are the most promising approaches to tertiary structure prediction, fold recognition and remote homology detection on a large genomic scale. *Ab initio* methods are not covered here because, to date, they are less application relevant, with the exception of recent approaches combining local sequence-structure similarities with *ab initio* conformation sampling (see Section 6.2.5). Homology-based modeling based on an alignment or on a rough structural model is discussed in Chapter 5 of this volume.

Bioinformatics approaches to protein structure prediction and the identification of homology and similarities involve the solution of search problems. As the range of possible protein sequences and folding conformations is astronomical, appropriate pruning of the search space is inherently necessary. The so-called Levinthal paradox [4] states that even real proteins cannot try out all the possible conformations during the time they fold into

their native structure. One possibility would be to concentrate on local sub-problems and assemble the subsolutions to form overall solutions. Fortunately, proteins are composed of highly regular recurring elements (secondary structures) such as helices or strands. The secondary structures, in turn, form certain supersecondary structure motifs. Unfortunately, building up a protein structure locally is not possible directly, as short peptide segments of a protein can fold into different structures in different protein environments. It is immediately apparent that the 3D structure brings amino acids that are distant in the sequence in close contact in the fold. Such long-range contacts often may overrule local conformational tendencies and have to be dealt with, e.g. via conformational search approaches [5].

For the detection of remote homologies it may be required to investigate complex evolutionary and functional relationships. *Homologues* are proteins that relate back to the same common ancestor in evolution. Evolutionary relationship often points to similar structure and function. Finding close homologues, such as protein family members or related (*paralogous*, see also Chapter 5) sequences from other species can be detected via similarities exhibited with standard sequence search methods. Such methods include sequence alignment [6–8] and heuristic statistical search methods such as BLAST [9–13] and FASTA [14–18]. Current profile and multiple alignment methods, HMMs [19–28] and iterative searches [29] allow to identify more distant and insecure homology (see also Chapter 2 on sequence analysis in this volume). Sequence, structure and function analysis problems are closely intertwined, thus, so are the associated prediction problems. Structure prediction problems range from aligning a sequence to a sequence of known structure across threading the sequence to structural environments without significant sequence similarity, across mapping the sequence to structural templates according to contacts between residues within protein structures [30], to mere functional similarity based on indicative functional motifs [31]. Typically, biological problems have fuzzy, non-local aspects which are invariably highly interconnected. This is also the case for protein structure prediction, which apparently looks like a simple optimization problem, but turns out to be a knowledge discovery task exploiting heterogeneous information with efficient algorithms in the most discriminative and critical way.

The practical problem of performing as detailed as possible a prediction for a given sequence usually involves a procedure of analysis steps (Figure 6.1), so that results from early steps can and should be used in subsequent steps, e.g. predicted secondary structure can help for selecting fold sets or improve sequence–structure alignments, multiple alignments can help to both improve secondary structure prediction and fold recognition, functional predictions can provide further evidence for top scoring threading models. In order to deal with various kinds of information in the individual steps, it is helpful to provide appropriate means to store features and results of prediction steps. One such approach to represent and visualize



intermediate results and features in a uniform way and to exploit them efficiently in subsequent prediction programs is to use an extensible (XML-based) protein description language (ProML [32, 33]).

It is probably fair to conclude that most of the improvements on prediction performance are due to efficient use of more evolutionary information (homologs) in individual methods and/or due to clever processes of analyses [317] trying to mimic the approach of human experts, which is still the state-of-the-art [34] in protein structure prediction.

### 6.1.1

#### Definition of terms

*Ab initio approaches* try to perform predictions based on first principles and the input sequence alone. In their pure form, they do not use database-derived potentials, knowledge-based approaches or transfer of information from homologous proteins [35]. Recently, *ab initio* methods have been combined with fragment searching and threading to yield a new fragment assembly approach, which appears to be quite successful for small proteins [5, 36, 37]. In contrast to homology prediction and protein threading completely new folds that have never been seen before can be computed with *ab*

#### ◀ Fig. 6.1

Overview of possible steps of a manual or automated hierarchical prediction process from a sequence to a (refined) model structure. Given a sequence a search against a (non-redundant) sequence database can be performed using appropriate searching methods (FASTA, BLAST, GAP-BLAST, etc). If a significant hit to a sequence of a protein with known structure is found, a structural model can be inferred. If a simple sequence search fails an iterative searching procedure with intermediate (probabilistic) sequence profiles can be applied to find more distant homologs. Again if significant hits can be identified they can be used for predicting a structural model. Otherwise, the sequence can be partitioned into its (predicted) domains and the analysis performed for the individual parts separately. This is especially important for the subsequent structure prediction steps. A secondary structure prediction yields additional information on the protein structure, which can also be used for fold recognition, full threading, combined

threading *ab initio* prediction (fragment assembly), and even pure *ab initio* approaches. The used (non-redundant) databases of sequences and structures and representative sets of sequences, folds, classifications, have influence on the searching procedures and the statistical significance of the hits. The different procedures are described in the main text except for pure *ab initio* approaches and comparative modeling and structure refinement that are dealt with in Chapter 5 of this volume. What is omitted from the figure are functional predictions or knowledge as well as additional experimental information (e.g., on accessibility, secondary or domain structure, distance constraints, functional or biochemical network context), which can be used in various ways in sophisticated structure prediction methods either via validating predicted models or in the model generation process itself (see Section 6.4). Procedures represented by shaded boxes are discussed in other Chapters of this volume.



*initio* methods, in principle. However, an overall solution of the *ab initio* protein structure prediction problem seems to be still out of reach.

*Threading* aligns a protein sequence (the target) to the sequence of a known protein structure (the template) producing a sequence-to-structure alignment by optimizing some objective function. If this function contains only sequence related terms threading reduces to sequence alignment, if structural and structure environment information are also considered threading tries to optimize the compatibility of a given sequence with a given structure. If the scoring functions contains pairwise interactions between spatially distant amino acids the problem of computing an optimal sequence–structure alignment becomes NP complete [38].

There is a subproblem of threading called *fold recognition*. Here, we only ask for identifying a suitable template structure among a set of candidate structures. This problem is easier than threading, as it can be observed that the correct fold can be identified even on the basis of an inferior alignment. In addition, it is much easier to rate the quality of fold recognition, since the best fitting template structure can be identified more readily than scoring the quality of an alignment. This makes fold recognition a popular version of the problem for benchmarking purposes.

*Homology modeling* takes a sequence-to-structure alignment and computes a full atom model for the target sequence. It does so by modeling gaps in the alignment of target and template which most often represent loops on the outside of the target protein. In addition, gaps in the template have to be modeled and side chains that differ between template and target have to be inserted. Homology-based modeling ends with a refinement step and yields a full-atom model, in general (see Chapter 5 for details).

Homology-based protein structure prediction methods are the methods of choice, in practice. The reason is that the number of protein folds occurring in nature is much smaller than the number of protein sequences, as nature conserves successful structural architectures much more than the exact protein sequence. On the sequence level much variation is found in orthologous proteins from different organisms with same structure and function. Thus, if we are looking for a fold of a natural protein, it is much more economical to test the limited set of folds realized by nature than to search in an astronomically large space of potential protein structures.

*Secondary structures* [39, 40] are regular local structure elements identified in the 1950s [41, 42] which are characteristic abstractions for proteins and believed to be of great importance for the 3D folding. Secondary structure elements are defined via characteristic main chain torsion angles. Usually, three discrete states,  $\alpha$ -helix (H),  $\beta$ -strand (E, extended), or other (L, loop) are distinguished [1] (see Chapter 5 for details).

The *fold* of a protein is usually represented by its secondary structure elements and their arrangement in 3D space. According to current fold classifications, i.e., SCOP [43] or CATH [44, 45], there are several hundreds of different folds. Classifying proteins into a hierarchy from class, fold,

superfamily to family (SCOP [318]) or class, architecture, topology, and homology (CATH [319]), is not a trivial task, probably because the fold space of proteins is not easily partitioned into a hierarchy of distinct fold clusters.

*3D structural coordinates:* The protein is composed of a chain of amino acids, which in turn are composed of atoms. The position of these atoms can be inferred from experimental measurements, most prominently, X-ray diffraction patterns or NMR spectra. Therefore, a very detailed description of a protein is given by the exact  $(x, y, z)$  coordinates of all the protein's atoms. It has to be kept in mind, however, that structure coordinates are still an abstraction of the protein in its actual environment interacting with solvent and/or other macromolecules. This is due to experimental measurement errors, crystallization effects, and the static picture implicated by the coordinates. In addition, it is not necessarily true, that the most accurate or comprehensive information, say regarding protein function, can be inferred best from the most detailed structural representation.

*Remote homology detection* identifies, for a given sequence, one (*candidate recognition*) or all (*related pair recognition*) of the evolutionarily related proteins in a set of candidates. This may or may not include the computation of a detailed structural model. Interestingly, it is often possible to detect remote homology with the help of structural criteria without actually being able to compute a reasonable alignment and structural model.

### 6.1.2

#### What is covered in this chapter

This Chapter discusses protein structure prediction methods and their use for homology-based modeling (Chapter 5), docking and design of ligands (Chapter 7), structural genomics, drug target finding (see also Chapter 6 of Volume 2), and function assignments. As outlined in the introduction, this Chapter focuses on homology-based or knowledge-based methods rather than physics-based (*ab initio*) approaches.

*Secondary structure prediction* assigns a local helical (alpha) or extended (beta) structure to amino acid chains. The problem has been approached in an *ab initio* manner [46–48] for some time and many prediction methods have been developed based on properties of amino acids. Major advances have been achieved by employing homology-based approaches to 2D prediction. Such methods using sequence family information or consensus formation from several prediction methods are discussed in Section 6.4.1.

In general *3D protein structure prediction* is an unsolved grand challenge problem in computational biology. It has been studied for decades starting with the seminal experiments of Anfinsen in 1961 [3]. Turn-key solutions for the overall problem of computing the complete 3D structure from an amino acid sequence are not in sight for the foreseeable future. Due to the increasing numbers of sequence and structure data and the more and more

complete coverage of sequence (see genome projects for bacteria [320] and higher organisms, e.g. *C. elegans*, *D. melanogaster*, *A. thaliana*, Rice, Rat, Mouse, Human [321]) and structure space (structural genomics projects in the US [322], in Europe [323], and in Japan [324]) homology-based methods are already quite successful. The different methods used for homology based tertiary structure prediction are described in the Methods, Section 6.4.

Related problems are *protein folding*, *molecular dynamics*, *sequence design*, and *protein design*. The *protein folding* problem involves predicting in more or less detail the sequence of steps leading from an unfolded protein chain to its final structure. A lot of theoretical work has been done in this field [4, 49–59]. Especially, lattice models of proteins [60–64] and the associated folding algorithms have been studied extensively, though the practical relevance of the results for genomics and pharmaceutical applications remains limited. All of these so-called lattice models have in common a highly simplified view of a protein as a chain inside a regular 3D lattice. Residues are abstracted to nodes in the chain that come to lie on vertices of the lattice. Only few residue properties are distinguished and modeled. The solution to the full atom protein folding problem would also solve the structure prediction problem. *Molecular dynamics* studies what happens to a protein structure under varying conditions, such as increasing temperature, unfolding, or interaction with other molecules (see Appendix). *Sequence design* asks for optimized sequences to fit a given structure or fold (it is also called the inverse folding problem), i.e. sequences which assume the given fold and have additional modified features such as a modified stability or active sites. *Protein design* involves rational engineering of a protein with a specific new fold or modifying an existing fold by changing certain amino acids to attain new or modified functions [65–69].

The rest of this Chapter is restricted to methods for homology-based protein structure prediction given a protein sequence that occurs in nature. The difference in dealing with natural and designed proteins is an essential one: natural sequences have evolved to fulfill a certain purpose and function. Evolution has challenged certain parts of the sequence and structure design and conserved the necessary ingredients. Traces of this evolutionary history can be detected and used for predictions within the territory of folded and functional proteins. Predictions can thus be made by similarity transfer: similarity of certain features can be analyzed with care to predict similarity of implied features. Such similarities are not likely to occur in the same fashion in artificially designed proteins.

Homology-based protein structure prediction in concert with experimental high-throughput methods for protein structure prediction within a global structural genomics effort might well effectively solve the prediction protein structure prediction problem before a satisfying solution to the protein folding problem is in sight. Evidence for this perspective is that progress in the protein structure prediction field is rapid and can be expected at an accelerating pace simply because of the amount of data gath-

ered both on sequences and structures. This will be even more the case as dedicated structural genomics projects [325] have started with the goal of charting the space of naturally occurring protein folds. Together with improved methods of protein classification a detailed map of the protein structure space will become available, which allows to assign a structure to a given sequence on the basis of the wealth of structural folds and their relations as well as the associated sequences already assigned to those structural classes.

In the following, we first describe the input data on which the protein structure problem is based to generate the output data (Section 6.2). Section 6.3 reviews the principal methods of structure prediction. Section 6.4 discusses the results obtained by structure prediction methods in the context of various application areas. Ways to assess and validate prediction results and the strengths and limitations of the overall approach are dealt with in Sections 6.5 and 6.6.

## 6.2

### Data

#### 6.2.1

##### Input data

In the simplest case the problem instance is defined by the amino acid sequence (*the target sequence*) only.

More detailed instances of the problem contain the available folds to be used as structure templates and the set of all available resolved 3D structures; the available sequence database and the sequences that are evolutionarily related to the target sequence, possibly including a multiple alignment of this protein family. It can also be helpful to distinguish between orthologous and paralogous members of the protein family. Today, tools such as PSI-BLAST (see Chapter 2) are used to locate the involved protein family, to collect the members of the family, and to generate the associated multiple alignment and/or sequence profile. Similarly, for each of the structurally resolved proteins from the set of templates evolutionary and/or structurally related proteins can be collected in order to determine the respective protein neighborhood and multiple alignment.

In addition, the availability of experimental or predicted secondary structure information on the query sequence as well as other experimental constraints such as knowledge on disulfide bridges, on distances or distance intervals between residues and functional groups within proteins and on accessibility of side chains could have large impact on the prediction methods and results as well as on the confidence in the predicted model structure.

In general, providing the information accompanying the protein sequence or any of the fold templates is subject of dedicated research and the quality of the structure prediction heavily depends on the quality of these derived input data.

### 6.2.2

#### **Output data**

The simplest output is an alignment of the query sequence onto the best (closest relative) template structure. Such an alignment is the starting point for several analysis and modeling steps.

A comparison of the alignment score of the target sequence with (many) different template proteins can be the basis of rating the significance of the produced structure model. *P*-values are a version of this approach that also applies to sequence analysis methods (see Chapter 2 of this volume), for which it has been studied extensively [9, 11, 70, 71]. In addition, the resulting models can be subjected to a wide variety of plausibility tests in order to ascertain their protein-likeness. Many of these checks will be made on the full-atom model of the protein that results from the procedures described in Chapter 5 of this volume.

Additional output data could be predicted secondary structure and accessibility annotation for the query sequence (see Section 6.3.1, [72]), a multiple alignment of related sequences and structures, structural models, and variance and mutation analysis of closely related proteins.

The following paragraphs list additional data resources relevant for structure prediction approaches. For knowledge-based methods it is essential to make the most efficient use of the relevant information to come up with a plausible prediction. Doing so seems to be the most promising approach to the problem.

### 6.2.3

#### **Additional input data**

Most protein threading methods rest on the input data mentioned in Section 6.2.1. In order to improve protein structure prediction we can benefit significantly from a wide variety of additional information that is contained in biological databases. We already mentioned the protein family information that is contained in a growing number of databases which exist both on the sequence [73–76] and structure level [43–45, 77–82]. This information can be used to produce profiles for both the target protein sequence and its family and the template fold database. Another approach is to exploit conserved motifs [31, 83, 84] and functional patterns [85–87] for performing the structure prediction, either via a post-processing step eliminating fold

candidates not compatible with necessary functional constraints, or via the selection of candidates or partial alignments on the basis of searching for motifs and patterns. This approach has been followed, for example in RDP (see Section 6.3.2, [32]).

An even more involved and innovative approach is to use knowledge or predictions of functional information on the protein and network information in the form of biochemical, e.g. metabolic and regulatory networks. Here the information that a set of proteins is contained in a certain pathway could provide additional constraints for candidate structures fitting into certain locations of that pathway, assuming that additional information on the functional and/or structural constraints of these pathway members is available. Again, such information can be used to select plausible folds to thread against, to filter conspicuous candidates from the ranked list of the prediction, or to use associated constraints in the prediction step itself [32]. In particular, networks generated from experimentally established (e.g. via large-scale yeast2hybrid screens [88–91]) and/or predicted interactions [92–96] of proteins can provide structural constraints to be of use for homology-based structure prediction.

In the next two subsections, we describe in more detail two of the main non-algorithmic ingredients of structure prediction methods, namely the structure classifications that represent something like the global view of protein structure space, and the scoring function rating a sequence–structure alignment. In many threading methods, these components are chosen by default and applied to all protein target sequences. However, a threading method can be improved by specially tailoring these components to the target sequence.

#### 6.2.4

##### **Structure comparison and classification**

In several ways, protein structure prediction methods rely on classifications of the known protein structures that are based on systematic comparison of the 3D structures. First, from such classifications overall principles constituting different protein folds can be inferred, which can be of use in protein structure prediction algorithms and/or associated scoring systems. Second, representative subsets of folds can be constructed which are exploited as the universe of possible structural templates for sequences to be predicted, e.g. as a list of folds against which to thread the sequence. Third, such classifications can be used to discriminate between the available choices. I.e. it is not only necessary to identify a probably close fold but to identify those folds that are the most probable models for the target sequence among all available fold types. Fourth, structure classifications are used as a standard-of-truth in benchmarks for fold recognition methods. Finally, they aim at providing a complete overview of the protein structure universe.

Structure comparison methods are a way to compare three-dimensional structures. They are important for at least two reasons. First, they allow for inferring a similarity or distance measure to be used for the construction of structural classifications of proteins. Second, they can be used to assess the success of prediction procedures by measuring the deviation from a given standard-of-truth, usually given via the experimentally determined native protein structure. Formally, the problem of structure superposition is given as two sets of points in 3D space each connected as a linear chain. The objective is to provide a maximum number of point pairs, one from each of the two sets such that an optimal translation and rotation of one of the point sets (structural superposition) minimizes the rms (root mean square deviation) between the matched points. Obviously, there are two contrary criteria to be optimized: the rms to be minimized and the number of matched residues to be maximized. Clearly, a smaller number of residue pairs can be superposed with a smaller rms and, clearly, a larger number of equivalent residues with a certain rms is more indicative of significant overall structural similarity.

Analogously to the structure prediction problem, for the structure comparison problem the first step of matching points in both sets is the crucial and quality-determining part. In fact, the second step of finding an optimal superposition of a given set of matched pairs of points can be solved efficiently and optimally [97–99]. For the first step a broad spectrum of methods has been proposed over the years: simple alignment, matching of features [100, 101], compactness and recurrence criteria [102], secondary structure matching (with or without obeying the chain order) [103, 104], to geometric hashing techniques completely ignoring the chain information of the residues [105–107]. For assessing the quality of 3D structure superpositions, besides the rms, the so-called soap film method of comparing protein structures [108] has been proposed. This method measures the distance between structures via the size of the area between the associated 3D curves of the protein backbones. In addition, assessment procedures for homology models have been developed during the CASP experiments. These can be classified into sequence dependent and sequence independent variants [109–111]. The former use the predicted alignment to identify equivalent residue pairs and compute the rms of the associated optimal superposition of model and experimental structure. The latter determines an appropriate set of matched pairs, given the model and native structure coordinates. These pairs are used for the superposition. The latter approach is a bit more tolerant to alignment errors, especially when an overall correct fold has been identified via a (partially) faulty alignment.

As mentioned above, several of these approaches have been employed to generate exhaustive clusterings of the protein structures into structural classes available via the resulting databases (DDD/FSSP/Dali [102], 3Dee [79], JOY/HOMSTRAD/DDBASE [112–114], structure cores LPFC [115]), These superposition-based classifications are complemented with classi-

fications based on other criteria (CATH) such as topology and orientation of secondary structure elements up to manual classification from visual inspection of the folds (SCOP). A recent systematic comparison of these classifications has been carried out by Hadley and Jones [116]. They suggest selecting a somewhat redundant larger set of threading templates that is compiled from clusters of all individual classifications and to base the evaluation on those examples on which all the classifications agree. Generally, the classifications agree on a large part (about two-thirds) of the protein structure data. There are some systematic inconsistencies among the different taxonomies due to the different approaches taken.

From classifications representative lists for threading as well as assessment criteria for prediction results can be obtained. In addition, involved prediction methods can take advantage of a global view of protein structure space to better discriminate between the available alternatives than can be done on alignment scores of a sequence to an individual fold alone. We expect more comprehensive global knowledge of protein structure space to significantly improve protein structure prediction accuracy, in the future.

**Domain assignment:** domains are basic building blocks of protein 3D structures. Several algorithms have been proposed to identify domains in a protein structure using criteria such as physico-chemical properties of amino acids in certain regions, e.g. hydrophobicity and accessibility, compactness of protein parts and their recurrence in other proteins [116–120].

For threading, often representative sets of protein structures are selected from domain fold libraries. Threading algorithms then align the sequence under consideration against a library of domains in order to detect the domain(s) constituting the unknown sequence. Due to the small signal-to-noise ratio, the performance could be increased by splitting the sequence into putative domains before threading them against the domain library. This can be done on the basis of the sequence via dedicated domain prediction methods [121–123] or by approximately matching the length of the respective structural domain in a straightforward manner.

### 6.2.5

#### **Scoring functions and (empirical) energy potentials**

A standard common assumption is that native structures of proteins are those conformations with minimal free energy [2, 57, 58]. Free energy cannot be determined exactly: many factors have to be taken into account, not all relevant factors are known, neither are their respective weightings. Threading has to estimate alignment conformations between sequence and structure and rate different alignments against each other. For this purpose it has to be taken into account that native structures and the model struc-



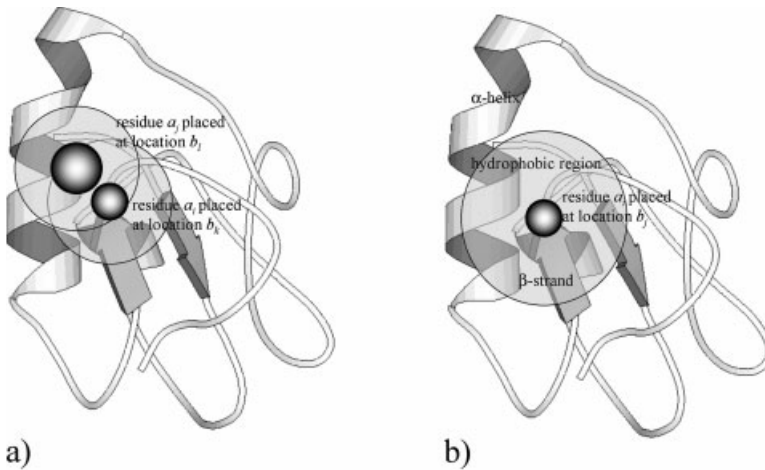
tures will be more or less different. Therefore, the energy evaluation via the scoring function has to keep a balance, in order to account for the expected differences of individual proteins and, at the same time, provide the necessary detail to achieve sufficient discrimination between different folds. In addition, almost always only partial protein models can be evaluated (e.g. due to incorporating the backbone conformation but no side-chain conformations, missing loop regions, or gaps). Usually the model and the template structure are different enough to prohibit the application of detailed atom-level physico-chemical potentials. Therefore, knowledge-based, i.e. database-derived potentials on different levels of abstraction have been successfully computed via database analysis and inverse Boltzmann statistics [30, 124] and applied to threading optimizations.

Inverse Boltzmann statistics rely on the relationship between the energy of a molecular state and the probability with which it is observed. This relationship is formulated in the Boltzmann law. Usually, the law is applied to derive the probability of observing a molecular state from its given energy. Here, we apply the law in the inverse direction. I.e., we count the frequency of observing a molecular state (or partial state) and derive from that a pseudo-energy of this state. We call the partial molecular states whose frequencies we are counting *events*. Event classes can be freely defined. We will give a few examples below. We collect the statistics on events by scanning through a representative set of protein structures. Using the Boltzmann law we deduce the contribution that this event makes to the total pseudo-energy of the protein structure. Contributions of all observed events in a protein structure are added up to yield the pseudo-energy of the protein structure.

This procedure is applied ubiquitously in protein structure prediction. Despite the caveats that the Boltzmann law is only applicable in thermal equilibrium and that it cannot readily be applied additively to partial molecular states scoring functions based on inverse Boltzmann statistics have proved very valuable in protein structure prediction.

#### 6.2.5.1 Pair-interaction potentials

Essentially there are two types of potentials that cater to two different sets of optimization methods. The more accurate potential functions score pair-interactions between amino acid residues in the protein (see Figure 6.2a). The residue pairs are grouped, for instance by the type of the two residues, their distance in the protein sequence, and their distance in the protein structure. E.g. residue A and L occurring at a distance of 6–15 residues apart in the protein sequence and at a distance between 3 Å and 6 Å in space may form such a group. The overall score of a sequence–structure alignment is the sum of all contributions determined by the inverse Boltzmann law of pairs of amino acids observed in the structure of the target sequence that is obtained by aligning the target sequence to the template structure as the alignment prescribes (see Figure 6.2). This kind of potential has been introduced by Sippl [30, 125].



**Fig. 6.2**

Pair potentials vs contact capacity potentials a) a pair potential attributes an energetic contribution to each pair of amino acids  $a_i$  and  $a_j$  that are mapped by a sequence-structure alignment onto locations of the amino acids  $b_k$  and  $b_l$  of the protein template structure. The contribution depends on the types of residues  $a_i$  and  $a_j$  as well as on geometric features of their locations  $b_k$  and  $b_l$ , such as their distance in space or in sequence. Often, only certain pairs of amino acid residues are scored, such as those that are contacting or close in space. The total energy is computed as the sum of all pair contributions. b) a contact capacity potential rates the energetic contribution of an amino acid  $a_i$  from the protein target

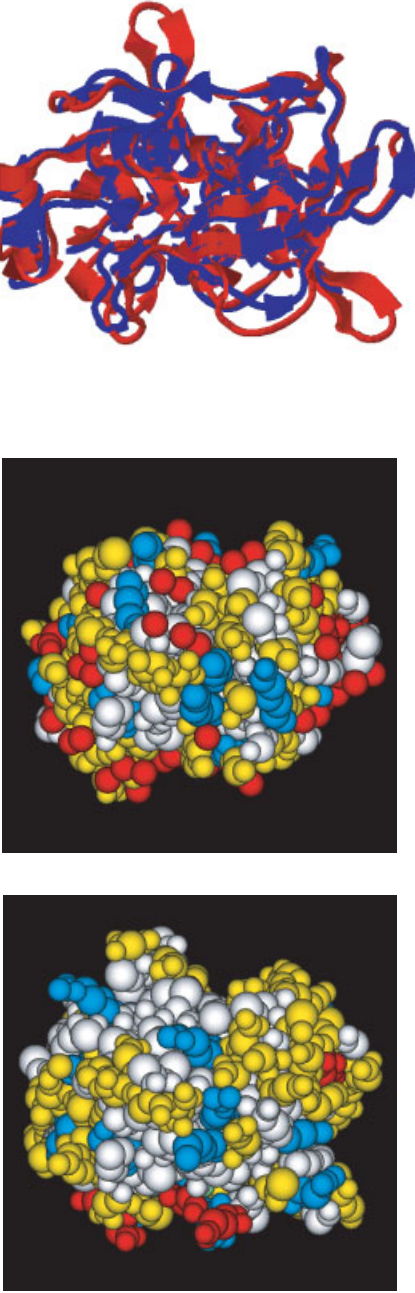
sequence A that is mapped onto a location  $b_j$  inside the protein template structure B. The energetic contribution is determined by the type of residue  $a_i$  and the chemical features close to the location  $b_j$  inside the protein template structure B. In the case of the figure,  $a_i$  comes to lie at the tip of a strand and next to a helix inside the protein core in a hydrophobic region. These pieces of information can be assembled in a fingerprint that characterizes the environment of location  $b_j$ . Note that the contact capacity potential is derived from the chemical makeup of the template protein exclusively. In both variations of potentials, the contribution of residue  $a_i$  as mapped onto location  $b_j$  is determined via inverse Boltzmann statistics.

Today, there is quite a variety of such potentials [125–132]. The performance of potentials for different tasks in structure prediction varies. Reviews and comparisons of database and molecular mechanics force fields as well as a discussion of their controversial relationship to free energy can be found in [133–135].

The involved alignment optimization problem, i.e. the problem of finding a sequence–structure alignment with minimum pair interaction score is NP-complete [38].

### 6.2.5.2 Contact capacity potentials

Because of the computational problems a different set of potential functions has been proposed that allows for more efficient optimization algorithms.



a) Elastase

b) Trypsin

c) Superposition of the backbones of elastase (red) and trypsin (blue)

**Fig. 6.3**

This figure shows that two proteins that have very similar structures can have widely differing chemical makeup. a) shows a full-atom model of porcine elastase (1est.pdb, 240 amino acids). The atoms are colored according to the chemical composition of the respective residue (red = positively charged, blue = negatively charged, yellow = polar, white = nonpolar). b) shows the same

picture of bovine trypsin (1tpp.pdb, 223 amino acids). Elastase and trypsin are both serine proteases and have a sequence identity of about 36% and 39%, respectively. c) shows a superposition of the backbones of elastase and trypsin, revealing that their structures are practically identical (222 superposed residues with an rms of 2.6.9). Still, the chemical composition revealed by the coloration of a) and b) is quite different.

Such potentials are sometimes called profiles, as the structure is abstracted into a linear representation (the profile) of positional score values for the 20 amino acids at the respective position. A profile does not explicitly score the energy of pair-interactions in the resulting protein structure. Rather the idea of the profile is based on the hypothesis, that the protein target sequence inherits not only the backbone conformation from the template structure but also the chemical composition. Then, e.g. we can assume hydrophobic patches in the target protein where they occur in the template protein, H-bond donor regions are preserved as are polar regions etc. (see Figure 6.2b) If we assume this hypothesis, then we have to score not pairs of amino acids but single amino acids, each against a given and constant chemical environment. Aligning, say, a residue  $a_i$  in the target sequence against a residue  $b_j$  in the template sequence would be scored according to how often a residue of type  $a_i$  is found within the chemical environment that the template protein structure presents for residue  $b_j$  (see Figure 6.2, page 252). The alignment optimization problem now becomes much simpler. Essentially, the sequence alignment methods described in Chapter 2 can be used. We only have to replace the amino-acid substitution matrix used for scoring in sequence alignment by the threading profile. Thus alignment optimization can be done with dynamic programming in polynomial time (see Section 6.3.2). In practice, in this case several optimal alignments can be computed per second on a workstation.

Of course, the hypothesis that the chemical composition be identical in the template and the target protein, is often false (see Figure 6.3). Thus one has to apply both kinds of potential functions in concert to make suitable structure predictions (see Section 6.3).

Bowie et al. first introduced profile scoring functions for protein threading [136]. They consider the type of residue and geometric aspects of its environment such as inside/outside or the secondary structure element in which the residue occurs. Later different variants of this scheme have been developed. They essentially differ in how they describe the chemical environment of the residue in the template structure. Contact capacity potentials (CCPs) are a variant used in the threader 123D (Alexandrov et al [137]). CCPs essentially count contacts to neighboring residues, irrespective of the type of residue. The 2D structure environment is considered in a version of CCPs and contacts are distinguished with respect to whether they involve residues that are close (short range) or far apart (long range) in the protein sequence. Effectively, contact capacity potentials aim at approximating generalized hydrophobicity measures and secondary structure preferences, at the same time.

### 6.2.5.3 Contacts in protein structures

Many potentials of both types rely on a particular definition of residue-residue contact. Starting with Sippl [30, 125] many authors have used the

distance between residue pairs to classify contacts. Distances below a certain threshold (e.g. less than 7 Å for the relevant  $C_{\beta}$  atoms) constitute a contact. This definition is debatable and often contradicts the actual geometry of a protein structure. E.g. a third residue can be located between both residues even though they are at close distance. We have introduced a more appropriate notion of residue-residue contact [138]. Here, residues are assumed to be in (direct) contact only if they share a contacting face of a certain minimum area. These *direct* contacts and the associated contact areas are determined via a so-called Voronoi tessellation of the protein. A Voronoi tessellation is a decomposition of the space occupied by the protein into so-called Voronoi cells, one cell for each atom. The Voronoi cell for an atom is composed of those points in space that are located closer to the particular atom than to all other atoms. There is analogous variant of Voronoi tessellation that is based on complete residues instead of single atom. An alternative tessellation on the residue level can be obtained by using the union of atom cells belonging to a residue as the representative for the residue cell. The potentials resulting from Voronoi tessellations have been demonstrated to improve the performance of fold recognition methods [138].

#### 6.2.5.4 Calibration of the scoring function

The definition and calibration of appropriate scoring contributions capturing relevant features of protein structure and sequence-structure compatibility has started with Margaret Dayhoff's derivation of her famous substitution matrices [139]. Various other attempts have been made to define better scoring terms for sequence compatibility [140–144], secondary structure preference and contact potentials (see above). Classification and calibration techniques such as linear programming, supervised learning, support vector machines have been used to systematically optimize substitution matrices [145], the discrimination of scores [146] and the relative weights of different individually optimized scoring contributions among each other [147–149]. E.g. the fold recognition performance of the ToPLign 123D\* threader using calibrated weights of 18 scoring contributions (sequence, secondary structure, gap insertion and extension, local and global contact capacity, each weighted individually in three secondary structure elements) has been improved by more than 40% as compared to the original 123D.

### 6.3 Methods

This section gives an overview of secondary structure and tertiary structure prediction methods for proteins. Methods for (secondary) structure prediction of other biomolecules, such as RNA are not discussed here.

## 6.3.1

**Secondary structure prediction**

Secondary structure prediction assigns a local structure to amino acid chains. Usually, local structure is characterized by one of three discrete states, helix (H), strand (E for extended), or other (C for coil). This measure of evaluation is also called three-state (Q3) measure. The problem has been approached first in an *ab initio* manner involving the target protein sequence without any additional information [48, 150–154], e.g. on the relevant protein family or homologous sequences, and many prediction methods have been developed based on properties of amino acids. These methods yield 60–65% performance on independent benchmark sets. Major advances have been achieved by employing a homology-based approach to 2D prediction. PhD [72, 155] exploits knowledge on homologous sequences by using multiple sequence alignments and the associated profiles of secondary structure assignments for training a neural network. This procedure does not need knowledge of the tertiary structure of proteins that are related to the target sequence. Such approaches profit significantly from the increased number of available sequences and, thus, larger and better multiple alignments, but also from much improved search procedures to recruit homologous sequences for the training. The collection of homologous sequences is done via iterative profile-based search methods, such as PSI-BLAST (see Chapter 2). A secondary structure method based on this approach is PsiPRED [156] achieving around 80% of correct three-state predictions on benchmark sets and on selected blind-prediction examples [157].

The second improvement in this area is the use of more involved consensus prediction methods. With a range of different methods at hand, it is possible to derive a consensus prediction as has been done in SOPMA [158], JPRED [159, 160], or CODE [161] by systematically optimizing the performance on a training set by using decision-tree methods or machine-learning approaches such as support-vector machines [162].

Having an improved secondary structure prediction available with, say, more than 70% expected correctness can improve 3D prediction. There are several approaches: such as matching the predicted secondary structures with the template structures [163, 164], using secondary structure prediction as a scoring term to evaluate sequence and threading alignments [147] or incorporating a respective scoring term into the objective function of a threading algorithm. From an algorithmic point of view all alternatives are easy to realize with negligible computational overhead. The evaluation of alignments requires training a neural network using an additional scoring term. The optimization involves adding another profile term to the objective function. Nevertheless, the use of secondary structure predictions generally improves the performance of remote homology detection as well as alignment quality [147, 165].

On the other hand, for really challenging instances of fold recognition the approach cannot always help: if no homologous sequences can be found, the secondary structure predictions tend to be of poorer quality; and using wrong secondary structure assignment can disturb the scoring function and mislead the alignment process.

**Assessment of secondary structure predictions:** these methods involve statistical analyses of applying the methods to known structures and recomputing their secondary structure as well as blind predictions (see also Section 6.5.1).

A state-of-the-art assessment of predictions requires an up-to-date set of secondary structure assignments and a measure to compare predictions with the correct assignments. Evaluation sets of non-homologous domains with secondary structure assignments have been proposed by Barton et al. [166, 326]. This dataset is an extension of other proposed test sets [155, 167]. It contains 496 non-homologous (SD cutoff of 5, which is more stringent than mutually less than 25% sequence identity) domains. The database contains 82 847 residues, 28 678 alpha-helix (H), 17 741 beta-strand (E), and 36 428 coil (C) residues.

The standard measure for accuracy is Q3 defined as the number of residues correctly predicted divided by the number of residues. Q3 can also be computed for the residues of individual proteins Q3P or of individual secondary structures QH, QE, and QC. Other methods use certain types of correlation coefficients or take the type and position of secondary structure elements into account (proposed by Rost [168] and further developed for the CASP experiments [169]). One example is the SOV (segment overlap) score, i.e. the average over all secondary structure segments of the ratio of the overlap of predicted and true secondary segment and the overall length of the two segments.

More measures and programs to compute some of these measures for comparison and evaluation of new methods can be found on the CASP website [327] and a summary of measures can be found at the EVA site [328].

### 6.3.2

#### **Knowledge-based 3D structure prediction**

*Knowledge or similarity-based methods* exploit similarities between a given sequence and (sequences of) proteins of known 3D structure: the underlying assumption is that the more similar the two sequences, the more similar are the two corresponding 3D structures. The similarity is often generalized to additionally include quasi-physical properties, such as preferences of amino acids to occur in certain states, e.g. buried inside the protein or

solvent-exposed at its surface. In general, similarity approaches try to align a given sequence to 3D structures from a library of structures (or folds) of known proteins, such that an alignment score quantifying the similarity is optimized. These search procedures find the fold from the library that is most compatible to the sequence in question.

Depending on the type of the scoring function, fast dynamic programming methods can be used to find optimal alignments or heuristic search procedures have to be employed to find approximate solutions. In practice, a two-step approach [170] can be applied to first identify a small set of candidate folds out of a large library using a simple scoring function and fast threading tool (e.g. CCP and ToPLign/123D) and then use a more detailed scoring function and a more involved alignment algorithm (pair-interaction potentials and ToPLign/RDP) to compute more accurate alignments and to distinguish among the remaining template candidates.

If there is significant structural similarity between the query sequence and proteins in the fold library used for threading, similarity searching followed by homology modeling is the method of choice, as it can yield plausible structures quickly. If the considered protein has a novel fold, i.e. a fold not yet represented in the fold library, the method will not be able to come up with the new fold as a structural model for the sequence, but at best can predict that none of the folds in the library applies.

The following section discusses methods for computing sequence-structure alignments for both types of scoring functions.

### 6.3.2.1 The algorithmic viewpoint on sequence-structure alignment and threading

In general any threading of a sequence of length  $n$  onto a structure with  $m$  residues can be represented by a simple alignment of the two sequences. As is well known the number  $f(n, n)$  of alignments of sequences of length  $n$  is finite but huge [171]:

$$f(n, n) \sim \frac{(1 + \sqrt{2})^{2n+1}}{\sqrt{n}}, \quad \text{as } n \rightarrow \infty.$$

Therefore, exhaustive enumeration of possible alignments followed by an evaluation of individual alignments is prohibitive. Fortunately, for scoring functions that take the form of threading profiles the optimal alignment among the exponential number of possible alignments can be found efficiently with dynamic programming methods as in sequence alignment (see Chapter 2 of this volume). Depending on the gap cost function, threadings with profiles can be computed in quadratic (linear and affine gap penalty functions) with the Gotoh [6] or in cubic time (general gap costs) with the Needleman and Wunsch [7] algorithm.



The protein folding process and its determining factors are still poorly understood. But even if all factors could be modeled appropriately, folding simulations can span only a tiny fraction of the actual time required to fold a protein which is on the order of milliseconds to seconds. In addition, the contributing factors need to be modeled extremely accurately in order to avoid error propagation during the vast amount of computation needed for completely folding a protein. Also the structure prediction problem appears to be difficult [172–174].

It is generally assumed that contact interactions between amino acids or their groups are important stabilizing contributions to protein stability. Such interactions preclude the use of dynamic programming methods for the optimization (see Section 6.2). In fact, since the involved optimization problem is NP-hard, optimal solutions are generally not computable within reasonable computing time and resources. Nevertheless, there are some approaches trying to do so, despite the fact that optimal solutions are likely to turn out not to be of much practical value due to the deficiencies of the scoring function, e.g. [175].

In contrast, knowledge-based discrimination methods can be often successful. This section discusses the two major approaches to knowledge-based 3D structure prediction: purely sequence based methods, and methods exploiting structural knowledge. The latter approaches can be classified into sequence methods using secondary structure information and methods using 3D structural templates (threading). An *ab initio* approach somewhat related to threading is fragment assembly, where fragments are identified via local sequence and structure conservation and then these fragments are assembled into known or new folds thereby minimizing some energy function [5].

Other theoretical structure prediction methods, such as simulating folding pathways, lattice models, simulations and predictions, inverse folding (i.e. designing the “best” sequence for a fold/structure), physical methods (based on finding minimum energy conformations for “realistic” physico-chemical potentials), are not discussed here.

#### 6.3.2.2 Sequence methods

Structure prediction can be achieved by considering sequence information alone: given a sequence a search is performed against a fold library by comparing the associated sequences of the folds with the query. If a significant hit is found, the fold is predicted as a model for the query sequence. The sequence alignment can be used to actually build a model from the identified template structure. Sequence alignment searches are certainly the methods most relying on homology information and profit a lot from the increase in sequence and family information. Therefore, sensitive sequence profile [176–181] search methods such as PSI-BLAST [9] and intermediate sequence search using, in addition, information from vast sequence

databases are competitive protein structure methods. In order to predict a structure, PSI-BLAST is used to search all sequence databases to collect related sequences, to compute a multiple alignment, to derive a consensus profile from it, and re-search the databases using the profile instead of the query sequence until a protein of known structure is hit. An E-value is used to indicate the confidence in the hits found.

Other successful structure prediction methods based on sequence alone are FASTA [15, 18], hidden Markov models (HMMs) [25, 26, 28, 179], intermediate sequence search [182], and iterative profile search [29]. Sequence analysis methods are discussed in detail in Chapter 2. The results of such methods in the CASP experiment are described in [157, 183, 184].

### 6.3.2.3 Sequence and secondary structure

Experimentally determined or predicted secondary structure information can successfully be phased into sequence-based methods for structure prediction. GenThreader uses profile sequence alignment and various other criteria evaluated on the resulting alignments including compatibility of predicted and template secondary structure using a neural network to come up with a confidence measure for the alignment [147, 157]. Koretke et al. [184] used consensus secondary structure predictions to align predicted and template secondary structure elements using the program MAP [164].

### 6.3.2.4 Tertiary structure-based methods

Structure-based methods are reviewed in [185] and can be classified into two types: profile methods based on dynamic programming (see Chapter 2 of this volume) for computing optimal alignments of structural environments and threading approaches using full pair-interaction potentials [30, 125, 138, 186] (see Section 6.2). The latter methods use optimal exhaustive search procedures [175], branch and bound approximation algorithms [187, 188] or other heuristic approaches for the optimization [189, 190]. Reviews and comparison of threading methods can be found in [186, 191, 192].

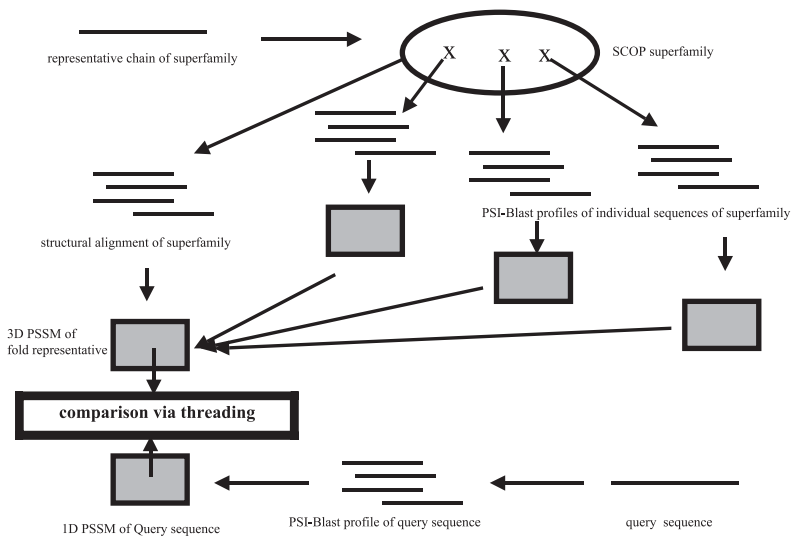
**Profile threading methods** As described in Section 6.2, algorithms that optimize threading profiles are the same as for pairwise sequence alignment and are based on dynamic programming. These methods have been introduced by Eisenberg et al. [136, 193, 194].

Several variants using multiple sequence information and sequence profiles generated for the query sequence for the threading process have been described and demonstrate to improve fold recognition performance [109, 195–203] similarly as has been observed for secondary structure prediction and sequence methods.

Combining secondary structure prediction with profile threading [165, 204] has been demonstrated to improve threading performance as well.

As methods based on dynamic programming cannot account for pairwise-interaction potentials the so-called frozen approximation approach [189] has been proposed. This method performs several iterations of profile environments. In the first iteration, the chemical environment is defined via the contact partners of the template. In subsequent rounds the aligned residues from the previous iteration replace the residues of the template. The idea is that target and template structure are similar enough such that the iterative process converges towards the optimal assignment.

A method employing 3D structural profiles for detecting remote homologies has been proposed by Sternberg et al. [205]. This method, called 3D-PSSM (three dimensional position specific scoring matrix), uses sequence profiles to search databases. The profiles are computed from multiple sequence alignments using the standard method used for generating PSI-BLAST profiles [9, 206]. In order to use structural information, the following approach has been proposed (see Figure 6.4): for any protein  $f$  in the fold library, determine the set  $S$  of related structures in the superfamily. For any sequence  $x$  in  $S$  compute the PSI-BLAST profile via a sequence



**Fig. 6.4** Overview of the 3D-PSSM approach for protein structure prediction: For any protein (superfamily) PSI-BLAST searches and profiles are computed and using the structural superposition of the (structurally related) family members. On the basis of

the supersposition a structural profile representing the family is compiled. Given a query sequence, a corresponding sequence profile is computed, again using PSI-BLAST, and both profiles are aligned using a dynamic programming based profile threading method.

search of  $x$  against an appropriate database yielding a profile  $P_x$ . The profiles for the sequences in  $S$  are then combined using the structural alignments for the superfamily and the implied equivalences of residues. This is possible, despite the fact that the proteins in  $S$  do not necessarily show a reliable sequence similarity. The reason is that these proteins are structurally superposable and thus reliable residue equivalences can be determined. The idea behind the approach is to use all the information available for a group of structurally similar proteins in order to do the sequence structure (threading) search. The fold library and the associated profiles can be computed beforehand such that the search for a query protein can efficiently be done via computing a sequence profile and matching this profile against all 3D-PSSMs from the library using standard dynamic programming methods. During this search, in addition, solvent accessibility and secondary structure preferences are evaluated. The secondary structure prediction is performed with the program PsiPRED [156]. Thus, the approach successfully combines evolutionary and structural information available in the sequence and structure databases using the most advanced search (PsiPRED) and prediction methods (PsiPRED) as well as established profile building methods (PSI-BLAST) and structure classifications (SCOP). The major disadvantage of the method appears to be that there is probably not one single profile representing the properties of any individual member of a protein superfamily. Nevertheless, the performance of the method has been shown to improve on sequence search alone on a benchmark of 136 homologous protein pairs that cannot be detected with PSI-BLAST (see [205] for details of the benchmark construction and the results obtained). Here, 3D-PSSM was able to detect 18% of the pairs. The program is available as a web service at the Biomolecular Modeling Laboratory of the Imperial Cancer Research Fund [205]. In the latest CASP-4 and CAFASP-1 experiments both this approach and the associated automated prediction server were among the most successful groups.

**Optimal threading with full pairwise/interaction potentials via exhaustive enumeration** Usually, algorithms aiming at optimal solutions address the fragment threading problem: here, the structure is partitioned into a number  $k$  of secondary structure elements, which are considered fixed fragments of a definite length where no gaps are allowed in any threading alignment. The problem reduces to finding a mapping of sequence positions onto the first positions of all the fixed fragments of the structure. This mapping has to observe the linear ordering of the sequence residues, the structure fragments and the loop length constraints. The basic assumption behind this model for sequence–structure relation is that the structure is composed of important and irrelevant parts, the important parts are the fragments, which have to be aligned to constitute the essential parts of the fold and the irrelevant part are loop regions, which are usually not conserved even among similar folds and,

thus, need not be aligned and the alignment score is not taken into account for the overall alignment score.

The fragment threading problem has a much smaller search space of size exponential in  $n$  and  $k$  instead of  $n$  and  $m$  for protein threading. In some cases, for small proteins or proteins with a small number of fragments an exhaustive enumeration of all mappings of sequence regions onto fragments is possible. Therefore, the optimal threading can be found for almost any conceivable scoring function, i.e. scoring functions that depend on the complete alignment. Nevertheless, to find meaningful threadings capturing shared biological and/or structural features in a true sequence-structure relationship between the two proteins, the scoring function has to model the respective features very accurately. As of yet, no scoring function appears to be accurate enough to discriminate between similar alignments and to select the 'correct' alignment from a large number of alternative alignments. For a broad class of empirical pair interaction potentials it has actually been demonstrated that almost always there exist alignments in the vicinity of the structurally closest alignments, that score better than the standard-of-truth according to the scoring function used. Such alignments are easily found via a greedy search of the alignment space or some simple perturbations of the correct alignment using a genetic algorithm [207].

**Branch-and-bound algorithms for fragment threading** An algorithm for finding the global optimum of gapped threadings between a query sequence and a structural core of consecutive fragments (fragment threading [208]) has been described by Lathrop and Smith [175]. The fragment threading formalization is contrasted to the full threading approach as addressed by recursive dynamic programming (RDP, see below). The method can deal with quite general scoring functions including full pair interaction potentials and imposes only some (length) restrictions on the loop regions. The method deals with sets of large numbers of threadings at a time using an efficient representation and estimates appropriate bounds for the maximum achievable score of all the alignments in the sets. Thus, large numbers of potential threadings can be eliminated by applying one of several bounding conditions. Alignments in the fragment threading formalization can be represented by defining sequence positions to be mapped onto the first structural position of any core element, i.e. a vector  $\langle t_1, \dots, t_n \rangle$  for  $n$  Core elements  $C_1, \dots, C_n$ . The amino acid chain and loop restrictions define additional lower and upper bounds for the amino acid to be placed onto the start of a structural segment. A set of alignments is then represented as a vector of lower ( $b_i$ ) and upper bounds ( $d_i$ ) for the segments  $C_i$  and contains all threadings compatible with the constraint that the amino acid aligned to  $C_i(1)$  is in the range  $(b_i, d_i)$ . The algorithm enumerates all possible threadings by selecting such a set and a segment and a so called split point  $t$  in  $(b_i, t_i)$  via recursively producing three subsets  $(b_i, t - 1)$ ,  $(t)$ , and  $(t + 1, d_i)$  and computing the associated bounds on the score of the subsets. If the set of alignments cannot improve the currently best alignment

score known, the whole set can be discarded. The algorithm guarantees to come up with the optimal threading alignment [175], given enough time. If stopped early, the algorithm will present the best scoring alignment found so far. Even after an optimal alignment has been found, the algorithm can be continued in order to produce alternative optimal alignments solutions or present the proof that the alignment is optimal, which can only be stated for certain, once the whole search space has been covered. Typically, it takes about ten times as long to provide the proof as compared to computing the optimal alignment.

The performance of the algorithm heavily depends on the stringency of the bounds of eliminated large sets of poorly performing solutions before actually evaluating them individually.

Considering the potentially huge search space, Lathrop and Smith report implicit evaluation of up to  $6.8 * 10^{28}$  alignments per second [175] (Table 6.1), which compares with about 140 explicit evaluations of the scoring function as observed by Bryant [209].

Despite the fact that, for a set of threading instances, the optimal alignment could be produced for a set of five commonly used scoring potentials, the relevance of globally optimal solution remains doubtful from a biological point of view, due to apparent deficiencies in the used scoring systems. Nevertheless, the approach demonstrated that optimal solutions can be produced for biologically relevant problem instances with, in some cases, structurally meaningful alignments that exhibit small deviations from the standard-of-truth alignment. Whether the algorithmic optimization or the tuning of the scoring functions is the most decisive factor for obtaining high quality alignments with a reasonable amount of computing resources remains to be seen.

Recently, Xu and Xu proposed a new branch-and-bound algorithm called PROSPECT [210–212] for the full threading problem which guarantees to find the globally optimal alignment for a scoring function incorporating pairwise contact terms (in addition to sequence terms, environment profile scores and gap penalties). The algorithm can avoid enumerating the complete search space by using appropriate bi-partitionings of the sequence and structure, such that topological complexity ( $TC$ ) is minimized, i.e. the parts of the partitions exhibit a minimum number of inter-partition links (contacts). As all legal combinations of link assignments have to be considered the  $TC$  is an important factor determining the running time of the algorithm. In biological examples, the  $TC$  has been observed to be small ( $<8$ ) in practice, and the algorithm has a time complexity of  $O(n^{TC} \cdot n^{TC/2})$ , where  $n$  is the target length and  $N$  the maximum allowed loop length (in the current version  $N < 20$ ). Again, including predicted secondary structure information improves the recognition performance of the approach. The PROSPECT method allows exploiting constraints on the target protein such as known disulfide bonds, active site information and long-range (NOE/NMR) distance constraints. The method has been evaluated on benchmark sets [210]

**Tab. 6.1**  
Protein structure resources and databases (coordinates, models, classifications, representative and benchmark sets) on the WWW  
Major protein structure archives and structure coordinate resources:

<b>Name</b>	<b>What</b>	<b>Where</b>	<b>Refs</b>
Protein Data Bank (PDB)	The main database of protein structures	<a href="http://www.rcsb.org/pdb/">http://www.rcsb.org/pdb/</a> Rutgers University, NJ, UCSD, and NIST, Washington	[245, 284]
Molecular Modelling Database (MMDB)	Structure data (ASN.1, Entrez), Visualization (cn3D) and structure comparison tool (VAST), Structure-Taxonomy DB	<a href="http://www.ncbi.nlm.nih.gov/Structure/NCBI, NIH, Bethesda, S. Bryant et al.">http://www.ncbi.nlm.nih.gov/Structure/NCBI, NIH, Bethesda, S. Bryant et al.</a>	[285, 286]
Molecules R US Utility	A full text search of the PDB database to identify structures (pdb ids) by name/keyword	<a href="http://molbio.info.nih.gov/cgi-bin/pdb">http://molbio.info.nih.gov/cgi-bin/pdb</a>	
PDB At A Glance	Hypertext-based classification of the PDB into biochemically meaningful contexts	<a href="http://cmm.info.nih.gov/modeling/pdb_at_a_glance.html">http://cmm.info.nih.gov/modeling/pdb_at_a_glance.html</a>	
PDBSUM	Summary and annotation of PDB entries	<a href="http://www.biochem.ucl.ac.uk/bsm/pdbsum/">http://www.biochem.ucl.ac.uk/bsm/pdbsum/</a>	[287]
Swiss-Model	Wide range of protein analysis tools and protein models	University College, London, R. Laskowski <a href="http://www.expasy.ch/swissmod/SWISS-MODEL.html">http://www.expasy.ch/swissmod/SWISS-MODEL.html</a>	[288]
ModBase	Automatically generated model structures (MODELLER)	Swiss Institute of Bioinformatics (SIB), Geneva, CH, M. Peitsch et al. <a href="http://guitar.rockefeller.edu/modeller/modeller.html">http://guitar.rockefeller.edu/modeller/modeller.html</a>	[243]
Protein Resource Entailing Structural Information on Genomic Entities (PRESAGE)	Resource for information on structural genomics projects and for keeping track of the current status (experimental structure status, structural assignments, models, annotations) for proteins	Rockefeller University, NY, A. Sali et al. <a href="http://presage.berkeley.edu">http://presage.berkeley.edu</a> UC Berkeley, S. Brenner et al.	[289]

Structural Genomics Projects	Resources from various structural genomics projects, Target and model proteins	<a href="http://www.structuralgenomics.org">http://www.structuralgenomics.org</a>	[246]
GeneCensus	Protein folds in (completely) sequenced genomes	<a href="http://www.x12c.nsls.bnl.gov/StrGen.htm">http://www.x12c.nsls.bnl.gov/StrGen.htm</a> <a href="http://bioinfo.mbb.yale.edu/genome/">http://bioinfo.mbb.yale.edu/genome/</a>	
Macromolecular Movements	Database of Macromolecular Movements with Associated Tools for Geometric Analysis	Yale University, M. Gerstein et al. <a href="http://bioinfo.mbb.yale.edu/MolMovDB/">http://bioinfo.mbb.yale.edu/MolMovDB/</a>	[290]
BIND	Database of macromolecular interactions	Yale University, M. Gerstein et al. <a href="http://www.bind.ca/">http://www.bind.ca/</a>	[291, 292]
DIP	Database of interacting proteins (Literature, yeast2hybrid)	Samuel Lunenfeld Research Institute, Toronto, C. Hogue et al. <a href="http://dip.doe-mbi.ucla.edu">http://dip.doe-mbi.ucla.edu</a> UC Los Angeles, I. Xenarios et al.	[92, 93]

Selection of databases on protein structure and protein family classification:

<b>Name</b>	<b>What</b>	<b>Where</b>	<b>Refs</b>
<b>Structural Classification</b>			
Structural Classification of Proteins (SCOP), Cambridge, UK	Major structural classification of protein structures and domains (Murzin et al.)	<a href="http://scop.mrc-lmb.cam.ac.uk/scop/">http://scop.mrc-lmb.cam.ac.uk/scop/</a> Cambridge University, UK, A. Murzin et al.	[245, 284]
Structural Classification of Proteins (CATH), Cambridge, UK	Major structural classification of protein structures and domains (Thornton/Orengo et al.)	<a href="http://www.biochem.ucl.ac.uk/bsm/cath/">http://www.biochem.ucl.ac.uk/bsm/cath/</a> University College, London, J. Thornton et al.	[44, 45]
Dali structure comparison and FSSP structure classification	Protein classification via structural alignment, Dali Domain Dictionary (DDD)	<a href="http://www2.ebi.ac.uk/dali/">http://www2.ebi.ac.uk/dali/</a> <a href="http://www2.ebi.ac.uk/dali/fssp/fssp.html">http://www2.ebi.ac.uk/dali/fssp/fssp.html</a> EBI, Cambridge, L. Holm et al.	[102, 293, 294]
3Dec	Database of structural domains	<a href="http://jura.ebi.ac.uk:8080/3Dec/help/help_intro.html">http://jura.ebi.ac.uk:8080/3Dec/help/help_intro.html</a>	
HOMSTRAD	Database of structure based alignments	EBI, Cambridge, G. Barton et al. <a href="http://www-cryst.bioc.cam.ac.uk/~homstrad/">http://www-cryst.bioc.cam.ac.uk/~homstrad/</a> K. Mizuguchi et al.	[112]



Tab. 6.1 (continued)

<b>Name</b>	<b>What</b>	<b>Where</b>	<b>Refs</b>
3d-ali	Database of structure based alignments	<a href="http://www.embl-heidelberg.de/argos/ali/ali.html">http://www.embl-heidelberg.de/argos/ali/ali.html</a>	[295, 296]
Database of 3-D Protein Structure Comparison and Alignment	3-D Protein Structure Comparison and Alignment via the Combinatorial Extension (CE) Method	S. Pascarella, P. Argos <a href="http://cl.sdsc.edu/ce.html">http://cl.sdsc.edu/ce.html</a> UC San Diego, CA, P. Bourne et al.	[297]
<b>Representative Sets of Structures</b>	40% sequence id	<a href="http://astral.stanford.edu/">http://astral.stanford.edu/</a>	[271]
PDB40	SCOP and derived classification information	Stanford University/UC Berkeley, S. Brenner	
pdb select	Non-redundant sets at various sequence id levels	<a href="ftp://ftp.embl-heidelberg.de/pub/databases/pdb_select/">ftp://ftp.embl-heidelberg.de/pub/databases/pdb_select/</a> , <a href="http://www.cmbi.kun.nl/whatif/select/">http://www.cmbi.kun.nl/whatif/select/</a>	[298]
culled pdb	Non-redundant sets at various sequence id levels	EMBL, Hobohm et al. <a href="http://www.fccc.edu/research/labs/dunbrack/culledpdb.html">http://www.fccc.edu/research/labs/dunbrack/culledpdb.html</a> Fox Chase Cancer Center, Philadelphia, R. Dunbrack	
FSSP/Dali	List of structurally related pairs according to Dali superposition	<a href="http://www.ebi.ac.uk/dali/fssp/TABLE2.html">http://www.ebi.ac.uk/dali/fssp/TABLE2.html</a>	[299]
nrdb	Non redundant sequence DB	EBI, Cambridge, L. Holm <a href="http://blast.wustl.edu/pub/nrdb/">http://blast.wustl.edu/pub/nrdb/</a> Washington University, W. Gish	
HSSP	Database of multiple alignments of sequences to known structures	<a href="http://www.sander.ebi.ac.uk/hssp/">http://www.sander.ebi.ac.uk/hssp/</a> EMBL/EBI, R. Schneider, C. Sander	[74]
<b>Protein Family</b>	Database of sequence patterns and profile indicative for protein function	<a href="http://www.expasy.org/prosite/">http://www.expasy.org/prosite/</a> Swiss Institute of Bioinformatics (SIB), A. Bairoch	[86, 87]
PROSITE			

BLOCKS	Ungapped sequence alignment blocks	<a href="http://blocks.fhrcr.org/">http://blocks.fhrcr.org/</a>	[31, 141]
EMotif	Database of significant sequence patterns for biochemical properties and biological functions	J. Henikoff, S. Henikoff <a href="http://motif.stanford.edu/emotif/">http://motif.stanford.edu/emotif/</a>	[84]
PRINTS	Database of family fingerprints	Stanford University, D. Brutlag et al. <a href="http://www.bioinf.man.ac.uk/dbbrowser/PRINTS/">http://www.bioinf.man.ac.uk/dbbrowser/PRINTS/</a>	[83]
PRODOM	Database of protein domains based on BLAST sequence alignment	Manchester University, UK, T. Attwood et al. <a href="http://www.toulouse.inra.fr/prodom.html">http://www.toulouse.inra.fr/prodom.html</a>	[75]
PFAM	Database of protein domains based on sequence alignment via HMM	INRA, FR, F. Corpet et al. <a href="http://www.sanger.ac.uk/Software/Pfam/">http://www.sanger.ac.uk/Software/Pfam/</a> Sanger Centre, Cambridge, UK, A. Bateman et al.	[22, 24, 73]
Interpro	Combination of the major protein family databases (PRINTS, PROSITE, Pfam, ProDom)	<a href="http://www.ebi.ac.uk/interpro/">http://www.ebi.ac.uk/interpro/</a> EBI, Cambridge, R. Apweiler et al.	[76, 300]
PROTOMAP	Clustering of the structure space	<a href="http://www.protomap.cs.huji.ac.il">http://www.protomap.cs.huji.ac.il</a> <a href="http://protomap.cornell.edu">http://protomap.cornell.edu</a> Cornell University, G. Yona et al.	[257, 301]
SMART	Database of multiple alignments of signalling domains	<a href="http://smart.embl-heidelberg.de">http://smart.embl-heidelberg.de</a> EMBL, P. Bork et al.	[302]
SYSTEMS	Clustering of Swissprot sequences via BLAST	<a href="http://www.dkfz-heidelberg.de/tbi/services/cluster/systersform">http://www.dkfz-heidelberg.de/tbi/services/cluster/systersform</a>	[303]
COGS	Clusters of orthologous groups (sequences occurring in several genomes)	DKFZ, Heidelberg, A. Krause, M. Vinbrun <a href="http://ncbi.nlm.nih.gov/COG">http://ncbi.nlm.nih.gov/COG</a> NCBI/NIH, E. Koonin et al.	[304]

and, for blind prediction, in the CASP experiment [213]. For achieving the results for the CASP targets, a neural network approach has been used to convert threading scores into a confidence measure based on a training set of structurally superposed proteins using SARF. In addition, the program has been run for a broad range of parameters using a supercomputer. The consensus predictions from these runs have been used to obtain higher confidence in the predictions [213].

#### **Heuristic search procedures for full threading based on simulated annealing and genetic algorithms**

Deficiencies of the scoring function obstruct the use of standard combinatorial search methods for finding the optimal threading, multiple alignment, and fragment assembly for protein structure prediction. Therefore, heuristic search procedures such as simulated annealing [36, 214–217], Monte Carlo methods [218–222], and genetic algorithms [173, 223–227] have been used. In general, these approaches are based on the Metropolis method [228]. Starting from an initial alignment, they generate alignment alternatives and accept the new alignments as replacement for the current alignment if they score better or otherwise only with a certain chance depending on the score and an annealing parameter (temperature). After a few steps of the procedure, the annealing parameter is changed according to a so-called cooling scheme and the procedure is continued until no further improvement on the current threading score can be achieved. Apart from the well known critical dependence on the particular cooling scheme to be employed, the approach is computer-intensive and will result in meaningful alignments only if the scoring function is very accurate.

The recent and very successful fragment assembly approach ROSETTA (sometimes called mini-threading method) [5, 37] has been proposed by Baker and co-workers [36]. Here, a simple simulated annealing procedure is used to assemble small segments with similar sequences from unrelated protein structures into possible protein-like structures, which are then subjected to an empirical Bayesian scoring function incorporating sequence, disulfide bonding, solvation, secondary structure packing, and residue pair interaction terms to select the most probable fold. Fragments are nine residues long and the search space is defined by using the conformations of nine residue fragments taken from an ensemble of 25 alternative structure fragments selected from a 25% non-redundant protein set via a sequence similarity criterion. Starting from the fully extended chain, a move substitutes the torsion angles from a randomly chosen neighbor fragment. Invalid (atom distances less than 2.5 Å) conformations are discarded, others are accepted according to a Metropolis criterion. A specific realization [37] of this approach employed for the CASP3 blind predictions constructs 1200 structures, each the result of 100 000 fragment angle substitutions in the simulated annealing procedure. This takes about 2 days of computing time on a fast workstation for a protein domain with 120 amino acids. The per-

formance of the method is evaluated via extensive simulation runs on randomized sequences and fragments, with and without information from multiple alignments. Scoring filters have been performed and the number of conformations within a small rms of the native structure during 100 simulation runs has been recorded to calibrate the method. The current version of the approach has been used with great success for blind prediction in the CASP experiments on both targets with known similar template structures as well as for novel folds [37, 229].

**Dynamic programming approaches, double dynamic programming** As has been mentioned above, dynamic programming (DP) approaches cannot guarantee the optimal solution. Nevertheless, DP methods have been used in several ways. They optimize simplified scoring functions ignoring individual pair contacts, such as threading profiles. In iterative approaches they take fixed interaction partners defined through the alignment of the previous iteration (frozen approximation) [189, 230]. Hierarchical variants, such as double dynamic programming [231, 232] employ DP to determine locally optimal structural environment scores (the first level matrix) for any structural position  $i$  matched with some residue  $a$  (from the sequence) by optimally aligning sequence residues to the contact partners of position  $i$ . They exploit this information from the first level matrix in a subsequent DP (second level matrix) step to find an overall alignment (see also [101] for more detail). The major disadvantage of double dynamic programming is that it requires a dynamic programming optimization of time complexity  $O(n * m)$  for any of the  $n * m$  initial matched sequence structure position pair resulting in a  $O(n^2 * m^2)$  overall time complexity. This is impractical already for medium sized proteins. Clearly, the method is too slow to be used in large scale threading experiments that align a sequence against a fold library of several thousand domains. Thus the method cannot be used to annotate large numbers of sequences. Moreover, the procedure is still a very rough approximation to the optimal solution, as dependencies of partial solutions cannot be handled fully.

**Core fragment threading** Core fragment threading or core-element alignment is a variant of the threading problem similar to the problem formulation used by Lathrop et al. for the optimal branch-and-bound algorithm [175, 208, 210–212, 233] (see Subsection 6.4.2.3.2 above). The idea is that the fold library is preprocessed into a library of structural cores. The query sequence is then aligned to the structural cores without allowing for gaps in the core elements and with some restrictions on the loop length connecting the core elements.

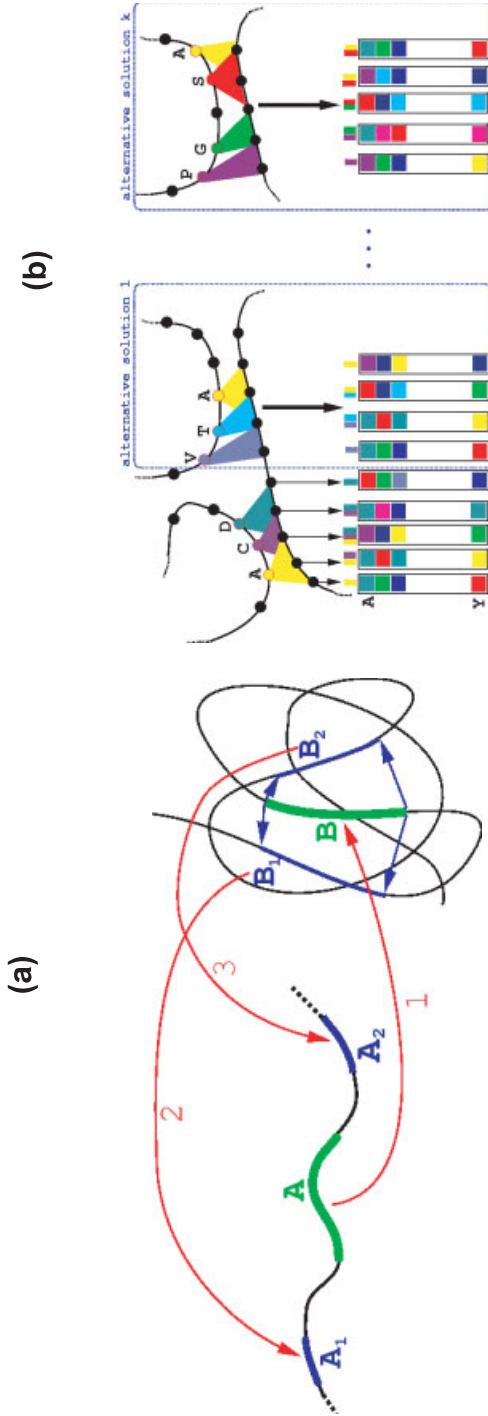
The core structures of template proteins are defined as consecutive segments of the structure that are conserved across multiple alignments of structural [195, 197]. Loop length restrictions are defined to account for the maximum length in observed for the structural neighbors and to avoid non-physical models.

The core-element alignment algorithm [234] starts with an initially random alignment of query sequence segments with the core elements. Given an alignment of the segments, alternative alignments are generated iteratively using preferences implied by the aligned fragments and an empirical pair potential [208] and other scoring terms via a Gibbs sampling strategy (see Appendix). Alternatives are produced by shifting or extending target sequence fragments. Gap penalties are not used aside from the requirement of satisfying the loop length constraints during the generation of alternatives. Similar to RDP (see below) this approach extends the “frozen approximation approach” [189] as the environment is dynamically updated during the Gibbs sampling optimization. The procedure is not guaranteed to converge to a unique and optimal solution. Therefore, a detailed estimation of the significance of a threading and its score is performed by a shuffling the query sequence followed by a re-alignment. Composition-corrected scores are computed as described in [208] and the probability is estimated that a random sequence would score higher than the query sequence assuming a normal distribution of scores and taking length dependencies into account [209].

The core fragment threading approach, using carefully defined structural core models, a calibrated scoring function, and a statistical estimation of  $P$ -values for the scores, appears to be one of the most successful methods in the CASP-3 experiments [196, 197].

**Recursive dynamic programming (RDP)** Another heuristic search procedure based on a branch-and-bound approach has been described in [188]. The approach tries to avoid the restrictions implied by a predefined structural core. Even if the structural core is correctly and completely defined, even the best alignment of the query sequence onto the structure core can be based on information of about half or less of the residues and structural positions of the template structure. Often, structures contain highly conserved loop regions, which if responsible for conserved functions (e.g. ATP binding loops [170]) are also conserved on the sequence level or contain detectable sequence motifs or functional patterns. Such loops can provide important initial hints on similarities and partial alignments exploitable for the assembly of overall alignments.

The recursive dynamic programming approach is based on this observation. It uses a number of so-called oracles to find initial hits of various origin, sequence conservation, conserved disulfide bonds, matching secondary structures, functional patterns, 123D hits, etc. Such an initial hit represents a partial alignment (see Figure 6.5a). From a partial alignment, the contacting regions in the structural template are determined. The already aligned parts then shed (see Figure 6.5b) some pseudo-energy preferences defined by the scoring system used onto the contacting positions. These preferences can be compiled into a structural preference profile for these



**Fig. 6.5**

This figure illustrates the RDP procedure: (a) illustrates the finding of initial hits and how to proceed with further searches based on the contact structure of the template and the remaining not yet aligned sequence parts. (b) illustrates the basic

search step given a partial alignment: the partial alignment implies contact profiles for the remaining positions. Optimal threadings can be found using dynamic programming of sequence parts onto the profiles.

positions. The initial hit also partitions the query sequence into aligned and not aligned parts. Now, the procedure uses dynamic programming to find the best possible match from the not aligned parts of the query with the structural regions represented by the contact profile. This results in more partial alignments extending the partial alignment under consideration. The procedure is continued recursively until no further significant partial alignment can be found in the still unaligned parts. The alignment is then complete and the rest of the structure and the query is considered to be insertions and treated as gaps in the alignment.

This recursive process starts with a set of initial partial alignments each partitioning the problem into a set of subproblems, which need to be solved to obtain an overall solution. From all solutions for each member of the set, the best overall solution(s) are considered as solutions to the overall problem.

Especially in the early phases of this process, many incorrect or sub-optimal partial alignments have to be expected due to the partial information used so far. Therefore many alternatives are considered using a variety of oracles.

Technically, the search space of RDP can be represented as an AND-OR tree with alternating levels of AND and OR nodes. An AND node contains a set of subproblems to be solved in order to solve the problem represented by the node. An OR node contains a set of problems, where the solution of any of them could be used as the solution of the overall problem represented by the OR node.

Usually, the search is organized using a priority queue, holding the nodes in the order of most promising subproblems first. The search selects the most promising subproblem and determines solutions for it in the context of the partial assignments already made.

A detailed description and discussion can be found in [188]. The approach has been shown to improve on profile threading methods on several comprehensive benchmark sets containing structural pairs of increasing difficulty.

The approach is fast enough to allow threading a query sequence against a small fold library of several hundred domains. For large scale threading applications using thousands of template structures or for predicting thousands of sequences, e.g. for whole genome analysis, the normal mode of operation is to apply fast threading methods such as TopLign/123D\* first to select a smaller set of candidates, against which the query is threaded with RDP to obtain additional evidence or more accurate alignments.

Additional functional information and constraints derived from experiments can be exploited in similarity methods to balance inaccurate scoring functions or excessive computing requirements in order to improve fold recognition and alignments of sequences to structures. Distance constraints provide orthogonal measures for scoring alignments and structural models and means for drastically reducing the search space, and thus computing and memory resources. There are essentially two ways to incorporate such

distance constraints into an improved prediction method: the first uses the constraints in evaluating the predicted models either via discarding invalid (i.e. incompatible with the measured constraints) models or by re-sorting the ranked list of models via a score combining threading score and fulfilled constraints. The second approach uses the constraints in the prediction algorithm itself thereby excluding partial model structures in conflict with the experimental constraints. Here a XML-based specification language ProML is employed to represent and access the additional information on the target sequence and the fold template in the prediction algorithm [32, 33].

The RDP method lends itself naturally to such an approach. The search procedure can be easily tailored to special needs via using specialized generating functions to compute the sets of partial solutions and specialized evaluation functions to select the best assembled solutions. If a set of distance constraints is given, the generating functions can be restricted such that only solutions respecting the set of constraints are generated, thereby focusing on consistent alignments in order to guide the search process towards valid alignments and discarding huge search spaces not containing any valid alignment. On the other hand, distance constraints can also be checked between aligned residues after having assembled partial solutions into overall solutions. Therefore, the evaluation function is also restricted with respect to the constraints in order to filter out invalid alignments. Again, the idea is to eliminate invalid alignments as early as possible to accelerate the computation and to improve the alignment quality of solutions.

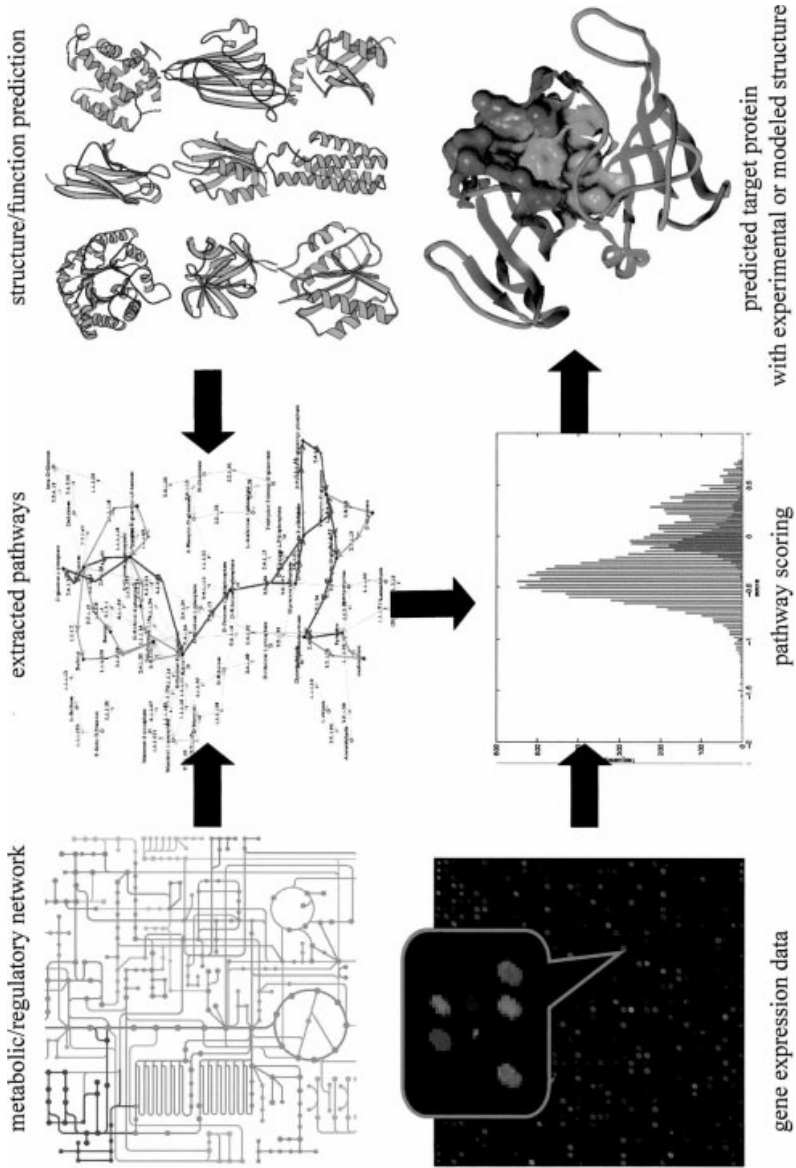
Experimentally measured distance constraints can be of valuable use in similarity methods for structure prediction. Even for moderate numbers (sequence length/10) of distance constraints, we could gain one-third more recognized folds by re-ranking 123D alignments and about two-thirds by employing better alignments on a representative benchmark set [32].

## 6.4

### Results

Protein structure methods as discussed in the previous sections can be employed in a variety of applications from whole genome analysis to detailed rational drug design projects. In basic research, protein structure prediction and the resulting structural models can help to assign possible functions of proteins and to better understand possible functional mechanisms and specificities, possibly in order to plan expression, mutation and recombination experiments, and understand genetic variation. In pharmaceutical and molecular medicine, research applications range from finding functional proteins in complex networks (probably with the help of new high throughput screening data, such as DNA expression measurements with arrays and chips) through the identification of possible drug target molecules, based on the protein function in metabolic and regulatory net-





work contexts, to modeling detailed structures for rationally guiding and designing new drug molecules. Figure 6.6 gives a schematic overview of structure-based methods in a larger functional genomics and target-finding setup.

The following sections briefly discuss some results and trends in using structure prediction for remote homology detection especially in the genomic context, to aid the structural genomics projects, to further whole genome annotation and to exploit the sequence-to-structure-to-function paradigm for functional predictions.

#### 6.4.1

##### Remote homology detection

For the detection of remote homologies based on sequences alone two methods are state-of-the-art: iterative sequence searching with profiles PSI-BLAST, iterative sequence search and hidden Markov models. Haussler et al. have performed a range of experiments and evaluations, which clearly demonstrates the superiority of methods exploiting evolutionary information in the form of profiles [20, 235, 236]. Based on a standard benchmark derived from the PDB40 set they could show that these methods could identify more than twice as many related pairs of proteins (in the PDB40 set) according to the SCOP classification. For distant homologs (less than 30% sequence identity) the HMM based SAM-T98 [236] method outperforms pairwise methods such as FASTA and GAP-BLAST by a factor of three for related pair recognition. With respect to the recognition of all related pairs of proteins in a set of targets these sequence based method currently rival the best structure based methods. Also HMMs have been applied with great success in the CASP experiments [183, 229, 237].

State-of-the-art methods for protein structure prediction are continuously discussed and evaluated in the bi-annual CASP experiments (see the accompanying special issues of the Journal Proteins [329]), the LiveBench [330] and EVA [331] experiments. The major available methods, programs and servers are collected in Tables 6.2 and 6.3.

#### ◀ Fig. 6.6

Protein structure prediction and analysis in a broader context of drug target finding and rational drug design. The figure illustrates one particular method of combining background information on biochemical networks, relevant metabolic or regulatory pathways, the mapping of available structural knowledge onto these pathways, and available expression measurements using a statistical scoring function. The

method produces a candidate list of most interesting target proteins within the context and data in question. In an ideal situation, the most promising target proteins have an experimentally solved or a predicted model of high quality attached to them. Such candidates could then be used not only in functional assays and biochemical experiments but in addition to computer aided rational drug design and experiment planning.

**Tab. 6.2**  
Selection of tools for structure analysis, superposition, classification, and visualization

<i>Name</i>	<i>What</i>	<i>Where</i>	<i>Refs</i>
<b>Secondary Structure Computation and Prediction</b>			
DSSP	3-state secondary structure annotation for protein structures	<a href="http://www.cmbi.kun.nl/swift/dssp/">http://www.cmbi.kun.nl/swift/dssp/</a> W. Kabsch, C. Sander (distributed via CMBI, NL, G. Vriend)	[305]
STRIDE	Secondary structure, accessibility, and hydrogen bonding information	<a href="http://www.embl-heidelberg.de/stride/stride.html">http://www.embl-heidelberg.de/stride/stride.html</a> EMBL, D. Frishman, P. Argos	[306]
TOPS	Topology cartoons of secondary structure arrangements of proteins from DSSP input	<a href="http://www.sander.embl-ebl.ac.uk/tops">http://www.sander.embl-ebl.ac.uk/tops</a> EBI, Cambridge, UK, D. Gilbert et al.	[307]
<b>Visualization and Structure Viewers</b>			
Swiss-PDBViewer	Structure, superposition, alignment and modelling visualization	<a href="http://www.expasy.ch/spdbv/">http://www.expasy.ch/spdbv/</a> SIB, CH, N. Guex et al.	[308]
Rasmol	Easy to use molecular structure viewer	<a href="http://www.bernstein-plus-sons.com">http://www.bernstein-plus-sons.com</a> R. Sayle	[309]
chime	Molecular structure viewer and plugin	<a href="http://www.mdlchime.com/chime/">http://www.mdlchime.com/chime/</a> MDL Information Systems Inc.	
Cn3d	Molecular structure viewer and plugin	<a href="http://www.ncbi.nlm.nih.gov/Structure/CN3D/cn3d.shtml">http://www.ncbi.nlm.nih.gov/Structure/CN3D/cn3d.shtml</a> NCBI, Bryant et al.	
kinemage	Interactive molecule presentation and annotation system	<a href="http://kinemage.biochem.duke.edu/">http://kinemage.biochem.duke.edu/</a> Duke University, J&D. Richardson	

Molscript	Production of publication quality graphics for protein structures	<a href="http://www.avatar.se/molscript/">http://www.avatar.se/molscript/</a>	[310]
Ligplot	Publication quality graphics for protein ligand complexes	<a href="http://www.biochem.ucl.ac.uk/bsm/ligplot/ligplot.htm">http://www.biochem.ucl.ac.uk/bsm/ligplot/ligplot.htm</a> University College, London, J. Thornton et al.	[311]
<b>Structural Alignment and Analysis</b>			
ProFit	Superposition of two pdb structures or regions	<a href="http://www.bioinf.org.uk/software">http://www.bioinf.org.uk/software</a> University of Reading, A. Martin	[294]
Dali	Multiple structural alignment and protein clustering (DDD, FSSP)	<a href="http://www2.ebi.ac.uk/dali/">http://www2.ebi.ac.uk/dali/</a> EBI, Cambridge, L. Holm	[101, 232, 312]
SSAP	Double Dynamic Programming based structure alignment	<a href="http://www.biochem.ucl.ac.uk/~orengo/ssap.html">http://www.biochem.ucl.ac.uk/~orengo/ssap.html</a> MRC and UCL, London, W. Taylor, C. Orengo	[103]
SARF2	Secondary structure element based superposition	<a href="http://www-lmmb.ncifcrf.gov/~nicka/sarf2.html">http://www-lmmb.ncifcrf.gov/~nicka/sarf2.html</a> NCI/NIH, N. Alexandrov	
VAST	Alignment of vectors of secondary structures	<a href="http://www.ncbi.nlm.nih.gov/Structure/VAST/vast.shtml">http://www.ncbi.nlm.nih.gov/Structure/VAST/vast.shtml</a> NCBI/NIH, Bryant et al.	
CE/CL	Structural alignment of SCOP sequences Combinatorial extension (CE) of optimal paths of local structural similarities/compound likeness (CL) searches	<a href="http://bioinfo.mbb.yale.edu/align/">http://bioinfo.mbb.yale.edu/align/</a> Yale University, M. Gerstein Lab <a href="http://clsds.sc.edu/ce.html">http://clsds.sc.edu/ce.html</a> UCSD, Bourne et al.	
Procheck	Checking protein quality and quality plots	<a href="http://www.biochem.ucl.ac.uk/~roman/procheck/procheck.html">http://www.biochem.ucl.ac.uk/~roman/procheck/procheck.html</a> University College, London, J. Thornton et al.	[313]
WHATIF/WHATCHECK	Structure quality check modules as part of the modelling package WhatIF	<a href="http://www.cmbi.kun.nl/whatif">http://www.cmbi.kun.nl/whatif</a> CMBI, Nijmegen, NL, G. Vriend	

Tab. 6.2 (continued)

Name	What	Where	Refs
Alpha shapes	General geometry package usable to analyse packing and accessibility	<a href="http://alpha-geomagic.com/alpha/">http://alpha-geomagic.com/alpha/</a>	
Voronoi tessellation	General geometry package usable to analyse packing and accessibility	NCSA, H. Edelsbrunner <a href="http://www.geom.umn.edu/locate/qhull">http://www.geom.umn.edu/locate/qhull</a> Qhull program, University of Minnesota, Geometry Center	[314]
Scoring Potentials, PDF (probability density functions) ProStar	Archive and evaluation of scoring potentials	<a href="http://prostar.carb.nist.gov/">http://prostar.carb.nist.gov/</a> CARB, Rockville, MD, J. Moulton et al.	

**Tab. 6.3**  
Structure prediction servers and programs

<b>Secondary Structure Prediction</b>	
PHD	Multiple alignment trained neural network prediction [72] <a href="http://www.embl-heidelberg.de/predictprotein/predictprotein.html">http://www.embl-heidelberg.de/predictprotein/predictprotein.html</a> EMBL/Columbia University, B. Rost
predator	multiple alignment based <a href="http://www.embl-heidelberg.de/argos/predator/predator_info.html">http://www.embl-heidelberg.de/argos/predator/predator_info.html</a>
PsiPRED	PSI-BLAST generated sequence profiles and NN [156] EMBL/MIPS/GSF, Munich, D. Frishman <a href="http://insulin.brunel.ac.uk/psipred/">http://insulin.brunel.ac.uk/psipred/</a>
JPred	Consensus of several predictions [159, 160] <a href="http://barton.ebi.ac.uk/servers/jpred.html">http://barton.ebi.ac.uk/servers/jpred.html</a> EBI, Cambridge, UK, G. Barton et al.
CODE	Decision tree based consensus of several predictions [161] <a href="http://cartan.gmd.de/ToPLign/ToPLign.html">http://cartan.gmd.de/ToPLign/ToPLign.html</a> GMD, Sankt Augustin, J. Selbig et al.
<b>Threading programs</b>	
PDB-BLAST	Produce sequence profile with PSI-BLAST and use it in PSI-BLAST run against PDB [283] <a href="http://bioinformatics.jcrf.edu/pdb_blast/">http://bioinformatics.jcrf.edu/pdb_blast/</a> A. Godzik, L. Rychlewski
TOPITS, Predict_Protein	alignment of secondary structure motifs [163] <a href="http://www.embl-heidelberg.de/predictprotein/">http://www.embl-heidelberg.de/predictprotein/</a> EMBL, B. Rost
threadder, threadder2	Threading with pair interaction potentials [190] <a href="http://insulin.brunel.ac.uk/threadder/threadder.html">http://insulin.brunel.ac.uk/threadder/threadder.html</a> Brunel University, UK, D. Jones
Bryant Gibbs Threading	Gibbs sampling for sequence-structure alignment with contact potentials [234] <a href="http://www.ncbi.nlm.nih.gov/Structure/RESEARCH/threading.html">http://www.ncbi.nlm.nih.gov/Structure/RESEARCH/threading.html</a> NCBI, NIH, S. Bryant et al.

Tab. 6.3 (continued)

GenThreader (mGenThreader)	Sequence (PSI-BLAST profile) alignment with neural network based evaluation of alignments with structural scores	<a href="http://insulin.brunel.ac.uk/psipred/">http://insulin.brunel.ac.uk/psipred/</a> Brunel University, D. Jones	[147, 157]
3D-PSSM	Sequence profiles are threaded against structure profiles using secondary structure preferences and solvation potentials	<a href="http://www.bmm.icnet.uk/~3dpssm">http://www.bmm.icnet.uk/~3dpssm</a> ICRF, London, M. Sternberg et al.	[205, 283]
BioinBGU	Consensus of several profile alignment methods with different scoring matrices and profiles	<a href="http://www.cs.bgu.ac.il/~bioinbgu/">http://www.cs.bgu.ac.il/~bioinbgu/</a> Ben-Gurion University, Israel, D. Fischer	[200, 283, 315]
FFAS	Produce sequences with PSI-BLAST and search against PDB	<a href="http://bioinformatics.burnham-inst.org/FFAS/">http://bioinformatics.burnham-inst.org/FFAS/</a> A. Godzik, L. Rychlewski	[20, 183, 237]
SAM-T99	Build HMM for sequence; use HMM to predict secondary structure and to search the PDB. Conversely, HMMs for PDB structure are used to score the sequence	<a href="http://www.cse.ucsc.edu/research/compbio/HMM-apps/T99-query.html">http://www.cse.ucsc.edu/research/compbio/HMM-apps/T99-query.html</a> UC Santa Cruz, CA, K. Karplus	
FUGUE	Library of profiles and substitution matrices from HOMSTRAD structural alignment DB. Sequence or profile is searched against library.	<a href="http://www-cryst.bioc.cam.ac.uk/~fugue/prfsearch.html">http://www-cryst.bioc.cam.ac.uk/~fugue/prfsearch.html</a> University of Cambridge, UK, K. Mizuguchi, T. Blundell et al.	
P-Map	PSI-BLAST approach	<a href="http://www.dnaming.com/Pages/products1.html">http://www.dnaming.com/Pages/products1.html</a> dnaMining informatics software LLC, Bruno	
ssPsi		<a href="http://www.sbc.su.se/~arne/sspsi/">http://www.sbc.su.se/~arne/sspsi/</a> Stockholm University, A. Elofsson	
FORREST	HMMs for structural families from CATH	<a href="http://abs.cit.nih.gov/foresst2/">http://abs.cit.nih.gov/foresst2/</a> Analytical Biostatistics Section, NIH, V. Di Francesco et al.	

loopp	Learning, Observing and Outputting Protein Patterns	<a href="http://ser-loopp.tc.cornell.edu/loopp.html">http://ser-loopp.tc.cornell.edu/loopp.html</a>	
frsvr	Threading with 3D profiles	Cornell University, R. Elber et al. <a href="http://www.doe-mbi.ucla.edu/people/frsvr/frsvr.html">http://www.doe-mbi.ucla.edu/people/frsvr/frsvr.html</a>	[199]
PROFIT	Dynamic programming method for contact potential threading	UCLA, D. Fischer, D. Eisenberg <a href="http://lore.came.sbg.ac.at/home.html">http://lore.came.sbg.ac.at/home.html</a>	
123D+	Profile threading with CCPs (see section 4) using secondary structure predictions	University of Salzburg, M. Sippl <a href="http://www-immb.ncircf.gov/~nicka/123D.html">http://www-immb.ncircf.gov/~nicka/123D.html</a>	[137]
123D*	Profile threading with CCPs (see Section 6.4) using PSI-BLAST sequence/structure profiles and optimized scoring term weighting	NIH/NCI, Frederick, MD, N. Alexandrov, R. Zimmer <a href="http://cartan.gmd.de/ToPLign/123D.html">http://cartan.gmd.de/ToPLign/123D.html</a>	[137, 138, 149, 273]
ToPLign	CCP (123D*) and pair contact potential threading with (recursive) dynamic programming (RDP, branch-and-bound algorithm), profiles, and <i>p</i> -values	GMD, Sankt Augustin, R. Zimmer et al. <a href="http://cartan.gmd.de/ToPLign.html">http://cartan.gmd.de/ToPLign.html</a>	[138, 149, 188, 273]
ROSETTA	Combination of threading/sequence features with <i>ab initio</i> simulation: assembly of structure fragments with sequence similarity using MC simulations	<a href="http://depts.washington.edu/bakerpg/">http://depts.washington.edu/bakerpg/</a> (Information only) Washington University, D. Baker et al.	[5, 36, 37, 316]
Swiss-model	Variety of alignment, analysis and modeling tools, automated modeling server	<a href="http://www.expasy.ch">http://www.expasy.ch</a> Swiss Bioinformatics Institute, Geneva, CH, N. Guex; M. Peitsch	

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## 6.4.2

**Structural genomics**

The goal of structural genomics projects is to solve experimental structures of all major classes of protein folds systematically independent of some functional interest in the proteins [238, 239]. The aim is to chart the protein structure space efficiently. Functional annotations and/or assignment are made afterwards.

Several estimates on the number of different folds have been presented. Some estimates have been based on the number of new and related folds and an analysis of structural classifications and their growth during the past years [240]. Other estimates can be derived from systematic predictions on complete genomes [241–243]. A recent analysis [244] of this type, classifying the new entries of the central structural depository (PDB [245, 332]), has revealed that the fraction of proteins establishing a new family, superfamily, or fold remains constant (at about 15%) over the years (1990–1997), as compared to new proteins belonging to a known family (15%), known proteins from different species (20%), and new (modified) structures from already resolved proteins (about 50%). In addition, new structural entries for sequences with no detectable sequence similarity to known folds (no significant hit via PSI-BLAST searches) yielded about 30% new folds, about 25% new superfamilies with an already known fold, and 45% new families homologous to a known superfamily.

Bioinformatics can support structural genomics by selecting targets for experimental high throughput structure determination, in order to make the best use of the solved structures; i.e. the chance of determining novel folds among the structures to be solved should be optimized. Several clusterings, based on a range of approaches, are used for the purpose of selecting and prioritizing appropriate targets for further X-ray or NMR investigation (see the protein structure initiative [333]) and for keeping track of structural knowledge on proteins, including models and predictions (Presence server [334]). One goal of the overall structural genomics endeavor is to have a solved structure within a certain structural distance to any possible target sequence, which would allow computing a reliable model for all target sequences. This requirement depends on the future enhancements of the modeling procedures on the one hand and on the metric of structural similarity underlying the structure space and the accompanying clusterings, on the other hand. Once a map of the structure space is available, this knowledge should provide additional insights on what the function of the protein is in the cell and with what other partners it might interact. Such information should add to information gained from high-throughput screening and biological assays.

So far, glimpses of what will be possible could be obtained by analyzing complete genomes or large sets of proteins from expression experiments

with the structural knowledge available today, i.e. more or less complete representative sets and a quite coarse coverage of structure space [243, 246, 247].

Recently, several successful individual structure-based functional predictions within the structural genomics projects have been reported: an ATPase or ATP-mediated switch in one example [248], a new NTPase in *M. jannaschii* in a second example [249]; and other test cases are the HIT family [250], *E. coli* ycaC [251], HdeA [252], and yjgF gene products [253]. For a recent review on the current capabilities and prospects of function predictions from structure for bona fide hypothetical proteins on a genome-wide scale, see [254].

#### 6.4.3

##### Selecting targets for structural genomics

The number of different folds is much smaller than the number of proteins. Structure is more conserved than sequence, as nature modularly re-uses successful architectural designs, which allow for some sequence variation. A (continuously increasing) fraction of the possible folds is known. Structural genomics projects aim at producing a representative set of folds. There are increasing chances of finding an appropriate fold for a sequence with unknown structure in the current database of representative folds. Determining a correct fold facilitates the application of homology-modeling techniques to produce a full-atom model structure.

In order to enable the structural genomics projects to focus on the most promising targets for structure determination, structure prediction methods have been applied to estimate the chances of particular proteins to adopt a novel fold, i.e. a fold not yet contained in the structure databases. This should be done for many sequences based on the sequence information alone. Two different approaches have been published. One is based on a clustering of proteins and the distribution of the clusters in sequence and structure space [255]. The other is based on an estimate of erroneous predictions as a function of the corresponding computed sequence-structure compatibility score resulting from a prediction experiment [256]. In this work, Eisenberg and coworkers use a threading program (sequence derived properties SDP [199]) to assign folds to all ORFs of the archaeobacterium *P. aerophilum*. For that purpose they determine a confidence level for an alignment describing the likelihood that it is correct. This confidence is derived from the probability distribution of incorrect matches as judged by DALI/FSSP alignments and z-scores (see Chapter 2 of this volume for the definition of z-score) using a test set of 3285 domains and 10 784 655 pairs achieving a high enough SDP z-score. For the calibrated confidence levels more than 40% of the non-trans-membrane proteins could be as-

signed a fold with 90% and more than 10% with 99% confidence. In addition, about 11% have an as yet unobserved fold with 90% confidence and for 99% confidence still 14 proteins have been predicted to adopt a novel fold.

Portugaly and Linial used the exhaustive PROTOMAP clustering of sequences [257] and the structural classification SCOP [81] to determine a combined sequence–structure classification of proteins. Based on PROTOMAP 2.0 (72 623 SwissProt sequences clustered into 13 454 clusters, 1999) and SCOP version 1.37 (2294 domains, 834 families, 593 superfamilies, 427 folds, Oct. 1997) and a mapping of structures to sequences, the sequence clusters are classified into vacant and occupied, depending on whether the clusters contain known folds. The PROTOMAP clustering structures the sequences space with a graph (the PROTOMAP graph), the edges between the sequence clusters indicate their proximity. Using several measures derived from the cluster space (distances between the clusters, density of clusters, maximal vacant volume) statistical models are constructed to characterize the vacant clusters containing yet undetermined known folds and vacant clusters with probably novel folds. The procedure is illustrated in Figure 6.7. The clusters having the highest probability for representing an as yet unobserved fold constitute promising targets for experimental structure determination.

#### 6.4.4

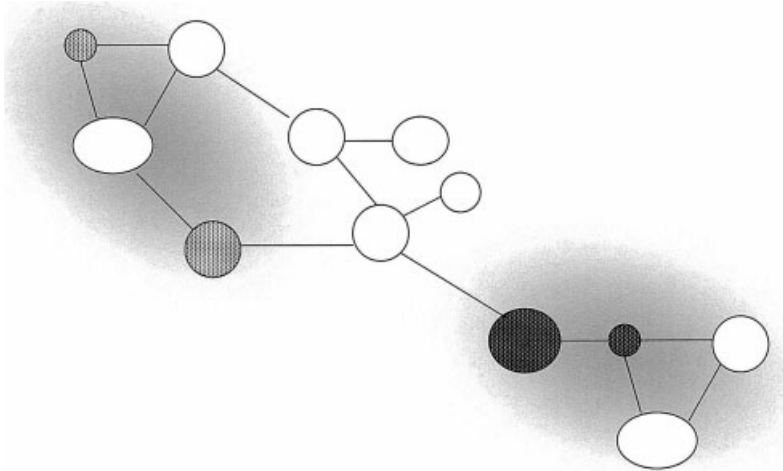
##### **Genome annotation**

Since only 20–40% of the protein sequences in a genome such as *M. genitalium*, *M. Janaschii*, *M. tuberculosis* have a sequence similarity that can exhibit their paralogy to proteins of known function [242, 258], we need to be able to make conclusions on proteins that exhibit much lower similarities to suitable model proteins. As the similarity between query sequence and model sequence decreases below a threshold of, say, 25% we cannot make safe conclusions on a common evolutionary origin of the query sequence and the model sequence any more. However, it turns out that, in many cases, we can still reliably predict the protein fold, and in several cases we can even generate detailed structural models of protein binding sites. Thus, especially in this similarity range, protein structure prediction can help to ascertain aspects of protein function [243, 246, 247].

#### 6.4.5

##### **Sequence-to-structure-to-function paradigm**

Having 300 000 sequences but only a few thousand structures in the databases requires a method to establish relationships of sequences to homologous proteins with known function and/or structure.



**Fig. 6.7**

Illustration of the PROTOMAP approach and an example of a PROTOMAP graph of protein structure space. Nodes represent sequence clusters; edges represent proximities between clusters in sequence. Lightly and darkly shaded nodes

are clusters that contain sequences known to fold into structure A and B respectively. Using these clusters the remaining protein space can be heuristically partitioned into areas that are likely to fold into known structures and those that are likely to fold into novel structures.

The relationship between structure and function is a true many-to-many relation. Recent studies have shown that particular functions can be mounted onto several different protein folds [85] and, conversely, several protein fold classes can perform a wide range of functions [259]. This limits our potential of deducing function from structure. But it is still possible to use aforementioned knowledge on the range of folds supporting a particular function and the range of functions implemented by particular folds in order to make functional prediction from structure.

Homologous proteins have evolved from a common ancestor and have nearly always a similar 3D structure. This does not necessarily imply similar function: [129, 259, 260]. Probabilistic sequence searches have their limits despite the significant improvements due to PSSMs [9, 203, 205], secondary structure information [156–7, 165, 184] and HMMs [25, 179, 261], which have been systematically evaluated by several authors [20, 129, 147, 195, 205].

If a protein has a known fold, which in addition is annotated with a particular or a few specific functions, the model derived for the sequence in question based on the respective fold can be evaluated with respect to compatibility with some of the proposed functions.

In general, this sequence-to-structure-to-function paradigm involves predicting a structural model for a given sequence (either by sequence and/or structural means, e.g. threading) and, afterwards, identifying a 3D pattern,

e.g. an active site, indicative for a particular function in the constructed model.

Especially for enzymes, common mechanisms and associated active sites are often observed. This allows searching for appropriate 3D patterns in a database of structures and for checking models to contain such a pattern (for a well known example, see SER-HIS-ASP serine protease catalytic triad [262]).

Sometimes ligand-binding pockets are identifiable on protein surfaces. With such a binding pocket hypothesis, docking tools can be applied in order to screen potential compounds to bind the protein site.

Protein-protein interactions can be predicted with various methods: analyzing known protein interfaces and inferring common motifs via surface and sequence conservation criteria [263–265], protein fusions [95, 266] or, less successful so far, correlated mutations [267].

A systematic assessment of the relationship between protein function and structure has been performed by Hegyi and Gerstein [259] via relating yeast enzymes classified by the Enzyme Commission (EC numbers) to structural SCOP domains. In this study it has been found that different structural folds have different “propensities” for various functions. Most versatile functions (hydrolases and O-glycosyl glucosidases) have been identified to be mounted onto seven different folds, whereas the most versatile folds (e.g. TIM-barrel and Rossmann folds) realize up to 16 different functions.

In addition to conventional sequence motifs (Prosite, BLOCKS, PRINTS, etc.), the compilation of structural motifs indicative of specific functions from known structures has been proposed [268]. This should improve even the results obtained with multiple (one-dimensional sequence) patterns exploited in the BLOCKS and PRINTS databases. Recently, the use of models to define approximate structural motifs (sometimes called fuzzy functional forms, FFFs [269]) has been put forward to construct a library of such motifs enhancing the range of applicability of motif searches at the price of reduced sensitivity and specificity. Such approaches are supported by the fact that, often, active sites of proteins necessary for specific functions are much more conserved than the overall protein structure (e.g. bacterial and eukaryotic serine proteases), such that an inexact model could have a partly accurately conserved part responsible for function. As the structural genomics projects produce a more and more comprehensive picture of the structure space with representatives for all major protein folds and with the improved homology search methods linking the related sequences and structures to such representatives, comprehensive libraries of highly discriminative structural motifs are envisionable.

Whether or not it is possible to infer protein function, however detailed, from a experimental structure, or from an approximate model, or even from only a partial fold model is a key question in the context of large-scale functional annotation. Currently, the situation is that sometimes such deri-

vations are possible; in more cases additional evidence for functions predicted with sequence methods or from functional motifs can be established; in most cases detailed functional classification from inaccurate structures cannot be guessed.

Though it is not easy to derive functions from resolved protein structures, the availability of structural information improves the chances as compared to relying on sequence methods alone.

## 6.5

### Validation of predictions

Theoretical predictions are risky. Therefore for almost all such prediction experimental validation is required. Nevertheless, often the models can indicate appropriate ways for validation or further experiments. These experiments can be expected to be time-consuming, and expensive. Furthermore, the protein actually needs to be available for the suggested experiments. All of this limits the applicability of experimental validation. Therefore, it is mandatory to reduce errors as much as possible and to indicate the expected error range via computer-based predictions. This is not a trivial problem for structure prediction, though. An estimation of the performance and accuracy of the respective methods can be obtained from large scale comparative benchmarking, from successful blind predictions and from a community wide assessment experiment (CASP [109, 229]/CAFASP [283]). These are addressed in turn in the following:

#### 6.5.1

##### Benchmark set tests

Structure prediction methods have to be validated using comprehensive and representative benchmark sets. A couple of benchmark sets have been proposed [188, 199, 270] consisting of a fold library and a list of structurally similar protein pairs without significant sequence similarity. Such a benchmark [335] involves using the method in question to thread one protein of each pair against the library and to count the number of successful recognitions of the structural match or the number of correct identifications of related protein pairs. A couple of methods have been evaluated this way.

More convincing are benchmarks representative for the currently known folds. Thus, given a structural classification of all the structures in the PDB (e.g. SCOP) and a list of proteins without high sequence similarities, say less than 40% pairwise identities (e.g. PDB40 [336], which can be obtained from, e.g. the ASTRAL server [271] or from a couple of other sources for such representative lists [337]), one can easily determine a complete list of

structural similarities from the structural classification. I.e. a pair of proteins from PDB40 are considered structurally related if they belong to the same structural class (depending on the level, the same superfamily or the same fold). Other lists of structurally related pairs (via structural alignment of the 3D structures) are available (e.g. from the DALI/FSSP database [338]). A comprehensive benchmark then involves threading each sequence of the PDB40 against all other members [20, 272, 273]. Two benchmarking protocols can be defined: 1. (*Fold recognition*) for each threading run identify the best fitting fold (usually the one with the best score, sometimes after applying filtering or reranking of alignments or models in a post-processing step) and predict it as the closest structural homologue in the fold library and use it as structure template for model building. Of interest in this protocol is the number or percentage of correctly predicted structures. 2. (*Related pair recognition*) for the all-to-all threadings and the respective scores define a score threshold above which threadings are considered predictions of structural similarity. As only one score threshold is applied for the individual threading runs, the scores need to be normalized with respect to the length and other features of the query protein in order to render the scores comparable to each other. Computing  $p$ -values from the scores is a method to achieve comparability among experiments and to assign a significance measure to each prediction [273]. This leads to a list of predicted pairs with associated  $p$ -values, which can be compared to the list of standard-of-truth pairs. In the simplest case this leads to the number of correctly predicted pairs ( $TP$ ), of incorrectly predicted pairs ( $FP$ ), and the number of missed correct pairs ( $FN$ ). Indeed, the ranked list of scored pairs contains much more information and allows yielding a complete overview of the performance of the method for all possible threshold values of the  $p$ -values. This is represented as a so-called sensitivity/specificity plot. Given a  $p$ -value threshold the following values of sensitivity  $Sens$  and specificity  $Spec$  can be easily read off the ranking list:

$$Sens = TP/(TP + FN) \quad \text{and} \quad Spec = TP/(TP + FP),$$

where  $TP$  = true positives,  $FN$  = false negatives,  $FP$  = false positives, the predicted pairs (positives) change according to the  $p$ -value threshold, the negatives remain constant. The sensitivity is the fraction of correct pairs above the threshold as compared to all correct pairs, and the specificity ( $1 - \text{error rate}$ ) is the fraction of correct pairs above the threshold as compared to all pairs above the threshold. Now the set of all possible ( $Sens$ ,  $Spec$ ) pairs for all possible thresholds can be simply obtained for the ranking list by iteratively decreasing the threshold from the maximum achieved  $p$ -value, to the next highest  $p$ -value, down to the lowest  $p$ -value observed. For each consecutive threshold value the ( $Sens$ ,  $Spec$ ) values are updated according to whether the new pair above the threshold is a true ( $TP$ ) or false ( $FP$ ) structural pair. Example plots for evaluating the fold recognition performance of 123D with five different significance criteria are shown in Figure 6.8. From

such a plot, the expected performance of a certain method using certain parameter settings can be read off for any given error rate to be acceptable for the user.

### 6.5.2

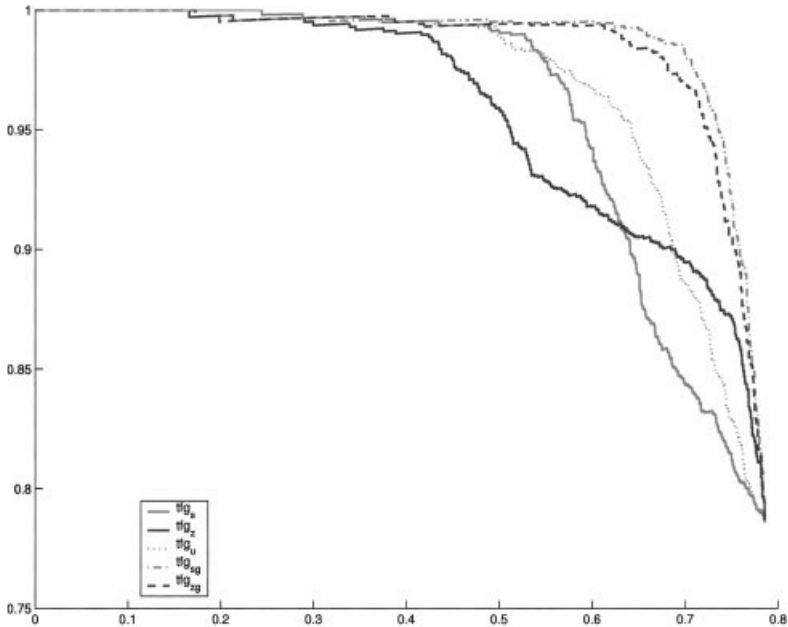
#### **Blind prediction experiments (CASP)**

A blind prediction is a computer-based construction of 3D models of a protein sequence for which no structure is known at the time. CASP (comparative assessment of structure prediction methods [109, 274–276]) is a worldwide contest of protein structure prediction that takes place every two years. During the CASP experiments, a set of automated numerical evaluation tools have been implemented [110, 111, 277, 278] to cope with the large number of predictions in a way that is as objective as possible. In fact, the experiment is also devoted to the research and development of such unbiased methods. However, there is still quite some controversy on the criteria to judge protein structure predictions and the corresponding models [339].

For the CASP2 experiment, Marchler-Bauer and Bryant defined a large number of quantitative criteria [110] to be used by the assessor to judge and rank the models and prediction teams. These criteria include several measures for fold recognition specificity, for threading alignment specificity, threading contact specificity, and model accuracy.

For an evaluation of the CASP3 predictions, Sippl used the ProSup rigid sequence-independent 3D superposition software [111, 279]. The basic assumption is that the best prediction corresponds to the best structural match between the predicted model and the experimental structure, but that the best structural match cannot always be determined without ambiguity automatically. Therefore, all possible structural matches between model and structure are considered and ambiguities are resolved in favor of the predictor to determine the closest correspondence between the prediction and structural matches. Ranking a set of structures according to their similarity to a reference fold or model defines a totally ordered similarity measure. Unfortunately, structural superpositions are characterized by two contradicting quantities, the number of equivalent residues and the best possible rms achieved by optimally superposing the equivalent residues, which can be individually optimized compromising the other. A solution proposed by Sippl for CASP3 follows a similar strategy as used in the protein superposition tool SARF [103, 280], i.e. considering only superpositions with a rms smaller than a constant value (say 3.0 Å) and maximizing the number of residues within that rms range. Usually, this leads to an easily interpreted measure of structural similarity and corresponding rankings of structures [111]. In order to evaluate a prediction, the maximum number of equivalent residues is determined for all possible structural matches between model and structure. The conclusion of the numerical evaluation is that it is difficult to clearly discriminate between appropriate and inappro-





**Fig. 6.8** Sensitivity-specificity plots. The figure shows example sensitivity/specificity plots for several fold recognition criteria (raw threading score, z-score,  $p$ -value,  $gap$ -score,  $gap$ -z-score) for a benchmark experiment of 123D\* threading on a PDB40 domain library using sequence profiles generated for the query sequence via PSI-BLAST. Such plots allow an

immediate overview of the recognition performance (sensitivity,  $x$ -axis) over the whole range of error rates (specificity,  $y$ -axis). In this case, one can achieve a maximum sensitivity of about 78% (at any error rate), more than 60% sensitivity for a specificity of more than 95% for all but one criterion, for which about 8% error (Spec = 0.92) has to be tolerated in order to recognize more than 60% of the folds.

appropriate structural matches as often the ranked list of PDB structures matching a target structure shows a continuum of matched residues. In addition, different criteria, such as equivalent residues according to the best structural match or according to the match producing the maximum number of equivalence, or minimum shift error of aligned residues, lead to a general trend of good and bad models, but in most cases also to debatable decisions on which models are best or better than others. The comparison of the results with structural matches obtained from other methods, e.g. the Dali structural alignment, where a  $z$ -score is computed from the two quantities and used for ranking, leads to quite different results [111].

Therefore, an alternative approach by Hubbard [277] avoids coming up with a single number qualifying a prediction, but rather a plot of rms vs coverage of a number of predictions. The plot is produced such that for any prediction there is a set of points (drawn as a line) representing for each coverage value the minimum rms out of a sampling of superpositions. Such

plots allow for a comparative general overview of all predictions: one can easily identify outstanding predictions for a given structural similarity (i.e. rms) or coverage (i.e. a given number or percentage of aligned residues) and, in addition, assess predictions for the whole range of structural similarities or coverages. The plots for the CASP3 targets and the submitted predictions are available at the CASP Web site [340].

Another approach used for CASP3 is sequence-dependent superposition – in contrast to the two approaches mentioned above which are sequence independent relying on the best structural superposition of the 3D model without reference to the natural 1:1 alignment between model and template. In difficult cases, sequence-dependent superposition iteratively excludes residues being modeled less accurate from the computation [278]. This procedure does not necessarily lead to unambiguous solutions. The method allows defining and displaying another set of evaluation criteria for fold recognition and individual alignment accuracy such as longest continuous segments and largest superimposable sets of residues. These criteria can also be restricted to certain subsets of the structure, such as secondary structure elements, buried or exposed residues etc. Results of these measures for the CASP predictions are also available from the CASP Web site.

Due to the debatable outcome of the various methods for numerical evaluation of protein structure prediction the final assessment of models based on the quantitative measures is kind of a creative task similar to the inherent ambiguities encountered in structural classifications [196]. This is especially true for the range of sequence and structure similarities encountered for CASP fold recognition targets, where similarities and partial similarities are hard to identify even for the best human experts in the field. Indeed, a slightly more general definition of the SCOP fold similarity has been used for the CASP3 assessment [196]. The overall result of the structure prediction experiment relevant for this Chapter (fold recognition), which can be considered as an approximation to the overall state-of-the-art in the field, is as follows (according to Table I in [196]): 22 of the 26 target domains show substantial similarities to known structures, three proteins are classified as new fold, one domain has not been classified. Eleven domains are likely to be distantly homologous to known structures. Nine of the 11 homologous relationships have been detected, four of which with accompanying overall correct alignments and models. The other 11 domains with known folds are predicted as follows: five folds identified, one correct topology predicted, two partial structure correctly predicted, and only three folds have not been identified by any method. Two of the three new folds have been predicted correctly as new. Overall, 19 of the target domains have been solved by one or more of the prediction teams, only six could not be solved, and one domain remains unclassified.

In accordance with the focus of this chapter on current proteins structure prediction methods, and despite some considerable improvements of *ab initio* methods (e.g. the fragment assembly approach ROSETTA [37],

sometimes called “mini-threading approach” and an approach based exclusively on a global optimization of a potential energy function [35]) as compared to CASP2, CASP3 has shown no emphasis on “physical methods”, i.e. methods concentrating on the balance of forces determining structure or folding pathways leading to native structures [109]. It is also apparent that currently the homology-based methods have much more predictive power and it seems to be like that for some time to come.

## 6.6

### Conclusion: strengths and limitations

#### 6.6.1

##### Threading

Threading allows, given a query sequence, to search for sequences with a similar fold but without apparent sequence similarity. Therefore, threading has three major objectives: first, provide orthogonal evidence of possible homology for distantly related protein sequences, second, detect possible homology in cases where sequence methods fail, and third, to improve structural models for the query sequence via structurally more accurate alignments.

Towards all three goals continuous improvements have been achieved during the up-to-date four rounds of the CASP experiment. The CASP experiments are the main indicator of the progress in protein structure prediction. Whereas the evaluation of progress is not as easy as first expected, it turns out that, for fold recognition, detectable progress can be observed from CASP1 to CASP2, where there is no significant progress between CASP2 and CASP3, but the same performance was achieved on more difficult targets. In CASP4, again, manually applied knowledge-based approaches to distant fold recognition outperformed automated methods and servers. Improvements have been made especially on the prediction of new folds where both manually (A. Murzin) and the *ab initio*/fragment assembly approach [5] performed well on a couple of targets with novel folds. The latter approach performed well on average also for fold recognition and comparative modeling targets [281]. But there appears to be a certain limit of current fold recognition methods, which is still well below the limit of detectable structural similarity (via structural comparisons). In addition, in CASP3 several groups produced reasonable models of up to 60 residues for *ab initio* target fragments, in CASP4, Baker could demonstrate good average performance of the fragment assembly approach.

We provide a summary of an estimate of the current state-of-the-art, in CASP3. Out of 43 protein targets, 15 could be classified as comparative

homology modeling targets, i.e. related folds and accompanying alignments could be derived beyond doubt. For more than half of the 21 more difficult cases, a reasonable model could be predicted by at least one of the participating prediction teams. In addition, the CAFASP subsection of the assessment has demonstrated that 10 out of 19 folds could be solved via completely automatic application of the best threading methods. The complete set of results obtained during the CASP experiments, including the recent CASP4 assessment (summer/winter 2000), can be viewed and analyzed through the CASP Web page at the prediction center [341].

Methods for refining rough structural models towards the true native structure of the query protein are not straightforward, but are an active area of research [282].

#### 6.6.2

##### **Strengths**

The strengths of structural predictions is that structural and in some cases functional classifications and annotations can be made, where such an annotation is not possible at all with pure sequence-based methods. In general, if there is detectable sequence similarity the corresponding annotation should overrule contradicting – even high confidence – structural predictions. In cases with insignificant sequence similarity, a significant threading alignment can provide additional evidence for an annotation. In addition to extending the range of remote homology searches, a knowledge-based structure prediction always results in a structural model, which can be inspected and evaluated with other means. In particular, such models can be completed to full atom models accessible to energy minimization and theoretical model validation procedures. The model can also trigger the validation and cross-checking with additional experimental facts and in turn can indicate further experiments either to validate the model or to check for putative functions of interactions with other partners.

In addition, experimental methods become available for checking predicted models in a cheap, fast and efficient manner. One such method uses chemical cross-links of a certain length and specificity and allows for validating distance constraints between residues and groups implied by a model. The method uses a protocol involving chemical cross-linking, digestion by specific hydrolases, measurement of peptide masses via (MALDI-TOF) mass spectrometry, inferring distance constraints for the protein under investigation and checking the experimental distances against the proposed model [32]. This appears to be a very promising approach to boost the significance and thus the practical application of theoretical structure prediction approaches, especially for proteins, which are not easily amenable to experimental structure determination, i.e. proteins difficult to crystallize, membrane proteins, or highly flexible proteins.

## 6.6.3

**Limitations**

Still structure prediction is an unsolved problem and the methods available are rough and heuristic. The outcome of predictions is not easily interpreted by the non-specialist and can easily be over-interpreted by biologists if taken too seriously. The computation of suitable p-values to allow for a straightforward judgment of the respective prediction remains problematic for many methods and parameter settings. In addition, models derived from similarities on the fold level often simply are not as structurally similar as one could hope for, i.e. not very usable for detailed structural analysis. Variations and differences apart from conserved structural cores have to be expected frequently even for highly significant structural predictions.

This is one reason why in general reliable function predictions cannot be inferred from structural models and mere significance of structural similarity at the level of folds. The current state-of-the-art of understanding structure-function relationships requires additional evidence to make such predictions. This, however, could be available via additional experimental measurements (e.g. expression data) or knowledge on biochemical and protein interaction networks.

**Protein interaction/fusion:**

1. Recently, the *Rosetta stone* method has been introduced. This method uses over 20 completely sequenced genomes and analyzes evolutionary correlations of two domains being fused into one protein in one species and occurring in separate proteins in another species. From these classifications the method establishes pairwise links between functionally related proteins [94] and elicits putative protein–protein interactions [95].
2. The *phylogenetic profile* method analyzes the co-occurrence of genes in the genomes of different organisms, for the same purpose [96].

One major reason for the currently not satisfying performance of knowledge-based prediction methods is the obvious deficiency in the currently used empirical scoring functions. As has been demonstrated there are many methods to improve the capabilities of the potentials by calibrating parameters or weightings of the scoring system [149]. This hints to clear deficiencies of the available potentials. As more data become available the database-derived potential can be expected to become more accurate.

The other major disadvantage today is that prediction methods do not efficiently exploit the available heterogeneous information on function, motifs, family information, network topology, sequence and structure space

topology during the prediction process. This is clearly demonstrated by the fact that the prediction performance and alignment accuracy of human experts armed with the above mentioned prediction methods has been much better than automated versions of the methods alone [109, 283]

As has been briefly mentioned above the sensitivity of current methods can be expected to be quite low for the detection of related structural pair recognition and can be estimated at a quite satisfying performance of about 50% for finding a suitable structural template. The rather high error rate of around 50% has to be tolerated to achieve these sensitivities. This seems to be acceptable if the correctly identified models are new targets and of sufficient interest for further investigation to compensate for the false hints. If appropriate experimental assays are available, probably even a hit rate of 1 in 10 could be acceptable in target finding applications. As mentioned above, together with additional theoretical evaluation and evidence, and appropriate experimental techniques and validation, the success rate can be expected to be much higher.

## 6.7

### Accessibility

Many of the resources, programs, databases, and servers discussed in this Chapter and relevant for protein structure prediction are freely available over the internet. The field, and thus the available resources, change quickly according to the progress made on collected experimental data and new methods. The Tables in the Appendix provide a current overview.

Table 6.1 contains a selection of protein structure resources and databases (coordinates, models, classifications, representative and benchmark sets) available on the WWW with the major protein structure archives and structure coordinate resources.

Table 6.2 lists a selection of tools for structure analysis, superposition, classification, and visualization.

Table 6.3 contains a list of structure prediction servers and programs. See also the CAFASP home page for further information and for results achieved for CASP4 targets and structures newly submitted to the PDB (check LiveBench [342] and EVA [343] for current updates and evaluations).

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## 7

# Protein–Ligand Docking in Drug Design

*Matthias Rarey*

### 7.1

#### Introduction

Since more and more three-dimensional protein structures become available due to X-ray crystallography, NMR spectroscopy, and homology modeling, software tools using this information in drug design projects become more important [1, 2]. The key problem these software tools have to tackle is the docking problem, namely predicting energetically favorable complexes between a protein and a putative drug molecule, also called ligand in this context.

Figure 7.1 (s. p. 317) shows the complex between the protein thrombin and a well-known inhibitor called NAPAP. Thrombin plays a central role in the blood coagulation cascade and has long been an important target in thrombosis research. Like many other enzymes, thrombin catalyzes a specific reaction by forming a complex with a substrate molecule, a peptide in this case. The inhibitor NAPAP is designed to bind to the same region on the protein surface, the active site, than the substrate but with a higher binding affinity. Therefore, in the presence of NAPAP, the natural reaction on the substrate molecule cannot be catalyzed by thrombin, the enzyme is blocked in its activity.

In the first steps of a drug design project the goal is to find a so-called lead structure, a small molecule which binds to a given target protein and can be further developed to a drug. If the three-dimensional structure of the target protein is known, docking algorithms can be applied to virtually search for potential leads. The two main questions arising are what the complex between a protein and a potential lead looks like and how strong the binding affinity of the lead is with respect to other candidates.

Several aspects make the docking problem hard to solve. First of all, one has to deal with the scoring problem [3–5], namely the calculation of binding affinity given a protein–ligand complex. Today, no general applicable scoring function is available giving an accurate prediction of the binding affinity. Secondly, a large number of degrees of freedom has to be consid-

ered during a docking calculation. The most important degrees of freedom are the relative orientation of the two molecules and the conformation of the ligand molecule. Beside these, the protein conformation may also vary, water molecules can be located between the molecules and the protonation states of the molecules can change. Finally, the various variables are subject to a complicated network of constraints. A small change in one variable may decide whether an overlap between the two molecules occur or a hydrogen bond between two groups can be formed or not. This implies that the scoring functions for protein-ligand docking typically contain lots of local minima and are difficult to optimize.

Although no general solution of the docking problem is available today, several algorithms for various kinds of docking problems have been developed and successfully applied. This chapter will give an overview of these algorithms, additional review articles about software for structure-based drug design can be found in the literature [1, 6–13].

Before we start our methods overview, we will summarize the types of docking problems and describe some typical application scenarios.

### 7.1.1

#### **A taxonomy of docking problems**

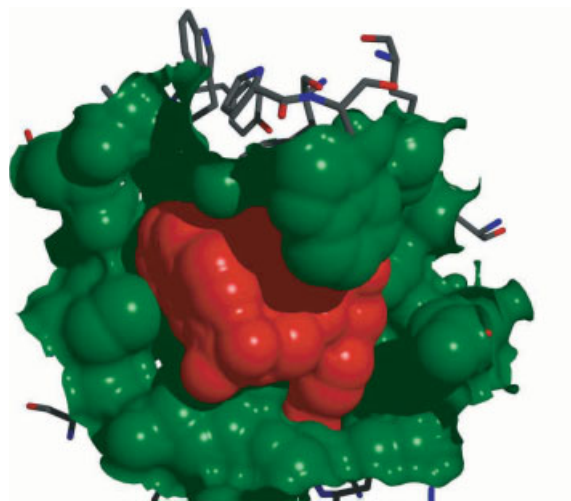
Docking problems can best be classified by the type of input molecules. In macromolecular docking, two macromolecules like proteins or DNA are docked. The major characteristics of these problems are that the complex typically has a large contact area and that the molecules – although still flexible – have a fixed overall shape. These features imply that methods based on geometric properties like shape complementarity alone can already be used efficiently for creating energetically favorable complexes. A survey of methods for protein-protein docking is given in Chapter 8 of this book and elsewhere [14].

The second class contains docking problems in which a small molecule is docked to a macromolecule. The macromolecule can be a DNA fragment or a protein, in the latter case the problem is called protein–ligand docking. In general, the small molecule is an organic molecule, for example a natural substrate or an inhibitor like NAPAP mentioned in the example above.

Small-molecule docking differs substantially from macromolecular docking in the fact that the ligand is typically not fixed in its overall shape. The conformational flexibility of the ligand molecule is of importance and geometric properties alone are often not sufficient to determine low-energy complexes. Even in cases where the molecule is more or less fixed in conformational space, the shape of the molecule is not characteristic enough to find low-energy complexes based on shape alone. Therefore, different algorithms are developed for small-molecule docking compared to macromolecular docking.

**Fig. 7.1**

Complex between the protein thrombin and one of its inhibitors NAPAP. Only the active site of thrombin is shown (atoms drawn with sticks colored by atom, molecular surface is shown in green, NAPAP is shown in red).



For various reasons pertaining to important drug properties like their bioavailability, most drugs are small molecules. Therefore, small-molecule docking is of great interest in pharmaceutical research. Because pharmaceutical targets are often proteins, most docking algorithms are developed for protein-ligand docking.

In principle, docking of small molecules to DNA can be handled by the same algorithms. Differences occur in the handling of DNA specific binding mechanisms like covalent binding or binding of so-called intercalators [15]. An intercalator is a small molecule that binds to the DNA between two subsequent bases. Since a structural change in the DNA is necessary for binding, conformational flexibility of the DNA must be taken into account. This can be done by docking into a distorted DNA structure [16]. A better way is to deal with the DNA flexibility directly. For example, Zacharias et al. [17] developed an algorithm based on normal mode analysis for handling structural changes of the DNA during docking calculations.

Protein-ligand docking problems can be further distinguished by the size of the ligand molecule. The typical ligand molecule occurring in drug design docking problems has about 5–12 rotatable bonds, thus, often the question of placing only a part of the ligand, a fragment, in the active site of the protein emerges. In contrast to a typical drug molecule, the conformational space of a fragment is quite small and must not necessarily be taken explicitly into account. Therefore, algorithms handling only the relative orientation of two molecules (rigid-body docking algorithms) can be applied. Docking fragments to proteins is a subproblem handled in several more complex docking algorithms. Some examples are the protein-ligand docking algorithms in which flexibility is handled by dividing the ligand into smaller fragments, *de-novo* design algorithms in which new molecules

are created from a fragment database, or combinatorial docking algorithms in which combinatorial libraries of molecules are docked by combining placements for individual building blocks of the library.

Orthogonal to this classification by the type of input molecules, the second important parameter for categorizing docking algorithms is the time spent per prediction. The number of molecules which have to be processed covers a range from single molecules to several ten thousands and depends on the concrete application scenario.

### 7.1.2

#### **Application scenarios in structure-based drug design**

Docking problems arise in several different scenarios in structure-based drug design. The overall goal is always to plan the next experiment in order to find or create a lead structure or optimize specific properties of it.

If compounds binding to the target protein are known, the three-dimensional structures of their complexes with the target protein are of interest. They may give hints as to the catalytic mechanism of the enzyme or opportunities for optimizing the binding affinity of a compound. Because only a small set of compounds can be inspected in detail, computing time spent for predicting a protein–ligand complex plays a minor role. Typically, creating a set of starting geometries with a protein–ligand docking algorithm is the first step succeeded by molecular dynamic simulations and detailed energy calculations.

In some cases, a medium sized set of compounds (a hundred up to several thousand) are considered as candidates for lead structures. This set can emerge for example from a screening experiment, from filtering databases with other software tools, or from varying a known inhibitor or lead structure in several ways. Here, the required output is a ranked list of the compounds giving the medicinal chemist a hint which compounds should preferably be investigated further.

Virtual screening is the *in silico* variant of high-throughput screening (HTS) used to find initial lead structures from large compound collections. A virtual screening experiment can be performed on the basis of a docking algorithm. Although the accuracy of computer-predicted binding affinities is much lower than of experimental determined affinities, virtual screening has some advantages. There is no uncertainty about the molecular structure tested, the calculation is cheaper than the real experiment, and, most importantly, the calculation can be performed on compounds that are not yet purchased or synthesized. Obviously, for virtual screening a fast docking algorithm is needed. Compound collections with up to half a million molecules are typical for screening exercises and it can be assumed that this number will increase in the next years.

If no lead structures are available yet, a *de novo* ligand design can be performed as an alternative to a screening experiment. The immanent problem with this approach is to create molecules which can be synthesized easily. In order to achieve this goal, fragment-based approaches are preferably used. The central step in these approaches is to dock a library of small fragments into the active site of the target protein. Because these fragment libraries can be quite large, time-efficient docking algorithms are necessary for *de novo* ligand design.

A few years ago, combinatorial libraries of small organic compounds were introduced to drug design [18, 19]. Combinatorial chemistry offers a way to create large collections of diverse compounds which can then be used in screening experiments. Although the chemistry behind combinatorial libraries can be quite complicated, the overall methodology is quite intuitive and simple. The starting point of a combinatorial library is a specific reaction or a small set of reactions and a set of molecules (reagents). The reactions connect two or more of the reagents in a well-defined way resulting in a single molecule (product) of the library. By combinatorially combining all reagents, an exponential set of products is synthesized. Usually the reactions result in a molecular fragment which is in common between all molecules of the library. This fragment is then called the core of the library, the fragments attached to the core are called R-groups.

Besides HTS libraries in which diversity and drug-likeness of the individual molecules are the main design criteria, targeted libraries play a role in drug design. A targeted library is designed such that the expected number of compounds binding to a specific target protein is increased. Docking algorithms can be used for solving various kinds of library design problems: Evaluation of a library (docking and scoring of all library molecules), pre-selection of R-group molecules to reduce the size of the library, or the *de novo* design of a library based on reactions and a database of available compounds. As in *de novo* ligand design, docking molecular fragments is a subproblem in several combinatorial library design problems. Due to the internal structure of a combinatorial library (the same set of fragments are connected in a well defined way in different combinations), drastic speed-ups can be achieved compared to docking a classical compound collection.

## 7.2

### Methods for protein–ligand docking

The following section contains a survey of algorithms applied to the various types of docking problems as well as a short summary on scoring functions. The methods are typically related to specific software tools also mentioned here.

## 7.2.1

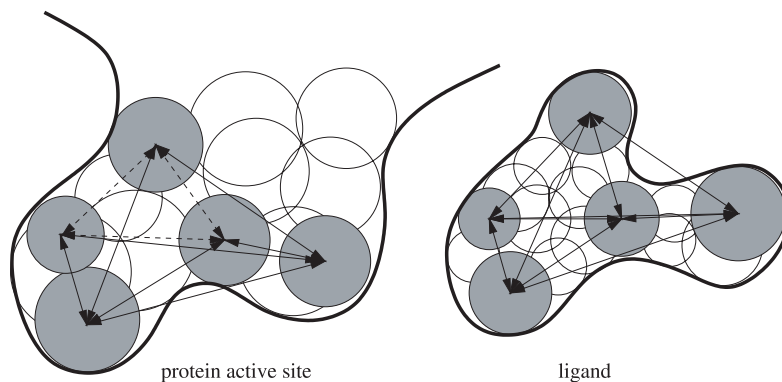
**Rigid-body docking algorithms**

Rigid-body docking algorithms have historically been the first approaches for screening sets of ligands by their fit to a given target protein. The protein as well as the ligand are held fixed in conformational space which reduces the problem to the search for the relative orientation of the two molecules with lowest energy.

**7.2.1.1 Clique-search based approaches**

The docking problem can be understood as a problem of matching characteristic features of the two molecules in three-dimensional space [20]. A match is an assignment of a ligand feature to a protein feature. Such a feature can either be a piece of volume of the active site of the protein or the ligand or an interaction between the molecules. The search procedure maximizes the number of matches under the constraint that they are compatible in three-dimensional space, i.e. that they can be realized simultaneously. In other words, compatibility means that a transformation can be found which simultaneously superimposes all ligand features to the matched protein features. In order to search for compatible matches, the following graph  $G$  is used: The vertices of  $G$  are all possible matches between the protein and the ligand, the edges connect pairs of vertices representing compatible matches. Mostly, compatibility is implemented as distance compatibility within a fixed tolerance  $\varepsilon$ : The matches  $(p_1, l_1), (p_2, l_2)$  are distance-compatible if and only if  $|d(p_1, p_2) - d(l_1, l_2)| < \varepsilon$ . A necessary condition for a set of matches to be simultaneously realizable is that all pairs of matches are distance compatible. Therefore, an algorithm for enumerating cliques (fully connected subgraphs) can be applied to  $G$ . By superimposing the matched features of a clique one obtains an initial orientation of the ligand molecule in the active site of the protein.

One of the historically first and today probably most widely used software tool for molecular docking, the DOCK program [21], is based on the idea of searching distance-compatible matches. Starting with the molecular surface of the protein [22, 23, 24], a set of spheres is created inside the active site as shown in Figure 7.2. The spheres represent the volume which could be occupied by the ligand molecule. The ligand is either represented by spheres inside the ligand or directly by its atoms. In DOCK, an enumeration algorithm searches for sets with up to four distance-compatible matches. Each set is used for an initial fit of the ligand into the active site. Then the set is augmented by further compatible matches and the position of the ligand is optimized and scored. Since its first introduction, the DOCK software has been extended in several directions. The matching spheres can be labeled with chemical properties [25] and distance bins are used to speed up



**Fig. 7.2** On the left, a protein active site is shown filled with spheres as they are used in the DOCK algorithm. On the right, a ligand covered with spheres is shown. An example for a distance compatible matching is highlighted in grey. All distances compared are shown by arrows.

the search process [26, 27]. Recently, the search algorithm for distance-compatible matches changed to the clique-detection algorithm introduced by Kuhl [20, 28].

An interesting algorithmic extension was introduced by Knegtel [29]. Instead of using a single protein, an ensemble of protein structures is used for the docking calculation. With this approach limited protein flexibility can be taken into account. Two of the flexible ligand docking approaches based on DOCK will be introduced below. Furthermore, several scoring functions are applied in combination with the DOCK algorithm [30–33].

While the initial DOCK algorithm uses volume as the feature to be matched, other approaches use chemical interactions. Mizutani et al. [34] presented the program ADAM in which hydrogen bonding is the feature used for matching. Possible matchings are enumerated and filtered based on distance compatibility.

Further examples for distance-compatibility as an initial screen for fragment orientations are the well known *de novo* ligand design program LUDI [35, 36] and the rigid body docking program CLIX [37]. While LUDI's placement algorithm is based on matching hydrogen bond vectors and hydrophobic points, CLIX takes energetically favorable regions for functional groups of the ligand as the underlying model. These regions are precalculated with the computer program GRID [38] which will be introduced in Section 7.2.5. CLIX uses only two points for an initial matching. After fitting the two matched sites, the rotation about the common axis between the matched sites remains as an open degree of freedom. This rotation is then sampled in regular intervals.

### 7.2.1.2 Geometric hashing

Geometric hashing [39] originated from the area of computer vision and was first applied to molecular docking problems by Fischer et al. [40, 41]. In computer vision, the geometric hashing scheme was developed for the problem of recognizing (partially occluded) objects in camera scenes. For simplicity, we explain the geometric hashing algorithm for the two-dimensional case first and describe its application to the three-dimensional molecular docking afterwards.

Given a picture of a scene and a set of objects which can occur therein (called models in this context) both represented by points in 2D space, the goal is to recognize some of the models in the scene. In a preprocessing phase, a hash table is created from the set of models: For each model, each pair of points define a so-called basis. Then, for each basis, every third point belonging to the model is expressed in coordinates relative to the basis. A tuple (model, basis) is stored in a hash table addressed by the relative coordinates of the third point. The reason for having several bases for a model instead of a single one is that it is unknown in advance, whether a part of the model is occluded in the scene.

In the recognition phase the scene is analyzed as follows: Every pair of points is considered as a basis. Once the basis is defined, all other points can be expressed by relative coordinates with respect to the basis resulting in a query for the hash table created before. The query votes for all matching tuples (model, basis) stored in the hash table. Finally, models with many votes are extracted, a transformation is calculated from the matching points, and the match is verified.

Two aspects make geometric hashing attractive for molecular docking problems: it is time-efficient and it deals with partial matchings standing for partially occluded objects in the terms of pattern recognition. The latter is extremely important because in most docking applications, not all of the ligand features are matched with the protein since parts of the ligand surface are in contact with bulk water.

In order to apply geometric hashing to molecular docking, Fischer et al. [40, 41] used the sphere representation of DOCK as the underlying model. Because docking is performed in 3D space, three points (here spheres or atoms) are necessary to define a basis. As a consequence, the number of hash table entries increases with the fourth power of the number of ligand atoms resulting in unacceptably large hash tables. Therefore, the basis is described by only two points leaving one degree of freedom open (rotation around the axis defined by the two points). With this model in mind, the geometric hashing approach can be directly applied to the molecular docking problem.

### 7.2.1.3 Pose clustering

Pose Clustering [42] is a different approach from pattern recognition applied to the molecular docking problem [43]. Pose clustering was originally



developed for detecting objects in pictures with an unknown camera location. The algorithm matches each triangle of object points with each triangle of points from the picture. From a match, a camera location can be computed such that the triangles superimpose. The camera locations are stored and clustered. If a large cluster is found, the object is identified and the orientation of the camera with respect to the object is determined.

For applying pose clustering to molecular docking, the LUDI model of molecular interactions [35, 36] is used as the underlying representation. For each interacting group, an interaction center and an interaction surface is defined (see Figure 7.3). The interaction surfaces of the protein are approximated by discrete points, which then form the scene in the pose clustering algorithm. The centers of the ligand interactions are the object points which have to be matched to the scene.

While in the pattern recognition problem each triangle of object points can be matched to each triangle of picture points, in the docking application, the matches are limited in various ways. First of all, the interaction types must be compatible and secondly, the triangle edges must be approximately of the same length. A hashing scheme is necessary to efficiently access matching protein interaction point triangles (picture points) for a given ligand interaction center triangle (object points). The hashing scheme stores edges between two points and addresses them by the two interaction types and the edge length. A list merging algorithm then creates all triangles based on lists of fitting triangle edges for two of the three query triangle edges. For a match, additional directionality constraints for the three interactions are checked. Then, a transformation is calculated that superimposes the two triangles. In the original application of pose clustering, the transformation is used to calculate the camera location. In molecular docking, the transformation directly describes the location of the fragment in the active site.

Finally, the transformations must be clustered. We use complete-linkage hierarchical clustering for this task. As a distance measure, the root-mean-square deviation of the atoms after applying the transformations is used. After a linear-time preprocessing phase, this quantity can be calculated from the transformations in constant time [43]. For each of the clusters generated, the typical post-processing steps are performed, like searching for additional interactions, checking for protein–ligand overlap, and scoring.

#### 7.2.1.4 Superposition of point sets

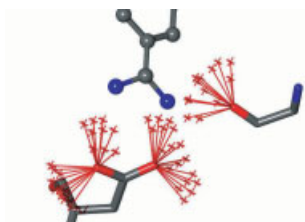
In each of the introduced algorithms for rigid-body docking, the superposition of point sets is a fundamental subproblem that has to be solved efficiently and therefore is discussed here shortly:

The superposition problem can be described as follows: Given two sets  $X, Y$  with  $n$  vectors each, find a transformation  $T = (\Omega, t)$  minimizing the root-mean-square deviation between  $X$  and the transformed vector  $Y$ :

$$RMSD_{X,Y}(T) = \sqrt{(1/n) \sum_i (x_i - \Omega y_i - t)^2}.$$



**Fig. 7.3**  
Interaction surfaces of three hydrogen bond acceptors in thrombin (shown in red). Protein atoms are drawn with sticks, ligand atoms with balls &



sticks. The left picture shows the interaction surfaces itself, the right picture shows the approximation by interaction points used in the pose clustering algorithm.

Let  $\Omega$  describe a rotation around the centroid of  $Y$ , then  $t' = (1/n) \cdot (\sum_i x_i - \sum_i y_i)$  minimizes  $RMSD_{X,Y}(T)$  for all rotations  $\Omega$ .

Optimizing  $\Omega$  is a more difficult task. Ferro & Hermans [44] and later Sippl and Stegbuchner [45] proposed iterative algorithms rotating subsequently around the  $x$ -,  $y$ -, and  $z$ -axis. If the axis is fixed, the optimal rotation angle can be determined analytically. Kabsch [46, 47] formulated the problem as a constrained optimization problem. Using Lagrange multipliers and the calculation of eigenvectors, Kabsch was able to solve the problem directly. Kabsch's algorithm is very time-efficient and contained in several codes for molecular docking today.

#### 7.2.1.5 Concluding remarks

Reviewing the development of more recent approaches to fragment docking, two enhancements were made. First, more elaborate algorithms for searching in three-dimensional space, often adapted from other disciplines, were applied. These methods allow a reasonable coverage of search space in short computing times. Second, more information about the physico-chemical properties of binding are included directly into the search process. Therefore, energetically unfavorable fragment placements are directly avoided, even if they make sense from a steric point of view. Both enhancements are essential for achieving a good performance for large variety of molecular fragments.

#### 7.2.2

##### Flexible ligand docking algorithms

The major limitation of the rigid-body docking algorithms is that the conformational flexibility of the ligand molecule is not considered. Often small molecules have large conformational spaces with several low energy states. Even for small molecules with only a few rotatable bonds significant differ-

ences between the bound conformation and the calculated low energy conformation can occur [48].

### 7.2.2.1 Conformation ensembles

In principle, ligand conformational flexibility can be incorporated by applying rigid-body docking algorithms to ensembles of rigid structures, each representing a different conformation of the same ligand. The size of the ensemble is critical, since the computing time increases linearly with the number of conformations and the quality of the result drops with increasing difference between the most similar conformation of the ensemble and the complex conformation.

Miller and Kearsley developed the Flexibase/FLOG docking algorithm based on conformation ensembles. Flexibases [49] store a small set of diverse conformations for each molecule from a given database. The conformations are created with distance geometry methods [50, 51, 52] which will be introduced later in the context of docking. Then a set of up to 25 conformations per molecule are selected by RMSD dissimilarity criteria. Each conformation of a molecule is then docked using the rigid-body docking tool FLOG [53] which is similar to the DOCK algorithm discussed above.

A different approach based on conformation ensembles was presented by Lorber and Shoichet [54]. Here, about 300 conformations per molecule are created on the average for a database of molecules. For each molecule, a rigid part, for example an aromatic ring system, is defined. The conformation ensemble is created such that the atoms of this rigid part are superimposed. Then, the DOCK algorithm is applied to the rigid part and all conformations are subsequently tested for overlap and finally scored. With this method, a significant speedup can be achieved compared to an independent docking of the conformations.

An important point in this scheme is the dependence between the conformation generation algorithm and the docking algorithm: The fewer conformations are created, the higher must be the tolerance in the matching phase of the docking algorithm. Therefore, the algorithms applied to the two subproblems are related making it necessary to tune parameters describing the coverage of conformational space in accord with the tolerance in protein–ligand overlap.

### 7.2.2.2 Flexible docking based on fragmentation

A possible way to handle conformational flexibility directly in a docking algorithm is by fragmentation. Here the ligand is divided into several fragments. Each fragment is either rigid or has only a small number of conformations which can be handled by a conformation ensemble. Obviously, the fragment-based docking approaches and *de novo* ligand design are closely related. The major difference is that while in the docking algorithm

the fragments stem from a single molecule, the *de novo* ligand design algorithm picks a fragment out of a database.

#### 7.2.2.2.1 Place & join algorithms

Looking at the way in which the fragments are reconnected during the docking calculation, we can distinguish between two classes of algorithms: In a place & join algorithm, multiple fragments are all docked independently. Then, placements for adjacent fragments in which the connecting atoms overlap are identified and reconnected.

The first algorithm of this kind was developed by Desjarlais et al. [55]. The ligand is manually divided into two fragments having one atom in common. Then, placement lists are created for each fragment using the docking algorithm DOCK. The algorithm searches through these lists for placement pairs in which the common atom is located approximately at the same point. Finally, the fragments are reconnected, energy minimized, and scored.

Sandak et al. [56–59] applied the geometric hashing paradigm to develop a place & join algorithm. As before, the ligand is divided into fragments with one overlapping atom, called the *hinge*. For each ligand atom triplet of a fragment, a hash table entry is created and addressed with the pairwise distances between the atoms. The entry contains a fragment identification as well as the location of the hinge. In the matching phase, protein sphere triplets are used to extract ligand atom triplets with similar distances. As a result a vote is counted for a hinge location for each match. Hinge locations with many votes are then selected, the fragments are reconnected accordingly and finally scored.

Place & join algorithms are advantageous in cases in which the molecule consists of a small set of medium-sized rigid fragments. If the fragments are too small, it is difficult to place them independently. Another difficulty is to generate correct bond lengths and angles at the connecting atom without destroying the previously found interactions of the fragments to the protein.

#### 7.2.2.2.2 Incremental construction algorithms

The second kind of fragment-based docking algorithms, and the one which is used most frequently today, follows the incremental construction method. Instead of placing all fragments of the ligand independently, the incremental construction algorithm starts with placing one fragment (called base or anchor fragment) into the active site of the protein. Then, the algorithm adds the remaining parts of the ligand to the already placed fragment iteratively. Thus, an incremental construction algorithm has three phases: The selection of base fragments, the placement of base fragments, and the incremental construction phase. An incremental construction algorithm can start with several base fragments, but in contrast to the place & join algo-

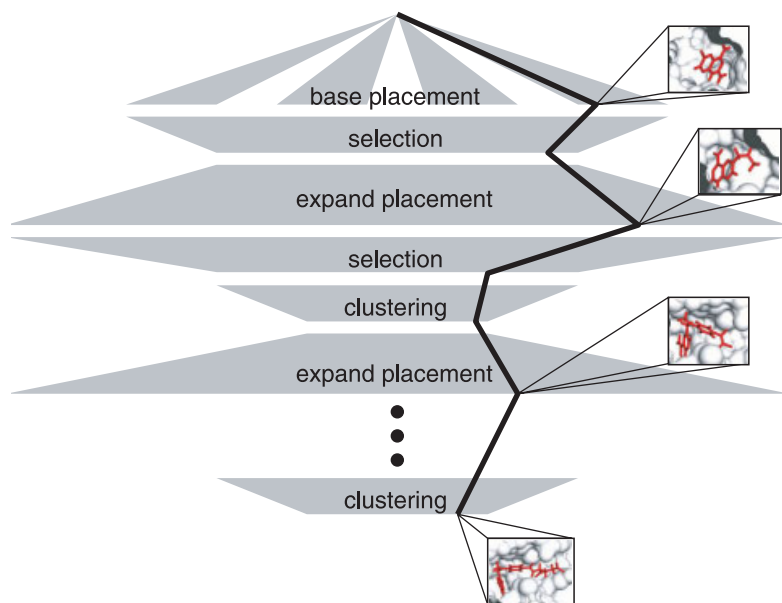
gorithms the placements are not combined but taken as anchoring orientations to which the remaining parts of the ligand can be added.

Incremental construction originated from the area of *de novo* ligand design. Moon and Howe [60] presented the peptide design tool GROW based on this strategy. The first docking algorithm based on incremental construction was developed by Leach & Kuntz based on the DOCK program [61]. First, a single anchor fragment is selected manually and docked into the active site using a variant of the DOCK algorithm which handles hydrogen bonding features in the matching phase. A subset of placements is selected for which the incremental construction phase is started. For this phase, a backtracking algorithm is used enumerating the space of non-overlapping placements of the whole ligand in the active site. After adding a fragment to the current placement, a refinement routine is used to eliminate steric strain and to improve hydrogen bond geometries. The final placements are then filtered, refined and scored with a force-field based approach. Although there are several manual steps in this procedure, the work demonstrated that the incremental construction idea can be applied to the docking problem.

Leach published a second docking algorithm [62] similar to incremental construction with respect to the way that the degrees of freedom are fixed sequentially. Degrees of freedom considered are the ligand orientation and conformation described by a discrete set as well as a set of rotamers for selected protein side-chains. Leach used a variant of a branch & bound scheme, called *A\* algorithm with dead-end elimination* (see Appendix), to search efficiently through the space of possible configurations.

The docking algorithm contained in the FLEX package [63–65] is also based on incremental construction. FLEX is a fully-automated approach to molecular docking developed for virtual screening purposes. In the first phase, a small number of base fragments are selected. An efficiently computable scoring function is used to select fragments which are suitable for placement. They should contain a reasonably high number of interacting groups and at the same time a relatively low number of low-energy conformations. A necessary condition for a successful calculation is that the selected base fragment binds to the protein and is not mostly exposed to water in the final protein-ligand complex. In order to ensure this, a small set of base fragments distributed over the ligand is selected.

For placing the base fragments, the pose clustering algorithm is applied. Base fragment conformations are enumerated within the placement algorithm. The advantage of the pose clustering algorithm is that it is based on the molecular interactions instead of the shape of the fragment. This allows the handling of much smaller fragments than in shape-based algorithms down to the size of a single functional group. All calculated placements up to given number form the input to the incremental construction phase. In contrast to Leach's algorithm, a greedy strategy is applied selecting always the  $k$  placements with the highest estimated score ( $k$  is about 800). Each



**Fig. 7.4**  
Flexible ligand docking by incremental construction. The search tree resulting after the steps of the algorithm as

described in the text are shown in grey. The black line illustrates the construction process for one placement of the complete ligand.

iteration of the incremental construction algorithm contains the following steps: adding the next fragment in all possible conformations to all placements from the previous iteration (or the base placement phase), searching for new protein–ligand interactions, optimizing the ligand position to improve the interaction geometries and reduce steric strain, selecting a subset of placements with high score, and clustering the placements to achieve a reasonable degree of diversity in the solution set. The overall search strategy is shown in Figure 7.4.

Ligand conformations within FLEXX are based on the MIMUMBA model [66]: To each rotatable bond, a set of low energy torsion angles is assigned, previously derived from a statistical analysis of the Cambridge Structure Database (CSD) [67]. Ring system conformations are precomputed using the 3D structure generator Corina [68, 69]. For scoring protein–ligand complexes, a variant of Böhm’s empirical scoring function [70] is used.

Several extensions of the FLEXX approach have been developed recently. The interaction model has been extended such that hydrophobic fragments can be handled with the pose clustering algorithm [71]. In order to place critical water molecules and metal ions located in the protein–ligand interface, the “particle concept” has been invented and integrated into the FLEXX software. The particle concept allows for automatic placement and energetic consideration of approximately spherical objects during the docking calculation [72]. FLEXE is an extension of FLEXX regarding the handling of

protein flexibility [73, 74]. Similar to Knegtels approach mentioned above, FLEXE takes an ensemble of slightly differing protein structures as input. However the structures are not numerically merged to a single description. FLEXE uses various graph algorithms to explicitly consider the different alternative conformations for protein parts like side-chains or small loop fragments and combine them to a single protein conformation best suited for the protein–ligand complex created during the docking calculation.

Two other approaches based on incremental construction have been published. The program Hammerhead [75] differs from FLEXX in the construction strategy. Instead of adding small fragments (cut between each rotatable bond), the ligand is divided into a small set of large fragments. During the construction phase, the next fragment is added such that the connecting atom (or bond) overlaps and interactions to the protein can be formed. Therefore, there is no discrete sampling at the torsion angle of the added fragment. However, since the bond angle at the overlapping atom may vary, high-energy conformations will also be generated and the situation in which a fragment does not interact directly with the protein is more difficult to handle.

Makino and Kuntz [76] presented a fully automated incremental construction docking algorithm based on backtracking. A single anchor fragment is selected maximizing hydrogen bonding features. During the incremental construction phase, the number of conformations for each fragment is limited to reduce the size of the search space. This method is called a limited backtrack search. For scoring, the AMBER force field [77, 78] is used with a modification allowing the handling of multiple protonation states.

Incremental construction algorithms are the basis for widely used docking tools like DOCK and FLEXX. The quality of the predicted structures strongly depends on the number of different placements considered in each iteration of the incremental construction process. Therefore, although the overall concept of incremental construction is simple, much effort must be put in the time-efficient analysis of partially placed ligands, for example the protein–ligand overlap test, the evaluation of the scoring function, and the postoptimization of the ligand placement.

In practice, incremental construction has been proven to be a reasonable compromise between accuracy and computing speed. This holds especially for virtual screening applications, in which computing speed is of central importance.

### 7.2.2.3 Genetic algorithms and evolutionary programming

Since the mid-1990s, genetic algorithms have been applied to the molecular docking problem in several approaches [79–83]. A genetic algorithm [84] is a general purpose optimization scheme which mimics the process of evolution. The individuals are configurations in the search space. A so-called fitness function is used to decide which individuals survive and produce offsprings.

Several elements must be modeled in order to use the idea of genetic algorithms for an application like molecular docking. First of all, a linear description of a configuration (the chromosome) is needed describing all degrees of freedom of the problem. Finding the chromosome description is the most difficult modeling part. A suitable description is free of redundancy and models constraints of the configuration space directly such that configurations violating constraints are never generated during the optimization.

Second, a fitness function has to be developed. The fitness function is closely related to scoring functions for molecular docking with one extension. Scoring functions normally work on 3D coordinates, therefore the chromosome of an individual has to be interpreted in order to apply the scoring function. This step is called the *genotype-to-phenotype conversion*. Since most of the computing time is spent for evaluating the fitness function, the conversion and the evaluation has to be done efficiently.

The optimization scheme itself is more or less independent from the application. Typically, several parameters have to be chosen like the population size, the number of generations, crossover and mutation rates, etc.. Here, it is important to achieve a reasonable trade-off between optimizing the fitness function and keeping the diversity in the population.

The genetic algorithm which is today probably most widely applied for molecular docking was developed by Jones et al. [79, 85] and is implemented in the software tool GOLD. A configuration in GOLD is represented by two strings. The first string stores the conformation of the ligand and selected protein side chains by defining the torsion angle of each rotatable bond. The second one stores a mapping between hydrogen bond partners in the protein and the ligand. For fitness evaluation, a 3D structure is created from the chromosome representation by first generating the ligand conformation. According to the mapping stored in the second string, hydrogen bond atoms are superimposed to hydrogen bond site points in the active site. Finally, a scoring function evaluating hydrogen bonds, the ligand internal energy as well as the protein–ligand van der Waals energy is applied as the fitness function.

Oshiro et al. [80] developed two variants of a docking method, both based on genetic algorithms and the DOCK approach. The variants differ in the way the relative orientation of the ligand to the protein is described. The first variant is similar to the GOLD algorithm and encodes the matching of ligand atoms to protein spheres in the chromosome. A superposition is used to generate the 3D orientation of the ligand. The second variant stores the relative orientation directly by a translation vector and three Euler angles. For scoring, a simplified version of the AMBER force field was used.

Gehlhaar et al. [82, 86] developed a docking algorithm based on evolutionary programming called EPDOCK. In contrast to a genetic algorithm, offspring are created from one parent by mutation only. Each member of a population is competing for survival in a so-called tournament. EPDOCK contains a self-developed scoring function based on atomic pairwise linear potentials for steric interactions and hydrogen bonding.



#### 7.2.2.4 Distance geometry

Distance geometry [51, 50] is a well-known technique from the area of structure determination via NMR technology. Instead of describing a molecule by coordinates in Euclidean space, it is described by a so-called distance matrix containing all interatomic distances. Based on distance matrices, a set of conformations can be described in a comprehensive form by calculating a distance interval for each atom pair.

The distance geometry methodology can be directly used in the docking algorithm based on clique search and distance compatibility (see Section 7.2.1.1). Two matches between protein site points and ligand atoms are compatible, if the site point distance lies within the distance interval of the ligand atom pair. The drawback of this approach is that the distance matrix is overdetermined. Fixing the atom-atom distance between a given atom pair to a single value causes other distance intervals to shrink. Since the exact new interval boundaries are difficult to calculate, the triangle and tetrahedron inequality are used to approximate them [87, 88]. In other words, only a very limited number of distance matrices can be converted back to 3D space.

Ghose and Crippen [89] first worked on this approach on a more theoretical basis. Later, Smellie et al. [90] applied this methodology to real test cases. Billeter et al. [91] combined the description of molecules by distance constraints with an efficient algorithm for constrained optimization.

The screening software Specitope developed by Schnecke et al. [92] uses distance matrix comparisons as a first filter step. However, flexibility of molecules is not modeled by distance intervals. Instead, a weighting scheme is defined to scale down the contributions of more flexible atom pairs in the overall score.

#### 7.2.2.5 Random search

Once a scoring function for evaluating protein-ligand complexes is available, random search algorithms can be applied to the docking problem. Random placements can be either created directly by randomly fixing all degrees of freedom or they can be derived from a (random) starting structure by random moves. In most cases the structure generation is combined with a numerical optimization driving the placements to the closest local minima. Most approaches of this kind are Monte Carlo algorithms discussed separately in Section 7.2.3.3.

Sobolev et al. [93] presented a random search algorithm in their docking program LIGIN. A large set of starting structures are created randomly and then optimized in two steps. The surface complementarity is optimized first, the hydrogen bond geometry in a second step. So far, LIGIN does not include ligand flexibility although the method is in principle able to handle it.

In order to avoid a repeated evaluation of very similar structures, Baxter et al. [94] use a method called *tabu search* in their docking software called

PRO\_LEADS. Starting with an initial random structure, new structures are created by random moves. In tabu search optimization, a list (the tabu list) is maintained containing the best and most recently visited configurations. Moves which results in configurations close to one in the tabu list are rejected except if they are better than the best scoring one. The tabu-list technique improves the sampling properties of the random search algorithm by avoiding to revisit configurations. (see also Appendix, “Tabu search”)

### 7.2.3

#### Docking by simulation

While all previous methods for the docking problem are based on some kind of combinatorial optimization algorithm, there are several approaches tackling the problem by simulation techniques. Instead of trying to enumerate a discrete low-energy subspace of the problem, these approaches begin their calculation with a starting configuration and locally move to configurations with lower energy.

##### 7.2.3.1 Simulated annealing

Simulated annealing [95] is a well-known simulation technique which is also frequently used for solving complex optimization problems without any physical interpretation of the simulation itself. The overall simulation routine iterates the following steps. Starting with a configuration  $A$  with an energy or score value  $E(A)$ , a random local move to a new configuration  $B$  with energy  $E(B)$  is calculated. The acceptance of the new configuration is based on the Metropolis criterion, which means that it is accepted if  $E(B) \leq E(A)$  or with probability  $P = e^{-(E(B)-E(A))/(k_B T)}$  otherwise where  $k_B$  is the Boltzmann constant. Over simulation time, the temperature  $T$  is reduced based on a so-called cooling schedule such that accepting configurations with increased energy becomes less likely. (see also Appendix, “Simulated annealing”)

The AUTODOCK program for protein-ligand docking developed by Goodsell et al. [96, 97, 98] is based on this strategy. For energy calculation, molecular affinity potentials [38] are precalculated on a grid. Yue developed a program for optimizing distance constraints for rigid-body docking based on simulated annealing [99].

##### 7.2.3.2 Molecular dynamics

In principle, molecular docking problems can be solved with molecular dynamics (MD) simulations. In fact, the earliest approaches for predicting protein-ligand interactions with the computer were based on MD calculations [100, 101].

In MD, a force field is used to calculate the forces on each atom of the simulated system. Then, following Newtonian mechanics, velocities and accelerations are calculated and the atoms are moved slightly with respect to a given time step. Introducing molecular dynamics and force fields is clearly beyond the scope of this article. However, some aspects of docking by MD simulations should be mentioned briefly. The simulation becomes more exact, the smaller the time step and the more atoms of the system are taken into account. Thus, MD simulations can become very time consuming and are therefore not appropriate for inspecting large sets of molecules. (see also Appendix, “Molecular dynamics”)

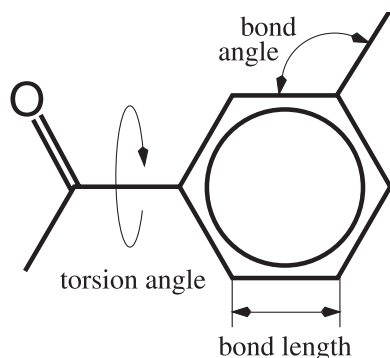
In order to avoid very long computing times, methods performing greater moves of the ligand in a single step have been developed. This decreases the dependence of the outcome of the calculation from the starting structure and allows a better sampling of the search space. DiNola et al. [102, 103] developed a technique for this purpose called ‘helicopter view’. For a limited time, the temperature of the system is increased for selected degrees of freedom (protein-ligand relative orientation) and the repulsive terms of the energy function are decreased. This enables the algorithm to escape from local minima in the energy function.

A similar method is used in a four-phase docking protocol developed by Given and Gilson [104]. First, a set of low-energy ligand conformations is created using MD with alternated heating (in order to perturb the structure) and cooling (in order to minimize the structure). Then the ligand is placed randomly into the active site and several times minimized. In the final phase, the most stable configurations are investigated further using MD with alternate heating and cooling. The goal of the last phase is to explore the search space around the stable conformations in more detail.

A frequently used technique to speed up MD simulations is the precalculation of force-field contributions from protein atoms on grids. The force on a ligand atom can then be efficiently calculated by a simple table-lookup instead of summing over all protein atom contributions. The potentials, however, can only be precalculated for atoms which do not change their orientation in space. In order to avoid completely neglecting protein flexibility, Luty et al. [105, 106] divided the protein into a rigid and a flexible part. Every atom sufficiently far away from the active site is considered as fixed in space, all force-field contributions of these atoms can be precalculated on a grid. During the MD simulation, only the active site atoms of the protein and the ligand atoms are allowed to move and have to be considered explicitly in the force-field calculation.

### 7.2.3.3 Monte-Carlo algorithms

In an MD simulation, the local movements of the atoms are performed due to the occurring forces. In contrast, in a Monte-Carlo (MC) simulation, the local moves of the atoms are performed randomly. The simulated annealing



**Fig. 7.5** Internal coordinate representation of a molecule. The three-dimensional structure can be constructed from bond lengths, bond angles, and torsion angles.

algorithm discussed above is one special variant of an MC simulation. Two components are of major importance in the development of an MC algorithm: The description of the degrees of freedom and the energy evaluation.

Concerning the degrees of freedom, the aim is to describe them such that high-energy states are avoided. A good example of how to realize this concept is the description of the conformational space of a molecule by internal coordinates (bond lengths, bond angles and torsion angles) shown in Figure 7.5 instead of by Cartesian coordinates for all atoms. With internal coordinates, each type of variable is related to a different energy scale: Changing a bond length is energetically more expensive than changing a bond angle, which in turn is energetically more expensive than changing a torsion angle. Using internal coordinates, it is possible to navigate through the low-energy conformational space by defining the amounts by which the variables of each type are changed. Thus, internal coordinates are the more appropriate description form for MC algorithms. (see also Appendix, “Monte Carlo” Methods)

Most of the computing time during an MC simulation is spent in the calculation of the energy (or score) of a state. Therefore, this step has to be as time-efficient as possible. Often, energy potentials are precalculated on a grid to speed up this step.

There are several examples for MC-based docking algorithms. Hart and Read [107] developed an MC scheme combined with simulated annealing. In the MC run, random orientations are created and moved such that protein-ligand overlap is reduced. The results are then optimized in a simulated annealing optimization scheme. Protein-ligand overlap and scores are calculated based on precalculated grids. McMartins and Bohaceks QXP program [108] implements a similar approach which can be applied to molecular docking and structural superposition. Wallqvist and Covell [109] also use MC for optimizing the final ligand orientation. Instead of a random set of starting structures, a surface matching algorithm is used.

Abagyan et al. [110] developed the software package ICM combining the MC algorithm with an internal coordinate description explained above. In

contrast to other approaches making purely random moves, ICM is able to make moves based on probability distributions for variable sets. For example, a probability distribution for a torsion angle can be defined such that low-energy torsion angles are more likely to occur than high-energy torsion angles. This biases the MC calculation towards low-energy states.

MC algorithms can be used to overcome the limitation of MD simulations to get stuck in local minima. In order to get low energy structures, MC can then be combined with energy minimization as described by Apostolakis et al. [111]: After a random creation of starting structures, a minimization with a modified van der Waals potential is performed. The energy function for the evaluation of the conformations is a sum of three terms: a force field energy, a hydrophobic solvation term which is proportional to the solvent-accessible surface of the complex and an electrostatic solvation term obtained from the solution of the linearized Poisson-Boltzmann equation. The best scoring configurations are further analyzed with a Monte-Carlo simulation method called MCM. MCM performs Monte-Carlo interleaved with minimization steps and uses the energy function mentioned above.

PRODOCK is an MC-based docking algorithm developed by Trosset and Scheraga [112, 113, 114]. The software uses either the AMBER IV or ECEPP/3 force field with a grid-based energy evaluation. For calculating energy values within the grid, Bezier splines were used allowing for a more accurate estimation of the energy value as well as information about the derivatives at this point. Like ICM, an internal description of the degrees of freedom is used. The MC simulation is interleaved with energy minimization steps like in Apostolakis' approach.

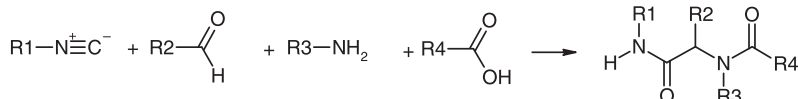
#### 7.2.3.4 Hybrid methods

Due to the complexity of the docking problem, all methods have their pros and cons. Fragment-based approaches and genetic algorithms achieve a wide coverage of the configuration space, however simulation-based methods outperform others in finding exact local minima, i.e. predicting an exact placement. Combining different methods is therefore a reasonable approach which can result in methods containing the best of each.

Two approaches have been published recently which combine rapid fragment-based searching techniques with sophisticated MD or MC simulations.

Wang et al. [115] developed a multi-step approach based on rigid-body docking and MD. First a set of low-energy conformers is created. Each conformer is docked rigid into the active site using DOCK. The high scoring placements are then optimized using an MD-based simulated annealing optimization in combination with the AMBER force field.

Hoffmann et al. [116] combined the incremental construction algorithm in FLEX with an MD-based procedure for post-optimization. First, FLEX is used to create a sample of a few hundred ligand placements. The goal of



**Fig. 7.6**  
Based on the Ugi reaction, combinatorial libraries with four different R-groups can be created. After the reaction, all library

molecules have the core shown on the right in common but differ in the four R-groups attached to it.

the second phase is to improve the overall ranking of the solutions and to identify the correct placement. The placements are first energy-minimized using the CHARMM force field. For re-ranking, the software package CAMLab [117] is used. The final score consists of three contributions, a force field energy, an electrostatic part, and a nonpolar part of the solvation energy. The electrostatic part of the solvation energy is calculated by solving the Poisson equation using fast multigrid methods, while the nonpolar part is approximated by the total solvent-accessible surface.

#### 7.2.4

#### Docking of combinatorial libraries

The development of combinatorial chemistry and its application to drug design [18, 19] has led to new search problems in the context of molecular docking. An example of a combinatorial library is given in Figure 7.6. The number of molecules which can be synthesized on the combinatorial chemistry bases has increased dramatically compared to classical methods. Therefore, any screening methodology has to face many more molecules.

Probably more important for the development of docking methods is the introduction of formal structure into this increased search space. If an unstructured compound collection is given, each molecule has to be analyzed independently in a screening experiment. Combinatorial libraries, however, follow a systematic build-up law for synthesizing molecules from a highly limited set of building blocks. This structure can be exploited to drastically reduce the runtime of virtual screening calculations.

In the context of combinatorial libraries, one can distinguish between three kinds of docking problems:

**Combinatorial Docking Problem** Given a library, calculate the docking score (and the geometry of the complex) for each molecule of the library.

**R-Group Selection Problem** Given a library, select molecules for the individual R-groups in order to form a smaller sublibrary with an enriched number of hits for the target protein.

**De novo Library Design Problem** Given a catalog of molecules, design a library (including the rules of synthesis) optimizing the number of hits for the target protein.

Methods for these problems have emerged from the area of molecular docking and *de-novo* ligand design (see [118] for an overview on combinatorial docking methods). In the first case, the docking algorithms are applied to individual molecule fragments like R-groups or the core of the library and the resulting information is then combined yielding placements for individual library molecules. The methods for combining the placements differ. Like in the case of docking algorithms, they can also be classified as either a place & join or an incremental construction method. In the second case, the *de-novo* ligand design is constrained by predefined rules of synthesis.

Early algorithms for the combinatorial docking problem analyzed the similarity in given ligand datasets in order to speed up the search process. The focus in these papers is to structurally relate ligands within the dataset. One approach to do so is to generate a minimal tree structure representing the whole ligand dataset [119]. Another approach is to speed up conformational searching based on clustering similar molecules [120]. In both cases, the derived hierarchy of molecules can then be used in an incremental construction docking method.

The combinatorial docking tools PRO\_SELECT [121] and CombiDOCK [122] are based on the incremental construction method. In both approaches, a library is formed by a template (or core) molecule with a set of attachment points to which one out of a predefined set of substituents can be connected. The template is then positioned in the active site without considering the substituents. Starting from a few orientations of the template, the substituents are placed into the active site of the protein independently. In case of PRO\_SELECT, substituents are then selected based on score and additional criteria like 2D similarity and feasibility of synthesis. CombiDOCK calculates a final score for whole library molecules by combining fragment scores.

A new algorithm for the combinatorial docking problem based on the FLEX incremental construction algorithm was recently presented [123]. The algorithm is part of a new combinatorial docking extension of FLEX called FLEX<sup>c</sup>. FLEX<sup>c</sup> handles the library as a rooted tree in its closed form. Molecules from the library required during the docking calculations are created on the fly. With this method libraries with a few hundred thousand molecules can be handled in main memory during a docking calculation. The tree representation allows for handling complex libraries consisting of several R-groups which can be arbitrarily linked with the one limitation that no ring closures over different R-groups are allowed. The recursive combinatorial library algorithm is a natural extension of the incremental construction method to multiple molecules: In each incremental construction step, all possible R-group molecules are added sequentially.

Several approaches based on ligand *de novo* design software have been published for R-group selection problems. Kick et al. [124] applied a variant of the BUILDER program [125] to the preselection of substituents for a

library targeted to Cathepsin D. Böhm [126] applied the LUDI program to the docking of two groups of fragments which can be connected pairwise in a single-step reaction to search for new thrombin inhibitors. In principle, all programs for fragment-based *de novo* ligand design can be applied in a similar way to the R-group selection problem.

Finally, we mention two methods for *de novo* library design. Caffisch [127] applied the MCSS technique generating fragment placements which are subsequently connected. The DREAM++ software [128] combines tools for fragment placement and selection. The selection process is done such that only a small set of well characterized organic reactions are needed to create the library.

### 7.2.5

#### Scoring protein–ligand complexes

A fundamental element of each software for molecular docking is the scoring function. The general question to answer is, given a protein and a ligand in aqueous solution, what is the ligand's free energy of binding to the protein? The scoring function topic is very complex, we will therefore give only a short overview about the main characteristics of the problem and the methods used. More comprehensive reviews can be found in [3, 4, 5, 129–132].

Computer methods are currently far from accurately calculating the free energy of binding. Typically, assumptions fixing some of the various degrees of freedom are made. Furthermore, in most approaches, it is assumed that the free energy of binding is dominated by a single, low-energy protein–ligand binding mode. Therefore, the energy is associated with a protein–ligand complex. Other aspects difficult to quantify like solvation effects, conformational changes in the protein, or loss of entropy during complex formation are either omitted or approximated only roughly.

With respect to the docking application, three kinds of problems can be defined which have to be solved by scoring functions with increasing complexity. The simplest problem is the ranking problem during a docking calculation. Here, different orientations of the same ligand in the same active site have to be evaluated and the scoring function has to distinguish energetically favorable orientations from less favorable ones. The most common problem in structure-based drug design is the ranking of a set of ligands with respect to the binding energy to a given protein. In contrast to the first problem, the scoring function has to deal not only with different orientations but also with different ligand structures. However, the protein is always the same and, at least in most cases, the ligands bind at approximately the same location, namely the active site. Also the necessary conformational changes of the protein during the binding of the ligand are likely to be similar simplifying the scoring problem. The most general case,



namely predicting the free energy of binding of individual protein–ligand complexes with different ligands and proteins, is of course the most complicated one.

In order to dock a single molecule, the scoring function has usually to be evaluated several thousand times. Therefore, only functions with short computing times can be applied. Currently, there are three types of scoring functions frequently used in molecular docking calculations: force-field scoring, empirical scoring, and knowledge-based (sometimes also called PMF for potentials of mean force) scoring.

Force fields (see [133] for an overview) can be applied directly to create a score for a protein–ligand complex. This scoring method is normally used in docking algorithms based on simulation (see above and [134]). Normally, some modifications are performed: First a grid is used to speed up the calculation. Second the Lennard-Jones potential is exchanged by a function that is not as steep, in order to account for small structural changes in the protein. While force field scoring performs quite well on selecting docking modes, it is difficult to compare different molecules on this basis. The main reason is that entropic and solvation effects are not considered. A combination of a force field score with an approximation of the solvation energy can improve the results but is computationally expensive [116].

A frequently applied program for force field scoring is GRID [38]. Instead of atom-based potentials, the software calculates interaction potentials on a grid for frequently occurring molecular groups based on a force field energy function. These maps can be either used for scoring or for detection and visualization of important interaction centers in the active site of the protein.

Empirical scoring functions are based on a correlation of geometric parameters of the protein–ligand complex to measured free binding energies. The overall score is a sum of terms representing different physico-chemical effects contributing to the binding energy like hydrogen bonding, hydrophobic contacts, etc.. For each term a functional form and a set of geometric parameters derivable from the protein–ligand complex are chosen. Usually, weights for the individual terms are determined from a set of protein–ligand complexes with known structures and binding energies using multiple linear regression or partial least squares.

The first function of this type was developed by Böhm [70] for the *de novo* design program LUDI [35, 36]:

$$\Delta G = \Delta G_0 + \Delta G_{rot} \times N_{rot} + \Delta G_{hb} \sum_{\text{neutral H-bonds}} f(\Delta R, \Delta \alpha) \\ + \Delta G_{io} \sum_{\text{ionic int.}} f(\Delta R, \Delta \alpha) + \Delta G_{lipo} |A_{lipo}|$$

Here,  $f(\Delta R, \Delta \alpha)$  is a scaling function penalizing deviations from the ideal geometry of hydrogen bonds and ionic interactions,  $N_{rot}$  is the number of

free rotatable bonds of the ligand, and  $A_{lipo}$  is the lipophilic contact area between the protein and the ligand. The terms  $\Delta G_0$ ,  $\Delta G_{rot}$ ,  $\Delta G_{hb}$ ,  $\Delta G_{io}$ , and  $\Delta G_{lipo}$  are adjustable parameters.

The Böhm function was further improved and customized for screening applications [70, 135, 136]. Since Böhm published his function, several other empirical scoring functions were developed [137–142]. The functions differ in the list of physico-chemical effects considered and the functional form for the individual terms.

Recently, three groups independently developed knowledge-based scoring functions for molecular docking [143–146]. This scoring method was previously used by Sippl [147, 148] for protein structure prediction and is successfully applied in this field. The idea underlying knowledge-based functions is that situations frequently seen in structures are energetically favorable. All three functions are based on inter-atom distances. For each atom type pair, the number of occurrences are counted depending on the distance. The resulting histograms are converted to an energy function. The score for a given complex is then calculated by adding the energy function values for all inter-atom pairs of the complex. Gohlke [146] combined the atom-pair score with a solvent-accessible surface term in order to take solvation into account. Mügge [145] added a volume correction factor to account for the ligand volume around each ligand atom which avoids interactions with the protein.

### 7.3

#### Validation studies and applications

All docking algorithms presented are heuristic algorithms giving no guarantee of the quality of the result. In addition, the underlying physico-chemical models are very rough. Therefore, an intensive validation of the docking algorithm is an essential requirement.

##### 7.3.1

#### Reproducing X-ray structures

The first test usually performed with a docking tool is the reproduction of complexes with known structures. The results achieved with docking algorithms are quite difficult to compare for several reasons. First, there is no unique benchmark set and every method is published with a small set of test results only. Second, different levels of *a priori* knowledge are used in the preparation of the input data and in the calculation. Examples for *a priori* knowledge are the definition of the active site, the placement of polar

hydrogens, the limitation of conformational or translational degrees of freedom, the removal of ligand parts located outside the active site, or simply the selection of considered test cases. Finally, it is a non-trivial task to reasonably quantify the quality of a docking calculation. For a single result, the RMSD (root mean square deviation) can be used. However, this value can be misleading if the molecule has nearly symmetric groups. In addition, the bound for what is called a 'good result' depends on the test case, important criteria are the size of the ligand and the overall hydrophobicity.

Two evaluation studies on large test sets can be found in the literature [85, 149]. Jones et al. applied the genetic algorithm GOLD to a set of 100 protein–ligand complexes taken from the PDB. Kramer et al. evaluated the FLEXX program with 200 protein–ligand complexes from the PDB. In general it can be said that about 65–70% of the test cases can be predicted in agreement ( $\leq 2.0 \text{ \AA}$  RMSD) with the crystal structure. This number varies with the flexibility of the ligand molecule: the larger and more flexible the ligand, the smaller is the chance to reproduce the crystallographic complex. This is not surprising, since the search space that has to be explored grows exponentially with the number of rotatable bonds in the ligand.

In both studies, the protein structure was taken in the complex conformation. One has to keep in mind that, in a real-case scenario, the protein can undergo conformational changes due to an induced fit during ligand binding. Obviously, the fraction of correct predictions will decrease with the amount of flexibility in the active site of the protein. In both papers, this problem is addressed in some additional calculations employing either protein structures crystallized without a ligand or structures of the same protein complexed with different ligands. A more detailed study on the influence of protein flexibility using the docking software PRO\_LEADS can be found in [150].

Finally, we mention two papers which directly compare different algorithms for protein ligand docking. Vieth et al. [151] compared genetic algorithms (GA) with Monte-Carlo (MC) and Molecular Dynamics calculations (MD). They conclude that GA works best within a given number of energy evaluations while MD gives more accurate predictions. Westhead et al. [152] compared four heuristic search strategies: a genetic algorithm, an evolutionary program, simulated annealing and tabu search. Their analysis showed that the genetic algorithm finds the placements with lowest energy but the tabu search created placements closer to the crystallographically observed complexes.

The results achieved with a specific method depend strongly on the modeling of the problem into the algorithmic framework. The individual approaches based on genetic algorithms presented above illustrate how much freedom for modeling the problem there still is although the overall optimization technique is fixed. Therefore, the comparisons cannot be considered as a general statement for or against an optimization method.

## 7.3.2

**Validated blind predictions**

Most protein–ligand docking programs are designed empirically. First, the scoring function is derived from experimental data. Second, the algorithms are designed with background knowledge about protein–ligand complexes. Therefore, either testing the program on a large set of examples or performing blind predictions is essential for evaluating the capabilities of the software. Testing a large set of examples is similar to a blind prediction in the sense that it is unlikely that information about most of the complexes has flown into the software development.

Performing a validated blind prediction is a difficult task and requires the availability of the software, a test scenario, and the experimental facilities for validating the study. For protein–protein docking, a blind prediction of the complex between TEM-1  $\beta$ -lactamase and the inhibitor BLIP was performed by 6 independent groups. All groups were able to identify the correct complex within 2 Å RMSD [153].

In CASP II (Critical Assessment of Methods of Protein Structure Prediction: Round II), a new prediction section on protein–ligand docking was introduced [154]. Several groups participated and submitted up to three models for some of the seven protein–ligand complexes (see [155–158] for reports). For each target, the 3D structure of the protein and the 2D structure of the ligand was given to the participants. All complexes were unpublished before the submission deadline.

The results show that accurate molecular docking is still a challenging problem. Although predictions within 3 Å were made for nearly all targets and within 2 Å for more than half of the cases, there is no method currently available that reliably and consistently predicts correct protein–ligand complex geometries. Dixon [154] concludes that, since the right solutions were often found but not ranked highest, there is a need to develop more accurate and reliable scoring functions.

## 7.3.3

**Screening small molecule databases**

The main application of molecular docking software is the virtual screening of compound databases for potential lead molecules. Testing docking software within this scenario is quite difficult and only a few papers discuss screening results with molecular docking software. Since software is often developed by groups not directly involved in drug design projects, experimental validation of the achieved screening results is not possible. Pharmaceutical companies that use docking software for preselecting compounds typically test the selected molecules such that a hit rate (the number of active molecules divided by the number of selected molecules) can be

calculated. However, the number of active molecules within the whole data set before the preselection is still unknown after this experiment such that selectivity (the number of actives in the selected subset) can be measured but sensitivity (the number of selected actives with respect to the number of active in the database) cannot.

A useful measure for evaluating methods for preselecting compounds is the enrichment factor which is the hit rate of the method divided by the random hit rate (which is the total number of active molecules divided by the size of the data set). The enrichment factor shows how much the method improves the screening result compared to a random selection. Note that for calculating the enrichment factor, one has to know the total number of active molecules which can only be guessed in test scenarios, i.e., the whole database except the known actives is assumed to be inactive. In the following, a number of papers containing different types of evaluations studies are summarized.

Burkhard et al. [159] developed a software tool for rigid-body molecular docking called SANDOCK. The method is based on matching surfaces with mapped properties and is not described in detail here. The authors applied SANDOCK to searching for new thrombin inhibitors. One of the hits was *p*-amino-benzamidine, a small molecule with ten non-hydrogen atoms, was experimentally validated. The structure predicted for the complex was in agreement with the crystallographic structure determined afterwards.

Shoichet et al. [33] presented an extension of the DOCK scoring function taking solvation effects into account. In order to test the function, a screening run of the ACD with about 150 000 molecules was performed for three different targets using the rigid-body variant of DOCK. Kramer et al. [149] applied FlexX for screening a small set of about 550 molecules against 11 targets with known nanomolar inhibitors. The screening data set was created by mixing ligands of 200 experimentally solved protein–ligand complexes with drug-related molecules selected from the CSD (Cambridge Structural Database) [67]. Knegt et al. [160] presented a screening application of DOCK for thrombin inhibitors and progesterone agonists. The compound collection used consists of about 1000 molecules. Since activity information was available for all molecules, enrichment factors could be calculated. They conclude that docking is effective in preselecting data sets, however the methods were not able to distinguish between true actives and similar inactives.

A comprehensive study on different scoring functions in combination with docking was recently presented by Charifson et al. [161]. In the study 13 different scoring functions in combination with two docking algorithms were applied to database searching. The database was formed from known inhibitors for three different targets mixed with 10 000 random molecules passing a drug-likeness filter. For each molecule, a set of conformers was generated and docked with DOCK and an in-house developed genetic algorithm called GAMBLER. The authors demonstrate that the number of hits

found depend strongly on the scoring function used and that there is not a single function with best performance. As a consequence, a consensus scoring method was developed showing better performance than the individual functions.

#### 7.3.4

##### **Docking of combinatorial libraries**

As in the case of database screening, the experimental validation of combinatorial docking methods is difficult to perform and therefore rarely found. The main reason is the lack of publicly available data on binding affinities for combinatorial libraries to proteins with known 3D structure.

Kick et al. [124] applied the *de novo* design program BUILDER [125] in a combinatorial fashion to the design of a focussed library for Cathepsin D. BUILDER is a fragment-based *de novo* design program related to the docking software DOCK discussed above. The program was used to evaluate putative R-groups for a given scaffold oriented in the active site of Cathepsin D. The created focussed library was synthesized and tested. Containing 30 compounds inhibiting Cathepsin D in the nanomolar range, the method improved the hit rate by a factor of 7.5 compared to a diverse library. The same library was used later to evaluate CombiDOCK [122].

A frequently used target protein for evaluating docking algorithms is thrombin already mentioned in the introduction. Böhm et al. [126] experimentally evaluated ligands proposed from a virtual combinatorial library by LUDI. They showed that five of ten proposed compounds exhibit binding affinities in the nanomolar range. Caffisch [127] and Murray [121] used human  $\alpha$ -thrombin for demonstrating their combinatorial docking and design tools MCSS-CCLD and PRO\_SELECT. They showed that ligands with well known binding patterns are created.

In order to evaluate FLEX<sup>c</sup> [123], we used three different data sets for thrombin and DHFR for which experimental binding data was available (thrombin: [162, 163], DHFR: [164]). Molecules in the DHFR data set were very closely related such that no correlation between predicted and experimental binding affinity was found. For the first thrombin data set, a correlation of 0.6 was achieved. For the second thrombin data set, a molecule with high binding affinity was known which could be identified at a high rank.

#### 7.4

##### **Molecular docking in practice**

In the previous sections, the algorithms for molecular docking and their performance on example data were discussed. In practical applications

however, several additional issues arise. Pre- and post-processing steps are necessary on the input and output data and a docking algorithm most appropriate for the application scenario has to be chosen.

#### 7.4.1

##### **Preparing input data**

Typically, the input data needed to perform a docking calculation is not in the appropriate format such that preprocessing steps are necessary. The situation is substantially different for the protein and the ligand molecule. All relevant application scenarios deal with only a single protein structure, the target protein, while typically large numbers of ligand molecules are involved. Therefore, manual intervention during protein preprocessing can be afforded while automatic tools are needed for the ligand molecules.

The most common starting point for the protein is the 3D structure in PDB [165] format containing coordinates for all heavy atoms. Additional information necessary for most docking programs are the location of the active site and the protonation states of polar atoms, and in some cases the location of the hydrogen atoms. In addition, ambiguities like alternate locations or conformations of amino acid side chains have to be resolved. Although automatic tools are available for most of these steps, manually driven approaches should be preferred since the exact definition of the protein structure has a large effect on the docking results. Often, many properties of the target protein are known or can be derived from already known ligands resulting in an improved model of the protein active site.

The ligand molecules are taken from a compound databases typically containing only structural formulae of the individual molecules. In order to use them in a docking calculation, 3D structures of the molecules have to be generated. For this step, very powerful tools like Concord [166] or Corina [68, 69] exist. In addition, atom hybridization and protonation as well as bond types have to be determined. Typically, this task is performed by a series of subsequently executed rule-based scripts. It should be noted that due to the complexity of this task, the resulting structures are not free of errors in the assignment of atom and bond types. The docking software should therefore rely as little as possible on this information.

#### 7.4.2

##### **Analyzing docking results**

The output of a docking program is a list of complex structures and an estimate for the binding affinity, the score, for each ligand. If the structures should be further examined, an energy minimization can be used for an initial refinement. Most docking programs are based on a discrete con-

formational model and therefore have to be tolerant against small overlaps between protein and ligand atoms. The minimization reduces these artefacts of combinatorial docking algorithms and moves the complex into the closest local minimum.

As already discussed in Section 7.2.5, estimating the binding affinity is a very complex task. The scores calculated by docking programs should therefore be seen only as a very crude measure. Depending on the number of ligand molecules, different techniques can be used for rescoring. If only few structures are of interest, time consuming methods like free energy perturbation can be applied to get a more detailed view on differences in binding energy between alternative ligands [167]. In most applications however, the number of ligands to be processed is too large for these techniques. One approach to overcome the limitations of single scoring functions is consensus scoring [161]. Depending on the screening performance achieved for a known set of ligands, a scoring function or a combination of functions can be chosen for calculating a final ranking for molecules with unknown binding affinity.

#### 7.4.3

#### **Choosing the right docking tool**

In Section 7.1.2, the most frequent docking scenarios occurring in drug design projects have been introduced. Since a variety of docking algorithms have been developed, the question arises which algorithm or software is most useful for the individual scenarios.

If only a small set of ligands should be docked, one can choose from the full spectrum of methods since then the computing time is only a minor issue. The most promising approaches are probably the hybrid methods combining a very efficient search technique in a first phase with a very exact scoring and optimization technique in the second phase. The first phase is of importance since it guarantees a broad sampling within the solution space. The second phase then tries to find the local minima and scores them as reliably as possible.

If the set of ligands increases, a tradeoff between accuracy and computing time has to be made. Rigid-body docking tools are most efficient, but they can only be used if the ligands of interest have only a small number of low-energy conformations. If this is not the case, the chance of missing hits due to missing conformations is too high. In the frequent case that the ligands are flexible (about more than five rotatable bonds), a tool handling conformational flexibility during docking must be used. Using a rigid-body tool as a prefilter is not helpful here. A good prefilter has a low false-negative rate resulting in a low number of hits removed. The false-positive rate is of less importance, since all molecules passing the filter will be analyzed in more detail in the subsequent screening step. A rigid-body docking tool however



has the opposite effect. The false-negative rate will be high since all molecules not tested in the binding conformation will be filtered out, while molecules tested in the binding conformation are kept resulting in a relatively low false-positive rate.

Besides computing time issues, known protein–ligand complexes with the target protein may also help in selecting the most promising docking algorithm. First of all, the degree of conformational flexibility in the protein while binding different ligands should be analyzed. If there are significant changes, a docking tool considering these effects is necessary. Docking tools handling protein flexibility are very rare and often not fast enough to be used in screening scenarios. Some of these tools are mentioned in Section 7.2. As an alternative, a docking algorithm can be run several times with different structures of the target protein as input. Of course, this method can only be applied if the protein conformations fall into a low number of separate classes.

Besides analyzing protein flexibility, the shape of the active site and the orientation of different ligands is of importance. Often, the active site has a deep cavity as it is the case in the thrombin example shown in the introduction. This is a favorable situation for incremental construction algorithms since they achieve the best results if they can place a first anchoring group deep into the active site and then continue growing to the outside. The situation becomes more difficult if the active site is very shallow or tunnel-shaped (e.g., in HIV-protease). If, in addition, the key interactions to the protein are spread over the whole ligand structure, the chance of missing the global minimum when using an incremental construction algorithm increases. Here, a more generic search technique like a genetic algorithm or a monte carlo algorithm could be favorable.

## 7.5

### Concluding remarks

Molecular docking tools are still far away from giving reliable results for all kinds of protein–ligand complexes. The complexity of the problem increases with the number of rotatable bonds in the ligand and the flexibility of the active site. In addition there is the lack of a reliable scoring function for the prediction of the free energy of binding.

The two main results of a docking calculation are the predicted complex structures for individual ligands as well as the ranking of binding molecules with respect to non-binding ones. Concerning structure prediction, docking software is able to generate a complex within a reasonable error bound (less than 2 Å RMSD) typically in about 70% of the test cases. In virtual screening, the current bottleneck is the scoring problem. Even if correct structures are generated, scoring functions often achieve only a weak correlation with

experimental binding affinities. Nevertheless, molecular docking is already a quite useful tool in the process of structure-based drug design. It allows to search in large virtual chemical spaces in a time- and cost-efficient way. Docking tools are successfully applied to prioritize compound databases for screening, designing targeted combinatorial libraries, and improving the binding affinities of initial lead structures.

Within the next years, we can expect a further increase in computing speed and probably an improvement in the quality of scoring functions. Looking at whole molecular libraries in a closed rather than an enumerated form has a great potential of speeding up docking calculations. Algorithms for docking combinatorial libraries as discussed above have shown this already. It can be assumed that once methods are available which overcome the sequential screening paradigm by new search techniques, a speedup of several orders of magnitude can be achieved.

Scoring functions are of central importance for virtual screening applications. They have to give reliable estimates of differences in binding affinity and must be efficiently computable in order to be used on thousands of structures. Reviewing the literature on this topic, improvements can be seen in the quality of scoring functions over the past years. Very recently, the idea of consensus scoring emerged combining the individual strengths of several scoring functions. While, in the past, most scoring functions were developed on small sets of complexes with known structure and binding affinity, the focus is shifting now towards more realistic screening scenarios as the basis for developing and calibrating scoring functions. Although no breakthrough is in sight, further improvement in the next years can be expected.

The docking problem in its full generality is likely to remain an open problem even in the long term. Too many subproblems are still unsolved, the most urgent ones being the accurate prediction of binding affinity and the handling of protein conformational flexibility.

## 7.6

### Software accessibility

Table 7.1 lists some available software packages for molecular docking discussed in this article. More information can be found on the corresponding Internet pages.

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Tab. 7.1  
Software for automated molecular docking

<i>Name</i>	<i>Refs.</i>	<i>WWW Homepage</i>
AutoDock	[96, 98, 97, 83]	<a href="http://www.scripps.edu/pub/olson-web/doc/autodock/index.html">http://www.scripps.edu/pub/olson-web/doc/autodock/index.html</a>
DOCK	[21, 26, 30, 25, 27, 28]	<a href="http://www.cmpfarm.ucsf.edu/kuntz/dock.html">http://www.cmpfarm.ucsf.edu/kuntz/dock.html</a>
DockVision	[107]	<a href="http://www.dockvision.com">http://www.dockvision.com</a>
FlexX	[63, 64, 65, 72, 71, 149]	<a href="http://cartan.gmd.de/FlexX">http://cartan.gmd.de/FlexX</a>
GOLD	[79, 85]	<a href="http://www.ccdc.cam.ac.uk/prods/gold.html">http://www.ccdc.cam.ac.uk/prods/gold.html</a>
ICM	[110, 168]	<a href="http://www.molsoft.com/icmpages/icmdock.htm">http://www.molsoft.com/icmpages/icmdock.htm</a>
LIGIN	[93]	<a href="http://swift.embl-heidelberg.de/ligin">http://swift.embl-heidelberg.de/ligin</a>
QXP	[108]	

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## 8 Modelling Protein–Protein and Protein–DNA Docking

*Michael J E Sternberg and Gidon Moont*

### 8.1 Introduction

#### 8.1.1 The need for protein–protein and protein–DNA docking

A description of many biological processes requires knowledge of the three-dimensional structure of macromolecular complexes. This structural information will provide insights into the specificity and can suggest lead compounds for the development of novel pharmaceutical agents. However, structural studies by crystallography and NMR often follow a reductionist approach so that the coordinates of the component molecules are available but the conformation of the complex is unknown. Indeed in the protein data bank [1], there is a large discrepancy between the number of different determined protein structures (c. 3000) and the number of protein–protein and protein–DNA complexes with proteins (c. 300). Thus computer algorithms are needed to predict the structure of macromolecular complexes starting from coordinates of the unbound components. This chapter describes computational strategies that can be employed to model the docking of two types of macromolecular complexes – protein–protein and protein–DNA.

The crystal structures of protein–protein and protein–DNA complexes have provided detailed descriptions of the interactions that lead to the specificity that is central to the biological activity of the system, such as the mechanism that leads to a disease. Currently we have details of a variety of protein–protein complexes including enzyme–inhibitor, antibody–antigen, hormones and their receptors and cell surface–cell surface proteins [2, 3, 4, 5]. The nature of the inter-protein recognition can be understood in terms of a general shape complementarity with specificity provided by particular spatial constraints from close packing and by hydrogen bonds and charge–charge ion pairings. From the structural information, one can start to alter one of the proteins to affect its recognition properties. For example, the de-

tails of a protein–protein complex can suggest that a particular loop is central to the interaction and this could lead to the design of lead compounds that could yield novel drugs. Indeed, there are several lead compounds that have been designed based on the structure of a protein receptor interacting with a small molecule ligand (e.g., see review by Colman [6]). With the increasing number of determined protein structures, the determination of protein–protein complexes by both experiment and modelling should also lead to suggestions of new pharmaceutical agents.

### 8.1.2

#### **Overview of the computational approach**

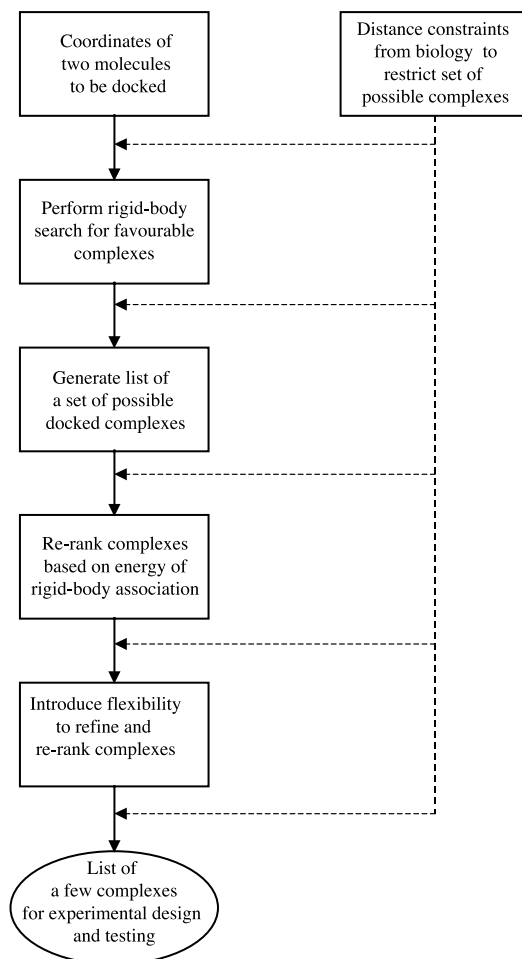
This chapter will describe computational strategies to predict the structure of protein–protein and protein–DNA complexes. The consensus approach to protein–protein docking is described in Figure 8.1. The precise order of implementing the steps can differ but essentially the key features are now overviewed.

- i. One starts with three-dimensional structures of the two unbound components and considers systems that are expected to have limited conformational change on macromolecular association.
- ii. The rigid-body approximation is then used which is that one can simulate the docking starting with the unbound components, given the limited conformational change.
- iii. A search is performed over all possible associations. If there is no biological information about which parts of the molecules interact then all possible associations must be considered. If however there are some constraints, these can be used either to limit the initial search or as a post search filter. The search will sample the space of possible associations and consequently there will be a lower limit on the difference in conformations between two docked complexes that determines the resolution of the search procedure.
- iv. A function is developed to score the quality of the docked complex often using simplified terms to evaluate shape and electrostatic complementarity. There are two reasons for a simple function at this stage. First, as a large number of different complexes are generated in global scan, the energy term must be computationally fast to evaluate. Second, one requires a soft potential that is not unduly sensitive to the conformational differences between the docked structures formed from unbound components and the true complex with the bound components.
- v. Ideally, the docking algorithm would thereby identify a single complex that is a close approximation to the true docked complex based on this complex having the best score (i.e., the lowest energy complex). In



**Fig. 8.1**

Overview of a typical docking procedure. This diagram summarizes the consensus approach based on rigid body docking followed by re-ranking and refinement. Approaches vary according to whether the steps are combined or performed independently in a particular program. Many strategies only perform some stages, particularly the initial docking and scoring or just subsequent re-ranking and refinement. Biological distance constraints can be included at different stages in the procedure.



practice, the current status of the algorithms is that they generate a limited list of possible complexes ranked on the score and the objective is that one member of the list should be close to the true complex.

- vi. At this stage, a re-ranking of the resultant complexes can be undertaken possibly using computationally more intensive calculations.
- vii. Conformational flexibility is generally introduced into the algorithm when there is only a limited number of complexes to consider. Perturbations to the structure of the docked complex are made and the energies of the resultant conformation is evaluated. The aims are both to improve the structural quality of the predicted complex and to improve the power of identifying the true solution from the set of false structures that have been generated.

A summary of the current status of protein-protein docking is that when this strategy has been applied to around ten test systems, one can generate a close model (2.5 Å rmsd for the C $\alpha$  atoms) to the true complex at a rank of ten or less for many of the systems studied.

### 8.1.3

#### **Scope of this chapter**

Given the reliance of most methods on starting by a rigid-body docking, the next section in this chapter will describe the extent of conformational change on protein–protein association. It will be shown that for many systems, the change is sufficiently limited to suggest that the rigid-body approach is viable.

Next, a computational strategy developed in our Laboratory for protein–protein will be described. The results will show that protein–protein docking can often yield a limited list of possible complexes one of which is close to the native. The chapter will then describe the application of this procedure to protein–DNA docking. Protein–DNA docking is harder than protein–protein docking given the more uniform shape of double-stranded DNA and the more extensive conformational change that tends to occur on association. The results will show that despite these problems, modelling protein–DNA complexes is viable. The description of our approach will serve to illustrate many aspects of the general strategy outlined above (Section 8.1.2) and thereby report the current status of protein docking.

We will then survey several alternative approaches for protein–protein docking. A true test of any predictive method is its success in blind trials since this prevents bias for over optimisation during the development of the approach. There have been two such blind trials and the results will be described. In addition to this chapter, the reader is referred to other reviews of the field [7–9].

## 8.2

### **Structural studies of protein complexes**

The first step in the development of a protein docking algorithm is to examine the known complexes to establish the principles of molecular recognition. Following earlier work [10], there are reviews that examined the interactions between hetero-protein complexes [4, 7]. These analyses have focussed on the static structure of the complexes. A major problem in protein docking is to cope with the conformational flexibility that occurs on complex formation. Accordingly, we have analyzed the conformational changes on complex formation for 39 pairs of structures of proteins in

complexes and in their unbound states [2]. The data set mainly consisted of enzyme–inhibitor and antibody–antigen complexes but also included other systems such as human growth hormone and its receptor.

The conformational differences were evaluated in terms of root mean square deviations (rmsd) of  $C\alpha$  positions and of side-chain positions. Residues were identified as exposed when their total relative side-chain accessible area (main-chain for Gly) was greater than 15%. Interface residues were defined as having at least one atom within 4 Å of the other component in the complex. To assess the significance of the differences between bound and unbound structures, it is necessary to identify the differences in coordinates that can occur simply from the crystallographic determination of the structure. Accordingly, 12 pairs of independently solved crystal structures of identical proteins were analysed. 95% of this data set (11 out of 12) had an rmsd for exposed  $C\alpha$  atoms of  $<0.6$  Å and for exposed side-chain atoms of  $<1.7$  Å. These values were taken as the control cut-offs. Four measures were adopted for overall conformational change – rmsd of  $C\alpha$  and of side-chain atoms for just the interface atoms and for exposed non-interface atoms. 19 out of the 39 proteins that were involved in complex formation do not have any of their four measures for conformational change above the cut off values. Many of the other proteins showed changes just above the control cut offs. Thus for many systems, there is limited overall conformational change on protein–protein association which suggests that the rigid-body approach should be widely applicable.

In addition to overall conformational changes, one or a few individual residues might show particularly large conformational changes and this could markedly affect the viability of the rigid-body approach. The control cut-offs for the movement of an individual residue are 3.0 Å for a  $C\alpha$  and 5.6 Å for side-chain displacements. Examination of the complexes showed that all large movements of exposed residues that were not in the interface can be explained by either their close proximity to the interface or by structural disorder. For a few of the systems, there are movements of individual or a set of residues in the interface that are above the control cut-offs. These shifts are intimately involved in the complex formation. Thus there are several complexes with substantial conformational change for one or a few residues whilst still there is overall a limited structural perturbation.

The general conclusion from this analysis is that protein–protein interactions are described by the induced fit model. However for many systems, the extent of conformational change is limited and the lock-and-key model is a valid first approximation. Accordingly, for many systems it is appropriate to develop a protein–protein docking algorithms that starts with the docking of the components as rigid bodies and then as a refinement considers limited conformational changes. The major caveat is that the systems analyzed were dominated by enzyme–inhibitor and antibody–antigen complexes; consequently other biological systems may exhibit a greater degree of conformational change on complex formation.

### 8.3

#### Methodology of a protein-protein docking strategy

In this section we describe a computational strategy developed in our Laboratory for docking. Our suite of programs is available from <http://www.bmm.icnet.uk> and models the docking two medium-sized proteins (50–500 residues). In outline (see Figure 8.2) the method is:

- i. a rigid-body search for docked complexes that are favourable in terms of shape complementarity and electrostatics,
- ii. the application of an empirical scoring function to re-rank the docked structures generated by (i),
- iii. the use of biological distance constraints, particularly details of the binding sites in one or both proteins, to screen the putative complexes from (ii),
- iv. the final refinement and further re-ranking of the rigid body structure by a consideration of side-chain conformational change.

#### 8.3.1

##### Rigid body docking by Fourier correlation theory

The first step in most approaches including ours is the rigid-body docking of the two molecules to generate a set of complexes. There are two major requirements:

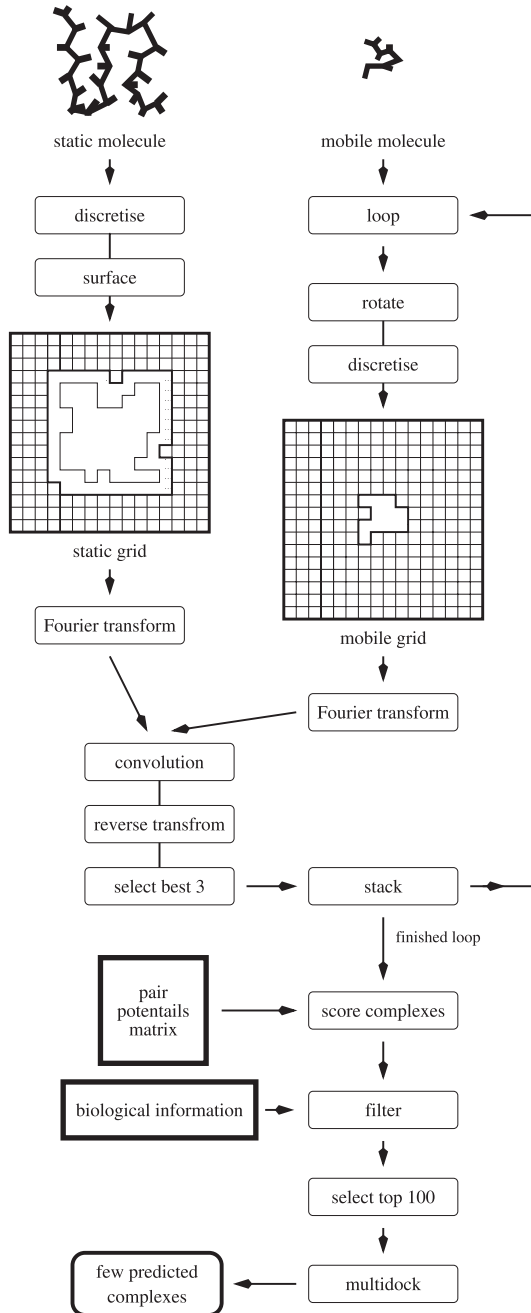
- i. The method must be able within a realistic computation time to generate and evaluate trial complexes that cover all relative orientations at sufficient resolution to yield at least one solution that is close to the true structure.
- ii. The approach must use scoring functions that are sufficiently soft to cope with a limited conformational change on association yet be able to identify a solution that is close to the true complex.

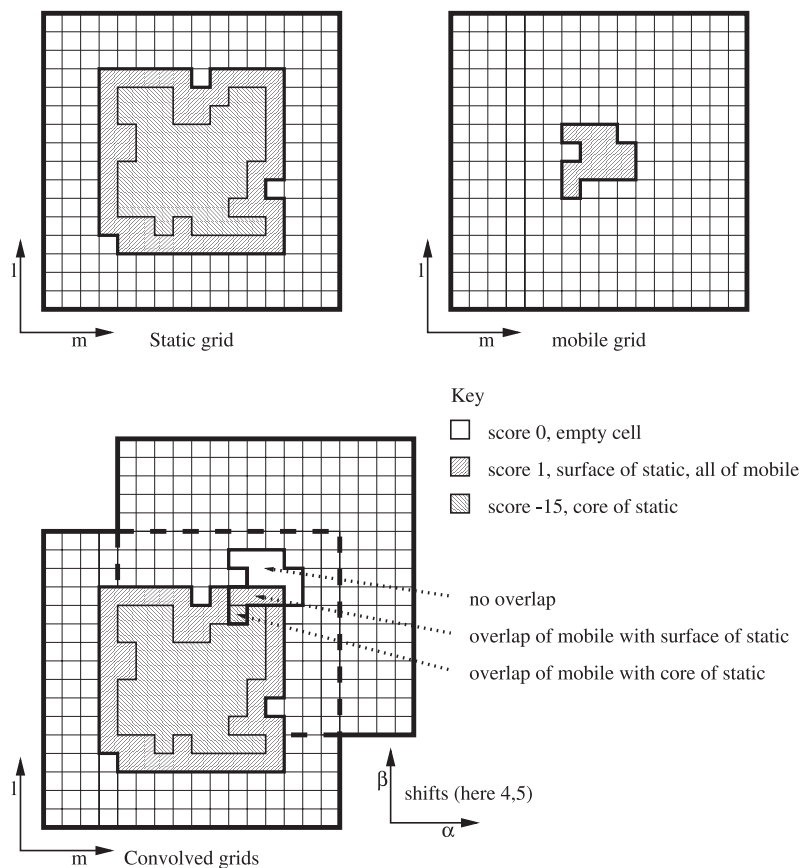
The Fourier correlation approach introduced by Katchalski-Katzir et al. [11] was developed to meet these requirements. We have augmented their method to include electrostatic effects that often are required to obtain results starting from unbound complexes. Our initial work (FTDOCK1) was described in Gabb et al. [12], but the approach has been re-implemented with minor changes and this algorithm (FTDOCK2) is now described (see Figure 8.2).

##### 8.3.1.1 The generation of the grid representation

The Katchalski-Katzir method is a grid based approach to evaluate the shape complementarity between two proteins (see Figure 8.3). Consider two mol-

**Fig. 8.2**  
Schematic of our strategy  
for generating and  
screening docked protein  
complexes.





**Fig. 8.3**  
Scoring shape complementarity using a grid-based scoring approach. The Figure illustrates the approach used in FTDOCK. The top two grids show the individual

discretisation of the static and mobile molecules. The lower grid shows the convolved grids with the calculation of shape complementarity.

ecules  $A$  and  $B$ . Let  $A$  be the larger of the two and taken to be the static molecule.  $B$  will be the smaller mobile component to be docked against  $A$ .

Each molecule is placed onto a 3-dimensional grid and the algorithm requires that both grids are the same dimension ( $N \times N \times N$ ). The size of grid cell must be sufficiently small to model the atomic structure of the molecules. However the computational time increases as the cell size decreases. We currently use a value of  $0.7 \text{ \AA}$  that is sufficiently small that generally only one atom (excluding hydrogen) is in an individual cell. To evaluate the required  $N$ , for each protein the maximum distance from any atom to the geometric centroid is calculated ( $DMAX_A$  and  $DMAX_B$ ). The value  $XN$  is calculated from:

$$XN = (2 \times (DMAX_A + DMAX_B) + 1.0)/0.7$$

$N$  is set to the nearest integer to  $XN$ . For ease of computation in the Fourier method,  $N$  should be even and so if  $N$  is odd, the next highest even integer is used. The result of this calculation of  $N$  ensures that the mobile molecule can always be placed to touch the static molecule and there will remain grid cells at the boundary unoccupied by the protein.

The discretization is done with each molecule at the centre of each grid. The empty space not filled by the molecule is necessary for the algorithm. To perform the discretization, each grid cell within which an atomic position is found is turned 'on'. Grid cells whose centre is within 1.8 Å of any atomic position are also turned 'on'. This value of 1.8 Å is chosen to approximate an effective van der Waals radius for an atom combined with any hydrogen atoms that are bound to it. Thus the surface of the resulting grids will represent the atomic surface of the molecules.

Next, the larger static molecule ( $A$ ) is assigned a surface thickness below its atomic surface in order to be able to calculate the quality of the fit. For algorithmic speed, this surfacing procedure is done to the static molecule, and accordingly it needs to be done only once by the program. The depth of this surface is 1.4 Å, so that the surface is never more than two cells thick. After the surface has thereby been assigned, the core cells still cover all the actual atomic positions, but the van der Waals radius has been effectively removed, so allowing closer interactions between the two components of the complex. This representation of the surface provides a model for complex formation in which there is some side-chain rearrangement followed by side-chain interdigitation.

### 8.3.1.2 Evaluation of shape complementarity

The grid values  $a_{l,m,n}$  for the static molecule at node  $l, m, n$  are given by

$$a_{l,m,n} = \begin{cases} 1 & \text{for grid points on the surface of the molecule} \\ \rho & \text{for the core of the molecule} \\ 0 & \text{for outside the molecule} \end{cases}$$

where  $\rho$  is negative (we use  $-15$ , see below).

For the second molecule ( $B$ ) the grid values

$$b_{l,m,n} = \begin{cases} 1 & \text{for the molecule} \\ 0 & \text{for outside the molecule.} \end{cases}$$

The two grids can then be superimposed and the movable grid ( $B$ ) translated by shifts  $\alpha, \beta, \gamma$  (see Figure 8.3). The value of

$$a_{l,m,n} \cdot b_{l-\alpha, m-\beta, n-\gamma}$$

gives the extent of shape complementarity for grid point  $l, m, n$  of the grid of molecule  $A$ . A value of 1 for the product indicates that the cell from molecule  $B$  is superimposed on the surface of  $A$  which is favorable shape complementarity. A value of  $\rho$  indicates a steric clash with the cell from  $B$  superimposed on the core of  $A$ . The value of  $\rho$  is chosen so as to penalise but not totally prevent steric clashes. A value of zero means that there is no overlap between the two molecules. Thus the total shape complementarity for the two superimposed grids  $c_{\alpha,\beta,\gamma}$  is evaluated from

$$c_{\alpha,\beta,\gamma} = \sum_{l=1}^N \sum_{m=1}^N \sum_{n=1}^N a_{l,m,n} \cdot b_{l-\alpha,m-\beta,n-\gamma} \quad (1)$$

### 8.3.1.3 Use of discrete Fourier transforms

The value of  $c$  is a convolution and its calculation requires approximately  $N^3$  multiplications and a summation for every  $N^3$   $\alpha, \beta, \gamma$  shifts. Katchalski-Katzir et al. introduced the use of discrete Fourier transforms to speed up the process of calculating  $c$ . First, calculate the discrete Fourier transforms of the discrete functions for the cell values  $a$  and  $b$  and let these transforms be denoted as  $\text{DFT}(a)$  and  $\text{DFT}(b)$ . Then, calculate the complex conjugate of  $\text{DFT}(a)$  which is denoted as  $\text{DFT}^*(a)$ . Then by Fourier theory, the discrete Fourier transform of  $c$ ,  $\text{DFT}(c)$ , is given by

$$\text{DFT}(c) = \text{DFT}^*(a) \cdot \text{DFT}(b)$$

Hence,  $c = \text{IFT}(\text{DFT}^*(a) \cdot \text{DFT}(b))$  where IFT denotes the inverse Fourier transform.

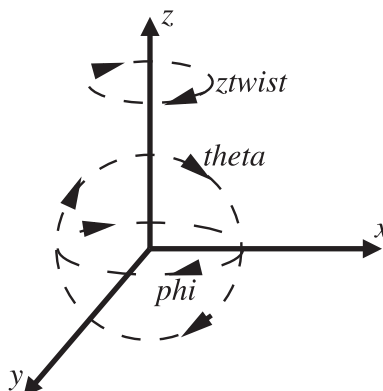
The discrete Fourier transform, if performed efficiently, takes  $N^3 \log N$ . This must be performed three times, but this does not increase the order of magnitude of the calculation. The pointwise multiplication takes  $N^3$ . Thus the total runtime takes  $N^3 \log N + N^3$ , which is of the same order as  $N^3 \log N$ . In contrast without the transform in real space the calculation would be of order  $N^6$ . As  $N$  is of the order of 100, the discrete Fourier transform decreases the time required by more than a factor of 10 000.

### 8.3.1.4 The global search

The movable molecule is rotated to sample all possible rotations in as fair a way as possible. In order to describe the orientation of a three-dimensional object in a three-dimensional space, three rotational angles are required. Therefore, in order to orient a molecule it is necessary to perform three rotations (see Figure 8.4). For FTDOCK2 the rotations were in order; around the z-axis (z twist), around the y axis (theta), and around the z axis again (phi). Since only integer values are used to describe the rotations, this limits



**Fig. 8.4**  
A schematic diagram of the rotational search.



the angle step used to being an integer factor of 180. Once the angle step had been chosen for the sampling rate, in order to create an even sampling, the angle step used for phi was calculated dependent on the value of theta. This results in a fair sampling of rotational space. For angle step of  $20^\circ$ , this method gives 1980 rotations. For an angle step of  $15^\circ$  it gives 4656, and for  $12^\circ$  it gives 9240.

#### 8.3.1.5 Electrostatic effects

Both shape complementarity and electrostatic effects are important in the recognition process in protein complex formation. Accordingly a treatment of electrostatics was introduced into the Fourier correlation approach. The charge–charge interaction is evaluated from point charges of the mobile molecule B interacting with the potential from static molecule A. This choice results in having to perform the potential calculation only once (for the static molecule) whilst the charge calculation (see below) is performed for every rotation of the mobile molecule. In the above treatment of rigid body docking based on shape complementarity, it is possible to place two charges closer together than would be allowed by van der Waals packing. Since the potential energy of two interacting charges depends inversely on their separation, this would result in an artificially highly favorable or very unfavourable interaction. These artificial terms are prevented by the method now described to calculate the potential.

Charges (see Gabb et al. [12]) are assigned to the atoms of molecule A and the electrostatic potential evaluated from

$$\phi_{l,m,n} = \sum_j \frac{q_j}{\varepsilon(r_{ij})r_{ij}}$$

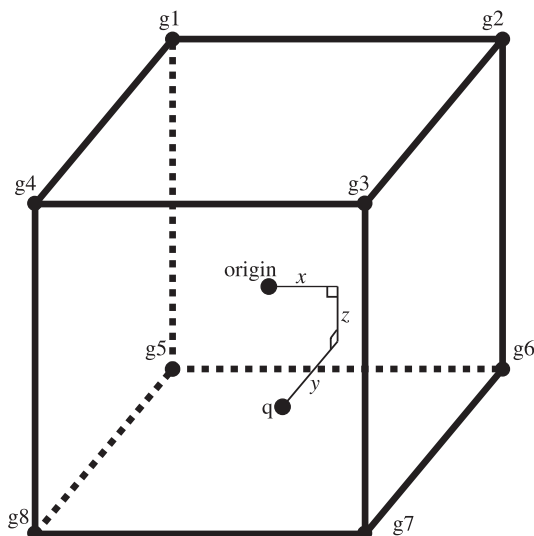
where  $\phi_{l,m,n}$  is the potential at node  $l, m, n$  (position  $i$ ),  $q_j$  is the charge on

atom  $j$ ,  $r_{i,j}$  is the distance between  $i$  and  $j$  (with a minimum value of  $2 \text{ \AA}$  to avoid artificially large values of the potential) and  $\varepsilon(r_{ij})$  is a distance dependent dielectric function. The sigmoidal function of Hingerty et al. [72] is used:

$$\varepsilon(r_{ij}) = \begin{cases} 4 & \text{for } r_{ij} \leq 6 \text{ \AA} \\ 38r_{ij} - 224 & \text{for } 6 \text{ \AA} < r_{ij} < 8 \text{ \AA} \\ 80 & \text{for } r_{ij} \geq 8 \text{ \AA}. \end{cases}$$

This function was introduced for modelling the effective dielectric between atoms in proteins. The rationale for this function is that at close separation ( $r_{ij} \leq 6 \text{ \AA}$ ) when there is no intervening water molecules, the effective dielectric is that of protein atoms and a value of 4 is appropriate. For separations of  $8 \text{ \AA}$  or more then the dielectric is dominated by the screening effect of the intervening water and the value for bulk water (80) is used. Between these two separations a linear interpolation is used. In FTDOCK, we do not use precise atomic positions, but still need to model the complex dielectric behavior of proteins in solvent and this function was used and found to perform well.

The potential  $\phi_{l,m,n}$  is only assigned to nodes outside and on the surface region of molecule A. Inside the core of molecule A,  $\phi_{l,m,n}$  is zero. For the mobile molecule B, the charges on the charged atoms are distributed amongst the closest 8 grid cells, see Figure 8.5. Given the atomic position of the charged atom normalised onto the centre of the 8 neighbouring cells as  $(x, y, z)$ , and the normalised centre of one of those closest 8 cells as



**Fig. 8.5**  
Schematic diagram of the approach to transfer an electronic charge to its nearest eight grid nodes.

( $X, Y, Z$ ), then the charge given to that cell is

$$(q_{l,m,n}/8) * ((x + X)/X) * ((y + Y)/Y) * ((z + Z)/Z)$$

The electrostatic interaction  $e_{\alpha,\beta,\gamma}$  for a shift of  $\alpha, \beta, \gamma$  is calculated from

$$e_{\alpha,\beta,\gamma} = \sum_{l=1}^N \sum_{m=1}^N \sum_{n=1}^N \phi_{l,m,n} \cdot q_{l-\alpha,m-\beta,n-\gamma} \quad (2)$$

Eq. (2) is analogous to Eq. (1) for shape complementarity and accordingly the Fourier correlation approach can be also used to speed up the calculation.

#### 8.3.1.6 Generation of putative complexes

In trials with FTDOCK, it was found that the electrostatic term worked best as a binary filter. Complexes with unfavourable electrostatics are discarded and the remaining docked structures ranked by shape complementarity. For a given orientation of the movable molecule, the shape complementarity correlation function  $c$  is examined and the three most favourable solutions stored. After all orientations are sampled, the top set (typically 10 000) of complexes are kept for subsequent examination.

#### 8.3.2

##### Use of residue pair potentials to re-rank docked complexes

After the global search for shape complementarity and favourable electrostatic interaction, a list of several putative complexes is generated. A model close to native (see below for quantitative measures) is rarely at the top of the list of complexes ranked by shape complementarity. It is necessary, therefore, to develop further scoring functions to identify good solutions from the list of alternatives. The strategy we have developed is to use a residue-residue scoring scheme to evaluate the docked complexes. Such an approach has an energy landscape that is smoother than an approach based on all (heavy) atoms but incorporates information that cannot be included in the grid-based first step. The terms to quantify the residue-residue interactions are derived empirically from known protein structures and these terms are often known as empirical pair potentials.

The theory of empirical pair potentials is that since they are derived from observations they will incorporate the dominant thermodynamic effects without explicitly having to partition and quantify the effects. Empirical pair potentials have been widely used in protein structure prediction, particularly in fold recognition following their introduction by Sippl [13] and their use

Tab. 8.1

Complexes used to generate the empirical scoring function. The protein data bank codes [1] of the complex are given.

---

1a14	1a1y	1a22	1a2x	1a2y	1a4y	1agr	1ahw	1aip	1ak4	1akj	1am4
1avw	1avz	1ay7	1azs	1b6c	1bd2	1bdj	1bgx	1bh9	1bi8	1bj1	1bkd
1blx	1bqg	1bvq	1d0g	1dan	1dfj	1dvh	1eai	1eay	1efn	1efu	1fdl
1fin	1fle	1gc1	1got	1gua	1hia	1hlu	1igc	1itb	1jrh	1kb5	1ld9
1lfd	1lpb	1mct	1mda	1mel	1mlc	1nfd	1nfi	1nmc	1noc	1oak	1osp
1qbk	1raa	1rrp	1seb	1sgp	1slu	1spb	1tab	1taw	1tx4	1uea	1ugh
1viw	1vpp	1wej	1yys	2jel	2pcc	2seb	2trc	3sic	4cpa	4htc	

---

by Jones et al. [14]. For a review, see Vajda et al. [15]. In addition, empirical functions have been used to evaluate protein-low molecular weight ligand complexes [16, 17, 18, 19]. Here we present the derivation and application of pair potentials specifically designed to re-rank docked protein-protein complexes.

The natural source to derive empirical potentials for protein docking is the complexes themselves. However at the time we first developed the potentials there was insufficient data to obtain the potentials and the statistics were taken from observations of residue-residue pairing within protein structures [20]. Recently, with the increase in the number of determined complexes, we have been able to generate the pair potential from these data and this approach is now described. By use of the SCOP (Structural Classification of Proteins) database [21] version 1.48, a set of 83 non-homologous complexes was obtained (see Table 8.1). In this dataset homology between two complexes (A-B) and (C-D) is defined as when both A is homologous to C and B is homologous to D, taking homology as being in the same SCOP superfamily.

Two residues were taken as pairing across the interface if any pair of atoms between them was closer than a cut off  $d_{cut}$ . A value of  $d_{cut}$  of 4.5 Å is used to represent two typical van der Waals radii (1.5 Å each) plus an additional 1 Å. Let the frequencies of observed pairings between residues of type  $a$  and  $b$  be  $F_{ab}$ .

These observed frequencies are compared to those for a random state. The model for the random state is based on random associations of exposed residues on one protein with exposed residues on another protein. To evaluate this one needs the frequencies of exposed residues in the database of proteins used to generate the observed pairings. From the coordinates of a protein, the program DSSP [22] estimates the extent to which each protein residue is accessible to solvent contact. This is represented by an accessibility score and a value of 10 units is equivalent to one water molecule in contact with the residue. We find that a value of an accessible score  $> 5$

units identifies residues that are not primarily buried within the protein core.

The expected number of random pairings of residue types  $a$  and  $b$ ,  $E_{ab}$ , is based on frequencies of the exposed residues. Let  $T$  be the total number of all pairings observed in calculating the observed residue-residue pairings,

$$T = \sum_{a=1}^{20} \sum_{b=1}^{20} F_{ab}$$

Let  $n_a$  and  $n_b$  be the total occurrences of exposed residues of types  $a$  and  $b$ . The fraction of residues that are of type  $a$  and type  $b$  are respectively

$$\frac{n_a}{\sum_{a=1}^{20} n_a}$$

and

$$\frac{n_b}{\sum_{b=1}^{20} n_b}$$

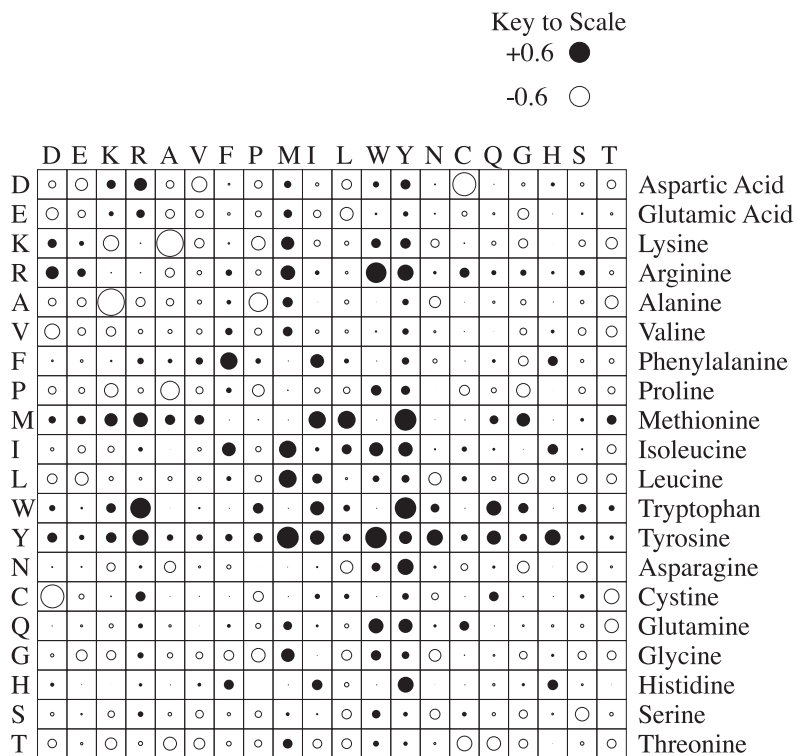
Thus the expected frequency for the  $a$ - $b$  pairing is given by

$$E_{ab} = T \cdot \frac{n_a}{\sum_{a=1}^{20} n_a} \cdot \frac{n_b}{\sum_{b=1}^{20} n_b}$$

A log-odds score for a pairing of residue types  $a$  and  $b$  is derived:

$$S_{ab} = \log_{10} \left( \frac{F_{ab}}{E_{ab}} \right)$$

These scores are shown schematically in Figure 8.6. As expected, pairings between hydrophobic residues and between residues with opposite charges are favourable. The total score for a complex is obtained by summing the  $S_{ab}$  values for all residue pairings between the two molecules with the distance less than  $d_{cut}$ . High scoring complexes are evaluated by this function to be more favourable. In many other applications of empirical pair potentials the log odds scores are converted to a potential of mean force by the application of Boltzmann's principle. However certain groups have questioned the validity of this approach [23, 24] and accordingly we simply treat the scores as a statistical measure of relative likelihood. This screening procedure is implemented in the program RPSCORE.



**Fig. 8.6**  
Empirical residue-residue docking potentials. The empirically derived residue-residue contact scores are

illustrated. The largest solid sphere represents a favourable score of +0.6 whilst the largest open sphere the most unfavourable score of -0.6.

### 8.3.3

#### Use of distance constraints

The location of the binding site can yield distance constraints to filter possible solutions and this is implemented in our procedure. An intermolecular residue-residue interaction is defined if any pair of atoms is closer than a 4.5 Å cut off. This cut off considers the probable error in conformation of any predicted complex. One feature of the Fourier correlation method implemented in FTDOCK, is that biological constraints on one molecule cannot be used to reduced the initial search. For any orientation of the movable molecule, it is placed at every translational position with respect to the static molecule. However, if constraints were available for both molecules, the range of orientations sampled by the movable molecule could be restricted.

### 8.3.4

#### Refinement and additional screening of complexes

The above strategy explores and ranks rigid-body associations of the molecules with the pair potentials operating at the level of residue-residue interaction. The next step is to allow for conformational change for side-chains and to consider the interactions between the proteins in the complex at the atomic level. We [25] have developed a procedure (MULTIDOCK) that models both side-chain conformational changes in a mean field approach together with limited rigid-body shifts between the component molecules. The energy of interaction is evaluated from a molecular mechanics function.

##### 8.3.4.1 Potential energy function

The proteins are represented at the atomic level by multiple copies of side-chains on a fixed peptide backbone modelled according to a rotamer library [26] that gives the commonly occurring side chain conformations. The model to be described considers protein-protein interactions. The terms for van der Waals interactions are taken from the AMBER force field [27] and atomic charges from the PARSE parameters [28]. A cut off of 10 Å is used in the calculation of the non-bonded interactions (van der Waals and electrostatic) between atoms. The dielectric screening between charges is represented by the widely-used distance dependent dielectric (dielectric  $\epsilon = \text{distance of separation in \AA}$ ). The effect of this dielectric model is that for close separations the value is low and represent the dielectric due to protein atoms. Thus at a separation of 4 Å the dielectric is 4. For larger separation the dielectric increases representing the greater electrostatic screening due to water molecules (not included explicitly).

This approach uses a rotamer library so side chains adopt a limited number of conformations rather than being able to sample every value for bond rotation. In addition, the starting model generated by FTDOCK (or another rigid body docking approach) may be a few Ångstrom rmsd ( $C\alpha$  atoms) from the true structure. The consequence is that there can be an unrealistic close approach of atoms that could distort the modelling due to high repulsive van der Waals interactions or electrostatic effects of large magnitude. To reduce this effect, van der Waals interactions are truncated to a maximum value of 2.5 kcal/mol. Similarly, an electrostatic interaction scheme was introduced in which a minimum allowed distance separation between two interacting charges  $q_i$  and  $q_j$  is set so that atom pairs that come closer than allowed are re-scaled to realistic values which is no greater than the approximate sum of their van der Waals radii. The minimum allowed distances for two charges are 3 Å for two heavy atoms, 2 Å for one heavy atom with hydrogen and 1 Å for two hydrogen atoms. Note that the treatment of the screening effect of water for larger separations and the prevention of anomalous electrostatic effects are also included in FTDOCK.

### 8.3.4.2 Refinement procedure

The object of the refinement procedure is to move from a completely rigid-body docking scheme to include both flexibility of the side-chains together with a limited re-orientation of the two interacting molecules. Thus the refinement procedure is an iterative two step approach repeated until convergence involving:

- i. optimisation of the protein side chain conformations by a self-consistent mean field approach [29, 30]. The resultant protein conformations then undergo
- ii. rigid-body energy minimisation to relax the protein interface.

The mean-field approach is now described. The side chain degrees of freedom are defined by a conformational matrix,  $CM$ , where each rotamer,  $k$ , has a probability of  $CM(i, k)$ , where the sum of the probabilities for a given residue,  $i$ , must be equal to 1. The potential of mean force,  $E(i, k)$ , on the  $k$ -th rotamer of residue,  $i$ , is given by;

$$E(i, k) = V(\chi_{ik}) + V(\chi_{ik}, \chi_{mc}) + \sum_{\substack{j=1 \\ j \neq i}}^N \sum_{l=1}^{K_j} CM(j, l) V(\chi_{ik}, \chi_{jl})$$

where  $V$  is the potential energy,  $\chi_{ik}$  are the coordinates of atoms in rotamer  $k$  of residue  $i$  and  $\chi_{mc}$  are the coordinates of atoms in the protein main chain.  $N$  is the number of residues in the protein and  $K_j$  is the number of rotamers for residue  $j$ . The first term models the internal energy of the rotamer whilst the second represents the interaction energy between the rotamer and all the main chain atoms. These two values are constant for a given rotamer on a given main chain. The third term models the interaction energy between the rotamer and all the rotamers of other residues weighted by their respective probabilities.

Given the effective potentials acting on all  $K_i$  possible rotamers of residue,  $i$ , the Boltzmann principle can be used to calculate the probability of a particular rotamer:

$$CM(i, k) = \frac{e^{-E(i, k)/RT}}{\sum_{k=1}^{K_i} e^{-E(i, k)/RT}}$$

where  $R$  is the Boltzmann constant and  $T$  the temperature. The values of  $CM(i, k)$  are substituted back into the equation describing  $E(i, k)$  and its new value recalculated. This process is repeated until values of  $CM(i, k)$  converge. The predicted structure corresponds to the rotamer of each residue with the highest probability. Trials showed that the procedure



converged to the same side-chain rotamers using a number of different schemes for initiating the starting probabilities in the *CM* matrix.

After a complete cycle of mean field optimization of side-chain conformation, a rigid-body minimization is performed on the resultant coordinates of the interacting protein molecules. Only interface residues whose  $C\beta$  atoms ( $C\alpha$  for Gly) are within 15 Å of the  $C\beta$  atom of the other molecule are included in the minimization. The larger molecule is kept stationary while the three rotational and three translational degrees of freedom of the smaller molecule are moved according to the path determined by the derivatives to minimize the intermolecular interaction energy. The steepest descents approach for minimization is used.

### 8.3.5

#### Implementation of the docking suite

The software for the strategy is available from our Web site under the home page <http://www.bmm.icnet.uk> and is free to academic users. The original distributed version of FTDOCK (FTDOCK1) had two implementations of the Fast Fourier Transforms (FFT) to search the translational binding space of two rigid molecules. One used the FFT routines from Numerical Recipes Software [31] and can be implemented on a variety of hardware platforms. The other implementation used the more efficient Silicon Graphics library functions.

Our recent implementation (FTDOCK2), is written in ANSI C and uses the “The Fastest Fourier Transform in the West,” developed by M. Frigo and S. G. Johnson (see <http://www.fftw.org/>). As this is a fast public domain Fourier Transform Routine, we can distribute the entire source code for FTDOCK (as FTDOCK2). FTDOCK2 has not been parallelised which facilitates its implementation on a wide variety of platforms, including Linux. The program requires at least 128 Mbytes of RAM to prevent excessive paging. On a single R10000 Silicon Graphics processor a typical docking took between 9 and 36 hours, depending on the size of the molecules. In addition, PERL 5.003 or later is required for pre-processing the protein data bank coordinates.

To run FTDOCK2, the user first runs a supplied program to pre-process the coordinate files. Then the user need simply to identify the coordinate files for the static and the mobile molecules although other parameters can be adjusted.

The RPSCORE program that re-ranks the generated solution using the empirical pairing score is written in ANSI C and the source code is available. The program will automatically use our chosen matrix but the user can input a matrix of their choice.

The FILTER program implements the distance constraints. The program takes a list of inter-molecular constraints which can either be residue to

residue, chain to residue, or chain to chain. Only one from a given list of constraints, needs to be satisfied for the program to accept a complex as passing the filter.

The code for MULTIDOCK is available as an executable for a Silicon Graphics computer. In our development of MULTIDOCK we included a version that explicitly introduced solvent molecules. The typical run times for MULTIDOCK to examine a single complex such as that of a serine protease – inhibitor system are 10 minutes with solvent and 3 minutes without solvent. For a larger system, an antibody docking to hen lysozyme, the corresponding run times are 40 minutes with solvent and 7 minutes without solvent. In an application of the procedure, it is essential to include a large number of trial solutions to ensure not missing the predicted complex that is close to the native. This can mean that up to 200 solutions may need to be screened (see Results section below). Thus for larger systems a solvent based screening could take several days. However, trials showed that inclusion of solvent does not turn a native-like solution with a poor *in vacuo* energy into a high ranking solution. At present we therefore distribute MULTIDOCK for use without explicit solvent molecules.

The total run time to examine a single protein-protein complex is on one processor about 2 days. Many users would be interested in how a particular pair of molecules will dock and then this run time is acceptable. In addition, one can evaluate the strategy for a number of test systems (typically 10–20) over about one month. However this length of run time prevents one exploring the use of the procedure to study problems such as all pairwise associations within a set of (say) 100 proteins.

#### 8.4

##### Results from the protein-protein docking strategy

We are currently benchmarking the protein docking procedure on a set of 15 complexes with coordinates available for both components in their unbound state. The results will be placed on our Web page ([www.bmm.icnet.uk](http://www.bmm.icnet.uk)). Preliminary inspection suggests the results of the new benchmark will be similar to our original benchmark. Here we report the results of the original benchmark. The approach used to obtain these results was (i) FTDOCK (as reported in Gabb et al. [12]), (ii) filtering on biological distance constraints [12], (iii) RPScore [20] and (iv) MULTIDOCK [25].

The benchmark consists of six enzyme/inhibitor and four antibody/antigen complexes. With the exception of the coordinates of two antibodies (HyHEL5 and HyHEL10), all the coordinates were from the unbound state. Table 8.2 presents the results. In each study, 4000 solutions were generated by FTDOCK1. The total number remaining after the application of the biological filter is shown in column 2 of Table 8.2 and forms set 1. All the en-

Tab. 8.2

Results of protein–protein docking.  $\alpha$ CHYN– $\alpha$ -chymotrypsinogen,  $\alpha$ -CHY– $\alpha$ -chymotrypsin, HPTI–human pancreatic trypsin inhibitor, BPTI–bovine pancreatic trypsin inhibitor, CHYI–chymotrypsin inhibitor, Subtilisin I–subtilisin inhibitor D1.3, D44.1, HyHEL5 and HyHEL10 are monoclonal antibodies. For further details of coordinates see Gabb et al. [12]. In the table some degenerate identical complexes included our earlier studies have been excluded.

System	Total no after FTDOCK & FILTR (set 1)	$N \leq 2.5 \text{ \AA}$ in FTDOCK list	Rank FTDOCK in set 1	Rank RPScore in set 1 (making set 2)	Rank MULTI- DOCK in set 1	Rank MULTI- DOCK in set 2 (i.e., after RPScore)	rmsd C $\alpha$ atoms (Å)
$\alpha$ CHYN–HPTI	93	1	3	2	2	2	2.0
$\alpha$ CHY–ovomucoid	85	5	11	6	1	1	1.1
Kallikrein–BPTI	349	16	128	13	2	2	1.0
Subtilisin–CHY I	26	2	8	1	12	2	2.0
Subtilisin–subtilisin I	–	–	–	–	–	–	–
Trypsin–BPTI	205	7	12	14	23	3	1.7
D1.3–lysozyme	636	2	149	75	211	38	1.8
D44.1–lysozyme	539	4	34	24	101	21	2.5
HyHEL5–lysozyme	498	2	218	36	29	29	1.5
HyHEL10–lysozyme	700	4	48	6	9	2	1.1

zymes were serine proteases and the distance filter was that at least one residue in the inhibitor must be in contact with the one of the catalytic triad (i.e., His, Ser or Asp). For the antibodies, the constraint was that the antigen must contact either the third complementarity determining region of the light or the heavy chain (CDR-L3 or CDR-H3). In this list there are at least one, but often several, complexes that are considered a good prediction. In this study a good prediction is defined as within 2.5 Å of the correct structure superposing C $\alpha$  all atoms of the predicted complex generated from unbound coordinates onto the X-ray bound complex. If just FTDOCK is used, the rank of the first good solution is given in column 4. In four out of the six enzyme/inhibitor complexes a good solution is obtained within 16 solutions. However no suitable complex is obtained for subtilisin docking to its inhibitor. The results for docking antibodies with antigens are less successful.

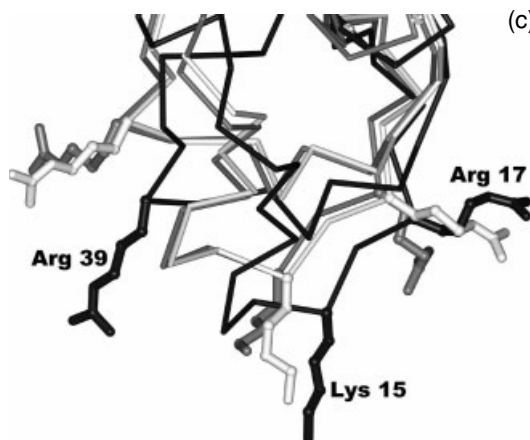
The results from FTDOCK can then be re-ranked using empirical scores (RPScore) to yield set 2. In general the rank at which one finds a good solution is decreased. A good solution is always found in the top 12% of the list of structures generated by FTDOCK (apart for subtilisin with its inhibitor). Set 2 can be further re-ranked using MULTIDOCK. The strategy was to take the top 25% of structures generated by RPScore, which is over double the fraction required in this benchmark. These are solutions are then ranked using MULTIDOCK *in vacuo* (see column 7). For the five serine proteases (i.e., excluding subtilisin and its inhibitor), a good solution is at rank 3 or better. The results for the antibody-antigen complexes are poorer with ranks as large as 38 that must be examined to ensure a good solution. Possibly the lower affinity for antibody-antigen complex formation compared to the enzyme-inhibitors studied [32] is reflected in the poorer discrimination in modelling their docking. Table 8.2 also presents the results of re-ranking the results of FTDOCK directly with MULTIDOCK (column 6). The approach of first using RPScore and then MULTIDOCK is always as good as, and generally superior, to using just MULTIDOCK for screening. Table 8.2 also gives the rmsd of the first ranked good solution. For about half of the simulations, the predicted structure is better than 2.0 Å rmsd of the bound coordinates.

Figure 8.7 illustrates some of the predicted complexes. Figures 8.7a and b show the C $\alpha$  trace of the predicted complex generated by FTDOCK superposed on the X-ray complex for trypsin-BPTI (bovine pancreatic trypsin inhibitor) and Fab D1.3-hen lysozyme. Figure 8.7c illustrates the results of side-chain refinement using MULTIDOCK. Three side-chain in BPTI that mediate the interaction with trypsin are shown. After MULTIDOCK two of the three move closer to the correct position than their original location following rigid-body docking using FTDOCK. In this benchmark, the agreement between predicted and native structure is sufficient to suggest further experiments, such as mutagenesis, to probe the interaction. Furthermore, the identification of the loop and key side-chains in an inhibitor

**Fig. 8.7**

Comparison of native and modelled protein-protein complexes. In the figures the black line denotes the X-ray structure of the complex and the grey lines the predicted complex. *Fig. a.* Superposition of the predicted and native complexes for bovine pancreatic trypsin inhibitor (BPTI) docking to trypsin. A  $C\alpha$  trace is shown. The predicted structure is generated by FTDOCK1 and the rmsd of the superposition is 1.7 Å. *Fig. b.* Superposition of the predicted and native complexes of Fab D1.3 and hen lysozyme. A  $C\alpha$  trace is shown. The predicted structure is generated by FTDOCK1 and the rmsd of the superposition is 1.8 Å.





(c) **Fig. 8.7** (continued)  
*Fig. c.* Refinement of the docking of BPTI to trypsin. Only BPTI is shown for clarity. The figure shows a C $\alpha$  trace with three side-chain that are central for the interaction of BPTI with trypsin. The grey lines are the initial results from FTDOCK1, the light lines, the prediction after refinement by MULTIDOCK. The black lines are the X-ray structure of the complex.

could be a valuable guide for the design of novel compounds of pharmaceutical use.

## 8.5 Modelling protein–DNA complexes

Knowledge of the structure of protein–DNA complexes provides insights into the structural basis of many fundamental biological processes [33]. For example an experimental structure of a transcriptional repressor with DNA will (generally) provide an understanding of why certain base sequences are required to recognise the repressor and of which residues in the repressor are responsible for interacting with DNA. This information provides both fundamental understanding and in the future opens the route for the design of drugs that specifically recognise certain DNA sequences. As for protein–protein complexes, there are often coordinates for an unbound protein that binds DNA but not for the protein–DNA complex. Thus predictive protein–DNA could be used to derive structural information, and the consequential biological insights, for such systems.

In the formation of many protein–DNA complexes there are gross conformational changes that would prevent employing a rigid-body approximation as the first step of the docking strategy. Nevertheless, there are some systems that often have only limited conformation change and to evaluate the effectiveness of predictive docking on these targets we have modelled transcriptional repressors recognising DNA [34]. The objective was to explore all possible association modes of the macromolecules. Previous work on protein–DNA docking focussed on exploring specificity once the protein

has been oriented in a correct recognition mode, such as the recognition  $\alpha$ -helix along the major groove [35, 36].

### 8.5.1

#### Method

The approach employs the first three stages of the protein–protein docking strategy amended to tackle protein–DNA complexes [34]. The steps are:

- i. rigid-body docking;
- ii. filtering with a protein–DNA scoring function and
- iii. the use of biological distance filters.

Eight repressor–DNA complexes were examined (see Table 8.3). They show a diversity of recognition modes. Five (CRO, GAL, LAC, LAM and PUR) involved the  $\alpha$ -helix/turn/ $\alpha$ -helix recognising the major DNA groove, two (ARC and MET) involved a two stranded antiparallel  $\beta$ -sheet recognising bases in the major groove and in TRP there is both major and minor groove recognition. The unbound DNA was constructed according to standard geometry. For seven of the eight complexes, the unbound repressor structure was used but for one (LAM), bound coordinates had to be employed.

In the study FTDOCK1 was used with a  $128 \times 128 \times 128$  grid that yielded a resolution of 0.51–0.87 Å. The DNA was kept as the fixed molecule. Rotational sampling was at  $12^\circ$ . A specific charge set was developed for DNA that exaggerates the partial charges on the chemical groups in the DNA helix groove and dampens the phosphate charges. The electrostatic potential is only calculated for grid nodes within 2 Å of a charged atom (on the DNA). For the protein, charges are assigned to the main-chain atoms and to Asp, Glu, Arg, Lys, Asn, Gln and His. The top 4000 structures generated by FTDOCK1 were stored.

Empirical potentials were derived to score amino-acid–nucleotide interactions. The parameters were derived from the structures of 20 protein–DNA complexes, none of which were homologous to the eight systems being modelled. An amino acid of type *a* was in contact with a nucleotide of type *b* if the distance between the  $C\beta$  atom and the base glycosidic N was less than a cut off  $d_{cut}$ . As for the protein–protein pairing scores, the random model was based on composition (i.e., a molar fraction). In a cross-validated trial on the eight complexes, the best parameters for screening were found to be  $d_{cut}$  of 12 Å and using a sparse matrix that only scored interactions with charged or polar amino-acids (C, D, E, H, K, N, Q, R, S, and T).

Two types of distance filtering were considered. In the first (filter 1), it was assumed that there is knowledge of the central two base pairs involved in recognition. Such information is commonly derived from experimental

Tab. 8.3

Results of modelling repressor docking to DNA. The complexes modelled with the Protein Data Bank codes of the bound and unbound components are: ARC (arc repressor, 1par, 1arr) CRO (cro repressor, 3cro, 2cro); GAL (CD2-GAL4 DNA binding domain, 1d66, 125d); LAC (lactose repressor, 1lbg, 1lqc); LAM (lambda phage operator repressor, 1lmb, 1lrp); MET (met repressor, 1cma, 1cmc); PUR (purR repressor, 1pnr, 1pru); TRP (trp repressor, 1tro, 2wrp). For coordinate details see Aloy et al. [34]. Filter 1 employs knowledge of the central two base pairs. Filter 2 employs filter 1 plus knowledge of some critical residues on the repressor. %CC is the percentage of correct contacts. rmsd is the deviation of the superimposed predicted and native complexes.

Complex	Ranking by shape complementarity			Ranking by empirical score				
	Rank after filter 1	Rank after filter 2	%CC	rmsd (Å)	Rank after filter 1	Rank after filter 2	%CC	rmsd (Å)
ARC	91	22	69	4.1	1	1	69	4.0
CRO	12	3	85	3.0	121	9	80	3.8
GAL	37	26	75	3.6	2	2	75	3.6
LAC	30	13	72	4.0	133	117	77	3.4
LAM	22	3	84	3.0	4	4	98	1.6
MET	-	-	-	-	-	-	-	-
PUR	9	2	68	4.3	28	28	92	2.3
TRP	4	2	65	4.2	1	1	67	3.2



work, particularly DNA footprinting, and historically this information was often available prior to determination of the structure of the complex. The second more restrictive distance constraint (filter 2) was that in addition to the base sequence, there was knowledge about one or a few repressor residues that contacted the DNA. Such information could be derived from mutagenesis experiments or can be predicted from identification of spatial clusters of conserved residues (Aloy and Sternberg, unpublished). To establish if there was a protein–DNA contact, each side-chain type was assigned an effective side-chain length ( $L$ ) ranging from 0.5 Å (Gly) to 6.0 Å (Arg) as detailed in Gabb et al. [12]. The distances ( $D$ ) between protein  $C\alpha$  atoms and the base glycosidic N atoms were calculated. There was a contact if any  $D < L + 4.5$  Å.

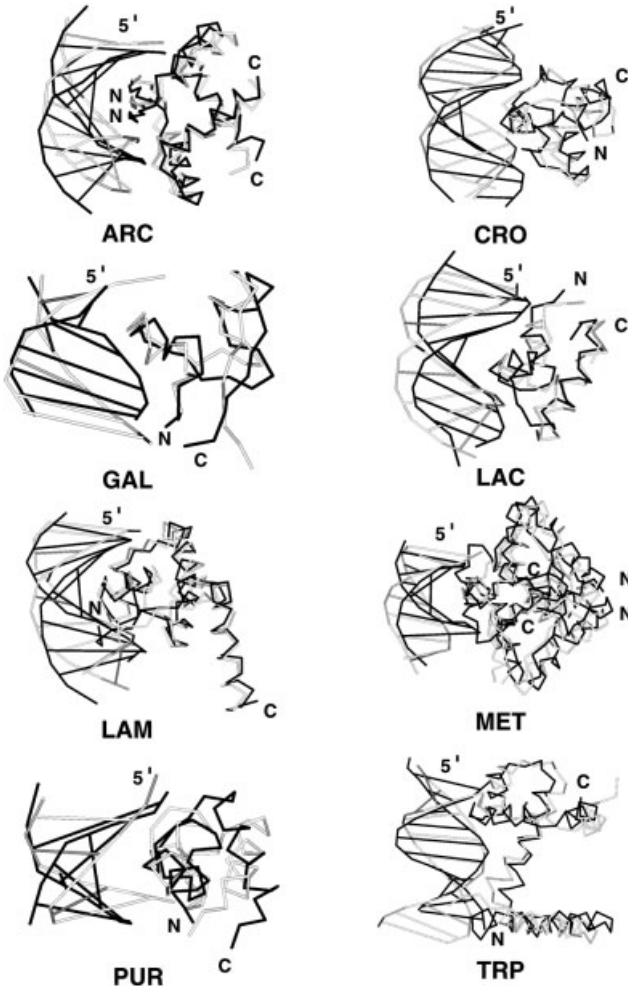
### 8.5.2

#### Results

The agreement between the predicted and native complex was quantified using the standard rmsd superposing representing the protein by  $C\alpha$  atoms and the DNA by  $C1'$  nucleotide atoms. However inspection of the results suggested that this did not provide a good guide to the quality of the prediction. Instead the percentage of correct contacts in the interface was used. First, the protein–nucleotide pairs in the native interface were defined as those pairs that had at least one non-hydrogen protein–DNA atom–atom contact of  $<5$  Å. For these pairs, the  $C\alpha$ – $C1'$  distances were measured in the native and the predicted complexes and if the difference in distance was  $<4$  Å, then a correct contact was assigned to the predicted structure. A “good” prediction was taken if the percentage correct contacts compared to the total number of interface contacts (%CC) was  $>65\%$ .

Table 8.3 gives the rank of the first “good” prediction in lists ordered by the scoring function after the two types of distance filters ranking by shape complementarity and by the empirical scoring scheme. Out of the eight systems only for MET did the approach fail to generate a “good” solution. After filter 2, shape complementarity gave 4 solutions with a rank in the top 5. When ranking by the empirical score, four complexes were in the top 5 rank after filter 1 or after filter 2. However other complexes were below the top 100 solutions. At present, the empirical score would be useful to generate a limited list (say 5) of suggestions which has about a 50% chance of including a “good” solution. In contrast, shape complementarity would be more suitable to generate a list for subsequent refinement. The predicted complexes superimposed on the native structures are shown in Figure 8.8. The first “good” solution ranked on shape complementarity is shown.

To our knowledge, this work is the only systematic study of a simulation that performed a global search of docking proteins to DNA. The results show a somewhat poorer level of discrimination that we and other groups



**Fig. 8.8**

Comparison of native and modelled repressor–DNA complexes. Superposition of native (black) and modelled (grey) complexes. The DNA is shown as the phosphate backbone with lines for the

base pairs. The repressor is shown as a  $C\alpha$  trace. The first correct structure from Table 8.3 is shown. For MET the best model is described. Reproduced with permission from Figure 8.2 of Aloy et al. [34].

are obtaining for protein–protein docking. The method therefore would have to be used cautiously, for example in systems where there is substantial experimental data or when experiments would be designed to evaluate the prediction. In addition, the repressor–DNA system studied has limited conformational change on complex formation and many systems exhibit far larger changes in structure. In general, one does not know of the extent of conformational change on complex formation in advance of knowing the structure.

## 8.6

### Strategies for protein–protein docking

Different groups have explored a variety of methods for protein–protein docking and we now review some of the major strategies that have been used, also see reviews [7–9]. We will emphasise the recent developments.

#### 8.6.1

##### Evaluation of the results of docking simulations

There are several difficulties in trying to compare the results of the different approaches for protein–protein docking. Firstly there are a variety of methods to quantify the rmsd between the predicted and native complexes. Normally this is expressed as the rmsd between superimposed C $\alpha$  coordinates of the predicted and a reference set of coordinates. There are however many approaches to calculate this rmsd. The reference molecule can be taken from the true (experimental) complex. However other groups first optimally superpose the unbound receptor onto the bound receptor in the complex and then the unbound ligand onto the bound ligand. The coordinates of the superposed unbound components is then taken as a reference as this represents the best structure one could achieve in rigid body docking. (This latter approach will be referred to here as rmsd from best attainable prediction.) Sometimes the optimal superposition of predicted receptor–ligand onto true receptor–ligand is evaluated. Other groups optimally superpose one of the molecules (typically the larger receptor) and then calculate the rmsd for the resultant predicted and true inhibitor positions. This second measure generally yields a substantially larger rmsd than the first. In addition, some groups will superimpose the entire molecules whilst others, particularly for larger molecules, will consider only residues in the interface.

One then needs also to define what rmsd constitutes a “good” prediction and groups use different cut offs. Generally with present day algorithms, the good structure rarely is consistently at top rank. Thus results cite the rank in a list at which a good model occurs. Comparison of ranks will depend on the fineness of sampling alternate docked arrangements and these vary between algorithms.

In evaluating algorithms, it is important to distinguish between results on docking starting with the components taken simply from the bound complex and studies that docking of both components starting from their unbound states. Although docking bound complexes might be helpful to develop an algorithm and to understand how it operates, there is a danger in over-optimising for stereochemical complementarity that will not occur in rigid body docking starting from unbound complexes. We therefore focus on results from unbound docking in this review. In addition, we note that in early work on docking, many of the algorithms were time consuming and the developers sometimes presented results on a very few systems. Results

on limited data sets can suffer from the problem of over optimising the approach to provide success on a few specific test cases.

### 8.6.2

#### Fourier correlation methods

The Fourier correlation method that is the basis of our program FTDOCK described above was originally introduced by Katchalski-Katzir et al. [11] and the original paper emphasised the application to docking bound complexes. One of the authors of this work was Vakser who has continued with this approach and explored its application to low resolution docking [36, 37]. In this approach, a low resolution grid (typically several Ångstroms) is used and this leads to representing only the major spatial features of the two molecules. The energy surface is therefore smoother, i.e., not at the atomic level, and the search far faster as the number of cells used to represent the two molecules far fewer. Recently Vakser [39] explored the use of low resolution docking on a database of 475 co-crystallized protein-protein complexes. Most of the database were multimeric proteins but it also included complexes of the type conventionally studied in docking. He considered the average separation between the mass centres of the receptor and the ligand in his database (48 Å). The study considered the binding region of the receptor which is defined as within a 10 Å arc traced out by the centre of mass of the ligand at this separation. Note that the orientation of the ligand is not considered, only the location of the binding site of the receptor. The docking program GRAMM was run at grid spacing of 6.8 Å and a rotational step of 20°. The receptor binding site was recognised in 52% of the complexes using this low resolution docking. For 113 complexes with a large interface area ( $> 4000 \text{ \AA}^2$ ), the success rate rose to 76%. Thus low resolution docking provides a valuable tool to delineate a putative binding site in a protein. This could then be used to provide a filter for results of higher resolution procedures.

A new approach for rapid protein docking using Fourier correlation, HEX, has been introduced by Ritchie and Kemp [40]. They employ spherical polar Fourier correlations which removes the time consuming requirement of FTDOCK and GRAMM for explicitly generating different orientations for the mobile molecule. They consider both shape complementarity and electrostatic effects. Note that their approach uses a Fourier correlation but does not use the discrete Fourier transform employed by GRAMM and FTDOCK. A full search on a single workstation takes around two hours. However, constraints on the location of the receptor binding site can readily be incorporated and this reduces the calculation to the order of minutes. In the conventional Fourier approach (FTDOCK and GRAMM), knowledge of the binding site in one of the two molecules (say the static) does not provide a constraint. Every orientation of the mobile molecule still needs to be sampled. Their program has been extensively benchmarked on unbound dock-

ing and yields results that suggest their program is a valuable tool for protein docking.

### 8.6.3

#### Other rigid-body docking approaches

Janin and coworkers [41] used spheres of different radii to replace residues and docked the resultant structures so as to maximise the buried surface area. These initial docked complexes were refined using a Monte Carlo search in which trial perturbations are generated randomly. A perturbation is always accepted if it reduces the energy and may also be accepted if the energy increases but is below a randomly chosen value determined by Boltzmann's principle. Finally a fully atomic representation (apart from hydrogen) is restored and an energy minimisation performed that allows the side-chains in the predicted interface to move. Note that the initial selection of associations to examine including flexibility is constrained by the rigid-body approximation. Cherfils et al. [41] tested the approach on two unbound docking systems and for one obtained a native-like solution at rank three.

The DOCK algorithm, developed by Kuntz and his group, is widely used in the community for the docking of low molecular weight ligands to protein receptor. It can however also be applied to protein-protein docking [42]. The DOCK approach fills the binding site of one protein (the receptor) with a cluster of overlapping spheres. Then the algorithm matches the sphere centres of this cluster with similar clusters from the ligand protein. Predicted complexes are then ranked in terms of a score for residue–residue contact. In the early study on docking starting with unbound complexes [42], individual atoms from the molecules had to be manually truncated to obtain good predictions. Although there were suggestions of which atoms may be mobile from the crystallographic thermal parameters present in the coordinate file, in other simulations one would not know in advance which atoms need to be truncated. Recently, the approach has been developed further for macromolecular docking [43]. The approach consists of three steps: defining the shape-based sites that define putative docking locations; docking using these site points; and scoring the docked complex. Complexes were scored using van der Waals and electrostatic interactions calculated from the AMBER program with united-atom parameters [27]. The method was benchmarked on re-docking several complexes starting from the *bound* coordinates. The approach was then used to suggest a model for the docking of human growth hormone to its receptor.

The group of Wolfson and Nussinov have developed an approach for docking approach based on matching critical points [44–46]. These points define the knobs and holes on the two interacting surfaces. Both surface points and surface normals are matched and the approach can be implemented speedily. A global search takes of the order of minutes. After the search, putative solutions are checked to penalise overlap. In addition, the

extent of hydrophobic packing across the interface is assessed. The method was recently tested on four different protease–inhibitor systems starting with unbound components and two bound antibody – unbound antigens complexes. The rmsd was taken between the true and predicted ligand after optimally superposing the receptor. Rankings of the first structure better than 5 Å rmsd were between 1 and 600, with several systems having rank 10 or better. For each system studied, several sets of coordinate were used in different trials. This showed that the method proved sensitive to the precise starting set of coordinates for a given biological complex. This highlights that for other algorithms (including FTDOCK) it is important to assess the dependency of the results on the precise starting coordinates.

Lenhof [47] has developed an approach for docking based on the identification of points on the surface of each molecule that could be equivalenced in a close-packed association. The search for possible rigid-body associations is then speeded up by considering which sets of three points on one molecule could be equivalenced to three points on the other molecule. Suitable docked complexes are then scored in terms of the geometric match between atoms followed by consideration of the chemical complementarity of the match. Trials with docking starting from unbound components showed that for several, but not all, systems the method yielded lists of a few complexes (<10) one of which was close (<4 Å rmsd) to the native. This method has been extended to include a treatment of side-chain flexibility in a subsequent screening that is described in 6.6 below [48].

Recently, Ausiello et al. [49] have developed a new docking procedure, ESCHER. The method starts with shape complementarity based on a slices along the Z axis of the protein surface mapped onto sets of polygons. Complementarity is assessed from the close approach of polygon vertices between the two docked molecules. Steric clashes and charge complementarity are then evaluated. The quality of the results is assessed from the rmsd of the predicted complex from the best attainable prediction. Only one true unbound docking system was studied (chymotrypsin–ovomucoid inhibitor) and good structures (rmsd for the complex <2.0 Å) was obtained at rank 3. In addition similar results were obtained for two complexes in modelling the docking of bound antibody to unbound antigen.

#### 8.6.4

#### **Flexible protein–protein docking**

Abagyan and Tortov [50] have developed a method that introduces side-chain flexibility early in the search procedure. The approach was applied to unbound docking hen lysozyme to the bound conformation of the combining site of antibody HyHEL5. A trial set of starting conformations are generated to sample space. Then a Monte Carlo search is performed starting with random rigid-body shifts (see 8.6.3 for a brief description of the Monte Carlo procedure). Following each random step, side chain torsion angles

were allowed to vary and the energy of the resultant complex minimised. This procedure was run to identify the set of conformations close to (within 20 kcal/mol) to the best structure. In this system, this leads to 30 trial structures that were then subject to extensive energy minimisation including both rigid-body shifts and side-chain rotation. The lowest energy structure was close the native complex – when the antibody coordinates were superimposed the rmsd for the C $\alpha$ , C and N backbone lysozyme atoms was 1.6 Å. The procedure was reported in 1994 and was time consuming taking 500 hours of computing time on a state-of-the-art workstation (AXP3000/400).

### 8.6.5

#### **Rigid-body treatment to re-rank putative docked complexes**

Evaluation of a scoring function to assess the stereochemistry of trial complexes is a central feature of all docking algorithms. However additional methods can be applied to re-rank putative complexes. An important distinction must be made between whether the re-evaluation treats the complexes as rigid bodies or if the procedure first includes conformational flexibility to refine the complex before re-ranking. This distinction is illustrated by the distinction in our protein-protein docking procedure described in sections 3 and 4. The scoring potentials implemented in RPSCORE are used to re-rank rigid-body complexes. In contrast MULTIDOCK refines side-chain conformation and then re-ranks the complexes.

Prior to the development of RPSCORE and MULTIDOCK, in our Laboratory we introduced a rigid-body re-ranking procedure referred to as the continuum model [51]. This approach evaluated the electrostatic and hydrophobic contribution to relative energies of a series of putative docked complexes treating the solvent macroscopically (i.e., as a continuum) rather than explicitly including the solvent atoms. The total electrostatic energy of binding involved the loss of interaction between the solvent and each of the protein components independently followed by the interaction between the two proteins. The system treated each protein as a low dielectric surrounded by a high dielectric solvent. The electrostatic contributions were evaluated using the program DelPhi [52]. This program maps the protein-solvent system onto a grid and then calculates the resultant electrostatic effects considering both the local dielectric (protein or solvent) and the local charge distribution. In addition, in the continuum model we optimized the position of polar hydrogens prior to the calculation of the electrostatic effects. The hydrophobic effect was quantified in terms of the change to the molecular surface (MS) when the two proteins dock [53]. Generally the hydrophobic effect is quantified in continuum modelling as being proportional to the change in solvent accessible surface (SAS) area [54], where SAS is the surface traced by the centroid of a hypothetical probe water molecule as it rolls along the surface of the protein [53]. Molecular surface can be considered as the surface representing the protein/solvent-probe interface. Pre-

viously, Jackson and Sternberg [55] suggested that MS provides a better model for the hydrophobic effect than SAS. The continuum method was applied to re-rank the results on three enzyme-inhibitor systems generated in unbound docking using DOCK by [42]. The continuum model was able to identify a near native solution as having a particularly low energy. We are planning to evaluate the use of this continuum model to screen the results of complexes generated by our docking suite after their refinement by MULTIDOCK.

Recently an approach to re-rank complexes treated as rigid bodies was developed by [56]. They developed a new soft mean-field potential derived from analysis of protein-protein contacts in crystal structures. The hydrophobic-hydrophobic atom potential was applied to screen docked complexes. The series of docked complexes were generated by the approach of Cherfils et al. [41], see above. Four systems were studied: a reconstitution of the bound components of barnase and barstar, two docking starting with unbound coordinates ( $\beta$ -lactamase- $\beta$ -lactamase inhibitor and chymotrypsin-ovomucoid) and one docking bound antibody coordinates to lysozyme. For the first three of these systems studied, the lowest free energy trial complex was a "good" prediction being within 2.5 Å of the true complex. In the fourth system, such a "good" solution was found at rank 2 on ordering by increasing energy. Thus the potentials are highly effective in screening for a "good" docked structure.

#### 8.6.6

##### **Introduction of flexibility to re-rank putative docked complexes**

A method to include side-chain flexibility into the refinement and re-ranking of docked complexes has been developed by [57]. They considered the three enzyme-inhibitor complexes generated using unbound docking by [42] and subsequently studied by the continuum model by [51], see Section 8.6.5 above. The conformation of inhibitor side chains buried in the docked complex with the enzyme were examined by exhaustive conformational search for energetically more stable positions using CONGEN [58]. The resultant complexes were then scored by a measure of their relative energetic stability. The function considering the electrostatic interaction between the molecules, desolvation and side-chain conformational entropy. Desolvation was evaluated using the rapid approach of being proportionality to the change in accessible surface area where the constant of proportionality depends on the nature of the atom [59]. Side-chain conformational entropy was assumed to be proportional to the change in solvent accessible surface [60, 61]. For each complex the procedure lead to the identification of a native-like structure as having the lowest energy of association. In addition, in general there was an improvement in the agreement between the native and predicted geometry of the side-chains whose conformation was adjusted.



A method that provides an extensive sampling of side-chain conformations has been recently reported [48]. The set of rigid-body docked complexes for screening were generated by the method of [47] described in Section 8.6.4. All side chains with rotatable bonds to non-hydrogen atoms that are part of the putative interface are considered. A combinatorial search for favourable orientations is undertaken using computational methods (dead-end elimination and branch-and-bound) to prune the search space and thereby speed up the calculation. The docked structures are evaluated in a method similar to the continuum model of Jackson and Sternberg [51]. They workers considered three enzyme-inhibitor complexes and for each the lowest energy conformation was close to the native complex.

## 8.7

### Blind trials of protein–protein docking

There are several problems in comparing the success of docking approaches from different groups on a set of previously known structures. We have above (8.6.1) described the variety of different measures used to report the agreement between a predicted and the true complex and the problems of the fineness of sampling the space of docked conformation. In addition, working on reproducing the docked structures of known complexes can lead to optimising an approach until it is successful. A developer might be aware of specific features of the stereochemistry of the known complexes and include these features in the algorithm leading to a bias towards known rather than unknown targets. In recognition of these problems, there have been two blind tests of protein–protein docking. In these tests, docking groups were supplied coordinates of the components and challenged to predict the structure of the complex prior to its structure being reported.

The first test, the Alberta challenge, was organised by James and Strynadka [62] and involved docking the coordinates of unbound  $\beta$ -lactamase to those of its unbound inhibitor. Six groups submitted entries (see Table 8.4). The results were impressive because all entrants identified as their favored suggestion a model that had an rmsd for superimposed  $C\alpha$  atoms of no more than 2.5 Å. However some groups submitted other entries that were far from the correct structure. Many of the entries used a biological filter requiring that the inhibitor docked to the known active site of  $\beta$ -lactase. The closest prediction to the true structure, submitted by Eisenstein and Katchalski-Katzir, had an rmsd of all superimposed  $C\alpha$  atoms of 1.1 Å which corresponds to an rmsd for the inhibitor  $C\alpha$  atoms of 4.6 Å when the enzyme was optimally superposed. Their successful approach employed the Fourier correlation method developed by Katchalski-Katzir et al. [11]. Their version only considered rigid-body shape complementarity which clearly was sufficient in this system for a successful prediction. However, in trials with

**Tab. 8.4**

Results of the Alberta Docking Challenge. The complex was the docking of unbound  $\beta$ -lactamase coordinates with unbound inhibitor coordinates. The value of rmsd-all is the rmsd of superposed main-chain atoms of the complex. The value of rmsd-inhibitor is the rmsd for the main-chain atoms of the inhibitor after optimally superposing the enzyme. Note that the best rank was often decided by the user rather than automatically by the algorithm.

<i>Group</i>	<i>No of models</i>	<i>rmsd-all of best ranked (Å)</i>	<i>rmsd-all of other models (Å)</i>	<i>rmsd inhibitor of best ranked (Å)</i>
Abagyan & Totrov	3	1.9	11.3–16.2	6.6
Duncan, Rao and Olson	14	1.9	2.0–17.7	4.5
Eisenstein & Katchalski-Katzir	3	1.1	13.4–14.1	3.4
Jackson & Sternberg	1	1.9	N/A	4.0
Janin, Cherfils & Zimmerman	4	2.5	2.5–16.0	6.1
Shoichet & Kuntz	15	1.8	2.3–18.7	3.8

FTDOCK, we found that for many other systems electrostatic effects need to be included.

In the Alberta challenge, four other groups all performed a global search and submitted a model between 1.9 Å and 2.5 Å from the true complex corresponding to an rmsd for the inhibitor of between 4.0 Å and 6.6 Å. The methods were diverse. Two methods were totally rigid-body dockings – the DOCK approach from Shoichet and Kuntz (see 8.6.3 above and Shoichet and Kuntz [42]) and the comparison of protein surfaces using a smoothed representation (by Duncan Rao and Olson [63]). The model submitted by Janin, Cherfils and Zimmerman (see Section 8.6.3 above and Cherfils et al. [41]) started with a rigid-body docking but then includes side-chain optimisation. Only one approach, from Abagyan and Totrov, used a procedure (see Section 8.6.4 and Totrov and Abagyan [64]) that incorporated flexibility at an early stage of the search procedure. These results suggest that for this system, the rigid-body approximation is appropriate for docking simulation. In addition, the different approach to match surfaces yield broadly similar results. In contrast to the other five submissions that performed a global search, the use of the continuum model (see Section 8.6.5 and Jackson and Sternberg [51]) for screening was evaluated by Jackson and Sternberg. They used the results generated from DOCK and were able to identify a single preferred complex that was close to the native.

The second test was as part of the second comparative assessment of protein structure prediction (CASP2) [65]. The target was an antibody-hemagglutinin complex and coordinates of unbound hemagglutinin and bound antibody were supplied. This was a difficult target given the size of the complex and only four groups entered. Multiple entries were allowed and confidences then had to be assigned so that the total was 100%. No

**Tab. 8.5**

Results of the CASP2 Docking Challenge. The complex was the docking of unbound hemagglutinin coordinates with bound antibody coordinates. The rmsd refers to rmsd for the interface Fab C $\alpha$  atoms after optimal superposition of the hemagglutinins. The interface atoms of the antibody are those C $\alpha$  atoms within 8 Å of the hemagglutinin–Fab interface. Correct residue–residue contacts are defined as inter-protein atom–atom distances less than the sum of the van der Waals radii plus 1 Å. There were a total of 59 contacts. The mean values refer to the weighted scores from all the predictions. Min and max are the minimum and maximum values.

Group	No of models	rmsd (Å)		No of correct contacts	
		mean	min	mean	max
DeLisi	2	18.3	15.1	4.5	5
Rees	2	32.3	30.6	0	0
Sternberg	8	20.2	8.5	1.8	8
Vakser	1	9.5	9.5	0	0

group submitted any entry that was close to the true complex (Table 8.4). The averaged best prediction, from Vakser, used the Fourier correlation approach. Only one model was submitted and this yielded an rmsd for the interface C $\alpha$  atoms of the antibody of 9.5 Å, calculated after optimally superposing the hemagglutinin. However this prediction did not have any correct contacts (defined as atoms that are separated by their van der Waals radii plus 1 Å). This submission was based on the Katchalski-Katzir Fourier method implemented for low-resolution search in Vakser's program GRAMM [37, 66]. The single prediction that was closest to the native (an rmsd of 8.5 Å for the interface antibody C $\alpha$  atoms) was also based on the Fourier correlation method as applied by the ICRF group of Sternberg, Jackson and Gabb using preliminary versions of FTDOCK and MULTIDOCK. However the approach when applied did not provide a single confident prediction and accordingly the group submitted eight entries. There were entries using two other approaches – a matching of surfaces using graph theory from Rees and co-workers [67] and the approach of DeLisi and coworkers (see 8.6.6 and Weng et al. [57]).

From only two blind trials, one cannot draw definitive conclusions. However, in both challenges the Fourier correlation approach of Katchalski-Katzir yielded the best submission which suggests that it must be considered as a valuable strategy for macromolecular docking.

## 8.8

### Energy landscape for protein docking

This chapter has considered predictive protein–protein and protein–DNA docking. However there remains a related question of how two molecules

can associate within the time observed biologically. The problem is that association rates for protein-protein docking would be  $\sim 10^3 \text{ M}^{-1} \text{ s}^{-1}$  if it were just governed by diffusion and a correction for orientational constraints. However observed rates are typically far faster being of the order of  $10^5\text{--}10^7 \text{ M}^{-1} \text{ s}^{-1}$ . This increase in speed is often attributed to long range effects, particularly long range electrostatic steering. Janin [68] has shown that long-range electrostatic steering can enhance association rates by up to  $10^5$  fold.

Recently, the energy surface near the native docked complex has been modelled in terms of empirical atomic contact energies and Coulomb electrostatic interactions [69]. The Coulomb interaction has a distant-dependent dielectric and a cut off of 20 Å. Thus the energy surface does not consider long-range effects. The study showed that the energy gradient provided by the surface provides a funnel towards the docked structure that increases the probability that an encounter will evolve into the stable complex by about 400-fold. Given the simplified treatment of the interaction energy, in real system energy funnels could provide even greater enhancements for the rates of association. Thus even without long-range electrostatic effects, energy funnels provide a possible explanation for the observed relatively rapid association rates. More generally, the role of funnels in directing protein folding and function has been reviewed by Tsai et al. [70].

## 8.9

### Conclusions

This review presents the current status of protein-protein and protein-DNA docking. The field will advance and new approaches will be proposed. To identify useful approaches, it is essential that results are presented on docking starting with unbound components. In addition, our experience is that good results can readily be obtained for a few systems and it therefore important to benchmark an algorithm on a large test set. Indeed part of the design principle for an algorithm should be that it sufficiently fast to be tested on a large data set. However, the best approach to compare approaches is via blind tests.

How useful is predictive protein docking in current work in structural bioinformatics? In general, the status of the better algorithms is that for systems that do not involve substantial conformational change, one can generate a solution close to the native structure ( $<3.0 \text{ \AA}$  rmsd for superposed interface C $\alpha$  atoms) and this solution can be ranked at 10 or better. However results are this type are not guaranteed – typically it will occur for only half the systems studied.

We will soon have the genome sequences for man and many pathogens. Structural genomics initiatives [71] will exploit this sequence information

and over the next few years the three-dimensional structures for many proteins in these genomes will be determined. There will remain the difficulty of obtaining structural information of how the proteins interact. Thus the application are for predictive docking will increase drastically over the next few years. The predicted structures of docked complexes will suggest further experimentation such as mutagenesis to probe activity and alter binding affinities between molecules. For example, the systematic modification of the sequence of the combining site of antibodies based on a predicted structure could lead to molecules that recognise antigens with improved or altered specificity. These engineered antibodies should prove to have a major role in both medical and commercial applications, for example in recognising tumors. However in many application, high molecular weight proteins are not suitable as pharmaceutical agents due to problems such as being metabolised rapidly or generating an allergic response. However predictive models of a protein–protein complex could suggest the design of low molecular weight compounds that mimic the recognition region these could provide the lead for drug discovery.

The development of the algorithms for protein-protein docking has been highly successful over the 1990s and now is a viable approach for modelling. Over the next decade, we envisage that will there be further advances in the methodology together with many application of predictive docking in both fundamental and applied research.

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## Appendix

Thomas Lengauer

### Glossary of algorithmic terms in bioinformatics

#### A\* algorithm

A heuristic procedure for navigating in large search spaces that are formally described by graphs. The algorithm originated in the field of Artificial Intelligence in the 1960s (Hart et al., 1968). The algorithm finds a cheapest path in a graph from a start vertex to one among a selected set of goal vertices. As usual with path search algorithms the path is extended away from the start vertex step by step. At each step the next most promising vertex  $v$  to proceed to is selected. The selection is based on a scoring function that encompasses two terms

- a term amounting to the cost of the initial path segment from the start vertex to  $v$
- a term estimating the cost of the optimal remaining path segment from  $v$  to the closest goal vertex.

If we require that the estimate underestimates the actual costs of the remaining path segment then we can prove that the A\* algorithm always finds a cheapest path. If the estimator is always zero, then the path search amounts to a normal breadth-first traversal of the graph (Russell & Norvig, 1995).

In bioinformatics, the A\* algorithm has been used in conformational search in docking a (rigid) ligand into a flexible protein structure ((Leach, 1994), see Chapter 7 of Volume I). The path assigns rotameric states to side chains in the protein binding pocket, one after the other. A goal vertex is one that corresponds to a complete assignment of rotameric states. The start vertex is the one where no rotameric states have been assigned.

The estimation of the cost of the optimal remaining path segment is performed by generalizing a process called *dead-end elimination* (Desmet et al., 1992). This process precomputes for each rotamer  $j$  of a side chain whether there is another rotamer  $k$  of the same side chain that leads to an

energetically better conformation, independent of rotameric choices for other side chains. If so, rotamer  $j$  is excluded from the search. Leach (Leach, 1994) generalizes this process from just excluding alternatives to computing a lower bound on path cost by considering optimum rotamers independently for each side chain yet to be placed and adding up the corresponding cost terms. This extension, which Leach also adapted to the placement of side-chains in homology-based modeling (Leach & Lemon, 1998), is described in detail in Chapter 5 of Volume I.

### Branch & bound

Search method in tree-structured search spaces (Cormen et al., 1990). The original problem instance corresponds to the root of the tree. Internal nodes of the tree are generated iteratively, beginning with the root, by decomposing a problem instance into alternatives. Each alternative becomes a child of the node representing the subdivided problem. The solution of any of the alternatives (children) solves the original problem. Problem instances that are not decomposed further correspond to leaf nodes of the tree. They can be solved directly without any further subdivision. The solution of the original problem then amounts to completing any path from the root to a leaf node. Each interior node represents a partial problem solution that amounts to the set of alternatives already decided upon that are represented by the nodes along the path.

In general the tree is very large but not very deep. The depth of the tree (the length of the longest path from a root to a leaf) corresponds to the number of decisions to be taken to assemble the solution, which is limited in general (linear or a small-degree polynomial in the problem size). In contrast, the number of leaves of the tree, i.e. the number of alternative solutions, is usually huge, often exponential in the problem size. Thus traversing the tree efficiently becomes a major problem.

Often, problem solutions (complete or partial) are labeled with costs. Then, an optimal solution is represented by a path from the root to a leaf with optimal cost. In general, in order to find an optimal solution, the whole tree has to be enumerated. Complete enumeration can be avoided if we can provide for each node  $v$  inside the tree a lower bound (for minimization, upper bound for maximization) on the cost of all leaves that are contained in the subtree rooted at  $v$ . If this bound exceeds the cost of any solution that we have found before in the tree traversal, the whole subtree rooted at  $v$  can be disregarded. The sharper the lower bound the larger the regions of the tree that can be eliminated in this branch & bound procedure.

When traversing the whole tree, a branch & bound algorithm is certain to produce an optimal solution. Even when stopped early, the algorithm generally has found some solution already that it can return. In addition, the

lower bounds computed so far can be used to generate a lower bound for the optimal solution of the problem and thus a quality certificate for the solution returned. The longer the algorithm runs the better the solution it returns and the closer the lower bound to the cost of the optimal solution. In typical cases, a branch & bound algorithm that runs to completion will spend about 10% to 20% of its time finding the optimal solution. The remaining 80% to 90% are spent on completing the tree traversal in order to make sure that there is no better solution.

A branch & bound algorithm for protein threading has been introduced by Lathrop and Smith (Lathrop & Smith, 1996). Here, the alternatives amount to different alignments of secondary structure blocks in proteins, and the cost function estimates the energy of the resulting protein structure. The algorithm is described in detail in Chapter 6 of Volume I. A branch & bound algorithm for docking ligands into protein structures with flexible side chains has been described in (Althaus et al., 2000), see also Chapter 8 of Volume I. The  $\rightarrow A^*$  algorithm with dead-end elimination is a variant of a branch & bound algorithm, as described in Chapter 5 of Volume I (for side chain placement).

## Cluster analysis

A basic technique for data classification. Cluster analysis is aimed at grouping items in a set into clusters. The item in each cluster should share some commonalities that justify to group them. The similarity or difference between different items is usually measured by a function of pairs of items. If we measure difference (similarity), the values decrease (increase) with increasing similarity between a pair of items.

Clustering procedures can be distinguished with respect to the structure of the item set. In bioinformatics, mostly, the items are points in Euclidean space of some dimension or else they are nodes in a graph or hypergraph.

Hierarchical clustering procedures iteratively partition the item set into disjointed subsets. There are top-down and bottom-up techniques. The top-down techniques partition can be into two or more subsets, and the number of subsets can be fixed or variable. The aim is to maximize the similarity of the items within the subset or to maximize the difference of the items between subsets. The bottom-up techniques work the other way around and build a hierarchy by assembling iteratively larger clusters from smaller clusters until the whole item set is contained in a single cluster. A popular hierarchical technique is nearest-neighbor clustering, a technique that works bottom up by iteratively joining two most similar clusters to a new cluster.

Algorithms for computing phylogenies are closely related to hierarchical clustering methods (Gusfield, 1997).

Nonhierarchical techniques do not build up a tree structure of clusters.

Probably the most popular nonhierarchical clustering technique groups the items – in this case points in Euclidean space – around a fixed number  $k$  of cluster centers. The clusters are first determined randomly and then the following two steps are iterated. First the center of each cluster is computed, then the items are reassigned to the cluster whose center is closest. The iteration proceeds until the clusters do not change any more. If we extend this method such as to recalculate cluster centers everytime that we reassign an item to a new cluster then we obtain the widely used  $k$ -means clustering algorithm (Kaufman & Rousseeuw, 1990).

In bioinformatics, clustering is used in the analysis of protein families – sequence families and multiple alignment (Chapter 2 of Volume I), structure families (Chapter 6 of Volume I), or families of proteins with the same function. Furthermore gene expression data are clustered with respect to similarity in the expression profiles (Chapter 5 of Volume II), clustering is used in grouping similar molecular conformations in protein modeling (Chapter 5 of Volume I) or molecular structures or interaction geometries in docking (Chapters 7 and 8 of Volume I). In proteomics, gel images are clustered (Chapter 4 of Volume II). Clustering is also a major approach to analyzing molecular similarity (Chapter 6 of Volume II).

### **Divide & conquer**

One of the basic algorithmic techniques, which solves a problem by subdividing it into a usually small set of subproblems, solving those problems and then assembling their solution to a solution of the original problem (Cormen et al., 1990). The subproblems are solved in the same fashion, unless they can be solved directly without further subdivision. In contrast to the tree search described above under  $\rightarrow$  branch & bound, in divide & conquer algorithms we need to solve not just one but all subproblems of a problem to solve the problem. The procedure is most effective, if the subdivision is balanced in the sense that all subproblems of the same problem have about equal size.

Divide & conquer algorithms for sequence alignment are described in Chapters 2 of Volume I and Volume II, applications to the analysis of molecular similarity are described in Chapter 6 of Volume II.

### **Dynamic programming**

Tabular method of discrete optimization. The basis of dynamic programming is the *principle of prefix optimality*. This principle states that the optimal solution to the optimization problem can be composed of optimal solutions of a limited number of smaller instances of the same type of problem. The tabular method incrementally solves subinstances of increas-

ing size. The solutions of larger problem instances are assembled from previously computed solutions of smaller problem instances. The method stops when the original problem instance has been solved. The runtime for solving each subinstance tends to be constant or linear in the problem size. The number of subinstances can often be limited to a low-degree polynomial in the size of the problem instance (Cormen et al., 1990). For hard optimization problems this number may grow exponentially, however.

Sequence alignment (Chapter 2 of Volume I) is a prime example of a problem in bioinformatics that can be solved efficiently with dynamic programming. Dynamic programming for sequence alignment is also discussed extensively in (Gusfield, 1997; Waterman, 1995). Other examples of dynamic programming algorithms occur in Chapters 3 and 6 of Volume I and Chapter 2 of Volume II.

### Energy minimization

Optimization method that aims at minimizing the energy of a molecular system that is usually described in the coordinates of the participating atoms. Since we usually deal with large molecules, the dimension of the space over which the energy is to be minimized is large (goes into the many thousands). Usually the energy function has a large number of local minima whose energy is close to or exactly the global minimum. In this context, finding the global minimum is very difficult (multi-minima problem). Depending on the type of energy function, different optimization methods are used. If we use local optimization methods that descend to the next local minimum we have to judiciously choose starting points for the optimization. Still, in the presence of many local minima, with local optimization we cannot hope to sample the solution space adequately to find the global minimum. Thus Monte Carlo methods such as  $\rightarrow$  simulated annealing are employed that manage to escape from local minima.

Often the energy minimization is hampered by the fact that the energy function used is inaccurate. In addition, nature actually minimizes the *free energy* as opposed to the internal energy. Accurate computational models for free energy scoring require sampling of many conformations and are hence burdened by either substantial statistical error or extensive computing resource requirements. A particular problem is the incorporation of water, which makes large entropic contributions to the free energy. If water is modeled as a large set of discrete molecules large computing requirements result. The alternative is to model water as a continuum which, however, means another approximation.

Energy minimization is a voluminous field with dominating physico-chemical and thermodynamic aspects (Christensen & Jorgensen, 1997; Church & Shalloway, 2001; Trosset & Scheraga, 1998).

## Fourier transform

A very important mathematical operation that transforms functions (continuous Fourier transform) or vectors (discrete Fourier transform) into the frequency domain (Bracewell, 1986). This means that the function is described in terms of the contributions of basic (sine) waves with different frequencies that compose the function. Besides its very fundamental physical relevance, the Fourier transform, especially its discrete version, has computational significance (Cormen et al., 1990). The reason is twofold. First, the Fourier transform of a vector of length  $n$  can be computed quite quickly, namely in time  $O(n \log n)$  by the so-called fast Fourier transform (FFT) algorithm (Cooley & Tukey, 1965). Second, the important but quite complex operation on vectors that is called convolution, can be reduced to the much easier pointwise multiplication, when transformed into the frequency domain. The convolution of two vectors of length  $n$  requires time  $O(n^2)$  when performed in a straightforward fashion. If we instead first Fourier transform the vectors, then pointwise multiply the result and then inverse Fourier transform, we spend  $O(n \log n)$  time doing the transforms and  $O(n)$  time for the multiplications, which amounts to a total of  $O(n \log n)$  time. The convolution operation is related to polynomial and integer multiplication and therefore these are operations that can be sped up by using the Fourier transform. In protein-protein docking (Chapter 8 of Volume I), the convolution operation comes up in the analysis of the steric complementarity of two rigid protein structures. Thus Fourier transform helps there, as well.

## Genetic algorithm

A class of algorithms that mimics nature's way of optimization by variation and selection (Goldberg, 1989). Genetic algorithms work on populations of solutions to an optimization problem. Each solution (phenotype) is encoded in a string form called the chromosome (genotype). Chromosomes are varied by mutation only (evolutionary algorithm (Fogel et al., 1966)) or by mutation and cross-over, i.e. merging parts of two chromosomes in analogy to the recombination process taking place during meiosis (Holland, 1973). A selection function is applied to the resulting phenotypes. This process is iterated through a number of generations. Typical population sizes rank from 10 to a few hundred. Generally, a few dozen generations are computed. The key ingredients of the development of an evolutionary algorithm are the suitable choice of the chromosomal representation of the solutions, the mutation and cross-over operators and the diverse parameters governing the execution of the algorithm.

In bioinformatics, genetic algorithms are popular as a tool for fast prototyping. For this reason genetic algorithms can be found in practically all



facets of bioinformatics. In contrast, exploiting all of the power that lies in the approach of genetic algorithms requires an expert, and is an involved task.

This book mentions genetic algorithms in Chapters 6 and 7 of Volume I. An overview of genetic algorithms in molecular recognition is given in (Willett, 1995). In addition, we mention here recent application of genetic algorithms to protein structure superposition (Szustakowski & Weng, 2000), protein folding (Schulze-Kremer, 2000), protein-protein docking (Gardiner et al., 2001), and drug selection from combinatorial libraries (Illgen et al., 2000).

### **Greedy algorithm**

A heuristic optimization strategy that operates on the same kind of tree-structured optimization problem as the  $\rightarrow$  branch&bound algorithm. Whereas branch&bound algorithms aim at covering the whole tree, greedy algorithms generate only one path from the root of the tree to a (hopefully) close to optimal leaf node. The path is extended from the root towards the leaf by selecting one alternative to proceed at each step. This selection is usually made heuristically. Therefore, we also speak of greedy heuristics. The notion “greedy” originates from the fact, that each decision for an alternative, when taken, is never reconsidered or taken back.

Since greedy algorithms generate only one path in the tree they are generally much faster than branch&bound algorithms, Therefore greedy algorithms are an alternative to branch&bound algorithms, whenever runtime efficiency is of prime importance or if no effective lower bounding procedures are available.

In contrast, greedy algorithms usually provide solutions of significantly lower quality than branch&bound algorithms. Also no quality certificates can be given for solutions computed by greedy algorithms Generally, greedy algorithms are used in an early stage of algorithm development. There is a subclass of optimization problem for which greedy algorithms exist that are guaranteed provide the optimal solution. These problems are called matroid problems (Cormen et al., 1990).

Greedy algorithms are explicitly or implicitly mentioned in many chapters of this book.

### **Gibbs sampler**

A specific, in some sense the simplest Markov Chain Monte Carlo algorithm ( $\rightarrow$  Monte Carlo methods). In Gibbs sampling one samples the variables of a solution space one after the other. Each variable is drawn from a conditional distribution on the values of the other variables that were pre-

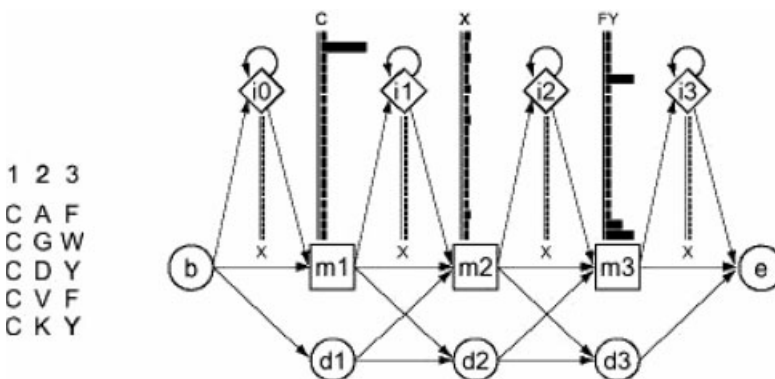
viously chosen. The conditional distributions must be accessible. The order in which variables are sampled one by one or in groups differs among different Gibbs sampling methods (Geman et al., 1984; Lawrence et al., 1993).

### Hidden Markov models

Hidden Markov models (HMMs) are a popular variant of stochastic models for analyzing biomolecular sequences. HMMs are used for characterizing protein families (Baldi & Chauvin, 1994), (multiply) aligning protein sequences (Krogh et al., 1994), detecting remote protein homologies (Karplus et al., 1998), protein threading (Karplus et al., 1997), predicting signal peptides (Nielsen & Krogh, 1998) and transmembrane domains (Sonnhammer et al., 1998) in proteins, finding secondary structures in proteins (Asai et al., 1993), finding and characterizing promoters in DNA sequences (Pedersen et al., 1996), and gene finding (Henderson et al., 1997; Kulp et al., 1996).

An HMM is essentially a Markov chain ( $\rightarrow$  Monte Carlo methods). Each state inside the Markov chain can produce a letter, and it does so by a stochastic Poisson process that chooses one of a finite number of letters in an alphabet. Each letter has a characteristic probability of being chosen. That probability depends on the state and on the letter. After the state produces a letter, the Markov chain moves to another state. This happens according to a transition probability that depends on the prior and succeeding state.

We exemplify HMMs with Figure 1, which is taken from (Eddy, 1998). Figure 1 depicts a Markov model for protein sequence alignment. Sometimes this specific topology of a hidden Markov model is called a *profile hidden Markov model*. There is a specific *start* state *b* and a specific *final* state *e*. The *Match* states (*m1* to *m3*) represent sequence positions. The *Insert*



**Fig. 1**  
A profile hidden Markov model (with permission of Oxford University Press).

(i0 to i3) and *Delete* (d1 to d3) states allow for inserting or jumping over sequence positions. Above each match state and below each insert state is a histogram of the probabilities with which each letter from the alphabet of amino acids is generated by the model. The model has been trained on the five three-letter sequences shown on the left.

The length of the hidden Markov model is the number of match states. A profile HMM stands for a family of – in this case protein – sequences.

Basically there are three algorithmic problems associated with an HMM.

1. **Training:** Set the probabilities of an HMM such that the protein sequences you want the protein family modeled by the HMM to belong to are generated with highest probability possible (Baum et al., 1970; Rabiner, 1989). Special measures are taken to avoid overfitting (see *Regularization* below). Training is by far the most complex of the three problems. It amounts to a minimization in a high-dimensional space with an irregular cost function with many minima. General solution methods boil down to local optimization procedures combined with a judicious choice of a limited set of starting solutions. Much progress can still be made on this problem.
2. **Classification:** Given a trained model and a protein sequence, evaluate the probability with which the model generates the sequence. If this probability is above a suitable threshold predict that the sequence belongs to the family modeled by the HMM. Otherwise predict the opposite. For profile HMMs and other special cases of HMMs this problem can be solved in linear time in the length of the model with  $\rightarrow$  dynamic programming methods. In general the algorithm requires time  $O(n^2t)$  where  $n$  is the number of states in the HMM and  $t$  is the length of the sequence to be generated by the HMM (Rabiner, 1989).
3. **Alignment:** Given a trained model and a protein sequence, determine, which path through the model is the most probable to generate the sequence. In the case of profile HMMs this path represents an alignment of the sequence to the model. The alignment can also be done in linear time in the length of the model with  $\rightarrow$  dynamic programming methods (Forney, 1973; Rabiner, 1989). Thus, by aligning many sequences to the model one after the other we can generate a multiple alignment in linear time in the length of the model times the number of sequences. This is in contrast to many other methods of multiple sequence alignment whose runtime grows exponentially in the number of sequences. Of course, HMMs transfer much of this complexity into the training phase.

A very nice general tutorial of HMMs is given in (Rabiner, 1989). An extension of HMMs from finite automata (which is the deterministic version of a Markov chain) to context-free languages leads to the concept of a *stochastic context-free grammar*. Such grammars have been used in an analogous way to predict RNA secondary structures (Grate et al., 1994).

## Molecular dynamics

In molecular dynamics methods the equations of motion of a molecular assembly are integrated numerically (Sprik, 1996). Thus we obtain not only a molecular structure but a trajectory of molecular motion. Due to high frequency atomic oscillations the time step for the numerical integration must be very small (about  $10^{-15}$  sec). This results in extraordinarily high computing requirements even for computing trajectories that span a few nanoseconds. In contrast, proteins fold in a few seconds. This difference in time scale cannot be overcome by sheer computing power. Thus methods are under investigation that can smooth out the high frequency oscillations without invalidating the final outcome (Elber et al., 1999).

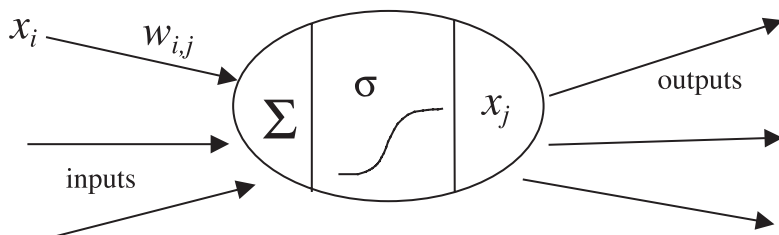
Molecular dynamics methods are primarily used for the refinement of structural models (Li et al., 1997) or the analysis of molecular interactions (Cappelli et al., 1996; Kothekar et al., 1998). In both cases the time scales to be simulated are within range of current computing technology. Another application area is the study of allosteric movements of proteins (Tanner et al., 1993). Molecular Dynamics approaches to protein-ligand docking are described in Chapter 7 of Volume I.

## Monte Carlo methods

General class of algorithmic methods that involve a stochastic element, i.e., that let the computer make random decisions (Binder & Landau, 2000). An important subclass, the so-called *Markov Chain Monte Carlo (MCMC) methods* can be understood as acting on Markov chains. A Markov chain is a stochastic finite automaton that consists of states and transitions between states (Feller, 1968). At each time we consider ourselves as resident in a certain state of the Markov chain. At discrete time steps we leave this state and move to another state of the chain. This transition is taken with a certain probability characteristic for the given Markov chain. The probability depends only on the two states involved in the transition. Often one can only move to a state in a small set of states from a given state. This set of states is called the *neighborhood* of the state from which the move originates.

A Markov chain is *ergodic* if it eventually reaches every state. If, in addition, a certain symmetry condition – the so-called criterion of *detailed balance* or *microscopic reversibility* – is fulfilled, the chain converges to the same *stationary probability distribution* of states, as we throw dice to decide which state transitions to take one after the other, no matter in which state we start. Thus, traversing the Markov chain affords us with an effective way of approximating its stationary probability distribution (Baldi & Brunak, 1998).

Monte Carlo algorithms pervade bioinformatics. Especially the chapters dealing with molecular structures mention this class of algorithms fre-



**Fig. 2**  
Neuron  $j$  in a neural network

quently, for instance, to generate diverse ensembles of molecular structures or to perform energy minimization. → Simulated annealing is a special instance of a Monte Carlo algorithm. Chapter 7 of Volume I contains a detailed overview of Monte Carlo approaches to protein-ligand docking.

### Neural network

A computational connection pattern that is formed in (simplified) analogy to biological (cortical) neural networks and used successfully for the classification of unstructured data. Neural networks have a broad range of applications in biology and chemistry (Schneider & Wrede, 1998).

In general, a neural network is a directed graph with nodes called neurons (see Figure 2). There may be a directed edge  $(i, j)$  from each neuron  $i$  to each neuron  $j$ . The directed edges  $(i, j)$  of the network are labeled with numbers  $w_{i,j}$  called *weights*. Each neuron is equipped with an activation function they use. The more powerful class of neural networks uses non-linear, e.g., sigmoid activation functions like  $\sigma(x) = 1/(1 + e^{-x})$ .

The neural network operates as follows. It is presented numbers  $x_1, \dots, x_n$  at distinguished input nodes. Each neuron applies its activation function to combine the numbers presented on its incoming edges and propagate the result along its outgoing edges. Specifically, the neuron  $j$  with, say,  $k$  incoming edges from neurons  $i_1, \dots, i_k$  computes the value  $x_j = \sigma(w_{i_1,j}x_{i_1} + \dots + w_{i_k,j}x_{i_k})$ . The numbers produced at the outputs of the network are the result of the computation of the neural network. Data classification can be performed, e.g., by thresholding the single output of a neural network or presenting a separate output for each class and comparing the numbers produced at each of a set of distinguished output nodes.

Neural networks can be distinguished by their topology. *Layered feed-forward* networks, the most popular type, are acyclic and organized in layers of neurons. They always produce outputs. The class of functions they can

compute increases with the number of layers of neurons. In contrast, neural networks with cycles are more complicated. They can become unstable or oscillate or exhibit chaotic behavior.

The training procedure for neural networks optimizes the weights along the arcs of the network in order to optimally reproduce the function to be learned. This function is implicitly represented by the training data. Here we have to take care to avoid overfitting ( $\rightarrow$  *Regularization*). The training problem is hard ( $\rightarrow$  *hidden Markov models*) and the training procedure approaches the problem heuristically, usually by performing some type of local optimization (Russell & Norvig, 1995).

The application of neural feed-forward networks has proved effective in protein 2D structure prediction (see Chapter 6 of Volume I, (Rost et al., 1994)). Other applications of neural networks in bioinformatics include the recognition of protein sequence features such as transmembrane domains (Jacoboni et al., 2001) and recognition of features in DNA sequences (promoters (Matis et al., 1996; Nair et al., 1995), coding regions (Cai & Chen, 1995) etc.) as well as the interpretation of NMR spectra for protein structure resolution (Hare & Prestegard, 1994) and the discrimination between drug-like compounds and those that are not drug-like (Sadowski & Kubinyi, 1998).

A particularly interesting kind of neural network for bioinformatics is the *Kohonen map* (Kohonen, 1990). This is a two-dimensional field of artificial neurons that affords the projection of adjacencies in high-dimensional Euclidean spaces down to two dimensions. Among other things, Kohonen maps are used to classify sets of organic compounds (Anzali et al., 1996; Cai et al., 1998; Kirew et al., 1998), to distinguish between different protein binding sites (Stahl et al., 2000), to investigate protein families (Ferran et al., 1994) or aspects of protein structure (Ding & Dubchak, 2001; Schuchhardt et al., 1996), to analyze the codon usage in different bacterial genomes (Wang et al., 2001) and to cluster gene expression data (Tamayo et al., 1999).

### Regularization

There is a general problem with methods that learn structure from limited amounts of training data, such as  $\rightarrow$  hidden Markov models,  $\rightarrow$  neural networks or  $\rightarrow$  support vector machines mentioned in this glossary. The problem is called *overfitting*. If we carry the training procedures mentioned in the respective sections to their extreme, we will memorize all that we have seen in the training data, including the regularities that generalize to other data for which we want to make predictions in the future, and the irregularities that are specific to the training data and carry no information on data seen in the future. In order to avoid overfitting, the training data have to be supplemented with information that accounts for the lack of our

knowledge on the data yet to be seen. In some sense, the classification function to be learned is smoothed out. It takes the training data into account, but also the lack of information on what is yet to come. Regularization is an involved concept that can be approached with theory (Vapnik, 2000). In practice, regularization is often performed heuristically, sometimes even in an *ad hoc* fashion.

The regularization procedure takes a different form for each method of statistical learning. When training hidden Markov models, the derived probabilities do not only take the observed sequences into account but also use so-called *prior* distributions that formulate some hypothesis on the occurrence of output characters in the case that we have no additional information (such as the observed sequences). An expected background distribution of amino acids or nucleotides is a natural starting point for such a prior distribution.

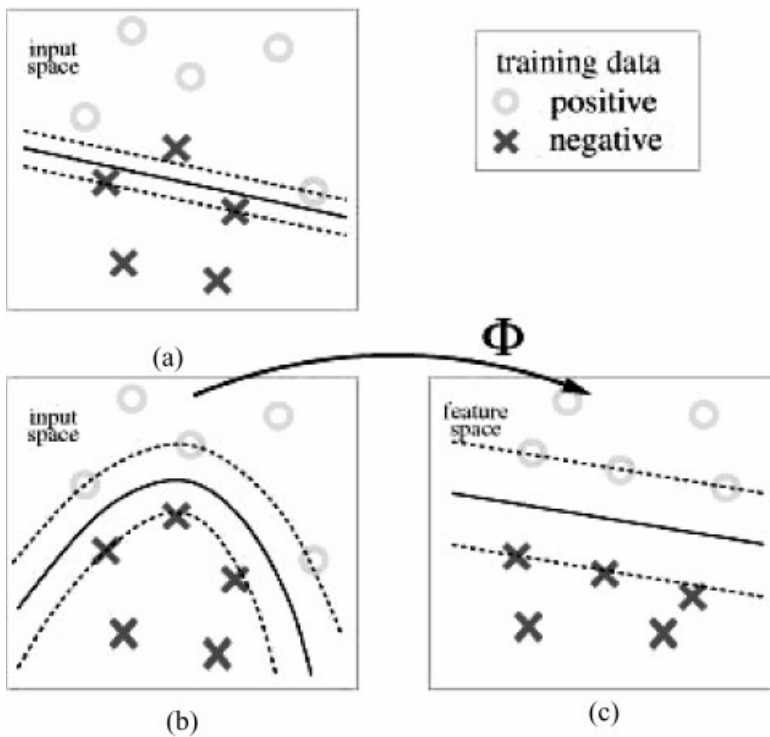
For → neural networks, a common implementation of regularization is to not carry the training procedure to the end but stop sometime before. → Support Vector Machines are automatically regularized by maximizing the margin of separation of the two classes of training data. The amount of regularization can be directly controlled by the parameter that guides the tradeoff between correct classification of the training data and the size of the margin.

### Simulated annealing

Another Markov Chain Monte Carlo (MCMC) algorithm. Here each state of the Markov chain is labeled with a “cost” or “energy”. Moves to neighboring states are accepted if they involve a decrease in cost. If they involve an increase in cost, they are only accepted with a certain probability. This probability is determined by the Metropolis criterion  $e^{-c/kT}$ , where  $c$  is the increase in cost,  $T$  is a *temperature* parameter and  $k$  is the Boltzmann constant (Metropolis et al., 1953).  $T$  starts out at a finite initial temperature and is then decreased according to a so-called *cooling schedule* until it approaches zero. This means that, as the algorithm proceeds, increases in cost become less and less likely. Partly because of its paradigmatic relationship with statistical physics, simulated annealing is a popular version of an MCMC algorithm. The algorithm has the facility of escaping from local minima reached during execution of the algorithm. It is frequently used in global optimization (Kirkpatrick, 1984). Simulated annealing cooling schedules are also very popular with Molecular Dynamics, e.g. for structural refinements using experimental constraints (Brunger et al., 1997) or for generating and refining molecular conformations in homology-based protein modeling (Chapter 5 of Volume I). Simulated annealing approaches for protein-ligand docking are described in Chapter 7 of Volume I.

### Support vector machine

Support vector machines are a technology developed by Vapnik and others for classifying unstructured data (Burges, 1998; Christianini & Shawe-Taylor, 2000) that has gained much attention in bioinformatics lately. Support vector machines classify points in multi-dimensional Euclidean space into two classes. They extend linear classifiers (see Figure 3 a) that classify the points by placing a hyperplane into the space that separates the space into two half spaces. Training the classifier amounts to selecting the hyperplane such that it best fits a subset of preclassified points. This means that the smallest number of preclassified points comes to lie in the wrong half space. In fact, this is not quite the case, since we want to avoid overfitting to the training data ( $\rightarrow$  *Regularization*).



**Fig. 3** Support Vector Machines: (a) linear classifier, (b) nonlinear classifier, (c) nonlinear classifier mapped to a high-dimensional space, where the classification becomes linear. Crosses and circles denote preclassified points on which the support

vector machine is trained. The dotted lines denote the margin between the separating surface and the closest preclassified points. The mapping from (b) to (c) is performed implicitly by a so-called *kernel function*  $\Phi$



Often hyperplanes are not sufficient for the classification of point sets. Support vector machines allow nonlinear separation between the two point classes (Figure 3 b). They do so by mapping the points nonlinearly into a higher-dimensional feature space and performing linear separation there (Figure 3 c). The mapping is done implicitly by a so-called *kernel function*, thereby making very high-dimensional feature spaces computationally affordable. The choice of the kernel function is the essential ingredient in the development of the support vector machine.

Support vector machines receive their name through their second feature. Rather than taking all preclassified points into account, they only consider those points that come to lie close to the separating surface. These points are called the support vectors. The surface is placed such that the distance to the closest support vectors is maximized (modulo preventing overfitting, → Regularization). Here we allow a limited number of misplacements.

Support vector machines have been applied to a wide variety of classification problems in bioinformatics including finding translation initiation sites in DNA sequences (Zien et al., 2000), classifying genes via their promoter regions (Pavlidis et al., 2001a), analyzing gene expression data (Brown et al., 2000; Furey et al., 2000), prediction of gene function (Pavlidis et al., 2001b), determining secondary structures of proteins (Hua & Sun, 2001), protein fold recognition (Ding & Dubchak, 2001), prediction of protein-protein interactions (Bock & Gough, 2001), and protein sorting (Cai et al., 2000).

### **Tabu search**

Meta-heuristic for local search strategies. The goal is to avoid getting caught in cycles while traversing the search space. Tabu search starts out by looking for local minima. To avoid retracing the moves just made the method protocols characteristic features of solutions that were recently visited in one or several tabu lists, each list for a certain class of solution attributes. Tabu search methods vary in the details of the management of the tabu lists (Glover, 1989; Glover, 1990). Tabu search is mentioned in Chapter 7 of Volume I. Additional material on tabu search in docking can be found in (Hou et al., 1999; Westhead et al., 1997).

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## Part II: Applications

# 1

## Integrating and Accessing Molecular Biology Resources

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### 1.1 Introduction

With biological research increasingly focusing on molecular biology and genetics more and more resources are becoming available for this research community to use. These resources include the data which is being collected and stored in databases and applications which operate on these data, producing yet more data. Databases range from simple sequence databases to complex metabolic pathway databases, as well as chemical compound, mutation and literature databases. The value of these databases can only be fully realised when they are integrated so that relationships between data can be found. There are also a vast range of applications available now, from similarity searches and sequence alignment programs to Hidden Markov Models. Results from these applications must also be easily integrated.

The growth of the data available from molecular biology resources has introduced several complexities. The main problem is obtaining a consistent, cross-referenced view of the data. Databases contain widely different data and hence are inevitably presented in a different way. Even databases containing similar data may be formatted totally differently. This leads to relatively simple tasks being quite difficult. For example, searching for similar text in multiple databases becomes an extremely difficult operation. More difficult tasks, such as finding relationships between different entries then become almost impossible. While some databases cross-reference entries to other databases, these explicit references are only a beginning to the many relationships possible among the data.

Further difficulties in integrating resources exist. The quality of the data must be considered, while keeping all the databases and resources updated is a considerable task. With databases and applications changing regularly, a practical way of maintaining and updating data must be found.

In this Chapter, we first examine the resources which are available to the molecular biology community, both databases and applications. We then



discuss approaches to dealing with the problems mentioned above. One resource which attempts to address the above issues is SRS. SRS is now a mature product and is widely used within academic institutes and commercial companies world wide. Our discussion first describes the way SRS tackles the above problems and then compares this approach with the approaches adopted elsewhere.

## 1.2 Molecular biology resources

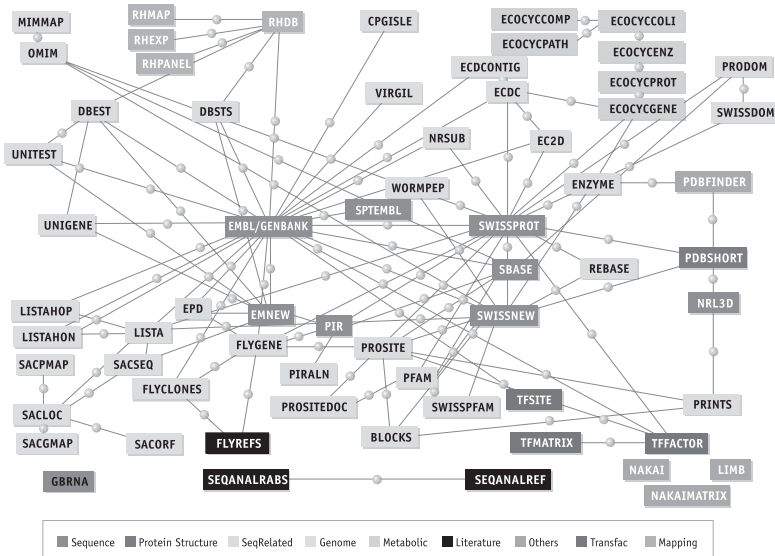
Molecular biology resources include, but are not restricted to, databases and applications. In this section we examine the databases and applications which have given rise to the need for bioinformatics solutions.

### 1.2.1 Databases

As mentioned in the introduction, there is now a considerable number of molecular biology and related databases available. While some are freely available, such as the DNA sequence collection EMBL [30] and GENBANK [18], others are only freely available to the academic community, such as the protein sequence database SWISS-PROT [15], and others are only available on subscription, such as the EST databases available from the company Incyte Genomics. Academics and pharmaceutical companies also have their own proprietary data which must be integrated into a system so that relationships with publicly available data can be found.

Molecular biology databases can be divided into groups, such as the sequences databases, which have actual DNA and protein sequence, sequence related databases, such as protein structure databases, whole species databases, metabolism databases and so forth. Figure 1.1 shows a representation of the main publicly available databases. The databases are represented as a network, with lines drawn between databases where a link has been defined between the databases. From this network it is possible to plot a path from a piece of data in one database, through to a related piece of data in another database. This link may sometimes be through the result of an application, which means that an application uses data from one database and produces some results which are included in the application result database and then linked to another database.

It is also necessary at this point to discuss another problem, that of the maintenance of databases. Some databases are maintained and annotated by humans, while other databases are generated. Errors in the humanly annotated databases, databases not being updated and badly maintained



**Fig. 1.1**  
The SRS universe at the EBI.

databases are just some of the problems which need to be addressed when it comes to the integration of these resources.

### 1.2.1.1 Sequence databases

The first databases to appear were DNA sequence databases, namely those from the EMBL (Europe), NCBI (USA) and the DDBJ (Japan), known as EMBL [30], GENBANK [18] and DDBJ [1] respectively. These are DNA databases of sequences and their annotations. These databases continue as a collaborative effort, with the three databases sharing their information. So all three databases contain identical data, albeit in a different format.

The EMBL database is a model nucleotide database to consider. Each entry in the database is given a unique ID and information about the entry is presented in a semi-structured way. The actual nucleotide sequence is presented at the end of the entry.

These databases contain genomes, genes, ESTs, variations, introns, exons – all pieces of DNA which have been sequenced and submitted by scientists around the world to the maintainers of the databases.

Each entry in the databases contains keywords and descriptions, which describe the function of the piece of data. Any publication on the sequence and the authors names also form part of the entry. However, the most interesting information is in the annotated feature tables. These contain

information on what part of the sequence are coding sequences, what they code for, which regions of the sequence are introns or exons, and any known function of other parts of the sequence.

Protein databases used to be from direct protein sequencing, but now they are made almost exclusively from the translation of ORFs (Open Reading Frames on DNA sequences). Both the European Bioinformatics Institute (EBI) and National Center for Biotechnology Information (NCBI) provide databases, TREMBL [15] and GENPEPT [19] respectively, which are automatic translations from the CDS features of the DNA in their nucleotide databases. This includes some automated annotation of the role which the protein plays.

SWISS-PROT [15] is a protein database which includes annotation by humans, as well as automated annotation. SWISS-PROT also includes links to many more databases than in other protein databases, and hence is a rich source of finding links between data.

#### 1.2.1.2 Sequence-related databases

The sequence related databases contain further information about the sequences in the sequence database. Often the information is derived computationally from the sequence data, although experimentally defined databases often contain more detailed information. Entries in sequence related databases almost always provide an explicit link to a sequence database.

One of the best known protein structure databases is the Protein Data Bank (PDB) [23]. PDB archives experimentally determined three-dimensional structures of biological macromolecules and contains atomic coordinates, bibliographic citations, primary and secondary structure information, as well as crystallographic structure factors and NMR experimental data

DSSP [24] and HSSP [17] are databanks of secondary structure, based on proteins in the PDB. HSSP contains homology-derived structures of proteins, with each protein in PDB providing the basis for including sequence homologies and then aligning those sequences with the sequence of known structure from PDB. This provides implied secondary and tertiary structures for many more proteins than could be found experimentally.

Taxonomy [19] is a database which contains the names of all organisms that are represented in the GENBANK with at least one nucleotide or protein sequence. The taxonomy database is organised as a tree, so the link from species to subspecies can be followed. Although taxonomy contains about 30 000 species represented in GENBANK, there is no comprehensive databank with all approximately 2 million known species. Other species data are available from projects such as Biodiversity and Species 2000.

There are also several protein family databases. Prosite [14] and Blocks [22] provide motifs or patterns for protein families, so that new proteins can be examined to see if they fit into a particular protein family. PFAM [13] and

DOMO [21] contain protein families and their alignments, so that changes in the proteins can be observed. Sequences also share domains, or modules, and the ProDom [20] and Smart [2] databases contain these domains and the proteins which fit those domains.

#### 1.2.1.3 Genome databases

The first genome database was the GDB (Genome Databank) which contained all known human loci. With the start of the human genome project this database was discontinued, but is now once again being updated. Currently there are several whole genome databases recently completed or nearing completion. These include many microbial genomes and several eukaryotic (including yeast and *C. elegans*), available at the Sanger Centre in the UK and from The Institute for Genomic Research (TIGR) in the USA. The Human Genome Mapping Project (HGMP) is an inter-governmental project started in 1995 to sequence the whole human genome. In June 2000 the privately listed company, Celera Genomics, which published the *Drosophila* Genome in April 2000 [27], together with the HGMP, announced a draft human sequence.

#### 1.2.1.4 Gene expression

Gene expression databases can be made from two sources, ESTs (Expressed Sequence Tags) or DNA chip expression data.

ESTs are short (200–500bp) DNA sequences generated from the 3' and 5' ends of randomly selected cDNA clones. Quite a large proportion of the sequences in EMBL and GENBANK are actually ESTs. Other proprietary data is available from a private company, Incyte Genomics.

There are currently no publicly available DNA chip expression databases, however the Microarray Gene Expression Database Group (MGED) [3] at the EBI aims to facilitate the creation of a gene expression data repository.

#### 1.2.1.5 Mutations and SNPs

Mutations occur at a DNA level and can give rise to a different protein product. This can be the cause of an inherited disease, such as breast cancer or cystic fibrosis) or indicate a condition.

Currently OMIM [19], from the NCBI, is the most comprehensive database of genetic diseases and their associated mutations. However there are many more locus-specific databases, such as the mutation databases for p53 and Cystic Fibrosis.

There are three types of mutations (Figure 1.2). Exchange, where a nucleotide is exchanged for another, insertion, where an extra nucleic acid is inserted into the sequence, or deletion, where a nucleic acid is deleted.

Exchange	Insertion	Deletion
ACTGTTACT ACAGTTACT	ACTGTTACT ACA TACT	ACT TACT ACAGTTACT

**Fig. 1.2**  
Three types of mutations.

There are, of course, several different ways of representing these mutations and creating a uniform way of searching for a specific mutation across dozens of mutation databases is a challenge. The work of Lehvaslaiho et al. [26] at the EBI on creating a unified access to mutation databases attempts to solve the problem of a lack of uniformity across the mutation databases.

SNPs (Single Nucleotide Polymorphisms) are the most frequent type of variation in humans. A SNP is defined as:

SNPs are single base pair positions in genomic DNA at which different sequence alternatives (alleles) exist in normal individuals in some population(s), wherein the least frequent allele has an abundance of 1% or greater. In practice, the term SNP is typically used more loosely than above. Single base variants in cDNAs (cSNPs) are usually classed as SNPs since most of these will reflect underlying genomic DNA variants. SNP datasets also typically contain variants of less than 1% allele frequency. The 'some population' component of the definition is limited by practical challenges of surveying representative global population samples. [4]

There are several SNP databases, such as MITSNP [31] and one from the SNP consortium, which contain the SNP and its associated effect.

Databanks such as SWISS-PROT and EMBL are also including variation information in their annotations.

#### 1.2.1.6 Pathway and chemical compounds

Databases such as Enzyme [12], Brenda [5], Ligand [28] and EMP [6] contain enzymes which are involved in a large number of reactions. Each enzyme with known enzyme function are catalogued and named by a nomenclature committee. Also included in the databases is information on the reaction and specificity of the enzyme and the various conditions the enzyme will be active under. Information about the structure and stability of the enzyme and the preparation of the reaction may also be included.

Reaction databases, such as MPW (or WIT) [7], KEGG [8] and EcoCyc [9] contain specific information about the pathways of a reaction, giving input and products as well as enzyme information. Links to various protein and chemical databanks are included in these databases.

Publicly available chemical compound databases, consisting of ligands which may include related information to sequence databases, are relatively small, and include Ligand [28] and Clotho (US). Proprietary databases, such as CAS, Beilstein and others, are much larger but are very expensive.

### 1.2.1.7 Transcription factors

There are several transcription factor databases. Transfac [32] and TFD [10] provide databases of transcription factors, their binding sites and the genes which regulate these transcriptions.

### 1.2.1.8 Others

Some of the most important data is not in databases, but in the literature. Medline (published by the National Library of Medicine) and Excerpta Medica (published by Elsevier) are two databases which provide abstracts of published papers. It is increasingly important to link these resources with the scientific databases listed above.

## 1.2.2

### Applications

Bioinformatics applications have for the most part revolved around similarity search engines such as BLAST, FASTA [29], and CLUSTALW.

BLAST and FASTA, and their derivatives, take a sequence as input and compare that sequence to a database of sequences (the search database). A list of sequences which are similar to the query sequence is then produced with a score of how well the sequences match. The search database is usually composed of all the sequences in a database such as SWISS-PROT or GENBANK, and hence the matched sequence in the search database contains information which may be useful in annotating the query sequence.

CLUSTALW is a sequence alignment tool, which takes as input a set of sequences and produces a multiple alignment of those sequences.

The results from these applications themselves form a database which contain links to other databases (namely the search databases). This is a rich source of finding relationships between sequences which a user may have and the public databases.

## 1.2.3

### Databases and applications world-wide

Catalogues of databases are limited to LiMB, which unfortunately is no longer maintained and out of data and DBCAT [11], from Infobiogen in France, which has information on 400 databases relevant to biology. The first issue of Nucleic Acids Research each year also provides a list of currently available databases.

However, the growth in the number databases implies that it is not always easy to find available resources. The Databank of Databanks [25], part of the SRS server at the EBI, provides a place to not only search for

Information on all databanks offered by each of the listed servers is regularly compiled into [Databanks](#)

**list** all representative entries of **Databanks**

**Full Search** **Databanks**

**Quick Search**

Search in:

for

**SRS Servers World Wide**

[WEHI, Melbourne, Australia](#)

[Vienna BioCenter EMBnet Node, Vienna, Austria](#)

[Belgian EMBnet Node \(BEN\), Brussels, Belgium](#)

[DBBM-IOC, Fiocruz, Rio de Janeiro, Brazil](#)

[CBR-NRC, Halifax, Canada](#)

[The Genome Mine, Bazed Bioinformatics, Canada](#)

[CBI EMBnet Node, University of Beijing, China](#)

[CSC, Otaaniemi, Espoo, Finland](#)

[DJFCBIOGEN, Villejuif, France](#)

[Institut Pasteur, Paris, France](#)

[LBMRPMMINRA-CINES, Auzeville, Toulouse, France](#)

[DKFZ, Heidelberg, Germany](#)

[EMBL, Heidelberg, Germany](#)

[GBF, Braunschweig, Germany](#)

[MIPS - MPG/GSF, Martinsried/Munich, Germany](#)

[Bioinformatics Centre, University of Pune, India](#)

[INCB EMBnet Node, Dublin, Ireland](#)

[Weizmann Institute BCD, Rehovot, Israel](#)

[CNR EMBnet Node, Bari, Italy](#)

[CRISCEB, Second University of Naples, Italy](#)

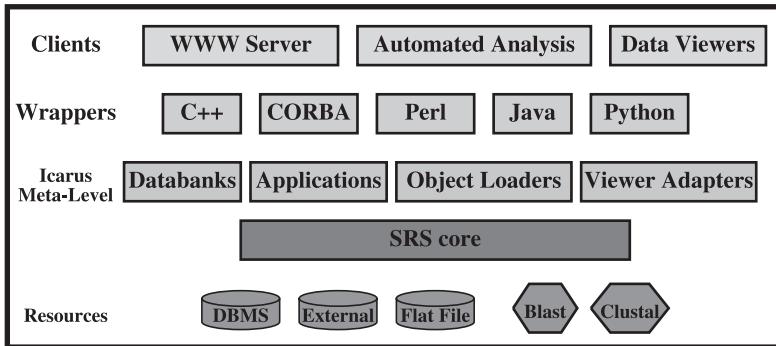
**Fig. 1.3**  
The SRS databank of databanks.

available databases but also to find the most recently updated version of a particular database.

Figure 1.3 shows a partial list of the publicly available SRS servers taking part in the Databank of Databanks project. Each night each SRS Server retrieves information about all the other servers, such as, the databases and applications which that server has. This is then combined to make a searchable database. It is then possible to search for servers which have a particular database (such as all databases which have “sequence and align” in their description, or the server with the most up to date version of a particular database).

### 1.3 An overview of SRS

SRS attempts to meet all the challenges listed in the introduction to this chapter. Molecular biology resources – databases and applications – can be quickly and easily incorporated into the system. Using the meta definitions of the format and syntax of the resources, SRS creates indices of the data (both databases and the output of applications) which can then be quickly and easily queried. Once a query is performed against one or more databases, either the whole database entry or just the information which the user requires can be extracted from the databases and displayed. Links (both explicit and implicit) between those data and other databases can be made and hence navigation between resources is possible.



**Fig. 1.4**  
The SRS system.

SRS consists of several layers, as shown in Figure 1.4. Central to SRS are the twin philosophies of:

- (1) leaving data in their original context
- (2) the use of meta-description to describe the data.

To achieve this SRS utilizes the special purpose programming language, ICARUS (Interpreter of Commands and RecUrsive Syntax), which is used both for the meta definition of biological resources and the syntactic description of the data in the resources. The SRS core includes programs which use ICARUS meta definitions and syntactic descriptions to index and retrieve information from resources. In the case of applications, the meta definition includes information on how to run the programs as well as the meta description of the syntax and structure of the output.

To index and retrieve data, SRS uses a *token server*. The meta definition is used to extract *tokens*, i.e. strings, from the database for indexing. When a query is performed, SRS searches for the query term against these indices and creates a set of entries which match that query. The fields of the entry which the user wants displayed are then extracted from the entry. This is done by using the connection made in the meta definition between a field and *tokens* received from the *token server*.

We now go onto describing the layers of the overall system, as shown in Figure 1.4, in more detail.

### 1.3.1

#### The meta-definition layer

Meta definition can be defined as a higher level definition and is widely used in SRS. Meta definition is used to describe both the resources to be



integrated *and* the format (or syntax) of the data in those resources. The data extracted using this syntax can then either be viewed or passed to another program. It also has several ancillary uses within SRS, such as describing the HTML pages which are generated from the web interface.

Meta definition has many advantages, foremost being that changes to the format of input and/or outputs only require minimal changes to the meta definition, and none to any core component of SRS.

In SRS, meta definition is used to describe objects which the SRS core uses. In the case of a database, a library object must be defined. This object contains the name of the library, what sort of library it is (i.e. what group of databanks it belongs to), the name and whereabouts of the flat files containing the data. It also contains a link to a file containing a list of rules which describe the internal syntax of the databank. These syntax rules will be described below.

From the meta definition layer, it is also possible to use alternative viewer adaptors. The view adaptor is given a meta definition, which describes the input that the viewer requires. This is currently used to allow graphical Java applets to use information from SRS for data visualization.

### 1.3.2

#### **The SRS core**

The SRS core consists of several programs which use the meta definition layer to allow the resources to be queried, data from these resources to be returned and, in the case of applications, other programs run on selected data. This includes the creation of indices from the data resources. It is the creation of these indices which makes the querying of the databases quick.

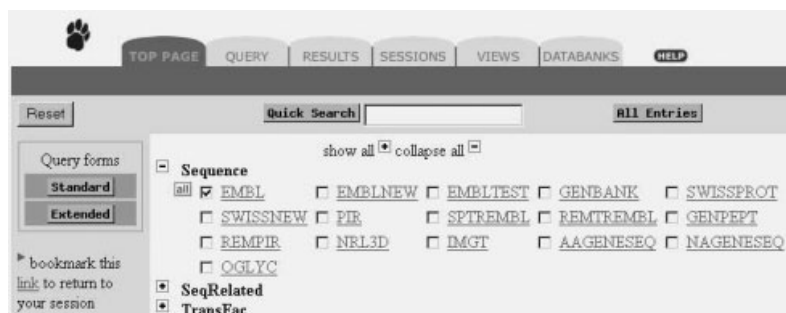
To create the indices, SRS uses the rules given in the syntax description to create *tokens*. In this case, a *token* is defined as a piece of text from the database entry. These tokens are stored in token tables which are linked to a particular field in the database by a meta definition in the library meta definition object.

These rules are also used to extract data from the database. An object loader, another object in the SRS system, can be defined in the meta definition layer and describes the way a token table relates to a field to be extracted and viewed.

### 1.3.3

#### **Wrappers**

Another advantage of the meta-definition approach which SRS has taken is that wrappers can be generated to provide a programming interface to SRS. This allows programmers to be able to link in the SRS system to their own



**Fig. 1.5**  
The library select page of the SRS web client.

programs (C++, Perl, Java or Python) or for a connection to a remote SRS system via the CORBA (Common Object Request Broker Architecture) server.

The wrappers provide full access to the SRS system. Queries can be done and data from the databases returned. It is also possible to get information about the resources and to run applications.

### 1.3.4

#### Clients

The top layer of the SRS system is where biologists come to retrieve data from the system and there are a number of ways in which SRS can be used. The most popular client is the web client available at numerous academic sites around the world or from a local installation (Figure 1.5).

Other clients could be data viewers, which extract information from the databases and present it graphically, or other programs which perform automated analysis may use SRS to extract data, perform analysis and write the results to another file.

## 1.4

### Integrating molecular biology resources

With the aim of integrating databases to offer sensible queries where only the pertinent information is retrieved, the structure of data is important.

Databases can be thought of as having three levels. All databases contain entries, which are made up of several fields within the entry and then each field has a micro-structure. For the purposes of both querying, and retrieving information from, a particular field, the micro-structure, or syntax, of the field must be known.

```

ID  AB000360 standard; DNA; HUM; 2582 BP.
AC  AB000360;
DE  Homo sapiens PIGC gene, complete cds.
KW  glycosylphosphatidylinositol-synthesis gene; PIGC.
SQ  Sequence 2582 BP; 694 A; 494 C; 581 G; 813 T; 0 other;
    ggatccctgc tgcagagggg gtaacgggtg ctggcttgcc tgtttgtgtc 50

```

Fig. 1.6

Sample piece of a typical database entry.

In this section we describe the relationship between the meta description of the library object, and other objects, and the syntax description of the fields.

#### 1.4.1

##### The SRS token server

For each library a syntax description of the data must be supplied. This syntax description must describe the data in the database and takes the form of a set of *rules*, also called a *grammar*. This list of rules is written in the SRS programming language ICARUS. The rules in this case provide the syntax part of the ICARUS language (the other part being Meta Definition) and are called *productions*. Productions are used to extract pieces of text from a database entry, *tokens* which are written to *token tables*. It is the data which is in these token tables that is indexed and used when querying the database, or extracting parts of the database for display.

All flat file databases are semi-structured, containing a list of entries, with each entry containing a list of data-fields (e.g. an Id, an Accession number, key words, a sequence etc.). Figure 1.6 shows a sample of a database entry. Each of the data-fields consists of *strings* or *tokens*. The set of productions for each database must describe how to divide the database into entries and then further into fields and then into the strings or *tokens* within that data field. It is these tokens within each field which are inserted into an index.

Figure 1.7 shows a subset of the rules for describing an entry in the EMBL database. The \$Wrt command is the instruction to write a string into a token table.

#### 1.4.2

##### Meta definition of molecular biology resources

The meta definition which describes the structure of the resource is also written using ICARUS. For the EMBL database this meta definition consists of the definition of several objects.

```

entry:      ~ id_line
            ac_line de_line kw_line sq_line end_line ~
id_line:    ~ 'ID' id ~
ac_line:    ~ 'AC' ac ~
kw_line:    ~ 'KW' (word)* ~
sq_line:    ~ 'SQ' sequence ~
end_line:   ~ '//' ~
id:         ~ word $Wrt ~
kw:         ~ (word $Wrt)* ~
            ..
word:       ~ /[a-zA-Z0-9-]*/ ~

```

Fig. 1.7

Subset of rules describing an entry in the EMBL database.

```

$EMBL_DB=$Library:[EMBL group:$SEQUENCE_LIBS
                  format:$EMBL_FORMAT
                  searchName: '*.dat'

```

Fig. 1.8

The EMBL library object.

The EMBL library object, (Figure 1.8), defines the EMBL library as being part of the sequence library group, with the format being described in the \$EMBL\_FORMAT object (Figure 1.9) and the files which make up the EMBL database being all those files with the extension 'dat' in the EMBL flat file directory.

The EMBL FORMAT object contains a list of fields which make up the database. In our example EMBL database the library has 5 fields.

The field information is related to a particular token table via the index-Token attribute in the \$Field object. Hence, when a particular piece of information from the description field of the EMBL databank is requested, SRS looks in the description field meta definition for the EMBL database

```

$EMBL_FORMAT=$LibFormat:[syntax:$EMBL_SYNTAX contains:$DNASEQ_DATA
                        fileType:{ $DAT_FILE $SEQ_FILE}
                        fields:{
                            $Field:[ $DF_ID index:id indexToken:id]
                            $Field:[ $DF_Accession index:str indexToken:acc]
                            $Field:[ $DF_Keywords index:str indexToken:key]
                            $Field:[ $DF_Description index:str indexToken:des]
                            $Field:[ $DF_DNASequence format:embl]
                        }
]

```

Fig. 1.9

The EMBL FORMAT object.

for the name of the token table which contains that information. The SRS token server then requests that that token table be built and the tokens, or strings, from the databank can be returned.

#### 1.4.3

##### **Indexing databases**

In the case of indexing databases, the token tables for each field which is to be indexed are constructed. This means that for each field defined in the \$LibFormat object, a token table is generated for that field based on the indexToken attribute in the meta definition. The values in the token table for each entry are then written to indices which are sorted and can be searched quickly.

As well as the usual indices, link indices can be built which allow for navigation between databases or cross-database queries. Two fields from different databanks must be specified as having a link between them.

#### 1.4.4

##### **Querying and linking databases**

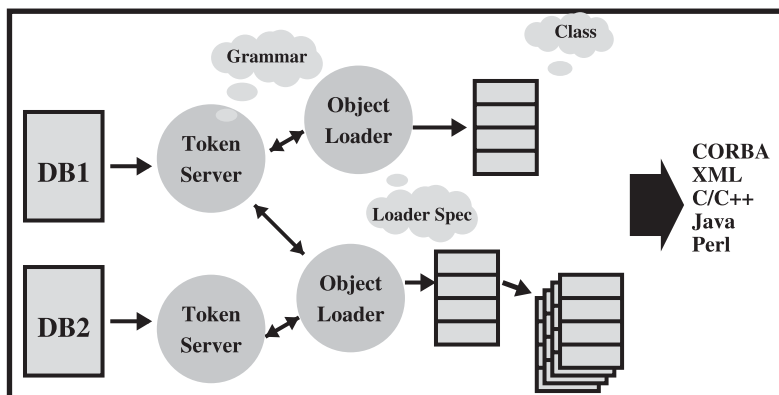
Queries are performed on databases based on fields. SRS provides a full query language for interrogating both strings and numeric fields and combining searches across multiple libraries and fields. SRS returns a *set* as the result of a query. Sets can be combined with the AND ('&'), OR ('|') and BUT NOT ('!') operators, which allow for the union, intersection and “and not” combination of two sets. In addition, two link operators are provided. These allow for 2 sets to be linked together and the returning set to either be the set of entries in set 2 which link to set 1 or the set of entries in set 1 which link to set 2.

#### 1.4.5

##### **Views and object loader**

The result of a query or link operation in SRS is a set of entries. This is just a collection of entry-IDs and does not contain any information from the entry in the databank. Views provide a way of viewing part of the information from a databank for those entries in the set. Moreover, related information from a different databanks can be included in the view, thus creating a virtual entry.

Object loaders [16] provide a new and more flexible way of obtaining the data for those entries in the set from the databases. Instead of retrieving a set of strings or text, the object loader retrieves an object containing the re-



**Fig. 1.10**  
Object loaders.

requested information. Views within SRS now use object loaders to obtain the data before printing it out in the requested way.

This provides much more flexibility, as the data objects returned by the object loader can be passed to other programs, such as those written in C, C++, Java Perl or Python, passed over the Internet via a CORBA server or used to create more interesting views, such as a view including an image made from data in the object (an example of this can be seen in Figure 1.11 further in this Chapter).

Figure 1.10 shows how objects containing information from multiple databases can be constructed. From the databases, DB1 and DB2, the token server can retrieve information via the grammars describing those databases as described above. An object loader can then be defined which relates an attribute in the object to a specific token table from the token server. The object loader can then take this meta-defined object and load information into it from the databases for each entry in the set.

#### 1.4.6

#### **Applications – analyzing data**

As mentioned in Section 1.2 describing the various Molecular Biology resources available, applications have played a large part in developing these resources. In fact some databases are entirely based on the results of applications.

The output from these applications themselves forms a database and often contains information related to entries in other databases. For example, if a BLAST run is performed against the SWISS-PROT database then the output will contain SWISS-PROT accession numbers and parts of se-

quences in the SWISS-PROT database. In SRS this output is parsed and indexed, so that it forms part of the SRS universe and can be used to link data in different databases.

## 1.5

### The SRS data warehouse

Keeping all relevant databases up to date, both internal and external, can be a difficult task. Databases can be updated irregularly and updating them is a routine but cumbersome task. Once the databases have been down-loaded from their repository, certain tasks must be performed. These include possibly reformatting the data, combining with other data to make new databases, indexing the databases.

While all this sounds routine, there are many things which can go wrong and there are many interdependent tasks. If, for example, one particular FTP site is down, then the downloading of the database may not occur. If the indexing of the database goes ahead anyway, then that database will be out of date and databases with links to that database may also contain incorrect information.

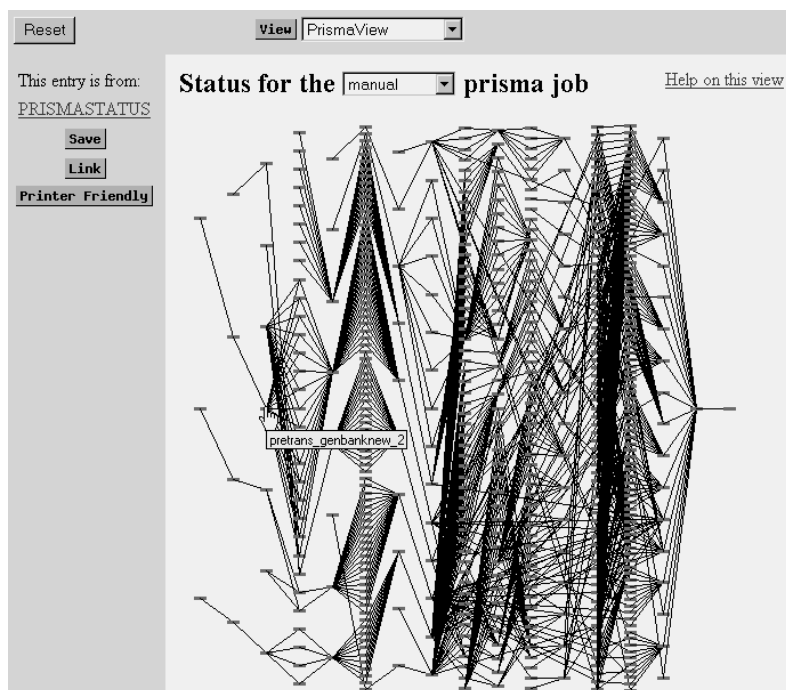
A new module of SRS, Prisma, automates this process. Prisma should not just do the down-loading, reformatting and re-indexing tasks normally associated with such a tool, but also the report the progress of the process. Figure 1.11 shows the status of the updating of the databases after a particular run. Each task in the process (such as downloading a file, reformatting a database, indexing the database etc.) is represented by a box and dependent tasks are depicted by a connecting line. This view allows for quick checking of the updating process with any failures, and their effect immediately apparent (if a process fails for some reason the box representing it will become red, while processes waiting on another task are shown in purple).

This extension to SRS has proved very successful at keeping data at both public sites, such as the EBI, as well as private companies up to date.

## 1.6

### Accessing integrated data

Integrating the data into a single system is only part of the problem. The interface presented to the scientist must be easy to use and provide several powerful features to enable easy manipulation and viewing of the data. It must also be possible to access the data and use it easily from other programming environments, such as other languages or across a network.



**Fig. 1.11**  
Prisma view.

### 1.6.1

#### The web interface

SRS provides an extremely powerful, but relatively simple, web interface to the databases mentioned above. It is possible to select a database that one wants to search and then retrieve the whole database, or search one or more fields within the database. Figure 1.12 shows the standard query page for the SWISSPROT databases. Users can either search for their search term in the ID, keyword, or other fields of the database, or search all text fields for their search term.

Once a set of entries which match the search terms has been retrieved, various operations can be performed on that set. The simplest is to view each whole entry in it, or to just view various fields from the entries. Pre-defined views, either part of the SRS package, or defined by the local SRS administrator, allow for the data to be presented in other formats. Figure 1.13 shows the default entry view of a whole SWISS-PROT entry. This looks considerably different to the text in the flat file, as shown in Figure 1.14, with the information easier to read. Either a pre-defined view of the entries can be used to view just some of the fields, or alternatively a view of the data



TOP PAGE QUERY RESULTS SESSIONS VIEWS DATABANKS HELP

Reset search SWISSPROT Info about field AllText

Submit Query

append wildcards to words

combine searches with AND

display entries in chunks of 30

Extended query form

separate multiple values by & (and), | (or), ! (and not)

AllText AllText AllText AllText

retrieve entries of type Entry

Use predefined view SequenceSimple

Create your own view

Display in  List  Table

sequence format swiss

ID  
AccNumber  
Description  
GeneName  
Keywords  
Date  
Organism

Fig. 1.12 Querying the SWISS-PROT database.

Test Entry | SwisEntry

Reset

This entry is from: SWISS-PROT

Save Link

Launch BlastP

Printer Friendly

General Description References Comments Links Featured Sequence

**General Information about the Entry**

Entry name SWISSPROT:CD22\_HUMAN

Prim. accession # P20273

Sec. accession # Q01665, Q92872, O95699, O95701, O95702, O95703,

Created Release 17, 1-FEB-1991

Last sequence update Release 38, 15-JUL-1999

Last annotation update Release 38, 15-JUL-1999

**Description and Origin of the Protein**

Keywords Glycoprotein, Cell adhesion, Transmembrane, Signal, B-cell, Immunoglobulin domain, Alternative splicing, Phosphorylation, Polymorphism,

Description b-cell receptor c422 precursor (Iu-14) (b-lymphocyte cell adhesion molecule) (bl-cam)

Gene name(s) c422

Organism source homo sapiens (human)

PIR A35648, A35648, JH0371, JH0371

MIM 107266, -

PFAM PF00047, ig, 5

**Features**

VARSPPLIC: 241 417, MISSING (IN CD22-ALPHA)

Key	Begin	End	Length	Description
signal	1	19	19	potential
chain	20	847	828	b-cell receptor c422
domain	20	687	668	extracellular (p-potential)

Fig. 1.13 The Entry Page for a SWISS-PROT Entry.

```

ID CD22_HUMAN STANDARD; PRT; 847 AA.
AC P20273; Q01665; Q92872; O95699; O95701; O95702; O95703;
DT 01-FEB-1991 (Rel. 17, Created)
DT 15-JUL-1999 (Rel. 38, Last sequence update)
DT 15-JUL-1999 (Rel. 38, Last annotation update)
DE B-CELL RECEPTOR CD22 PRECURSOR (LEU-14) (B-LYMPHOCYTE CELL
ADHESION
DE MOLECULE) (BL-CAM).
GN CD22.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata;
Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
RN [1]
RP SEQUENCE FROM N.A. (CD22-BETA).
RC TISSUE=TONSIL;
RX MEDLINE; 91086838.

```

**Fig. 1.14**

A part of the equivalent SWISS-PROT text entry.

can be built and used to view the data. The data can also be down-loaded onto the user's local disk.

From the set of entries, it is also possible to link to other databases, and find related entries, enabling navigation across all the available databases. It is also possible to run applications, such as BLAST and FASTA, on retrieved data. The results are indexed and immediately available for querying.

The web interface provides an easy way for scientists to access the SRS functionality. Examples of the SRS web interface can be seen at [srs.ebi.ac.uk](http://srs.ebi.ac.uk) or [www.celera.com/celeraScience](http://www.celera.com/celeraScience) and other public web sites.

## 1.6.2

### The application programmers' interfaces (APIs)

The API interfaces to SRS provide an object oriented programmers interface into an SRS system. This allows programmers to access the databases in their SRS system from their programs written in C++, Perl, Java or Python. The databases can be queried and information retrieved about the set of entries. The SRS system itself can be interrogated about the databases available and the fields which can be queried within those databases.

Below (Figure 1.15) is a perl script for searching the SWISS-PROT database for the entry with ID "C931\_SOYBN". A set of entries is returned, this time with just one entry in it. The entry can be retrieved using the *getEntry* method of the *set* class, and the *fieldToString* method of the *entry* class is

```

use lib $ENVSRSEXE;
use srsperl;
$sess = new Session;
$set = $sess->query("[swissprot-id:C931.SOYBN]", "");
$numEntries = $set->size();
print "The query produced ", $numEntries, " entries";
$e = $set->getEntry(0);
print $e->fieldToString("", "", "", "");
print "Now print just the Sequence field";
print $e->fieldToString("Sequence", "", "", "");

```

**Fig. 1.15**

An example Perl program performing a query.

used to print the whole entry and just the sequence field respectively. Alternatively loaders can be used to retrieve an object with the fields defined in the loader available to the programmer.

### 1.6.3

#### **Other interfaces**

In addition to the APIs and the web interface, there are other ways of accessing the SRS system. Possibly the most powerful is by using the ICARUS programming language itself. This provides an interface for performing queries and running applications similar to the APIs but also allows for parsing and easier definition of object loaders etc. Another interface to SRS is via the unix command line and the “getz” program, which allows for querying of databases and extraction of fields.

### 1.7

#### **Other approaches**

Other attempts to solve the problem of integrating Molecular Biology resources can be divided into two possible approaches, either using relational databases to store and retrieve data or to use database specific programs to parse flat files.

The National Center for Biotechnology Information (NCBI) presents their series of databases, GENBANK, GENPEPT, etc. on the web with a simple query form. The query form allows users to think about the data differently to how SRS works, with users searching for a DNA or protein sequence rather than searching a specific database. This allows scientists and occasional users to ask simple questions very quickly. However it is not possible

to search a specific field in a database or for the user to build their own view of the data.

The NCBI does use a flat file approach to parse and retrieve the data in their databases and present it on the web. While the NCBI continues to add databases, there are not as many databases available as some SRS servers, and hence it is difficult to find relationships that may exist between the data displayed and data in other databases. Since the NCBI present their data on their web site it is also not possible for other academic institutions or companies to bring the software in-house for integration of their proprietary data.

The use of Relational Database Management Systems (RDBMS) to store genomic data is particularly wide spread in pharmaceutical companies where security is of significant importance. While RDBMS' provides great security and maintenance of databases they can be very slow when querying the data, especially when asked novel questions linking data in many different tables. With finding links between data in different databases becoming more important, queries can take a very long time. Scalability is also a big problem as the number of databases increases. However with pharmaceutical companies wanting to keep their data in a RDBMS and still look for links in the public databases, it will become imperative for a tool such as SRS to allow the integration of RDBMS and public data into the one system.

## 1.8

### Conclusions

In this Chapter, we have detailed the Molecular Biology resources available and the way in which this data may be integrated to allow scientists to ask novel questions and investigate the relationship between data in different databases.

While there are many and varied approaches to this problem, the approach which SRS uses allows new databases to be quickly integrated into the system. Users then have the flexibility to follow a path from data in one database to data in another database and then to view the data in both which may interest them.

### References

- 1 <http://www.ddbj.nig.ac.jp/>.
- 2 <http://www.smart.embl-heidelberg.de/>.
- 3 <http://hgbase.cgr.ki.se/hgbase/snpinfo.htm>.
- 4 <http://www.ebi.ac.uk/microarray/MGED/>.
- 5 <http://www.brenda.uni-koeln.de/>.

- 6 <http://www.biobase.com/EMP/>.
- 7 <http://www.cme.msu.edu/WIT/>.
- 8 <http://www.genome.ad.jp/kegg/>.
- 9 <http://ecocyc.PangeaSystems.com/ecocyc/>.
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## 2 Bioinformatics Support of Genome Sequencing Projects

*Xiaoqiu Huang*

### 2.1 Introduction

The genome of an organism is the “book of life”. It encodes the complete set of genetic instructions for the development of the organism. The structure of a genome is a linear sequence of nucleotides. Determination of the sequence of a genome lays the foundation for understanding biology at the molecular level. With the current biotechnology, it is a challenging task to determine the sequence of a genome. A sequencing machine can read the sequence of a piece of DNA for up to 1000 bp (base pairs). However, genomes are very huge. For example, the genome of the bacterium *E. coli* is about 4 Mb (million base pairs) in size, the genome of the nematode *C. elegans* is 100 Mb in size, and the human genome is 3 Gb in size. The inability to produce long sequences by sequencing machines requires that long sequences be produced from short sequence reads. A shotgun sequencing strategy is widely used to determine the sequence of a long segment of DNA. In this strategy, multiple copies of the DNA segment are randomly cut into small pieces. The sequence of each piece is read by an automated sequencing machine. The sequence of the large DNA segment is reconstructed by a computer program from short sequence reads. The sequence assembly problem is to assemble short reads into long sequences. What makes the sequence assembly problem non-trivial is that there is no information about how short sequence reads are ordered with respect to the DNA segment.

In addition to the huge size of the genome, there are a number of difficulties with solving the sequence assembly problem. First, the genome contains repetitive regions (*repeats*), different parts of the genome that are highly similar in sequence. Methods for reconstructing the sequence of a DNA segment from short reads are based on finding overlaps between reads and using overlaps to guide assembly of reads into long sequences. Two reads have an *overlap* if a region of one read is highly similar to a region of the other read. If two reads come from overlapping parts of the DNA segment, then they have a true overlap. On the other hand, if two reads

come from two repetitive regions, then they have a false overlap. Repeats in the DNA segment lead to false overlaps between reads, which cause sequence assembly methods to make mistakes. Large genomes such as mammalian genomes contain more repeats than small genomes such as microbial genomes.

Second, the orientation of a read is unknown. The DNA segment consists of two complementary strands. It is not known whether a read is from the plus strand or the minus strand of the DNA segment. Third, sequencing machines make errors in reading DNA sequences. A good portion of a read contains errors at a rate of 1% to 5%. Much more errors occur at the ends of a read. Fourth, a read sometimes contains a contaminant at its left end, the sequence of a bit of foreign DNA, which is not from the DNA segment to be sequenced. This bit of foreign DNA comes from a cloning vector or other sources. Fifth, a read occasionally consists of two pieces which are from different parts of the DNA segment. This read is called a *chimeric read*. Sixth, the genomes of different individuals are not identical. For example, it is estimated that the genomes of two different persons differ at a rate of 1 in 1000 bp. This kind of differences is called *polymorphism*. Regions of a genome with a high rate of polymorphism pose a problem to assembly methods.

Two strategies are currently used in genome sequencing projects. In the first strategy called clone-based shotgun sequencing, a *physical map* of the genome is first constructed before sequencing. Here multiple copies of the genome are broken into DNA segments of sizes from 40 kb to 200 kb. Each DNA segment is characterized by a number of markers. A *marker* on a DNA segment could be any identification feature for the segment. Two DNA segments overlap if they share some markers. The DNA segments are ordered using overlaps between segments. A physical map of the genome is a collection of ordered DNA segments that cover the complete genome. Once a physical map of the genome is constructed, DNA segments from the map are selected for sequencing. The sequence of a DNA segment is determined by the shotgun sequencing strategy. The sequence of the genome is constructed from the sequences of the DNA segments according to the physical map. The current cloning vector of choice for carrying a DNA segment is the *bacterial artificial chromosome* (BAC). The advantage of this approach is obvious. It breaks the large genome into segments that can be easily handled by the current sequencing technology. While the genome may contain repetitive regions longer than DNA segments, DNA segments can only contain short repetitive regions. On the other hand, a major difficulty with this approach is in ordering the DNA segments. Because the markers for characterizing DNA segments are not precise and the genome contains long repetitive regions, it is very difficult to order the DNA segments with the markers. The genomes of the bacterium *E. coli*, the yeast *S. cerevisiae*, and the nematode *C. elegans* were sequenced using this clone-based approach with physical mapping [1, 2, 3].

The clone-based strategy is currently used to sequence the human ge-

nome by the public efforts called the Human Genome Project ([www.nhgri.nih.gov](http://www.nhgri.nih.gov)). Because it was not possible to construct a physical map of the human genome before sequencing, the project went directly to sequencing without a prior physical map. In order to produce human sequences quickly and make them immediately available to the scientific community, the Human Genome Project decided to produce draft sequences of 99% accuracy first. To generate draft sequences, DNA segments were selected and sequenced at low coverage. Here every position of a sequenced DNA segment is covered by three to five reads on the average. Reads from a DNA segment were assembled into a number of contigs (continuous assemblies of DNA sequence). Consensus sequences of contigs were made immediately available on the web.

The initial phase of generating draft human sequences with the clone-based strategy was recently completed [36]. A total of 29,298 DNA segments were selected for sequencing. Of the 29,298 segments, 21,021 segments were sequenced at a low coverage depth of 2 to 5 and the rest at a high coverage depth of 8 to 12. A physical map of DNA segments was constructed simultaneously with the sequencing efforts. The map was used to place the sequenced DNA segments on the genome. Contigs from overlapping segments were merged. The clone-based strategy produced a total of 87,757 scaffolds and a total of 149,821 contigs. Half of all nucleotides of the genome are within a contig of at least 82 kb and within a scaffold of at least 274 kb. The Human Genome Project has moved to the next phase of producing human sequences of high quality (99.99% accuracy) by 2003.

An important issue in the cloned-based strategy is how to produce the complete sequence of the human genome without a physical map. A walking approach has been proposed to address the issue [4]. In this approach, a number of DNA segments are randomly selected and fully sequenced. Then the genome is walked by iteratively selecting and sequencing DNA segments that minimally overlap a fully sequenced DNA segment. To find DNA segments that overlap a fully sequenced DNA segment  $f$ , two reads are produced for every DNA segment, one from each end of the segment. A database of all end reads is constructed. The sequence of segment  $f$  is compared with the database of end sequences. A DNA segment overlaps segment  $f$  if an end sequence of the DNA segment is highly similar to a region of the sequence of segment  $f$ . A major problem in the cloned-based strategy is that the walking process can only be carried out in a small number of steps because it takes a month to fully sequence a DNA segment [5]. One way of avoiding many walking steps is to start with a large number of DNA segments as seeds and perform walking from the seeds in parallel. Parallel walking, however, causes a problem of redundant sequencing; DNA segments with large overlaps are sequenced at the same time. Some chromosomes of the plant *Arabidopsis* were sequenced using the walking method [6].

In the second strategy called whole-genome shotgun sequencing (WGS), the shotgun sequencing method is applied to the whole genome. Multiple



copies of the genome are broken into pieces. Both ends of every piece are read by an automated sequencing machine to produce two sequence reads, one from each strand of the piece. The size of the piece is measured, which is the distance between the two reads on the genome. The two reads along with their distance and orientation information form a constraint. Short reads are assembled into contigs. Constraints are used to order and orient contigs from unique and short repetitive regions of the genome into scaffolds. A scaffold is a collection of contigs that are ordered and oriented with respect to the genome. The advantage of the WGS method is that the method can quickly produce a draft picture of the genome that shows the order and orientation of unique and short repetitive regions of the genome. The disadvantage is that it cannot handle repetitive regions that are longer than constraints. A major challenge in the WGS strategy is development of a powerful sequence assembly program that can handle tens of millions of reads and use constraints to sort out short repeats such as the Alu repeat. In 1995, the WGS method was used to sequence the 1.8-Mb genome of the bacterium *H. influenzae* [7].

The WGS method was used to sequence the genome of the fruit fly *D. melanogaster* [8]. A total of 3.16 million reads were generated in the WGS sequencing phase. After trimming to an accuracy level of 98%, the average length of reads is 551 bp. Of the 3.16 million reads, 1.15 million pairs of reads form constraints, 57% of which have an average distance of 2 kb and the rest have an average distance of 10 kb. A total of 491 000 reads including 12 152 pairs of BAC end reads with an average distance of 130.2 kb, were produced by the public efforts using the clone-based method. The joint data set was assembled by the Celera Assembler into 838 firm scaffolds, where a *firm* scaffold contains a contig from a unique region of the genome. Of the 838 firm scaffolds, 134 scaffolds are considered to be the preliminary reconstruction of the euchromatic sequence. There are 25 scaffolds of sizes greater than 100 kb, which cover more than 95% of the assembled sequence. The largest scaffold is 24.3 Mb in size. The assembly is in excellent agreement with the maps of the *Drosophila* genome produced by the clone-based method. The assembly took less than a week on an eight-processor suit of Compaq Alpha ES40s with 32-Gb memory.

Draft sequences of the human genome were produced by the WGS method at Celera Genomics [37]. A random shotgun data set of 27 million reads with a coverage depth of 5 was produced at Celera. After quality and vector trimming, the average length of reads is 543 bp and the average accuracy level of reads is 99.5%. The data set contains constraints of three distance classes: 2 kb, 10 kb, and 50 kb. The random shotgun data set was combined with a synthetic shotgun data set of 16 million reads. The synthetic shotgun data set was produced by shredding consensus sequences of BAC contigs from the Human Genome Project. No constraint information about the synthetic data set was generated. The combined data set was used for assembly by the WGS method. The WGS method produced a total of

118,968 scaffolds and a total of 221,036 contigs. The average length of scaffolds is 23,938 bp and the average length of contigs is 11,702 bp. The total amount of assembly work was about 20,000 hours for one processor. The computer platform for the assembly task consists of 10 Compaq ES40 servers, each with 4 processors and 4 Gb of main memory, and a Compaq GS160 computer with 16 processors and 64 Gb of main memory.

A localized assembly approach was also pursued by Celera. In this approach, consensus sequences of BAC contigs from the Human Genome Project were used to partition the random and synthetic reads into components such that the reads in a component come from a megabase region of the genome. Each component was assembled by the WGS method. The localized assembly approach produced slightly better results than the WGS method.

Because of major improvements in automated sequencing machines, hundreds of reads can be produced by a sequencing machine in a day. Computers are also much faster and have much more memory. It appears more cost-effective and efficient to produce a draft picture of a mammalian genome with the WGS method first. Then the clone-based method is used to walk the regions of the genome with long repeats, which cannot be handled by the WGS method. The combination of the WGS method and the clone-based method can work if a large portion of the genome are unique or contain only short repeats. The two methods complement each other nicely. The WGS method can produce correct assemblies for unique and short repetitive regions of the genome, which avoids the redundant-sequencing problem with the clone-based method. For every repetitive region of the genome that can not be handled by the WGS method, the clone-based method can sort it out in a few walking steps.

Throughout this chapter we will expect basic familiarity with approaches to the analysis and alignment of biological sequences. This knowledge can be drawn from Chapter 2 of Volume 1 of this book. In the following, we will use notation that is compatible with that chapter.

This Chapter focuses on DNA sequence assembly, a major phase in genome sequencing. The problem is to assemble short reads into long sequences. A *read* is a short DNA sequence. As mentioned before, some bases of a read are less accurate than others. One way of addressing sequencing errors in a read is to give to the assembly program the degree of accuracy of each base in the read. This enables the program to treat bases differently according to their degree of accuracy. The *quality value* of a base in a read is  $q = -10 \times \log_{10}(p)$ , where  $p$  is the estimated error probability for the base [9]. The way in which quality values are defined allows small positive integers to cover a wide range of error probabilities. For example, a quality value of 10 corresponds to an error rate of 1 in 10 and a quality value of 40 corresponds to an error rate of 1 in 10 000. Reads along with their quality values are produced by a base-calling program such as Phred [10]. Phred translates a trace of digital signals from a sequencing machine into a read

and computes a quality value for each base of the read. Input to the assembly program includes a set of reads, their quality values, and constraints.

Reads are assembled into contigs by the assembly program. A *contig* is an ordered list of overlapping reads. A contig is shown in two ways. A layout of all reads in a contig provides an overview of the contig. A multiple alignment of all reads in a contig gives a detailed view of the contig at the base level. The sequence of a contig is a consensus sequence of a multiple alignment of reads in the contig. Output from the assembly program includes the sequences of all contigs and their quality values. A small assembly example is given in this Chapter.

A number of DNA sequence assembly programs have been developed [11, 12, 13, 14, 15, 16, 17, 18, 19, 20]. These programs are for BAC and microbial genome assembly. We have developed three versions of the CAP assembly program [13, 21, 22]. In this Chapter, we describe the assembly techniques used in the CAP3 sequence assembly program [22], which is for BAC and microbial genome assembly. The CAP3 program includes a number of improvements and new features. A capability to clip 5' and 3' low-quality regions of reads is included in the CAP3 program. Base quality values are used in computation of overlaps between reads, construction of multiple sequence alignments of reads, and generation of consensus sequences. Efficient algorithms are employed to identify and compute overlaps between reads. Forward-reverse constraints are used to correct assembly errors and link contigs.

## 2.2 Methods

The sequence assembly problem is to reconstruct the sequence of a DNA segment from short reads, where each read is approximately the sequence of a short piece of the DNA segment. Note that reads contain sequencing errors and some reads are chimeric. A mathematical formulation of the problem, called *shortest common superstring problem* (SCS), is to compute a shortest sequence such that each read is exactly identical to a region of the sequence. SCS is a simplification of the assembly problem because it does not address sequencing errors, chimeric reads, or repeats. Nevertheless, it has been shown that SCS is NP-hard, that is, it is likely that there is no efficient algorithm for solving SCS exactly [23]. An approximation algorithm with guaranteed performance has been developed for the SCS problem [24]. The algorithm is guaranteed to produce a sequence of length  $O(n \log n)$  for an optimal sequence of length  $n$ . Note that constraints are not included in the SCS formulation. A mathematical formulation of the sequence assembly problem that includes constraints, called SCS with constraints, is to compute a shortest sequence such that each read is very sim-

ilar to a region of the sequence and the maximum number of constraints are satisfied by the sequence. Because SCS is a special case of SCS with constraints, it is likely that there is no efficient algorithm for solving SCS with constraints exactly. For SCS with constraints, no approximation algorithm is known that guarantees its performance in terms of percent identity between a sequence produced by the approximation algorithm and an optimal sequence. In practice, an engineering approach has been taken to address the assembly problem. A sequence assembly program is developed, evaluated, and improved on real data. The program is evaluated by comparing the sequences produced by the program with the answer sequence that is verified by experimental laboratory techniques. Evaluation of the program on a large number of real data is very helpful for finding and addressing deficiencies of the program.

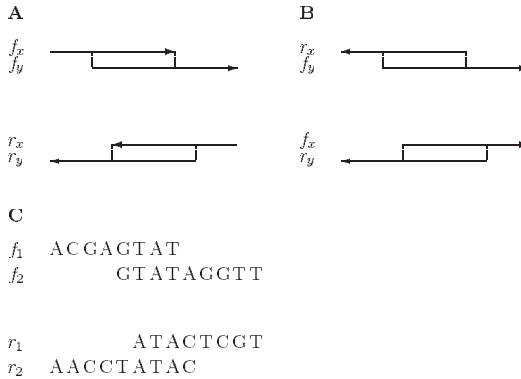
The assembly algorithm of CAP3 works in three major phases. In the first phase, pairs of similar reads are identified. Poor end regions of each read are identified and removed. Overlaps between reads are computed. False overlaps are identified and removed. In the second phase, reads are joined to form contigs in decreasing order of overlap scores. Then forward-reverse constraints are used to make corrections to contigs. In the third phase, a multiple sequence alignment of reads is constructed and a consensus sequence along with a quality value for each base is computed for each contig. Note that base quality values are used in computation of overlaps and construction of multiple sequence alignments. We describe each step in detail below.

### 2.2.1

#### Fast identification of pairs of similar reads

Pairs of similar reads are identified using a fast database search technique. Let  $f_1, f_2, \dots$  be all the input reads in given orientation and let  $r_x$  be the reverse complement of read  $f_x$ . The goal of this step is to find quickly pairs of similar reads  $f_x$  and  $f_y$  with  $x < y$  and pairs of similar reads  $r_x$  and  $f_y$  with  $x < y$ . Note that each identified pair of reads represents two symmetric pairs. Because reads  $r_x$  and  $r_y$  are the reverse complements of reads  $f_x$  and  $f_y$ , respectively, a similarity between reads  $f_x$  and  $f_y$  is complementary to one between reads  $r_x$  and  $r_y$  and a similarity between reads  $r_x$  and  $f_y$  is complementary to one between reads  $f_x$  and  $r_y$ . Two types of complementary similarities are shown in Figure 2.1.

First consider how to determine quickly if two reads are similar. An alignment between two reads is simplified as an ordered chain of segment pairs, where each segment pair is an ungapped portion of sufficient length of the alignment. A part of the alignment between two adjacent segment pairs is considered as a gap in the chain. Two reads are similar and hence have a potential overlap if the reads contain a chain of similarity score

**Fig. 2.1**

Pairs of complementary overlaps. Read  $r_x$  is the reverse complement of read  $f_x$  and read  $r_y$  is the reverse complement of read  $f_y$ . **(A)** A similarity between two reads is indicated by a pair of broken lines. The

similarity between reads  $f_x$  and  $f_y$  and the similarity between reads  $r_x$  and  $r_y$  are complementary, and **(B)** so are the similarity between reads  $r_x$  and  $f_y$  and the similarity between reads  $f_x$  and  $r_y$ . **(C)** An example pair of complementary overlaps.

greater than a score cutoff. The score of a chain is the sum of scores of each segment pair and each gap in the chain. Segment pairs between two reads are computed by extending exact matches between the two reads [25]. See section 2.4 in Chapter 2 of Volume 1 of this book for details of computing segment pairs in the BLAST program. Segment pairs are combined into high-scoring chains by a dynamic programming algorithm as follows [26, 27, 28]. The last antidiagonal of a segment pair is the sum of the end positions of the two segments in the pair. The segment pairs are considered in increasing order of their last antidiagonals. The maximum score of chains ending at the current segment pair is computed. In order to obtain independent high-scoring chains, chains are partitioned into classes such that all chains in a class begin with the same segment pair. For each class, a chain with the maximum score in the class is selected as the representative for the class.

Next consider, for each read  $f_x$ , how to find all reads  $f_y$  with  $x < y$  such that  $f_x$  and  $f_y$  are similar (have a potential overlap). One approach to addressing this problem is to go through each read  $f_y$  with  $x < y$  and determine if  $f_x$  and  $f_y$  are similar. However, this approach is not efficient because it has to go through every read  $f_y$  that is not similar to read  $f_x$ . Below we describe a hashing technique that allows us to find directly reads  $f_y$  that are similar to read  $f_x$ .

The sequences of all reads  $f_1, f_2, \dots$  are concatenated together with a special character inserted at every read boundary. The resulting sequence is called the *combined sequence*. A *hash table* is constructed for the combined sequence, where for every word of length  $w$ , the table gives the positions of

each occurrence of the word in the combined sequence. Each read  $f_x$  is compared with the combined sequence to find pairs of reads  $f_x$  and  $f_y$ ,  $x < y$ , with a potential overlap as follows. A word of length  $w$  at each position of read  $f_x$  is considered. The hash table is used to find each occurrence of the word in the combined sequence. Note that reads  $f_y$  that have no common word of length  $w$  with read  $f_x$  are no longer considered. All exact matches of length  $w$  between read  $f_x$  and the combined sequence are extended into longer segment pairs. Segment pairs are combined into high-scoring chains by the dynamic programming method described above. The special boundary character is used to make sure that only segment pairs from the same read in the combined sequence are combined in a chain. Every high-scoring chain between read  $f_x$  and the combined sequence indicates that read  $f_x$  is similar to a region of the combined sequence. We need to determine the corresponding read  $f_y$  from which comes the region of the combined sequence. This is done as follows. Let  $j$  be the position of the region in the combined sequence. Let  $E$  be a list such that for each index  $z$ ,  $E(z)$  is the end position of read  $f_z$  in the combined sequence. A binary search for position  $j$  is performed in the list  $E$  to find the smallest index  $y$  such that  $j \leq E(y)$ . Since read  $f_x$  is similar to the region of the combined sequence, which comes from read  $f_y$ , reads  $f_x$  and  $f_y$  are similar and have a potential overlap. Note that the chains between read  $f_x$  and any read  $f_y$  with  $x \geq y$  in the combined sequence are ignored. Similarly, each read  $r_x$  is compared with the combined sequence to find pairs of reads  $r_x$  and  $f_y$ ,  $x < y$ , with a potential overlap.

The criterion for selecting a value for the word length  $w$  is that the hashing method performs well and requires a reasonable amount of memory. The number of entries in the hash table is  $4^w$ , the number of all words of length  $w$ . We suggest that  $w$  be set to a value such that  $4^w$  is close to the total length of all reads. This choice requires a linear amount of memory for the hash table and allows the hashing method to perform effectively because each entry in the table is likely to contain a small number of word occurrences. The default value for  $w$  in CAP3 is 11.

For a pair of reads with a potential overlap, as indicated by high-scoring chains of segment pairs, the overlap can be computed by the Smith–Waterman algorithm described in Chapter 2 of Volume 1. However, it is slow and not necessary to perform the computation over the full matrix because the overlap is of strong similarity and is located in a small area of the matrix. Here we describe a way of determining the small area of the matrix for the overlap using high-scoring chains computed previously for a pair of reads. A *diagonal* of the matrix is named by a number  $k$ , such that the diagonal consists of all entries  $(i, j)$  with  $j - i = k$ . A segment pair beginning with position  $i$  of one read and position  $j$  of the other read is said to occur on diagonal  $j - i$ . A *band of diagonals* from numbers *low* to *upp* with  $low \leq upp$  consists of diagonals  $low, low + 1, low + 2, \dots, upp$ . A band of diagonals in the matrix covers a chain of segment pairs if each segment

pair in the chain occurs on a diagonal inside the band. For a pair of reads with high-scoring chains of segment pairs, a minimum band of diagonals from *low* to *upp* is determined, where *low* is the small number of diagonals on which segment pairs in the high-scoring chains occur and *upp* is the largest number of diagonals on which segment pairs in the high-scoring chains occur. Clearly, the band from diagonals *low* to *upp* covers all the high-scoring chains of segment pairs between the reads.

For each pair of reads with a potential overlap, a minimum band of diagonals is determined to cover all the high-scoring chains of segment pairs between the reads using the method mentioned above. Later the computation of the overlap between the reads is carried out over the band of diagonals in the matrix.

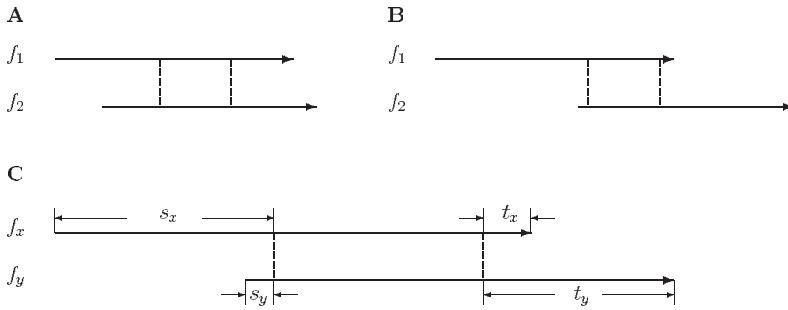
### 2.2.2

#### Clipping of poor end regions

Both base quality values and sequence similarities are used to identify poor end regions of full-length reads. Using base quality values alone is not sufficient for finding the exact positions of poor end regions since poor end regions of chimeric reads may have high base quality values. Any sufficiently long region of high quality values that is highly similar to a region of another read is defined to be good. In addition, any sufficiently long region that is highly similar to a good high-quality region of another read is defined to be good. The 3' clipping position of a read is the maximum of 3' end positions of good regions of the read. The 5' clipping position of a read is the minimum of 5' end positions of good regions of the read.

The high-quality region of each read is identified by computing a region of the read with the maximum quality score using a one-dimensional dynamic programming algorithm. The quality score of a base is its quality value minus a quality value cutoff and the quality score of a region is the sum of quality scores of each base in the region. The quality value cutoff is often an integer between 10 and 20, which corresponds to an error rate between 10% and 1%.

Regions of reads with a strong sequence similarity to other reads are identified by computing the start and end positions of an optimal local alignment for each pair of similar reads. An *overlap* between two reads is defined as an optimal local alignment between the reads, that is, a local alignment with the maximum score. However, if there are two optimal local alignments between the reads as shown in parts A and B of Figure 2.2, which of the two alignments is likely to represent the true overlap? The alignment in part B of Figure 2.2 is likely to represent the true overlap because in the true overlap, an end of a read is likely to be similar to an end of the other read and therefore there is likely to be a short terminal region

**Fig. 2.2**

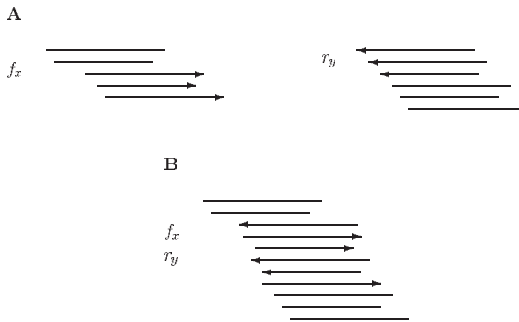
Definition of an overlap as a local alignment. (A) An optimal local alignment between  $f_1$  and  $f_2$  with two long terminal regions at each end. (B) An optimal local alignment between  $f_1$  and  $f_2$  with a short

terminal region at each end. (C) A local alignment between reads  $f_x$  and  $f_y$ . Let  $s_x$  and  $t_x$  be the lengths of the terminal regions of read  $f_x$  before and after the alignment, respectively.

before and after the overlap. The score of a local alignment is defined as follows so that an optimal local alignment (one with the maximum score) represents the real overlap between the two reads. Our strategy is to reflect the lengths of the terminal regions before and after an alignment in the score of the alignment. The score of a local alignment is the sum of scores of each match and each difference in the alignment plus the 5' and 3' gap penalty scores for the terminal regions not included in the alignment. Let  $term(s) = -(go + ge \times s)$  be the penalty score of a terminal region of length  $s$ , where non-negative integers  $go$  and  $ge$  are gap open and gap extension penalties. The values  $go$  and  $ge$  should be much smaller than the gap open and extension penalties used to score gaps within the alignment in order to tolerate terminal regions due to contamination. For the alignment in part C of Figure 2.2, the 5' gap penalty score is the smaller of  $term(s_x)$  and  $term(s_y)$ , and the 3' gap penalty score is the smaller of  $term(t_x)$  and  $term(t_y)$ . The algorithm of Smith and Waterman [29] is modified to handle penalties for terminal regions. For efficiency, computation with the modified algorithm of Smith and Waterman [29] is restricted to a band of diagonals in the dynamic programming matrix [30, 19]. Recall that our fast method for finding each pair of reads with a potential overlap reports a band of diagonals for the pair. The band is expanded in both directions by a certain number of bases, which can be specified by the user. A local alignment computation is performed over the expanded band of diagonals.

Base quality values are incorporated into the local alignment algorithm. Scores of matches and differences are weighted by the quality values of the bases involved. Specifically, the score of a match or a difference involving two bases of quality values  $q_1$  and  $q_2$  is weighted by  $\min(q_1, q_2)$ .





**Fig. 2.3**  
Merging of two contigs into a new contig. (A) The current overlap involves read  $f_x$  in one contig and read  $r_y$  in the other contig. (B) A new contig is obtained by merging the two contigs.

### 2.2.3

#### Computation and evaluation of overlaps

Overlaps between clean reads are computed as follows. An overlap between two reads is defined as a global alignment of the reads with the maximum similarity score. Match and difference scores are weighted by base quality values. For efficiency, the computation of an overlap between two reads is restricted to a band of diagonals centered at the start position of the optimal local alignment computed previously. The width of the band is the number of diagonals required to cover the optimal local alignment plus the band expansion size specified by the user.

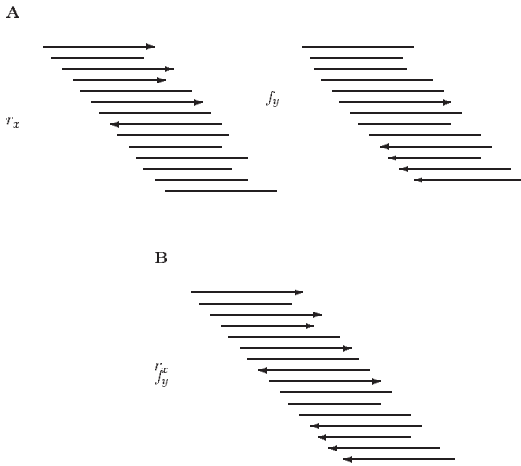
Each overlap is evaluated by several criteria. Any overlap that does not satisfy any of the requirements is not considered in construction of contigs. Requirements are imposed on the length, percent identity, and similarity score of the overlap. In addition, if the overlap contains a sufficient number of differences at bases of high quality values, then the overlap is not used.

### 2.2.4

#### Construction of contigs

A greedy method is used to construct contigs as follows. Initially, each read is a contig by itself. The overlaps are considered in decreasing order of their scores. Consider the current overlap involving reads  $f_x$  and  $r_y$ . If reads  $f_x$  and  $r_y$  are in separate contigs and the contigs can be merged into a longer contig using the overlap, then the two contigs are merged. Otherwise, no action is taken. Figure 2.3 shows an example of merging two contigs into a longer one. The step is repeated until all the overlaps are considered.

Forward-reverse constraints are used to correct errors in the construction of contigs and to produce a partial order of contigs. A *forward-reverse constraint* consists of two reads, and two integers that specify a range for the



**Fig. 2.4** Correction to contigs. **(A)** An unused overlap between reads  $r_x$  and  $f_y$  with each indicated by a line with an arrow head is supported by four unsatisfied constraints. The eight remaining lines with arrow heads indicate the eight reads involved in the four constraints. **(B)** A new contig is obtained by making a correction to the contigs in **A**. The contig with read  $r_x$  is disconnected right after  $r_x$ , the contig with read  $f_y$  is disconnected right before  $f_y$ , and the portion with  $r_x$  and the portion with  $f_y$  are merged into the new contig.

distance between the reads. Constraints are produced by sequencing both ends of each subclone. A constraint is satisfied by a contig if the two reads occur in the contig, the upstream read is in forward orientation, the downstream read in reverse orientation, and the distance between the two reads is within the given range. Otherwise, the constraint is unsatisfied. An overlap is unused if the overlap is not used in any contig. A group of unsatisfied constraints support an unused overlap if, after a correction is made to the current set of contigs to use the overlap in a resulting contig, all the constraints in the group are satisfied with respect to the resulting contig.

The algorithm for using constraints consists of the following steps. In step 1, each constraint is evaluated with respect to the current contigs to determine if it is satisfied or not. Unsatisfied constraints are partitioned into groups, where all constraints in a group support an unused overlap. In step 2, an unused overlap supported by the largest number of unsatisfied constraints is selected for consideration. If the number of unsatisfied constraints supporting the unused overlap is sufficiently larger than the number of satisfied constraints against the unused overlap, then a correction is made to the current set of contigs to implement the unused overlap. Figure 2.4 shows how a new contig is obtained by making a correction to existing contigs. If no correction is made for the selected overlap, then other unused overlaps are considered until a correction is made to the current set of

contigs or no more unused overlap is available for selection. In step 3, if no correction is made in the last step, then the algorithm is terminated. Otherwise, steps 1 and 2 are repeated.

Contigs are ordered using constraints as links. Unsatisfied constraints are partitioned into groups, where all constraints in a group link a pair of contigs. The groups are considered in decreasing order of group size. For each group of constraints that are associated with a pair of contigs, if the number of constraints in the group is sufficiently large and neither of the two contigs is already linked to other contigs at the corresponding end, then a link between the contigs is established.

### 2.2.5

#### **Construction of consensus sequences**

A multiple sequence alignment of reads is constructed for each contig. The construction is performed by repeatedly aligning the next read with the current alignment. The reads are considered in increasing order of their positions in the contig.

We consider aligning one read with the current alignment of reads. For clarity, the current alignment is called a block. Although the block may be very long, a major portion of the block remains unchanged for the rest of the construction. Thus, only the 3' portion of the block that gets changed is used for alignment. This 3' portion of the block is replaced by the resulting alignment. In the following, the block means the 3' portion. To produce an accurate alignment, base quality values are used in the construction. In this scheme, seven average quality values are computed for each column of the block: five for substitution, one for deletion, and one for insertion. Of the five average quality values for substitution, four values are for the four regular base types and one value is for the ambiguous base type. The average quality value for a base type is the signed sum of quality values of each base in the column divided by the number of bases in the column. The signed sum is computed by considering every base in the column. If the base is of the base type, its quality value is added to the sum. Otherwise, its quality value is subtracted from the sum. The average quality values of the block are used along with the quality values of the read to weight match and difference scores. A global alignment of the block and the read with the maximum score is computed in linear space using a divide-conquer technique [31, 32, 33]. Since the pair-wise alignment computation is performed at most once for each read, it is affordable to carry out the computation over the entire dynamic programming matrix for best results. The average quality values for the block are precomputed so that each entry in the dynamic programming matrix is calculated in a constant time.

After an alignment is constructed, a consensus sequence along with a quality value for each base is computed for the contig. For each column of

the alignment, a weighted sum of quality values is calculated for each base type and the base type with the largest sum of quality values is taken as the consensus base for the column. The quality value for the consensus base is the sum of quality values for the consensus base type minus the sum of quality values for every other base type. However, if the column contains two base types, each with a very large sum of quality values, then a very low quality value is assigned to the consensus base. The assignment of the low quality value indicates a potential problem often caused by polymorphism or collapsing of highly similar copies of a repetitive element.

## 2.3

### An example

We present a sequence assembly example. The data set in this small example contains 6 real reads of lengths from 799 bp to 1022 bp. The reads are named R1 through R6 with R+ denoting read R in given orientation and R- denoting the reverse complement of read R+. Read R- is obtained by reversing read R+ and exchanging bases A and T, and exchanging bases C and G. Each read consists of a sequence of bases and a sequence of quality values. The base sequence and the quality value sequence are of the same length and the accuracy of a base at position  $i$  in the base sequence is indicated by a quality value between 0 and 99 at position  $i$  in the quality sequence with 0 being the lowest quality and 99 the highest quality. An exception is that quality value 98 is reserved for an ambiguous base call indicated by the letter N [19]. This exception provides a way for the user to inform the assembly program that some bases are ambiguous. In the assembly program, bases of quality value 98 are not treated as bases of good quality. The base sequences of the reads are in one file and the quality values are in another file. The sequence and quality files are available at <http://genome.cs.mtu.edu/cap/data>.

The six reads were assembled into one contig by the CAP3 program, with three reads in given orientation and the other reads in reverse orientation. Below we go through the assembly procedure on the example in detail.

A high-quality region of each read was computed using the quality values of the reads. The quality value cutoff was set to 10. Table 2.1 shows the start and end positions of the high-quality region of each read and the length of the read. Fifteen pairs of reads with a similarity were identified. Table 2.2 shows the start and end positions of the similar regions of the reads in the proper orientation for each pair. For example, the first pair in Table 2.2 involves the region from bases 2 to 427 of read R1-, which corresponds to the region from bases 434 to 859 of read R1+. The 5' and 3' clipping positions of each read were computed from the pairs of similar regions in Table 2.2. For example, the five pairs of regions involving read R1 give the following

**Tab. 2.1**

The start and end positions of high-quality read regions.

<i>Read</i>	<i>Start</i> <sup>a</sup>	<i>End</i>	<i>Read length</i>
R1+	5	849	860
R2+	55	888	1022
R3+	55	870	918
R4+	4	790	799
R5+	17	599	920
R6+	70	789	850

<sup>a</sup> Positions are relative to raw reads in given orientation.**Tab. 2.2**

The start and end positions of similar read regions.

<i>Read</i>	<i>Start</i> <sup>a</sup>	<i>End</i>	<i>Read</i>	<i>Start</i>	<i>End</i>
R1–	2	427	R4+	374	799
R1–	2	239	R5+	390	628
R1–	2	311	R6+	518	827
R1+	12	829	R2+	55	873
R1+	12	829	R3+	55	874
R2–	27	545	R4+	282	799
R2–	34	357	R5+	304	628
R2–	27	429	R6+	426	827
R2+	55	838	R3+	55	838
R3–	4	441	R4+	364	799
R3–	90	253	R5+	464	628
R3–	4	325	R6+	508	827
R4+	46	611	R5+	52	628
R4+	18	683	R6+	152	827
R5+	52	628	R6+	181	755

<sup>a</sup> Positions are relative to raw reads in specified orientation.

good regions of read R1+: bases 434–859, 622–859, 550–859, 12–829, and 12–829. The 5' clipping position of read R1+ is 12, the smallest start position of the good regions, and the 3' clipping position of read R1+ is 859, the largest end position of the good regions. Table 2.3 shows the clipping positions of each read in forward orientation. The poor regions of each read were removed. The resulting reads are called clean reads.

For each pair of similar regions, an overlap between the clean reads was computed. The overlaps were evaluated. No overlap was removed. Table 2.4 shows the overlaps in decreasing order of scores. A containment is an overlap where one clean read is entirely similar to a region of another clean read. The overlaps were considered in decreasing order of scores for construction of contigs.

Tab. 2.3

The start and end positions of clean regions.

<i>Read</i>	<i>Start</i> <sup>a</sup>	<i>End</i>	<i>Read length</i>
R1+	12	859	860
R2+	55	996	1022
R3+	55	915	918
R4+	18	799	799
R5+	52	628	920
R6+	152	827	850

<sup>a</sup> Positions are relative to raw reads in given orientation.

Tab. 2.4

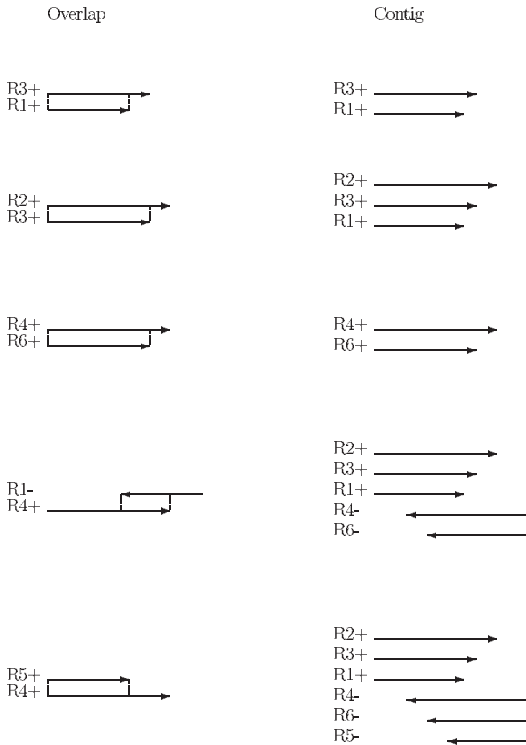
The positions and scores of overlaps.

<i>Read</i>	<i>Start</i> <sup>a</sup>	<i>End</i>	<i>Length</i> <sup>b</sup>	<i>Read</i>	<i>Start</i>	<i>End</i>	<i>Length</i>	<i>Score</i>
R1+	1	848	848	R3+	1	850	861	53378
R3+	1	861	861	R2+	1	861	942	51554
R1+	1	848	848	R2+	1	849	942	50668
R6+	1	676	676	R4+	1	666	782	36206
R4+	357	782	782	R1-	1	426	848	19848
R4+	265	782	782	R2-	1	519	942	19133
R4+	347	782	782	R3-	1	438	861	18011
R5+	1	577	577	R4+	29	594	782	10311
R6+	275	676	676	R2-	1	403	942	10308
R5+	1	577	577	R6+	30	604	676	10211
R6+	367	676	676	R1-	1	310	848	10047
R6+	357	676	676	R3-	1	322	861	8516
R5+	246	577	577	R2-	1	331	942	3021
R5+	339	577	577	R1-	1	238	848	2473
R5+	327	577	577	R3-	1	250	861	1871

<sup>a</sup> Positions are relative to clean reads in specified orientation.<sup>b</sup> Length is the length of a clean read without poor end regions.

A layout of reads in a contig is represented by an ordered list of containment trees. A child read of a parent read in a containment tree means that the child read is completely covered by the parent read in the layout. The root read of a tree is not covered by any other read in the layout. The root reads of two adjacent containment trees overlap in the layout. A doubly linked list of root reads is used to represent both strands of a contig, one direction for one strand.

Initially, each read was a tree by itself. The overlap between reads R1+ and R3+ was first considered. This overlap shows that read R1+ is completely covered by read R3+. Read R1+ was made a child of read R3+ and



**Fig. 2.5**  
The current overlap and the resulting contig after the overlap is processed. The overlaps are considered in the top-down order.

R1- a child of R3-. The overlap between reads R3+ and R2+ was considered next, which shows that read R3+ is completely covered by read R2+. Read R3+ became a child of read R2+ and R3- a child of R2-. No action was taken on the overlap between reads R1+ and R2+ since reads R1+ and R2+ were already in the same contig. Because of the overlap between reads R6+ and R4+, read R6+ became a child of read R4+ and R6- a child of R4-. The overlap between read R4+ and R1- was next considered. Since R1- was not a root, the root of the tree containing read R1- was located, which was read R2-. A search was performed to see if there was an overlap between R4+ and R2-. Since there was an overlap between R4+ and R2-, a link from R4+ to R2- and a link from R2+ to R4- were set. No action was taken on the overlap between reads R4+ and R2- since they were already in the same contig. There was no action on the overlap between reads R4+ and R3-. The next overlap caused read R5+ to become a child of read R4+ and R5- a child of R4-. No action was taken for each of the remaining overlaps. Figure 2.5 shows the current overlap and the resulting contig for each assembly step.

No forward-reverse constraints were used for this data set. One direction of the doubly linked list of roots was selected for construction of a multiple

```

R2+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R3+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R1+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R4-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R5-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R6-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
consensus  CCCTCGTGTGCGCGCACCCGGGCCAGCCCTATAGAAACATCTGAGGAGTCACTTCCTC-C

R2+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R3+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R1+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R4-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R5-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R6-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
consensus  CATGACTCTCGCCCGCCCGGCCGGCTGGAGTCGGCTCCTGGCAAGCTTCAGGCACCTCAG

R2+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R3+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R1+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R4-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R5-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R6-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
consensus  TTGTCTGAATACACACAGCACCCCTTTCCTTACTGAAGCCCTGAGAGCCTCCAGTTCTC

R2+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R3+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R1+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R4-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R5-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R6-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
consensus  CCTCCTTGCT-CACCCAGACTTCCTCCCTCCCGGCCCTTCAGCGTTCCAAAGGGTAAAT

R2+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R3+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R1+   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R4-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R5-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
R6-   .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .      .
consensus  -GTGGCTGGGAAGACAGCTCAGCAGTGGAGAAATCGGGTGCAGG-TCAGTGTGTGTCAT

```

Fig. 2.6

A portion of a multiple alignment of reads and a consensus sequence produced by CAP3 on the example data set.

alignment of reads in the contig, say, the direction from read R2+ to R4-. The final list consists of two roots R2+ and R4-. The tree rooted at R2+ has the edges from child R1+ to R3+ and from child R3+ to R2+. The tree at R4- has the edges from child R5- to R4- and from child R6- to R4-. The list is shown in Figure 2.5.

A multiple alignment of reads in the contig was constructed. Initially read R2+ was an alignment by itself. Then read R3+ was aligned with read R2+. Read R1+ was next added to the alignment. Then reads R4-, R5-, and R6- in this order were added to the alignment. A portion of the multiple alignment is shown in Figure 2.6. A consensus sequence along with its quality values was produced from the multiple alignment.



## 2.4

### Other assembly programs

We briefly describe computational techniques used in four other sequence assembly programs: Celera Assembler [8], GAP4 [34], Phrap [19], and TIGR Assembler [18]. Then we indicate the strengths of each program.

Celera Assembler is designed to use many computers to assemble millions of reads into long sequences. In Celera Assembler, a BLAST-like method is used to find quickly pairs of reads with a potential overlap. Overlap and coverage information are used to partition all the reads into unique reads and repetitive reads, where unique reads are from unique regions of the genome and repetitive reads are from repetitive regions of the genome. Unique reads are assembled into contigs called *unitigs*. Uniques are linked into scaffolds of contigs using constraints. Some gaps between unitigs are filled by repetitive reads linked by constraints to unique reads, where the proper place for a repetitive read is determined according to the location of the unique read and the distance in the constraint. Features of Celera Assembler are that it can handle millions of reads and that it makes extensive use of forward-reverse constraints to address the problem of repeats.

The GAP4 program is a comprehensive sequence assembly package. It supports directed and interactive assembly in addition to random shotgun assembly. A few external assembly programs (CAP2, FAKII, and Phrap) can be used in GAP4. Below we describe the internal assembly program in GAP4. Low-quality ends of reads are determined using quality values and removed in a preprocessing step. The clean reads are given as input to the assembly program in GAP4. In the assembly program, the overlap computation and construction of contigs are interleaved. The reads are processed one at a time as follows. The current read is compared with the consensus sequences of the current contigs. If the read overlaps well with some contigs, then a best overlap between the read and a contig is identified and the read is added to the contig by aligning the read with the consensus sequence of the contig. Otherwise, a new contig consisting only of the read is formed. The consensus sequence of a contig is obtained from a multiple alignment of reads in the contig using quality values. A major difference between the internal assembly program in GAP4 and the other assembly programs is that overlap computation and construction of contigs are interleaved, which is suitable for directed and interactive assembly. In the other assembly programs, overlap computation is before construction of contigs, which is suitable for random shotgun assembly.

The Phrap program takes as input full-length reads and their base quality values. The program works in three phases. In the overlap computation phase, a banded Smith–Waterman algorithm is used to compute overlaps between reads. In the contig construction phase, the overlaps are considered in decreasing order of scores. For the current overlap between reads  $f$  and  $g$ ,

if the contig containing read  $f$  overlaps the contig containing read  $g$ , then the two contigs are merged into one. Ends of the two contigs may need to be clipped in order to make the merging possible. Each contig is represented as a profile. In the last phase, a sequence along with its quality values is generated for each contig. The sequence is a mosaic of read regions of the highest quality values. In other words, the sequence consists of many short pieces, each of which is a region of a read with the highest quality values. A multiple alignment of reads in a contig is constructed by aligning each read with the sequence of the contig. Unique features of Phrap are that poor regions of reads are dynamically determined and removed during construction of contigs and that the sequence of a contig is constructed without using a multiple alignment of reads in the contig.

TIGR Assembler takes as input clean reads, which are produced from raw reads in a preprocessing step. The assembly program works in three phases. In phase 1, overlaps between reads are computed using a fast method based on counting of exact short word matches between reads. In phase 2, contigs are constructed one at a time as follows. A seed read is selected to start a new contig. The contig is grown by repeating the following step. A read that has a best overlap with the current contig is chosen and merged with the contig by aligning the read with a profile of the contig with the Smith–Waterman algorithm. Forward–reverse constraints are also used in the selection of a read to add it to the current contig. In phase 3, for each contig, a consensus sequence is generated from the profile of the contig. TIGR Assembler is the first program to perform whole-genome assemblies of microbial genomes.

Celera Assembler can handle a large mammalian genome. The other assembly programs are for small genomes. There are few performance comparisons of the assembly programs. Performance comparison of CAP3 and Phrap showed that Phrap often produces fewer, longer contigs than CAP3 [22]. Evaluation of CAP3, Phrap, and TIGR Assembler on public ESTs without quality values showed that CAP3 produces more accurate assemblies of ESTs [35]. No comparison of the assembly programs on their efficiency has been published. However, we found that Phrap is a few times faster than CAP3 on genomic BAC data sets and that Phrap requires much more memory than CAP3 on EST data of very deep coverage. CAP3 takes 1–3 hours on a genomic BAC data set [22]. Phrap works well on full-length reads with quality values. TIGR Assembler works well on clean reads with or without quality values. CAP3 works well on full-length reads without quality values. On data sets with constraints, CAP3 and TIGR Assembler make more use of constraints than Phrap. GAP4 works well on directed assembly. The strengths and availability of the assembly programs are summarized in Table 2.5. Because improvements are continually made to the assembly programs, heavy users of assembly programs should obtain the latest version of each program and evaluate the programs on their data. For instance, the fourth version of the CAP program, CAP4, has been de-

Tab. 2.5

The strengths and availability of the sequence assembly programs.

<b>Program</b>	<b>Strength</b>	<b>Availability</b>
CAP3	Assembly of ESTs	<a href="http://genome.cs.mtu.edu/cap/cap3.html">genome.cs.mtu.edu/cap/cap3.html</a>
Celera Assembler	Assembly of large genomes	<a href="http://www.celera.com">www.celera.com</a>
GAP4	Interactive assembly	<a href="http://www.mrc-lmb.cam.ac.uk/pubseq">www.mrc-lmb.cam.ac.uk/pubseq</a>
Phrap	Assembly of reads with quality values	<a href="http://www.phrap.org">www.phrap.org</a>
TIGR Assembler	Assembly of clean reads	<a href="http://www.tigr.org">www.tigr.org</a>

veloped. On five BAC and microbial data sets with quality values and constraints, CAP4 produced fewer, longer contigs than Phrap and was close to Phrap in running time ([www.paracel.com](http://www.paracel.com)).

## 2.5

### Conclusion

The genome community has made tremendous progress in sequencing the genomes of humans and model organisms. High quality sequences of the model organisms and draft sequences of the human genome have been produced. Efforts are under way to sequence the genomes of a number of important animals and plants. However, with the current sequencing technology, it will take a few years to produce high quality sequences of the human genome. Continued improvements to the sequencing technology are required to meet the challenges in those genome projects. Sequence assembly software, an important component of the sequencing technology, can have a major impact on a genome project, as demonstrated by Celera Assembler. We think that it is necessary to make continued improvements to the efficiency and accuracy of the existing sequence assembly programs in order to produce quickly high quality sequences of a large genome.

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## 3

### Analysis of Sequence Variations

*Christopher S. Carlson, Mark J. Rieder, and  
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#### 3.1

##### Introduction

Genetics may broadly be defined as the study of how traits, also known as phenotypes, are transmitted from generation to generation. While the basics of dominant and recessive inheritance (also known as Mendelian inheritance) have been known for better than a century, the genetic and biochemical basis of most traits has been difficult to determine. Nonetheless, family studies have shown that many traits of medical importance cluster in families, from common traits such as obesity and high blood pressure to less frequent traits such as sickle cell anemia and epilepsy. In the past two decades, unprecedented genetic tools have become available, which have facilitated the identification of the underlying molecular sequence variation causing many rare disorders, including disorders of drug metabolism [1] (Table 3.1). These tools include linkage analysis, positional cloning and association analysis, all of which rely on the identification and scoring of common variations in DNA sequences known as polymorphic markers. While linkage analysis is a proven technique for dissecting the genetics of rare Mendelian traits, the optimal study design for understanding common genetic disease and drug response is likely to be association analysis.

#### 3.2

##### Sequence variation

There are several major types of DNA sequence variation in the human genome. One type comprises repeated sequences with variations in the number of repeat units, such as short tandem repeat polymorphisms (STRPs, also known as microsatellites) in the form of mono-, di-, tri-, and tetra-nucleotide repeats, or more complex sequence repeats such as variable number tandem repeats (VNTRs) [2–4]. Repeat polymorphisms are relatively

**Tab. 3.1**

Steps in genetic mapping.

- 
- Identify a phenotypic trait of interest.
  - Evaluate the complexity of the genetic contribution to phenotype (Mendelian, non-Mendelian).
  - Design analysis based on genetics (linkage study for Mendelian trait, association study for non-Mendelian trait).
  - Collect samples, phenotypes, medical histories, DNA.
  - Perform whole genome scan or a candidate gene study (genotype the collected samples)
  - Statistical analysis to evaluate linkage to trait (linkage or linkage disequilibrium analysis)
  - Identification of gene or region(s) linked or associated with trait variability.
  - Identification of functional sequence variation within a region(s) or gene(s) relative to trait variability and direct testing of these relationships.
- 

rare, occurring every 50–100 kilobase pairs (kbp), and tend to occur in non-coding regions of the genome, but have many alleles at each marker. Because repeat polymorphisms have many alleles, they are very likely to be heterozygous in any given individual, which is ideal for linkage studies. The development of microsatellite marker maps and their refinement into a semi-automated fluorescence-based format greatly aided in the routine application of linkage analysis to a wide variety of phenotypes [5, 6].

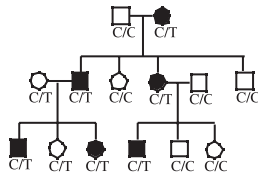
The other major type of genomic variation arises from discrete changes in a specific DNA sequence, including small but unique base insertions or deletions but more frequently single nucleotide substitutions, also known as single nucleotide polymorphisms (SNPs). SNPs are the most abundant form of DNA sequence variation in the human genome, occurring approximately 1 in every 1000 bp when any two copies of a chromosome are compared. Based on their high natural frequency in both coding and non-coding regions, single nucleotide substitutions are probably the molecular basis of most genetically determined phenotypes. Therefore, the identification of SNPs will play an increasingly important role in analyzing genotype–phenotype correlations [7].

### 3.3

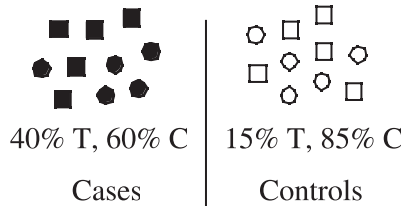
#### Linkage analysis

Linkage analysis looks at the correlation between trait and genetic marker within a pedigree (Figure 3.1). Linkage analysis for human traits blossomed in the mid-1980s and early 1990s with the availability of low density genetic maps of the human genome. The earliest low density maps were composed

## Linkage Analysis



## Association Studies\*



Correlate trait and marker allele within a family	Correlate trait and marker allele within a population
~300 marker map	30,000-500,000 marker map
Optimal for Mendelian Traits	Optimal for complex traits

\* Statistics shown are for the ApoE4 polymorphism and late onset Alzheimer's Disease[92]

**Fig. 3.1**  
Strategies for genetic analysis.

of several hundred restriction fragment length polymorphisms (RFLPs) but these were rapidly replaced when more informative microsatellite maps became available. Application of low density maps in linkage studies led to the identification of regions containing genetic variants which cause hundreds of rare Mendelian diseases, such as cystic fibrosis and Huntington's Disease [8].

Genetic linkage is a phenomenon where genotype at two markers is not inherited independently within a pedigree. This reflects proximity between markers on a chromosome such that the probability that recombination will occur between the markers during meiosis is less than 50%. Linkage regularly extends across tens of megabases in the human genome. Assuming two loci (A and B), each with two alleles (A1/A2 and B1/B2 respectively), a parent heterozygous for both loci has 25% probability of transmitting each of the following pairs of alleles: A1B1, A1B2, A2B1, or A2B2. However, if the loci are linked some allelic combinations will be more likely than others. The specific allelic combinations which are favored will be specific to each pedigree and not consistent throughout the affected population.

In linkage analysis researchers attempt to identify non-random patterns of inheritance within pedigrees for a phenotype and any single polymorphic marker from a set of markers spread relatively evenly across the genome (a low density genetic map). The phenotype is used as a proxy for genotype at the functional variant. Thus, statistical power to detect linkage is greatest for phenotype strongly correlated to underlying functional variation. Other factors affecting statistical power in linkage analysis are recombination between assayed marker and functional variant (less is better) and number of alleles at the assayed marker (more is better).



In classical linkage analysis, for each trait studied it was necessary to ascertain extended pedigrees in which the trait was segregating before linkage could be established. More recently, a number of strategies have been developed for using smaller pedigrees such as affected sibling pairs for mapping traits of biological or medical interest [9]. There are a number of programs available for linkage data analysis based on study design and complexity of the phenotype being explored (<http://linkage.rockefeller.edu/soft/list.html>) [10].

Once a trait is mapped, positional cloning allows researchers to identify the causative molecular variants of genes within linked regions by identifying functionally important variation which segregates with the trait. After linkage analysis is used to identify a critical region of the genome which contains a functional variation, a physical map consisting of clones of the DNA spanning the critical region is constructed. Candidate genes are identified from the sequence data of the overlapping clones which make up the physical map. The candidate genes are then surveyed in the linked pedigrees for functional variants which are infrequent in the general population and/or make sense in light of known biochemistry of the disease pathology.

Positional cloning has led to remarkable insight into the pathology of Mendelian disorders. The increasing availability of sequences across the human genome is greatly enhancing the speed with which candidate genes are identified and functional variants described. Thus, many projects are turning directly to databases to collect information about the sequences and candidate genes found in a disease-linked region to identify causative functional variation [8]. Unfortunately, the success of low density sets of polymorphic markers in mapping rare Mendelian diseases with high phenotypic penetrance has not extended to linkage scans for common genetic disease (e.g. diabetes [11], hypertension [12], asthma [13], multiple sclerosis [14, 15]).

Relative risk is a measure of the probability that an individual with a given genotype will develop a phenotype, relative to the probability that any individual in the population as a whole will develop a phenotype. Although twin and adoption studies consistently demonstrate that most common diseases have a significant heritable component, linkage analysis has generally failed to identify loci with high levels of relative risk (greater than four-fold increase in risk). The statistical power of linkage analysis is greatest for detecting high relative risk loci, where variation in a single gene is both necessary and sufficient to cause disease, such as cystic fibrosis and Huntington's Disease. Linkage analysis is much less powerful for detecting genetic variants conferring low relative risk, even when such variants are common [16]. The failure of linkage analysis to identify major genetic components of common disease has led many groups to believe that multiple loci are involved, each conferring a low level of relative risk. Low levels of relative risk can also reflect interaction between genotype at multiple loci

(epistasis), interaction between genotype and environment, or truly stochastic factors such as X chromosome inactivation or allelic exclusion.

### 3.4

#### Association analysis

Linkage disequilibrium (LD) is a phenomenon where genotype at two markers is not inherited independently within a *population*, not just within pedigrees. This reflects much closer proximity between markers on a chromosome than linkage. LD rarely extends further than 100 kilobases. As with linkage, assuming two loci (A and B), each with two alleles (A1/A2 and B1/B2 respectively), a parent heterozygous for both loci has 25% probability of transmitting each of the following pairs of alleles: A1B1, A1B2, A2B1, or A2B2. However, if the loci are in linkage disequilibrium, some allelic combinations will be more likely than others throughout the population. In association analysis researchers attempt to identify non-random patterns of association throughout a population for a phenotype and an allele of a single polymorphic marker.

Association studies have greater power to detect variants with low relative risk than linkage studies, but require very dense sets of polymorphic markers that can rapidly be typed on large numbers of samples (Figure 3.1) [16]. A dense marker map is required because these methods rely on linkage disequilibrium, and linkage disequilibrium extends over much shorter regions than linkage. As in linkage methods, linkage disequilibrium methods assume that we do not know which polymorphism in a region is functionally relevant. While a functionally important polymorphism will show association with phenotype, other markers in linkage disequilibrium with such a functional polymorphism will also show association with the phenotype. The strength of the genotype-phenotype association will depend on both the level of linkage disequilibrium between the marker assayed and the functional polymorphism, as well as the relative risk associated with the functional polymorphism. Thus, the optimal number of markers required for a full genome scan using association methods will depend on the extent of useful levels of linkage disequilibrium in the genome.

Theoretical models suggest that maps as dense as one SNP every 3–6 kb will be necessary for association studies [17], which would mean a map with more than 500 000 SNPs. Experimental evidence suggests that useful levels of linkage disequilibrium regularly extend up to tens of kb, with significant variance across the genome [18, 19]. Nonetheless, even using a highly optimistic estimate of one SNP every 30 kb, an optimized map of 100 000 SNPs will be required for a full genome scan. Currently, this is far beyond the routine genotyping capacity of most laboratories, assuming several hundred

samples and 100 000 SNPs, or on the order of tens of millions of genotypes per study. Although genome-wide association studies are not yet feasible (from the standpoint of cost and throughput), with current assay capacities it is possible to rapidly perform meaningful analyses of candidate genes and these are generating new insights into common human genotype-phenotype associations [20, 21].

### 3.5

#### **Why is genetic analysis shifting to single nucleotide polymorphisms (SNPs)?**

While linkage analysis using low density genetic maps of repeat polymorphism markers has proven utility for simple Mendelian trait analysis, it is increasingly clear that association studies will be required to identify loci conferring modest relative risk for common genetic disease. Single nucleotide polymorphisms (SNPs) have become the practical marker of choice for large-scale association studies for a number of reasons:

1. Single nucleotide substitutions (the most common type of SNP) are the most abundant form of polymorphism in the human genome. Although their frequency does vary across the genome, conservative estimates suggest that SNPs with rare allele frequency greater than 5% occur once every 1000 bp. This translates into an extremely large and informative potential source of markers from the human genome: greater than 3 000 000 markers assuming a genome of approximately 3 billion bp [22–24]. In this regard, SNPs are the only type of polymorphism frequent enough to generate a marker set dense enough for whole genome association analysis [17].

2. SNPs are diallelic, which makes them easy to assay using a variety of automated approaches that have simple binary read-outs. Therefore, SNP genotyping can be fully automated, and the prospects for large scale genotyping are improving [24–26].

3. SNPs are found in both coding and non-coding regions of the genome. Detailed molecular analysis of the changes responsible for rare Mendelian traits has revealed that single-nucleotide substitutions are the most frequent form of disease-causing mutation [27]. Nucleotide substitutions in coding regions can lead to amino acid substitutions that affect protein function (e.g.  $\beta$ -globin and sickle cell anemia) or changes that introduce a stop codon and truncate the protein product (e.g. BRCA1 and early onset breast cancer) [27]. Many mutations with biological consequences have also been identified as nucleotide substitutions in non-coding regions of a gene, including splice site variants (e.g.  $\beta$ -globin and  $\beta$ -thalassemia), mutations in promoter regions (e.g. EPM1 and epilepsy) and remote regulatory regions such as in locus control regions [28]. Based on this evidence, it is reasonable to assume that frequent nucleotide substitutions with more subtle biological consequences are also present in the genome and may be associated with in-

creased susceptibility to common disease. At present, the limited data on common variable sites in genes makes these hypotheses difficult to test which has stimulated the development of SNP markers from the human genome.

### 3.6 SNP Discovery

A number of strategies are being applied to identify SNPs from the human genome; and most of these rely on gel-based sequencing for either direct discovery of SNPs or confirmation of SNPs from indirect methods. In direct discovery strategies, overlapping sequences from multiple individuals are computationally aligned to identify high quality mismatches [29–33]. At these sites, information about alignment depth, sequence context, and read quality are used to predict a statistical likelihood that the site is an actual polymorphism and not a sequencing error. This process of ‘*in silico*’ SNP detection has been augmented tremendously by the development of rigorous base calling quality analysis programs such as *Phred* [34, 35], that determine quantifiable differences in the quality of the sequence reads. Several programs are available to automate SNP detection in sequence traces, including PolyBayes (<http://polybayes.wustl.edu/>) and PolyPhred (<http://droog.mbt.washington.edu/>).

Using the primary data generated by the Human Genome Project 45% of the 1 500 000 SNPs presently in the dbSNP database (<http://www.ncbi.nlm.nih.gov/SNP/>), were identified *in silico* from overlapping regions of genomic clones. These SNPs are clustered in 10–50 kb regions, but the clusters are fairly randomly distributed across the genome. A much smaller fraction of dbSNP (~4% of all dbSNP entries) comes from similar data mining efforts using single pass sequence reads from cDNAs (known as ESTs) as the raw data for *in silico* detection [36–38]. Because the EST SNPs are clustered in processed transcripts, they are more likely to occur in coding regions which can be tested for functional consequences. However, many other molecular mechanisms exist for the control of transcription, RNA splicing, and mRNA stability that can affect the final expressed phenotype. Thus, SNPs occurring in noncoding sequences such as 5′ promoter sequence, splice site junctions, and regulatory/enhancer elements (intronic, 5′, and/or 3′ regions) can also have significant functional effects [39].

Most of the remaining SNPs in dbSNP have been identified by the SNP Consortium using Reduced Representation Sequencing [40]. In this approach, pooled DNA from a group of unrelated individuals is digested extensively with restriction enzymes and size fractionated on an agarose gel. Fragments approximately 1.5 kb in size are cloned, representing the same small percentage of the genome (~1%) from all individuals. The clone

library is shotgun sequenced to 2–5× coverage and the sequence traces aligned and compared to screen for mismatches (SNPs) where clones happen to overlap.

The bulk of SNPs currently available come from large clone overlap and reduced representation sequencing. Although studies using these approaches have proven useful in identifying SNPs en masse, the major drawback of random clone overlap strategies is that they do not provide information on the population frequency of alleles at each SNP. The relative frequency of a SNP can significantly affect statistical power to detect associations through linkage disequilibrium mapping. In addition, estimates for the allele frequencies of these sites are unreliable because many SNPs are identified based on the overlap between just two clones, which represents a comparison of just two chromosomes. The population minor allele frequency of such SNPs cannot be predicted; and it varies from very rare to 50%. Therefore, all the sites must be genotyped in additional samples to determine which ones would be useful in association studies. Furthermore, many modestly frequent SNPs are missed because so few chromosomes are being compared.

To increase the number of chromosomes compared during polymorphism discovery and to directly obtain genotypes as well as allele frequencies, targeted SNP discovery methods rely on DNA amplification by PCR to obtain similar regions of the genome from multiple individuals. Once a specific region is amplified by PCR it is possible to directly sequence the region across many individuals and compare for polymorphism as in the direct discovery methods above. Alternatively, there are many indirect techniques available to scan for SNPs in PCR products obtained from multiple individuals. Most indirect approaches for SNP identification are two-tiered, where SNPs are indirectly detected and then verified by direct DNA sequencing to identify the nature and location of the SNPs found during the primary scan [41–44]. Indirect detection methods include denaturing gradient gel electrophoresis [41], denaturing high performance liquid chromatography (DHPLC) [43], chemical or enzymatic heteroduplex cleavage [45–47], single-stranded conformational polymorphism analysis (SSCP) [48, 49], and gel migration heteroduplex analysis [48, 49]. DNA sequencing verification is important because nearly all the approaches for high throughput genotyping require sequence context information of a SNP for assay development [24, 25]. Therefore, direct high throughput sequencing remains a popular SNP discovery tool, even for targeted SNP discovery [50–52].

### 3.7

#### **Genotyping technologies**

In general, SNP discovery and SNP genotyping take place on separate technological platforms. This is because SNP discovery technologies are

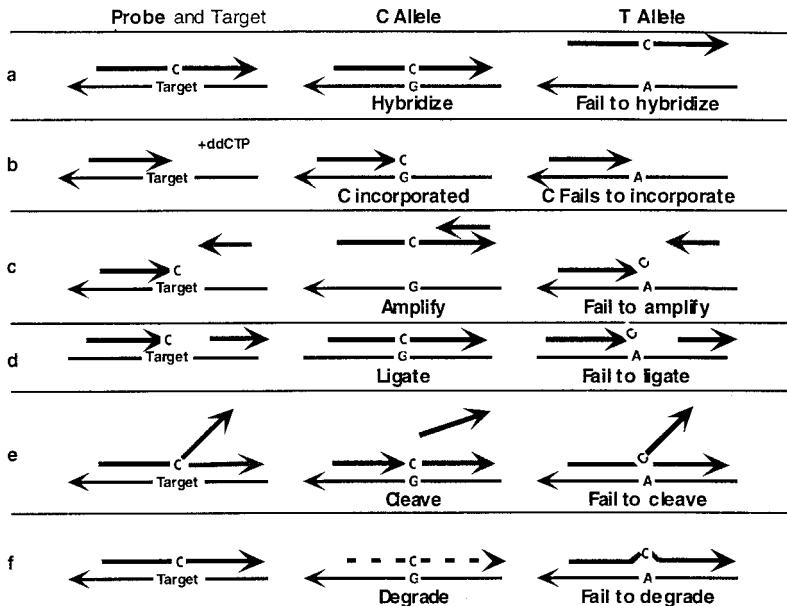
optimized to identify variation at any position in a given target sequence for a few individuals, while SNP genotyping technologies are optimized to detect variation at a single position in many individuals. While some technologies (e.g. DHPLC and SSCP) may be used for both discovery and genotyping, they tend to be sub-optimal in both throughput and sensitivity.

The ideal genotyping system for SNPs should be robust, cheap, homogeneous (i.e., a self-contained assay requiring little or no processing for sample analysis), highly multiplexed (i.e. many SNPs tested in the same assay) and fully automated. Unfortunately, none of the existing technologies currently meets all of these goals. However, many formats provide sufficient throughput to generate thousand of genotypes per day in a semi-automated fashion. Costs for SNP genotyping currently range from 20 cents/SNP to 80 cents/SNP (even the most highly multiplexed format, the human SNP chip, costs approximately 60 cents/genotype) and remain a factor in scaling assays to level required for a whole genome scan. This is stimulating work on the development of highly multiplexed and miniaturized formats, that make genotyping of large numbers of SNPs feasible and cheap [24, 53, 54]. However, until more scalable platforms for genotyping arrive, SNP analysis in candidate genes and pathways containing such genes is likely to remain at the forefront of genotype-phenotype analysis in mammalian genomes.

The most basic genotyping assays simply rely on differential hybridization of a pair of allelic probes to amplified target (Figure 3.2a). These include allele specific oligonucleotide assays (ASO) [55] and reverse blot assays [56], which differ only in terms of whether the target or the probe is immobilized. Newer solution based hybridization systems based on hairpin probe assays (e.g. molecular beacons) are also emerging [57]. All hybridization assays must be performed under conditions that favor perfect match binding over mismatch binding. In most cases, the polymorphic site is placed centrally within the probe. Genotype is determined by examining which probes bind perfectly matched target, and assay specificity depends on the hybridization kinetics of the probe/target system. Optimization of each hybridization system requires significant effort, but subsequent genotyping is quite efficient, so these technologies are optimal for studies of a small number of SNPs in a large number of individuals.

Minisequencing is a genotyping method where unique primers anneal immediately 5' of the SNP and initiate a single-base extension using DNA polymerase and labeled dideoxynucleotides (Figure 3.2b). Genotype is scored by determining which dideoxynucleotides were incorporated at the polymorphic site, and specificity is determined by the accuracy of nucleotide incorporation by the polymerase [58, 59]. This technology appears to require the least optimization, and therefore should be optimal for large-scale studies with many hundreds of SNPs [53, 60].

Allele specific PCR uses three primers: one allele specific primer for each allele with the polymorphic site at the 3' end, and a common reverse primer (Figure 3.2c) [61, 62]. Using a polymerase without 3'-5' proofreading activity, samples are amplified twice, once using each allele specific primer.



**Fig. 3.2**  
SNP genotyping technologies.

Genotype is scored for each sample based on which of the allele specific amplifications were positive. Specificity is determined by the amplification kinetics of each assay and the differential efficiency of the polymerase in extending 3' matched and 3' mismatched primers. This assay format fails to multiplex and requires significant optimization of each assay, but uses cheap unmodified oligos and relatively cheap instrumentation.

Oligonucleotide ligation assays (OLA) [63] also use three specific primers to type a SNP with two allele-specific oligonucleotides and one joining oligonucleotide (Figure 3.2d). After the oligos have had time to hybridize to the target sequence, a DNA ligase is used to join the adjacent oligos (an allele-specific with a joining oligonucleotide). The fusion occurs only if the 3' nucleotide at the ligation site complements the target DNA site. Genotype is scored by determining which of the allelic ligation reactions were successful. OLA enables multiplex genotyping of a large number of individuals for multiple SNPs [64] but requires expensive custom oligonucleotide syntheses.

Flap cleavage (Invader, Figure 3.2e) [65] and 5' nuclease discrimination (Taqman, Figure 3.2f) [66] are both assays that use enzymatic cleavage to identify the presence of a SNP in a sample. During flap cleavage, an enzyme is used to cleave the structure formed when two overlapping oligonucleotides hybridize to a target DNA strand, and cleavage is dependent on a 5' match between the cleaved oligo and target. This assay requires rela-

tively cheap instrumentation and limited optimization, and can achieve reasonable levels of throughput. Alternatively, in the 5' nuclease assay a complementary probe bound to the target DNA strand is cleaved if and only if it is perfectly complementary to the SNP site. Using oligonucleotides complementary to both the wild-type and the polymorphic base, the presence of both kinds of target DNA (both SNP alleles) can be detected even in a complex mixture. The 5' nuclease assay requires expensive fluorescent oligos, and frequently requires optimization, but provides excellent throughput once assays are developed.

Colorimetric or fluorescent readout from genotyping reactions is generally performed in microtiter plates, although some array technologies are showing great promise for highly multiplexed readout [53, 60], and mass spectrometry readouts are showing promise for very high throughput low-multiplex genotyping [67, 68]. Whichever assay is used, the readout from all SNP genotyping technologies is a binary score for whether a given allele is present or absent. Combining the binary scores from both alleles of a SNP yields genotype for a sample.

### 3.8

#### **SNPs are frequent in the human genome and their organization complex**

New information is emerging from the analysis of single SNPs and patterns of SNPs in candidate genes. It is revealing that the amount and organization of the sequence variation found in genes is generally more complex than expected [52, 69–71]. For example, in a gene of average size (30 kb) with average sequence diversity (0.1% sequence variation between any two copies of the gene) more than 100 SNPs with minor allele frequency above 5% are likely to be present. The vast majority of these sites will not lead to amino acid substitutions in the protein product of the gene, either because they are in non-coding regions (introns) or because they are at redundant positions of codons (synonymous SNPs). Although polymorphisms that lead to amino acid changes (known as cSNPs or nonsynonymous SNPs) can clearly lead to functional changes in the gene product, only a small portion of all SNPs in genes will occur at nonsynonymous sites. Nonsynonymous SNPs also generally have lower minor allele frequencies than synonymous and intronic SNPs [44, 72], which has been hypothesized to represent greater selective pressure on amino acid substitutions. Therefore, although cSNPs can be directly tested for function, their use as markers for association to other functional sites may be limited because of their relatively low frequencies in the population [44, 72].

Also, many other molecular mechanisms exist for the control of transcription, RNA splicing, and mRNA stability that can affect the final expressed phenotype. SNPs occurring in noncoding sequences such as 5'

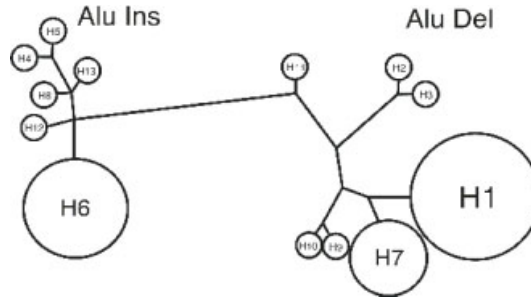


promoter sequence, splice site junctions, and regulatory/enhancer elements (intronic, 5', and/or 3' regions) can have functional effects that need to be evaluated since little is known about the impact of this type of sequence variation [39, 73, 74]. Since we cannot predict which variants may have functional significance with regard to common phenotypes, it is difficult to select the sites that need to be examined. This combined with the fact that there are so many common sites present in every gene makes the issue of site selection for large-scale genotyping an important one.

Although it is possible to measure the association between genotype and phenotype for any single site in a gene, such studies are hardly conclusive. First, in these studies it is likely that many real effects due to variants in a candidate gene are missed because the wrong SNP was assayed. Second, spurious relationships can be identified due to such factors as population stratification and inadequate correction for multiple testing. A number of recent studies have shown that even in the most complex areas of the genome, a greater understanding of the organization of the variable sites within genes can help define the sites that maximize the potential for uncovering associations between genotype and phenotype [52, 70, 75]. However, this requires prior knowledge of SNP frequency and linkage disequilibrium relationships in the population(s) of interest. One way to describe the patterns of linkage disequilibrium within a region is to use haplotypes. A haplotype is the pattern of alleles at multiple loci along a single chromosome. Given  $n$  diallelic loci there are  $2^n$  possible haplotypes. However, for markers showing significant linkage disequilibrium far fewer haplotypes will actually be observed in the population, and it is fairly straightforward to derive the evolutionary relationships between the observed haplotypes. For site selection, much of the information needed is available in the haplotype structure of a gene within a population [76]. An example of this is the angiotensin-converting enzyme (ACE). This enzyme catalyzes the conversion of angiotensin I to the physiologically active peptide angiotensin II, which controls fluid-electrolyte balance and systemic blood pressure. Because of its key function in the renin-angiotensin system, many association studies have been performed with ACE, and because it is technically easy to genotype, most of these studies have used an alu repeat element insertion polymorphism marker. Many studies have associated the presence (insertion, I) or absence (deletion, D) of the alu element with levels of circulating enzyme or cardiovascular pathophysiology [77–80]. Many epidemiological studies suggest that the D allele in ACE confers increased susceptibility to cardiovascular disease; however, other reports have found no such association or even beneficial effects [81–83].

When the sequence polymorphisms in ACE are resolved into haplotypes and a gene tree constructed [52], the pattern of variation in ACE reveals the presence of two major clades that are detected by the Alu indel (Figure 3.3). Because the deletion clade has been consistently associated with cardiovascular disease pathology including myocardial infarction, arterial hyper-

**Fig. 3.3**  
A consensus parsimony gene tree for ACE haplotypes. The relative frequency of each haplotype is indicated by the area of the circle.



tension, and left ventricular hypertrophy [79], this subdivision could be of potential medical importance.

The haplotype structure in a variety of genes has shown distinct clustering with only a few major clades [52, 69]. These results are consistent with evolutionary theory, which indicates that polymorphic sites are generated in a stepwise mutational fashion producing closely related haplotypes. Evolutionary trees based on haplotypes should be informative for generating a minimal panel of polymorphic sites that identify major substructures or relationships among the variable sites [76].

Thus, linkage disequilibrium (LD) mapping is likely to be an iterative process that requires site selection at different stages of the process. In the initial stages of a candidate gene scan LD mapping can be used to test phenotypic associations with the major structure of a tree (such as the sites associated with the I/D polymorphism in ACE). To fine-map an association, sites associated with subclades should prove useful, as with sites distinguishing the two major haplotypes within the ACE deletion clade, H1 and H7 (Figure 3.3).

### 3.9

#### Pooling strategies

Because of the required number of assays, a full genome association study is not a tractable problem with any current genotyping technology. One way to reduce the number of assays required for a full genome scan is to multiplex samples instead of markers. In the simplest pooled study design two sample pools are generated, representing affected and unaffected individuals [84, 85]. After accurate measurement of allele frequency in each pool, the frequencies are compared for significant differences. The magnitude of the allele frequency difference is dependent on several parameters of the risk allele: allele frequency, mode of inheritance, and relative risk. It is also important to note that genetic interaction between loci and haplotype spe-

cific risks will confound the power of pooling analysis, although the likelihood of these effects remains to be determined.

Pooling samples requires a quantitative readout of genotype frequency, and the accuracy and precision of allele frequency measurements will have significant impact on the power of pooled study designs to detect disease associated loci. At the very least, the error in pooled frequency estimates should be less than 2% across a broad spectrum of allele frequencies. Promising preliminary results have been published for pooled frequency estimation using mass spectrometry minisequencing [86], DHPLC minisequencing [87], SSCP [88], allele specific PCR [89], chip array minisequencing [53], and even direct sequencing [90].

The utility of pooled approaches remains to be proven, especially in light of recent studies which suggest that the relative risk associated with a SNP is context-dependent on local haplotype [20, 91]. Haplotype information is lost in pooling strategies because the chromosomes are not observed independently; and consequently haplotype specific effects would be missed.

### 3.10

#### Conclusions

Although hundreds of mutations causing rare Mendelian diseases have already been identified, the impact of sequence variation analysis on understanding the patterns of normal and pathologic variation in human phenotypes has just begun. Using current technologies it is possible to comprehensively study the impact of sequence variation in candidate genes on phenotype. This will facilitate the process of weeding through the weak linkage hits and biochemically implicated genes to identify those genes with significant effect on medically important phenotypes, for both complex genetic disease and drug safety/efficacy. The development of new technologies for very high throughput single nucleotide polymorphism identification and genotyping, well characterized dense genetic maps, and analytic tools for whole genome association analyses are likely to provide new insights into the molecular basis of common genetic disease.

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## 4

**Proteome analysis**

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## 4.1

**Introduction and principles**

Today, genome sequencing projects are generating, with very high throughput capabilities, a huge amount of information in the form of nucleotide sequences. These sequences are being stored in specific databases. This information has to be analysed in order to complete its annotation. Annotation means description of the coding sequences, of the precursor elements, of the position of the DNA recognition sites, etc. In addition to this structural information, functional data has to be annotated, such as levels of expression of the genes, organ specificity, functionality of the genes, etc. The quality of this information represents a crucial aspect in the study of the expression products of the genes, i.e. the proteins.

Proteomics covers the study of the proteins expressed by a genome in a biological sample, such as an organism, an organ, an organelle, a biological fluid [1, 2]. This involves the identification of the proteins contained in this sample. The complete sequences of active proteins can only be partially deduced from their corresponding gene sequence. In fact, during and after the transcription and the translation processes leading to an active gene product, alterations often occur, such as alternative splicing, N-terminal truncation, post-translational modification (PTM)<sup>1)</sup>. Therefore, proteomics involves

- |   |   |
|---|---|
| <p>1) Post-translational modifications are affecting the protein sequences between the time they are produced by the ribosomes and the time they are biologically active, that means after the translation. The amino acid sequence as obtained by a translation of the coding sequence of the corresponding gene can be modified in different ways. Beside alternative splicing and N-terminal truncation, some amino-acids of the</p> | <p>proteins can be modified by the addition of chemical entities. Some of them are called homogeneous, as the added functional group is of discrete and unique mass. Phosphorilation is, in that respect, the addition of a phosphate group to the side chain of an amino-acid, such as typically serine or tyrosine. Others are methylation, sulfation, farnesylation, and so one. Others are named heterogeneous, as the chemical</p> |
|---|---|

also the description of the events generating the modifications these proteins carry as functional entities.

In addition, the function and the activity of a protein depend on its concentration and more generally on its environment. To be active, a protein might have to interact with other proteins, with non-peptidic cofactors, with substrate molecules, at certain pH; it might have to be transported from a cell compartment to another, its PTMs can dynamically be controlled as function of the cell state, etc.

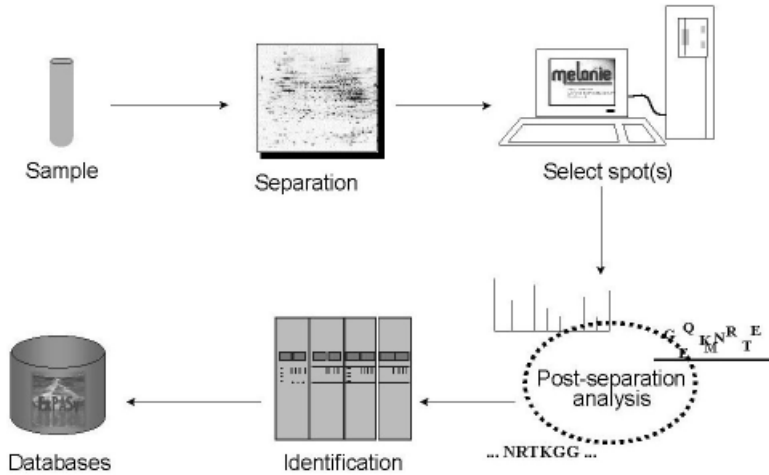
Proteomics has to deal with much higher complexity than genomics. Proteins occur in very variable concentrations: the dynamic range between transcription factors and albumin can be at least  $10^{12}$ . Single genes can be expressed in more than 20 protein forms in a single tissue: as an example, at least 22 different protein forms matching alpha-1-antitrypsin have been described in human plasma. The physico-chemical properties of proteins are extremely variable. Their molecular masses range from a few thousand to over a million Daltons. Solubility and isoelectric point<sup>2)</sup> (pI) can vary a lot in proteins, and their load of PTMs can be very heterogeneous. Therefore, the analysis and the understanding of the suspected half a million to one million proteins in human, expressed by a number of genes that is estimated to be around 40 000, represent a real challenge, both from the technological and the bioinformatics point of view. Methods involving high-resolution protein separation, parallelization of sample preparation, automation of experimental processes and of database comparison, as well as powerful and specific visualization tools need to be developed and integrated.

The functions of the identified and then characterized proteins have also to be determined. Proteins often interact with each other in protein complexes, or with other important chemical entities such as DNA in protein–DNA interaction systems. Their activities involve also the interaction with smaller molecules such as cofactors or hormones. Therefore a number of methodologies have to be adapted to understand the behavior of proteins in their biological environment and the implications of a molecular dysfunction in a disease. The development of new diagnostic tests or the discovery

group added can be made of chains of subgroups, and therefore can have very variable masses. These are glycosylation and lipidation. There are currently more than 200 different PTMs described in the literature and annotated in the SWISS-PROT protein database (<http://www.expasy.org/sprot/>). The PTMs are functionally very important as, in a number of cases, they can trigger the activity of a protein, or they can help the binding of a protein to another one in a

complex, or they can be of crucial importance in the folding process of a protein. This is particularly the case of the disulfide bonds, that bind covalently the sulfur atoms of two cysteine residues.

- 2) The protein isoelectric point (pI) value is the pH at which the sum of acidic amino-acid negative charges and of basic amino-acid positive charges equals zero. It ranges between less than 3 and more than 12.



**Fig. 4.1**  
Schematic representation of a proteomics analysis pathway.

of new drug therapies in the future will depend on the capacity to analyze complex systems. Proteomics offers the possibility to observe the global influence of a drug in a complex mixture of proteins. This will be possible only if one can describe the identity, the occurrence and the interaction of each individual component of this mixture.

The analysis of a proteome, described as the ensemble of the proteins expressed by a genome in a given tissue, for a given organism at a given time, requires to use and to combine a number of procedures, both experimental (wet-lab experiments) and bioinformatics (dry-lab experiments). Due to the chemical and physical complexity of proteomes, various methodological approaches have to be considered. Nevertheless, a consensus principle of proteome analysis can be described as in Figure 4.1. This linear pathway includes most of the wet- and dry-lab steps required for the complete analysis of a proteome.

- First the sample is to be chosen in an adequate manner. It can be a raw biological fluid, a cell extract, a pre-fractionated sample, etc. The choice of the sample is crucial, as it is strongly dependent on the type of separation to be applied. The sample has to be compatible with the complexity both in number of components and in the dynamic range the separation can handle. Methodological aspects are discussed in Section 4.2 of this Chapter.
- The proteins contained in this sample have to be separated. In proteomics, one-dimensional electrophoresis (1-DE) and mainly two-dimensional

- electrophoresis (2-DE) techniques are preferred. The technique will be described in Section 4.2 [3, 4].
- The next step is the analysis of the separation and the choice of the proteins to be identified. The separation result is visualised using image analysis software. This software allows for visualizing gel images, for comparing gel images, and for performing a number of comparative analyses that enable tracking statistically significant changes in protein expression between populations of gel/samples. It helps the user to choose protein spots of interest. The description of the computer analysis of gel images will be extended in Section 4.3.
  - To proceed further, the selected proteins have to be submitted to a post-separation analysis. This step requires experimental determination of highly specific protein attributes, such as amino acid composition, sequence information, or peptide mass fingerprints following endoproteolytic cleavage. In endoproteolytic cleavage the proteins are typically incubated with an enzyme that recognizes particular amino acids and cuts specifically the polypeptide chains. The reaction produces shorter peptides that are fragments of the so-called digested proteins. Today's standard procedure involves a protein digestion step with trypsin and the analysis of the obtained fragments generated with a crucial tool in proteomics, mass spectrometry, and more precisely Matrix Assisted Laser Desorption Ionization (MALDI) mass spectrometry or Electrospray Ionization (ESI) mass spectrometry<sup>3)</sup> [5].
  - These protein attributes are then submitted to a database search. This search identifies a protein by looking at the best match between experimental data and data obtained by in-silico processing and “digestion” of a protein sequence database. The identification and characterization procedures using bioinformatics tools will be the topic of Section 4.4.
  - Comprehensive sequence databases are a prerequisite for successful protein identification, and the identified protein data is in turn used to
- 3) MALDI and ESI refer to two different possibilities of ionising molecules in a mass spectrometer. In a MALDI-TOF mass spectrometer (MS), The analyte is mixed with a solution of usually an aromatic organic acid (a matrix solution). The dried mixture is apposed on a metal plate. A laser shoots on the formed crystals and its energy, that is thought to be transferred to the analyte through the matrix, ionises the target material. The ions are accelerated under an electric field in a vacuum tube. The record of the mass over charge value of the ions is made by measuring the time of flight (TOF) needed by the ions to reach the detector. In an ESI MS the analyte is reaching the machine in solution through a capillary. At the end of the capillary the droplets are dried and ionised under an electric field. The ions are accelerated in an analyzer, that can be a TOF, a quadrupole or an ion trap. A quadrupole is a system that deviates the ions under a perpendicular electric field on an ionizing detector. The deviation is function of the mass over charge of the ions and of the strength of the field. An ion-trap keep the generated ions in a small volume under an well controlled electro-magnetic field. They are then released specifically on a detector.

populate specific secondary databases. A short description of proteomics databases will be the topic of Section 4.5.

This linear representation of a proteome analysis pathway represents and orders the different steps required to perform the identification of proteins from crude biological sample. This pathway can be applied as a way to generate a systematic description of a proteome. The result of such an analysis is often concretized by the creation of an annotated database. Databases like SWISS-2DPAGE [6] and other federated 2-DE databases are typically populated with the result of systematic identification of all protein spots observed on a 2-DE image. From all possible combinations of methods currently available, a gold standard is widely adopted. It combines the very high separation power of 2-DE and the very high sensitivity and accuracy of mass spectrometry. However, while it is agreed that most of the proteins can be separated with this technique, this approach does not allow for identifying and characterizing all proteins present in a biological sample. Some proteins will not appear on the gel because of their extreme physico-chemical properties, such as exceedingly high or low molecular weight, exceedingly high pI, extreme hydrophobicity or insufficient abundance. Other techniques should therefore be applied to complete the picture, such as one-dimensional electrophoresis, pre-treatment of the starting material, pre-fractionation, early digestion of the protein mixture followed by RP-HPLC, and others. Some of these methods are described in [5].

Proteome analysis is however not limited to the systematic identification of all proteins present in a biological sample. The described pathway allows the creation of databases useful for being searched or compared with other real samples. This information can be used to compare various samples. In this case, the annotated 2-DE image is being compared with a non-annotated image in order to correlate positions and intensities of protein spots. From all the spots in the studied image only some spots of dedicated interest can then be chosen and further analysed with identification methods. A very widely used method to search for biological markers of specific diseases is the comparison of a statistically significant number of 2-DE images from samples of healthy and diseased patients or samples, treated or not treated with target drugs [7]. The images are compared, clustered and a number of protein spots appear to be differently expressed. These become the spots of interest to be further identified. In this approach most of the effort is concentrated in the generation of the 2-DE gels and in the comparison of the generated images. Section 4.2 will discuss these aspects with an application in diabetes. Section 4.3 will describe the possibilities of proceeding such an analysis using dedicated software.

In addition, beyond the identification efforts, it might be of highest interest to describe and understand the modifications carried by the active gene products. This implies the search for splicing variants, amino acid

mutations or PTMs. Mass spectrometry is very often used in a so-called MS/MS mode<sup>4)</sup> to unravel these difficulties [5, 8]. The interpretation of the spectra generated by this process reveals structural information on the amino acid sequence and the description of PTMs attached to the studied peptides. Besides mass spectrometry, other methods allow to highlight the presence of specific PTMs in proteins, such as immunological staining (also called immunoblotting). In this latter approach, antibodies specific to particular PTMs can be used to visualize the presence of PTMs in proteins separated by electrophoresis methods (see also below, in Section 4.2).

As the number of proteins to be identified in a single sample can reach a few hundreds to a few thousands, it becomes necessary to automate the whole identification process. The goal of automation is to yield a higher throughput as well as a better reproducibility of the experimental procedure. This requires to develop sample tracking procedures specific to proteomics, to create new bioinformatics tools to pipeline the data management, and to implement a data management system allowing to store the huge amount of information generated, and to be queried for later analyses. Section 4.6 will cover these automation aspects, together with the presentation of an alternative to the standard robotised approach of protein identification, known as a molecular scanner [9, 10, 11].

## 4.2

### Protein separation

Qualitative and quantitative comparison of proteomes under different conditions can now help to further unravel complex biological processes. This is due to drastic improvements in massively parallel protein separation, identification and characterization as well as in bioinformatics.

#### 4.2.1

#### Experimental aspects: description of the 2-DE technology, a wet-lab technique, from a wet-lab point of view

The central tool today to display and evaluate the proteome complexity of any organism is 2-DE independently developed by O'Farrell and Klose in

4) The MS/MS mode is a fragmentation mode. In such a mode, one peptide of interest is selected in a first MS separation. Then it is specifically fragmented in a cell that adds energy to the ions, by either physical shock with Argon or a pulsed electromagnetic field.

The peptides are then cleaved more or less specifically, depending on the fragmentation hardware, at the amide bond between amino acids. This produces series of ions, the mass of those are differing by a number of distinct amino acid residues.

1975 [3, 4]. It has been designed in a way that 2000–3000 polypeptides are separated on the basis of different molecular properties in each dimension; charge (pI) in a first dimension and molecular mass ( $M_r$ ) in a second. Isoelectric focusing (IEF) is the electrophoretic technique in which the proteins are fractionated according to their pI through a pH gradient. In the process of IEF with carrier ampholytes (CA), the pH gradient is created, and maintained, by the passage of an electric current through a solution of amphoteric compounds which have closely spaced pIs, encompassing a given pH range. Their electrophoretic transport causes the CA to stack according to their pIs, and a pH gradient, increasing from anode to cathode, is established. After this stacking process is completed, some CA still enter zones of higher or lower pH, by diffusion, where they are no longer in isoelectric equilibrium. But as soon as they enter these zones, the CA become charged and the applied voltage forces them back to their equilibrium position, and so on in a kind of pendulum movement. This phenomenon affects drastically the reproducibility of protein separation. In the process of IEF with immobilized pH gradients (IPG), the pH gradient, which exists prior to the IEF run itself, is co-polymerized, and thus insolubilized, within the fibers of a polyacrylamide matrix. This is achieved by using, as buffers, a set of weak acids and bases named Immobilines<sup>TM</sup>. When the electric field is applied only the sample molecules and any non-grafted ion migrate. Upon termination of the electrophoresis the proteins are separated into stationary isoelectric zones [12]. The majority of the laboratories running 2-DE gels are currently using 3.5–10 IPG strips (i.e. strips with a pH range from 3.5 to 10) to display a wide range of proteins. However, the use of 1 pH unit narrow range IPG (i.e. strips the pH range of which extends to 1 pH unit, like 5–6 for instance) permits higher protein concentration loading and the investigation of a larger proteome “window”. In fact, these IPG strips are also called “zoomed” IPG strips, as they allow a better separation of protein spots that are very near. In addition, the total amount of protein load can be increased. These strips allow for increasing the number of visible protein spots in a pH window, therefore also for increasing the total amount of protein forms to be detected. After the first dimensional separation, the IPG strips are transferred on classical vertical or horizontal SDS<sup>9</sup> slab gels designed by Laemmli et al. [13] to dissociate all proteins into their individual polypeptide sub-units. The most common dissociating agent used is the ionic detergent sodium dodecyl sulphate (SDS). In addition, heating at 100 °C in the presence of excess SDS and of a thiol reagent to cleave disul-

- 5) The folding of polypeptide chain regions into regular structures defines the secondary structure of a protein. The tertiary folding between these regions involves both covalent disulfide bonds and non-covalent bonds. The quaternary structure exists in proteins that contain several polypeptide chains “aggregated” on a multimer. Conditions that reverse the quaternary, tertiary and secondary structure are termed denaturant. Sodium dodecyl sulfate (SDS) is an anionic detergent that denatures proteins.



vide bonds denatures the protein mixture. Under these conditions, most polypeptides bind SDS in a constant weight ratio (1.4 g of SDS per gram of polypeptide). The intrinsic charges of the polypeptide are insignificant compared to the negative charges provided by the bound detergent, so that the SDS-polypeptide complexes have essentially identical charge densities and migrate in polyacrylamide gels of the correct porosity strictly according to polypeptide size. Today, silver staining<sup>6)</sup> is probably the most popular non-radioactive protein detection as it is 50-fold more sensitive than Coomassie Brilliant Blue<sup>7)</sup> or Amido black staining. However, other procedures that include S<sup>35</sup> or P<sup>32</sup> radiolabeled samples can also be used. Quantitative 2-DE image analysis should be performed with a specialised software such as Melanie 3 (see Section 4.3) that contains all the basic functions for 2-D PAGE image analysis as well as links to comprehensive proteome databases [14].

#### 4.2.2

#### The use of 2-DE as a tool towards diagnostics and disease description

The development of new diagnostic and predictive tools as well as successful therapies for complex polygenic diseases including diabetes, cancer and cardiovascular diseases will need the understanding of the fundamental biological mechanisms implicated in these disorders. This can be achieved under defined environmental conditions with strategies that combine genetic and proteomic tools. Proteome analysis has the ability to detect and identify polypeptides that correlate with disease states and further lead to the discovery of potential molecular markers and therapeutic targets. Furthermore, 2-DE analysis can display the pharmacological and toxic effects of candidate drugs on a disease process. There is a high relationship between drug treatment, protein expression and resulting physiological effects. Most of the time, pharmacological mechanisms entail the secondary regulation or modulation of gene product expression, in a similar way that complex disease processes alter global protein expression. From this, we can assert that the best drug should be the one that restores global protein expression of a disturbed organism to a normal state. In addition, it is quite unusual that a drug only modulates gene products implied in the disorder. Most of the time, it also causes perturbations in the expression of proteins that are not involved in the disease process. This leads to side effects of drugs. Leigh

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| <p>6) Silver staining of proteins after their separation by electrophoresis is based on binding of silver ions to sulfhydryl and carboxyl groups of proteins. The proteins are detected as black precipitate of silver. The sensitivity of this method is 20–100 times more sensitive than Coomassie blue staining.</p> | <p>7) Coomassie blue staining of proteins after their separation by electrophoresis is based on non-specific binding of Coomassie dye to proteins. The proteins are detected as blue spots on a clear background.</p> |
|---|---|

Anderson pioneered this approach by investigating the effects of Aroclor [15], methapyrilene [16] and peroxisome proliferators [17] on mouse liver proteins using 2-DE and therefore by comparing globally the changes in protein expression patterns.

Another example is described here. Type II diabetes (non-insulin-dependent diabetes mellitus, NIDDM) is a heterogeneous disorder with several metabolic abnormalities including hyperglycemia, hyperinsulinemia and dyslipidemia which are the results of the combination of insulin resistance (acquired) and impaired insulin secretion (genetic) [18]. The first approach for the treatment of patients with NIDDM is to reduce metabolic abnormalities through diet control and exercise [19]. When they fail, additional therapeutic approaches to improve glycemic control are used to increase insulin secretion and diminish insulin resistance. Thiazolidinediones (Thzs) are oral antidiabetic agents that improve insulin action and decrease plasma glucose, insulin and triglycerides content. Unfortunately, the mechanisms of action of Thzs are not well defined. As Thzs have higher effects in obese animals, we treated lean vs. genetically obese C57 Bl/6 mice<sup>8)</sup> with troglitazone. Pharmacological and toxic effects of troglitazone could be efficiently investigated simultaneously using 2-DE analysis. Figure 4.2 shows an enlargement of the silver stained 2-DE separation of pancreatic islets from treated and non-treated lean and obese mice. The surrounded polypeptide is over-expressed in both obese and lean treated animals. This might indicate a side effect of troglitazone on pancreatic islets.

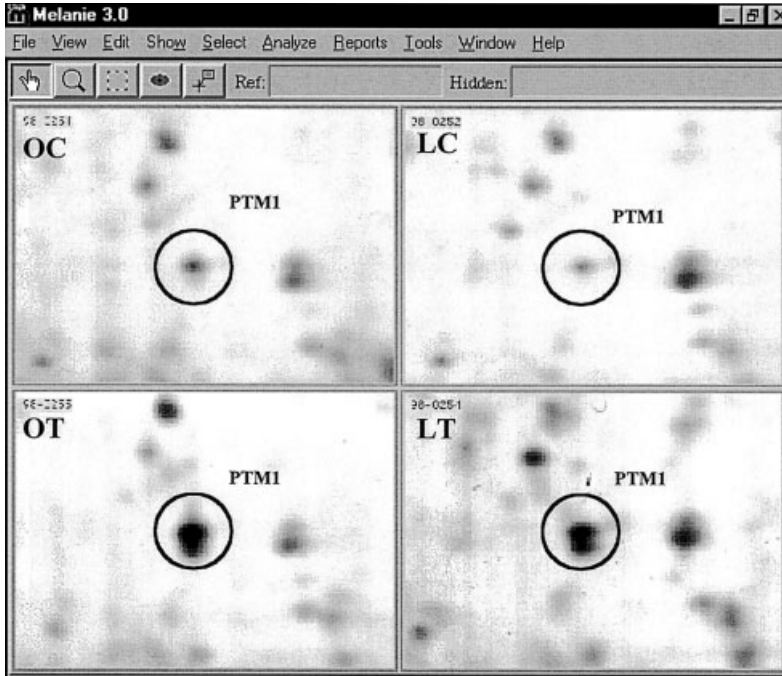
In general, the quality of a comparison of protein expression levels between two states depends on statistical facts. Variations of about up to 40% of spot volume can be expected between two gels of identical starting material. Therefore, changes that are lower than a ratio of 1 to 3 can be subjected to some doubt. To be able to describe small intensity changes with no ambiguity and with statistical significance, one ideally has to run 3–5 gels from identical material and sample. In a statistical analysis, one has also to choose more than one sample per biological state. One should select two (or more) populations of samples that will be run in several gels as starting material for a stable computer analysis that might reveal minor changes in protein expression patterns.

### 4.3

#### Computer analysis of 2-DE gel images

Once proteins in a sample have been separated by two-dimensional gel electrophoresis (2-DE), the main remaining task is to analyze the resulting

8) Name of one of the most widely used laboratory mouse model.



**Fig. 4.2**

Effect of troglitazone on pancreatic islets proteins expression. Comparison of the expression level of the polypeptide PTM1 in 2-DE gels under four conditions: **A) LC**: lean control c57Bl/6 female mice (125 islets: 95% of the islets were small, 5% were large), **B) LT**: lean treated c57Bl/6

female mice for one week with 300 mg/kg of troglitazone (135 islets: 95% small, 5% large), **C) OC**: obese ob/ob controls c57Bl/6 female mice (200 islets: 20% small, 80% large), and **D) OT**: obese ob/ob treated c57Bl/6 female mice for one week with 300 mgs/kg of troglitazone (200 islets: 40% small, 60% large).

2-DE images and to extract biologically relevant information. As researchers in the Life Sciences do not use the 2-DE technology with only one goal in mind, there is obviously more than one way to consider the analysis of 2-DE images. In general, one may consider three main approaches that vary in the globality or specificity of the proteins under study:

1. The creation of proteome maps. This is an important step towards the discovery of a full proteome. It is the systematic analysis of all (or as many as possible) spots contained on one 2-DE image of a given sample, in order to identify the corresponding proteins (see Section 4.4). The 2-DE image is then annotated with the proteins' names as well as additional related data, and the resulting protein map is added to a two-dimensional electrophoresis database, such as the SWISS-2DPAGE (<http://www.expasy.org/ch2d/>) [20, 6], the HSC-2DPAGE [21] or the Siena-

2DPAGE [22] databases. Such databases, often available on-line on the Web, allow for complex queries in order to retrieve data related to proteins and their location on the 2-DE maps [23]. Figure 4.3 shows the annotated 2-DE image of *Escherichia coli*, the related 2-DE image as it can be retrieved from SWISS-2DPAGE, as well as one of the textual database entries (in this case Elongation factor TU P02990). Most 2-DE databases available on the Web follow the rules for federated databases [24], that allow, from each entry in the database, to follow active hypertext links to the corresponding entries in other proteomic or genomic databases, such as the SWISS-PROT protein sequence [25] or the EMBL nucleotide sequence [26] databases (Figure 4.4) (see also Section 4.5).

2. The monitoring of the expression of one or a few gene products over a set of 2-DE gels. This approach encompasses the pharmacological or toxic effects of compounds on a disease process, either over a given time span or under various compound concentrations, as well as, more generally, when one is interested to measure the variation of the expression of a protein over a series of 2-DE gels of a given sample. To achieve such measurements, a set of 2-DE images is considered, and the analysis focuses on quantitative variations limited to the proteins of interest.
3. The global investigation of the differential expression of proteins between 2-DE images of two or more populations. A typical example is the identification of polypeptides that correlate with disease states. In this case two sets of 2-DE gels from disease versus control are compared to identify which proteins are consistently differentially expressed between the two sets of images. Alternatively, one compares samples from two or more disease states or from any biological conditions whose differential analysis might be of interest. In general, this global approach thus entails the comparison of the spot intensities in two or more (often large) sets of 2-DE images, leading to the further exploration of a few to several tens of gene products that could be of potential biological importance, such as, typically, potential disease markers or therapeutic targets. In given situations, researchers may also be interested in the investigation of only one set of 2-DE images, in order to discover patterns or subsets of samples that could possibly be linked to specific biological conditions or disease states. In any case, each variant of this approach necessitates the use of powerful data analysis tools, such as statistical programs to identify the proteins that are differentially expressed. This part of proteome analysis is very often followed by the identification and partial characterization of the proteins that have been pinpointed, using the techniques described in Section 4.4.

The three approaches of 2-DE analysis described above differ mainly in the number of 2-DE images that are considered in one given study and in the kind of image or protein comparisons that are performed. In the first approach, typically one 2-DE image is considered at a time, while in the

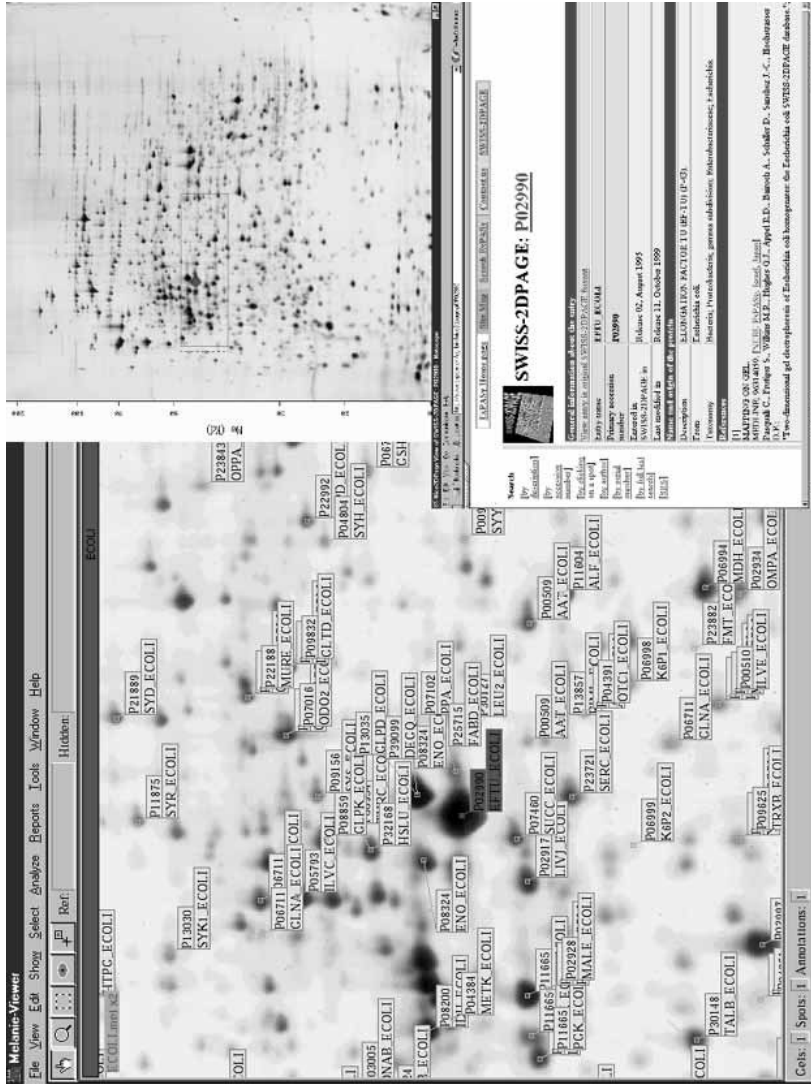


Fig. 4.3 Annotated image of a silver stained *E. coli* 2-DE sample in Melanie3 and related links to the SWISS-2DPAGE databases.

View of SWISS-2DPAGE: EFTU\_ECOLI - Netscape

Location: [http://www.expasy.org/cgi-bin/ht2d-search-ac?etu\\_ecol](http://www.expasy.org/cgi-bin/ht2d-search-ac?etu_ecol)

by full text search

[SES]

General information about the entry	
Entry name	EFTU_ECOLI
Primary accession number	P02990
Secondary accession number(s)	Nonc
Entered in SWISS-PROT in	Release 01, July 1986
Sequence was last modified in	Release 1, March 1989
Annotations were last modified in	Release 3E, July 1999

Name and origin of the protein	
Description	ELONGATION FACTOR TU
Gene name(s)	(TUFA OR B3339) AND EF-TU
From	<i>Escherichia coli</i> [TaxID: 562]
Taxonomy	Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae; Escherichia

**References**

[1] MAPPING ON GEL  
MEDLINE=96314039, PubMed=8740179, J. Biol. Exp. Syst. Eel. Israel, Japan

274539, TRYP ID ECUW69 standard: DNA, PRO; 175182 EP, XX  
 • SPOT ID=00117 XX  
 1962, 44; 1965, 1 AC  
 XX  
 XX  
 • MAPPING MA XX  
 XX  
 XX  
 IT 22-SEP--993 (Rel. 37, Created)  
 IT 00 MAY --008 (Rel. 55, Last updated, Version 5)  
 XX  
 XX  
 LE L. COLL chromosomal region from 39.2 to 92.6 minutes.

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**Cross-references**

SWISS-PROT	P02990; EFTU_ECOLI
ECODBASE	E042.0; 6TH
CC	Escherichia coli
OC	Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
CC	Escherichia coli

**2D PAGE maps for unidentified proteins**

- How to interpret a protein map
- You may obtain an estimated location of the protein on various 2D PAGE maps, provided the whole amino acid sequence

Document: Done

**Fig. 4.4**  
The entries in a federated 2-DE database like SWISS-2DPAGE contain active hyperlinks to cross-referenced databases, like SWISS-PROT and GeneBank.

third one, up to several thousands of 2-DE gels may be examined simultaneously, and thus the expression levels of sometimes several hundred thousands of proteins must be analyzed. This approach can only be carried out with the assistance of powerful 2-DE analysis software packages.

#### 4.3.1

### 2-DE analysis software

#### 4.3.1.1 Short history of 2-DE analysis by computer

In the early days of two-dimensional electrophoresis, researchers would visually examine and compare their 2-DE images in order to detect spots that might indicate differentially expressed proteins. However it became apparent quickly that only the development of specialised computer software would provide the necessary means to efficiently extract and analyse the huge amount of data contained in even the simplest of these images. In the early 1980s, a handful of academic groups started to develop computer software to analyze 2-DE images. All software packages were able to perform basic 2-DE image analysis, that is background and streaks subtraction, followed by spot detection and quantitation, as well as some kind of automatic comparison between two or more 2-DE gels. One of the first such software systems, Tycho, was developed by Leigh Anderson and Norman Anderson using specialized hardware [27], soon followed by Gellab [28], Elsie [29], Quest [30], Hermes [31], and Melanie [9, 14]. These six groups pioneered the computer analysis of 2-DE images, improving and extending their software packages for many years, thus giving the possibilities to many research labs to carry out automatic or semi-automatic analysis of 2-DE gels. However, all these software packages, while valuable academic research projects, were often difficult to use and required expensive hardware.

The advent of proteomics and the increasingly extensive use of the two-dimensional electrophoresis technology has been adopted by a continuously growing number of research groups around the world. Some of them are processing larger and larger number of 2-DE images simultaneously. Obviously thus, the need for robust software tools arose that could be run on standard computer hardware. By the end of the 1990s, the development of most of the academic packages had been discontinued, and research labs now mainly use commercially available software packages. One exception is the Melanie system, and in particular Melanie 3, the third generation of Melanie software packages for 2-DE analysis. While commercially distributed and supported by one international company<sup>9)</sup> in order to make it

9) Geneva Bioinformatics (GeneBio) S.A.,  
Geneva, Switzerland.

available to the largest possible audience, it continues to be extended and further developed by Daniel Walther and his team at the Swiss Institute of Bioinformatics (SIB – <http://www.isb-sib.ch>). This development involves a close collaboration with the other groups of the Geneva Proteome Network – in particular the research groups of this Chapter’s authors – as well as Geneva Bioinformatics (GeneBio) SA. GeneBio SA is the Geneva based bioinformatics company that distributes the SIB’s databases and proteomics software tools, including Melanie 3 (<http://www.genebio.com>)

In the remaining part of this Section, we describe some of the fundamentals of 2-DE image analysis by computer, illustrated using the Melanie 3 software package, release 3.0.

#### 4.3.1.2 Basic analysis of 2-DE images

First, two-dimensional gels must be digitized. In most cases, this is achieved by the use of a laser densitometer, a CCD camera or a phosphor imager [32]. These pieces of equipment produce images of typically around  $2000 \times 2000$  pixels or more, and a depth of 12 or 16 bits, thus providing a dynamic range of 4096, respectively 65 536 grey levels.

Once loaded into the software, spot detection and quantitation is performed automatically, producing a repository of all protein spots contained in the 2-DE images, as well as related quantitative data, such as the spots’ optical densities, area and volume (integration of the optical density over the area). Relative measures of these values are also given, as for example the relative volume calculated as the absolute volume divided by the total volume of protein in the whole 2-DE gel. Relative values allow to partially compensate for variations in sample load or staining. Using such relative quantitative values provides better reproducibility of data. Automatic 2-DE image comparison is then performed on user-selected gels, by a pattern-matching algorithm. This creates a unique mapping of all the protein spots in all considered 2-DE gels to a reference gel. The latter can either be one of the original 2-DE images or a synthetically produced image containing all (or a selection of) the spots contained in the analyzed gels. Melanie 3 provides several means to manipulate and display the gels, such as gel stacking, various zoom modes, customisable grids, “flicking” back and forth between two gels, or a transparency mode to visually inspect the result of matching. This makes it easy to browse through the data extracted from the 2-DE images. Figure 4.5 shows some of the display and navigation facilities of Melanie 3. Eight gels have been loaded. Four are visible, four are hidden and their names shown on the top right status line of the screen. On the four visible gels, one spot has been selected, the corresponding spot being displayed in green on all four gels. The top left part of the image shows two zoomed in, stacked gels (ECOLI and 94-005). Their spots are displayed in transparency mode, vectors allowing to examine the paired spots. A pop-up



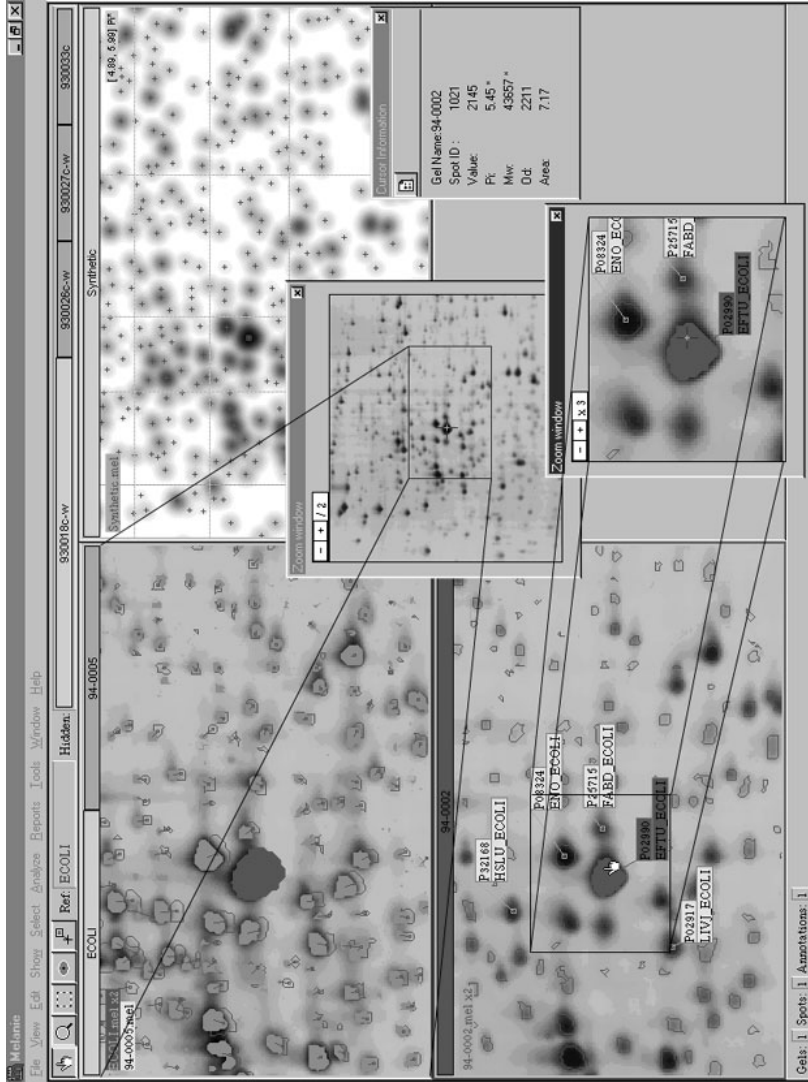


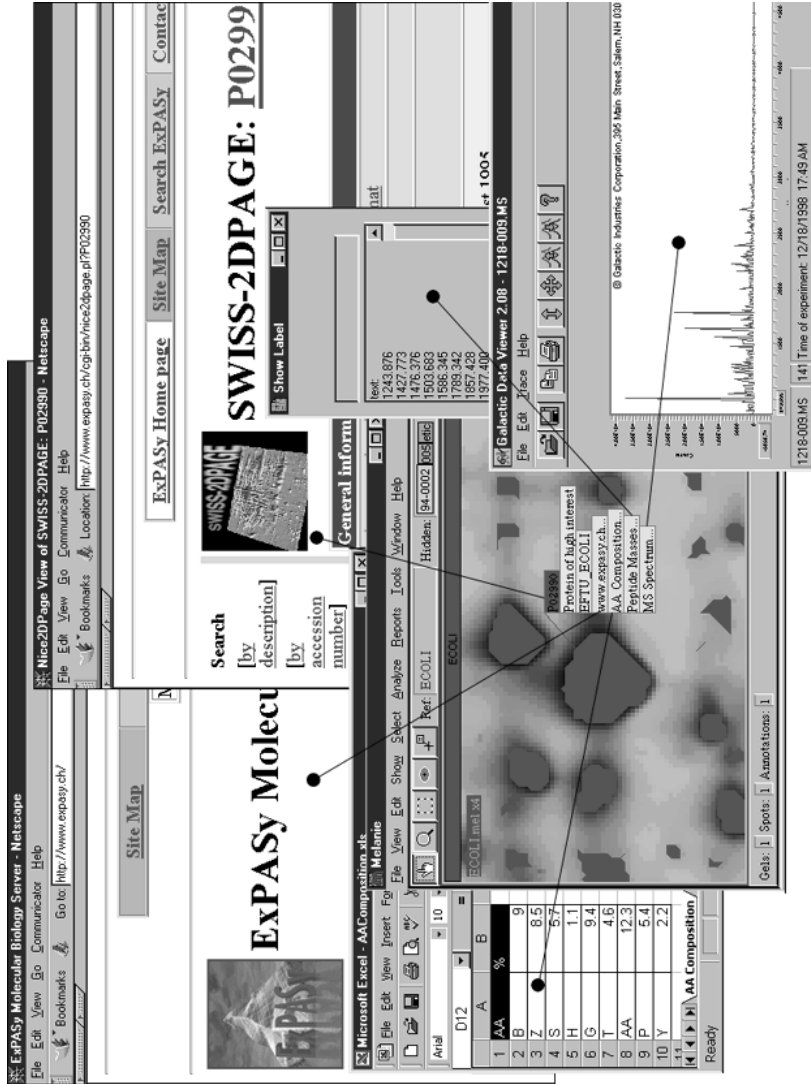
Fig. 4.5 Display and navigation facilities in Melanie. For details see text.

zoom window shows the front gel zoomed out, a rectangle indicating what part of the gel is currently displayed. The bottom left gel displays some annotations (protein names and accession numbers). A second zoom window shows the same area, but zoomed in three times. An information window (bottom right) displays selected data related to the cursor position (in the current setting: gel name, spot ID, grey level value,  $pI/M_r^{10}$  calculated from the reference gel, optical density in the spot centre and spot area). Finally the top right gel is the synthetic gel resulting from merging the other three. It is displayed with a  $pI/M_w$  grid. The current  $pI$  and  $M_w$  values are shown in the top right, respectively bottom left part of the gel (only the  $pI$  range is visible on this Figure: [4.89, 5.99]).

#### 4.3.1.3 Annotations and databases

After extensive analysis of the protein spots has been carried out either by Melanie 3 or through external programs, including protein identification as detailed in Section 4.4, the 2-DE images may be annotated. Melanie 3 provides comprehensive annotation capabilities in order to include into the gel image all related data and information that has been acquired. Each point and each spot in a 2-DE image may given an annotation. Annotations can be added either manually, or they may be imported from an external database, for example through a Laboratory Information Management System (LIMS). Any kind of annotations may be attached to a spot or pixel, as for example the protein ID, a SWISS-2DPAGE ID, a landmark, calibration values ( $pI$ ,  $M_r$ , intensities) or comments, as well as links to external files such as text files or MS spectra and also Internet links. Figure 4.6 shows a zoomed-in portion of an *E. coli* 2-DE gel, a spot corresponding to Elongation factor TU (EFTU\_ECOLI) being selected and highlighted in green. An annotation with seven fields (labels) is attached to it. By double clicking on the various labels, the corresponding piece of information is displayed. In the case of the Accession number (P02990), it launches the default Web browser and requests the corresponding database entry from the user-defined database (here SWISS-2DPAGE on the ExpASY Web server). The database may be on the Web or it can be local. The label «[www.expasy.ch](http://www.expasy.ch)» links to the corresponding URL. A double-click on “AAComposition” opens an Excel file containing the amino acid composition of the protein. “Peptide Masses” displays the list of peptide masses obtained from an MS spectrum, while “MS Spectrum” launches an external program that displays the original spectrum.

- |  |  |
|--|--|
| <p>10) <math>M_r</math> describes the apparent molecular mass of a protein. That means the position on a calibrated molecular weight</p> | <p>scale where the protein of interest is migrating. <math>M_w</math> means molecular weight, in a general manner.</p> |
|--|--|



**Fig. 4.6** The annotations in Melanie are of various formats and can include active links to various types of information, such as hyperlinks to database entries, to spectra, to textual information, to tables, etc.

Annotations may also be exported to external, non-proprietary databases. For example, one can export the accession numbers of all identified proteins and automatically upgrade the SWISS-2DPAGE database.

#### 4.3.1.4 Data analysis

Once basic analysis has been performed on the 2-DE gel images, and they have been compared, the main goal when searching for variation in protein expression is to carry out data analysis. As there are probably as many ways to perform data analysis on 2-DE gel data, as there are research labs, several flexible data analysis tools and statistical programs have been provided in Melanie 3. All produce reports that can either be printed or exported to other external data analysis software packages or imported into databases, such as typically a database linked to a Laboratory Information Management System (LIMS) in which differentially expressed proteins are stored. An important feature of Melanie 3 is that data analysis tools and reports are interactively linked and may thus be used recursively. For example, let us consider the eleven 2-DE images in Figure 4.7. These are ten gels (*93007ve-w.mel* to *930037c-w.mel*) from smooth muscles cells of rat samples, out of which five are from newborn and five from aged rats [7]. The eleventh image (*master1*) is the synthetic composite gel image produced by merging the other ten images. All ten original images have been matched to the composite reference gel. Two classes have been defined containing each five images (marked NEWBORN, respectively AGED on the labels in Figure 4.7). We first select all groups of spots in each of the two classes, then produce a spot report for each class (Figure 4.8) that shows, among other values, the Median and the Mean Standard Deviation (MSD) computed on the spots' area. A group is a set of spots that have been matched across all gel images, thus representing the same protein. Sorting the spots by their MSD value allows us to select from this table only those spots that have similar quantitative values, thus eliminating outliers, i.e. spots that were possibly mismatched, whose shapes were badly detected or that are simply not representative of their class. A class report will then show various statistics about the groups in the two classes, such as Student *t*-test, Wilcoxon test or the Kolmogorov–Smirnov test [33]. This lets the user select and highlight those protein spots that are most differentially expressed between the two classes. Figure 4.5 shows the result of one of these tests, highlighting the group of spots that were ranked highest in term of separability between the two classes. Figure 4.5 also displays the histograms of the top three spot groups. Other classification programs built into Melanie 3 allow to carry out global data analysis on the whole set of gels, such as correspondence analysis (Figure 4.9a) or heuristic clustering (Figure 4.9b).

Once specific proteins of interest have been selected through careful data analysis, further analysis may be carried out to identify or partially characterize those proteins.

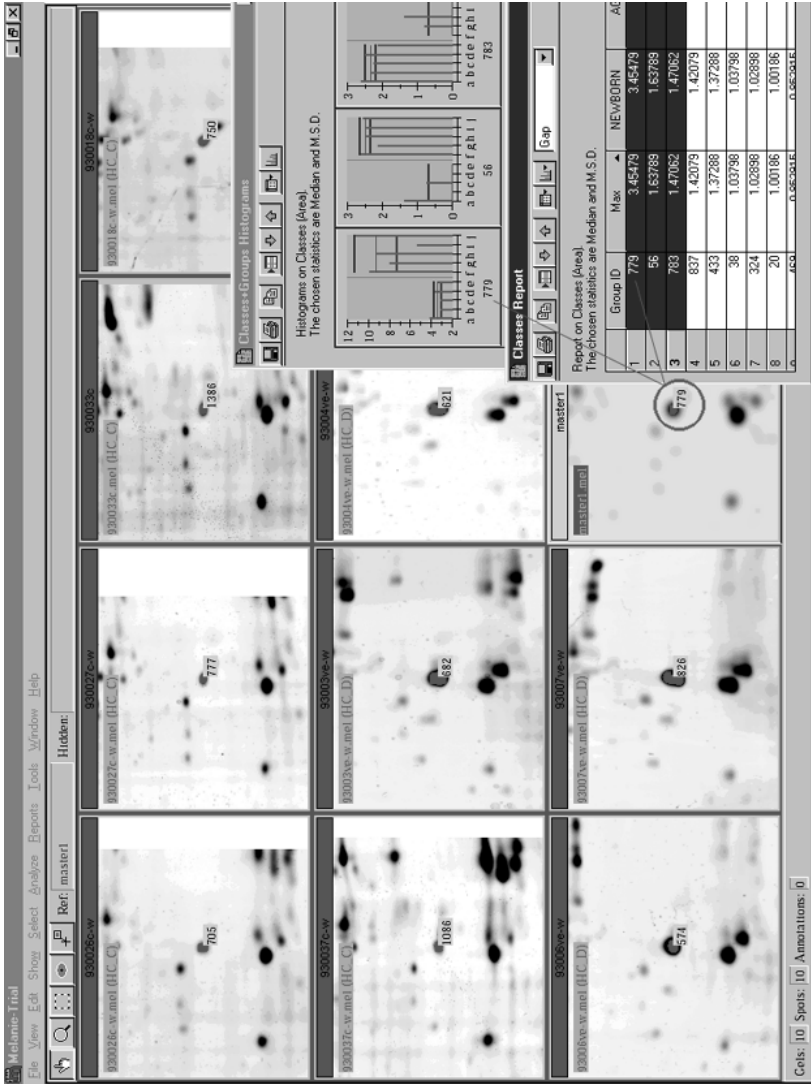


Fig. 4.7 Protein gels analysed with Melanie 3, including a composite image. See text for details.

**Groups Report**

Choose the statistics.  
The sliders restrict the statistics to the central values.

Central tendency

Mean  Midrange  Reference gel values

Dispersion

Mean squared deviation  Mean absolute deviation  Half-range size

Additional values

Coef. Variation  
 Range ratio  
 Separability  
 Correspondence Analysis

OK Cancel

**Groups Report**

Report on Groups (Area).  
The chosen statistics are Median and M.S.D.

	Group ID	Median	M.S.D	Variation	930026c-w	930027c-w
1	297	1.19563	0.0999932	0.0836321	1.29960	1.24762
2	647	1.55952	0.138516	0.0888194	1.66349	1.61150
3	276	1.24762	0.143310	0.114867	1.55952	1.24762
4	391	1.87142	0.163564	0.0874007	1.66349	1.87142
5	158	1.29960	0.172411	0.132665	1.29960	1.29960
6	610	1.87142	0.172411	0.0921285	1.71547	1.97539
7	653	1.40357	0.187431	0.133539	1.40357	1.61150
8	401	2.39126	0.194506	0.0813404	2.13134	2.13134
9	325	0	0.197265	1.000000	0	0
10	416	1.81944	0.197265	0.108421	1.81944	2.28730
11	600	1.71547	0.205977	0.120070	1.87142	1.55952
12	608	1.66349	0.211799	0.127322	1.66349	1.55952

**Fig. 4.8**  
Analysis of single spots from the gels shown in Fig. 4.7.  
See text for details.



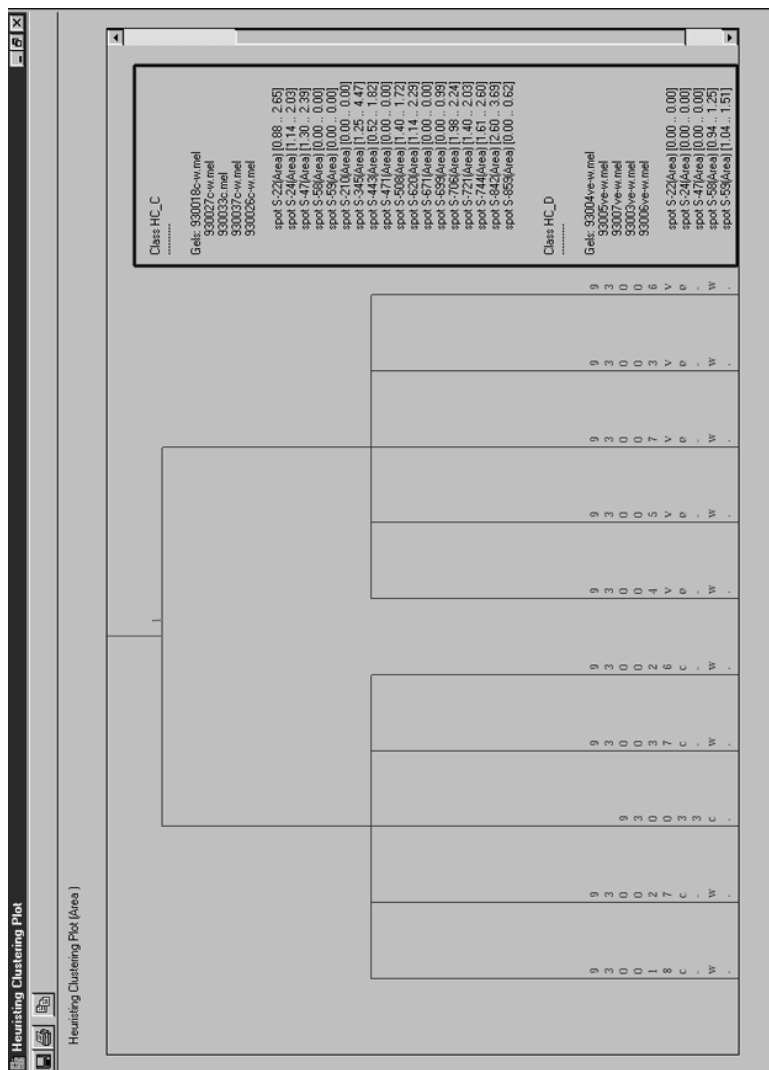


Fig. 4.9 (continued)



#### 4.4

### Identification and characterization of proteins after separation

#### 4.4.1

##### Introduction

Once the spots of interest in a 2-DE gel are selected, the next step is to identify the corresponding proteins in a database. This is another challenge for bioinformatics to design tools adapted to match experimental data with those in sequence databases. Even if the amino acid sequence of a protein can be predicted with a reasonable degree of confidence, post-translational protein modifications cannot always be predicted from the DNA sequence; and their presence or absence can be of paramount importance for the final structure, as well as for the function or dysfunction of a protein. Powerful protein identification tools therefore have to take into account information about known post-translational modifications wherever possible.

Typically, identification tools in proteomics focus on one or more experimentally obtained protein properties and try to achieve protein identification by matching their values against the corresponding theoretical values computed for all sequences in a protein sequence database. Characterization tools are using a more predictive approach in the sense that they do not only match experimental data with data extracted from a database. They perform a predictive calculation to propose PTMs, splicing events or other features.

Various methods are available to determine such protein properties, including amino acid composition analysis [34, 35, 36, 37], partial Edman sequencing [38], peptide mass fingerprinting and MS/MS sequencing (for more details see below).

One of the currently most common approaches for protein identification combines two-dimensional gel electrophoresis, image analysis of the 2-DE gels, extraction of potential target proteins and identification of these targets by mass spectrometry. Hereby, peptide mass fingerprints (technique described by [39, 40, 41, 42, 43]) can be determined by means of matrix-assisted laser desorption ionization time-of-flight mass spectrometry (MALDI-TOF MS). This technique is often limited to the creation of peptide maps. Therefore other MS methods have been proposed to obtain sequence information from peptides signals. These include liquid chromatography/electrospray ionization mass spectrometry (LC/ESI MS) and electrospray quadrupole time of flight (QTOF) [5].

In this Section, we describe a number of protein identification and characterisation tools available on the World-Wide Web (WWW). All the tools described here are accessible from the ExPASy tools page <http://www.expasy.org/tools/>, which is constantly updated. Particular emphasis will be given to the programs provided on the ExPASy proteomics server [23, <http://www.expasy.org>]. Note that currently the Geneva ExPASy server ([www.expasy.org](http://www.expasy.org)).

expasy.org) has five mirror sites in Canada (<http://ca.expasy.org>), Australia (<http://au.expasy.org>), Taiwan (<http://tw.expasy.org>), China (<http://cn.expasy.org>) and Korea (<http://kr.expasy.org>). The tools available on ExPASy use the SWISS-PROT and TrEMBL databases [25] and take the annotations contained in these databases into account wherever possible. In order to facilitate the use and interpretation of the complete and non-redundant set of databases formed by SWISS-PROT and TrEMBL, TrEMBL has been designed to use the same format as SWISS-PROT and to share its system of accession numbers. In particular, SWISS-PROT and TrEMBL use the same conventions in terms of taxonomy annotation (namely the NCBI taxonomy, <http://www.ncbi.nlm.nih.gov/Taxonomy/tax.html>); and they also share the principle that the sequence shown in a database entry is the sequence of the unprocessed precursor. Known post-translational modifications or processing events are documented in the feature table, which is parsed by the ExPASy proteomics tools. In particular, this implies that ExPASy tools first remove signal sequences, propeptides and transit peptides, and consider each mature chain or peptide as a separate 'entry' to which all subsequent computations, such as  $pI/M_w$  calculation or theoretical enzymatic digestion, are applied.

One of the disadvantages of so-called non-redundant databases such as OWL [44] or NCBI nrdb (<http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=Protein>) – apart from the fact that they are not really non-redundant (see Section 4.5) – is that each of the protein sequence databases integrated into such 'non-redundant' databases has its own format and its own conventions. Such format differences include different use of control vocabularies and keywords (e.g. for taxonomic ranges), as well as the decision whether to display the unprocessed or the mature form of the protein with all its co- and post-translational modifications. In particular, molecular weight and isoelectric point can vary significantly between the precursor sequence and the modified mature protein. In the context of protein identification with methods like peptide mass fingerprinting and subsequent database matching, this means that restriction of a search to a taxonomic range, or filtering of results by specifying approximate values for molecular weight and isoelectric point, may compromise the reliability of the result and should be handled much more carefully if data from different databases is used.

#### 4.4.2

#### The tools

##### 4.4.2.1 Identification with amino acid composition: AACompIdent

The AACompIdent tool [37, 45, <http://www.expasy.org/tools/aacomp/>] compares the experimentally determined amino acid composition (the numbers of each amino acid present in a molecule) of an unknown protein with theoretical compositions of all proteins in SWISS-PROT/TrEMBL. A score

quantifying the difference between the compositions of query and database proteins is used to rank the candidate proteins. Estimated pI and  $M_w$ , as well as the name of an organism or a taxonomic range, and a keyword which will be matched against the annotation in SWISS-PROT can be used to filter the results. In order to further increase confidence in the identification results, the amino acid composition of a calibration protein analyzed in parallel, and/or a short sequence tag of up to six amino acids known to match the sequence of the unknown protein can be specified if known. Several amino acid constellations are available, accounting for different experimental methods of obtaining amino acid compositions and the sets of amino acids quantified by these methods.

#### 4.4.2.2 Protein identification with sequence tags: TagIdent

TagIdent [46, <http://www.expasy.org/tools/tagident.html>] makes use of the high specificity of short amino acid sequence tags in molecularly well defined organisms with small proteomes and a low degree of post-translational modifications and, in particular, few N-terminally blocked proteins. The tool can match a short sequence tag of up to six amino acids, pI and  $M_w$  against all SWISS-PROT/TrEMBL sequences from a species or taxonomic category. If more than one protein satisfies the user-specified tag and pI/ $M_w$  ranges, TagIdent produces an unranked list of all candidate database entries.

Alternatives to TagIdent, which do not make use of pI and  $M_w$  estimates but might be used for sequences longer than six amino acids, are ScanProsite (<http://www.expasy.org/tools/scnpsit2.html>) or PHI-BLAST [47, [http://www.ncbi.nlm.nih.gov/cgi-bin/BLAST/nph-psi\\_blast?form=1](http://www.ncbi.nlm.nih.gov/cgi-bin/BLAST/nph-psi_blast?form=1)]. Both programs allow a higher degree of ambiguity in the sequence tag, which can be specified in regular expression format.

#### 4.4.2.3 Peptide mass fingerprinting: PeptIdent, SmartIdent, ProteinProspector, MOWSE, Mascot, and PROWL

Increasing reproducibility of available separation techniques and sensitivity and affordability of mass spectrometers, as well as the desire and need to automate the identification process, have caused peptide mass fingerprinting and MS/MS sequencing to gain importance and to become the method of choice for many proteomics laboratories. Several tools are available to assist users in the interpretation of mass spectrometry data. PeptIdent (<http://www.expasy.org/tools/peptident.html>) on the ExPASy server follows the concept of the other tools from the ExPASy proteomics suite, in that it takes into account annotation available in the SWISS-PROT/TrEMBL database, in particular as post-translational modifications and processing are concerned. The user can paste peptide masses (monoisotopic or average) into the PeptIdent form, but peptide mass data can also be uploaded from a file on the user's local computer. Supported file formats are '.pkm'

format, a format produced by a number of peak detection algorithms implemented in software like Voyager 4 (Applied Biosystems, Framingham) controlling Applied Biosystems MS instruments or PerSeptive Grams (Galactic Industries Corporation), and Sequest format. In particular, the first column of each line contains a peptide mass in Daltons; and example files can be found in the PeptIdent documentation on ExPASy. Further parameters can be specified describing the experiment which produced the peptide mass fingerprint data, such as modification of cysteine and methionine residues, the enzyme used for digestion and the quality of this digestion (maximum number of missed cleavage sites to be considered), the ion mode, and peptide mass tolerance. Species filtering, that is, restriction of the database search to one species or to a taxonomic category can considerably reduce the number of candidate proteins, which can be of enormous help in the interpretation of identification results. Further filtering can be done by specifying pI and  $M_w$  estimates, and by giving a minimum number of peptides required to match for a protein to be suggested as a candidate. Candidate proteins are ranked by the number of matching peptides (modified or unmodified). Another tool, SmartIdent [48], which uses artificial intelligence methods to search databases has been developed and is available from the tools page of ExPASy (<http://www.expasy.org/tools/>). It takes into account several other characteristics of a candidate protein and its matching peptides. Properties such as mass errors, the type of modification that a peptide carries, parameters related to AA composition or intensity of the MS signal are weighted in a multilevel system that allows to discriminate between highly probable and less probable peptide hits. This allows for a better discrimination between relevant and irrelevant matches, and for automation of the identification process. In addition, SmartIdent does not require that the MS spectra are internally calibrated. The program corrects for non-constant mass shifts by using a robust regression system. This allows for selecting appropriate and non-appropriate masses for every identified protein. A PeptIdent or a SmartIdent result provides, for each suggested protein, direct access to the FindMod, GlycoMod and Findpept programs for further characterisation and prediction of post-translational modifications (see below), and to the BioGraph tool (<http://www.expasy.org/tools/biograph/bio/>) for graphical representation of matched and unmatched peaks.

A number of related tools are available, for example:

- The ProteinProspector Server (<http://prospector.ucsf.edu>) from the University of California San Francisco mass spectrometry facility. This suite of programs includes various tools for mining sequence databases in conjunction with mass spectrometry experiments, such as MS-Fit, MS-Tag, MS-Product and others.
- MOWSE (<http://www.seqnet.dl.ac.uk/Bioinformatics/Webapp/mowse/>) at the SERC Daresbury Laboratory, a peptide mass fingerprinting tool searching the OWL database.

- Mascot ([http://www.matrixscience.com/cgi/index.pl?page=/search\\_form\\_select.html](http://www.matrixscience.com/cgi/index.pl?page=/search_form_select.html)) at Matrix Science Ltd. which provides identification resources for peptide mass fingerprinting, MS/MS ion search and MS/MS sequence queries.
- PROWL (<http://prowl.rockefeller.edu/>) at Rockefeller University. The PROWL environment comprises ProFound, a peptide mass fingerprint identification program which can also be configured to identify components from a mixture of up to four proteins, and PepFrag, an identification tool for fragmentation mass spectrum data.

Given the variety of different identification programs available (see also [49]), we have developed CombSearch (<http://www.expasy.org/tools/CombSearch/>), a program which provides a unified search engine for a prototype query with peptide mass fingerprinting data, and automatically submits the user-entered parameters to all of the above-mentioned identification programs. This enables users to simultaneously query several programs and databases, and to take advantage of the strengths of each of these programs without having to fill in multiple submission forms. By this means, scientists might also find their 'favorite', which, among the available programs, best suits their specific application scenario.

#### 4.4.2.4 Cross-species identification with multiple parameters: MultiIdent

Despite the rapid growth of protein sequence databases, the majority of the information contained in these databases concerns a relatively small number of model organisms which are targets of genome projects and extensive study (see Section 4.5). If the organism under study is not well characterized, proteome data from evolutionarily related model organisms can be used for the identification, provided that the parameters used in this process are well conserved across species boundaries [1]. Identification based solely on the MS-derived masses of peptide resulting from an enzymatically digested protein cannot always be regarded as highly reliable, as peptide masses are usually not very well conserved across species boundaries. However, amino acid composition data have been shown to be very similar for homologous proteins from closely related species. Hence, to identify proteins from organisms which do not have their genomes sequenced and are thus not well represented in the databases, peptide mass fingerprinting can be used in combination with amino acid composition data, as well as with isoelectric point and molecular weight estimates.

This is the concept of the MultiIdent tool [50, <http://www.expasy.org/tools/multiident/>]. MultiIdent first generates a list of best-matching proteins by amino acid composition, whose sequences are then digested theoretically and the resulting peptide masses matched against the experimentally obtained peptide mass fingerprint of the unknown protein. Similarly to

AACompIdent, several amino acid compositions are available, and a sequence tag of up to six amino acids can be used to increase confidence in the identification results.

#### 4.4.2.5 Protein characterization and discovery of post-translational modifications: FindMod and GlycoMod

As discussed above, peptide mass fingerprinting can identify a protein if the genome is available in a database. However, a considerable portion of the peptide masses often remains unmatched and can be assumed to represent post-translationally modified forms of predicted peptides. The FindMod tool [51, <http://www.expasy.org/tools/findmod/>] examines mass differences between empirical and theoretical peptides. If such a mass difference corresponds to a mass difference known to be induced by a discrete post-translational modification, FindMod consults a set of rules in order to predict which amino acid residues in the peptide might carry that modification (if any). FindMod can currently predict 27 modifications, that are known as amidation, farnesylation, methylation, palmitoylation, phosphorylation, and many more. At least one rule has been established for each of these modifications by carefully examining the relevant annotation in the SWISS-PROT and PROSITE [52] databases and the literature, with respect to the type of organism, position and amino acid at which the modification can be observed. The same algorithm is applied to suggest possible single amino acid substitutions in peptides from differences between experimental and theoretical peptide masses. In this way, the tool assists scientists to characterize a gene product and its possible interactions, activities, and role in normal and disease states.

Glycosylation is one of the most abundant forms of covalent protein modification. The range of monomers of which carbohydrate structures in glycopeptides can be composed is broad, and these monomers can be joined in many ways to form linear and branching structures. The diversity of carbohydrate compositions and structures results in a very large number of different potential post-translational modifications that by far exceeds the relatively small number of modifications considered in FindMod. This diversity, as well as a variety of parameters specific to glycosylation, requires a specialized tool, GlycoMod (<http://www.expasy.org/tools/glycomod/>) [53]. Mass spectrometry of glycopeptides can provide useful information on the composition of the attached oligosaccharides: if a peptide is presumed to be glycosylated, the mass of the oligosaccharide can be deduced as the difference between the empirical peptide mass and the mass predicted from the peptide sequence. The potential oligosaccharide compositions corresponding to this mass difference can be calculated from a combination of the masses of possible monosaccharide constituents [54]. GlycoMod considerably facilitates this assignment of mass to composition. The program is

linked to SWISS-PROT so that peptide masses of known proteins can be accessed, screened for potential glycosylation sites (both N-linked and O-linked) and used in the identification of potential glycopeptides.

## 4.5

### Proteome databases

#### 4.5.1

##### Introduction

The science community has clearly established the essential role which protein and DNA sequences play in the understanding of biological systems. The sequences themselves are informative: Indeed, many software tools are available which allow to “make sense” of the primary sequence information. Take those that analyse protein sequences for domains and active sites, perform similarity and homology searches, or predict the three-dimensional structure or physico-chemical parameters. However, raw sequences contain insufficient information, per se. One cannot infer any description or understanding of the level of expression of the active proteins, the content of post-translational modifications (PTMs), the tertiary structure and, what is perhaps the most relevant information, a protein’s function. Like the sequences themselves, all these “added value” data need to be captured in various databases. These databases have to be queried by different types of users in proteomics and should therefore be easily searchable by software tools and be inter-linked in order to document the correspondences between the type of information provided by the different databases.

This Section briefly describes the data resources available and relevant for researchers working in proteomics. The databases can be classified according to the type of information provided, i.e. protein sequence, nucleotide sequence, pattern/profile, 2-DE, 3-D structure, PTM, genomic, and metabolic. A more extensive description and list has been published by Bairoch [55]. A reasonably exhaustive and constantly updated list of databases can also be found at <http://www.expasy.org/alinks.html>. Here we focus on sequence databases, protein domain databases, 2-D PAGE, and PTM databases.

#### 4.5.2

##### Protein sequence databases

The most comprehensive source of protein information is found in protein sequence databases. These can be divided into universal databases, which

store protein information from all types of biological sources, and specialised databases, which concentrate their efforts on restricted groups of protein families or organisms. Universal protein sequence databases can be categorised in databases that are simple repositories of sequence data, mostly translated from DNA sequences, and in annotated databases. The latter demand the assistance of biologists (“annotators”) who screen the original literature, review articles as well as electronic archives.

Here we will mainly describe SWISS-PROT, an annotated universal sequence database, and TrEMBL, an automatically generated sequence database with repository character, which supplements SWISS-PROT.

#### 4.5.2.1 SWISS-PROT, TrEMBL and their integration in proteomics

SWISS-PROT [25, <http://www.expasy.org/sprot/>] is a curated protein sequence database which provides a high level of annotation, such as the description of a protein’s function, its domain structure, PTMs, conflicts between literature references and variants. It also provides a minimal level of redundancy, a high level of integration with other biomolecular databases, and an extensive external documentation. Created in 1986, it now includes more than 100 000 entries from more than 7000 different species. SWISS-PROT is available from more than 100 WWW servers world-wide; and its main host is the ExpASY server.

Each entry consists of different line types (Figure 4.10). A first section represents the description part. It includes, among others, the accession number (a unique entry identifier), the update dates, the protein description (its “name”), the gene name, and the taxonomic origin. This information heading is then followed by a reference section. Each of the reported references includes the type of experimental work contributing to the entry (sequencing, 3-D structure determination, mutagenesis studies, etc.), the author list, and the literature references. Entries are not only populated by literature work: a continuously increasing number of sequences are directly submitted to the database and published this way. The following section is a comment section. Here a variety of textual comments, classified into topics are listed. Such topics include function, subunit, similarity, PTM, MS source, etc. This section is followed by the database cross-reference section, which provides active links to other biological databases. In this way, SWISS-PROT is currently linked to more than 30 different databases. After a keyword field, the next section is the so-called feature table. It describes regions or sites of interest in the sequence and documents PTMs, binding sites, active sites, secondary structures, variants, conflicts, and the like. The entry ends with the amino acid sequence itself. It is important to note that the sequence shown is that of the unprocessed precursor of the protein, before post-translational modification and processing. Tools that identify proteins by using isoelectric point (pI), molecular weight ( $M_w$ ) or peptide fingerprints should ideally make use of the information held within the



```

ID  ACTG_HUMAN      STANDARD;      PRT;   375 AA.
AC  P02571; P14104; P99022;
DT  21-JUL-1986 (Rel. 01, Created)
DT  21-JUL-1986 (Rel. 01, Last sequence update)
DT  15-JUL-1999 (Rel. 38, Last annotation update)
DE  Actin, cytoplasmic 2 (Gamma-actin).
GN  ACTG1 OR ACTG.
OS  Homo sapiens (Human), Mus musculus (Mouse), Rattus norvegicus (Rat),
OS  Bos taurus (Bovine), Trichosurus vulpecula (Brush-tailed possum), and
OS  Anser anser anser (Western graylag goose).
OC  Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC  Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
RN  [1]
RP  SEQUENCE FROM N.A.
RC  SPECIES=Human;
RX  MEDLINE=88246448 [NCBI, ExPASy, Israel, Japan]; PubMed=2837653;
RA  Erba H.P., Eddy R., Shows T., Kedes L., Gunning P.;
RT  "Structure, chromosome location, and expression of the human gamma-
RT  actin gene: differential evolution, location, and expression of the
RT  cytoskeletal beta-and gamma-actin genes.";
RL  Mol. Cell. Biol. 8:1775-1789(1988).
RN  [2]
RP  SEQUENCE FROM N.A.
...
CC  !- FUNCTION: ACTINS ARE HIGHLY CONSERVED PROTEINS THAT ARE INVOLVED
CC  IN VARIOUS TYPES OF CELL MOTILITY AND ARE UBIQUITOUSLY EXPRESSED
CC  IN ALL EUKARYOTIC CELLS.
CC  !- SUBUNIT: POLYMERIZATION OF GLOBULAR ACTIN (G-ACTIN) LEADS TO A
CC  STRUCTURAL FILAMENT (F-ACTIN) IN THE FORM OF A TWO-STRANDED
CC  HELIX. EACH ACTIN CAN BIND TO 4 OTHERS.
CC  !- SUBCELLULAR LOCATION: CYTOPLASMIC.
CC  !- MISCELLANEOUS: IN VERTEBRATES 3 MAIN GROUPS OF ACTIN ISOFORMS,
CC  ALPHA, BETA AND GAMMA HAVE BEEN IDENTIFIED. THE ALPHA ACTINS ARE
CC  FOUND IN MUSCLE TISSUES AND ARE A MAJOR CONSTITUENT OF THE
CC  CONTRACTILE APPARATUS. THE BETA AND GAMMA ACTINS CO-EXIST IN MOST
CC  CELL TYPES AS COMPONENTS OF THE CYTOSKELETON AND AS MEDIATORS OF
CC  INTERNAL CELL MOTILITY.
CC  !- SIMILARITY: BELONGS TO THE ACTIN FAMILY.

```

**Fig. 4.10**

Example of a SWISS-PROT entry in its textual format.

"..." indicates cuts in the entry. Underlined text is associated with hyperlinks on the ExPASy server.

feature table. One can then achieve an optimal approximation of the protein in its mature state. Hence, before computing  $pI$  and  $M_w$ , the signal sequence should be removed. On the ExPASy server, SWISS-PROT (and TrEMBL; see below) entries can be accessed either in their original format, with html links added (e.g. <http://www.expasy.org/cgi-bin/get-sprot-entry?P00750>), or in a more user-friendly visualisation mode, called "NiceProt View" (e.g. <http://www.expasy.org/cgi-bin/niceprot.pl?P00750>).

On the one hand, genome sequencing projects are generating a dramatically increasing number of sequences to be incorporated in SWISS-PROT. On the other, the number of annotators, who screen literature and sequence databases to populate SWISS-PROT with the high quality annotations, is limited. However, despite the increase in available raw sequence data it was not judged appropriate to automatically populate SWISS-PROT with data of lower quality standards. This is where TrEMBL (Translation of EMBL

```

DR   EMBL; M19283; AAA51579.1; -. [EMBL / GenBank / DDBJ] [CoDingSequence]
...
DR   EMBL; M26111; AAA49315.1; -. [EMBL / GenBank / DDBJ] [CoDingSequence]
DR   PIR; A28098; ATHUG.
...
DR   PIR; A55001; A55001.
DR   HSSP; P02570; 2BTF. [HSSP_ENTRY / SWISS-3DIMAGE / PDB]
DR   SWISS-2DPAGE; P99022; MOUSE.
DR   GeneCards; ACTG1.
DR   MIM; 102560; -. [NCBI / EBI]
DR   MGD; MGI:87906; Actg.
DR   INTERPRO; IPR000279; -.
DR   PFAM; PF00022; actin; 1.
DR   PRINTS; PR00190; ACTIN.
DR   PROSITE; PS00406; ACTINS_1; 1.
DR   PROSITE; PS00432; ACTINS_2; 1.
DR   PROSITE; PS01132; ACTINS_ACT_LIKE; 1.
DR   PRODOM [Domain structure / List of seg. sharing at least 1 domain]
DR   BLOCKS; .
DR   DCMQ; .
DR   PROTOMAP; .
DR   PRESAGE; P02571.
DR   DIP; P02571.
KW   Multigene family; Structural protein; Methylation; Acetylation;
KW   Cytoskeleton.
FT   PROPEP      1       1       REMOVED IN MATURE FORM.
FT   CHAIN       2       375      ACTIN, CYTOPLASMIC 2.
FT   MOD_RES     1       1       ACETYLYATION.
FT   MOD_RES     2       2       ACETYLYATION.
FT   MOD_RES     73      73       METHYLATION.
FT   CONFLICT    316     316      E -> K (IN REF. 3).
FT   CONFLICT    344     344      S -> F (IN REF. 3).
SQ   SEQUENCE   375 AA; 41793 MW; 54D08F986964EFD5 CRC64;
MEEFIAALVI DNGSGMCKAG FAGDDAPRAV FPSIVGRPRH QGVMVGMGQK DSYVGDAAQS
KRGILTLKYP IEHGIVTNWD DMEKIWHHTF YNELRVAPEE HPVLLTEAPL NPKANREKMT
QIMFETFNTF AMYVAIQAVL SLYASGRITG IVMDSGDGVV HTVPIYEGYA LPHAILRLDL
AGRDLTDYLM KILTERGYSF TTTAEREIVR DIKEKLCYVA LDFEQEMATA ASSSLEKSY
ELPDGQVITF GNERFRCPFA LFPQPSFLGME SCGIHETTFN SIMKCDVDIR KDLYANTVLS
GGTMYPGIA  DRMQKEITAL APSTMKIKII APPERKYSVW IGGSIILASL TFQQMWSIKQ
EYDESGPSIV HRKCF

```

Fig. 4.10 (continued)

Nucleotide Sequence Database [26]) steps in. TrEMBL was created in 1996 and consists of computer-annotated entries in SWISS-PROT-like format. It is populated by protein sequences translated from the coding sequences (CDS) in EMBL and is a supplement to SWISS-PROT. In a way, it can be considered as a preliminary section of SWISS-PROT; indeed, once the manual annotation is performed, the entries move on to SWISS-PROT.

Apart from text searches, both SWISS-PROT and TrEMBL can be queried by sequence analysis tools as well as protein identification and characterization tools. Tools like PeptIdent or FindMod (see Section 4.4) take advantage of the annotation of SWISS-PROT/TrEMBL to improve their capacities of identifying and characterizing active chains and proteins annotated to be post-translationally modified. This is better than basing their computations and predictions merely on the precursor protein sequences.

For many applications, such as protein identification or sequence similarity searches, it is necessary to use a database as complete and non-redundant as possible. SWISS-PROT, and to some extent TrEMBL, aim to

be non-redundant by maintaining only one consensus sequence record for each protein – even if several copies or subfragments of the same protein have been sequenced. Polymorphisms (sequence variants), alternative splicing events, and sequencing conflicts are documented in the feature table of the retained copy. As a consequence, so-called “non-redundant” databases, which use strict sequence matches as exclusion criteria, contain a higher number of entries than the combination of SWISS-PROT + TrEMBL. The drawback, though, is that they contain multiple copies of the same protein and are thus not really non-redundant. Examples are OWL, and nrdb (Non-Redundant DataBase provided by the NCBI, built from SWISS-PROT and GenPept, an automatic translation of the GenBank nucleotide sequence database).

The links provided by the SWISS-PROT entries to other proteomics databases like SWISS-2DPAGE, PROSITE/InterPro and about 30 other databases allow for rapid access to experimental proteomic data, like position and number of protein spots on a 2-D gel, other members of the same family, or the 3-D structure of the protein, etc.

#### 4.5.2.2 Specialized protein sequence databases

There are many such databases available. Their contents vary a lot in terms of the range of interest, number of entries and type of information. As it is impossible to be exhaustive here, we recommend readers to consult the updated list on the World-Wide Web, maintained by Amos Bairoch on the ExpASY server (<http://www.expasy.org/alinks.html>).

#### 4.5.3

##### Nucleotide sequence databases

Nucleotide sequence databases are the main source of sequence information for protein sequence databases. There is an international collaboration for the production, the maintenance and the distribution of nucleotide sequence information. The organisations involved are the European Bioinformatics Institute (EBI) in Great Britain distributing EMBL Nucleotide Sequence Database, the National Centre for Biotechnology and Information (NCBI) in the USA distributing GenBank™ [56], and the National Institute of Genetics in Japan distributing DDBJ (DNA Database of Japan) [57]. In the scope of this collaboration, EMBL, GenBank and DDBJ all contain the same information – the nucleotide sequence database is in fact one single database distributed under three different names, and in three different formats.

EMBL/GenBank/DDBJ is an archive or repository database. In that respect, no attempt is made to reduce the size of the database by merging entries submitted by different groups and which correspond to the same

stretches of genome or mRNA. As stated above and for historical reasons, their format is not unified, but they share the same organizational principles as described for SWISS-PROT, i.e. a header containing the name of the sequence, the species of origin, followed by the references, a feature table, and the sequence data.

The nucleotide sequence databases can be screened by a number of sequence similarity tools and some protein identification applications, such as SEQUEST [58]. As it is difficult to predict coding sequences from raw genomic data, and in some cases the authors have not submitted the information to the database, the tools propose protein hits from translated raw DNA sequences.

In addition, a powerful approach to search DNA sequences exclusively in their coding region is to search databases populated with Expressed Sequence Tags (ESTs) [59]. ESTs are short nucleotide sequence of 250–400 base pairs generated by random, single-pass sequencing of cDNA libraries. In September 2000, public EST databases contained more than 5.6 million entries, about 40% of which were human sequences. They are highly redundant and contain many sequencing errors. However, in combination with expert analysis or searches in other sequence databases, they provide a highly valuable source of proteomic information and can be searched with mass spectrometric fragmentation data [40, 60].

#### 4.5.4

##### **Databases for protein families, domains and functional sites: InterPro**

Given the rapid release of raw, uncharacterized sequence data from genome sequencing projects, there is an increasing need for automatic methods to identify protein family traits and to assign functions to unknown sequences. A number of signature databases are available, built around the idea that certain regions which are vital for a protein function are usually well conserved. An analysis method is chosen which allows to describe the conserved motif, fingerprint or signature in a way that enables software tools to scan against unknown protein sequences in order to identify, rapidly and reliably to which known protein family (if any) a new sequence belongs. Such databases are: PROSITE [52] for regular expression patterns and profiles, Pfam [61] for hidden Markov models (HMMs), PRINTS [62] for fingerprints (groups of aligned, unweighted motifs) and Blocks [63], for aligned weighted motifs or blocks. ProDom [64] is an automatic compilation of homologous domains, derived by clustering of SWISS-PROT and TrEMBL. As each of the above-mentioned databases has its strong points and its weaknesses, it became increasingly difficult for scientists to perform a comprehensive and reliable functional analysis of their unknown sequences. Indeed, each database has its own format, search tools and search outputs. In an attempt to address these difficulties, the authors of the sig-

nature databases cited above formed the InterPro consortium and developed InterPro [65], an integrated documentation resource for protein families, domains and functional sites. InterPro is accessible for text- and sequence-based searches at <http://www.ebi.ac.uk/interpro/>, thus providing an integrated view of all member databases and an intuitive search interface.

#### 4.5.5

##### 2-DE databases

2-D PAGE databases, along with PTM databases (see sub-section 4.5.6) are among the most recent proteomic databases available on the web. They are also pioneers in data integration on the WWW. They collect information on proteins identified on 2-DE and consist of two major components: image data and textual information. The image data are a collection of reference gel maps. More than one map per biological sample may exist. Such an image is the representation of a stained gel scanned optically. Apparent spots represent the position of focused protein forms and are linked to the textual information component of the database. The latter includes, essentially, data on apparent pI and Mw of the spots, the name and description of the protein, the identification method, bibliographical references, and cross-references to SWISS-PROT and other databases. Access to the data and navigation within the databases are performed through simple operations, such as clicking on a spot in an image, selecting a spot/protein identifier from a list, or keyword searches.

##### 4.5.5.1 SWISS-2DPAGE

The SWISS-2DPAGE [20, 6] database was created and is maintained at the Swiss Institute of Bioinformatics in collaboration with the University Hospital of Geneva. In March 2000, it contained 26 reference maps from human, mouse, *Saccharomyces cerevisiae*, *Escherichia coli* and *Dictyostelium discoideum*. More than 700 entries document over 3000 identified “spots”. The proteins represented by those spots were identified either by matching with other gels, by amino acid composition, by Edman sequencing, by immunoblotting, and recently over 700 of them were identified by mass spectrometry. The text format is similar to the one in SWISS-PROT. It includes specific fields, such as the type of master gel from which the protein spot has been identified, the list of gel images associated with the entry, as well as other 2-DE specific data, such as the mapping procedure, the spot identifier, the experimental pI and Mw, the peptide mass fingerprint and the amino-acid composition – if experimentally determined. The database can be accessed from SWISS-PROT by a hyperlink anchored to a cross-reference. If no identified spot exists in SWISS-2DPAGE for a given entry in

SWISS-PROT, an image is generated that highlights the theoretical position of the corresponding protein.

#### 4.5.5.2 Other 2-D PAGE databases

The number of 2-D PAGE databases is continuously increasing. An up-to-date list can be found in WORLD-2DPAGE (<http://www.expasy.org/ch2d/2d-index.html>), an index of 2-D PAGE databases and services. More than 25 species are represented in about 100 2-D maps all over the world. The databases are established in various formats. However, an increasing number of groups have joined the principle of federated 2-DE databases [24], according to which the organization of and access to a database must comply with five rules.

#### 4.5.6

##### Post-translational modification databases

In an era where more and more complete genomes are being sequenced, the challenge of understanding proteins and proteomes is moving towards the understanding protein modifications that cannot be predicted from the nucleic acid sequence. Most proteins are indeed the target of PTMs and are not functional unless they are modified. While SWISS-PROT, as a universal database, places a considerable emphasis on the documentation of post-translational modification within the sequence records, several specialized databases have been set up recently to cater for this emerging field.

**RESID** [66] is a general database of protein structure modifications (<http://www-nbrf.georgetown.edu/pirwww/dbinfo/resid.html>), maintained by the National Biomedical Research Foundation in the USA and the Protein Information Resource group (PIR). The database contains descriptive, chemical, structural, and bibliographical information on 283 (Release 22.1, July 2000) types of modified amino-acid residues. Apart from text-based searches, RESID can also be queried by molecular weight: an average or monoisotopic mass can be entered (together with a mass variance) to search for all modified amino acid residues in the database with masses similar to the input mass.

Apart from this general PTM database, some databases specialized in one particular type of PTM have been created. **GlycoSuiteDB** (<http://www.glycosuite.com>) is an annotated database of glycan structures. The database is provided by Proteomesystems Ltd and contains information about most published O- and N-linked glycans. If glycan structures are known to be attached to proteins, cross-references to SWISS-PROT/TrEMBL are given. GlycoSuiteDB can be queried by mass, by attached protein, by oligosaccharide composition or different modes of textual queries (taxonomy, biological source, etc.).

Two more examples of PTM databases are **O-GLYCBASE** [67], a database of O-glycosylated proteins (<http://www.cbs.dtu.dk/databases/OGLYCBASE/>), and **PhosphoBase** [68], a database of phosphorylation sites in proteins and peptides (<http://www.cbs.dtu.dk/databases/PhosphoBase/>). Both databases are maintained by the Center for Biological Sequence Analysis (CBS) in Denmark, which also provides prediction servers for both types of modifications (NetOGlyc and NetPhos).

Although the current number of PTM databases is quite small, we expect them to multiply, with an increasing amount of data on PTM structures becoming available, as well as the amino-acid sequence patterns where they can be found.

#### 4.5.7

#### **Conclusion**

It is clear that the databases described above do not cover all the aspects of interest for proteomics researchers. There are databases that use the sequence databases to perform calculation and analysis, such as sequence clustering, phylogeny, or profile searching, and thus create added-value databases. Other databases report results from functional studies and mutational experiments, or from 3-D structure determination, or describe metabolic pathways. Although it is impossible to list them all here, it is of interest to know that they exist. Some of them are permanently updated, some of them have only a short existence, some of them are not publicly available. The existence of databases in proteomics is following a dynamic and continuously developing model, and simply reflects the dramatic evolution the field of proteomics is witnessing.

#### 4.6

#### **Automation in proteome analysis**

##### 4.6.1

#### **Introduction**

As already explained, analysing a proteome first involves the isolation of the biological sample of interest. Then the contained proteins have to be separated in order to allow their individual identification, characterization and possibly quantification (Figure 4.1.). The number of proteins to analyse is very often large, ranging from a few hundreds in low prokaryotes to possibly 500 000 to one million in human samples [2].

In order to enable the analysis of entire proteomes, one has to develop scalable technologies. This often means that standard methods have to be automated and, if possible, parallelized.

We already know from the first Sections of this Chapter that the separation of all proteins from a complex biological sample using 2-DE is a well established and widely used technique [2]. It produces thousands of distinct protein samples ready to be further analyzed, and it does so with very few manipulations and in a parallel manner, (see Section 4.2).

From 2-DE gel spots, protein identification can be performed using various methods, as described in Section 4.4. All of these approaches imply a serial process, starting from a single-step or multiple-step chemical processing of the entire proteins and followed by the measurement and the interpretation of the produced protein attributes, then by specific queries in protein sequence database. The outputs of such searches are interpreted and used, first, to annotate the 2-DE image with the identification results and, second, to focus on the biological relevance of the expression level of the interesting protein spots.

A next step in proteome analysis is the characterization of all proteins, i.e. the description of the processing events occurred, of the presence and position of post-translational modifications or of amino acid substitutions.

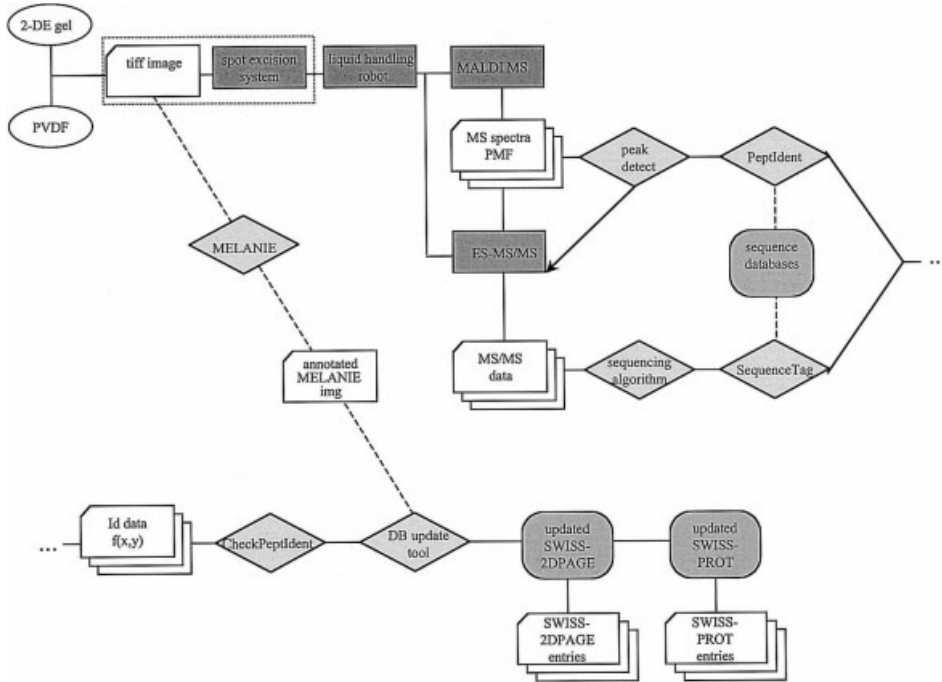
In this Section we talk about automation. Automation first means “hands-free” work. It involves reduction of the technician’s work, better reproducibility, reduction of potential human mistakes, an integrated approach, and a sample-tracking system. Very often automation is also correlated with high-throughput and therefore continuously faces bottleneck issues. It has to be noticed that both aspects are complementary and have to be considered seriously. Very high throughput methods do not afford the possibility to interpret and validate all detailed aspects of an analysis. In terms of protein identification and database update, this means that there exists a conflict between making use of all detailed annotations in the databases and validating manually all results on the one hand, and producing massive amount of very high quality computer interpreted data on the other. It is a real challenge for the bioinformatics developers to optimize the gel analysis software as well as the identification and characterisation tools and to release them as a powerful and complementary package. Indeed, these tools have to handle a high volume of data, often in a very short time. They need to be of the best quality and accuracy. The results of the analyses have to be visualized and interpreted in a global way, while providing access to all details.

#### 4.6.2

#### **Robotized protein identification using peptide mass fingerprinting**

Manual protein identification from 2-DE gels is a time consuming technique that starts with spot excision and ends with the update of 2-DE databases. Nevertheless, this repetitive and exhaustive procedure could be mechanized using robots, thus increasing protein identification throughput.





**Fig. 4.11**  
Flow chart describing a robotised approach for automated protein identification using 2-DE and mass spectrometry.

In the process of automating the procedure using robotics, one has to pay attention to different elements to be integrated. These constitutive elements include components that are used in a manual approach and are complete with dedicated elements: 1) machines to prepare samples, 2) mass spectrometry equipment to perform the analysis of the digested sample, 3) bioinformatics tools dedicated to identification, 4) databases to be queried, 5) validation and interpretation software, 6) launching algorithms and integration software 7) data acquisition and archiving hardware system. These elements are integrated in a system that can be represented as a flow chart, relating to the experimental protocol.

Figure 4.11 shows a representation of the system integrated in our laboratories. After the 2-DE separation, the gels (or the electroblotted membranes) are stained. An image is scanned. The Melanie image analysis software allows to detect and choose protein spots of interest. A spot excision system (in our case the ARRM214BR, ARRM, Sydney, Aus) cuts the gel pieces and deposits them in microtiter plates. The plates are processed by a first destaining step, followed by an incubation with a trypsin solution and a peptide extraction procedure, using a liquid handling system (in our case

the SymBiot Workstation, Perseptive Biosystems, PE Corp., Framingham, MA). This system is able to load different MALDI MS sample plates with peptide solutions and matrix solution. The samples are then ready to be measured, in an automated mode, with a mass spectrometer (in our case, Voyager Elite MALDI-TOF MS, Perseptive Biosystems, PE Corp., Framingham, MA). After the submission of all MS data to an improved peak detection and calibration algorithm, the obtained peak tables are sent on-line, together with input parameters, to a protein identification tool (in our case PeptIdent and SmartIdent). These tools provide html result files (see Section 4.4), interpreted by a validation software that allows the user to attribute identification assignment to protein spots (CheckPeptIdent, C. Hoogland, personal communication). The update of a 2-DE database, like SWISS-2DPAGE, is performed with high degree of automation by the database manager.

The major advantages of this robotised approach are the limitation of human interventions, thus an increase in reproducibility and a decrease of the number of potential human errors. In our laboratory, this procedure currently yields a throughput of 400 fully analyzed spots per person and per week. The bottleneck is the validation step, because each individual protein identification has to be validated manually before being introduced into the SWISS-2DPAGE database.

Other similar approaches are under development in various companies. We give a non-exhaustive list here. Bruker Daltonics (<http://www.bruker.com/>) integrates a MAPII robot and a BiflexIII or a ProflexIII MS and developed AutoXecute software to handle the system, that uses MS Biotools supporting MASCOT ([www.matrixscience.com](http://www.matrixscience.com)) identification program. Micromass (<http://www.micromass.co.uk>), together with Bio-Rad (<http://www.bio-rad.com/>) implement Proteome Works System to handle an ARR214BR spot excision system, a TOF Spec-2<sup>E</sup> MALDI-TOF MS or a Q-TOF MS, and a MassLynx-ProteinLynx software package. Genomic Solutions (<http://www.genomicsolutions.com>) prepares a system including a Flexys Proteomics Robot, a ProGest Digestion Station, a Pro-MS Robot and a MALDI-TOF MS. The Protein Solution 1 package (Applied Biosystems, <http://www.appliedbiosystems.com>) integrates the Symbiot workstation with various MS instruments (the Voyager STR MALDI-TOF MS, the Voyager DE-PRO or even the Mariner API-TOF LC-MS) and the Protein Prospector (<http://prospector.ucsf.edu>) identification tools.

#### 4.6.3

##### **The molecular scanner**

Even though the robotized approach is automated and thus requires minimal human intervention, the digestion process is partially sequential, and therefore does not require that all parameters in the manipulation of

numerous protein spots are fully identical. The approach is also time-consuming and involves expensive sample handling, due to material costs. In addition, since the size of a sample is limited by the size of the excized spot, problems occur when overlapping spots are present on a gel.

We can then think of a method that parallels the protein digestion step, that reduces sample handling and that decreases the analyzed sample size to that of the MALDI-TOF MS laser beam impact (a spot of a few tens of microns in diameter).

In that respect, and to offer a flexible and powerful proteomic visualization tool, we developed a highly automated method that can create a fully annotated 2-D map starting from a 2-DE. This technology is called “molecular scanner”. It combines parallel methods for protein digestion and electro-transfers (using the one-step digestion-transfer (OSDT) or the double parallel digestion (DPD) techniques as described by Bienvenu et al. [10]) with peptide mass fingerprinting approaches to identify proteins directly from PVDF membranes, the surface of which is scanned with MALDI-TOF MS. Using a set of dedicated tools, this procedure allows the creation, analysis and visualization of a set of virtual 2D or 3D images. These images reveal particular aspects of the proteome of interest, whose complexity can be described as a multi-dimensional image (see below for the explanation of the multi-dimensionality of the proteome) [11].

#### 4.6.3.1 The principle of the method

The method can be divided into four main sections (Figure 4.12).

- A) Separation and digestion of the proteins. First, the proteins are separated on 1-DE or 2-DE gels. The proteins are digested and simultaneously electrotransferred onto a polyvinylidene difluoride (PVDF) membrane. To do that, a hydrophilic membrane, to which trypsin is covalently bound, is inserted between the gel and the collecting PVDF membrane in the electro-blot sandwich. This method is called One-Step Digestion-Transfer. The collecting PVDF membrane thus contains sets of digestion products of all proteins, each of them concentrated at discrete positions on the membrane surface.
- B) Acquisition of the peptide mass fingerprinting data. The collecting PVDF membrane is then sprayed with a matrix solution and stuck on a modified MALDI sample plate using high vacuum grease as glue. The MS then scans the membrane, i.e. a mass spectrum is acquired at each position of a predefined array. The distance between separate MS acquisition ranges between 0.2 mm and 0.5 mm. Mass spectra are acquired on a Voyager™ Elite MALDI-TOF mass spectrometer (Perseptive Biosystems, Framingham, MA, USA) equipped with a 337 nm nitrogen laser and a Delayed Extraction device. The set of co-ordinates of the laser

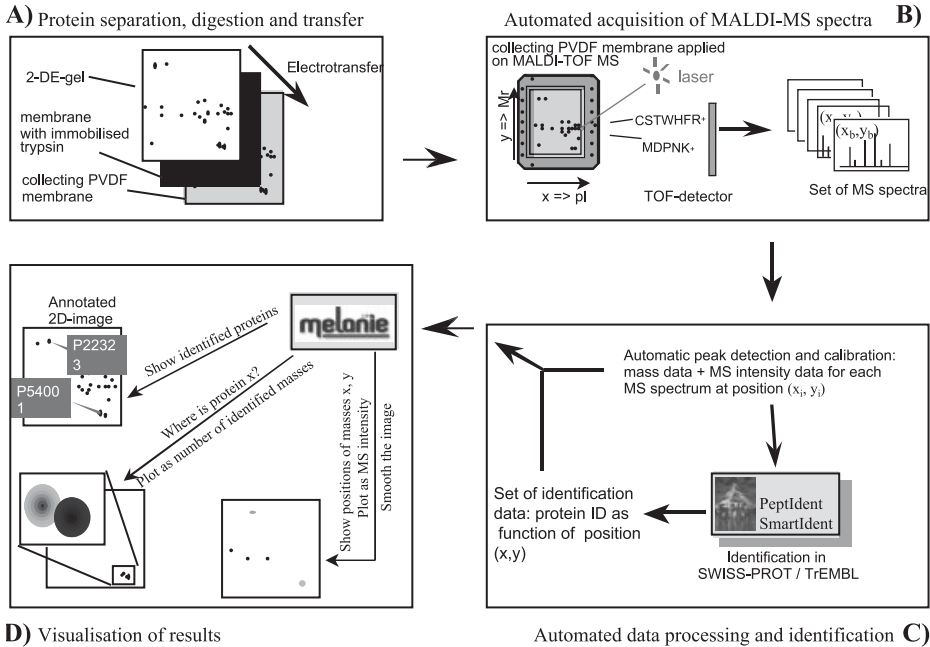
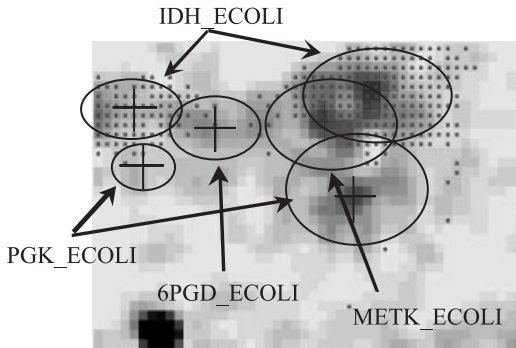


Fig. 4.12

Schematic representation of the molecular scanner process.

shots on the sample plate and the naming of the MS files is controlled by dedicated software.

- C) Processing of the MS data and protein identification. A flexible and interactive tool has been developed to automatically perform peak detection and calibration on all spectra. The positions on the sample plate are converted to apparent molecular mass ( $M_r$ ) and pI values. The PMF data of all spectra, together with the calculated pI and  $M_r$  and other user defined parameters (such as mass tolerance, considered chemical modifications, species, etc.), are automatically submitted to PeptIdent for identification
- D) Analysis of the results: creation of virtual maps. The identification results of PeptIdent are represented as an annotated image that can be read by the Melanie image analysis software. The data process program generates a first virtual “2-D-map”, i.e. a 3-D image where the  $x$  and  $y$  co-ordinates relate to pI and  $M_r$  values, respectively. The  $z$  values reflect in gray scales the intensity of the MS spectra, as defined by the sum of the intensities of the MS signals in the considered MS spectra. The image also contains the identification results, which can be highlighted as labels in Melanie. In addition, this image includes, other protein attributes that reflect the number of dimensions of the image, i.e. pI,  $M_r$ ,



**Fig. 4.13**

Interpretation of results obtained with the molecular scanner. 250  $\mu\text{g}$  of *E. coli* were separated on a mini-2DE on a narrow range pI gradient (5–5.5). The gel was submitted to DPD (full in-gel digestion followed by a through trypsin digestion).  $1.3 \times 0.9 \text{ cm}^2$  were scanned with a Voyager Elite MALDI-TOF MS. The protein identifications were performed with on-line

calls to Peptident. The area represents a pI range of about 5.1–5.25 and a  $M_r$  range of about 25–45 kDa. The grey scale intensity of this MS image represents the intensity of the MS spectra. The white dots show the positions at which the IDH\_ECOLI protein was identified. Similarly, the position of PGK\_ECOLI, 6PGD\_ECOLI and METK\_ECOLI are shown in circles.

identification labels (SWISS-PROT or TrEMBL AC numbers, ID labels), peptide masses, MS intensities as well as accesses to the detailed Peptident annotations (number of missed cleavages, annotated modifications, chemical modifications of Cys and Met residues, peptide sequences).

The user can choose to filter part of the information and visualise only particular aspects of the multi-dimensional image. Proteins or peptides can be visualized by searching the positions at which they have been identified. The  $z$  intensity can represent the number of peptides found to match the queried protein in the identification process, for instance, or the sum of the MS intensities of the peptide masses matching the queried protein (for examples of data interpretation and visualisation see Figure 4.13).

#### 4.6.4

##### Other techniques

The peptide mass fingerprinting identification method is a very accurate and quick approach, particularly using MALDI MS as an analytical tool. This method is optimal for the identification of proteins that are annotated in sequence databases. There are, however, a number of limitations to this technology. A number of small proteins are not producing a reasonable number of peptide ions to allow for unambiguous identification only by their masses. Very basic, very small (< 15 kDa) and highly hydrophobic

proteins are difficult to separate with 2-DE technology. Other type of gel-based separation can help in these cases, such as Tris-Glycine gels or 1-D SDS gels. Due to the complexity of the mixture, it is often reasonable to pre-fractionate the biological sample of interest [69].

To complement the peptide mass fingerprinting approach, a variation of protein identification by mass spectrometry, called MS/MS, is often used. In this approach, the peptides generated after endoproteolytic cleavage of the proteins are separated in a mass spectrometer equipped with a fragmentation device. From all ions of the peptide mass fingerprinting spectra, selected ions are further fragmented. In a simplistic description, these fragments produce a set of masses in the so called MS/MS spectra, the mass differences of which correspond to masses of amino acid residues. The peptide sequence can therefore be deduced. In this approach, only a few spectra of good quality are needed to unambiguously identify a protein in a database. From all available machines capable of MS/MS, the ion-trap MS and the Q-TOF MS are those whose controlling software allows for the most automatic and data-dependent sample handling. Two interpretation approaches are derived from the MS/MS technique. The peptide-sequence-tag approach [8] requires the user to interpret the MS/MS spectrum, i.e. to “read” a sequence tag associated to the spectrum. The database is searched using the sequence tag and the mass of the entire ion. Yates and colleagues [58, 60] introduced a more automated approach. Their SEQUEST program compares uninterpreted MS/MS spectra with a library of predicted MS/MS spectra from a chosen database. Those two algorithmic approaches and others can be automated and linked to a launching system. SEQUEST is integrated in a package provided with the Finnigan Corporation (<http://www.finnigan.com>) ion-trap MS instruments. A very similar approach is provided in the MASCOT software (Matrix Sciences Ltd, <http://www.matrixsciences.com>). The other programs can be reached through the ExPASy tools page (<http://www.expasy.ch/tools>). For a review on protein identification by mass spectrometry see Corthals et al. [5]. For a review on protein identification tools see Binz et al. [49].

#### 4.7

#### Conclusion

Today’s advances in technology allow us to analyze complex systems such as entire proteomes. Also, automation in proteomics generates a huge amount of heterogeneous data. This requires the integration of a Laboratory Information Management System (LIMS) connected to a full sample-tracking concept. Global solutions are still not commercially available for the proteomics field and do not cover a full system. A number of companies are currently developing such systems, like Applied Biosystems (Proteomics

Solution I and SQL LIMS) or Micromass-Bio-Rad (Proteome Works System), among others.

Comprehensive interpretation of the experimental proteomic data is only reachable in a step-by-step approach. This means that the experimentalist must have the possibility to come back to the original data at any time, in order to complete its characterization and in order to perform efficient comparison studies. In fact, as protein sequence databases are not complete, a number of identifications are failing, especially in eukaryotes. As the databases are regularly updated, identification must be regularly re-run with the incorporated data. This must regularly and automatically launch new calls to the identification tools and then update the results and the interpretation.

Moreover, even if all proteins are identified, there is the need to characterize them further. The importance of post-translational modifications in biological activity is proven, as well as that of amino-acid substitutions. These properties are often difficult to detect automatically, as they are often missing information in databases. They are, however, of potentially huge importance for the understanding of biological activity and of the presence of a diseased state.

It is now well established that combination of genomics and proteomics technologies will have a major impact in the future in: i) identifying gene products related to disease states that will further speed up the discovery of potential molecular markers and therapeutic targets, ii) understanding the mechanisms of action of drugs, including their pharmacological and toxic effects that may reveal much about the mechanisms of the disease, and iii) the discovery and design of new drugs that improve the diseased state.

In this chapter we have described a range of both wet-lab, experimental and dry-lab, bioinformatics approaches to extract biologically relevant information from one or several proteomes. In order to solve medical questions, such as “what is the molecular mechanism leading to a diseased state?” or “what kind of diagnostic test do I have to perform to decide for a diseased state?” or “what are the side effects of the absorption of a specific drug on the general state of a patient?”, the presented methods might be appropriate. Very often, though, the question cannot be answered without combining proteomics with transcriptomics (i.e. the study of the RNAs, the transcription products of genes), genomics and more classical chemistry methods. In all these cases, the researcher often generates a huge amount of experimental data. These data, alone or in combination with other researchers' experiments, contain a limitless source of information. The huge variety of databases available on the web and the amount of information included are to be compared with the experimental data. There are big challenges for bioinformatics to handle very heterogeneous data formats, to correlate highly variable types of biological information, to extract their relevance for a particular question, to perform the appropriate analyses, and to provide results in a form that is valuable for high quality interpretation.

This is the technical challenge that will lead to discoveries in prognostics, diagnostics, and drug therapies.

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## 5

# Target Finding in Genomes and Proteomes

*Stefanie Fuhrman, Shoudan Liang, Xiling Wen, and Roland Somogyi*

### 5.1

#### Introduction

The development of novel drug therapies is vital to the advancement of human health. As we move into the twenty-first century, the aging of the population will bring with it a significant rise in degenerative diseases, in particular, Alzheimer's Disease (AD) [1]. Current therapies for AD have only limited effects [2, 3, 4, 5], while the degenerative process eventually results in severe disability. In addition, no method yet exists for the regeneration of tissue.

Recently developed large-scale gene expression assay technologies such as DNA microarrays [6], SAGE (Serial Analysis of Gene Expression) [7], and RT-PCR [8], gene sequence databases such as LifeSeq® database services (Incyte Genomics, Inc.) and GenBank, and gene expression databases such as LifeExpress™ Target (Incyte Genomics, Inc.), now offer the opportunity to discover the logic of tissue degeneration and regeneration. By combining measurements for large numbers of genes expressed in parallel with appropriate data analysis methods, it should be possible to hypothesize functions for newly-discovered genes, redefine functions for familiar genes, and discover the logic of the genetic network.

In this Chapter, we will first discuss why the collection of *temporal* gene expression data is necessary for the development of therapies for degenerative disorders. We will then describe a straightforward information theoretic method for identifying drug target candidates from among thousands of genes expressed over a time course. Gene expression clustering will be discussed in the context of assigning functions to genes, and a method will be proposed for the combined use of information theory and clustering in therapeutic drug development. Finally, we will discuss the reverse engineering of genetic networks.

## 5.2

### Experimental design for large-scale gene expression studies and drug target identification

Current drug therapies for degenerative diseases such as Alzheimer's and Parkinson's Disease are of limited effectiveness, addressing only symptoms while degeneration continues. This should be no surprise given the manner in which degeneration has been approached experimentally, with a focus on individual genes at single stages of disease progression. This approach denies the very nature of degeneration as a *process* – a series of events. The examination of individual time points provides the equivalent of snapshots of the degenerative process.

Only by collecting data as time series can we observe the whole process and obtain a more complete picture of gene interactions. The technologies and analytical tools needed to accomplish this task are now available. It is now possible to measure gene expression for hundreds or thousands of genes at a time using DNA microarrays [6], SAGE [7], and large-scale RT-PCR [8].

Microarrays, of which there are different types, contain thousands of DNA probes spotted on a surface; each probe is specific for a different gene or EST. Fluorescent-tagged cDNAs from an experimental and a control tissue sample can be hybridized to the probes on the array. The cDNAs of the experimental sample are tagged with a fluorescent probe different from that of the control sample, so that the ratio of experimental to control cDNA can be determined based on color differences.

SAGE [7] involves the concatenation of short nucleotide sequence tags (9–10 bp), each representative of a different gene. The concatenated tags are then cloned and sequenced. The prevalence of a particular tag corresponds to the abundance of the corresponding mRNA in the sample.

RT-PCR can be performed in a semi-quantitative manner [8, 23] for large numbers of genes. This requires a control mRNA, which is reverse-transcribed and amplified along with the sample. Analysis of the resulting PCR products resolved with polyacrylimide gel electrophoresis, involves calculating the ratio of the sample to the control band on the gel. In this way, a relative measure of gene expression can be obtained.

The main benefit of DNA microarrays is their extremely large scale – an entire genome can be represented on a single microarray, while the advantage of SAGE is that only a short sequence is needed to identify each gene. RT-PCR provides the greatest reliability, sensitivity, and dynamic range of all the available large-scale gene expression measurement techniques; its only drawback is the problem of scaling it up with automation.

These techniques are easily employed in generating temporal expression patterns. The only requirement is the proper design of experiments. In that context, appropriate animal models of degenerative diseases must be used,

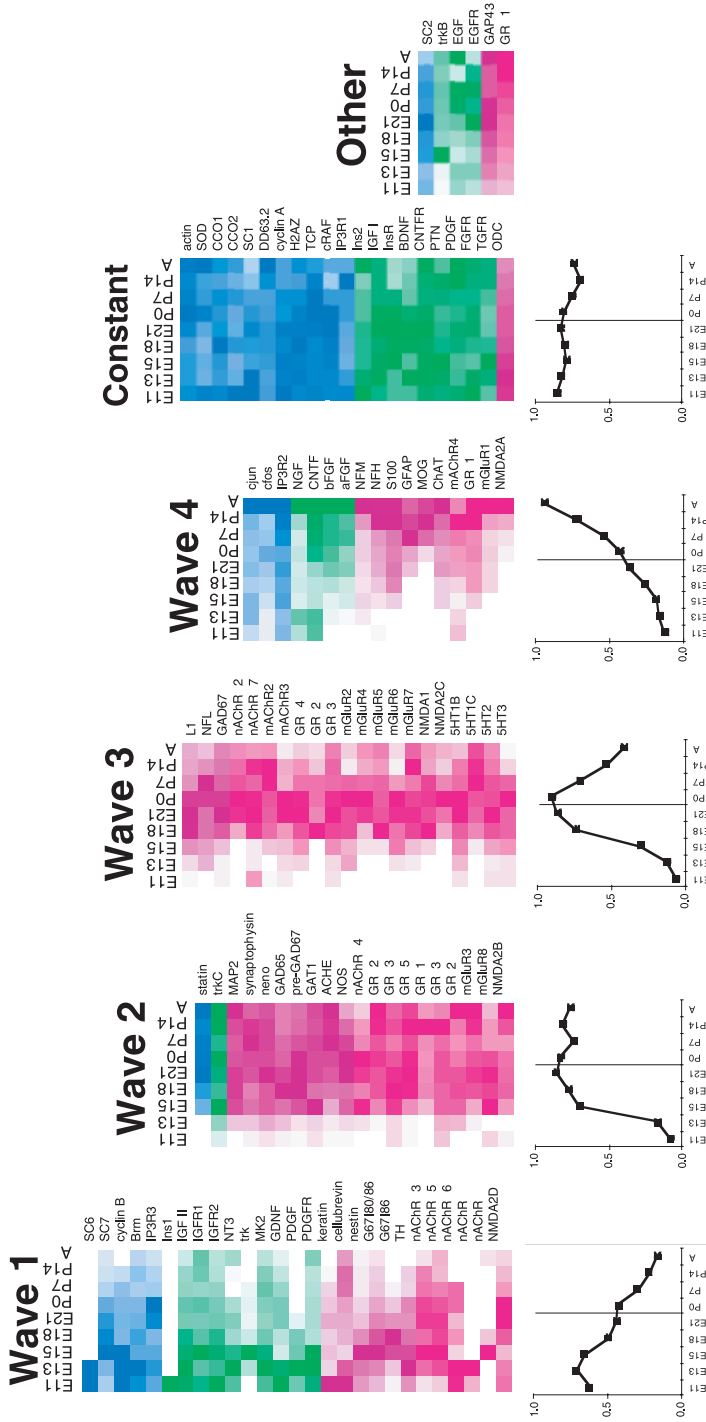
and tissue sampled at different stages of disease progression. The time intervals may be based on the results of published studies that indicate significant changes in gene expression at particular stages of the degenerative process.

Once these multiple-gene, multiple time point data have been collected, it will be necessary to organize them into an intelligible form. This can be accomplished with the right computational tools. Some relatively simple mathematical methods already in use in other fields can be adapted for the analysis of gene expression data. Two such methods discussed here include clustering [8, 9, 10], and Shannon entropy [11, 12].

Figure 5.1 contains an example of a 112-gene temporal expression pattern from the developing rat spinal cord [8], assayed using reverse-transcription polymerase chain reaction (RT-PCR). Genes from a number of recognized functional categories are represented, including a large number of genes involved in intercellular signaling (neurotransmitter receptors, peptides and peptide receptors). The data in Figure 5.1 are displayed so that they may be appreciated by visual inspection. Greater color intensity reflects higher levels of expression. These data have been normalized to maximal expression, so that for each gene, the highest expression detected is 1.0 (maximum color intensity).

The nine time points range from the earliest appearance of the spinal cord at embryonic day 11, to adult (postnatal day 90). Time intervals are based on the rate of development of the rat CNS, which slows as development progresses. It is important to note that time intervals should be spaced appropriately for the system being studied. According to [13], the embryonic rat CNS shows significant anatomical changes at 12 to 24-hour intervals; therefore, to attempt a time series with shorter intervals would be a waste of effort. Similarly, it would be inappropriate to space the time points too far apart, in which case important gene expression events could be missed. In general, published studies demonstrating significant gene expression changes at individual time points can be used as a guide in selecting intervals.

As in the case of normal development, disease progression or recovery from injury may be studied by collecting time series data. This approach is well-suited to the study of animal models of degenerative diseases. It is important to select appropriate animal models – i.e., those that provide a gradual degeneration, rather than the instantaneous destruction of tissue – to mimic human degenerative disease. Similarly, time series of regenerating animal tissue may be studied for clues to inducing this effect in human tissues that do not regenerate.



**Fig. 5.1** Clustering temporal patterns for 112 genes expressed in the developing rat spinal cord. Each data point is the average of data from three animals. For each gene, time series data are normalized to maximal expression. The higher the color intensity, the higher the expression value. White means undetectable expression. Colors distinguish among recognized functional categories: red, neurotransmitter receptors; purple, neuroglial markers; green, peptide signaling; blue, diverse. Genes are clustered into “waves,” with the average temporal expression pattern for all genes in a wave graphed below (see the later discussion of clustering). From Wen et al. [8].

## 5.3

### Computational analyses in drug target discovery

#### 5.3.1

#### Shannon entropy

Although it may be tempting to examine individual gene expression patterns and note that those patterns concur with the literature, a large data set becomes much more useful when examined in a global way using computational analysis. A certain amount of analysis can be accomplished by visual inspection, but the larger data sets such as those provided by DNA microarrays require more computing power.

One method of organizing temporal gene expression data is by applying a measure known as Shannon entropy, originally developed for analyzing telegraphic signals [11]. This is an information theoretic measure that provides the complexity or information content of a series of events. Fuhrman et al. [12] have proposed its application to gene expression data as a way of identifying drug target candidates from among thousands of genes expressed in parallel. Shannon entropy is a very direct way for pharmaceutical scientists to deal with large-scale gene expression data, and is defined as  $H = -\sum p_i \log_2 p_i$ , where  $p$  is the probability (frequency) of an event.

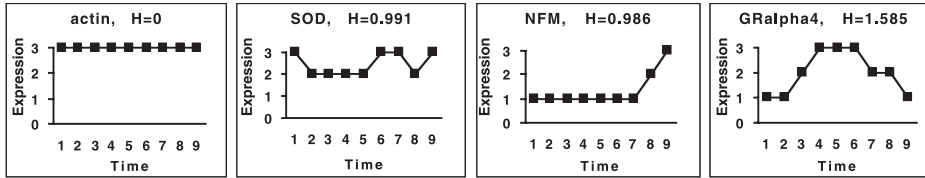
Shannon entropy is a measure of variation or change over a time series, and this is the same criterion used by biologists to determine which genes are the most interesting. Genes that exhibit significant changes are regarded as good drug target candidates, while those that remain relatively invariant in expression are more often ignored. In most cases these changes are observed in single-time point studies, such as comparisons of non-diseased versus diseased tissue. For data sets containing multiple time points, however, Shannon entropy is useful in selecting out genes with the greatest variation in expression over an entire time course.

Figure 5.2 explains how Shannon entropy works. Expression data must be binned for this calculation, which can be performed with spreadsheet software. Fuhrman et al. [14] contains instructions for calculating Shannon entropy in a computer spreadsheet. The number of bins is determined by the number of time points in the series, and should be  $\leq \log$  (base 2) of the number of time points (Bruce Sawhill, Santa Fe Institute, unpublished). So, for fewer than eight time points, two bins may be used; for eight or more time points, three bins; for 16 or more, four bins, etc. It is preferable to use a large number of time points, in part to help avoid the problem of binning artifacts caused by expression values that lie close to bin boundaries.

Once the calculation of entropy is completed, the genes can be rank-ordered. In terms of rank-ordering of genes, Shannon entropy distinguishes itself as a measure of complexity when applied to a temporal series with at least eight time points; with fewer than eight time points, Shannon entropy provides the same rank-ordering as the variance. Genes with the highest



$$H = -\sum p_i \log_2 p_i$$



**Fig. 5.2**

Shannon entropy applied to gene expression time series. The equation for Shannon entropy ( $H$ ) is shown above. For each gene, entropy accounts for the probability,  $p$  (or frequency), of a level of gene expression,  $i$ . Data from [8] (see Figure 5.1) are binned into three expression levels. SOD, superoxide dismutase; NFM, neurofilament medium; GRAlpha4, GABA receptor alpha 4 subunit. Actin has zero entropy (zero information content) because its expression is

invariant over the nine time points.

GRAlpha4 has the highest entropy because it exhibits all three levels of expression, spread out as evenly as possible over the time series (three at level 1, three at level 2, and three at level 3). A bias toward any level will reduce the entropy, as in the case of SOD (biased toward level 2) and NFM (biased toward level 1). Entropy accounts only for the frequencies of expression values, not their positions in the time series; therefore, the shapes of these patterns have no effect on entropy.

entropy can then be considered as possible drug targets for the condition being studied. In some cases, entropy may assign a low score to genes that could be considered good drug target candidates. For example, entropy will be low for genes that show a spike in expression at the beginning of a disease process, with invariant expression thereafter. There is also the possibility of false high entropy values from binning artifacts; these can be eliminated by visual inspection of expression patterns once a short list of putative drug targets has been selected. Most important, however, is that entropy provides a way of extracting essential information from an overwhelming number of gene expression profiles, allowing limited drug development resources to be focused on a small number of genes.

In neuroscience, we generally consider neurotransmitter receptors to be ideal drug targets, not only because of their accessibility, but also because of their functional specificity. The results of our entropy analysis [12] of the spinal cord data [8], are consistent with this assumption. Of the various functional categories of genes, the ionotropic neurotransmitter receptors are three-fold over-represented at the highest entropy level. At the second highest entropy level, the metabotropic neurotransmitter receptors are two-fold over-represented. Growth factors (peptides) and their receptors (receptor tyrosine kinases), on the other hand, are over-represented at several low entropy values. This suggests that neurotransmitter receptors are the major participants in spinal cord development.

Entropy can also be used to select out toxicity or diagnostic markers. In Fuhrman et al. [12], using DNA microarray data we demonstrated that certain genes are affected by toxic doses of several compounds. As in the analysis of spinal cord data, certain functional categories of genes were over-represented at high entropy. Other genes from these high entropy functional categories can be used as a focus of further toxicological research.

Another way of applying entropy is in the analysis of anatomical data. This is particularly relevant in neuroscience given the complexity of central nervous system anatomy. With the help of microarrays and other large-scale assay techniques, it is now possible to assay thousands of genes in multiple brain regions. Entropy can be applied to these data by substituting anatomical regions for time points, providing a measure of anatomical complexity that will vary from one gene to another. For example, for nine brain regions and three levels of gene expression, a gene would have the highest possible entropy if it occurred at all three levels, with each level occurring in three regions (three 1s, three 2s, and three 3s). As with time series, the shape of the distribution makes no difference. In contrast, a gene that showed no variation in expression across the nine regions would have zero entropy. The high entropy gene could be considered a major player in determining the complexity of brain anatomy. However, even in the case of anatomical studies, the time component must be considered. A gene with low entropy in its anatomical distribution could have high entropy at a later time if the brain is changing as a result of disease, injury, or normal development. Anatomical studies are therefore best approached as time series, such as development.

The use of Shannon entropy for analyzing gene expression data is a departure from traditional approaches. First, it is intended for time series, not individual data points. Further, entropy ignores a point traditionally regarded as important in the interpretation of gene expression data: the *n*-fold increase or decrease. Entropy accounts for the complexity of a time series without regard to the distance between expression levels. For example, a gene that exhibits two levels of expression could go from level one to level three, while another could go from level two to level three, yet both can have the same entropy. In regard to this, it could be argued that absolute differences in mRNA concentrations are important biochemically, but do not necessarily reflect the relative importance of genes within the system as a whole. It is reasonable to assume that each gene functions normally at its respective concentration(s), and that differences in the quantities of mRNAs or proteins reflect differences in biochemical efficiency.

Methods such as Shannon entropy will be needed in future studies of degenerating and regenerating tissue. Entropy may be used to select out the most likely drug target candidates from data on animal models of Alzheimer's and other degenerative diseases. Similarly, we can study the regeneration of animal tissues that fail to regenerate in humans (e.g., spinal

cord), and use entropy to determine which genes are the major effectors of regeneration. Some of these genes will undoubtedly be familiar – receptors for which drugs are already available – and could be tested with experimental therapeutic drugs in animal models.

### 5.3.2

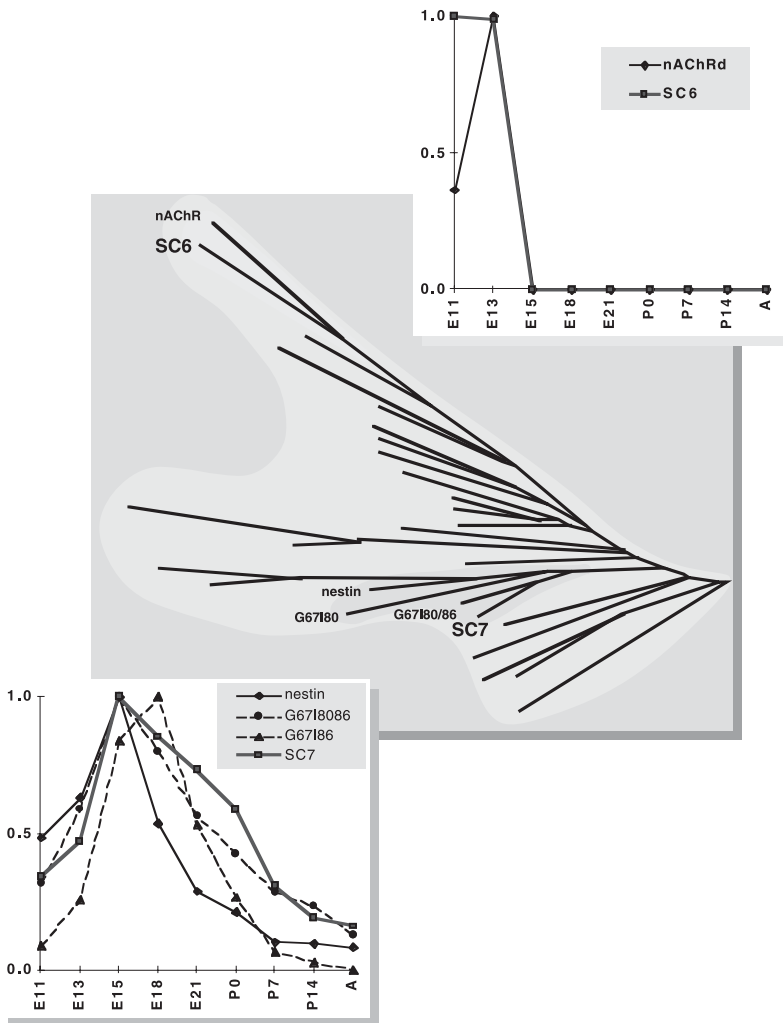
#### Clustering

Clustering is a method for grouping patterns by similarities in their shapes. This method is now being applied to large-scale gene expression patterns from eukaryotic cell cultures [15, 16, 17] and tissues [8, 17]. This technique can be used to suggest functions for newly discovered genes that have no known homologs, or to assign new functions to known genes. For example, if a new gene clusters with (has a temporal expression pattern similar to) a group of peptide receptors, the new gene may share a function with those receptors. In Figure 3, genes of unknown function are shown clustered in a dendrogram with familiar genes expressed during rat spinal cord development (also compare to wave 1 in Figure 5.1).

Before gene expression patterns can be clustered, the “distance” between each gene and every other assayed gene must be determined. This distance is based on the difference in expression level at each time point. So, as with Shannon entropy, the entire time series is taken into account. In Figures 5.1 and 5.3, the euclidean distance was used. Euclidean distance for a pair of genes is calculated by squaring the differences in expression levels between the two genes at all time points, adding the squares, and taking the square root of the sum,  $D = \sqrt{\sum (a_i - b_i)^2}$ . The completed calculations result in a “mileage chart” for all the genes assayed, and serve as the inputs for a clustering algorithm.

Clustering is based on “distance,” placing genes with highly similar patterns adjacent to each other in a dendrogram. By displaying the results in a dendrogram, one can quickly assess whether two or more genes have similar temporal expression patterns without having to search through large sets of graphs. Displays of this type become more important with very large data sets containing thousands of genes. Large cluster trees could be examined on a computer screen so that it will be possible to zoom in on a selected tree branch.

Euclidean distance is essentially a measure of positive, linear correlations; however, other similarity measures may be used for clustering. For example, *mutual information*, an information theoretic measure, may be used to capture positive, negative, and non-linear correlations all at the same time. A pictorial explanation of the concept of mutual information along with instructions on doing calculations can be found in [18]. Mutual information is based on Shannon entropy ( $H = -\sum p_i \log_2 p_i$ ; see above explanation of entropy) and is calculated as follows:  $M(X, Y) = H(X) + H(Y) - H(X, Y)$ ,



**Fig. 5.3**  
Clustering of novel with known genes [8]. The ESTs (expressed sequence tags), SC6 and SC7 cluster with neurotransmitter-related genes. Upper right panel: SC6 has a temporal expression pattern similar to that of nAChRd (nicotinic acetylcholine receptor delta subunit). Lower left panel:

SC7 shares a pattern with nestin and two splice variants of the neurotransmitter-metabolizing enzyme, GAD (glutamic acid decarboxylase). Middle panel: These genes are shown in a branch of the cluster tree for 112 genes expressed in the developing rat spinal cord.

where  $H$  is Shannon entropy, and  $X$  and  $Y$  are the two gene expression patterns being compared. Let's say that gene  $X$  has a binary expression pattern 011111000 (at each time point the gene is either on or off), and that gene  $Y$  has an expression pattern 0001100111. Because  $X$  has 40% 0s and 60% 1s,  $H(X) = -0.4 \log(0.4) - 0.6 \log(0.6) = 0.97$ .  $Y$  has 50% 0s and 50% 1s, so  $H(Y) = -0.5 \log(0.5) - 0.5 \log(0.5) = 1.00$ .  $H(X, Y)$  refers to the frequency at which each 0 or 1 in  $X$  corresponds to a 0 or 1 in  $Y$  at the same time points in the series. Zero corresponds to zero in 10% of the series, 1 in  $X$  corresponds to zero in  $Y$  in 40% of the series, zero in  $X$  corresponds to 1 in  $Y$  30% of the time, and 1 corresponds to 1 20% of the time, so  $H(X, Y) = -0.1 \log(0.1) - 0.4 \log(0.4) - 0.3 \log(0.3) - 0.2 \log(0.2) = 1.85$ . Finally,  $M(X, Y) = 0.97 + 1.00 - 1.85 = 0.12$ . Mutual information would be at a maximum if  $H$  were at a maximum for both  $X$  and  $Y$  (if both  $X$  and  $Y$  were 50% 0s and 50% 1s), and if there were a total consistency of matching between the two patterns (e.g., if  $X$  were 0011100110 and  $Y$  1100011001).

The benefit of mutual information is that it accounts for the possibility that two or more genes may respond in different ways to the same genetic regulatory input. In other words, one gene may be up-regulated and another down-regulated even though both are receiving the same input. These two regulated genes may be considered functionally related despite their divergent responses. A comparison of agglomerative clustering using a mutual information and Euclidean distance measure is presented in [10]. In conclusion, the results obtained by clustering are largely determined by the distance measure used.

There are many algorithms for clustering. The most popular methods for gene expression analysis are *agglomerative clustering* and the *k-means algorithm*. In the agglomerative clustering method, one first defines the similarity measure between two gene expression profiles, which is usually the Euclidean distance of normalized gene expression. The algorithm groups genes most similar to each other in a pairwise fashion. First, two genes closest to each other among all pairs of genes are joined together to form a new node (the distance between the genes determines the length of the line joining them in the dendrogram). The distance between the new node and all other genes are updated. Next, the two genes that are closest to each other are selected and joined together. This process continues, each time the closest pair is chosen, until all the genes are exhausted. This process results in a tree with branches that provide a hierarchical classification for the gene expression data.

Another popular algorithm is the *k-means* method, which partitions genes into  $n$  clusters. Each gene is initially assigned to a cluster. This assignment is then adjusted by reassigning a gene to the cluster whose center is closest to it. Since the center of the cluster also changes, the process is iterative. One can determine the right number of clusters by performing this procedure with different numbers of clusters until a certain clustering quality criterion is optimized.

A criticism sometimes heard about this approach is that cluster boundaries are arbitrary. Naturally, someone must decide where on a dendrogram one cluster ends and another begins. These decisions are based on non-computational judgements, as are decisions about whether a particular clustering method provides a result that an observer would have produced by visual inspection alone. In that context, we have found that by including in our calculations the slopes between data points, the clustering results “improve” – the results are more similar to what one would expect based on visual examination of the expression patterns (unpublished observations).

One method for assessing the quality of a clustering method is to establish the correlations between expression clusters and known functional families, gene sequence families, or gene clusters from other experiments (Figure 5.1). If the clusters capture genes known to be related according to other functional criteria, one may conclude that the clustering algorithm has identified meaningful biological groupings. However, different clustering methods may provide differential matchings with alternative systems of classifying genes, e.g. sequence families, functional families, shared promoter elements (unpublished observations). The truth is that computational analysis of any kind is only a guide for interpreting data, with human judgment as the final arbiter.

Clustering allows us to form new hypotheses about gene function by grouping genes not previously associated. However, this technique alone is not sufficient for identifying new therapeutic drug target candidates. In that context, it may be useful to combine clustering with other analytical methods, such as Shannon entropy.

### 5.3.3

#### **Combining analytical methods in the development of experimental therapies**

The development of new therapeutic drugs is no easy task given the amount of testing required to bring a drug to market. According to the Pharmaceutical Research and Manufacturers of America (1996), only about 1 in 1000 drug candidates ever makes it past the preclinical stage of testing. Considering the effort required to discover and develop new drugs, the importance of using highly efficient methods for drug target identification becomes clear. This is especially true in the development of combinatorial therapies, in which two or more drugs are administered simultaneously. Large-scale gene expression surveys provide a wealth of possible drug targets, any of which might be used in combination therapies. The result of all these possibilities would be a combinatorial explosion – too many combinations to deal with in the laboratory.

As we explained previously, Shannon entropy is a first step in narrowing the range of drug target candidates from hundreds or thousands of genes expressed in parallel down to a more manageable number for laboratory

research. With this method, there is no need to assume that any gene perturbed by a disease or condition is a good drug target. Entropy narrows the field to those genes with the most complex expression patterns over the entire course of a disease, with high entropy genes being the biggest participants in the disease process. A previous study demonstrated that the number of high entropy genes in a perturbed system is quite small (about 5–8%) (12).

At this point, clustering may be applied to the high entropy short list. In the case of a 10 000-gene microarray assay, this list might contain approximately 500 genes. Clustering would identify groups of high entropy (drug target candidate) genes that may function in parallel. In that context, it is reasonable to assume that genes that cluster together have a high probability of functioning together as drug targets. The coupling of Shannon entropy with clustering may therefore be useful in identifying drug targets for use in combinatorial therapies.

#### 5.3.4

##### **How these methods were selected**

In selecting analytical methods, the most important consideration is whether the mathematics fits the biological problem. In the case of applying Shannon entropy to drug target discovery, the *primary* consideration is the *biological definition* of a drug target. Shannon entropy has a similar definition, and is suitable for application to time series. Similarly, in the case of clustering, the biological assumption that genes that fluctuate in parallel may share a function is the basis for using a method that groups genes by expression similarities. The continued generation of large-scale data sets will provide biologists and computer scientists with ample opportunities to collaborate on the application of these and other techniques.

#### 5.3.5

##### **Welcome to the future: reverse engineering of genetic networks**

Is it possible to solve the “ultimate question” of inferring the major connections of the genetic network of molecular interactions based on well-designed experiments using large-scale gene expression data? Such a global reverse engineering strategy would revolutionize the elucidation of disease pathways and concomitant drug target discovery. Current research in this area is very encouraging; it has been demonstrated that this strategy is theoretically feasible [18], and results from initial experimental studies appear plausible [19].

Beginning from first principles, genomic sequence information is the evolutionary memory of an organism or species that allowed it to execute

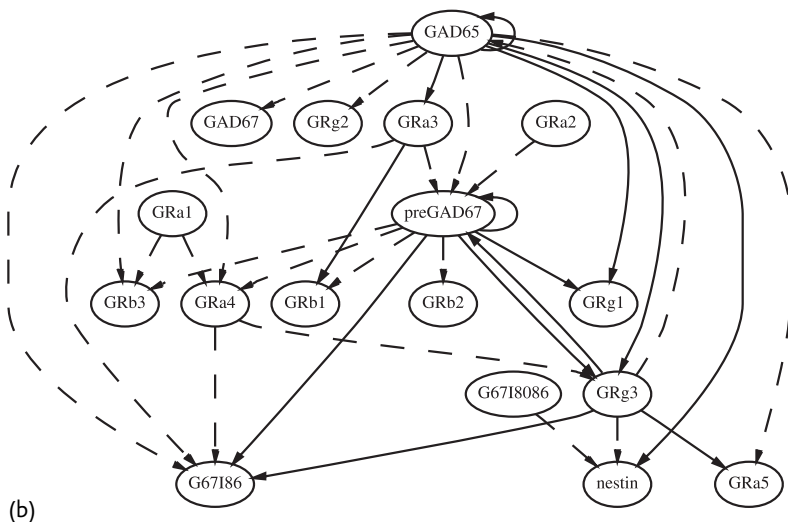
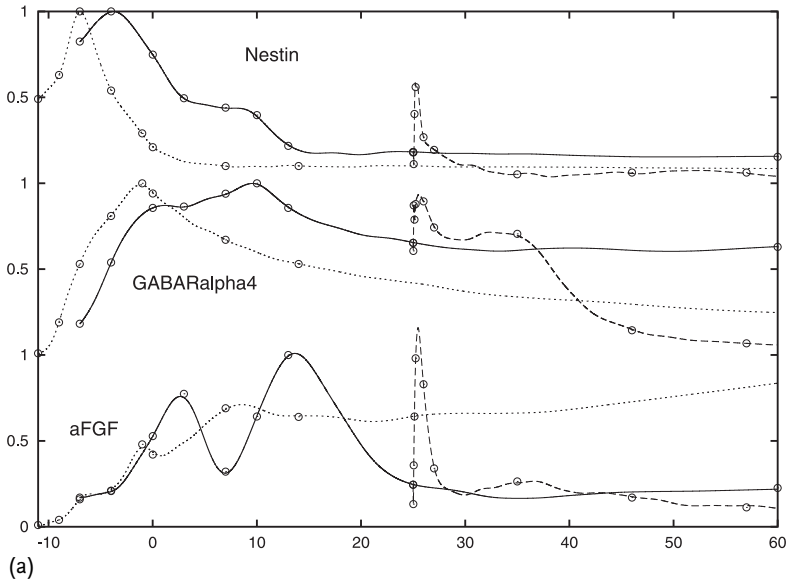
the functions required to persist in its environment. Essentially, this “static” information is expanded into the “dynamic” information of the molecular (genetic) network underlying development and many of its behavioral responses. While only the complex “hardware” of the cell is able to correctly interpret the genetic programs encoded in sequence information, we may consider an inverse approach of inferring the connections and dynamic rules of molecular interactions through their activity patterns. This defines the term “reverse engineering.”

Reverse engineering has been demonstrated to work in principle for model genetic networks of binary genes connected through logical rules [18]. A key issue is the data requirement necessary to provide sufficient information to capture the complexity of the molecular network. In model networks it has been shown that only a tiny subset of all possible behaviors need to be known in order to infer network architecture with accuracy [18], provided that the network exhibits significant constraints (biomolecular networks are far removed from randomly connected networks) [20].

Reverse engineering has been successfully applied to relatively simple biomolecular systems. Using a combination of cross-correlation analysis and multidimensional scaling, the glycolytic pathway was reconstructed from metabolite activity data from an *in vitro* enzymatic reactor system [21]. A complete spatio-temporal model of developmental gene expression in *Drosophila* was constructed for a small gene set based on models of differential equations and protein expression data [22].

There are too many unknowns to currently assess whether reverse engineering will work with certainty on large-scale gene expression data that is just becoming accessible with current technologies. In the empirical tradition, this should by no means preclude earnest attempts. One such study employed a simple linear model of gene interaction to supply a sufficient number of constraints that make parameter fitting possible given the present limitation on data complexity. Data complexity not only involves the numbers of genes assayed, but the diversity of conditions under which these activities are measured. In this example (Figure 5.4), temporal expression data from CNS development (two central nervous system areas) and CNS injury (one brain area) covering 27 time points was used to fit a linear gene interaction matrix using multiple regression [19]. This strategy proved successful in that the original data were accurately reproduced by the model (Figure 5.4a), and that major functional gene interactions could be proposed (Figure 5.4b). Interestingly, the conclusion of a positive feedback interaction for two of the genes studied (GAD65 and preGAD67) was reached independently in another work on the developmental modeling of these genes [23]. On the whole, these predictions provide important clues for further experimental investigation. Improving the reverse engineering strategy through acquisition of more informative data and testing of different models holds much promise for a systematic discovery of gene interactions, therapeutic targets and drug function. Overall, the implications of





**Fig. 5.4** Reverse engineering of a CNS genetic network using a linear model. Experimental gene expression data (circles; development and injury), and simulation using a linear model (lines). Dotted line: spinal cord, starting at day -11 (embryonic day 11). Solid line: hippocampus development, starting at day

-7 (embryonic day 15). Dashed line: hippocampus kainate injury, starting at day 25. b) Gene interaction diagram for the GABA signaling family inferred from developmental gene expression data (spinal cord and hippocampus data). Solid lines correspond to positive interactions, broken lines suggest inhibitory relationships. Reprinted with permission from [19].

the reverse engineering strategy for a systematic and efficient therapeutic design could be revolutionary.

### 5.3.6

#### Genomes and proteomes

Gene expression (mRNA) data may be limited in that we generally lack the corresponding protein expression or protein activity data, and few systematic investigations exist regarding whether RNA expression and protein activity are well-correlated [24]. This will undoubtedly have to wait for the development of better technologies for large-scale protein assays. For the present, we may make the parsimonious assumption that there is a positive correlation between gene and protein expression. As technologies advance for the measure of protein expression and activity, it will be possible to apply the same analytical methods used for genomics data.

## 5.4

### Concluding remarks

We now have the technological capacity to conduct experiments that address whole physiological processes, rather than mere snapshots of biological events. However, taking full advantage of this technology requires the proper experimental designs and data analysis methods. While it is possible to use large-scale gene expression assay methods for purposes far below their capacity, this would be a waste of an opportunity to study biomedical problems that were unapproachable prior to the genome projects.

Making efficient use of large-scale gene expression surveys for drug target identification is an exercise in selecting out the essentials of the resulting data. This requires computational methods that address time courses as well as large numbers of genes. Collaborations between biologists and computer scientists will be essential for the development of future data analysis methods.

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## 6

### Screening of Drug Databases

Martin Stahl, Matthias Rarey and Gerhard Klebe

#### 6.1

##### Introduction

The discovery of new and innovative lead compounds is one of the key elements in a drug development project. In consequence much effort has been devoted to various techniques to support and improve this procedure. At present, the establishment of compound collections at pharmaceutical companies, augmented by parallel and combinatorial chemistry, provides large libraries of molecules that can be searched for leads<sup>1)</sup>. High-throughput facilities for compound screening can generate biological activity data for many targets in very short time [1]. In the early stage of high-throughput screening this new technique has been greeted with much enthusiasm and the end of any rational and knowledge-based approaches was predicted [2]. Today, several years later, a more realistic view has been accepted. Despite the introduction of highly sophisticated robotics and advanced engineering techniques to automate high-throughput screening, such biological testing is not without problems. False positives and unspecific binding of test candidates together with low solubility, non-specific reactions with the protein material resulting in surface adhesion or protein precipitation are only some of the problems that puzzle scientists. Success rates reported for the translation of apparent hits from HTS into leads to be used for lead optimization are quite depressing. At the 1999 Cambridge Healthtech Institute meeting on high-throughput technologies, no pharmaceutical company could report a successful lead candidate from an HTS hit [3]. From a knowledge-based point of view, however, it is most disappointing that high-throughput screening can hardly contribute to our understanding of why and how the detected hits act upon the target. Any increase in the under-

1) A *lead compound* is a structure that marks the first step in the development of a drug. *Lead finding* is the process of selecting a lead compound from a compound collection or library. *Lead*

*optimization* is the process of modifying the lead compound in order to enhance properties such as specificity, bio-availability, and others.

standing of drug action on a molecular level is produced only once experimental structural biology or molecular modeling come into play to detect structural similarity or possible common binding modes among the obtained actives. Such analyses are prerequisite for a rational approach to lead finding and for any further optimization of leads to potential drugs.

As an alternative and complementary approach to experimental high-throughput screening *virtual* screening has emerged. This term comprises a large variety of computational tools to select potentially active and bio-available compounds from libraries [4, 5]. These tools range from fast filter criteria and crude molecular similarity measures to sophisticated molecular docking and scoring techniques. Which of these tools are applied in a given project depends on the knowledge assembled about a given target and the binding properties of its ligands. Furthermore, the usage of individual tools is restricted by the size of the databases to be screened.

While experimental high-throughput screening already relies on an enormous stock of compounds available to automated test robots, virtual screening could possibly screen any imaginable molecule. The universe of possible drug-like molecules is a tremendously large compound space – there may be around  $10^{100}$  drug-size organic molecules [5] – from which virtual libraries of any imaginable size could be generated. For reasons of fast substance acquisition and easy validation of the initial computer hits, however, libraries of existing or easily accessible molecules are screened first. A prominent example is the Available Chemicals Directory (ACD) [6], a collection of more than 200 000 commercially available compounds. Other databases, for example the World Drug Index (WDI) [7] or the MDL Drug Data Report (MDDR) [8], were compiled from compounds synthesized and studied for their action upon a biological target. The Beilstein database [9] contains a collection of several million entries of organic molecules for which the chemical synthesis has been described. Another fruitful application of virtual screening tools is the selection of building blocks of combinatorial libraries. Each combinatorial library offers fast synthetic access to a large variety of compounds out of a limited set of building blocks. Computers afford generating, handling and recording the vast variety of possible members of combinatorial libraries. Accordingly, the design and selection of a targeted combinatorial library can ideally be combined with criteria derived from virtual screening experiments.

As a first stage in a virtual screening experiment it is reasonable to apply fast filter criteria to eliminate compounds with undesired physicochemical properties. Many drug development projects have failed and still fail in a later state due to insufficient pharmacokinetic properties determining drug absorption, distribution, metabolism, excretion, and toxicity (ADMET parameters) of the developed drug candidates [10]. At present, the various criteria determining these properties are still poorly understood. However, an increasing body of knowledge is collected and allows for the definition of structural restraints that molecules have to meet for being a potential

drug. Often, the structure-activity relationships corresponding to the various ADMET parameters are orthogonal to the ones responsible for drug binding to the target under consideration. Accordingly, criteria to select and filter compound libraries to focus on candidate molecules corresponding to acceptable ADMET parameters should be applied at an early stage of the computer screening process. Examples of simple filters are molecular weight thresholds, the total number of donor and acceptor groups [11], or neural networks that have been trained to recognize structural patterns increasing the general “drug-likeness” of compounds [12].

Subsequent steps of compound selection in virtual screening are based on knowledge about the criteria responsible for binding to a particular target and involve concepts of molecular *similarity* or *complementarity*. Methods directly involving the complementarity of receptor and ligand are referred to as structure-based methods, because they require that the three-dimensional structure of the target be known, for example, by the determination of its crystal structure, by NMR experiments or via homology modeling. If this is the case, the geometry of the binding-site can be used as a “mold” for calculating possible binding modes of putative candidate molecules and to select molecules based on their fit to the binding pocket. Usually, this is done by means of fast flexible docking algorithms (see Chapter 7 of Volume 1). If no 3-D structure of the target is available, the bioactive conformation of a ligand can likewise be used to search for active compounds by means of algorithms that produce relevant superpositions with respect to a predefined pharmacophore. Both docking and superpositioning methods require fast approaches to detect the features that are responsible for topologically diverse candidate molecules being similar with respect to their recognition in the same portion of a binding pocket. At each step of this process a reliable *score* of the achieved placement of a candidate molecule into the binding site or onto a reference ligand has to be computed. Accordingly, an essential prerequisite for the success of any virtual screening approach is a robust scoring function to rank computer-generated binding modes or to estimate the degree of similarity obtained by a mutual superposition of a candidate molecule onto a reference ligand.

Molecular docking and structural alignment methods are based on three-dimensional structures of candidate molecules that can be generated by rule-based methods such as CORINA [13, 14] or CONCORD [15]. However, methods based on three-dimensional structures are computationally quite demanding and cannot routinely be applied to databases of hundreds of thousands of compounds today. For this reason, a number of fast methods to determine molecular similarity have been developed that operate solely on connectivity and atom types of molecules. Such tools allow rapid pre-screening of very large databases and avoid a conformational analysis of each candidate molecule.

In this chapter we give an overview of currently used virtual screening methods. The performance of some of these tools is then illustrated by

means of test libraries consisting of a subset of the WDI and collections of compounds active at several pharmaceutically relevant targets. In a first test scenario, a single active compound per target is used to retrieve similar compounds from a database. By this strategy, the performance of a number of fast similarity algorithms is compared. In the second scenario, it is assumed that only the receptor structure and its binding site are known. Test databases are docked into the corresponding binding sites and ranked according to the score computed for each candidate molecule. The results of such database rankings give an impression of strengths and weaknesses of current docking and scoring tools.

## 6.2

### Methods for virtual screening

In this section we give an overview of tools frequently used for virtual screening of drug databases. Based on whether the three-dimensional structure of the target protein is available or not, protein structure-based and ligand similarity-based screening methods are distinguished. In similarity-based methods compounds are selected by their similarity to a reference molecule known to bind to the target protein. In contrast, in structure-based methods compounds are selected according to their fit to the active site of the target protein (see Chapter 7 of Volume 1 for an introduction to docking techniques).

#### 6.2.1

##### Ligand similarity-based virtual screening

The experimental determination of protein structures is a difficult task and is not routinely possible. Especially membrane-bound proteins are difficult to crystallize [16]. Therefore, the situation that the target protein structure is unknown is common at an early stage of a pharmaceutical project. In such cases, only the structure of molecules already known to bind to the target protein, for example an endogenous ligand, can be used for virtual screening.

When a molecule binds to the protein, it adopts a conformation that sterically and electrostatically fits into the active site. In order to detect compounds possibly binding into the same active site, one can search for molecules that can adopt conformations similar to the given active molecule. This similarity is defined with respect to size, shape and spatial distribution of functional groups and can be regarded as “chemical similarity” [17–19]. In principle, however, we are interested in describing “biological similarity” in the sense that molecules exhibiting the same biological



response will be considered as similar. Biological similarity, however, depends on many unknown factors such as details of the protein structure and, therefore, cannot be computed. Chemical similarity can be regarded as a necessary but insufficient criterion for similar biological response of compounds. Fortunately, searching for chemically similar compounds has been demonstrated to be highly successful in many applications [20–22].

Methods for similarity-based virtual screening can be classified by the kind of information used to define similarity. While topological descriptors are based on the 2-D structure (connectivity, bond orders and atom types) only, 3-D descriptors take into account the spatial arrangement of atoms or their associated properties with respect to each other.

#### 6.2.1.1 Topological (2-D) descriptors

The 2-D structure of a molecule can be used to compute bit strings describing the topology of a molecule in a piecewise manner. Originally introduced to solve the problem of fast substructure searching [19], today, bit strings are the most widely applied descriptors for fast similarity searching. Two approaches to create bit strings can be distinguished: structural keys and fingerprints. In a structural key, each bit represents a molecular fragment that is set to 1 if the fragment is present in a molecule [23]. Although this method performs very well in practice, it has the limitation that a reasonable set of fragments has to be predefined. In order to avoid this lack of generality, molecular fingerprints were introduced [24, 25]. The patterns for a molecule's fingerprint are generated from the molecule itself. Each bit represents a sequence of connected atoms having specified atom types, called a *path*. The fingerprinting algorithm generates a list of all paths up to a specified length. For each path, a hash function calculates the location of the bit that has to be set. This means that a particular bit represents not just one specific bond path, but several paths. The length of the fingerprint can be reduced by a process called folding. The shorter the length of the fingerprint, the more different paths are represented by the same bit and the less specific information is encoded. Examples for programs based on bit string descriptors are structural keys by MDL [26], Daylight fingerprints [25], and UNITY 2-D screens by Tripos [27]. The latter two are used in the application study presented in this paper. Therefore they will be described in more detail.

**Daylight fingerprints** (DF) [25] are bit strings generated from bond paths of length zero to seven. The length of the fingerprints can be folded until a specified density of set bits is reached. Alternatively, the fingerprint length can be set to a fixed value. The similarity index used is the Tanimoto coefficient, which is the number of bit positions set to 1 in both strings divided by the number of bit positions set to 1 in at least one of the strings. If a set bit is

considered as a feature present in the molecule, the Tanimoto coefficient is a measure of the number of common features in two molecules [28, 29].

**UNITY 2-D screens** (UF) [27] are a mixture of fingerprints and structural keys. Unity allows the user to specify the individual components of the bit string, which can be paths within a specific length range or a variety of sub-structures. For this study, the default 988-bit strings were used. About 94% of the bits in the string are reserved for encoding paths of lengths 2 to 6 using atom type and bond type information. Another 3% of the bits in the string are reserved for atom counts of frequently occurring hetero atom types and the remaining 3% are used for important fragments such as benzene rings and heterocycles. The Tanimoto coefficient is used as a similarity index.

Several other descriptors based on the topology of molecules were developed and have been successfully applied. Reviews on descriptor technology can be found in [19, 20, 22, 30]. The following discussion focuses on three recently developed descriptors used in the application study in section 3. The selection of descriptors was based on availability and speed of the software.

**Correlation vectors** (CV) [31] are created by first assigning a generalized atom type to each atom. In the implementation applied here [32], five atom types were used: hydrogen-bond acceptor (A), hydrogen-bond donor (D), positively charged (P), negatively charged (N) and lipophilic (L). In a second step, each pair of atoms in the molecule is classified by their assigned generalized atom types and the number of bonds by which they are separated. The correlation vector, a string of integers, is a histogram representation of how often each possible pair of atom types is found at each distance. For bond paths with up to  $n$  bonds this results in correlation vectors of  $n$  times 15 integers. In order to achieve a size-independent description, each integer is divided by the total number of non-hydrogen atoms in the molecule. The Euclidean distance between vectors is used as a similarity index. Values of  $n = 6$  through  $n = 10$  were used in this work and did not give significantly different results. Therefore,  $n = 6$  was chosen for comparison with other methods.

The methods described so far have in common that they create a one-dimensional representation of the molecule from its 2-D structure, either a bit string or an integer vector. The advantage of this representation is that, in order to calculate a similarity index, there is no need to assign parts of two molecules or molecule representations to each other. This results in simple and extremely fast comparison algorithms. However, since molecules are three-dimensional objects, substantial information is neglected and, in consequence, some accuracy is sacrificed.

The following two descriptors are not based on a one-dimensional representation and, therefore, need an assignment procedure for the similarity calculation. The first descriptor is based on a matrix representation, the

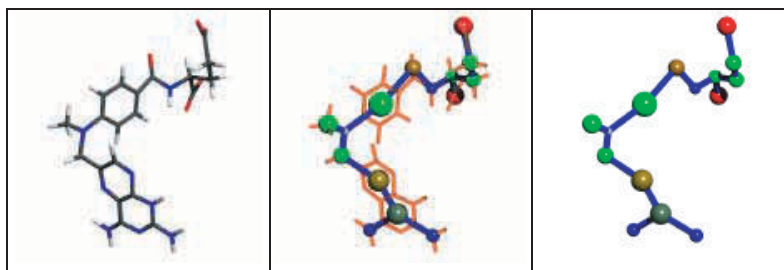
second is based on a tree representation. The two methods also have in common that they read a single 3-D conformation of each input molecule.

**Topological pharmacophores** (TP) [33] are based on data generated by the force field MAB [34, 35]. Hydrogen-bond strengths are assigned to atoms on a continuous scale. The atoms are then classified as hydrogen-bond donors, hydrogen-bond acceptors or hydrophobic. Hydrophobic atoms are grouped together according to a fixed maximum inter-atom distance criterion. A molecule is described by a set of donor, acceptor and hydrophobic centers and a matrix containing the through-space distances between these centers. Topological pharmacophores thus depend on the input conformation to some extent. To arrive at a similarity index for two molecules, centers of the same type are matched onto each other, leading to several alternative pairings for all centers. For each of these pairings, a similarity score between zero and one is calculated based on hydrogen-bond strength and the differences in mutual distances of the matched centers. The final similarity index is the highest similarity score calculated for one of the pairings.

**Feature trees** (FT) [36] describe the overall topology of a molecule with a tree structure. They are independent from the input conformation of the molecule. Starting from the molecule graph, the corresponding feature tree is created by shrinking rings as well as terminal atoms with their neighbors to single nodes (see Figure 6.1 for an example). A steric and a chemical feature are assigned to each node, describing the part of the molecule represented by the node. The approximated van der Waals volume is used as a steric feature. The chemical feature is an interaction profile summarizing which interactions can be formed by the molecule part. In order to calculate a similarity index for two feature trees, nodes of the trees have to be assigned to each other. This is done by the so-called split-search algorithm, which divides and assigns subtrees in a recursive fashion. In this way, the algorithm calculates an assignment between nodes by optimizing the similarity index.

#### 6.2.1.2 Three-dimensional descriptors

While the previously discussed topological descriptors are based on the molecular graph only, a three-dimensional descriptor uses information about the spatial arrangement of the atoms or functional groups of a molecule. Since the process of complex formation between a receptor and a ligand is mainly driven by steric and electrostatic surface complementarity, methods based on 3-D information seem to be more appropriate than topological features, at first sight. However, in practice it has often been observed that topological descriptors outperform 3-D descriptors [20, 30, 37]. A general problem with 3-D descriptors is handling conformational space. In principle, all low-energy conformers of a molecule would have to



**Fig. 6.1** A molecule and its corresponding feature tree representation. Feature tree nodes are colored by the interaction profile. Red: hydrogen bond acceptor, blue: hydrogen bond donor, green: hydrophobic. Mixed colors correspond to mixed profiles.

be taken into account, but conformational flexibility is often neglected and the descriptor is calculated for a single conformation only. On the other hand, it is likely that most data sets used for benchmarking similarity measures are biased towards methods based on 2-D features, because inhibitors are often designed and synthesized in terms of series comprising common structural elements.

As in the 2-D case, there is a wide range of different descriptors (see [38–40] for reviews). Most of them are one-dimensional representations, i.e. they project 3-D information into a bit- or integer string. In most cases the analysis of the 3-D structure is reduced to so-called pharmacophore elements or “hot spots” in the ligand structure that are most relevant for specificity and activity. Hydrogen bond donors, acceptors, and hydrophobic centers as well as their mutual distances are commonly used to address a bit in the string. This pair approach can be extended to triplets, where a characteristic value of the formed triangle (circumference, wing area, etc.) instead of the distance is used for addressing [41]. The approach can also be applied to conformation ensembles by disjunction of bit strings created from different conformations [42]. As a logical extension of such 3-point pharmacophores, 4-point pharmacophores have been used [43].

In contrast to the previous approaches, some other descriptors are based on superimposing 3-D structures [44–46], leading to more time-consuming comparison algorithms. An example are so-called field descriptors. They are derived from the analysis of superimposed molecules on a rectangular grid, whose elements contain local properties of each molecule [47]. Other similarity calculations are based on fragment superposition [48, 49]. An interesting new approach for similarity calculations is the determination of affinity fingerprints [50, 51]. Here, the affinity to a reference panel of proteins is determined for each compound in the database – either experimentally or by means of docking. Similarity indices are then calculated by comparing the affinity vectors. The underlying idea is that compounds

showing similar binding properties to the reference panel of proteins will also display similarity with respect to unknown targets.

## 6.2.2

### Structure-based virtual screening

In structure-based virtual screening, compounds are selected based on their fit into the active site of a protein. Two components are necessary for structure-based virtual screening, a docking algorithm and a scoring function.

#### 6.2.2.1 Fast docking methods

Docking algorithms require the 3-D coordinates of the target protein and of a candidate molecule from the database (the ligand) as input and predict the geometry of the formed molecular complex. The algorithm has to determine several degrees of freedom, namely the relative orientation and the conformation of ligand and protein. It is assumed that the protein conformation does not change significantly upon binding, since the protein is large and the active site has a concave surface. Furthermore, the structure of a ligand-bound complex is often used as a starting point. Experience shows that conformational properties are conserved much better among ligand-bound protein structures than among the apo (unbound) form and any ligand-bound case. However, the ligand conformation can undergo drastic changes compared to the solvated state. Algorithms considering the ligand conformation in addition to the relative orientation of the molecules are classified as flexible docking algorithms. Some algorithms handle protein flexibility simultaneously, but they have not yet been used for screening purposes.

A wide range of different methods has been applied to solve the docking problem. A review on protein–ligand docking algorithms is given in Chapter 7 of Volume 1 or elsewhere [52–58]. Here only the FlexX method used in the application study is outlined very briefly. FlexX [59–65] is based on an incremental construction algorithm. The method is based on the assumption that conformational flexibility of a ligand can be handled by dividing the ligand into smaller fragments at its rotatable bonds and reassembling the fragments within the active site. The algorithm consists of three phases. First, a set of so-called base fragments is selected. A base fragment is a part of the ligand which is not too flexible and is able to bind to the protein via directed interactions like salt bridges, hydrogen bonds, or directed hydrophobic interactions. The base fragments are then placed into the active site of the protein. For this step a pattern recognition algorithm is used to directly create energetically favorable placements. Finally, the remaining fragments of the ligand are successively connected to the base fragments.

Each time a new fragment is added, it is rotated around the single bond generated in the previous addition step to explore its conformational space. Then a set of high-scoring partial placements is selected for the next build-up step.

### 6.2.2.2 Scoring protein–ligand complexes

The second integral part of structure-based virtual screening is the scoring function. It operates on the coordinates of the protein–ligand complex and determines a score based on the generated ligand binding conformation and orientation. In an optimal case the score shows a high correlation to the observed binding affinity of the ligand to the protein. Unfortunately, the objective to reliably predict free energies of binding can hardly be achieved. The main reason for this limitation is the fact that important contributions to the binding affinity such as solvation effects or changes in entropy-determining ordering parameters upon complex formation are only poorly understood [66]. Nevertheless, scoring functions available today allow for achieving a good separation of true actives and inactive compounds when they are applied in virtual screening. Overviews on scoring functions can be found in the literature [67–72]. Here only the two scoring functions used in the application study in Section 6.3 will be described.

For scoring protein–ligand complexes in FlexX, a slightly modified version of the Böhm function [73] is used. The function belongs to the class of regression-based empirical scoring functions [71]. Empirical scoring functions are linear combinations of terms, each of which is a simple geometrical function derived from the ligand and receptor coordinates. The terms are chosen such that they intuitively cover important contributions to the total binding free energy. The modified Böhm function is a linear combination of seven terms, namely the entropy loss due to immobilizing torsional degrees of freedom in the ligand upon complex formation, neutral and charge-assisted hydrogen bonds, interactions of aromatic ring systems with other groups, lipophilic and ambiguous contact surfaces:

$$\begin{aligned} \Delta G = & \Delta G_0 + \Delta G_{\text{rot}} N_{\text{rot}} + \Delta G_{\text{hb}} \sum_{\text{neutral H-bonds}} f(\Delta R) f(\Delta \alpha) \\ & + \Delta G_{\text{io}} \sum_{\text{ionic interactions}} f(\Delta R) f(\Delta \alpha) \\ & + \Delta G_{\text{aro}} \sum_{\text{aromatic interactions}} f(\Delta R) f(\Delta \alpha) + \Delta G_{\text{lipo}} \sum_{\text{lipo atom contacts}} f^*(\Delta R) \\ & + \Delta G_{\text{ambig}} \sum_{\text{ambig. atom contacts}} f^*(\Delta R) \end{aligned}$$

Each term sums individual contributions scaled by geometric penalty functions  $f$ . For example, the contribution of a hydrogen bond falls into ranges

from  $\Delta G_{\text{hb}}$  to zero, depending on its deviation from idealized hydrogen-bond geometry (length and angle at the hydrogen atom). The relative weights of the individual contributions (the  $\Delta G$  values) have been calibrated on protein–ligand complexes with known structure and binding affinity.

An alternative approach is realized in the DrugScore function [74], which belongs to the class of knowledge-based scoring functions. The basic idea of knowledge-based scoring is that interatomic contacts occurring more frequently than in a mean reference state are energetically favorable. Based on this assumption, pair potentials can be generated from occurrence frequencies counting the distribution of atom-type pairs as a function of their inter-atom distances. The DrugScore function consists of two terms, the inter-atomic potentials and a solvent accessible surface (SAS) term:

$$\Delta W = \gamma \sum_k \sum_l \Delta W_{\text{type}(k), \text{type}(l)}(r_{kl}) + (1 - \gamma) \left[ \sum_k \Delta W_{\text{type}(k)}^{\text{lig}}(S_k^c, S_k^u) + \sum_l \Delta W_{\text{type}(l)}^{\text{prot}}(S_l^c, S_l^u) \right]$$

$\gamma$ : adjustable parameter

$\Delta W_{\text{type}, \text{type}}$ : pair potential

$S_{\text{atom}}^{\text{form}}$ : SAS of an atom in complexed or uncomplexed form

$\Delta W_{\text{type}}$ : SAS single atom potential

$k$ : index for ligand atoms

$l$ : index for protein atoms

The first term is the actual pair potential yielding a distance-dependent score contribution for each protein–ligand atom pair within a given distance interval. The SAS-term contains single atom contributions covering solvent effects. In the application described in the following, the DrugScore function was not used in the incremental construction phase, but only to re-score the 500 best scored FlexX solutions for each protein–ligand complex. In addition, the FlexX entropy penalty term was added to the score for each ligand.  $\Delta G_{\text{rot}}$  was set to 14 000 to adjust to the order of magnitude covered by the knowledge-based scores.

## 6.3

### Practical virtual screening

As outlined in the introduction, virtual screening in pharmaceutical industry is a tool to find new lead compounds for a target. In a realistic situation,

one or several ligands of the target may be known – from screening, from the literature or from patents – and the goal is to find different compounds with similar inhibitory properties. Alternatively, instead of or in addition to known inhibitors, the target 3-D structure may be available. The similarity measures or docking methods introduced in the previous section can then be used to rank order compound databases according to the likelihood that an individual compound will be an inhibitor. Since a complete and unambiguous separation of active and inactive compounds is not achievable, the term *enrichment* is used here. A database search can be considered successful if the top ranks of the ordered database are *enriched* with active compounds, i.e. more densely populated with active compounds than can be expected from a random distribution of hits. In the next two sections of this chapter, the ability of individual methods to enrich active compounds is assessed by means of test databases comprising known active and inactive compounds.

## 6.4

### First test scenario: application of fast similarity searching algorithms

In this test scenario, a single active compound per target is used to retrieve similar compounds from a database. Five fast algorithms are used for this purpose. Results are compared and possible ways to combine the results of searches with different algorithms are discussed.

#### 6.4.1

##### Library generation

Diverse sets of inhibitors were compiled manually for eleven well-established pharmaceutical targets. The main – subjective – selection criterion was to cover as many different compound classes as possible for each target. Compounds were retrieved from the WDI and from available complex crystal structures in the PDB [75]. Where patent literature was consulted in addition, references are given in the following list. The final data set consisted of:

- 58 ACE inhibitors [76]
- 43 angiotensin II antagonists [77]
- 33 aldose reductase inhibitors [78, 79]
- 32 cyclooxygenase inhibitors [80–82]
- 55 H1 antihistamines [83]
- 41 HIV protease inhibitors [84]
- 21 HIV reverse transcriptase inhibitors [85]



- 19 monoamine oxidase A inhibitors [86, 87]
- 23 gelatinase A inhibitors [88]
- 36 thrombin inhibitors [89, 90] and
- 37 topoisomerase II inhibitors [91].

All compounds were stored as SMILES and converted to single 3-D conformations in Sybyl mol2 format<sup>2)</sup> by means of CORINA [13, 14]. A C routine was used to generate the molecular structure most likely to be the dominant species at neutral pH under physiological conditions. Hydrogen atoms were removed from acidic groups to generate the anionic form, and were added to aliphatic amines and acyclic amidine nitrogens to generate the charged cationic form.

A subset WDI database was prepared by stepwise removal of

- compounds containing one of the mechanism keywords “ACE”, “aldose-reductase”, “ampa”, “angiotensin”, “antihistamine”, “beta-lactamase”, “cyclooxygenase”, “hiv”, “mao”, “thrombin”, “topoisomerase” (in order to ensure complementarity to the individual sets of active compounds in this and other studies),
- compounds with molecular weight greater than 800 or less than 200 (because drug-like molecules should fall in this molecular weight range)
- compounds with saturated carbon chains longer than seven carbon atoms (in order to avoid non-drug-like alcohols and fatty acids)
- compounds without at least one oxygen (very few drugs contain no oxygen atom)
- compounds without at least one hydrogen (to exclude poly-halogenated compounds)
- compounds containing elements other than C, N, O, P, S, and halogens (again to ensure the drug-likeness of all compounds)

The remaining WDI compounds were clustered according to Daylight fingerprint similarity by means of the nearest-neighbors clustering algorithm [94] implemented in the Daylight toolkit [95] considering the 14 nearest neighbors and requiring at least 8 cluster members per cluster. One compound per cluster was selected, resulting in a database of approximately 10 000 compounds. Further selection steps included the removal of macrocycles with larger than 12-membered rings, approximately 70% of the molecules with a steroid-type scaffold (because they were over-represented in the selected subset) and of molecules with more than 13 pharmacophore centers as determined by the TP software described in Section 6.2.1. This step essentially serves to remove large molecules with many polar func-

2) Both SMILES [92] and mol2 [93] are specific formats to store small molecule structures. SMILES encode the 2-D structure of a molecule in a text string,

whereas the mol2 format assigns a specific atom type class to each atom according to its environment and stores 3-D coordinates.

tional groups that are unlikely drug candidates. The final size of the database was 7528 compounds, which were stored as a SMILES list and converted to Sybyl mol2 format [96, 97] by means of CORINA. Protonation states were set as described above for the sets of active compounds.

Although care has been taken to remove those compounds of the WDI that are inhibitors of one of the targets used in the following calculations, this removal is certainly not complete. Nevertheless, to a good approximation the WDI compounds can and will be regarded as inactive in all calculations presented in the following.

#### 6.4.2

##### **Computational details**

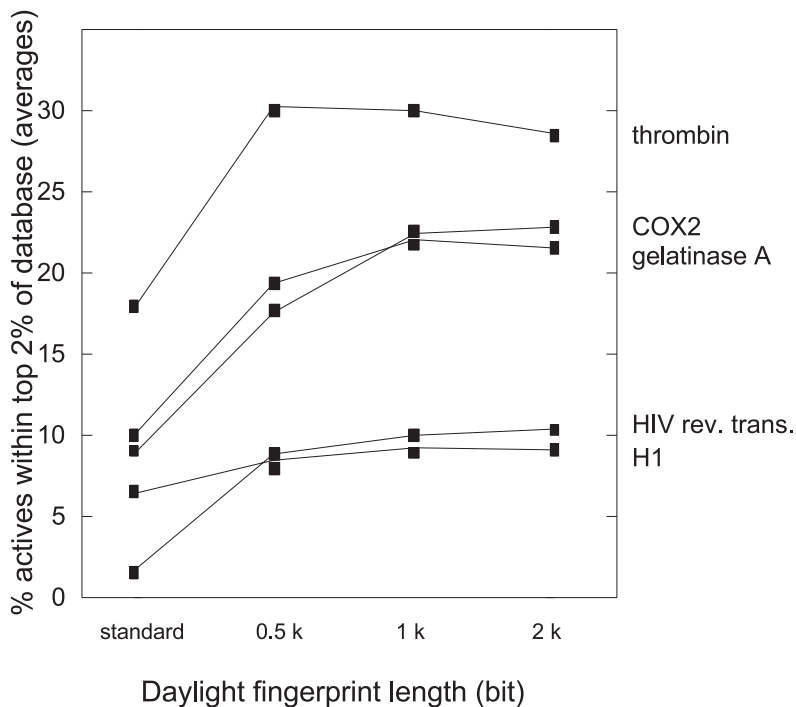
The 398 molecules in the 11 inhibitor classes were successively selected as query molecules. In each run, the complete database of 7925 compounds (7528 WDI compounds plus the remaining 397 active compounds) was ordered from high to low similarity with respect to the query molecule according to the five different algorithms described in Section 6.2: Daylight fingerprints, UNITY 2-D screens, correlation vectors, topological pharmacophores and feature trees. This leads to five different ranking lists per molecule that were subsequently analyzed with respect to the distribution of compounds belonging to the same class as the query molecule (“active compounds”).

The Daylight software was used with four different settings regarding the length of the fingerprints (default, 512 bit, 1 kbit and 2 kbit). In general, we have observed that a performance increase for longer fingerprints becomes visible at low similarity levels (Tanimoto coefficient < 0.9). Fingerprint length is a critical issue in the assembled database. Standard fingerprints (whose size varies from molecule to molecule until a fixed density of set bits is achieved) performed significantly worse than those of constant length, as shown in Figure 6.2 for five selected targets. Fingerprints of 1 kbit length were chosen for comparison with the other methods, because increasing the size to 2 kbit does not improve performance. Standard Unity 2-D screens also have a length of approximately 1 kbit.

#### 6.4.3

##### **Discussion of screening results**

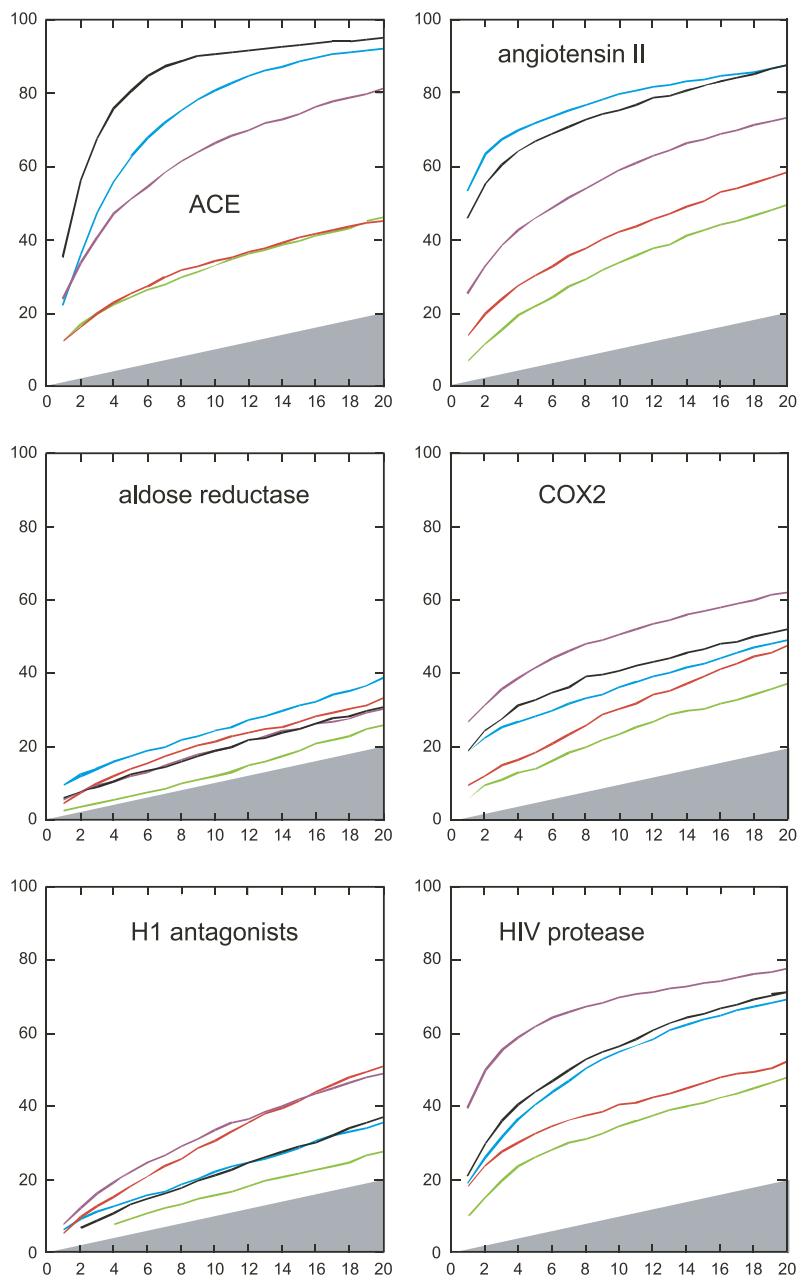
The performance of the five methods was compared by averaging, over all members of the class, the percentage of active compounds contained in each percentile of the ranked database. The results, graphically depicted in Figure 6.3, vary strongly depending on the inhibitor class. For some inhibi-



**Fig. 6.2** Performance of Daylight fingerprints of different length in retrieving active compounds related to a given query molecule. Fingerprints of 1kbit length are a reasonable choice for the present test scenario.

tor classes several algorithms generate very satisfying results, while for others all algorithms perform poorly.

On average, many active compounds receive top ranks in the ACE, angiotensin II and HIV protease classes, while enrichment is very poor for aldose reductase, H1 antagonists, HIV reverse transcriptase and MAO A. Results of intermediate quality are obtained for COX2, gelatinase A and thrombin. The two fingerprint-based methods (DF and UF) and the feature trees software (FT) outperform correlation vectors (CV) and topological pharmacophores (TP) in database ranking. The CV method performs worst in all cases. However, there is no single method that performs best for all eleven inhibitor classes. The FT method seems to be superior to the other methods if correct matching of hydrophobic groups connected by single bonds is important. For example, this is the case with COX2 inhibitors, the majority of which consists of three planar ring moieties whose centers span an angle of about 60°. As a point in case, consider the three structurally related COX2 inhibitors 1–3 in Figure 6.4. The differences in the central ring lead to different sets of bond paths and thus to different fingerprints



**Fig. 6.3**  
Average percentage of active compounds versus percentage of the ranked database plotted for eleven targets and five different fast similarity algorithms.

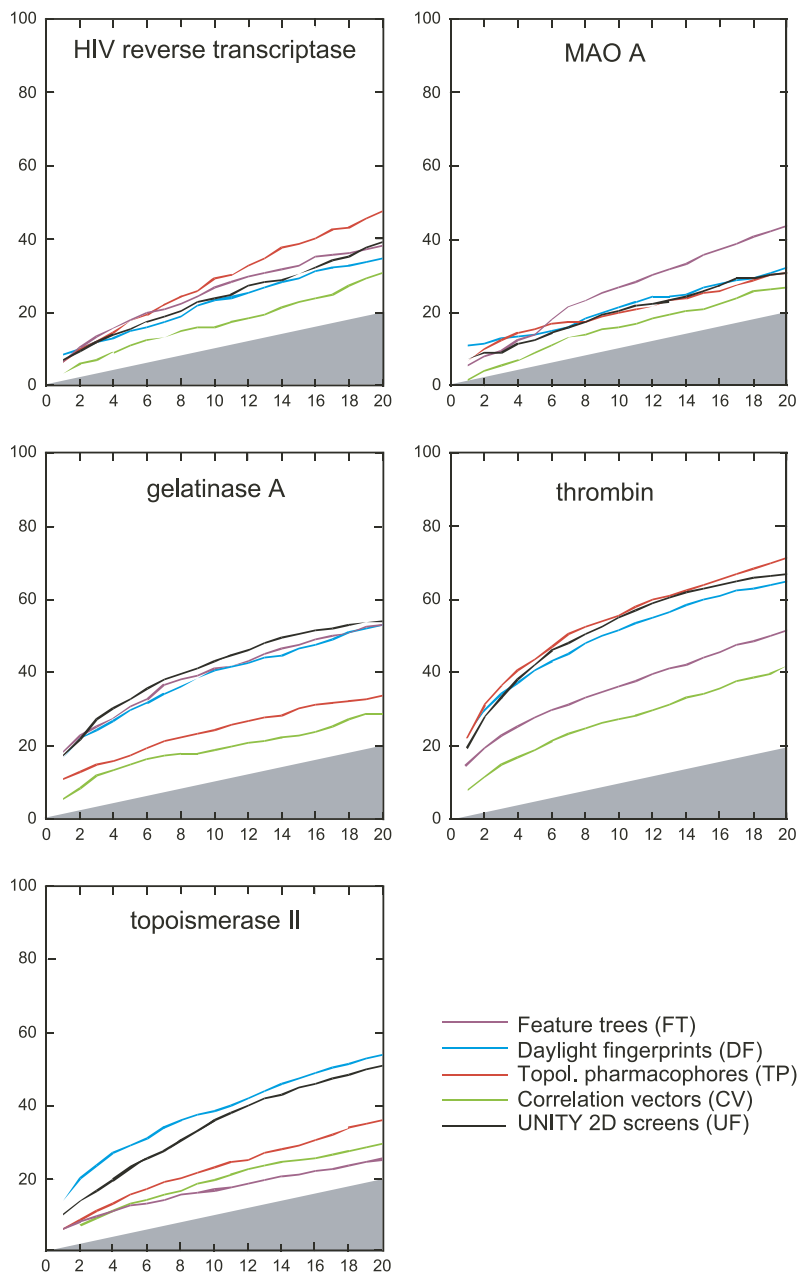
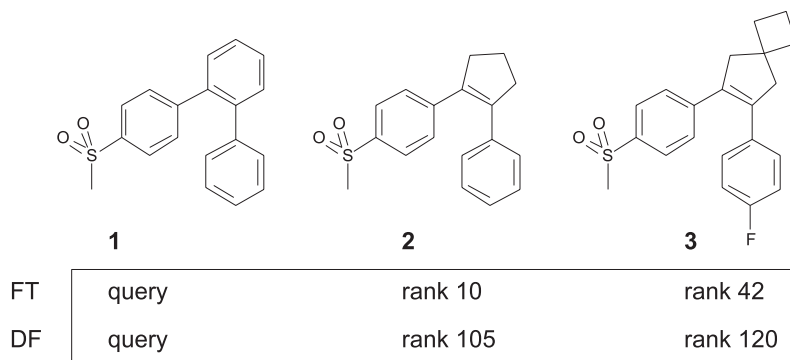


Fig. 6.3 (continued)

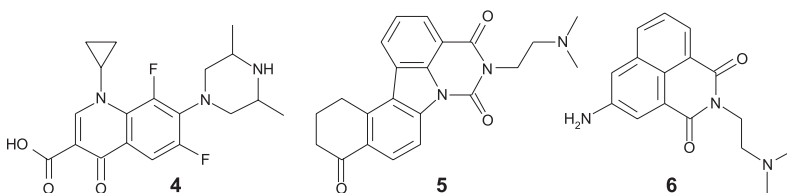
**Fig. 6.4**

Three structurally related COX2 inhibitors. Ranks for compounds **2** and **3** were calculated using the FT and DF methods with compound **1** as a query molecule.

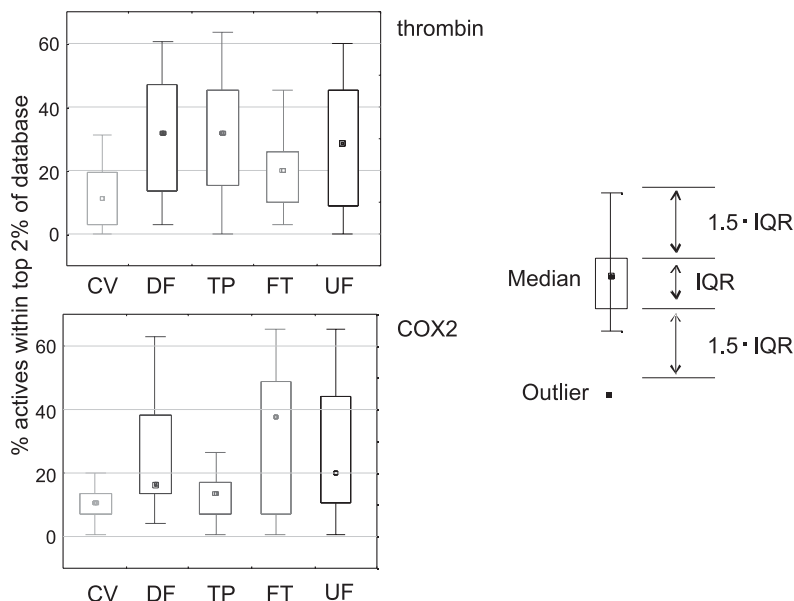
whereas, for the FT software, this difference is of minor importance, since the relative orientation and polarity remains the same for all three compounds. The comparison of fused aromatic and heterocyclic compounds is an inherent shortcoming of the FT method, as is obvious from the topoisomerase II results. To impart an impression of topoisomerase II inhibitors, three representatives of this class are shown in Figure 6.5. Finally, TP emphasizes matches of hydrogen-bond donors and acceptors more than the other methods do. For this reason, the TP method performs well for sets of polar molecules like thrombin inhibitors.

Plots of average enrichment, for example as shown in Figure 6.3, can be misleading, because they can easily be influenced by outliers. They do not give information on the variation of enrichment achieved for different members of a target class. This variation is high for all methods and target classes: A search with a single active molecule can retrieve many related active compounds or no actives at all. Small structural changes in the query molecule can significantly change the database ranking.

How strongly the enrichment varies depending on the query molecule can be illustrated by means of Box plots [98]. As illustrated in Figure 6.6 for

**Fig. 6.5**

Three topoisomerase II inhibitors



**Fig. 6.6** Box plots showing the variation of the percentage of active compounds contained in the top 2% of the ranked database. Results are shown for the COX2 and

thrombin libraries only. In the legend on the right, IQR is the inter-quartile range, i.e. the interval of a data set that is centered around the median and contains 50% of the data points.

thrombin and COX2, the percentage of active compounds within the top 2% of the database varies between zero and very high values. For thrombin, the average values shown in Figure 6.6 seem to indicate that the order of performance for the five methods is TP, DF, UF > FT > CV, but from the Box plots it is apparent that the difference between CV and FT is actually very small. In fact, it is insignificant at an error level of 5%, as can be deduced from statistical analyses of the data by means of the Tukey test [99]. Similarly, the plots for COX2 in Figure 6.3 suggest the order FT > UF, DF > TP > CV. Consulting the Box plots one must conclude that the median percentage of actives retrieved is almost as low for DF as it is for CV and TP. According to the Tukey test, at an error level of 5% the only robust statements about the COX2 data are that FT performs better than CV and TP.

From the above, it can be concluded that it is difficult to predict which of the methods will work best for a particular query molecule. To increase the success rate, it is therefore advisable to perform searches with two or more different methods. But how should the results be combined? Two possible strategies to select compounds from two searches employing methods A and B are:

1. Selection of all molecules that receive high ranks with either method A or method B. For each compound occurring in both the A and B rank lists, the position with the higher rank is selected and the combined list is sorted again according to rank. In cases of conflict, compounds are placed in arbitrary order. For this *union strategy* to be successful, methods A and B should emphasize different aspects of similarity and should therefore retrieve different active molecules on top ranks. It is then likely that more hits will be found in the union rank list than in any individual rank list.
2. Selection of all molecules that receive high ranks with both method A and method B. Such a procedure is successful if a common set of active molecules is ranked high by both methods A and B, while the top-ranking inactive compounds differ. This *intersection strategy* will be followed if it is important to reduce the number of false positives – even at the cost of reducing the number of true positives.

For the present test libraries, the union strategy was tested by counting the number of actives among the top 2% of the individual rank lists in comparison to those among the top 2% of the union rank list. On average, the union strategy leads to significantly more hits than the weaker individual method, but it is rarely superior to the better performing method. Table 6.1 illustrates this observation for the ACE sublibrary. The union strategy thus does not retrieve more actives, but, since it is difficult to estimate in advance which method will perform better, it leads to more robust results. A generally valid recommendation is to combine one of the fingerprint-based methods with any of the other three algorithms.

An example for the application of the intersection strategy is given in Table 6.2 with the antihistamine 7 (Fig. 6.7) as a query molecule.  $C_{\text{tot}}$  is the

**Tab. 6.1**

Results of the union strategy for the set of ACE inhibitors: Table entries are the average numbers of active compounds contained in the top 2% (158 compounds) of the union rank lists. Values on the diagonal (combinations of each method with itself) are the average number of actives in the top 2% of the individual rank lists.

	<i>FT</i>	<i>TP</i>	<i>CV</i>	<i>DF</i>	<i>UF</i>
<i>FT</i>	19				
<i>TP</i>	17	9			
<i>CV</i>	18	12	10		
<i>DF</i>	22	17	18	21	
<i>UF</i>	27	23	24	28	32



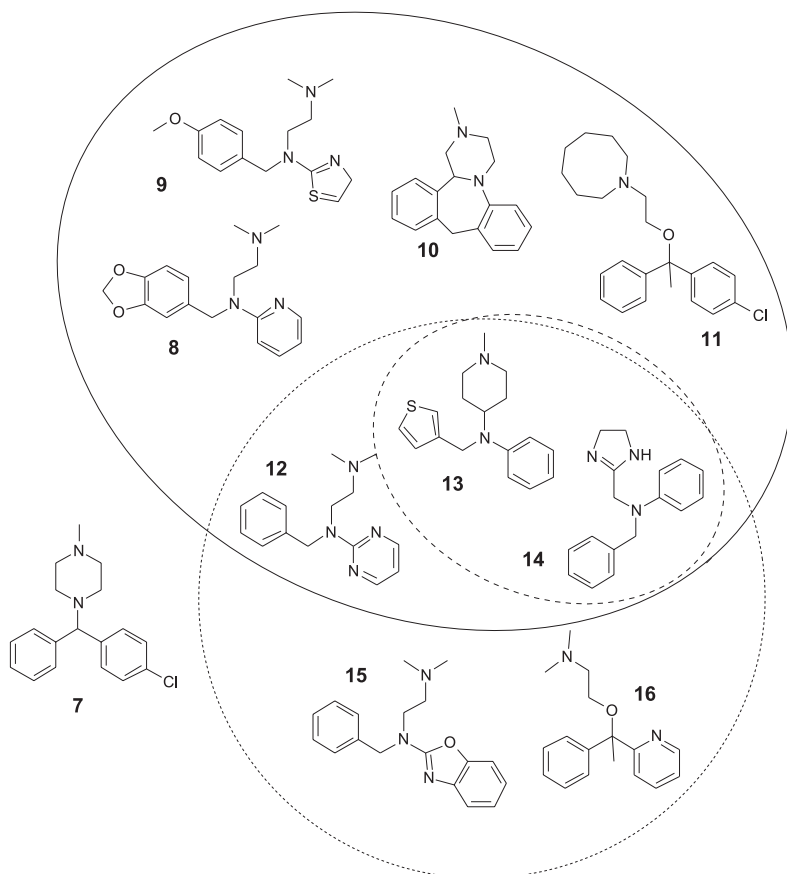
Tab. 6.2

Results of the intersection strategy for query molecule **7** in Figure 6.7:  $C_{\text{tot}}$  is the total number of common molecules in the top 2% of both rank lists,  $C_{\text{act}}$  is the number of H1 antagonists contained therein.  $I_{\text{act}}$  is the number of active molecules contained in the  $C_{\text{tot}}$  top ranking molecules of the library ranked according to the method given in the leftmost column. Diagonal elements of this table give the number of active molecules within the top 2% of the ranked database for each method (where  $C_{\text{act}} = I_{\text{act}}$  and  $C_{\text{tot}} = 158$ ).

	FT			TP			CV			DF			UF		
	$C_{\text{tot}}$	$C_{\text{act}}$	$I_{\text{act}}$	$C_{\text{tot}}$	$C_{\text{act}}$	$I_{\text{act}}$	$C_{\text{tot}}$	$C_{\text{act}}$	$I_{\text{act}}$	$C_{\text{tot}}$	$C_{\text{act}}$	$I_{\text{act}}$	$C_{\text{tot}}$	$C_{\text{act}}$	$I_{\text{act}}$
FT	158	13	13	19	4	5	8	3	3	12	6	4	16	7	5
TP	19	4	2	158	8	8	3	1	0	9	4	1	12	3	1
CV	8	3	0	3	1	0	158	5	5	11	3	0	13	3	0
DF	12	6	2	9	4	1	11	3	2	158	8	8	90	7	6
UF	16	7	2	12	3	2	13	3	2	90	7	5	158	8	8

number of molecules occurring in the top 2% of both rank lists; and  $C_{\text{act}}$  is the number of active molecules contained therein. From Table 6.2 it can be seen that the proportion of active molecules in the intersection lists is relatively high compared to the top 2% of the individual rank lists (values on the diagonal of Table 6.1). This comparison is not fair, however, since most of the active molecules should be found on the highest ranks of the individual rank lists. Considering only a single rank file, one would select only very few top-ranked compounds in order to keep the number of false positives to a minimum. Therefore,  $C_{\text{act}}$  is best compared to the number of actives among the  $C_{\text{tot}}$  top ranked compounds of each individual ranking list. This number is given as  $I_{\text{act}}$  in Table 6.2.

In the example in Table 6.2,  $C_{\text{act}}$  is larger than  $I_{\text{act}}$  for most combinations, which means that the intersection strategy identifies more active molecules than any individual algorithm. The efficiency of the intersection strategy is smallest for the combination of the two fingerprint methods DF and UF, because the two rank lists overlap significantly. For all other combinations, between 1/4 and 1/3 of  $C_{\text{tot}}$  are active compounds. Figure 6.7 shows results of the combination of UF and FT. Seven active compounds (**8–14**) are contained within the intersection list. Compounds **13** and **14** are among the top 16 compounds of both the UF and the FT rank lists. In addition, the top 16 compounds of the FT rank list also contain compounds **12**, **15** and **16**. The latter two compounds are not on the intersection list, since they are on rank positions beyond 158 on the UF rank list. The example presented in Table 6.2 is representative for the data used in this study. On average, the intersection strategy significantly reduces the number of false positives. Best results are achieved if two methods are combined that perform well sepa-



**Fig. 6.7**  
Results of the intersection strategy for the H1 receptor antagonist **7** as a query molecule. Compounds **8–14** are contained within the 16-membered intersection list of the DF and FT rank lists. Compounds **13** and **14**, encircled by a dashed line, are

within the top 16 molecules of the FT and DF rank lists, meaning that they are regarded as highly similar to the query molecule by both methods. The top 16 molecules of the FT list also include **12**, **15** and **16**.

rately. The combination of FT and one of the fingerprint methods can be generally recommended.

## 6.5

### Second test scenario: docking as a tool for virtual screening tool

In this scenario, it is assumed that only the receptor structure and its binding site are known. Test databases are docked into the corresponding binding sites and ranked according to the score computed for each candidate

molecule. The enrichment of active compounds on top ranks is analyzed and reasons for performance differences between the test targets are discussed.

### 6.5.1

#### Library generation

Five of the 11 sets of inhibitors described in Section 6.4 were used again in the second scenario, namely the thrombin, gelatinase A, HIV protease, HIV reverse transcriptase, and COX2 data sets. A set of 29 p38 MAP kinase inhibitors was assembled from PDB crystal structures and Ref. [100]. A set of 40 estrogen receptor ligands was taken from the Protherics web site [101]. These two data sets were converted to Sybyl mol2 format and protonation states were adjusted as described above.

### 6.5.2

#### Docking procedure

For each of the six targets, protein structures were retrieved from the PDB or the Roche in-house collection of protein structures. Binding pockets were defined manually using the interactive modeling program MOLOC developed at Roche. For thrombin the PDB complex 1dwd was selected and the water molecule adjacent to Tyr 228 in the S1 pocket included as part of the binding site. For gelatinase A, the only available X-ray structure was that of a proenzyme mutant (1ck7). Since only minor structural changes can be expected upon complexation and the active site region can be well superimposed onto other MMP structures, the structure was used in an “*in silico* activated” form. The N-terminal propeptide was removed and residue 404 mutated to Glu as in the wild-type enzyme. For HIV protease, the PDB complex 9hvp was selected. The flap water bound to Ile50A and Ile50B was optimized by means of the MAB force field [102] and hydrogen atoms were added by MOLOC. Docking was performed both neglecting and including this water molecule as part of the binding site. PDB structures were also selected for reverse transcriptase (1klm) and the estrogen receptor (1ere and 1err). For p38 MAP kinase as well as COX2, Roche in-house crystal structures were used.

For each target, the corresponding set of inhibitors was combined with the WDI subset and the resulting library docked into the binding site. All docking calculations were performed with the program FlexX using standard parameter settings, except where noted otherwise. Calculations were run in parallel on 10 SGI R12k processors, typical runs for 7600 compounds taking 10–15 hours of CPU time. The structure with the lowest score was saved for each compound and its score was used for database ranking.

**Tab. 6.3**

Results of database ranking by means of docking for six targets. Table entries are the percentage of all active compounds retrieved within the top 2%, 5% and 10% of the ranked databases consisting of the WDI subset and one collection of active compounds.

<i>target</i>	% <i>actives within top</i>		
	<b>2%</b>	<b>5%</b>	<b>10%</b>
thrombin	57	81	89
gelatinase A	21	42	61
p38 MAP kinase	17	24	55
COX2, standard	0	0	13
COX2, lipo scoring	13	19	25
HIV protease, no flap water	8	8	17
HIV protease, flap water	14	28	41
estrogen rec. (1err), FlexX	35	38	43
estrogen rec. (1err), DrugScore	35	45	63
HIV reverse transcriptase	5	10	14

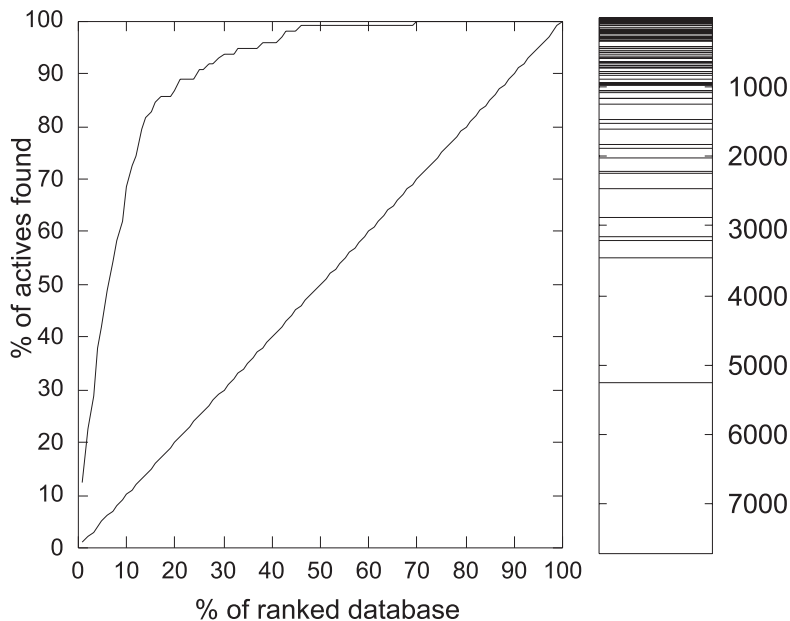
Compounds for which no docking solution was obtained were appended to the sorted rank list in arbitrary order.

### 6.5.3

#### Discussion of docking results

Results of the docking calculations are summarized in Table 6.3, which contains the percentage of active compounds within the top 2, 5, and 10 percent of the ranked databases. Docking results for p38 MAP kinase are also graphically depicted in Figure 6.8. For thrombin, p38 MAP kinase and gelatinase A, excellent enrichment of active compounds is obtained, while for HIV protease results are moderate and for COX2 (standard scoring) and HIV reverse transcriptase the distribution of inhibitors within the rank lists does not deviate significantly from a random selection. In the following, reasons for the obvious performance differences are discussed in more detail.

For thrombin, an almost quantitative separation between known active inhibitors and the remaining database is achieved. Visual inspection of the structures reveals all top-ranked molecules in the library to contain a benzamidine group binding to Asp 189 at the bottom of the S1 pocket, which contributes significantly to the calculated score. The dominant influence of this charged interaction to achieve significant enrichment of thrombin inhibitors is easily demonstrated by artificially removing contributions of interactions to Asp 189 to the final score. The percentage of top-ranked

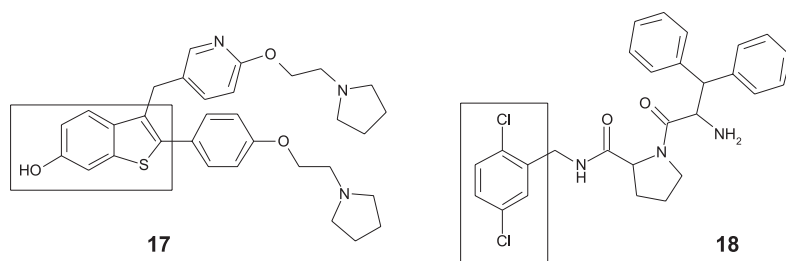


**Fig. 6.8**  
Accumulated percentage of p38 MAP kinase inhibitors plotted versus percentage of the ranked database subset and a bar diagram showing the rank of each inhibitor as a horizontal line.

inhibitors then drops dramatically to 8.3% at 2% of the database (22.2% at 5%, 38.9% at 10%). This observation appears to be generally valid: Wherever specific (for example hydrogen bond or metal contact) interactions are of importance for affinity and whenever these interactions are detected by the docking algorithm, significant enrichment of active compounds can be obtained. It is important to notice, however, that for thrombin charged ligand groups are not necessarily required in the S1pocket. Potent compounds with lipophilic S1 substituents, such as the Ely Lilly compound **17** and the Merck compound **18** shown in Figure 6.9 (S1 moieties inside the box), have been discovered. They are placed on relatively low ranks (between ranks 1200 and 1500) in the present docking study.

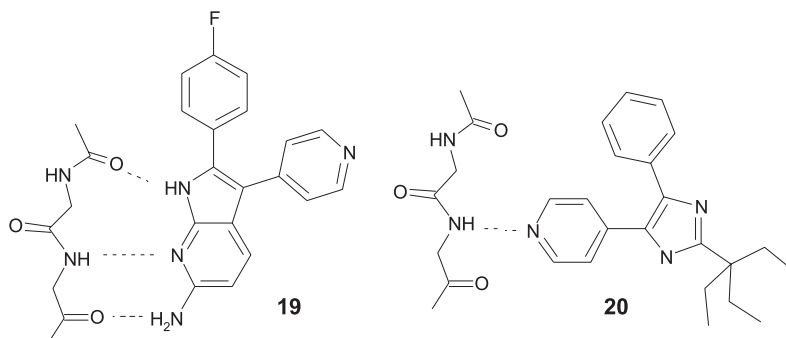
Specific interactions also play an important role in the case of gelatinase A and p38 MAP kinase. The binding site of gelatinase A is valley-shaped with a deep S1' pocket, which can accommodate lipophilic groups as large as biphenyl moieties. The active site is flanked on either side by beta strands allowing for hydrogen bonds to the ligand. These interactions and contacts of inhibitor carboxylate or hydroxamate groups with the catalytic zinc atom result in high ranks once the ligands are correctly placed.

ATP binding sites of kinases are lipophilic clefts formed mainly by ali-

**Fig. 6.9**

Two examples for thrombin inhibitors that do not possess a basic S1 moiety.

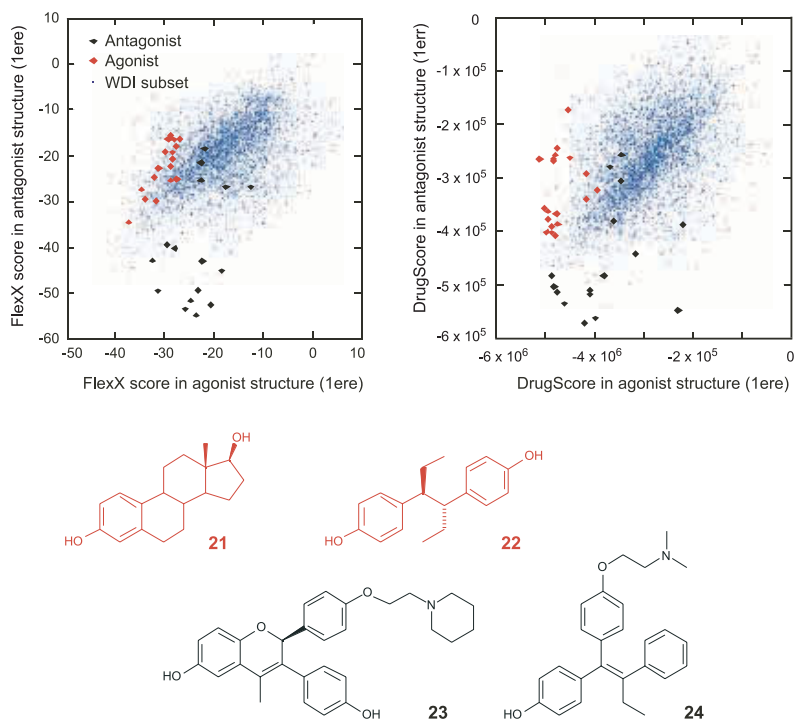
phatic amino-acid side chains. In the adenine-binding region, binding sites are narrow and can accommodate only planar conjugated systems. A special feature of the p38 MAP kinase is an extended, deeply buried lipophilic cavity adjacent to the adenine-binding region. The confined nature of this binding pocket allows FlexX to discard many WDI compounds solely based on shape criteria. One rim of the adenine binding pocket is marked by a beta strand orienting a backbone NH group into the binding site. Kinase inhibitors invariably form a hydrogen bond to this NH group and in addition frequently another bond to one of the two flanking carbonyl groups. Again the dominant contribution of the hydrogen-bond score on the total score is obvious. Inhibitor **19** (Figure 6.10), forming three hydrogen bonds to the protein, is placed on rank 20, although FlexX generates a binding mode reverse to crystallographic evidence. In the FlexX generated mode in which the fluorinated phenyl ring is oriented towards the solvent instead of being located in the buried lipophilic pocket. Conversely, inhibitor **20** is placed on a very low rank (> 3000), since it forms only one hydrogen bond to the receptor.

**Fig. 6.10**

Schematic representation of inhibitor-protein hydrogen bonds in two classes of p38 MAP kinase inhibitors.

The COX2 binding site is a narrow, completely buried lipophilic cavity. At its entrance the side chains of Arg 120 and Glu 524 form a mobile ion pair. Some inhibitors form hydrogen bonds to the guanidinium group of Arg 120, others place lipophilic groups in the same region. Hydrogen bond formation is thus not a predominant feature of inhibitor binding. Accordingly, many WDI compounds with docking solutions involving several spurious hydrogen bonds with the protein are ranked better than the known COX2 inhibitors. When the FlexX solutions for the WDI + COX2 database are re-scored with a modified scoring function in which the weight of lipophilic interactions has been arbitrarily doubled ( $\Delta G_{\text{lipo}} = -0.34 \text{ kJ/mol \AA}$ ) and charged hydrogen bonds receive no extra weight (charge factor = 1), an enrichment increase is observed (“lipo scoring” entry in Table 6.3).

The estrogen receptor binding site is a large lipophilic cavity with acceptor groups at either end that can form hydrogen bonds with ligand hydroxyl groups for example as present in the agonists **21** and **22** or the antagonists **23** and **24** (Figure 6.11). For both agonists and antagonists, lipophilic interactions largely determine the binding energy. The majority of antagonists, however, differ from the agonists in an additional side chain bearing a tertiary amino group. This difference is reflected in the bound structures of the receptor. In the agonist-bound state the binding pocket is not accessible to solvent, whereas in the antagonist-bound state it opens up and allows the positively charged antagonist side chain to form a salt bridge with the carboxylate group of Glu 351. The WDI subset and library of estrogen ligands were docked into both agonist (1ere) and antagonist (1err) forms of the receptor. FlexX scores obtained from both structures are plotted against each other in Figure 6.11. The narrow binding pocket and relatively rigid nature of the ligands restrains possible binding modes significantly. Accordingly, it can be assumed that FlexX is capable of generating reasonable solutions likely to be in agreement with experiment. Therefore we can expect the present example to represent a valuable test for scoring functions. It is observed that antagonists are clearly separated from the WDI compounds, whereas the agonists receive moderately high ranks in the agonist structure and are ranked among the bulk of the WDI entries in the antagonist structure. The reason for this finding is the large contribution of the surface-exposed salt bridge formed with Glu 351 to the total score. In the antagonist form, many WDI molecules also form such a salt bridge, resulting in a better score for many WDI molecules. For this reason – as in the thrombin or p38 MAP kinase examples discussed above – ligands not forming an equivalent salt bridge are not ranked as favorably. Almost the same result as with the FlexX score is obtained with the Chemscore function [103, 104] by Protherics [101]. The new DrugScore function developed by Gohlke et al. [74] performs better in this situation. Using this scoring function, applied as described in Subsection 6.2.2.2, results shown in Figure 6.11 are obtained. The agonists are separated significantly better from the WDI subset when docked into the agonist structure. More importantly, about half of the



**Fig. 6.11**

Plots of docking score in the antagonist form of the estrogen receptor (PDB code 1ere) against docking score in the agonist form of the same receptor (PDB code

1ere) for 20 agonists, 20 antagonists and a WDI subset. Compounds **21** and **22** are typical agonists, compounds **23** and **24** are typical antagonists of the estrogen receptor (for details see text).

agonists are also among the 10% top ranked molecules in the database when they are docked into the open, antagonist structure.

Finally, HIV protease and HIV reverse transcriptase are difficult targets for quite different reasons. Most of the HIV protease inhibitors are relatively large molecules that must be placed accurately into the tunnel-shaped active site to prevent clashing with protein atoms. The FlexX dihedral angle increments do not always permit such an accurate placement. Therefore, several HIV protease inhibitors were docked to regions outside the presumed active site. Furthermore, one must distinguish inhibitors binding to the structural water molecule in the flap region from others replacing it. Out of the 41 inhibitors presently regarded, 12 replace the flap water, 29 do not. Enrichment data given in Table 6.1 were calculated separately for these two groups. In the case of HIV reverse transcriptase, considerable induced fit is observed for most of the inhibitors. This is reflected by the observation that the native ligand corresponding to the crystal structure used receives a rather high rank (11), but all other active compounds are distributed almost



randomly among the ordered database. Furthermore, many inhibitors form water-mediated hydrogen bonds to the protein. Even if the position of essential water molecules can be predicted correctly, the question how to score these contacts has yet to be solved. FlexX has an option to place water molecules during the construction of the ligand [63]. This feature was tested for the structure 1klm, but no improved enrichment was observed.

## 6.6

### Conclusions and outlook

The first test scenario presented in this paper allows for the conclusion that fast similarity measures perform remarkably well when clustering compounds with similar biological properties. Performance increases once the results of searches based on different algorithms emphasizing different aspects of similarity are combined. However, a certain degree of chemical similarity expressed in common skeleton topology must be present for satisfactory performance. Compounds rated as diverse that still bind to a receptor in a similar mode can probably only be identified by means of 3-D superpositioning algorithms including full conformational analyses of each candidate, which is prohibitive for searches in large databases.

The identification of active compounds by means of docking, addressed in the second test scenario, is routinely successful for those targets and inhibitor classes forming a significant number of hydrogen bonds to the receptor. These directed interactions alleviate the generation of correct inhibitor binding modes and result in high ranks for active compounds. In cases where hydrogen bonding plays a minor role, scoring becomes difficult. Supposedly, lipophilic contacts have to obtain higher weight and a more detailed description of lipophilic interactions in the scoring function should be advantageous. As demonstrated for the example of the estrogen receptor ligands, this appears to be reflected better in the recently developed DrugScore function [74]. More detailed investigations will have to show whether DrugScore and other knowledge-based scoring functions [105, 106] are generally superior to traditional empirical ones. A common observation among all screening runs described here is that top ranks are contaminated by structures showing geometrically unreasonable binding modes in the binding pocket. For example, these involve unsatisfied buried donor groups, multiple hydrogen bonds to solvent exposed groups of the protein or unoccupied lipophilic cavities at the receptor-ligand interface. One possible approach to eliminate such unlikely solutions is the development of additional penalty terms and further filters [107]. An interesting approach is the combination of several functions to a consensus scoring function to eliminate unreasonable solutions [108]. This strategy is very similar to the intersection approach described in Section 6.3.3. The placement and scoring of

water-mediated interactions as well as an appropriate treatment of induced-fit phenomena remain on the list of unresolved problems. In addition, the uncertainty about protonation states has to be mentioned. Depending on the original  $pK_a$  value of a functional group in aqueous solution, significant  $pK_a$  shifts can be witnessed upon ligand binding, either of the ligand's functional groups or the protein residues. This finding can be attributed to local changes of the dielectric conditions inside the binding pocket. With respect to docking the change of a donor to an acceptor functionality due to modified protonation states has important consequences. Accordingly, improved docking, and scoring algorithms will have to use a more detailed and flexible description of protonation states.

It has to be kept in mind that the groups of active molecules used in this study mostly comprise high-affinity ligands. However, at the outset of a new drug development project, one can optimistically expect only weakly binding ligands. Furthermore, whatever virtual screening tools are applied, they can only identify active compounds for a new target if they are contained in the libraries that are to be screened. Intelligent assembly of diverse virtual libraries is therefore an important prerequisite of screening.

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## 7

### Future Trends

*Thomas Lengauer*

The previous Chapters of this two-volume series have gone into substantial technical detail of many aspects and sub-problems of bioinformatics, especially as they pertain to pharmaceutical applications. This concluding Chapter addresses the following questions:

1. *How does the progress in genomics and bioinformatics change our view on biology and medicine?*
2. *What are the main coming challenges for bioinformatics?*
3. *What are the well founded expectations and intrinsic limitations of bioinformatics?*

We will address each question in turn and, in line with the purpose of this book, confine our remarks to the medical and pharmaceutical field.

#### 7.1

##### **How does the progress in genomics and bioinformatics change our view on biology and medicine?**

The ability of sequencing has revolutionized the view we have on biological systems. Eric Lander [1] drew the rather fitting comparison with the discovery and consolidation of the periodic table of elements in chemistry, of about 130 years ago. That process reduced the seemingly limitless variety of different forms of matter to a rather small set of then less than 100 elements. This insight, in some sense, completed the vocabulary of chemistry and set the starting point of a remarkable development of the field of chemistry towards a quantifiable discipline in which rational approaches gradually gained ground from the “alchemists” approach of trial and error and in which understanding processes became a significant aspect of discovery.

It seems that we are at a comparable point in biology these days. Granted, the vocabulary is much larger. Genomes have up to billions of base pairs, there are tens of thousands of genes in complex organisms; and the number of different proteins probably goes into the millions, if post-translational modifications are taken into account. Yet, the finiteness of the genome is one of the central insights that come with the new biology. If the genome does not have a certain gene, it simply is not there; and the organism has to and can get by without it and achieve those respective goals by other means. Thus, insights into genes which do not exist in genomes can be a starting point for entirely new lines of biological reasoning.

The other central insight that is afforded by whole genome sequencing is the idea of a genomic map of the organism. For the first time, we can globally lay out the structure and components of a genome. The rough locations of genes, of repeat regions, of gene duplications become available; and they can be studied within a single genome and even in comparison with other genomes. This opens the time dimension of evolutionary development of the species, which is another rich source of essential biological information. Expression data add another dimension of information that can distinguish between cell types and cell states. SNP data allow for distinguishing between individuals of a species.

This global view affords us with two new options. First, we can try networking the detailed local knowledge that we have on a number of biological subsystems. Second, we can try to address biological questions without a preset hypothesis in mind. This approach is quite new to biology, where experiments traditionally are designed to validate or falsify hypotheses. Now, the ability to screen through the complete set of genomic data affords what has recently been called “discovery-based research”, research that does not aim at validating or falsifying an *a priori* hypothesis, but mines the genome-wide data in search of a suitable hypothesis. Discovery-based research not only requires a comprehensive set of data but also highly developed and very strong computer modeling and analysis methods that allow for forming relevant hypotheses based on the data. For reasons that we will discuss later in this Chapter, bioinformatics is not very far yet along the road of providing such analysis methods. As an example, note that protein structure prediction still needs to be equipped with a set of hypotheses – in the form of previously resolved protein structures that are used for templates – in order to make accurate predictions (See chapters 5 and 6 of volume 1). There are other examples that show that providing hypotheses is essential for rendering bioinformatics analyses effective, see e.g. [2].

The gradual change from hypothesis-driven to discovery-based research also weakens the dominance of reductionism that we have observed in biology in the past. The reductionist approach thinks of a biological system as decomposed into subsystems that are studied independently. The finer the decomposition, the smaller the subsystem and the easier it is to form



hypotheses on it and test them experimentally. As we are looking at the genomic data, we begin to see how important the interdependencies, links and communications between the various subsystems really are. The systemic view of the organism gains increasing importance; and this has strong repercussions on the view of bioinformatics. Bioinformatics has to become the basic tool for facilitating this integrated view on the organism. This means that bioinformatics has to offer integrated software and data, an issue that we will elaborate on further in this Chapter.

We have to develop the experimental and bioinformatics methods in order to render discovery-based research effective. This will be an incremental and continuous process that will take a lot of time after the revolutionizing event of providing the human genome sequence. The consequences for biology will be dramatic, and they are already taking effect. Biology is becoming an information handling discipline. Bioinformatics methods have already become the third major pillar of this scientific field, beside laboratory experimentation, and theory. All biologists will need to have basic understanding of this part of biology in order to exercise successful science. This has been recognized to a different extent in different countries. Several countries have started sizeable projects in bioinformatics education. In some of them, the resulting explosive growth of the field is putting the general capabilities of the system of scientific education to the test.

Thus computer-based information handling will be a basic aspect of biology. At the same time, the biological view of an organism will increasingly become that of an information handling system. After all, structure and dynamics is what characterizes biological systems. Organism-wide handling of information – be it gene-regulatory systems, in the immune system or in the nervous system – will be one of the main topics of research in coming years. This is all the more the case, as many diseases are based on malfunctions of these systems.

In summary, the high level attention that has been given to the biological and the informatics part of sequencing the human genome is justified. At the beginning of the postgenomic era we are experiencing a paradigm shift in biology. The field is preparing for it, and part of this preparation is to inject the field with information-handling expertise. This book contributes to this effort.

## 7.2

### **What are the main coming challenges for bioinformatics?**

There are several different alleys for approaching this question. The first is via new kinds of experimental data that are showing up on the horizon and that have to be supplemented with computer methods. The second is by the

kinds of methods that allow for more accurate analysis of the existing data. The third is by finding integrated views on the biological data in order to extract more global insights of the workings of an organism.

### 7.2.1

#### **New experimental data**

In former days, (up to the beginning of the nineties) the bulk of bioinformatics work was directed at analyzing and comparing – mostly aligning – biological sequences, DNA (mostly genes), RNA or protein (see Chapters 2 and 3 of volume 1). Databases like Genbank, the EMBL nucleotide sequence database, and SWISSPROT were the central publicly available data repositories. Work on protein structure snowballed in the early to mid nineties, as the number of significantly different protein structures deposited in the Protein Data Bank grew into the low hundreds [3, 4] and thus was beginning to allow for the derivation of useable potential functions for protein structure prediction (see Chapters 5 and 6 of volume 1). The first whole bacterial genomes appeared in the mid-1990s and immediately – even with the first publication of the genome sequence – attempts at genome-wide annotation with bioinformatics methods were made [5]. In close succession, different genomes were compared with each other, see e.g. [6]. Still today, the annotation of whole genomes and their comparison remains one dominating challenge that drives the development of bioinformatics methods.

The next data to enter the scene were mRNA expression data (see Chapter 5 of volume 2). These data appeared in appreciable volumes in the mid nineties, first as ESTs [7] and then in various forms of micro arrays [8]. It is fair to say that, today, from the application point of view a large part of scientific and industrial bioinformatics is preoccupied with analyzing these data. Transcending simple clustering approaches, the developed methods are now entering their second generation. Yet expression data analysis is still hampered by three issues:

- The measured data are quite noisy. (Technological advances may help that.) Normalization of expression data and comparability between different experiments is a difficult issue. (This problem has to be addressed by both experimental and computer advances.)
- Appropriate methods for interpreting the data have yet to be developed. (Methods are being refined.)
- Since we are measuring on the mRNA level the expression levels contain only indirect information about the concentration of the final protein product. Therefore, links between expression levels and gene function can be expected to be indirect. (This problem may be alleviated partially with time, as we are deepening our understanding about the dependencies

between mRNA expression and protein function. It will probably not go away completely, because the dependencies are expected to be very involved.)

It is hard to predict the future of mRNA expression measurement. What we can be sure of is that proteomics will gain ground from genomics as increasing amounts of data on expression levels of mature proteins become available. By the same token, mRNA expression measurement will probably continue to have its applications, e.g. in diagnostics, especially since genomics experiments are and can be expected to remain cheap and easy compared with proteomics experiments.

We see two major appearing frontiers for new kinds of molecular data. The first is *proteomics* (See Chapter 4 of volume 2) and *metabolomics*. With a combination of 2D gel, mass spectrometry, protein microarray and yeast-two-hybrid methods, a large amount of protein sequence, expression, and interaction data will be produced on a cell-wide level. On the one hand, bioinformatics has to address the challenge of interpreting these data. On the other hand, especially the protein interaction data will provide an interesting basis for probing deeper into the details of regulatory networks. Such data are collected in special protein interaction databases such as DIP [9, 10] and BIND [11].

The first metabolic databases, among them KEGG [12], ENZYME [13], Ecocyc [14], and BRENDA [15] were started in the mid-1990s. Today, they harbor data on well over 100 000 different catalyzed reactions. First, these databases had records, one for each catalyzed metabolic reaction, then a graphic view was added showing the topology of pathways and metabolic cycles. Recently some databases are increasingly being equipped with analysis tools that allow for pathway-oriented navigation and searching.

What is still largely missing is data on enzyme kinetics and metabolite flow. Currently, several groups are starting to gear up for projects that perform cell-wide measurements of metabolites. The corresponding data are supposed to provide the foundation for cell-wide modeling of metabolic networks. The corresponding machinery for simulating metabolic networks is in more than just a nascent state of development. Linear and nonlinear approaches have been introduced [16, 17]. Originally, these methods have only been exercised on comparatively small networks, such as the glycolysis or the pentose-phosphate pathway [18]. Today, metabolic networks are analyzed on an organism-wide scale, at least with respect to their topology. Linear and convex methods of control flow analysis have been applied for this purpose. This includes the analysis of minimal substrate requirements needed to support the production of the essential metabolic products for the organism and the criticality of certain genes for this purpose [19]. The software system METATOOL is a generally available basis for this kind of analysis [20]. Nonlinear analysis for the simulation of enzyme kinetics, as offered by the software system GEPASI [21, 22], is also gearing up for this

kind of scale-up. The lack of kinetic data has curbed the development of this field so far. This obstacle is now on the verge of being removed. This poses an acute demand on the development of mathematical methods, especially, since the field of modeling metabolic networks reaches deeply into the theory of nonlinear systems.

Databases on regulatory relations and pathways are in an earlier stage of development than those on metabolic relations. The first databases on regulatory relations appeared in the late nineties. Today databases such as Transpath [23] and CSNDB [24] contain a few thousand such relations.

The second kind of data that we are starting to be confronted with in large number is data on genetic variations. Stemming from sequencing projects, these data mostly take the form of single-nucleotide polymorphisms (SNPs, see Chapter 3 of volume 2). The wide-spread availability of data on genetic variation will prompt the methodological aspects of bioinformatics and statistical genetics to merge. These two disciplines have not had much interaction, in the past. Bringing them together will be both challenging and very rewarding. It will provide a basis for pharmacogenetics – where we try to analyze and eventually predict the differential effect of a drug on different patients based on the genotypic differences among the patients – and pharmacogenomics – where we aim at selecting the right drug therapy among different alternatives for a single patient, again involving genotype patient data. A first large-scale project underway in this context is the deCODE project that genotypes large parts of the Icelandic population with the aim of understanding the correlation between drug action and effectiveness and genotype as well as obtaining more insights into the genetic basis of complex multi-factorial diseases such as diabetes and coronary syndromes [25]. A necessary ingredient for the latter task is ample genealogical data, which the people of Iceland can supply. Furthermore, a certain genetic homogeneity such as the Icelandic population is hoped to exhibit help in uncovering such correlations. This project has stirred large controversy and raised not only scientific but also political and ethical issues [26].

### 7.2.2

#### **New analysis methods**

The picture of what a bioinformatician does used to be dominated by the BLAST search. Well, maybe, it is more PSI-BLAST, today. In short, a bioinformatician was envisioned doing turn-key analysis of biological data using a very small set of programs, sometimes even a single program. In recent years bioinformatics has become much more than that and, at the same time, not reached the level of expectation yet with which it is confronted by biologists.

The great variety of bioinformatics tools and data is best pointed to in Chapter 1 of volume 2. There are literally dozens of analysis problems for

data from hundreds of molecular biology databases. The whole picture can only be shaped if we exercise these methods on these data in an integrated fashion. Bioinformatics has become a very complex task.

At the same time both the methods and the data are in an early stage of development. Methods make analyses and predictions whose outcomes are not certain to be correct. Databases are more or less cured. In general, they contain a lot of undetected contradictions and errors. Improvement of this situation encompasses method refinement, curation of databases and the integration of heterogeneous data and methods. The former is a mostly scientific topic, the latter two have a sizeable organizational component.

#### 7.2.2.1 Method refinement

Today, analysis and prediction methods have mostly a statistical flavor. They are trained on classified data and make new classifications or predictions based on statistical models. In some sense, all Chapters of Volume 1 present such methods. For instance, the methods for homology-based protein structure prediction in Chapter 5 and 6 of volume 1 learn from a set of observed structures rules that predict alike structures.

In fact, it is reasonable to say that this way of going at things is characteristic for today's method spectrum of bioinformatics and sets it off against other fields in computer-based modeling, which use other approaches. Due to the discrete quality of genomic data and the kinds of statistics involved, the analysis methods are mostly discrete, i.e. relying on models that are based on sets of integers or finite objects such as graphs. There are exceptions to this rule, e.g. in the areas of analyzing expression patterns, simulating metabolic pathways, and detailed modeling of energy. But the respective fields and methods can be viewed as peripheral to bioinformatics as we experience the field today. (This may change in the future.)

If we can clean up the structural database and extend the set of data on known structures, we can increase the accuracy of structure predictions. We can also do so by advancing the methods of statistical learning that underlie the prediction machinery. The progress in both areas will add up to improve the predictions.

We cannot be sure of the answers provided by statistical methods. But if they are developed appropriately, they provide a significance estimate, such as the *P-value* that was the main claim to fame of the BLAST program and its real innovation. This value tells us how much we can believe an analysis or prediction, in the case of BLAST a local sequence match witnessed by an alignment. Recently several other bioinformatics analysis methods have been equipped with theoretically founded or heuristically fitted significance estimates (see Chapters 2 and 6 of Volume 1). Only few bioinformatics tools come with this kind of significance estimate today, and we need more of them.

Establishing high significance of a prediction still does not explain this

prediction. One disadvantage with learning from experience is that it is hard to make such explanations. In order to explain things, the model has to be based on some kind of theory on the structure or process to be predicted. Models that are the basis of learning from experience can have different degrees of diagnosability. E.g. a general feed-forward neural network has no discernable internal structure that reflects biological fact. Thus it is hard to explain why the network makes the predictions that it does. It just happens to be trained that way. On the other hand, a profile hidden Markov model (see Appendix) reflects the internal structure of a biomolecular sequence. The parameters that are trained in the HMM show where secondary structure elements are most likely to be located and where insertions of loops are to be expected. There is some higher degree of diagnosability to hidden Markov models than to neural nets. The force fields that are used in many molecular dynamics simulations are hardly trained at all. Their structure is derived from (some simplification of) the structure of the basic laws of kinetics. The respective parameters are often deduced from quantum-mechanical simulations of small molecules. Mostly, a few parameters remain to be calibrated. Also, there is some heuristic element to the simplification of the structure of the force laws, e.g., by eliminating or grouping multi-body terms. Thus, even *ab initio* calculations have a heuristic element. Yet, it is evident that, as we inject more solidly based theory into the analysis and reduce the purely heuristic or phenomenological aspects of the model, we can hope to better explain what the model is predicting.

Bioinformatics has a long way to go along this road. As we have mentioned above, even the estimation of the significance of a prediction is a rare event in bioinformatics analyses today. Putting more theory into the model is going to be a long and laborious process that relies on the incremental progress that is going to be made in the understanding of biological systems. This process is going to require the advances of the whole field of biology. Put concisely, we are at a very early scientific stage for theorizing about biological systems. Propelling this process is going to be one of the most exciting points of interaction between experimental and computer-based aspects of the field.

#### 7.2.2.2 Curing databases

This is a very difficult issue. Many people are even desperate about it. First, we cannot start a new world from scratch. We have to live with the imperfect databases of today and manage their cleanup as best possible. Simultaneously we have to cope with their dramatic growth. Second, the problem involves an issue of standardization. Nomenclature is inconsistent in basically all parts of the field. The same protein has numerous different names, for instance. Even worse, sometimes different proteins share the same name. Standardization issues involve multinational cooperation and very firm management. The Object Management Group (OMG) has created a

task force to develop a database interoperability standard on the basis of CORBA [www.omg.org/homepages/lsrc/] (see also Chapter 1 of Volume 2). Furthermore, multinational groups are working on the development of ontologies that are both generally accepted and supported by appropriate software [27]. Curing and linking databases is definitely one of the main bottlenecks for the whole field of bioinformatics.

### 7.2.2.3 Integrating heterogeneous data and methods

In some sense, this is at the center of the field of bioinformatics. Adding a new and powerful analysis component will continue to be helpful, but the majority of biological innovation will come in bringing methods and data together towards a common goal of answering a complex biological question. To provide the software platform for this quest is a gigantic effort. Chapter 1 of Volume 2 presents a workable solution for today, but database and method integration have a strong need for further development. The resources necessary for this process will exceed what can be put together in a scientific group. Thus such systems must be expected to come forth from bioinformatics companies. The role of scientific groups is to provide analysis components that fit the bioinformatics toolscape and answer essential biological questions, and to help in providing clean data.

### 7.2.2.4 Data visualization

This is another very important aspect of bioinformatics. We have workable solutions for visualizing the three-dimensional structures of molecular complexes and browsing genome sequences. However, the analysis of gene expression data leads to point clouds in high-dimensional spaces that are not easily visualized. The basic methods of visualization that are based on variants of principle component analysis or hierarchical clustering methods are often missing essential messages that are contained in the data [28]. The problem of appropriate visualization of high-dimensional scientific data is one that transcends bioinformatics, and it is a great challenge to computational science. Another yet unsolved problem is how to appropriately visualize complex metabolic and regulatory networks in a way that reveals interesting insights to the inspecting biologist.

## 7.2.3

### Integrated views on biology

In general, bioinformatics shows a tendency to progress from the solution of subproblems that are more directly related to the genome sequence towards those that are more distant from it and more closely related to phenotype. Here, the notion *phenotype* can be interpreted in different ways.

Again, a line can be followed extending from interpretations that lie closer to the genomic sequence to those that are more distant from it. Different versions of phenotype are three-dimensional molecular structure (e.g. of proteins and RNA) or molecular function (as witnessed by cellular localization, molecular binding partners and participation in certain metabolic and regulatory contexts). One step further the notion of phenotype applies to a whole cell state or even a multi-cellular system, both in structure and dynamics. Figure 1.7 of Chapter 1 in Part 1 can be adapted to visualize this development (see Figure 7.1). The progress in bioinformatics extends counterclockwise around the development cycle shown in Figure 7.1. Dark shaded areas represent comparatively well-developed fields. Light shaded areas represent fields in a comparatively less developed state. As a consequence, future activities in bioinformatics and future attention by the application fields are going to focus on the lighter areas in Figure 7.1.

The integration of heterogeneous data and methods are the basis for moving from genotype to phenotype in scientific analysis. One prime example for this is annotating genome sequences with protein function. There are several ways of deducing knowledge on protein function from genomic data.

- *Homology*: If the protein is homologous to a protein of known function, this homology can be an indication of the function of the protein. Homology can be deduced by sequence search methods, e.g. BLAST. One has to take care in order to separate orthologous proteins (that share the same function) from paralogous proteins (that generally have different function, even though the functions may be related). This distinction is not trivial, and it may be better addressed with databases of proven orthologs than with automatic prediction methods [29].
- *Similar phylogenetic profiles*: If the protein co-occurs with another protein in several organisms and is absent together with the other protein in other organisms, and if the co-occurrence pattern runs in subtrees of the phylogenetic tree of the organisms, then the two proteins are said to have similar (or the same) phylogenetic profiles. Similar phylogenetic profiles are indicative of related function [30], e.g. as binding partners or as components of the same biochemical pathway.
- *Rosetta stone method*: If two proteins occur separated in one organism and homologous proteins are fused in another organism, the former two proteins are candidates for being binding partners in the first organism [31].
- *Co-expression*: If the mRNA expression levels of two proteins are similar, then the proteins could have correlated function (see Chapter 5 of Volume 2 for more detail).
- *Functional motifs*: If a protein exhibits a sequence (or structure) motif that is characteristic for a specific binding site, the binding partner of the protein could be deduced. There are several databases of motifs that are equipped with tailor-made search methods. The oldest motif database is



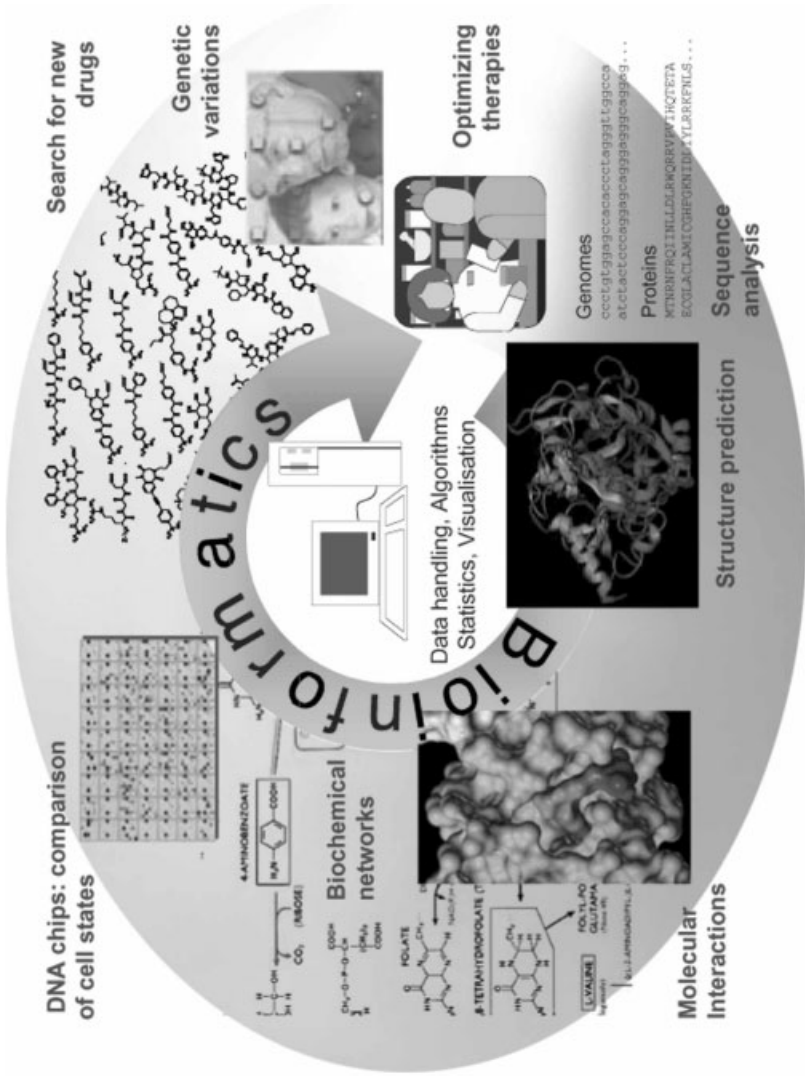


Fig. 7.1 The development of bioinformatics. Darkly shaded areas represent comparatively well-developed fields. Lightly shaded areas represent fields in a comparatively less developed state.

probably PROSITE [32]. PROSITE motifs are good for detecting whether a protein belongs to a protein family and help uncover its domain structure, but they are not always specific enough to predict protein function. A newer and very specific motif search method is the Emotif method [33]. There also is a motif database generated by this method [34]. PRATT [35] is another motif search method that has been extended to find 3D structural motifs with the variant SPRATT [36]. Another approach to defining functionally relevant structural motifs is that of fuzzy functional forms [37].

The power of each method taken by itself is more or less limited. E.g. as we have discussed, mRNA expression patterns reflect protein function only quite indirectly. Sequence homology searches are hampered by the necessity of distinguishing orthologs from paralogs. Exercised in concert, the methods can go much further. An integrated use of the first four methods has been reported in [38]. This is an example for the kind of analysis that is becoming possible with the amount heterogeneous data growing and that is able to address more comprehensive biological questions.

Such questions arise amply in the context of resolving the metabolic and regulatory function of proteins, and the specific role of pathways in disease. Towards this end, we need more information than has been tapped in the protein function studies above. The metabolic databases described in the previous section can be of significant help, here. Methods are being developed that use this kind of knowledge to sharpen the prediction of protein function using mRNA expression data [2, 39].

In the case of regulatory networks the situation is not quite as fortunate. The structure of such networks is much more diverse than that of metabolic networks. Also, as we have mentioned, databases on regulatory networks are just coming into existence, and their data volume is much smaller. Therefore people are trying to extract some of the relevant biological knowledge for the literature. The publicly available resource of the Medline Database [<http://www.ncbi.nlm.nih.gov/PubMed/>] has many millions of abstracts online that can be searched with more or less sophisticated language parsing methods. This approach is to be taken with great caution, because

- Only abstracts are available. (This can be remedied by resorting to electronic versions of complete papers after an initial selection.)
- The quality of the literature is limited. We already alluded to problems originating from inconsistent nomenclature and the lack of generally agreed computer-digestible ontologies. Furthermore, experimental conditions vary with the group that does the experiments, such that results cannot easily be compared. Also, reportedly, the frequency of outright error in the literature cannot be neglected.

- Finally, the computer-assisted parsing methods are far from fail-safe, especially if one resorts to the reduced versions that are able to handle large volumes of data.

Several approaches have been investigated. Marcotte et al. [40] deduce from the frequency of occurrences of certain words in the abstract whether a paper discusses a certain protein-protein interaction. Shatkay et al. [41] analyze the similarity of direction of different documents by providing models for a scientific topic. They train a stochastic model describing the frequency of occurrences of certain words on a given set of documents based on the scientific topic in question. Then they select related literature based on this model. Other systems go beyond counting words. Some systems invoke rules based on the structure of simple sentences [42]. More involved language parsing is performed in [43], [44].

In general the false-positive rate of text-mining analyses can be expected to be very high. Nevertheless, if the output is hand-cured, such analyses can still open new alleys of research and relevant biological insight. One interesting scenario for such methods is to uncover inconsistencies and errors in the literature database and the literature. Bioinformatics methods have already helped greatly to reduce the number of errors in sequence and structure databases. Finally, once appropriate ontologies have been worked out, one can help improve text mining by asking authors of papers to submit not only a list of keywords but rather a formal or semiformal account of the content of their paper that exhibits not only terms but also relations from the content. Ideally, this information should be collected in suitable databases. Parsing this information can both simplify and improve the quality of the mining procedure.

In most of the current discussion – and this is also reflected in Figure 7.1 – bioinformatics stops at the cell level. Of course, this is entirely, artificial. As we are beginning to conquer simulation and modeling of single cells [45], interaction and communication between cells is going to become an issue of increased attention by bioinformaticians. Cell division, differentiation and development, the cellular basis of tissue development, e.g. inside the nervous system, such will be topics that will move closer to the center of attention quite soon.

### 7.3

#### **What are well founded expectations and intrinsic limitations of bioinformatics?**

With the advent of the human genome sequence a certain amount of hype has surrounded the field of bioinformatics. The non-scientific media have

shown a tendency to post genomics and bioinformatics as the holy grail that harbors all solutions to biology and medicine and that will provide the basis for progress towards a longer life in health. These statements have been partly prompted by the large investments of pharmaceutical industry into genomics (in the hundreds of millions of dollars for each of several pharmaceutical companies). The developments of the bio-sector at the stock exchange have done their part to inflate the issue.

Yet, what is evident is that genomics and bioinformatics do present a paradigm shift in biology, and this will undoubtedly propel the field significantly into a new direction that will be characterized by more rational approach to diagnosis and therapy of diseases. In order to discuss in a little more detail what this means let us talk about the character of the field of bioinformatics, pointing out limitations and expectations alike.

Bioinformatics is an instrument for suggestions, no more and no less. For a long time to come, Nature will be too complex to be accurately modeled and simulated by computer. This means that we will have to cope with rough and inaccurate models that lead to predictions that are not fail-safe.

I do not think that there is anything to worry about here. Many examples have shown that even with crude methods one can make relevant predictions (see, e.g. the enrichment of the selection of drug candidates from a compound database by heuristic ranking based on docking or drug comparison, Chapter 6 of Volume 2). As the accuracy of predictions rises the guiding beam of bioinformatics will become more and more focused. Until then, false positives can be weeded out by hand curation or laboratory experimentation, unless their number is overwhelming. It is the false negative rate that we have to worry about more, because if this is high we are missing many solutions. Even here, finding one new solution that gives you the edge over existing approaches can be the basis of a new therapy, so we can start, while the analysis methods are being developed further.

Modeling is getting progressively harder as we transcend from single molecules or molecular interactions to networks of interactions. As data on proteomics and metabolomics mount, we can expect this question to move into the center of attention. Some over-optimistic scientists predicted that the first simulation of a bacterial cell could be working in the year 2005. I think, we will take much longer to have the first executable cell model that stands up to the scrutiny of scientific standards.

The situation in microelectronics, where chips are being designed by computer and the first manufactured silicon version often works, will not be attained in biology for some time to come. It probably is not even desirable to work towards this state of affairs. The lab with *in vitro* and *in vivo* experimentation is always going to test bioinformatics suggestions. The computer is there to limit the amount of experimentation, to guide the scientific mind into promising directions and to interpret the data that are produced in the laboratory in intermediate stages of the research and development process.

After the exciting days of sequencing the human genome the process of increasing biological understanding based on the genomic data will be much more gradual. But there will be progress. After all, after the development of the periodic table in chemistry, progress did not come immediately either. In some sense, the field of chemistry is still in the process of development that was started 130 years ago. By the same token, the field of biology will need time and resources to chart the territory of data that is becoming accessible through genomics and proteomics.

There is a certain component of healing that the molecular approach to medicine will probably never cover. Those are the psychological, sociological and personal aspects. In fact, by increasing our knowledge of the molecular basis of disease we are in danger of neglecting these very important parts of healing. Healing is always going to be an art and require the attention of the doctor to the patient. Notions like trust, ambience, and devoting time to the patient are going to remain central issues in therapy. Methods of healing that are not footed to the molecular viewpoint are experiencing a dramatic resurgence among the population just as we are forming the basis for the molecular approach to therapy. This is a clear message to science and medicine. Healing will only be complete if both the molecular, the systemic and the personal aspect of disease are covered appropriately.

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