Methods in Modern Biophysics

Bengt Nölting

# Methods in Modern Biophysics

Second Edition With 267 Figures



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Alan R. Fersht Robert Huber Manfred Eigen Kurt Wüthrich

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#### Preface

This second edition presents new chapters on (a) the utilization of mutants as highresolution nanosensors of short-living protein structures and protein nanophysics (Chap. 11) and (b) the recently developed method of evolutionary computer programming (Chap. 12), respectively. In the latter method, computer programs evolve themselves towards a higher performance. In contrast to simple selflearning programs, the code of the evolved program differs significantly from that of the original "wild-type" program. In applications on protein folding and structure, evolutionary programming has been shown to yield results many orders of magnitude faster and more efficient than traditional methods. The method is applicable on a wide range of complex problems, e.g., in the fields of nanooptics and adaptive optics (Sects. 12.4, 12.5).

The author gratefully acknowledges Max F. Perutz († 2002) for many inspiring discussions regarding methods for the study of the extreme efficiency of protein folding. These discussions were the strong inspiration for the development of the self-evolving computer programs.

Berlin, June 2005

Bengt Nölting

#### Preface to the first edition

In the recent years we have seen a remarkable increase of the interest in biophysical methods for the investigation of structure-function relationships in proteins, cell organelles, cells, and whole body parts. Biophysics is expected to answer some of the most urgent questions: what are the factors that limit human physical and mental abilities, and how can we expand our abilities. Now a variety of new, faster and structurally higher-resolving methods enable the examination of the mysteries of life at a molecular level. Examples are X-ray crystallographic analysis, scanning probe microscopy, and nanotechnology. Astonishingly large molecular complexes are structurally resolvable with X-ray crystallography. Scanning probe microscopy and nanotechnology allow to probe the mechanical properties of individual biomolecules. Near-field optical microscopy penetrates Abbe's limit of diffraction and enables sub-200 nm resolution. Electron microscopy closes the gap between methods with molecular resolution and cellular resolution. Other methods, such as proteomics, mass spectrometry and ion mobility spectrometry, help us to study highly heterogeneous analytes and to understand extremely complex biological phenomena, such as the function of the human brain. Detailed mechanistic knowledge resulting from the application of these physical and biophysical methods combined with numerous interdisciplinary techniques will further aid the understanding of biological processes and diseases states and will help us to find rational ways for re-designing biological processes without negative side effects. This knowledge will eventually help to close the gap between humans and machines under consideration of all drawbacks, and to find cures for diseases and non-native declines of performance.

This book was mainly written for advanced undergraduate and graduate students, postdocs, researchers, lecturers and professors in biophysics and biochemistry, but also for students and experts in the fields of structural and molecular biology, medical physics, biotechnology, environmental science, and biophysical chemistry. The book is largely based on the lecture "Biophysical Methods" given by the author at the occasion of a visiting professorship at Vienna University of Technology. It presents a selection of methods in biophysics which have tremendously progressed in the last few years.

Chap. 1 introduces fundamentals of protein structures. Proteins have evolved to become highly specific and optimized molecules, and yet the class of proteins may be seen as the biomolecule class with the largest variety of functions. Surely the understanding of biological systems much depends on the understanding of protein structure, structure formation, and function. The next chapter (Chap. 2) presents important chromatographic methods for the preparation of proteins and other biomolecules. Many biophysical studies require this form of sample preparation and often a lot of time can be saved by using optimized procedures of sample purification. Mass spectrometry (Chap. 3) is important for the quality control in preparations of biomolecules, but also has a variety of further analytical applications. Chaps. 4-7 focus on methods for the chemical and structural characterization of biomolecules. X-ray crystallography (Sect. 4.1.2) probably offers the highest resolving power for large biomolecules and biomolecular complexes, but it requires the preparation of high-quality crystals. Cheaper is infrared spectroscopy (Chap. 5) which may also comparably easily be applied in the fast time scale. Electron microscopy (Chap. 6) is particularly suitable for the structural resolution of complex biological systems at the size level of cells, cell organelles, and large molecular complexes. Different types of scanning probe microscopes (Chap. 7) can generate images of geometrical, mechanical, electrical, optical, or thermal properties of biological specimens with up to sub-nm resolution. In Chap. 8 (biophysical nanotechnology) we find novel methods for the mechanical characterization of individual biomolecules and for the engineering of novel nanotechnological structures and devices. The next two chapters (proteomics, Chap 9; and ion mobility spectrometry, Chap. 10) concentrate on two types of analytical methods for the characterization of complex samples such as human cells or bacteria. Finally Chap. 11 deals with some novel developments regarding the interaction of electromagnetic radiation with humans. Kinetics methods in biophysics were not much emphasized throughout the book since many of them can be found in the monograph "Protein Folding Kinetics" (Nölting, 2005). The reader may refer to this monograph for more information on protein structure, transitions state theory in protein science, and on a variety of kinetic methods for the resolution of structural changes of proteins and other biomolecules.

Prof. Dr. Alan R. Fersht supported the development of a variety of modern biophysical methods in our extremely fruitful collaboration at Cambridge University. Prof. Dr. Robert Huber and Prof. Dr. Max F. Perutz initiated highly inspiring discussions regarding modern applications of protein X-ray crystallography. I am particularly indebted to Prof. Dr. Calvin F. Quate, Prof. Dr. Steven G. Sligar, and Prof. Dr. Joseph W. Lyding for an introduction into the AFM technology.

I am indebted to Prof. Dr. Joachim Voigt, Prof. Dr. Martin H. W. Gruebele, Prof. Dr. Kevin W. Plaxco, Dr. Gisbert Berger, and Dr. Min Jiang for proofreading the manuscript, and to Dr. Marion Hertel for processing the manuscript within Springer-Verlag.

Berlin, July 2003

Bengt Nölting

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## Symbols

|       | arrow indicating a process or a          |  |  |
|-------|--|--|--|
|       | coordinate axis                          |  |  |
| →     | arrow pointing to a label or             |  |  |
|       | indicating a distance                    |  |  |
| Å     | angström (10 <sup>-10</sup> m; 0.1 nm)   |  |  |
| AC    | alternating current                      |  |  |
| ADC   | analog-to-digital converter              |  |  |
| AFM   | atomic force microscope                  |  |  |
| ATP   | adenosine triphosphate                   |  |  |
| BESSY | (Berlin Electron Synchrotron             |  |  |
|       | Storage Ring)                            |  |  |
| bp    | base pair                                |  |  |
| BSA   | bovine serum albumin                     |  |  |
| BSE   | bovine spongiform                        |  |  |
|       | encephalopathy                           |  |  |
| °C    | degree Celsius (kelvin-273.15)           |  |  |
| С     | speed of light in vacuum                 |  |  |
|       | $(2.99792 \times 10^8 \text{ m s}^{-1})$ |  |  |
| CBMS  | chemical-biological mass                 |  |  |
|       | spectrometer                             |  |  |
| CCD   | charge coupled device                    |  |  |
| CD    | circular dichroism                       |  |  |
| CJD   | Creutzfeldt-Jacob disease                |  |  |
| cm    | centimeter $(10^{-2} \text{ m})$         |  |  |
| СМ    | carboxy methyl                           |  |  |
| CNS   | central nervous system                   |  |  |
| CO    | carbon monoxide                          |  |  |
| CsI   | cesium iodide                            |  |  |
| CTP   | chain topology parameter                 |  |  |
| Da    | dalton (g mol <sup>-1</sup> )            |  |  |
| DC    | direct current                           |  |  |
| dCTP  | 2'-deoxycytidine 5'-triphosphate         |  |  |
| DSC   | differential scanning calorimetry        |  |  |
| DEAE  | diethyl-amino-ethyl                      |  |  |
| DNA   | deoxyribonucleic acid                    |  |  |
| dsDNA | double-stranded DNA                      |  |  |
| DTGS  | deuterated triglycine sulfate            |  |  |
| е     | elementary charge                        |  |  |
|       | $(1.6022 \times 10^{-19} \text{ C})$     |  |  |

| eV             | electron volt (1.6022 × $10^{-19}$ J)       |  |  |
|----------------|---|--|--|
| FPLC           | fast performance liquid                     |  |  |
|                | chromatography                              |  |  |
| FTIR           | Fourier transform infrared                  |  |  |
| FTMS           | Fourier transform mass                      |  |  |
|                | spectrometer                                |  |  |
| GC             | gas chromatography                          |  |  |
| GPS            | global positioning system                   |  |  |
| h              | Planck constant                             |  |  |
|                | $(6.6261 \times 10^{-34} \text{ J s})$      |  |  |
| HPLC           | high pressure liquid                        |  |  |
|                | chromatography                              |  |  |
| i              | imaginary number $(i \equiv \sqrt{-1})$     |  |  |
| IHF            | integration host factor                     |  |  |
| IMS            | ion mobility spectrometer                   |  |  |
| IMU            | inertial measurement unit                   |  |  |
| IR             | infrared                                    |  |  |
| $k_{\rm B}$    | Boltzmann constant                          |  |  |
|                | $(1.3807 \times 10^{-23} \text{ J K}^{-1})$ |  |  |
| KBr            | potassium bromide                           |  |  |
| kDa            | kilodalton (kg mol <sup>-1</sup> )          |  |  |
| kJ             | kilojoule (1 kJ = 240 cal)                  |  |  |
| kp             | kilopond (9.8066 N)                         |  |  |
| kV             | kilovolt $(10^3 \text{ V})$                 |  |  |
| kW             | kilowatt ( $10^3$ W)                        |  |  |
| 1              | liter $(10^{-3} \text{ m}^3)$               |  |  |
| Laser          | light amplification by stimulated           |  |  |
|                | emission of radiation                       |  |  |
| LD             | linear dichroism                            |  |  |
| LIDAR          | light detection and ranging                 |  |  |
|                | (measurement of light                       |  |  |
|                | backscatter)                                |  |  |
| μm             | micrometer $(10^{-6} \text{ m})$            |  |  |
| MΩ             | megaohm $(10^6 \text{ V A}^{-1})$           |  |  |
| MALDI          | matrix-assisted laser desorption            |  |  |
|                | ionization                                  |  |  |
| MCT            | mercury cadmium telluride                   |  |  |
| m <sub>e</sub> | electron rest mass                          |  |  |
|                | $(9.1094 \times 10^{-31} \text{ kg})$       |  |  |

| milliliter $(10^{-6} \text{ m}^3)$                   | RNAse  | ribonuclease  |
|--|--|---|
| millimolar (6.0221 × $10^{20}$ liter <sup>-1</sup> ) | μs   | microsecond $(10^{-6} \text{ s})$   |
| $6.0221 \times 10^{23}$                              | SAXS   | small angle X-ray scattering  |
| millivolt $(10^{-3} \text{ V})$                      | SDOCT  | spectral domain optical   |
| molecular weight                                     |  | coherence tomography  |
| mass-to-charge ratio                                 | SICM   | scanning ion conductance  |
| nanoampere $(10^{-9} \text{ A})$                     |  | microscope  |
| nanometer $(10^{-9} \text{ m})$                      | SNOM   | scanning near-field optical   |
| nuclear magnetic resonance                           |  | microscope  |
| nanonewton $(10^{-9} \text{ kg m s}^{-2})$           | SPM  | scanning probe microscope   |
| near-field scanning optical                          | ssDNA  | single-stranded DNA   |
| microscope - see SNOM                                | STEM   | scanning transmission electron  |
| optical coherence tomography                         |  | microscope  |
| open reading frame                                   | SThM   | scanning thermal microscope   |
| picoampere $(10^{-12} \text{ A})$                    | STM  | scanning tunneling microscope   |
| polymerase chain reaction                            | TEM  | transmission electron   |
| picogram $(10^{-12} \text{ g})$                      |  | microscope  |
| isoelectric point                                    | TGS  | triglycine sulfate  |
| piconewton $(10^{-12} \text{ kg m s}^{-2})$          | TIR  | total internal reflection   |
| part per billion volume $(10^{-9})$                  | TNT  | trinitrotoluene   |
| polyvinyl chloride                                   | TOF  | time-of-flight mass spectrometer  |
| Py-MS, pyrolysis mass                                | UV   | ultra-violet  |
| spectrometry   | VIS  | visible   |
| root mean square                                     | VUV  | vacuum ultra-violet   |
| root mean square deviation                           |  |   |
|  | milliliter $(10^{-6} \text{ m}^3)$<br>millimolar $(6.0221 \times 10^{20} \text{ liter}^{-1})$<br>$6.0221 \times 10^{23}$<br>millivolt $(10^{-3} \text{ V})$<br>molecular weight<br>mass-to-charge ratio<br>nanoampere $(10^{-9} \text{ A})$<br>nanometer $(10^{-9} \text{ m})$<br>nuclear magnetic resonance<br>nanonewton $(10^{-9} \text{ kg m s}^{-2})$<br>near-field scanning optical<br>microscope – <i>see</i> SNOM<br>optical coherence tomography<br>open reading frame<br>picoampere $(10^{-12} \text{ A})$<br>polymerase chain reaction<br>picogram $(10^{-12} \text{ g})$<br>isoelectric point<br>piconewton $(10^{-12} \text{ kg m s}^{-2})$<br>part per billion volume $(10^{-9})$<br>polyvinyl chloride<br>Py-MS, pyrolysis mass<br>spectrometry<br>root mean square<br>root mean square deviation | milliliter $(10^{-6} \text{ m}^3)$ RNAsemillimolar $(6.0221 \times 10^{20} \text{ liter}^{-1})$ $\mu \text{s}$ $6.0221 \times 10^{23}$ SAXSmillivolt $(10^{-3} \text{ V})$ SDOCTmolecular weightstatemass-to-charge ratioSICMnanoampere $(10^{-9} \text{ A})$ nanometer $(10^{-9} \text{ m})$ nuclear magnetic resonancesNOMnanonewton $(10^{-9} \text{ kg m s}^{-2})$ SPMnear-field scanning opticalssDNAmicroscope – see SNOMSTEMoptical coherence tomographyopen reading framepicoarpere $(10^{-12} \text{ A})$ STMpolymerase chain reactionTEMpicogram $(10^{-12} \text{ gg})$ isoelectric pointroot met on $(10^{-12} \text{ kg m s}^{-2})$ TIRpart per billion volume $(10^{-9})$ TNTpolyvinyl chlorideTOFPy-MS, pyrolysis massUVspectrometryVISroot mean squareVUVroot mean square deviation |

#### 1 The three-dimensional structure of proteins

#### 1.1 Structure of the native state

The human body contains the astonishing number of several 100,000 different proteins. Proteins are "smart" molecules each fulfilling largely specific functions such as highly efficient catalysis of biochemical reactions, muscle contraction, physical stabilization of the body, transport of materials in body fluids, and gene regulation. In order to optimally fulfill these functions, highly specific protein structures have evolved. The performance of humans, animals, and plants crucially depends on the integrity of these structures. Already small structural errors can cause diminishings of performance or even lethal diseases.

Proteins generally consist of thousands of atoms, such as hydrogen (H), carbon (C), nitrogen (N), oxygen (O), and sulfur (S). The van-der-Waals radii are about 1.0-1.4 Å for H, 1.6-2.1 Å for  $-CH_3$ , 1.4-1.8 Å for N, 1.4-1.7 Å for O, and 1.7-2.0 Å for S. Typical sizes of proteins range from a few nm to 200 nm. Since representations with atomic resolution of the whole molecule (Fig. 1.1a), or only its backbone (Fig. 1.1b), would be quite confusing for most proteins, it has become common to represent the protein structure as a ribbon of the backbone (Fig. 1.1c).

Multiple levels of structure are distinguished (see Nölting, 2005): The most basic is the primary structure which is the order of amino acid residues. The 20 common amino acids found in proteins can be classified into 3 groups: nonpolar, polar, and charged. Some physical properties of amino acids are given in Table 1.1. For the hydrophobicity of amino acids see Nölting, 2005. A typical protein contains 50-1000 amino acid residues. An interesting exception is titin, a protein found in skeletal muscle, containing about 27,000 residues in a single chain. The next level, the secondary structure, refers to certain common repeating structures of the backbone of the polypeptide chain. There are three main types of secondary structure: helix, sheet, and turns. That which cannot be classified as one of these three types is usually called "random coil" or "other". Long connections between helices and strands of a sheet are often called "loops". The third level, the tertiary structure, provides the information of the three-dimensional arrangement of elements of secondary structure in a single protein molecule or in a subunit of a protein molecule. The tertiary structure of a protein molecule, or of a subunit of a protein molecule, is the arrangement of all its atoms in space, without regard to its relationship with neighboring molecules or subunits. As this definition implies, a protein molecule can contain multiple subunits. Each subunit consists of only one polypeptide chain and possibly co-factors. Finally, the quaternary structure is the arrangement of subunits in space and the ensemble of its intersubunit contacts, without regard to the internal geometry of the subunits. The subunits in a quaternary structure are usually in noncovalent association. Rare exceptions are disulfide bridges and chemical linkers between subunits.



Fig. 1.1 The three-dimensional structure of the saddle-shaped electron transport protein flavodoxin from *Escherichia coli* (Hoover and Ludwig, 1997). (a) Space-filling representation of the complete molecule. (b) Ball-and-stick representation of the protein backbone. (c) Ribbon representation: ribbons, arrows, and lines symbolize helices, strands, and other, respectively. Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). The figure was generated using MOLSCRIPT (Kraulis, 1991)

Most proteins have only a marginal stability of 20–60 kJ mol<sup>-1</sup> and can undergo conformational transitions (Nölting, 2005). Small reversible conformational changes on a subnanometer scale occur very frequently. Reversible or irreversible molecular movements in the subnanometer or nanometer scale are essential for the function of many proteins. However, occasionally proteins irreversibly misfold into a non-native conformation. This can have dramatic consequences for the organism, especially when misfolded protein accumulates in the cell. A well known example of such a process is the misfolding of the prion protein (Figs. 1.2 and 1.3; Riek et al., 1996, 1998; Hornemann and Glockshuber, 1998). According to the "prion-only" hypothesis (Prusiner, 1999), a modified form of native prion protein can trigger infectious neurodegenerative diseases, such as Creutzfeldt-Jacob disease (CJD) in humans and bovine spongiform encephalopathy (BSE).

| Amino acid    | Molecular mass (Da) <sup>a</sup> | Partial molar volume<br>(cm <sup>3</sup> mol <sup>-1</sup> ) <sup>b,c</sup> | Partial molar volume<br>of residue in protein<br>$(cm^3 mol^{-1})^b$ |
|---------------|----------------------------------|---|--|
| Alanine       | 89.09                            | 60.4  | 54.7   |
| Arginine      | 174.20                           | 126.9   | 121.2  |
| Asparagine    | 132.12                           | 77.3  | 71.6   |
| Aspartic acid | 133.10                           | 74.3  | 68.6   |
| Cysteine      | 121.16                           | 73.5  | 67.7   |
| Glutamic acid | 147.13                           | 89.7  | 84.0   |
| Glutamine     | 146.15                           | 93.9  | 88.2   |
| Glycine       | 75.07                            | 43.2  | 37.5   |
| Histidine     | 155.16                           | 98.8  | 93.1   |
| Isoleucine    | 131.17                           | 105.6   | 99.9   |
| Leucine       | 131.17                           | 107.7   | 101.9  |
| Lysine        | 146.19                           | 111.4   | 105.7  |
| Methionine    | 149.21                           | 105.4   | 99.6   |
| Phenylalanine | 165.19                           | 121.8   | 116.1  |
| Proline       | 115.13                           | 82.2  | 74.8   |
| Serine        | 105.09                           | 60.7  | 55.0   |
| Threonine     | 119.12                           | 76.9  | 71.1   |
| Tryptophan    | 204.23                           | 143.9   | 138.2  |
| Tyrosine      | 181.19                           | 123.7   | 118.0  |
| Valine        | 117.15                           | 90.8  | 85.1   |

Table 1.1 Physical properties of natural amino acids

<sup>a</sup> (Dawson et al., 1969; Richards, 1974; Coligan et al., 1996; Nölting 2005)

<sup>b</sup> At 25 °C in water (Kharakoz, 1989, 1991, 1997)

<sup>c</sup> For the standard zwitterionic state



**Fig. 1.2** Structure of the mouse prion protein fragment PrP(121-231) (Riek et al., 1996). The displayed secondary structure is strand<sub>1</sub> (128–131), helix<sub>1</sub> (144–153), strand<sub>2</sub> (161–164), helix<sub>2</sub> (172–194), helix<sub>3</sub> (200–224), coil (124–127, 132–143, 154–160, 165–171, 195–199). The figure was generated using MOLSCRIPT (Kraulis, 1991)



**Fig. 1.3** A hypothetical mechanism of autocatalytic protein misfolding: with a low rate, the native helical conformation (**a**) spontaneously changes (misfolds) into a  $\beta$ -sheet conformation (**b**); contact of the misfolded protein with further correctly folded protein molecules (**c**) catalyzes further misfolding (**d**, **e**)

In soluble proteins, hydrophilic sidechains (that of aspartic acid, glutamic acid, lysine, arginine, asparagine, glutamine) have a higher preference for a location at the surface. Hydrophobic sidechains (that of alanine, valine, leucine, isoleucine, phenylalanine, tryptophan) are preferentially located inside the so-called hydrophobic core (Fig. 1.4). In contrast, the surface of membrane proteins often contains hydrophobic patches (Fig. 1.5).

Examples of the astonishing diversity of protein tertiary structure are shown in Figs. 1.6–1.8. Many proteins attain complicated multimeric structures. Fig. 6.18 in Chap. 6 shows an example of a complex assembly, the GroEL. For further details on the structures of proteins see Nölting, 2005.



**Fig. 1.4** In soluble proteins, charged and polar sidechains prefer a location at the surface. The sidechains of hydrophobic amino acids do not like to reside in an aqueous environment. That is why these sidechains are preferentially buried within the hydrophobic core



**Fig. 1.5** Typical distribution of hydrophobic and hydrophilic sidechains in membrane proteins. The sidechains of hydrophobic amino acids are preferentially buried within the lipid portion of the membrane. Hydrophilic sidechains prefer contact with the bulk water outside the membrane

*Next page:* **Fig. 1.6** Examples of proteins with mainly helical secondary structure. (a) 1ACP: acyl carrier protein (Kim and Prestegard, 1990); (b) 1HBB: human hemoglobin A (Fermi et al., 1984); (c) 1BCF: iron storage and electron transport bacterioferritin (cytochrome  $b_1$ ) (Frolow et al., 1994); (d) 1MGN: sperm whale myoglobin (Phillips et al., 1990); (e) 1QGT: assembly domain of human hepatitis B viral capsid protein (Wynne et al., 1999); (f) 2ABD: acyl-coenzyme A binding protein (Andersen and Poulsen, 1992); (g) 1FUM: the *Escherichia coli* fumarate reductase respiratory complex comprising the fumarate reductase flavoprotein subunit, the fumarate reductase iron-sulfur protein, the fumarate reductase 15-kDa hydrophobic protein, and the fumarate reductase 13-kDa hydrophobic protein (Iverson et al., 1999). Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). The figure was generated using MOLSCRIPT (Kraulis, 1991).



(f) acyl-coenzyme A binding protein

(g) fumarate reductase respiratory complex



**Fig. 1.7** Examples of proteins with mainly sheet-shaped secondary structure. (**a**) 1CSP: major cold shock protein (CSPB) from *Bacillus subtilis* (Schindelin et al., 1993); (**b**) 2PTL: an immunoglobulin light chain-binding domain of protein L, (Wikström et al., 1995); (**c**) 1NYF: SH3 domain from fyn proto-oncogene tyrosine kinase (Morton et al., 1996); (**d**) 2AIT:  $\alpha$ -amylase inhibitor tendamistat, (Kline et al., 1988); (**e**) 1FNF: fragment of human fibronectin encompassing type-III (Leahy et al., 1992). Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). The figure was generated using MOLSCRIPT (Kraulis, 1991)



(a) HPR protein



(c) domain of streptococcal protein G



(b) domain of procarboxypeptidase B



(e) domain of the U1A protein

(f) signal transduction protein CheY

Fig. 1.8 Examples of proteins with significant amounts of helical and sheet-shaped structure. (a) 1HDN: histidine-containing phosphocarrier protein, (van Nuland et al., 1994); (b) 1PBA: activation domain from porcine procarboxypeptidase B, (Vendrell et al., 1991); (c) 1PGB: B1 immunoglobulin-binding domain of streptococcal protein G (Gallagher et al., 1994); (d) 1UBQ: human erythrocytes ubiquitin, (Vijay-Kumar et al., 1987); (e) 1URN: RNA-binding domain of the U1A spliceosomal protein complexed with an RNA hairpin, (Oubridge et al., 1994); (f) 3CHY: signal transduction protein CheY, (Volz and Matsumura, 1991). Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). The figure was generated using MOLSCRIPT (Kraulis, 1991)

#### 1.2 Protein folding transition states

A considerable number of studies has been devoted to the resolution of folding transition states, see, e.g., Nölting, 2005. The structure of the folding transition



Residue number

**Fig. 1.9a** Inter-residue contact map for the main folding transition state of the monomeric protein src SH3 domain (Nölting and Andert, 2000). The sizes and fillings of the circles indicate the magnitudes of structural consolidation, measured by the so-called  $\Phi$ -value (Nölting, 2005). The diagonal of the plot displays secondary structure contacts, and tertiary structure contacts are contained in the bulk of the diagram. Usually, high  $\Phi$ -values (large full circles) indicate a high degree of consolidation of structure and about native interaction energies, and  $\Phi \approx 0$  (small open circles) are diagnostic of little, if any, formation of stable structure at the individual positions in the inter-residue contact space. Moderate magnitudes of  $\Phi \approx 0.2-0.8$ ) suggest different probabilities of the consolidation of structure. Because of the possibility of the occurrence of non-native interactions in the transition state, only clusters of several contacts (for  $\Phi$  around 0.5 usually at least 5 contacts) may be used to draw statistically significant conclusions about the presence or absence of a significant degree of structural consolidation. The positions of helices and strands of  $\beta$ -sheets in the native state are indicated by bars, H1, H2, ..., and bars, S1, S2, ..., respectively. For further details on transition state structures see also Chap. 11

state is the structure of which formation represents the rate-limiting step in the folding reaction, i.e., the reaction of formation of the native conformation which usually starts with the unfolded polypeptide chain. Knowledge of transition state structures is important to understand the high efficiency of such folding reactions. The structures of many transition states of monomeric and also some dimeric and multimeric proteins provide evidence for a nucleation-condensation mechanism of folding in which structure growth starts with the formation of a diffuse folding nucleus which catalyzes further structure formation (Nölting, 2005; Chap. 11).



**Fig. 1.9b** Inter-residue contact map for the main folding transition state of chymotrypsin inhibitor 2 (CI2) (Nölting and Andert, 2000). The sizes and fillings of the circles indicate the magnitudes of structural consolidation, measured by the so-called  $\Phi$ -value (Nölting, 2005; Chap. 11). For further explanation see the legend for Fig. 1.9a on p. 9

Fig. 1.9 a-d displays the structural consolidation of the transition states of four proteins. In these maps, the magnitudes of  $\Phi$ -values are a measure or probability of structure formation at the corresponding locations in the inter-residue contact space. For example, large filled circles on the diagonal indicate consolidation of

secondary structure contacts, and large filled circles in the bulk of the diagrams indicates consolidated tertiary structure contacts in the transition state (Nölting, 1998). The high structural resolution of the main transition states for the formation of native structure of these four small monomeric proteins (src SH3 domain, chymotrypsin inhibitor 2, barstar, barnase) and of the dimeric Arc repressor (not shown here) reveals that the most consolidated parts of each protein molecule in the transition state cluster together in the tertiary structure, and these clusters contain a significantly higher percentage of residues that belong to regular secondary structure than the rest of the molecule (Nölting and Andert, 2000). For many small monomeric and some dimeric proteins, the astonishing speed of protein folding can be understood as caused by the catalytic effect of the formation of clusters of residues which have particularly high preferences for the early formation of regular secondary structure in the presence of significant amounts of tertiary structure interactions (Nölting and Andert, 2000).



**Fig. 1.9c** Inter-residue contact map for the main folding transition state of barstar (Nölting and Andert, 2000). The sizes and fillings of the circles indicate the magnitudes of structural consolidation, measured by the so-called  $\Phi$ -value (Nölting, 2005; Chap. 11). For further explanation see the legend for Fig. 1.9a on p. 9



**Fig. 1.9d** Inter-residue contact map for the main folding transition state of barnase (Nölting and Andert, 2000). The sizes and fillings of the circles indicate the magnitudes of structural consolidation, measured by the so-called  $\Phi$ -value (Nölting, 2005; Chap. 11). For further explanation see the legend for Fig. 1.9a on p. 9

#### 1.3 Structural determinants of the folding rate constants

For the further understanding of the mechanism and extreme speed of protein folding, and for the rational design of artificial proteins and re-engineering of slowly-folding proteins with aggregating intermediates it is important to resolve, with subnanometer resolution, the question how contacts build up in the reaction (Nölting et al., 1995, 1997a; Nölting, 1998, 1999a, 2005), and how this consolidation of structure relates to the speed of folding (Goto and Aimoto, 1991; Fersht et al., 1992; Dill et al., 1993; Karplus and Weaver, 1994; Orengo et al., 1994; Abkevich et al., 1995; Govindarajan and Goldstein, 1995; Hamada et al., 1995; Itzhaki et al., 1995; Nölting et al., 1995, 1997a; Fersht, 1995a, b; Gross, 1996; Kuwajima et al., 1996; Unger and Moult, 1996; Wolynes et al., 1996; Gruebele,

1999; Forge et al., 2000; Griko, 2000; Niggemann and Steipl, 2000; Nölting and Andert, 2000; Nölting, 2005).



**Fig. 1.10** Example for the formation of intramolecular contacts. Here the contacting residues in the folded conformation with the largest sequence separation are residues number 10 and 30. The set of distance separations in sequence between all contacting residues in space is called chain topology. It is an important determinant of the folding rate constant of the protein (Nölting et al., 2003)

One of the key questions is about the interplay between local and non-local interactions in the folding reaction (Tanaka and Scheraga, 1975, 1977; Gromiha and Selvaraj, 1997, 1999; Goto et al., 1999). In a number of studies it has been shown that the folding rate constants,  $k_{\rm f}$ , of proteins depend on the contact order which is a measure of the complexity of the chain topology of the protein molecule (Fig. 1.10; Doyle et al., 1997; Chan, 1998; Jackson, 1998; Plaxco et al., 1998; Alm and Baker, 1999; Baker and DeGrado, 1999; Muñoz and Eaton, 1999; Riddle et al., 1999; Baker, 2000; Grantcharova et al., 2000; Koga and Takada, 2001). Proteins with a complicated chain topology, i.e., of which the native structure and the structure of the transition state contains many contacts of residues remote in sequence (Figs. 1.11 a, b; 1.12 a, b) have orders of magnitude lower folding rate constants,  $k_{\rm f}$ , than proteins with a simple chain topology, i.e., of which the native structure and the structure of the transition state is dominated by contacts of residues near in sequence (Figs. 1.11 c, d; 1.12 c, d). Within the range of  $10^{-1} \text{ s}^{-1} \le k_f \le 10^8 \text{ s}^{-1}$ ,  $-\log k_f$  correlates well with the so-called chain topology parameter, *CTP*, with a correlation coefficient of up to  $\approx 0.87$ :

$$-\log k_{\rm f} \sim CTP , \qquad CTP = \frac{1}{L \cdot N} \sum \Lambda S_{i,j}^2 , \qquad (1.1)$$

where *L* is the number of residues of the protein (chain length), *N* the number of inter-residue contacts in the protein molecule,  $\Delta S_{i,j}$  the separation in sequence between the contacting residue number *i* and *j*, and "~" marks a linear correlation (Fig. 1.13; Nölting et al., 2003).



**Fig. 1.11a** Chain topologies (Nölting et al., 2003) of three proteins and a peptide with vastly different folding times: (**a**) acylphosphatase (Pastore et al., 1992), (**b**) FK506 binding protein (FKBP-12) (van Duyne et al., 1991), (**c**)  $\lambda$ -repressor dimer bound to DNA (Beamer and Pabo, 1992), and (**d**) the hairpin forming peptide from protein G (41–56) GEWTYDDATKTFTVTE (Achari et al., 1992; Muñoz and Eaton, 1999). Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). *Continued on the following pages* 

The only important difference of the definition of *CTP* to the definition of the contact order is the quadratic dependence on  $\Delta S_{i,j}$ , and yet the fit is more stable and valid over a much larger range of rate constants and valid for both  $\alpha$ -helix proteins and  $\beta$ -sheet proteins. The relation  $-\log k_f \sim CTP$  can also reasonably well predict folding times of peptides. For various cut-off distances from 3.5 Å to 8.5 Å, the correlation coefficient, *R*, for  $-\log k_f \sim CTP$  is 0.80–0.87 (Nölting et al., 2003; Fig. 1.14). Ignoring the inter-residue contacts involving hydrogen atoms which generally have less precisely known or fluctuating positions in the protein molecule causes only little if any effect on *R* (Nölting et al., 2003). When ignoring the data points for the small peptides, the *R* for  $-\log k_f \sim CTP$  is still 0.75–0.81 for this range of cut-off distances.



**Fig. 1.11b** Chain topology (Nölting et al., 2003) of FK506 binding protein (FKBP-12). Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). *For further chain topologies see pp. 14, 16, and 17* 



**Fig. 1.11c** Chain topology (Nölting et al., 2003) of  $\lambda$ -repressor dimer bound to DNA. Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). *For further chain topologies see pp. 14, 15, and 17* 

A further important determinant of the speed of folding is the occurrence of some single strong interactions in the protein molecule. For example, some fast-folding proteins of thermophilic organisms contain a relatively large content of asparagine residues and salt bridges. These interactions can affect the rate of folding by a couple of orders of magnitude.  $-\text{Log } k_{\rm f}$  correlates also with the number of residues belonging to  $\beta$ -sheets. This may be due to the larger number of long-range secondary structure contacts in sheets than in helices.

The  $-\log k_f \sim CTP$  is inconsistent with a zipper-like model for folding where the time of folding would be roughly proportional to the zipper length (sequence separation between zipper beginning and end). Obviously this relation is also inconsistent with a random-search mechanism where  $-\log k_f [s^{-1}] \approx L - 9$ .



**Fig. 1.11d** Chain topology (Nölting et al., 2003) of the hairpin forming peptide from protein G (41–56) GEWTYDDATKTFTVTE. Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). *For further chain topologies see pp. 14–16* 

The protein folding problem, i.e., the understanding of the astonishing speed, complexity and efficiency of folding (Nölting et al., 1995, 1997a; Nölting and Andert, 2000; Nölting, 2005) has gained a large and still increasing importance in the context of folding-related diseases (Bellotti et al., 1998; Ironside, 1998; Brown et al., 1999; Gursky, 1999; Kienzl et al., 1999; Brown et al., 2000; Gursky and Alehkov, 2000), but also in the context of a variety of other exciting questions, such as macromolecular crowding inside the cell (Ellis and Hartl, 1999; van den Berg et al., 2000), high level expression of proteins (Hardesty et al., 1999; Kohno et al., 1999; Kramer et al., 1999), thermostability (Backmann et al., 1998; Williams et al., 1999) and packing problems (Efimov, 1998; Grigoriev et al., 1998; Light), and packing problems (Efimov, 2000b).



(a) acylphosphatase;  $k_{\rm f} = 0.23 \text{ s}^{-1}$ 



(c)  $\lambda$ -repressor;  $k_f = 5,000-100,000 \text{ s}^{-1}$ (bound DNA is also shown)

(b) FKBP-12;  $k_{\rm f} = 4.3 \text{ s}^{-1}$ 



(d) hairpin;  $k_{\rm f} = 200,000 \, {\rm s}^{-1}$ 

**Fig. 1.12** Structures of the three proteins and a peptide with vastly different folding rate constants,  $k_{\rm f}$ : (**a**) acylphosphatase (Pastore et al., 1992), (**b**) FK506 binding protein (FKBP-12) (van Duyne et al., 1991), (**c**)  $\lambda$ -repressor dimer bound to DNA (Beamer and Pabo, 1992), and (**d**) the hairpin forming peptide from protein G (41–56) GEWTYDDATKTFTVTE (Achari et al., 1992; Muñoz and Eaton, 1999). Coordinates are from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). The figure was generated using MOLSCRIPT (Kraulis, 1991)



Fig. 1.13 The measured folding rate constants,  $k_{\rm f}$ , of 20 proteins, a 16-residue  $\beta$ -hairpin and a 10-residue helical polyalanine peptide as a function of the chain topology expressed by the chain topology parameter,  $CTP = L^{-1} N^{-1} \sum \Delta S_{i,j}^2$ , where L is the number of residues of the macromolecule, N the total number of inter-residue contacts in the macromolecule, and  $\Delta S_{i,i}$  the sequence separation between the contacting residues *i* and *j* (Nölting et al., 2003). The fit provides  $\log k_f = 7.56 - 0.895$  CTP with a correlation coefficient of 0.86. Within the range of  $10^{-1}$  s<sup>-1</sup>  $\leq k_f \leq 10^8$  s<sup>-1</sup>, predictions of the folding rate constants of peptides and proteins are accurate to typically a couple of orders of magnitude. The relation between structure and rate of folding is so important because it tells us a lot about the mechanism of protein folding and helps to solve the so-called folding paradox (see Nölting et al., 2003; Nölting, 2005). Inter-residue contacts were calculated at a cut-off distance of 4 Å, and no contacts of hydrogen atoms were included in the calculations. Coordinates of the proteins and the  $\beta$ -hairpin were taken from the Brookhaven National Laboratory Protein Data Bank (Abola et al., 1997). For the choice of coordinates see Nölting et al., 2003. Coordinates of the 10-residue helical polyalanine peptide were calculated with the program FoldIt (Jésior et al., 1994). 18 rate constants from ref. (Jackson, 1998) and the  $k_f$  of the 16-residue  $\beta$ -hairpin were chosen as previously selected in ref. (Muñoz and Eaton, 1999). The  $k_{\rm f}$  of the 10-residue helical polyalanine peptide was estimated using data in (Williams et al., 1996; Gruebele, 1999; Zhou and Karplus, 1999; Nölting, 2005). Embedded in a lipid membrane, similar helices in folded proteins undergo intense vibrations with a frequency of  $10^7 \text{ s}^{-1}$  and several 0.1 Å elongation (e.g., Voigt and Schrötter, 1999). The  $k_{\rm f}$  for the thermostable variant of  $\lambda$ repressor and for the engrailed homeodomain,  $\approx 50,000 \text{ s}^{-1}$ , and 37,000 s<sup>-1</sup> are from (Burton et al., 1996, 1997), and (Mayor et al., 2000), respectively (Nölting et al., 2003)

Studies on protein folding have contributed to the better understanding of hydrophobic interaction (Drablos, 1999; Garcia-Hernandez and Hernandez-Arana, 1999; Chan, 2000; Czaplewski et al., 2000), hydrophilic interaction (Jésior, 2000),



**Fig. 1.14** Correlation coefficient for  $-\log k_f \sim CTP$  for different cut-off distances for the calculation of the contacts, as indicated. No contacts of hydrogen atoms were included in the calculations. Including these contacts leads to a slightly higher correlation coefficient (Nölting et al., 2003)

charge interaction (Åqvist, 1999; de Cock et al., 1999), sidechain association (Galzitskaya et al., 2000), and disulfide formation (Chang et al., 2000a, 2000b).

Speeding up folding was achieved by design of sequences with good folding properties (Irbäck et al., 1999) and facilitating folding with helper molecules, so-called chaperones (Csermely, 1999; El Khattabi et al., 1999; Itoh et al., 1999; Kawata et al., 1999; Yamasaki et al., 1999; Gutsche et al., 2000a, 2000b), and taking carbohydrates as templates for *de novo* design of proteins (Brask and Jensen, 2000).

Protein folding has gained interest also regarding RNA folding energy landscapes (Chen and Dill, 2000), the interpretation of multi-state kinetics (Bai, 1999, 2000; Goldbeck et al., 1999), interpretation of differential scanning calorimetry (DSC) data towards cooperative formation of a folding nucleus (Honda et al., 1999; Honda et al., 2000), the evolution of structure formation (Chan, 1999; D'Alessio, 1999a, 1999b), protein secretion (Chambert and Petit-Glatron, 1999; Berks et al., 2000), and protein structure prediction (Crawford, 1999).

## 1.4 Support of structure determination by protein folding simulations

Theoretically, the structure of the native state of a protein can be determined by calculating the energies of all conformations of the molecule. This is true even if the native conformation does not correspond to the global energy minimum. For example, with a few additional experimentally obtained distance constraints one could decide which is the native structure. Unfortunately, the number of possible

conformations of a polypeptide chain is astronomically large. For example, as judged by the entropy, for a protein comprising 100 residues it is of the order of  $10^{100}$  (Nölting, 2005). There are some more optimistic estimates which are based on mechanistic considerations, but still the number of conformations is astronomically large. A further problem is that there are large positive and negative contributions to the protein stability: The stability of the molecule is given by the difference of two large almost equal numbers (Nölting, 2005). In order to calculate the global energy minimum or a folding pathway with sufficient precision, these two numbers would need to be known with about 3–4 significant digits. Currently the theory of molecular energies is not precise enough to meet this requirement. That is why it has not yet been possible to calculate the global energy minimum of an average-sized protein without significant approximations and profound simplifications. Only recently, groundbreaking molecular dynamics simulations on a 23-residue mini-protein found the energy minimum in 700  $\mu$ s of simulation (Snow et al., 2002).



**Fig. 1.15** Support of structure determination by simulation of protein folding. (a) Step 100 of the simulation: initial collapse to a non-native conformation. (b) Step 400: formation of a molten-globule-like state. (c) Step 4,480 and (d) step 17,990: further condensation and reorganization of the molten-globule intermediate. (e) Step 38,174: formation of a native-like state. Each circle represents an amino acid residue of the protein



Fig. 1.16 Hydrophobic potential used for the folding simulation shown in Fig. 1.15

Due to their extreme simplicity, lattice models for the protein structure and statistical energies have become especially prominent (see, e.g., Shakhnovich et al., 1996; Shakhnovich 1997; Mirny and Shakhnovich, 2001). In these models, often the amino acid residues are represented by spheres and the possible angles of the backbone are significantly restricted, e.g., only  $0^{\circ}$  and  $\pm 90^{\circ}$  are allowed. Surprisingly, these simple approaches often yield reasonable results.

Fig. 1.15 exemplary shows lattice simulations which could fold small proteins into native-like structures. The hydrophobic potential used for these simulations is similar to the potential described by Casari and Sippl (1992), but has a strong repulsion at very short distances (Fig. 1.16). For the attractive component, the same relative factors for pairs of amino acids were used as given by Casari and Sippl (1992) in Table 2. The start conformations are random combinations of the structural elements helix, sheet and random coil. The use of not purely random start conformations, but start conformations that contain fluctuating secondary structure elements speeds up the simulation by several orders of magnitude. The aim was not to calculate a unique native structure, but is to find a set of lowenergy conformations. Experimental constraints are then used to rule out the wrong conformations and to determine the native conformation. Important features of the folding reaction are resembled: the initially expanded conformation collapses to a molten-globule-like state after 400 simulation steps (Fig. 1.15b) which reorganizes after a total of 38,174 simulation steps to a native-like conformation (Fig. 1.15e).

#### 2 Liquid chromatography of biomolecules

Proteins, peptides, DNA, RNA, lipids, and organic cofactors have various characteristics such as electric charge, molecular weight, hydrophobicity, and surface relief. Purification is usually achieved by using methods that separate the biomolecules according to their differences in these physical characteristics, such as ion exchange (Sect. 2.1), gel filtration (Sect. 2.2), and affinity chromatography (Sect. 2.3).

#### 2.1 Ion exchange chromatography

In ion exchange chromatography, the stationary solid phase commonly consists of a resin with covalently attached anions or cations. Solute ions of the opposite charge in the liquid, mobile phase are attracted to the ions by electrostatic forces. Adsorbed sample components are then eluted by application of a salt gradient which will gradually desorb the sample molecules in order of increasing electrostatic interaction with the ions of the column (Figs. 2.1-2.3). Because of its excellent resolving power, ion exchange chromatography is probably the most important type of chromatographic methods in many protein preparations.

The choice of ion exchange resin for the purification of a protein largely depends on the isoelectric point, pI, of the protein. At a pH value above the pI of a protein, it will have a negative net charge and adsorb to an anion exchanger. Below the pI, the protein will adsorb to a cation exchanger. For example, if the pI is 4 then in most cases it is advisable to choose a resin which binds to the protein at a pH > 4. Since at pH > 4 this protein is negatively charged, the resin has to be an anion ion exchanger, e.g., DEAE. One could also use a pH < 4 and a cation exchanger, but many proteins are not stable or aggregate under these conditions. If, in contrast, the protein we want to purify has a pI = 10, it is positively charged at usually suitable conditions for protein ion exchange chromatography, i.e., at a pH around 7. Thus, in general for this protein type we have to choose a cation ion exchange resin, e.g., CM, which is negatively charged at neutral pH.

The capacity of the resin strongly depends on the pH and the pI of the proteins to be separated (Fig. 2.4; Table 2.1), but also on the quality of the resin, the applied pressure, and the number of runs of the column (Fig. 2.5). To improve the life of the resin, it should be stored in a clean condition in the appropriate solvent and not be used outside the specified pH range and pressure limit.

For the separation of some enzymes which may lose their activity by contact with metals in the wall of stainless steel columns, glass-packed columns may be more appropriate. The chromatographic resolution mainly depends on the type of biomolecules, type and quality of the resin, ionic strength gradient during elution, temperature, and the geometry of the column.



Fig. 2.1 Example of ion exchange chromatography. (a)–(c) Loading the column: mobile anions (or cations) are held near cations (or anions) that are covalently attached to the resin (stationary phase). (d)–(f) Elution of the column with a salt gradient: the salt ions weaken the electrostatic interactions between sample ions and ions of the resin; sample molecules with different electrostatic properties are eluted at different salt concentrations, typically between 0-2 M. (g) Interaction of sample molecules with ions attached to the resin: at a suitable pH and low salt concentration, most of the three types of biomolecules to be separated in this example reversibly bind to the ions of the stationary phase


CM: pK<sub>a</sub> = 4.0 cation exchanger → binds to positively charged molecules (cations)





**Fig. 2.2** Two ion exchangers: diethyl-amino-ethyl (DEAE) and carboxy methyl (CM). The positive charge of DEAE attracts negatively charged biomolecules. CM is suitable for purification of positively charged biomolecules



**Fig. 2.3** Example for the salt concentration during adsorption of a sample to an ion exchange column, subsequent elution of the sample, and cleaning of the column. Example of a purification protocol: First the solution of biomolecules and impurities in buffer contained in a syringe is loaded onto the column. The biomolecules and some of the impurities bind to the ions attached to the resin. Loading is completed and non-binding molecules are partly rinsed through the column with some further buffer. The next step is to apply a salt gradient with a programmable pump which mixes buffer with extra salt-containing buffer. The steep salt gradient at the beginning elutes most of the weakly binding impurities. At a certain salt concentration, the biomolecules to be purified elute from the column. Elution is monitored with an absorption detector at 280 nm wavelength and the sample fraction collected. After each run the column is cleaned with 1-2 M KCl. This removes most of the strongly binding sample impurities



**Fig. 2.4** Charge properties of anion and cation exchangers. DEAE has a significant capacity at low and medium pH; CM is highly capacious at high and medium pH

| Functional<br>group | Type of exchanger               | pH range |
|---------------------|---------------------------------|----------|
| -N+-CH <sub>3</sub> | Quaternary amine (strong anion) | 1 – 11   |
| -NH <sub>2</sub>    | Primary amine (weak anion)      | 1 - 8    |
| -NH-                | Secondary amine (weak anion)    | 1 - 7    |
| -N<br>/             | Tertiary amine (weak anion)     | 1 - 6    |
| -COO <sup>-</sup>   | Carboxylic acid (weak cation)   | 6 – 14   |
| -SO <sub>3</sub>    | Sulfonic acid (strong cation)   | 1 – 14   |

Table 2.1 Properties of some important ion exchangers

The experimental set-up (Fig. 2.6) often just consists of a bottle with buffer, a bottle with buffer with salt, a programmable FPLC or HPLC pump, the column, a detector and recorder of absorption at 280 nm, or occasionally at 220 nm, and a sample collector. If the right conditions for protein preparation are unknown, a pre-run is performed with a small fraction of the sample. Attention should be paid not to overload the column in preparative runs since this can shift peak positions and lead to substantial sample losses. In many cases of modern high expression of recombinant proteins, it is possible to obtain a protein with 99% purity with a



Fig. 2.5 Change of the capacity of ion exchange columns due to usage. High performance columns operated at the appropriate pressure and pH can last many 1000 runs



**Fig. 2.6** Typical setup for chromatographic purification of proteins with ion exchange FPLC. The pump mixes the salt gradient for sample elution after the sample was loaded, e.g., with a syringe

single ion exchange chromatographic step. However, in case of comparably low expression levels and substantial sample contamination, ion exchange chromatography alone may not be sufficient. Subsequent gel filtration chromatography (Sect. 2.2) can significantly further improve the protein purity.

## 2.2 Gel filtration chromatography

This type of chromatography is a variant of size exclusion chromatography (molecular exclusion chromatography), and is also known as gel permeation chromatography. It lacks an attractive interaction between the stationary phase



**Fig. 2.7** Gel filtration chromatography. When the sample passes through the porous gel, small sample molecules can enter the pores, causing them to flow slower through the column. Large molecules which cannot enter the pores, pass through the column at a faster rate than the smaller ones. Correct pore sizes and solvents are crucial for a good separation

(gel) and solute. The sample solution passes through the porous gel separating the molecules according to their size. The smallest molecules enter the bead pores, resulting in a relatively long flow path and long retention. Large molecules cannot enter the pores and have to flow around them, resulting in a relatively short flow path (Figs. 2.7-2.10).



**Fig. 2.8** Band broadening in a column with too long a geometry. The so-called effective part of the column is sufficient for separation. Excessively long columns do not improve purity, but just cause dilution of the sample by band broadening



**Fig. 2.9** Band broadening in a column with too large a diameter. Despite the column length is about right to separate the two bands, significant sample dilution and possibly contamination occurs due to inhomogeneous loading of the column

Gel filtration chromatography is also an auxiliary method for assessing the molecular weight of biomolecules (Fig. 2.11). Although there are more precise methods, e.g., mass spectrometry (see Chap. 3), gel filtration chromatography is important for the measurement of monomer-multimer equilibria at about  $\mu$ M-concentrations of biomolecules.



**Fig. 2.10** (a) Chromatogram of the separation of a mixture of myoglobin and insulin with multichannel circular dichroism (CD) detection. The multiplex advantage of the multichannel detection prevents distortion of the shape of the spectra (see Nölting, 2005). (b) CD spectra of myoglobin and insulin for comparison



**Fig. 2.11** Molecules of known molecular weight enable an estimate of the molecular weight of the unknown molecule. In this case, two peaks of the investigated molecule indicate a monomer-dimer equilibrium

## 2.3 Affinity chromatography

Affinity chromatography is a method enabling purification of biomolecules and other macromolecules with respect to individual structure or function. It utilizes the highly specific binding of the macromolecule to a second molecule which is attached to the stationary phase. The principle of operation is as follows: (a) the sample is injected into the column; (b) buffer is rinsed through the column, so that sample molecules with no affinity to the stationary phase are eluted from the column, but sample molecules with a high affinity for the stationary phase are retained in the column; (c) the retained sample molecules are eluted from the column by buffer with a high salt concentration or a different pH or a different solvent composition (Fig. 2.12). The preparation of the protein can be performed by using a number of protein tags. The tags should not cause artificial interactions and should not alter the conformation of the tagged protein. Very common are poly-histidine tags that are attached to the protein by genetic engineering (Fig. 2.13). The tag typically consists of 8-12 histidine residues. It binds to nickel compounds at the surface of the chromatography beads. Fig. 2.14 illustrates a somewhat different variant of affinity chromatography in which misfolded proteins are continuously refolded by chaperones and eluted with buffer.



**Fig. 2.12** Purification of antibodies with affinity chromatography: The antigen is chemically bound to the beads of the column and the mixture of antibodies is rinsed through the column. Antibodies with high binding constants bind to the antigen and are eluted later with a buffer with a high salt concentration



**Fig. 2.13** Attachment of a protein to a bead of an affinity column with a histidine tag. About 10 histidine residues were attached to the protein by genetic engineering, e.g., by polymerase chain reaction (PCR) mutagenesis (see, e.g., Nölting, 2005). The histidine residues strongly bind to the bead made from a nickel chelate resin



**Fig. 2.14** Refolding of expensive, poorly folding proteins: Folding chaperones, also known as chaperonins, are attached to the beads and the unfolded or misfolded protein is rinsed through the column. The chaperone interacts with the sample protein and catalyses its folding into the correct conformation

### 2.4 Counter-current chromatography and ultrafiltration

A relatively old method of chromatography is the Craig counter-current distribution apparatus (Fig. 2.15). Nowadays it serves for the large-scale purification of some chemicals for which other chromatographic methods are too expensive. As in other types of counter-current chromatography, both stationary and mobile phase are liquids and separation is based on sample partition between the two liquids. It may, e.g., function as follows (Fig. 2.15): (a) A certain biochemical has a higher solubility in phase A than impurities of the biochemical, but has a lower solubility in phase B than the impurities. (b) Phase B with a high concentration



**Fig. 2.15** Craig counter-current distribution apparatus: both stationary and mobile phases are liquids. Sample separation is based on its partition between the two liquid phases (see text)



**Fig. 2.16** Ultrafiltration device (supplied, e.g., by Amicon Inc., Beverly, MA). Pressurized nitrogen from a nitrogen flask presses the protein solution against the membrane. Small molecules pass the membrane and are collectable at the outlet. Large molecules stay in the ultrafiltration vessel

of impurities is transferred to the next apparatus and fresh phase B is transferred from the previous apparatus to the shown apparatus. (c) Phases A and B are mixed and separated again, and the process continues with step (a). During suc-



**Fig. 2.17** Side view of a spiral cartridge concentrator (e.g., Millipore Corporation, Bedford, MA). Pressure is applied by centrifuging the concentrator. Similarly to the pervious ultrafiltration device (Fig. 2.16), small molecules pass the membrane and large molecules are retained

cessive cycles, different chemicals move through a chain of counter-current distribution apparatuses with different speeds, and are collected, e.g., at the end of the chain.

Strictly speaking, ultrafiltration (Figs. 2.16 and 2.17) is not a chromatographic method. However, it should be mentioned here since it is an extremely useful tool of sample preparation prior to chromatography and can sometimes even substitute chromatography. It is applicable for (a) protein purification, (b) buffer exchange, and (c) concentrating protein solutions. Purification of a protein with a particular molecular weight,  $M_w$ , requires two steps: (a) First, one runs the ultrafiltration apparatus with a membrane with a cut-off higher than  $M_w$  and collects the solution leaving the vessel. (b) Then, one runs the apparatus with a membrane with a cut-off lower than  $M_w$  and collects the solution remaining in the vessel.

# 3 Mass spectrometry

Mass spectrometry is an incredibly important analytical technique for the identification of molecules by way of measuring their mass-to-charge ratios, m/z, in the ionized state. It is particularly useful for the detection and analysis of traces of macromolecules down to less than 1 pg ( $10^{-12}$  g). The general design of a mass spectrometer comprises sample injector, sample ionizer, mass analyzer and ion detector (Fig. 3.1). First the sample is injected into the ionizer which ionizes sample molecules. Then sample ions are analyzed and detected. To prevent collisions with gas molecules, sample ionizer, mass analyzer and ion detector are generally operated in vacuum.



Fig. 3.1 General design of a mass spectrometer

The ion separation power of mass spectrometers is described by the resolution, R, defined as:

$$R = \frac{m}{\Delta m},$$
 (3.1)

where *m* and  $\Delta m$  are the ion mass and mass difference between two resolvable peaks in the mass spectrum, respectively. *R* typically ranges between 100 and 500,000.

### 3.1 Principles of operation and types of spectrometers

According to their mass analyzer designs, there are five important types of mass spectrometers (MS): (a) magnetic and/or electric sector MS (Figs. 3.2 and 3.3), (b) quadrupole MS (Fig. 3.4), (c) ion trap MS (Fig. 3.5), (d) time-of-flight MS (Figs. 3.6–3.9), and (e) Fourier transform MS (Fig. 3.10). Time-of-flight mass spectrometers (TOFs) often are less expensive than other types of mass spectrometers and have, compared to quadrupole MS and many sector MS, the advantage of recording the masses of all ions injected into the analyzer without scanning, contributing to a high sensitivity. TOFs usually have a smaller mass range and resolving power than Fourier transform mass spectrometers (FTMS).



3.1.1 Sector mass spectrometer

**Fig. 3.2** Single magnetic or electric sector mass spectrometer with a single channel (**a**) and multichannel (**b**) detector, respectively. Ions leaving the ion source are accelerated and passed through the sector in which the electric or magnetic field is applied perpendicular to the direction of the ion movement. The field bends the ion flight path and causes ions with different m/z to travel on different paths. In scanning mass analyzers (**a**) the electric or magnetic field strength is varied and only one mass detected at a time. In non-scanning mass analyzers (**b**) all masses are recorded simultaneously within a limited mass range with the help of a multichannel detector



**Fig. 3.3** Advanced virtual image ion optics with high transmission in a benchtop singlesector mass spectrometer (GCmateII from JEOL Ltd., Tokyo; Matsuda et al., 1974; Matsuda, 1976, 1981)



3.1.2 Quadrupole mass spectrometer

**Fig. 3.4** Quadrupole mass spectrometer. The ion beam is accelerated to a high velocity by an electric field and passed through the quadrupole mass analyzer comprising four metal rods. DC and AC potentials are applied to the quadrupole rods in such a way that only ions with one mass-to-charge ratio (m/z) can pass though the analyzer at a time. To scan different m/z, DC and AC potentials are varied

#### 3.1.3 Ion trap mass spectrometer



Fig. 3.5 Ion trap mass analyzer. With the help of different radio frequency signals applied to the ring electrode and the endcaps, all ions are trapped in the cavity and then sequentially ejected according to their m/z

#### 3.1.4 Time-of-flight mass spectrometer

In time-of-flight mass spectrometers, a uniform starting time of ions is caused, e.g., by a pulse of an ionizing laser (Fig. 3.6) or a voltage pulse to an electric shutter (Fig. 3.7). After passing through the accelerating potential difference, V, the kinetic energy, E, of an ion with the charge, z, mass, m, and velocity, v, is:

$$E = zV = \frac{mv^2}{2} \tag{3.2}$$

For a length,  $l_{\text{TOF}}$ , the time of flight, *t*, is:

$$t = l_{\text{TOF}} \sqrt{\frac{m}{2zV}}$$
(3.3)

For example, for  $l_{\text{TOF}} = 0.1$  m,  $z = e = 1.602 \times 10^{-19}$  C, m = 10 kDa = 10 kg /  $6.0221 \times 10^{23}$ , and V = 100 V we obtain t = 72 µs. A mass resolution of 1 Da requires in this example a time resolution of 3.6 ns. Unfortunately, not all ions start to move at the same time and not all ions have the same velocity. The differ-



**Fig. 3.6** Linear time-of-flight mass spectrometer (TOF) with matrix-assisted laser desorption ionization (MALDI). The linear configuration of TOFs represents the simplest implementation of the time-of-flight technique. The typical mass range lies between 0 and 100 kDa, and the typical mass resolution  $m/\Delta m$  is 300-2000

ences in velocity are called chromatic aberration. Due to the chromatic aberration and the differences in the starting time, the requirements for a very high resolution are hard to meet in the simple design of a linear TOF (Fig. 3.6). In reflectron TOFs (Figs. 3.7-3.9), the ion optics reverses the flight direction of the ions and reduces chromatic aberration.



**Fig. 3.7** A simple reflectron time-of-flight mass spectrometer (e.g., Bryden, 1995). The reflector enhances mass-spectrometric resolution: it increases the time of flight and can focus ions. Here a voltage pulse at the shutter electrode causes a uniform starting time of the ions



**Fig. 3.8** A reflectron time-of-flight mass spectrometer with orthogonal ion inlet (e.g., BioTOF II from Bruker Daltonik, Bremen, Germany)



Fig. 3.9 A high-resolution reflectron time-of-flight mass spectrometer. For further details see, e.g., IonSpec Corp., Irvine, CA; JEOL USA, Inc., Peabody, MA; Micromass and Waters Corporation, Milford, MA; Thermo Finnigan, San Jose, CA; Varian Instruments, Walnut Creek, CA



#### 3.1.5 Fourier transform mass spectrometer

**Fig. 3.10** Principle of operation of a Fourier transform mass spectrometer (FTMS), also called "ion cyclotron resonance mass spectrometer" (see, e.g., IonSpec Corp., Lake Forest, CA; Bruker Daltonik, Bremen, Germany). (a) Ions are injected into the analyzer cell of the spectrometer. The magnetic field forces the thermal ions on orbits with small radii that depend on their mass-to-charge ratio. (b) An applied radio frequency pulse resonantly moves the ions to higher orbits. (c) The radio-frequency signal generated by the cyclotroning of the ions is measured and Fourier-transformed. For the method of Fourier transform see also Sect. 4.1.1. The striking characteristic of FTMS is the high resolution, R, typically in excess of 100,000

#### 3.1.6 Ionization, ion transport and ion detection

Common methods of ionization are electrospray (Fig. 3.11), MALDI (Fig. 3.6), electron bombardment, ion bombardment, and chemical ionization. Ions are mainly guided by electrostatic lenses and quadrupole or octopole ion guides (Fig. 3.12). With the exception of FTMS, the ion signals emerging from the mass analyzer of the MS are commonly detected with an electron multiplier (Fig. 3.13). In FTMS the cyclotroning ions are indirectly detected by measuring and Fourier-transforming the voltage signal they induce into receiver electrodes.



Fig. 3.11 Electrospray ionization method. Analyte solutions delivered by liquid chromatography or a syringe pump are sprayed through the narrow, heated capillary leading into the mass spectrometer. A voltage of typically 200 V - 5 kV is applied between capillary and orifice in front of the electrostatic lenses. Ions form in vacuum by evaporation of the analyte solution of charged droplets



**Fig. 3.12** High frequency octopole ion guide for the injection of ions into an ion trap MS. Compared with a quadrupole ion guide, it enables a higher precision of guidance



**Fig. 3.13** Electron multiplier for a MS. The first dynode converts the ion current into an electron current. Further dynodes amplify the electrons by a total factor of typically  $10^3-10^8$ , largely dependent on the electron accelerating voltage between the dynodes, the number of dynodes, and the dynode composition. The last dynode is connected with an ammeter (not shown)

### 3.1.7 Ion fragmentation

Significant enlargement of the information content of spectra is achieved by fragmenting the sample, e.g., in a collision chamber (Fig. 3.14) or a helium-containing cavity of an ion trap mass analyzer (Fig. 3.5; see also Sect. 3.2).



Fig. 3.14 High resolution sector MS with a collision chamber

#### 3.1.8 Combination with chromatographic methods

For the study of highly complex systems, such as complete cells, MS is often combined with chromatographic methods, such as HPLC (high pressure liquid chromatography; Fig. 3.15), FPLC (fast performance liquid chromatography) and gas chromatography (GC; Figs. 3.16 and 3.17). The two types of connectors between chromatography and MS shown in Figs. 3.15 and 3.18 are both applicable for HPLC and FPLC. Two-dimensional spectra are obtained through



Fig. 3.15 Double sector MS in combination with HPLC

combination of mass spectrometry with chromatographic methods (e.g., Fig. 3.17). The resolution in two dimensions greatly enhances the analyzability of complex mixtures with a large number of components. For example, ion exchange chromatography on a crude cell extract with a resolution of 100 combined with mass spectrometry with a resolution of 10,000 can result in a total resolution of almost 1,000,000 for small and medium-sized soluble cellular proteins for which both methods are often largely independent from each other.

Buffer interference which is occasionally observed in MS can usually be prevented by increasing the sample concentration, decreasing the buffer concentration, or changing the buffer (Fig. 3.19).



Fig. 3.16 GC/MS. The combination of mass spectrometry with gas chromatography can greatly enhance the resolution of complex samples



Fig. 3.17 Example of a two-dimensional GC/MS spectrum



**Fig. 3.18** FPLC/MS connector. In several stages the solvent is removed from the analyte solution by application of dry nitrogen and vacuum. The quadrupole ion guide leads the ions to the mass analyzer of the mass spectrometer



**Fig. 3.19** Mass spectrogram of barstar, the 89-residue inhibitor of the ribonuclease barnase (Nölting et al., 1995, 1997a). *Left:* a number of side-peaks indicate the binding of buffer ions to the highly charged protein. *Right:* a measurement with a lower buffer concentration and higher protein concentration at a pH closer to the pI of the protein yields a cleaner mass spectrogram

## 3.2 Biophysical applications

A considerable interest in the fast point detection of toxic and non-toxic biological materials, such as certain bacterial strains, viruses, and proteins led to the development of portable mass-spectrometric biological detectors (e.g., Figs. 3.20 - 3.27;



**Fig. 3.20** Mass-spectrometric detector of biological agents, comprising a virtual impactor, a pyrolyzer, and a mass spectrometer: The two-stage virtual impactor selects particles of a certain size range, e.g.,  $1-10 \mu m$  (see also Fig. 3.21). These particles are then decomposed by pyrolysis (see Fig. 3.22) and analyzed by an ion trap mass spectrometer (see Figs. 3.24– 3.27). For a similar design see the CBMS (chemical-biological mass spectrometer) from Bruker Daltonik, Bremen, Germany



**Fig. 3.21** Principle of operation of a single-stage virtual impactor. The device splits the total flow of the aerosol into the minor and the major flow. Because the direction of the major air flow is perpendicular to the original direction of air flow, only particles with sizes smaller than the cut-off size can enter the major flow. In contrast, all large particles with a large inertia remain in almost the original direction of flow and join the minor flow containing also some of the small particles (Marple and Chein, 1980; Marple et al., 1998)

Williams et al., 2002). The sensitivity of some of these detectors is better than 1 biological agent particle per liter of air at a detection time of less than 3 minutes.

Further important biophysical applications of MS are the detection of mutations in DNA and DNA sequencing (e.g., Köster, 2001a, 2001b; Hung et al., 2002; Fig. 3.28), detection of mutations and post-translational modifications of recombinant proteins (e.g., Lee et al., 2002a; Sect. 9.1.2 in Nölting, 2005; Fig. 3.29), diagnosis of diseases (e.g., Anderson et al., 2002), protein identification (e.g., Aitken and Learmonth, 2002; and Chap. 9), protein purity control (e.g., Stephenson et al., 2002), peptide sequencing (e.g., Shimonishi et al., 1981; Katakuse et al., 1982; Chen et al., 2002; Nemeth-Cawley and Rouse, 2002; Shevchenko et al., 2002; Stoeva et al., 2002; Figs. 3.30 and 3.31), proteome analysis (e.g., Giffin et al., 2001; Nyman, 2001; Kersten et al., 2002; Lim et al., 2002; and Chap. 9), protein folding investigations (e.g., Favier et al., 2002; Grandori et al., 2002), protein protein interaction comparisons (e.g., Powell et al., 2002; Zal et al., 2002), and the search for extraterrestrial life (e.g., Fig. 3.32; Schwartz et al., 1995).



**Fig. 3.22** Principle of operation of pyrolysis: The pyrolysis tube is loaded sequentially and a radio-frequency current passed through the pyrolysis coil. The current rapidly heats the ferromagnetic foil up to the Curie point where the foil reversibly ceases to exhibit ferromagnetic properties and further heating stops. Suitable ferromagnetic materials are, e.g., iron-nickel alloys. Pyrolysates generated in vacuum are then transferred to the mass spectrometer. The pyrolysis mass spectrometry (PyMS) method (Aries et al., 1986; Berkeley et al., 1990; Goodacre, 1994; Freeman et al., 1995; Goodacre and Kell, 1996) is particularly useful for the detection and analysis of biological agents



**Fig. 3.23** Fragmentation of the pyrolysate by electron bombardment, typically in the energy region around 25 eV, further increases the information content concerning the nature of the biological agent under investigation (see, e.g., Ikarashi et al., 1991)



**Fig. 3.24** Detection of biological contamination and differentiation between different biological agents by comparison of the mass spectra of the pyrolysate with m/z in the range 50-200 (see, e.g., Goodacre, 1994; Goodacre and Kell, 1996)

Especially for the analysis of highly complex biological systems, such as bacterial spores, the combination pyrolysis-MS (PyMS) is extraordinary useful. In this method the sample is partially decomposed in its components prior to mass-spectrometric analysis. The mass spectra of pyrolyzed biological systems may contain more than 100 lines, enabling a very sensitive differentiation of different samples. PyMS is used for the detection of bacteria, bacterial spores and viruses and the differentiation between different species of bacteria and viruses (Freeman et al., 1990, 1997; Snyder et al., 1990, 2001; Sisson et al., 1991; Sultana et al., 1995; Goodfellow et al., 1997; Helyer et al., 1997; Leaves et al., 1997; Magee et al., 1997; Timmins and Goodacre, 1997; Taylor et al., 1998; Barshick et al., 1999;

Goodacre et al., 2000; Tripathi et al., 2001), for the analysis of forensic samples and the personal identification of humans (Ishizawa and Misawa, 1990; Kintz et al., 1995; Armitage et al., 2001; Sato et al., 2001), and for biotechnological applications (Goodacre and Kell, 1996). PyMS spectra may be analyzed by using neuronal networks (Fig. 3.26; Goodacre et al., 1996, 1998a, 1998b; Nilsson et al., 1996; Kenyon et al., 1997). Ion trap mass spectrometers are particularly suitable for the pyrolysis-MS identification of biological agents since they can directly measure multiple fragmentation (Fig. 3.27).



**Fig. 3.25** Detection of biological contamination and differentiation between different biological agents by comparison the mass spectra of the pyrolysate with m/z in the range 50–200 at different degrees of fragmentation. The fragmentation is achieved, e.g., by collision with helium atoms in the cavity of an ion trap MS (see also Fig. 3.27)



**Fig. 3.26** Example of a neuronal network for the analysis of PyMS spectra (Kenyon et al., 1997; Goodacre et al., 1998a, 1998b). In this simplified example the network consists of only 10 input, 5 hidden, 3 output nodes, a bias, and weighted connections. The actual number of nodes in the PyMS input layer is usually equal to 150 (one for each m/z from 51 to 200). The hidden layer may actually contain 8–20 nodes. Preferentially the weights of the connections and the bias are set by supervised learning using the hazardous substances to be detected or simulants of these hazardous substances



Fig. 3.27 Example of mass spectrometry with multiple fragmentation  $(m^3)$  in an ion trap mass spectrometer



**Fig. 3.28** DNA sequencing with the Sanger dideoxynucleotide termination method (Sanger et al., 1977; Sanger, 1988) and mass-spectrometric detection. *Top:* amplification of the template DNA with termination (only one of the four reactions is shown). *Middle:* for comparison to the mass-spectrometric method: conventional analysis of the reaction products with polyacrylamide gel electrophoresis. *Bottom:* analysis of the products of reaction A by mass spectrometry: the pattern of mass peaks shows the positions of adenine (A) in the sequence



**Fig. 3.29** Mass spectrogram of the 10,212-Da protein barstar. This protein preparation contains a fraction with a molecular weight 131 Da higher than expected. This is due to an N-terminal methionine which is not properly cleaved after protein synthesis



**Fig. 3.30** Sequencing of the peptide SDLHQTLKKELALPEYYGENLDALWDCLTG by proteolytic digestion and mass spectrometry. This peptide corresponds to the helix<sub>1</sub>-helix<sub>2</sub> peptide of the protein barstar (Nölting et al., 1997a). In this example with only three proteases, only some parts of the peptide sequence can unambiguously be identified



Fig. 3.31 Peptide sequencing using Edman degradation and mass spectrometry (Shimonishi et al., 1981; Katakuse et al., 1982): reaction products of sequential degradation are mass-spectrometrically identified



**Fig. 3.32** NASA plans to send rovers outfitted with MS to Mars. This sketch illustrates a mobile with mini-TOF searching for extraterrestrial life. Already now mass-spectrometry is utilized to search for extraterrestrial bacteria in meteors

# 4 X-ray structural analysis

## 4.1 Fourier transform and X-ray crystallography

X-ray crystallography is the method with the highest currently available resolution power for structures of large macromolecules and macromolecular complexes. Since the technique of Fourier transform is central to this method, we first deal with some essential aspects of this technique:

#### 4.1.1 Fourier transform

Mathematically the Fourier transform and inverse Fourier transform convert between two domains (spaces), e.g., the domain r (e.g., space or time) and the domain k (e.g., momentum or frequency):

Fourier transform: 
$$F(k) = (2\pi)^{-0.5} \int_{-\infty}^{\infty} f(r) e^{-2\pi i r k} dr$$
 (4.1)

Inverse Fourier transform:

$$f(\mathbf{r}) = (2\pi)^{-0.5} \int_{-\infty}^{\infty} F(\mathbf{k}) e^{2\pi i \mathbf{r} \mathbf{k}} d\mathbf{k}$$
 (4.2)

Fig. 4.1 illustrates a one-dimensional Fourier transform: (b) Represents the decomposition of the signal from (a). This decomposition was calculated from the Fourier transform (c). The inverse transform of (c) yields back exactly (a). (d) Is the inverse Fourier transform of the signal of (c) with all phases set to zero instead of using the correct phases. The comparison of (d) with (a) illustrates the importance of the phases in Fourier transform: in order to be able to correctly obtain back the original signal by inverse Fourier transform, both the amplitudes and the phases have to be known.

Three examples in Figs. 4.2–4.4 demonstrate the method of two-dimensional Fourier transform: Fig. 4.2b represents the Fourier transform of the hexagonal arrangement of peaks of Fig. 4.2a. In Fig. 4.3a some of Fourier components with low amplitudes are set to zero, and yet the inverse Fourier transform (Fig. 4.3b) shows that essentially all information is still preserved. Fourier transforming a noisy object, then substituting certain low-amplitude parts of the Fourier transform by zeros, and then inverse-transforming the modified Fourier transform, is an efficient method for noise reduction. Figs. 4.4a and 4.4b show the result when using only a small slice of the Fourier transform for the calculation of the inverse

Fourier transform: significant distortions are observed along the coordinate for which too few Fourier components were utilized for reconstruction.



**Fig. 4.1** Example for the effect of the loss of phase information on the inverse Fourier transform: With the correct phases, the four frequency components (**b**) add to the sum shown in (**a**). However, when adding the four components with the wrong phase 0, we obtain the wrong sum (**d**). (**c**) Represents the Fourier transform of (**a**). In (**a**), (**b**), (**c**) only a fraction of the function is shown; the complete function is periodical in  $(-\infty, \infty)$ 



(b) |F1(h,k)|

**Fig. 4.2** Example of a two-dimensional Fourier transform: (b) is the Fourier transform of (a). Only the absolute of the function is drawn in (b). However we must keep in mind that the complete function contains an amplitude and phase for each coordinate point (compare with Fig. 4.1)



(b) |f2(x,y)|

**Fig. 4.3** Example of a two-dimensional Fourier transform: In the corners, (**a**) is identical to Fig. 4.2b, but has the components with low amplitude in the middle of the coordinate space set to zero. (**b**) Is the inverse Fourier transform of (**a**). (**b**) Is found to be almost identical to Fig. 4.2a showing that the deleted low-amplitude components did not contain much information. Note that in this figure only the absolutes of the functions are drawn


(b) |f3(x,y)|

**Fig. 4.4** Example of a two-dimensional Fourier transform: (a) Is a thin slice of the function in Fig. 4.2b plus a thick slice with all components set to zero. (b) Represents the inverse Fourier transform of (a): some features of the function in Fig. 4.2a are still preserved in (b), but much information is lost. Note that in this figure only the absolutes of the functions are drawn



Fig. 4.5 Example of a diffraction experiment on a crystal. The X-ray diffraction pattern of the crystal is recorded with an area detector. The pattern consists of a large number of discrete spots

Why is the Fourier transform so important for X-ray crystallography? This is because the diffraction pattern of a crystal (Fig. 4.5) or any other physical object is the Fourier transform of its structure (see also later Fig. 4.10).

To understand why the diffraction pattern of a physical object is its Fourier transform let us consider the diffraction of a wave by a single object (Fig. 4.6) and two point-sized objects separated by  $\mathbf{r}$  (Fig. 4.7): the scattering vector,  $\mathbf{S}$ , is defined as  $\mathbf{S} = \mathbf{s}/\lambda - \mathbf{s}_0/\lambda$ , where  $\mathbf{s}$ ,  $\mathbf{s}_0$ , and  $\lambda$ , are the vector of the incident wave, vector of the diffracted wave, and wavelength, respectively (Fig. 4.6). Then the phase difference in units of wavelengths between the two waves in Fig. 4.7 is given by:  $\mathbf{rs}/\lambda - \mathbf{rs}_0/\lambda = \mathbf{rS}$ . Constructive interference of the two waves occurs in case of  $\mathbf{rS} = 0, \pm 1, \pm 2, ...$ ; destructive interference, i.e., extinction, is observed at  $\mathbf{rS} = \pm 1/2, \pm 3/2, ...$ . The diffraction pattern,  $F(\mathbf{S})$  of the two points is then given by  $F(\mathbf{S}) = e^{-2\pi i \mathbf{rS}}$ , with *i* being the imaginary number defined as  $i \equiv \sqrt{-1}$ .



Fig. 4.6 Diffraction of a wave by a single part of an object



Fig. 4.7 Diffraction of a wave by two objects of equal scattering power

For a the diffraction, F(S), of a macroscopic object consisting of many diffracting points with varying diffraction power,  $\rho(\mathbf{r})$ , we have to integrate all scattered waves:

$$F(\mathbf{S}) = \int_{-\infty}^{\infty} \rho(\mathbf{r}) \, \mathrm{e}^{-2\pi i \mathbf{r} \mathbf{S}} \, \mathrm{d}\mathbf{r}$$
(4.3)

This equation has the form of a Fourier transform (compare with Eq. 4.1). Hence the electron density and structure of a protein can be obtained from the inverse Fourier transform of its diffraction image.



**Fig. 4.8** A lens projecting the image of an object onto a screen performs an inverse Fourier transform of the diffraction pattern of the object

In microscopes the inverse Fourier transform is performed by lenses (Fig. 4.8). Unfortunately, currently there is no X-ray microscope with sufficient resolution and sensitivity. X-ray mirrors do not provide sufficient resolution, and because of radiation damage, we would not obtain a satisfactory resolution for a single protein molecule anyway. That is why we have to record the diffraction pattern of a protein crystal and to calculate the inverse Fourier transform of the diffraction pattern of an object with the help of a camera, all phase information is lost. With other

words, we do not record the complete Fourier transform, but only a fraction of it. The consequences of this serious problem were illustrated in Fig. 4.1. Thus, additionally to the recording of the diffraction pattern, one needs a special technique to recover the phase information. The currently most important method to recover phase information in protein crystallography on new structures is the technique of heavy atom replacement (see Sect. 4.1.2.4).

A specifics of the diffraction of macroscopic crystals is that not a continuous diffraction pattern is obtained, but discrete spots. To understand this behavior, consider the structure of a crystal (Fig. 4.9):



**Fig. 4.9** Mathematically a protein crystal can be described as the convolution of the crystal lattice with the unit cell (one or a few protein molecules)



**Fig. 4.10** The Fourier transform of the crystal lattice is the so-called reciprocal lattice. It determines the maximum number and positions of the observed diffraction spots

The protein crystal can be described as the convolution of the crystal lattice with the unit cell (Fig. 4.9): crystal = lattice \* unit cell. The unit cell is the smallest unit from which the crystal can be generated by translations alone. It

usually contains one or several protein molecules. According to the convolution theorem, the Fourier transform, FT, of two convoluted functions  $f_1(\mathbf{r})$  and  $f_2(\mathbf{r})$  is the product of their Fourier transforms:

FT 
$$(f_1(\mathbf{r}) * f_2(\mathbf{r})) = FT (f_1(\mathbf{r})) \cdot FT (f_2(\mathbf{r}))$$
 (4.4)

Thus, the diffraction pattern of a protein crystal is the Fourier transform of the unit cell times the Fourier transform of the crystal lattice. The latter is called reciprocal lattice (Fig. 4.10). Since the reciprocal lattice is zero outside its lattice points, the crystal diffraction pattern corresponds to the Fourier transform of the unit cell sampled at the points of the reciprocal lattice.

A second way to explain the occurrence of discrete spots in the diffraction pattern of macroscopic crystals, and to evaluate the information from the intensity of theses spots, is to think of the diffraction as a reflection on the X-ray at the lattice planes of the crystal (Fig. 4.11). These lattice planes are described by the Miller indices (Fig. 4.12).



**Fig. 4.11** Reflection of X-rays at the lattice planes of a crystal. Diffraction is viewed as reflection of the X-ray on the lattice planes



Fig. 4.12 Example for the nomenclature of Miller indices, hkl. Miller indices are defined as the smallest integer multiple of the reciprocal axis sections in which 1/0 is set to 0



**Fig. 4.13** Overview of X-ray crystallographic analysis of proteins: From the measured diffraction pattern of suitable native and, if necessary, heavy atom replaced crystals, an initial electron density and atomic model is calculated. The initial model is refined, e.g., by modifying it till its calculated diffraction pattern matches the measured pattern.

# 4.1.2 Protein X-ray crystallography

### 4.1.2.1 Overview

In 1934 Bernal and Crowfoot discovered that pepsin crystals give a well-resolved X-ray diffraction pattern (Bernal and Crowfoot, 1934; Bernal, 1939). It took three decades and the development of computers to obtain the first 3-D structures of proteins (Kendrew et al., 1960; Perutz et al., 1960). Many thousands of native protein structures have been solved since then. Examples are found in Figs. 1.6–1.8. A few structure determinations were even made under artificial conditions, e.g., in organic co-solvents (Schmitke et al., 1997, 1998).

# 4.1.2.2 Production of suitable crystals

For X-ray diffraction we must have a single crystal of suitable geometry and size (Fig. 4.13 on the previous page and Figs. 4.14–4.16). Commercial crystal screening kits, containing the most prominent buffers for protein crystallization, may be obtained, e.g., from JenaBioScience (Jena, Germany). Important parameters for coarse-screening and fine-adjustment are protein concentration, salt types and concentrations, pH, type and concentration of surfactants and other additions, temperature, and speed of crystallization.



**Fig. 4.14** Suitable protein and virus crystals are transparent and do not have inhomogeneities of color or refractive index. Crystals with cracks, intergrown crystals and crystals with cloudy inclusions are generally unsuitable for X-ray crystallography. Totally unsuitable are stacks of plate-like crystals or needle-like fibers and mosaics



**Fig. 4.15** Hanging drop method. The solvent of a small drop of protein or virus solution attached to a cover slide slowly evaporates partially. At the right conditions, a single crystal of suitable size grows



**Fig. 4.16** Crystallization robot for hanging drop crystallization. The computer-controlled mixer draws different solutions from reservoir bottles, mixes them with various ratios, and places the mixtures on a glass plate

One generally starts with a protein concentration of about  $2-50 \text{ mg ml}^{-1}$ . Usually, the protein or virus must not contain a significant amount of contaminants, such as other proteins or viruses, protein or virus fragments, unfolded or misfolded protein, particulate matter, chemical additions unnecessary for stability or solubility. In most cases compact proteins that do not contain floppy ends, such as histidine tags or native unstructured peptides, crystallize better. Suitable crystals have sizes of a few 0.1 mm.

#### 4.1.2.3 Acquisition of the diffraction pattern

For the acquisition of the crystal diffraction pattern (Figs. 4.17–4.19), multi wire area detectors or CCD area detectors (Fig. 4.20) are commonly used. With the example of a linear CCD, Fig. 4.21 illustrates the basic principle of operation of CCDs.

The most common X-ray sources for protein and virus crystallographic analysis are rotating anode generators (Fig. 4.22) with typically 5-25 kW electrical power and synchrotrons (Figs. 4.23 and 4.24). Synchrotrons are comparably expensive, but have a higher brightness enabling shorter measuring times. Reduction of the exposition time often results in a better quality of the diffraction pattern since decomposition of the crystal due to radiation damage is reduced.



**Fig. 4.17** X-ray diffraction pattern of a protein crystal (Norledge et al., 1996). The highlighted section is referred to in Fig. 4.26



Fig. 4.18 General setup for the acquisition of the diffraction pattern



Fig. 4.19 Setup for acquiring the diffraction pattern with an area detector (see, e.g., area detectors from Rigaku, The Woodlands, TX). The crystal is cooled with nitrogen from the cryogenic system nozzle. Cooling the crystal reduces radiation damage, but somewhat changes the intermolecular distances. Diffraction of the X-rays from the X-ray source by the crystal are recorded with the area detector with typically 2048 × 2048 or 4096 × 4096 pixels (see next figure)



**Fig. 4.20** A CCD area detector used for recording of X-ray diffraction patterns. For reduction of the dark current, this CCD is operated at -40 °C, allowing it to detect single photons. The fiber optic taper serves also for blockage of X-rays and thereby prevention of radiation damage to the sensitive CCD array. At a pixel size of 20  $\mu$ m × 20  $\mu$ m, the full well capacity is typically several 100,000 electrons per pixel, enabling the necessary high dynamic range



**Fig. 4.21** Linear charge coupled device (CCD). The sensor elements generate electrons by absorption of photons and store the electrons in potential wells. After a certain period of time, the collected electrons are transferred to the analogue shift register and read out. The symbols are: DD, drain of the output amplifier; DS, source of the output amplifier; DG, gate of the output amplifier; RD, drain of the reset transistor; R, clock gate of the reset transistor (Nölting, 1991)



**Fig. 4.22** Rotating anode generator. An electron beam is focussed onto the rotating anode. It knocks out electrons from the inner electron shells of the anode metal. Reoccupation of the vacant shells by electrons from higher level shells involves the emission of X-ray radiation. The interaction of the electron beam with the anode metal generates also a large amount of heat which is quickly dissipated by rotating the anode below the spot of incidence of electrons



Fig. 4.23 Design of a synchrotron. Ions or electrons are accelerated to a speed close to the speed of light and forced on a curved trajectory. A broad spectrum of radiation is produced along the curved sections of the beam. For protein crystallography, a certain wavelength, e.g., 1 Å, is selected by a monochromator



Fig. 4.24 Emission of the Berlin Electron Synchrotron Storage Ring (BESSY I)



**Fig. 4.25** Diffraction pattern of a poorly scattering crystal: only a few spots near the center are observed, any high resolution information is absent. The dashed circle indicates the area corresponding to a resolution of 2.5 Å. If diffraction spots would be visible up to this circle, the resolution of the obtained structure would be 2.5 Å. One can see that the resolution is much lower in this example

Already superficial inspection of the diffraction pattern provides a lot of information about the quality of the crystals: since the information about fine details of the protein structure is found at large diffraction angles, the absence of spots far outside the center of the diffraction pattern shows that only a low resolution will be obtained (Fig. 4.25).

#### 4.1.2.4 Determination of the phases: heavy atom replacement

As mentioned earlier, after measurement of the diffraction pattern, determination of the phase information is required. If we do not have information from molecules with a similar structure, or anomalously scattering atoms in the molecule, the method of choice may be the heavy atom replacement: the diffraction pattern of the original (native) crystal is compared with crystals that contain a single or a few heavy atoms at fixed positions. Those crystals can be prepared, e.g., by diffusing a solution of a heavy atom salt into the protein crystal.

Fig. 4.26 depicts sections of the diffraction pattern of the native protein crystal and heavy atom derivatized crystal, respectively. The diffraction spots labeled with "++" are significantly increased in intensity for the heavy atom derivative. This shows that they belong to phases with a large magnitude. How can we make this conclusion? See Fig. 4.27 which, in the upper part, shows the intensity of an interferogram of two waves as function of phase: when we introduce a small shift

to one of the waves (lower part), a large increase of intensity of the interferogram is found for large positive phases. Essentially no change of intensity occurs at phases around zero. Thus, analogously we can conclude that diffraction spots which increase in intensity only slightly between native crystal and heavy atom derivative belong to phases around zero or  $\pi$ . So, by comparing the intensities of the spots between native crystal and the heavy atom derivative we can estimate the phases of the individual diffraction spots. With only one heavy atom derivative, an uncertainty of two possibilities remains for each spot, but this can easily be removed with a further, different heavy atom derivative of the protein crystal.



**Fig. 4.26** Section of the diffraction pattern of a protein crystal. *Left:* "native" crystal. *Right:* heavy atom derivative

Another way of showing the importance and meaning of phases in crystallography is illustrated in Fig. 4.28: Atoms with different phases and relative positions may cause a diffraction spot at the same position. Thus, without information from heavy atom replacement, or from diffraction patterns of proteins with similar structure or other information, we cannot deduce the protein structure from the diffraction pattern. Theoretically one could also try out all possible phases and see if it leads to a meaningful structure, but currently for macromolecules the computational effort would be much too high.

It should be noted that the problem of loss of phase information occurs only in the common methods of recording the crystal diffraction, such as with a photographic film or a semiconductor detector. The use of lenses or mirrors to produce an image like in an microscope would prevent this loss of information (see p.65). Unfortunately, currently we cannot build a lens which is sufficiently suitable for focussing X-rays of less than a few Å wavelength: the surface of a conventional lens would not be smooth enough and the bulk of the lens would act like a non-regular grating. Further, it is also very difficult to build highly precise X-ray mirrors (Figs. 4.29 and 4.30). X-ray mirror microscopes using soft radiation currently reach only a few 10 nm resolution. More importantly, the radiation damage would prevent atomic resolution of a single protein molecule or virus.



**Fig. 4.27** Interferogram of two waves: within the phase interval  $[-\pi/2, \pi/2]$ , a small phase shift causes a large negative amplitude change,  $\Delta I$ , for large negative phases (**a**), and a large positive amplitude change for large positive phases (**c**), but almost no amplitude change for zero phases (**b**). Thus, e.g., from a large amplitude increase of a diffraction spot upon application of a small phase shift by an additional heavy atom, we can conclude that the phase of the spot has a large magnitude. Analogously one can estimate the phases from the observation of various intensity changes of diffraction spots upon derivatization of the crystal with a heavy atom. For the complete phase interval,  $(-\pi, \pi]$ , there are still two phases for each amplitude change (not shown). This uncertainty is removed by using data from a second heavy atom derivative



**Fig. 4.28** Different phases and relative positions may cause a diffraction spot at the same position: the atom pairs (1,2) and (2,3) with completely different relative locations cause positive interference at the same position



Fig. 4.29 Common X-ray mirrors are only suitable for low angles of incidence



Fig. 4.30 A pair of X-ray mirrors with grazing incidence focuses an X-ray beam to a spot

The mathematics behind the method of heavy atom replacement is exemplary illustrated in Figs. 4.31–4.34. Fig. 4.31a represents an array of atoms. Fig. 4.31b is the absolute of the Fourier transform of this array. From the imaginary and real parts (Fig. 4.32) of this Fourier transform, the phase was calculated (Fig. 4.33). Fig. 4.34a represents the same array as above, but with one additional heavy atom causing a small change of the Fourier transform. The difference of the absolutes of Fourier transforms between native array and heavy atom derivatized array is shown in Fig. 4.34b. Comparing this difference of the absolutes of Fourier transforms with the absolutes of the phases of the native array (Fig. 4.33b), we find a connection between phases (Fig. 4.33b) and amplitude differences (Fig. 4.34b). This connection allows the magnitude of the phase angles to be determined. As mentioned, the remaining ambiguity of sign is removed by including the data from a second isomorphous heavy atom derivative.



(a) f1(x,y)

FT



(b) |F1(h,k)|

Fig. 4.31 (a) Representation of an array of atoms. (b) Absolute of the Fourier transform of (a)



(a) Im(F1(h,k))



(b) Re(F1(h,k))

Fig. 4.32 (a) Imaginary part of the Fourier transform of the array of Fig. 4.31a. (b) Real part of the Fourier transform of the array of Fig. 4.31a



(b) |Phase(F1(h,k))|

**Fig. 4.33** Phase (**a**) and absolute of the phase (**b**) of the Fourier transform of the array of Fig. 4.31a, calculated from imaginary and real parts of the Fourier transform



(a) f2(x,y)



(b) |F1(h,k)| - |F2(h,k)|

**Fig. 4.34** (a) Representation of the array from Fig. 4.31 with an additional heavy atom. (b) Difference of the absolutes of the Fourier transforms of native array (Fig. 4.31a) and heavy atom replaced array (Fig. 4.34a). When we compare Figs. 4.33b and 4.34b, we can see a connection between phases and amplitude differences. Note that  $|F| = ((Im(F))^2 + (Re(F))^2)^{0.5}$ ; phase(F) = arctan(Im(F)/Re(F)), where "Im" and "Re" stand for imaginary and real parts, respectively

# 4.1.2.5 Calculation of the electron density and refinement

Software for the calculation of the initial electron density from the diffraction data and the refinement of structures is being rapidly developed by several academic institutions and often supplied for free. It may be found on the internet, e.g., by searching with the keywords "protein crystallography software".

#### 4.1.2.6 Cryocrystallography and time-resolved crystallography

Short-living conformational intermediates in the microsecond and nanosecond time scale have been resolved by time-resolved crystallography (Srajer et al., 1996; Genick et al., 1997; Fig. 4.35) and cryocrystallography (Schlichting et al., 2000; Petsko and Ringe, 2000; Wilmot and Pearson, 2002; Fig. 4.36). Time-



Fig. 4.35 Example for time-resolved crystallography. 100 ns after initiation of a conformational change, the electron density indicates the occurrence of two conformations,  $C_1$  and  $C_2$ 



Fig. 4.36 Example for cryocrystallography. The CO is flashed off the heme group of the heme protein. This initiates a conformational transition which is detected, e.g., at -196 °C

resolved crystallography interprets time-dependent electron density maps and can offer detailed structural information on short-lived intermediates under nearphysiological conditions. In cryocrystallography, reactions are induced and measured at a low temperature. At the very low temperatures of flash photolysis and acquisition of the diffraction pattern in the experiment shown in Fig. 4.36, the reaction kinetics of the conformational changes is slowed down by many orders of magnitude. This enables to determine the coordinates of structural intermediates that would normally be too short-lived to be resolved by X-ray crystallography.

# 4.2 X-ray scattering

#### 4.2.1 Small angle X-ray scattering (SAXS)

Small angle X-ray scattering serves for the elucidation of microstructural information in amorphous materials on length scales ranging from a few Å to a few  $\mu$ m (Figs. 4.37 and 4.38). Fig. 4.39 is an illustration of the setup for SAXS. Significant effort is undertaken to enable the measurement at very small angles. Since there is an about reciprocal relationship between distance separation of scattering points ( $\Delta x$ ) and the scattering angle ( $\theta$ ), this measurement is essential to obtain sufficient information in the relatively large length scale compared with the wavelength ( $\lambda$ ) of the X-rays ( $\Delta x \approx 0.5\lambda \sin^{-1}(\theta/2)$ ). For details on X-ray optics see also the previous section.



**Fig. 4.37** Automatic SAXS and wide angle X-ray scattering analysis of biological samples. A large number of vials is automatically sampled and production faults immediately are detected and responded to

SAXS measurements revealed that

(a) an unliganded aspartate transcarbamoylase adopts a T-quaternary structure (Fetler et al., 2002),

(b) the axial period of collagen fibrils is  $65.0 \pm 0.1$  nm in healthy human breast regions, and 0.3 nm larger in cancer-invaded regions (Fernandez et al., 2002),

(c) flax cellulose microfibrils probably have a cross section of  $10 \times 50$  Å<sup>2</sup> (Astley and Donald, 2001),

(d) microfibrils with an axial repeating period of approximately 8 nm are present in the major ampullate silk from the spider *Nephila* (Miller et al., 1999; Riekel and Vollrath, 2001), and

(e) the ATPase domain of SecA has dimensions of approximately 13.5 nm  $\times$  9.0 nm  $\times$  6.5 nm (Dempsey et al., 2002).

SAXS revealed information regarding the conformational diversity and size distribution of unfolded protein molecules (Kamatari et al., 1999; Panick et al., 1999a; Garcia et al., 2001; Choy et al., 2002), and was used in a large number of protein-folding and peptide-folding studies to obtain information about size changes (e.g., Chen et al., 1998; Panick et al., 1998, 1999b; Arai and Hirai, 1999; Segel et al., 1999; Kojima et al., 2000; Russell et al., 2000; Aitio et al., 2001; Canady et al., 2001; Katou et al., 2001; Muroga, 2001; Tcherkasskaya and Uversky, 2001). SAXS is one of the very few methods which can directly monitor structural changes of small virus particles (Sano et al., 1999; Perez et al., 2000).



Fig. 4.38 Diffraction pattern of a cell suspension. SAXS can serve to obtain a "fingerprint" of a biological specimen which helps to identify unknown biological samples





Distance constraints derived from SAXS measurements can be used to filter candidate protein structures for the purpose of protein structure prediction (Zheng and Doniach, 2002). In some cases even low resolution solution structures of proteins were obtained solely from SAXS data (Chacon et al., 1998; Shilton et al., 1998; Bada et al., 2000; Maruyama et al., 2001; Scott et al., 2002) or SAXS data combined with neutron scattering data (Egea et al., 2001). SAXS can reveal the structure of bones (Rinnerthaler et al., 1999) and structural changes in bones due to diseases (Grabner et al., 2001). The method was used to obtain information about conformational changes of bacterial cell wall enzymes upon binding to a substrate (Schönbrunn et al., 2001). SAXS results on human dentin, which is a complex composite of collagen fibers and carbonate-rich apatite mineral phase, are consistent with nucleation and growth of an apatite phase within periodic gaps in the collagen fibers (Kinney et al., 2001).

#### 4.2.2 X-ray backscattering

The property of X-rays to penetrate materials is used in many biophysical applications, ranging from for the purpose of determination of the molecular weight of



**Fig. 4.40** Detection of biological and other organic material behind a metal layer with X-ray backscattering: since the absorption of biological material is much smaller than that of metal, the biological material is difficult to detected in single-wavelength X-ray absorption measurements. X-ray backscattering provides much better contrast in this application. However, the quantum efficiency of X-ray scattering is low and thus relatively large expositions and sensitive cameras must be used

proteins to X-ray backscattering for the purpose of detection of organic material hidden in metal containers (see, e.g., Fig. 4.40).

A problem of detection of organic material, such as illicit drugs and explosives, by X-ray absorption is their low absorption coefficient compared with metals and the possibility to camouflage the material, e.g., by embedding it in other organic material, such as flour or sugar. X-ray backscattering offers a good contrast for the detection of such powdery material (Fig. 4.40). The main disadvantage is the low backscattering coefficient compared with transmission coefficient of most organic samples. Thus, a significantly higher exposure compared with X-ray transmission is usually required.

# 5 Protein infrared spectroscopy

Infrared spectroscopy is based on the infrared absorption of molecules and is, compared with crystallography, a relatively simple and inexpensive tool for the global characterization of molecular conformations and conformational changes of proteins and other biomolecules. Depending on the measurement technique, scanning infrared (IR) spectrometers, Fourier transform infrared (FTIR) spectrometers, and single wavelength infrared apparatuses are distinguished (see Sect. 5.1). Typically the most interesting spectral region for biomolecules is v = 400 - 1004000 cm<sup>-1</sup>, where the wavenumber, v, is defined as  $v \equiv 1$ /wavelength. Infrared activity requires a change of dipole moment upon excitation (Fig. 5.1). For proteins the amide chromophore absorption in the region of 1500 cm<sup>-1</sup>-1700 cm<sup>-1</sup> ( $\approx 6 \,\mu m$ wavelength) is particularly important for the assessment of secondary structure content and structural changes. Regarding the resolution of protein secondary structure, the information content of IR and FTIR spectroscopy is comparable with that of circular dichroism (see, e.g., Nölting et al., 1997b; Nölting, 2005), and regarding the resolution of features of the tertiary structure of proteins, IR and FTIR are often inferior, and yet IR is much easier to apply on a fast time scale and for remote sensing (see, e.g., LIDAR in Sect. 5.1.3).



Fig. 5.1 Example of infra-red active and non-active vibrations. Note that infra-red activity requires a change of dipole moment

# 5.1 Spectrometers and devices

#### 5.1.1 Scanning infrared spectrometers

Early IR spectrometers (Fig. 5.2) were constructed similarly to scanning UV/VIS absorption spectrometers. The emission of the source, e.g., a thermal source operated at 1000 °C, is passed through a monochromator selecting a single wavelength. The monochromatic beam is split into two beams – one having the sample in the path. A shutter passes through only one of the two beams at a time. Both beams are alternatingly detected by an IR detector, e.g., a pyroelectric detector, and compared which each other. The optical density of the sample is calculated from the logarithm of the intensity quotient. The use of light modulation is quite indispensable since the problem of background radiation is much more severe than in UV/VIS spectrometers. Spectra are recorded by scanning the wavelength region of interest. This scanning principle of operation is still widely used in IR spectrometers with time resolutions in the femtosecond to nanosecond region, where infrared lasers serve as IR source (see Nölting, 2005).



**Fig. 5.2** Example of a scanning infrared (IR) spectrometer. The monochromator separates the radiation of the IR source into its different wavelengths and selects one wavelength at a time. A beam splitter separates the monochromatic beam into sample beam and reference beam. The absorption coefficient, according to the chemical and structural properties of the sample molecules, is calculated using the detected intensity quotient between both beams, the pathlength, and the sample concentration

#### 5.1.2 Fourier transform infrared (FTIR) spectrometers

FTIR spectrometers (Figs. 5.3–5.7) use the technique of Michelson interferometry and have the advantage of using a larger part of the emission of the IR source during the measurement of a spectrum, compared with scanning IR spectrometers

that are based on monochromators which select only one wavelength at a time. The better usage of radiation improves the inherent signal-to-noise ratio, especially for strongly absorbing samples for which the measurement may be photon shot noise limited. Also the spectral resolution of FTIR spectrometers, which is limited by the path length of the moving mirror, is often better than that of scanning IR spectrometers.

In FTIR spectrometers (Fig. 5.3) the beam of radiation from the IR source is focused on a beam splitter constructed such that half the beam is transmitted to a moving mirror and the other half is reflected to a fixed mirror. Both the moving mirror and the fixed mirror reflect the beam back to the beam splitter which reflects the half of both beams to the detector where they interfere according to their phase difference. The light intensity variation with optical path difference, called interferogram, is the Fourier transform of the incident light spectrum (light intensity as a function of the wavenumber). Absorption spectra are obtained by measuring interferograms with a sample and with an empty sample cell in the beam and inverse Fourier transforming the interferograms into spectra (Figs. 5.4-5.6).



**Fig. 5.3** Typical design of FTIR spectrometers. The lamp, e.g., a thermal source, emits a beam of infrared radiation. A Michelson interferometer, consisting of a beamsplitter, a fixed mirror and a moving mirror, splits the beam into two beams and generates an interference of them. The sample inserted in one of the beam paths changes the interference. Interferograms with and without sample are recorded and the absorption of the sample is calculated by inverse Fourier transform (see Fig. 5.6)

Fig. 5.4 shows three examples of interference of the two monochromatic light beams of the interferometer resulting in different intensities of the interferogram. Eq. 5.1 describes the intensity of the interferogram,  $I_{interferogram}$ , for the interference of two polychromatic beams of equal intensity in the FTIR spectrometer:

$$I_{\text{interferogram}}(\delta) = const \times \int_{0}^{\infty} I_{\text{beam}}(v) \cos(2\pi\delta v) \, dv \,, \tag{5.1}$$

where  $\delta$  is the phase difference of the two beams, *const* a constant,  $I_{\text{beam}}$  the intensity of the beams, and v the wavenumber. From the interferogram, the intensity of the beams can be calculated by inverse Fourier transform:

$$I_{\text{beam}}(v) = const \times \int_{-\pi}^{+\pi} I_{\text{interferogram}}(\delta) \cos(2\pi\delta v) d\delta$$
(5.2)

Analogously, the intensity of the beam with the sample in the path is calculated from the corresponding interferogram. The absorption is given by the logarithm of the intensity quotient of blank to sample.



**Fig. 5.4** Interference of two monochromatic light waves with equal intensity. *Top:* both beams have the same phase; their interference yields the maximum of the interferogram, i.e., the sum of both intensities. *Middle:* at a phase difference of  $\lambda/4$ , the intensity of the interferogram equals the intensity of the interfering beams. *Bottom:* at a phase difference of  $\lambda/2$ , both beams extinguish each other



Fig. 5.5 Example of an interferogram of two polychromatic light beams



**Fig. 5.6** Principle of operation of a FTIR spectrometer. IR intensities at the detector are recorded both for the sample cell filled with solvent and for the sample cell filled with sample. Inverse Fourier transform of the two interferograms yields the IR intensities. The IR absorption spectrum is calculated using the logarithm of the intensity quotient



Fig. 5.7 Sample cell for FTIR experiments. The transparent walls of the cell are made from silicon wafers supplied by a manufacturer of electronic chips

A very suitable material for the manufacture of sample cells, sample holders, and windows is silicon (Fig. 5.7). Polished silicon wafers of 0.5 - 1 mm thickness are sufficiently transparent from 400 to 4000 cm<sup>-1</sup> (25 – 2.5 µm wavelength) (Jiang et al., 1996). Only the fragility and the high refractive index of this material might be problematical in some experimental set-ups. Used infrared sources are often thermal sources operated at about 1000 °C. Beam splitters made from a thin germanium film evaporated on a potassium bromide (KBr) or cesium iodide (CsI) slide are transparent down to about 400 cm<sup>-1</sup> (25 µm wavelength) and 200 cm<sup>-1</sup> (50 µm wavelength), respectively. Liquid nitrogen cooled mercury cadmium telluride (MCT) detectors and deuterated triglycine sulfate (DTGS) pyroelectric detectors are frequently applied for infrared detection. For an excellent introduction into the instrumentation of FTIR spectroscopy see Perkins, 1986.

#### 5.1.3 LIDAR, optical coherence tomography, attenuated total reflection and IR microscopes

IR spectroscopy is exquisitely suitable for remote sensing of clouds of biological agents (Fig. 5.8). The IR LIDAR set-up consists of a pulsed IR laser and an IR detector which senses the backscattered light from the laser. Since the light travels extremely fast, the detector senses the return echo before the next pulse is sent. The time it takes for the laser pulse to travel down and back is a measure of the distance. Mobile commercial LIDAR systems quite often employ an integrated global positioning system (GPS) to determine the own position.

Equipped, e.g., with an optical modulator which rapidly changes the direction of the beam, and mounted on top of a roof, the IR LIDAR can scan the 360°-environment at distances of 0 to several 10 km. This method has importance, for example, for early warning systems of smog in large cities and for three-dimensional analysis of forest structure and terrain (Fig. 5.9). Remote sensing of changes in forest structure utilizes the information of time and intensity of multiple reflections from leaves and branches. Effects of environmental pollutants and pests are quickly detectable in vast areas and economic damage is largely reducible.



**Fig. 5.8** Remote sensing of environmental changes, e.g., a cloud of biological material, with an IR LIDAR (light detection and ranging; measurement of light backscatter)



**Fig. 5.9** Remote survey of forest structure and terrain with IR LIDAR technology. The plane is equipped with a GPS and an inertial measurement unit (IMU). The latter contains several gyroscopes and an accelerometer and can determine the position and angle of tilt with some accuracy during periods of failure of the GPS



of the probe beam reflected from the sample is interfered with light of the reference beam, and the interference is detected by the Fig. 5.10 Optical coherence tomography (OCT). The IR light from a light emitting diode is split into reference and probe beams. Light photodiode. The pathlength of the reference beam is modulated by stretching an optical fiber with a piezoelectric transducer. Light from the sample which has traveled the same distance as the reference beam interferes constructively. Its signal is extracted from the interference intensity by a lock-in amplifier (Duncan et al., 1998)



**Fig. 5.11** Spectral domain optical coherence tomography (SDOCT) (Andretzky et al., 1998; Häusler and Lindner, 1998). Polychromatic backscattered light from different depths interferes with polychromatic light of a reference beam. The interference of the beams is analyzed with a polychromator and a multichannel detector. From the spectral changes due to interference, information about the depth of the scattering layer is obtained

Another important variant of IR spectroscopy on biological samples is optical coherence tomography (OCT). OCT (Figs. 5.10 and 5.11) utilizes echoes of infrared light waves backscattered off the internal microstructures within biological objects to obtain images on a  $\mu$ m scale. In the design of Fig. 5.10, IR radiation backscattered from the sample is interfered with a reference beam. Light from a scattering layer in the sample with a certain depth has the same phase as the reference beam and thus interferes constructively, i.e., produces a high interference intensity. Light from slightly deeper or shallower scattering layers cause a lower interference intensity. By modulating the phase of the reference beam and detecting the interference intensity with a lock-in amplifier, the signals from layers with different depths are extracted from the interference intensity (Duncan et al., 1998). Fig. 5.11 depicts a second design variant of optical coherence tomography (Andretzky et al., 1998; Häusler and Lindner, 1998). Here the information on depth is gained by analyzing the spectrum of the backscattered light.


Petersen, 2002). The internally total-reflected light slightly leaves the waveguide and so can probe the sample molecules on the outside of the waveguide. The part of the light wave which leaves the waveguide at the total reflection points is called evanescent wave. Only very little Fig. 5.12 Flow cell for attenuated total reflection (ATR) infrared spectroscopy (Fringeli et al., 1998; Feughelman et al., 2002; Snabe and sample is needed. Using a large number of reflections can lead to a more than 100-fold amplification of the measured signal

The next IR spectroscopic technique to be mentioned is attenuated total reflection (ATR) infrared spectroscopy (see, e.g., Fringeli et al., 1998; Ding et al., 2002; Feughelman et al., 2002; Snabe and Petersen, 2002; Figs. 5.12 and 5.13). Here the coefficient of internal total reflection of an IR beam in a waveguide is changed by a sample deposited on the surface of the waveguide. An advantage of ATR on thin layered samples is the dramatic increase of the effective optical pathlength and sensitivity through multiple reflections compared with conventional transmission spectroscopy on such a sample.



**Fig. 5.13** Attenuated total reflection (ATR) infrared spectroscopy on membrane proteins (see, e.g., Ding et al., 2002)



**Fig. 5.14** Scanning IR microscope. The focussed beam from an IR laser is passed through the sample and detected. *Left:* simple microscope with planar resolution, especially suitable for thin layer samples. *Right:* microscope with three-dimensional resolution: for acquisition of the image, the sample is moved in xyz-directions

Because of the significantly lower scattering of IR light relative to light of shorter wavelength, IR microscopes (Fig. 5.14) enable the inspection of most strongly scattering samples. Computer aided image processing allows two- or three-dimensional resolution. More complicated microscopes may utilize step-scan interferometry for photoacoustic depth profiling, monochromators for spectral analysis and polarizers/analyzers for linear dichroism (LD) analysis.

### 5.2 Applications

One of the biophysical main applications of FTIR is the characterization of the structure and conformational changes of proteins (Barrera et al., 2002; Butler et



**Fig. 5.15** Isotope-edited FTIR spectroscopy (see, e.g., Li et al., 2002; Barth, 2002). Since the spectrum of the isotope-labeled part of the protein molecule is significantly shifted, it can be distinguished from the spectrum of the non-labeled part. A change of the protein IR spectrum upon binding of the substrate to the protein shows which part of the molecule the substrate binds to. In this example, the magnitude of a peak in the spectrum of the isotope-labeled part of the protein has changed upon binding of the substrate. This shows that the substrate binds to the labeled part of the enzyme



**Fig. 5.16** Apparatus to monitor protein unfolding under high pressure with IR. Since the volume of unfolded protein is less than that of folded protein, high pressure favors transition to the unfolded state



Fig. 5.17 Protein molecule in organic solvent: only a few strongly bound water molecules remain attached to the protein molecule

al., 2002; Castellanos et al., 2002; Dong et al., 2002; Hilario et al., 2002; Moritz et al., 2002; Mui et al., 2002; Noinville et al., 2002), of peptides (Bianco et al., 2002; Gordon et al., 2002; Huang et al., 2002; Torres et al., 2002), and of DNA (Lindqvist and Graslund, 2001; Malins et al., 2002). In some cases, interactions were resolved at the level of individual amino acid residues (Kandori et al., 2002; Mezzetti et al., 2002; Zhang et al., 2002a).

Isotope-edited FTIR is particularly useful for the structural characterization of specific macromolecular regions (Fig. 5.15): e.g., the three phosphate stretching

vibrations of the phosphate calcium ATPase complex were detected at a background of 50,000 protein vibrations in an isotope exchange experiment (Barth, 2002).

Time-resolved step-scan FTIR spectroscopy enables the monitoring of conformational changes of proteins in the microsecond time scale (Bailey et al., 2002).

FTIR spectroscopy allowed to map out the nucleotide binding site of calcium ATPase (Liu and Barth, 2002). IR and FTIR spectroscopy are two of the only few methods suitable to monitor conformational changes of proteins under high pressure (Fig. 5.16; Dzwolak et al., 2002). FTIR spectroscopy on bacterio-rhodopsin revealed a pre-melting conformational transition at 80 °C (Heyes et al., 2002). FTIR is also suitable to investigate the structure and hydration shell of protein molecules in organic solvents (Fig. 5.17; Costantino et al., 1995). Further, IR and FTIR spectroscopy was used for the characterization of irradiated starches (Kizil et al., 2002), and the determination of dihedral angles of tripeptides (Schweitzer-Stenner, 2002). Molecular changes of preclinical scrapie can be detected by IR spectroscopy (Kneipp et al., 2002). FTIR spectroscopy can serve as an optical nose for predicting odor sensation (van Kempen et al., 2002) and for chemical analysis of drinks (Coimbra et al., 2002; Duarte et al., 2002).

FTIR microscopy at a spatial resolution of 18  $\mu$ m resolved single cells (Lasch et al., 2002). IR spectroscopy is also a tool for discrimination between different strains or types of cells (Gaigneaux et al., 2002).



**Fig. 5.18** Decomposition of a FTIR spectrum into three components corresponding to helical structure, sheets and non-regular structure, respectively. Percentages of structure content and structural changes, e.g., due to protein denaturation, are quantifiable



**Fig. 5.19** FTIR spectrum of a single molecular monolayer of A126C sperm whale myoglobin (Jiang et al., 1996). The peaks around 1660 cm<sup>-1</sup> and 1530 cm<sup>-1</sup> correspond to the amide I and amide II bands, respectively. The spectrum was acquired with a BioRad FTIR spectrophotometer equipped with a TGS detector

Fig. 5.18 shows an example for the decomposition of a FTIR spectrum of a protein into the components corresponding to helical, sheet-like and random coillike (non-regular) structures, respectively. Such decompositions can be calculated, e.g., by fitting a linear combination of the base spectra for the secondary structure components to the measured spectrum.

Fig. 5.19 illustrates the amazing sensitivity of FTIR spectroscopy. The sample was only two monolayers of a protein. Since at very low sample absorbances it is quite difficult to avoid the sharp lines of water-vapor absorption, these measurements were taken in a nitrogen-filled chamber at two different, very low concentrations of water, and later the water spectrum was subtracted. With this procedure, average artifact and noise levels were reduced to less than 0.00003 absorbance units.

# 6 Electron microscopy

### 6.1 Transmission electron microscope (TEM)

Transmission electron microscopy utilizes the wave properties of moving electrons to generate highly resolved images of specimens.

#### 6.1.1 General design

In 1986 the Nobel prize in physics was awarded by one half to Ernst Ruska for his fundamental work in electron optics, and for the design of the first electron microscope (EM), and by one half to Gerd Binnig and Heinrich Rohrer for their design of the scanning tunneling microscope (see Chap. 7). In some aspects, the



Fig. 6.1 Transmission electron microscope (see text on pp. 107 and 109)



Fig. 6.2 A more complicated design of a transmission electron microscope with an analyzer which can remove inelastically scattered electrons (see, e.g., LEO Elektronen-mikroskopie GmbH, Oberkochen, Germany)



**Fig. 6.3** In order to resolve the two points  $P_1$  and  $P_2$  of an object, the objective lens has to catch the first diffraction maximum of the two points. It appears in the direction where the diffracted (scattered) waves from the two diffracting (scattering) points have a phase difference,  $\Delta d$ , of one wavelength. Eq. (6.1) was derived from this condition. A typical objective lens has a bore of 2 mm and a focal length of about 1–2 mm

operation of a transmission electron microscope (TEM) is comparable with that of a slide projector (Figs. 6.1 and 6.2): Electrons from the electron gun pass through condenser lenses that focus the electrons onto the sample. The electron beam shines through the specimen. Objective lenses and projector lenses magnify the transmitted beam and project it onto the fluorescent viewing screen. Impact of electrons excites the screen and produces a visible magnified image of the sample. This image is recorded with various detectors, such as a CCD camera.

#### 6.1.2 Resolution

Electron microscopes enable significantly greater magnification and greater depth of focus than conventional optical microscopes. High-resolution TEMs permit spatial resolutions around 0.1 nm (1 Å) at acceleration voltages of 50–600 kV. Because of the wave nature of the electrons, the resolution limit, *d*, is given by the diffraction theory of coherent imaging:

$$d = \frac{\lambda}{n\sin(\alpha)} , \qquad (6.1)$$

where  $\lambda$ , *n*, and  $\alpha$  are the vacuum wavelength, index of refraction of the medium (=1 in TEMs), and aperture half angle of the objective lens, respectively (Fig. 6.3). The de Broglie relation provides the wavelength,  $\lambda$ , of the electrons:

$$\lambda = \frac{h}{mv} , \qquad (6.2)$$

where  $h = 6.6261 \times 10^{-34}$  J s, *m*, and *v*, are the Planck constant, electron mass, and electron velocity, respectively. For the relativistically high velocities of the electron beam we have to use Einstein's equations:

$$m = \frac{m_{\rm c}}{\sqrt{1 - \frac{v^2}{c^2}}}, \quad E = mc^2, \tag{6.3}$$

and obtain:

$$\lambda = \frac{hc}{\sqrt{2E_0\Delta E + (\Delta E)^2}} = \frac{hce^{-1}}{\sqrt{2m_0c^2e^{-1}V + V^2}} \approx \frac{1.24 \text{ nm kV}}{\sqrt{(1022 \text{ kV}) \cdot V + V^2}}, \quad (6.4)$$

where  $e = 1.6022 \times 10^{-19}$  C is the elementary charge,  $m_e = 9.1094 \times 10^{-31}$  kg the electron rest mass,  $c = 2.99792 \times 10^8$  m s<sup>-1</sup> the speed of light in vacuum,  $E = E_0 + \Delta E$ ,  $E_0 = m_e c^2$ ,  $\Delta E = V \cdot e$  the kinetic energy of the electrons, and V is the applied acceleration voltage, typically 200 V – 200 kV. For a voltage of, e.g., 100 kV we find  $\lambda = 0.0037$  nm. In contrast to most light microscopes, TEMs have small objective lens apertures of typically  $\alpha = 1-2^\circ$ , and thus according to Eq. 6.1 the limit of spatial resolution is 0.1–0.2 nm in this example.

#### 6.1.3 Electron sources

Thermionic electron guns (Fig. 6.4) and cold field emission guns (Fig. 6.5) are



Fig. 6.4 Thermionic electron gun (e.g., Structure Probe, Inc., West Chester, PA)



Fig. 6.5 Cold field emission gun focusable with an electrostatic lens comprised of two apertures with different electrostatic potentials



Fig. 6.6 In the production of an image of an object by an optical system, the product of area, A, and aperture solid angle,  $\Omega$ , remains about constant

common electron sources. A considerable concern is the brightness and size of electron sources. Fig. 6.6 illustrates why this is important: For a light beam passing through an optical system, the product of area and aperture solid angle of radiation remains constant. Thus, a large source can be focussed on a small spot only by using a large aperture angle of the optical system. Considering the limited aperture angles of electron lenses, a source of small size and high brightness is required to obtain a sufficiently bright picture of the sample.

#### 6.1.4 TEM grids

TEM grids (Fig. 6.7) should not get charged during measurement which would distort the electron path. Usually they are made from conductive chemically inert non-gassing materials suitable for high vacuum, such as platinum and platinum-iridium alloys.



**Fig. 6.7** TEM grids. (**a**), (**b**) Some common patterns of TEM grids made from a variety of materials, e.g., platinum, silver, tungsten, molybdenum, stainless steel, or titanium. Among these materials, platinum is the chemically most inert, but expensive. (**c**) Silicon nitride "grid" with a single window (from SPI Supplies, West Chester, PA)

#### 6.1.5 Electron lenses

There are magnetic (Figs. 6.8 and 6.9), electrostatic (Fig. 6.5) and compound lenses (Figs. 6.10 and 6.11). Electron lenses have some similar characteristics like optical lenses, such as focal length, spherical aberration, and chromatic aberration.



**Fig. 6.8** Ernst Ruska's pohlschuh lens: the circular electromagnet is capable of projecting a precisely circular magnetic field in the region of the electron beam



Fig. 6.9 Function of a magnetic electron lens (pohlschuh lens). One of Ernst Ruska's major achievements was the development of electron lenses



**Fig. 6.10** Compound lens made from magnetic and electrostatic lenses: the magnetic field forces the electrons on spiral-shaped trajectories; the electric field further decreases the diameter of the electron beam. Additional coils may serve for the correction of spherical aberration (not shown)

One important difference of electron optics compared with photon optics is the mutual charge interaction of electrons in the beam. That is why electron optics is often designed for beam paths with few if any intermediate crossovers.



Fig. 6.11 Compound lens made from magnetic and electrostatic lenses

In the beam booster technique (LEO Elektronenmikroskopie GmbH, Oberkochen, Germany; Fig. 6.12) a high energy beam is generated, passed through the condenser column of the microscope, and then decelerated and passed through the sample. The high energy electrons are less affected by stray magnetic and electric fields. Also the propagation of the electrons in the column is independent from the selected electron probe energy.



**Fig. 6.12** Beam booster (LEO Elektronenmikroskopie GmbH, Oberkochen, Germany). The electrons are accelerated to a high energy, passed through condenser lenses and filters, and then decelerated prior to interacting with the sample. This technique largely protects the electron beam against stray magnetic fields in the column of the microscope

#### 6.1.6 Electron-sample interactions and electron spectroscopy

There are different sources of chromatic aberration: (a) inelastic scattering of the electrons by the sample changes their energy (Fig. 6.13), and (b) the electrons leave the electron source with slightly different energies. The dispersion of electron energy is measured with energy filters (Fig. 6.14). Similar dispersive elements serve for the reduction of chromatic aberration, i.e., the selection of monochromatic electrons (Figs. 6.14 and 6.15)



**Fig. 6.13** Interaction of the electron beam with the sample. Inelastically scattered electrons have changed both direction and energy and may generate a diffuse contrast-reducing background image unless these electrons are eliminated by energy filtering (see Fig. 6.15). Elastically scattered electrons interfere with another and with unscattered electrons to produce a phase contrast image



**Fig. 6.14** Determination of electron energies. The voltage applied between the two hemispherical electrodes disperses the electrons with respect to their velocity. In order to record a full spectrum for a large range of electron energies, the applied voltage is swept



Fig. 6.15 Two types of dispersive elements for energy filtering of the electron beam to eliminate inelastically scattered electrons, removing the diffuse background and thereby enhancing contrast, or to perform a chemical analysis of the sample. *Left*:  $\Omega$ -filter. *Right*: filter using magnetic and electric fields

Phase differences due to differences in the optical pathlength and electron scattering contribute to the contrast (Fig. 6.16). Often it is quite difficult to generate sufficient sample contrast at very high resolutions. A common method to visualize very small biological structures, such as single protein molecules, is negative staining: the sample is embedded in a stain with strong electron-optical properties (Fig. 6.17). Important innovations towards better contrast were the introduction of a technique for enhanced resolution (Haider et al., 1998) and the nanofabrication of solid-state Fresnel lenses for electron optics (Ito et al., 1998).

For biological samples a further important reason for the limitation of the resolution of TEMs is radiation damage, i.e., the destruction of the sample by inelastically scattered electrons. Since a certain number of electrons is necessary to obtain an image, this limit depends on the ratio of inelastically to elastically scattered electrons. Practically the resolution of frozen protein molecules is restricted by this reason to worse than about 5 Å. Negatively staining (Fig. 6.17) may provide some improvement, nevertheless atomic resolution of proteins is still beyond reach. It was suggested the theoretical possibility of a neutron microscope, for which the ratio of elastically to inelastically scattered particles may much better for isotope-exchanged proteins (Henderson, 1996). Another theoreti-



**Fig. 6.16** Generation of amplitude contrast. The electron beam is weakened at different points to a different degree by scattering and interference: elastically scattered electrons, i.e., those which have changed direction but not energy, interfere with each other and with unscattered electrons to produce a phase contrast image



Fig. 6.17 Edge-on view of a negatively stained sample: the TEM senses volumes of lower density in the stain

cal possibility to overcome the problem of the decay of the structure of sample during the measurement might be the use of ultrashort electron flashes on deeply frozen samples: if the duration of the flash is shorter than the time of mechanical movement of the protein molecule, its chemical decomposition would affect the obtained micrograph to a lesser degree.

#### 6.1.7 Examples of biophysical applications

Fig. 6.18 demonstrates the resolution power of TEM for large protein complexes (Roseman et al., 1996; White et al., 1997; Ranson et al., 1998; Rye et al., 1999; Saibil, 2000a). Clearly differences between two conformations of GroEL/GroES are resolved. The TEM structure is consistent with the crystal structure.

Electron microscopy resolved the structure of the bacteriophage  $\Phi 29$  packaging motor (Simpson et al., 2000) and visualized the filamentous phage pIV multimer (Linderoth et al., 1997). Electron microscopy contributed to the understanding of conformational changes connected with the opening of an ion channel through a membrane (Saibil, 2000b), and with connexin trafficking (Gaietta et al., 2002).

In groundbreaking experiments Terry G. Frey and coworkers succeeded in the 3D-visualization of cell organelles using electron tomography. In this method the three-dimensional structure is calculated from a series of electron micrographs of samples tilted over a range of angles (Dierksen et al., 1992; Perkins et al., 1997a, 1997b; Frey and Mannella, 2000).



**Fig. 6.18** *Top:* electron micrograph of two conformations of GroEL/GroES at 30 Å resolution (Roseman et al., 1996; White et al., 1997; Ranson et al., 1998; Rye et al., 1999; Saibil, 2000a). *Bottom:* the X-ray crystal structure of GroEL at 2.4 Å resolution for comparison (Braig et al., 1994; Boisvert et al., 1996). The latter figure part was generated using MOLSCRIPT (Kraulis, 1991)

## 6.2 Scanning transmission electron microscope (STEM)

In contrast to TEMs, scanning transmission electron microscopes use an electron beam with only a few Å or nm diameter to scan the sample area (Fig. 6.19). The resolution is generally limited by the diameter of the electron beam at the location of the sample and radiation damage.



**Fig. 6.19** Scheme of a scanning transmission electron microscope: The objective lens focuses the electron beam onto a small sample section. Scattered electrons are detected with the elastic dark field detector. The STEM image is generated by moving the focussed beam over the specimen

Although the STEM was pioneered already in the thirties of last century, mainly by adding scan coils to a TEM (von Ardenne, 1940), significant developments have taken place in the last years: electron optics has been significantly improved and the resolution increased by several orders of magnitude. Nowadays high resolution STEMs offer unprecedented capability for the characterization of biomolecules, allowing structure to be determined with up to sub-nm resolution.

Similarly to TEMs, the STEM can employ various energy filtering techniques for chemical analysis and improvement of resolution, e.g., by removing unscattered electrons in inelastic dark field imaging (Fig. 6.20). Many STEM have both capabilities, elastic dark field imaging (Fig. 6.19) and inelastic dark field imaging (Fig. 6.20). A third mode is bright field detection where electrons are collected through a small aperture placed on the optical axis and an energy



**Fig. 6.20** Example of a scanning transmission electron microscope with an energy filter: inelastically scattered electrons, i.e., those which have changed both energy and direction upon interaction with the specimen, are collected yielding the inelastic dark field image. Electrons with different energies are separable by their trajectories with different curvatures in the electric field applied perpendicular to the flight direction

filter removes those electrons that have lost energy, i.e., low-angle elastically scattered and unscattered electrons are collected to produce the image.

# 7 Scanning probe microscopy

Scanning probe microscopes generate a highly-resolved image of the specimen by scanning it with a small mechanical, electrical, optical, thermal, or other probe.

### 7.1 Atomic force microscope (AFM)

The AFM was invented by Gerd Binnig, Christoph Gerber, and Calvin F. Quate in the mid-eighties (Binnig et al., 1986), and is one type of the so-called scanning



**Fig. 7.1** Principle of operation of an atomic force microscope. A very sharp tip attached to a tiny cantilever probes the sample surface. An optical system comprised of diode laser and detector, e.g., a diode array or a position-sensitive diode, senses the bending of the cantilever and thereby the distance-dependent tip-sample interaction force. For scanning the surface, the sample is moved by the piezoelectric scanner (Binnig et al., 1986)

probe microscopes (SPMs) which also include scanning tunneling microscope (STM; Sect. 7.2; Binnig et al., 1982a, 1982b, 1983; Binnig and Rohrer, 1987), scanning near-field optical microscope (SNOM; Sect. 7.3), scanning thermal microscope (SThM; Sect. 7.4), and the scanning ion conductance microscope (SICM; Sect. 7.4). The AFM is used in both industrial and fundamental research to obtain atomic-scale images of metal surfaces and nanometer-scale images of the three-dimensional profile of the surface of biological specimens. It is a very useful tool for determining the size and conformation of single molecules and aggregates adsorbed on solid surfaces. The AFM scans the sample with a tiny tip mounted on a small cantilever (Fig. 7.1). It measures the small force of interaction between tip and sample surface by sensing the reflection changes of a laser upon cantilever movement caused by interaction with the sample. An image of the sample surface relief is recorded using piezoelectric translation stages that move the sample beneath the tip, or the tip over the sample surface, and are accurate to a few Å.

Note the similarity of the AFM (Fig. 7.1) to the stylus profilometer (Fig. 7.2) and to the STM (Fig. 7.19). Actually, the idea of AFM is based on the design of stylus profilometers, but the AFM can reveal the sample relief with subnanometer resolution.



**Fig. 7.2** Stylus profilometer for comparison with the AFM. A set of styli probes the sample which is drawn below the set of styli. The small motions of the styli are transformed into an electrical signal by linear, variable transducers. Step heights of down to a few 10 nm are resolvable

The force of tip-sample interaction (Fig. 7.3) has a magnitude of typically only a few pN - nN. That is why the cantilever must have a small mass, and the

weight-bearing parts of the AFM (Figs. 7.4 and 7.5) have to be rigid and equipped with a good vibrational damping. Three support posts in the design of Fig. 7.4 reduce wobbling. The xyz-translation stages for coarse adjustment of the piezoelectric scanner with the sample on top are engineered for little wobbling as well. The whole AFM is placed on a rubber support preventing transmission of high frequency vibrations from the laboratory (not shown). A low force of interaction is crucial for high resolution force microscopy on soft biological specimens. Low spring constants of the cantilever may facilitate this purpose at the expense of resolution, but the most common way of gentle measurement is to reduce the intensity and duration of contact by oscillating the cantilever, as will be explained later (Fig. 7.13).



**Fig. 7.3** When a small tip approaches a surface, it experiences the van-der-Waals force which is attractive at a distance of a few Å, but repulsive at very short distances (see, e.g., Chap. 3 in Nölting, 2005). Additional Coulomb forces may play a role when the AFM tip was charged



**Fig. 7.4** Design of an AFM. The sample spot of interest is positioned near the tip by coarse xyz-translation stages. The piezoelectric scanner (see also Fig. 7.5) then heightens the sample position further till the tip starts to interact with the sample. It allows motion control of the tip with subnanometer precision. A photodiode detects the reflection changes of a laser beam from the cantilever upon approach of tip to sample. In this example, the cantilever is mounted to a dithering piezo element which excites oscillations of the cantilever. The lock-in amplifier detects changes of these oscillations due to tip-sample interactions. The sample surface is scanned by sample movement in horizontal direction by the piezoelectric scanner. The scanner also adjusts the relative height of the cantilever during scanning to avoid crashes of the tip with the sample surface. Such crashes can damage the tip and then cause artifacts (see Fig. 7.6)



**Fig. 7.5** Principle of operation of the xyz-piezoelectric scanner, a ceramic positioning device which changes its size in response to a change in applied voltage. A voltage change at the x- or y- electrodes causes bending in the horizontal plane; contraction and expansion are generated by simultaneous application of x- and y-voltage



**Fig. 7.6** Artifacts caused by different shapes of the AFM tip. Blunt tips and broken tips give rise to an seemingly flattened sample relief which may be difficult to recognize as an artifact



**Fig. 7.7** Tungsten tip, made by sharpening a tungsten wire by dragging it over a plate coated with alumina. This self-made tip may also be used for STMs (see Sect. 7.2)

Obviously a robust and sharp single tip is essential for this method. Typical apex radii of commercial tips are  $\approx 10-20$  nm. Fig. 7.6 depicts common types of artifacts observed when using worn out tips, broken tips, or probes with more than one tip.

One can make tips themselve by grinding a tungsten wire on a sheet covered with alumina (Fig. 7.7). These tips are also suitable for STM (Sect. 7.2), but the tip shape is not very reproducible and tungsten is not very hard.



Fig. 7.8 Sharpened pyramidal silicon nitride tip. SiN is extremely hard, and tips can be engineered with radii of only a few nm

Silicon nitride tips are the currently available tips with highest robustness (Fig. 7.8). Sharper tips with quite reproducible shape are made from silicon which is relatively fragile, however (Fig. 7.9a). Even sharper tips for application on samples with particularly deep structures are manufactured by attaching a high

density carbon fiber to a silicon tip (Fig. 7.9b). Sharp cantilevers for the examination of very rough surfaces (Fig. 7.10) and cantilevers with trigonal design for the purpose of high resistance against torsion (Fig. 7.11) are supplied, e.g., by Olympus Optical Co. (Tokyo).



Fig. 7.9 (a) AFM tip made from silicon. (b) Silicon tip with a high density carbon fiber attached to it



Fig. 7.10 Sharp cantilever geometry for very rough samples (e.g., Olympus Optical Co., Tokyo)



**Fig. 7.11** A trigonal design of the cantilever (e.g., Olympus Optical Co., Tokyo) causes a better stability against torsion, compared with rod-shaped cantilevers

The life of the AFM tip decreases very rapidly with applied force (Fig. 7.12). High aspect-ratio tips made from silicon or carbon fiber are generally less durable than low aspect-ratio tips made from silicon nitride.



Fig. 7.12 Example for the wear of two AFM tips due to surface load



**Fig. 7.13** (a) Contact mode: the cantilever follows the height profile of the sample. (b) Dynamic mode: the cantilever has only intermittent contact or oscillates above the sample. In the latter mode, oscillations are excited by a piezo crystal and the forces between tip and sample are very small. This mode permits truly atomic resolution (Giessibl, 2000)

There are two common modes of operation of AFMs (Fig. 7.13): the contact mode and the dynamic force mode. In the contact mode, the probe tip is in continuous contact with the sample surface. The force the cantilever exerts on the substrate in contact mode may perturb the surface of soft biological materials. In the gentler dynamic mode, the probe tip only oscillates up and down as it is scanned over the sample surface. Two sub-modes may be distinguished for the dynamic force mode, the non-contact sub-mode in which the distance between tip apex and sample surface is always larger than the van-der-Waals distance, and the tapping sub-mode in which the tip has intermittent contact.

As pointed out, a high degree of protection against external high-frequency vibrations is obviously crucial for the operation scanning probe microscopes with atomic resolution. Fig. 7.14 shows a further solution to this problem. Here the AFM is made from very thick and short plates of steel and the AFM is placed on three rubber balls that do not transmit fast vibrations. Another technique of efficient vibrational damping is to hang the AFM on a rubber string (Fig. 7.15).



**Fig. 7.14** Robust design of an AFM with atomic resolution. The vibrational damping is attained by a very rigid construction and an elastic support in form of three rubber balls



Fig. 7.15 "Hanging AFM": vibrational isolation of an AFM by hanging it on a rubber string

Important biological applications of AFMs were the direct observation of the structure of DNA (Lindsay et al., 1989) and the monitoring of actin filament dynamics in living cells (Henderson et al., 1992). The direct visualization of a DNA glycosylase searching for damage shows that the glycosylase interrogates DNA at undamaged sites by introducing drastic kinks (Chen et al., 2002b). Intramolecular triplex DNA formation results in a kink in the double helix path (Tiner et al., 2001). A sharp DNA bend is induced by binding of integration host factor (IHF) to the region between the upstream regulatory sequence and the promoter sequence (Seong et al., 2002). Single DNA molecule force spectroscopy can discriminate between different interaction modes of small drug molecules with DNA by measuring the mechanical properties of DNA and their modulation upon binding of small drug molecules (Krautbauer et al., 2002) and dye molecules (Kaji et al., 2001). A decrease of the ionic strength from 50 mM to 1 mM resulted in a change of the number of nodes (crossings of double helical segments) of a supercoiled 3000-bp piece of DNA from a 15 to one or two nodes (Cherny and Jovin, 2001). High resolution fluorescence imaging of  $\lambda$ -phage DNA molecules, intercalated with the dye YOYO-1, by a SNOM/AFM (SNOM, scanning nearfield optical microscope; see Sect. 7.3) resolved the distribution of the dye (Kim et al., 2001).

AFM proved to be a very useful tool for the study of proteins, yielding some unique insights into structure and physical properties:  $\beta$ -Lactoglobulin forms fine-stranded aggregates at pH 2 with the diameter of strands being ca. 4 nm (Ikeda and

Morris, 2002). AFM technology was used to map out the electrostatic potential of the transmembrane channel OmpF porin (Fig. 7.16; Philippsen et al., 2002). AFMs gave crucial topological information of blood cell adhesion on different sensor materials (Hildebrand et al., 2001). Ac-GWWL(AL)nWWA-Etn peptides induce the formation of extremely ordered domains in some biologically relevant membranes (Rinia et al., 2002). The heads of bacteriophage  $\Phi KZ$  and T4 have different compressibilities (Matsko et al., 2001). Atomic force microscopy resolved fusion pores in the apical plasma membrane in live pancreatic cells (Cho et al., 2002) and visualized the growth of Alzheimer's *β*-amyloid-like fibrils (Goldsbury et al., 2001). Cardiac muscle and skeletal muscle exhibit different viscous and elastic properties as determined by atomic force microscopy. Cardiac cells are stiffer (elastic modulus =  $100 \pm 11$  kPa) than skeletal muscle cells (elastic modulus =  $25 \pm 4$  kPa; see Mathur et al., 2001). Atomic force microscopy allowed to visualize the structure of biomolecules, e.g., the native chaperone complex from Sulfolobus solfataricus, in solution under physiological conditions providing a nanometer resolution topographic image of the sample (Valle et al., 2001). It is also an excellent technique to study the initial events of mutual cell adhesion (Razatos, 2001). An AFM image of a monomolecular film of bovine serum albumin shows individual monomers and dimers (Fig. 7.17; Gunning et al., 1996; Morris et al., 1999).



**Fig. 7.16** Imaging the electrostatic potential of the transmembrane channel OmpF porin (Philippsen et al., 2002). Different apparent shapes of the porin are observed at different ionic strengths. These differences reflect changes of the electrostatic potential which is experienced by the charged tip of the AFM



**Fig. 7.17** AFM image of a monomolecular film of the protein bovine serum albumin (BSA,  $M_{w,monomer} = 66$  kDa) adsorbed at an oil/water interface (Gunning et al., 1996; Morris et al., 1999). Individual monomers and dimers of BSA can be seen

AFMs are also very useful for the manipulation of macromolecules: proteins may physisorb to the AFM tip and can then be lifted and manipulated (Fig. 7.18). The sensitivity of the AFM cantilever, to forces in the pN range, was exploited to measure folding-unfolding forces within single protein molecules and breakaway forces between different biomolecules (Jiao et al., 2001; Allison et al., 2002). Atomic force microscopy has yielded tantalizing insights into the dynamics of protein self-assembly and the mechanisms of protein unfolding (Furuike et al., 2001; Yip, 2001). For further, similar applications of AFM technology see Chap. 8.



**Fig. 7.18** Manipulation of a protein molecule with an AFM: The tip is lowered till it touches the macromolecule. Due to the attractive action of the van-der-Waals interaction, the macromolecule sticks to the tip and can be lifted and moved to a different place

## 7.2 Scanning tunneling microscope (STM)

In 1986 Gerd Binnig and Heinrich Rohrer were awarded the Nobel Prize for Physics for the groundbreaking invention of the STM. It was the first member of the family of scanning probe microscopes (SPM) that can characterize surface morphology with atomic resolution. In contrast to the AFM, its principle of operation (Fig. 7.19) requires electrically conductive samples. A sharp tip located on a flexible cantilever is used to probe the distance between the tip and sample surface, as judged by the tunneling current (Fig. 7.20). Since the tunneling current also depends on the chemical nature of sample and tip, the STM also serves for characterization of electronic properties of solid samples.

Significant complications on the way towards higher resolution of proteins are the undesired excitation of the soft biological material by the high current of STMs, typically pA-nA, and the distribution of conductivity within the sample distorting the image (Fig. 7.21). Low currents and stable attachment of the sample to the support are required for high resolution images of biological macromolecules.



**Fig. 7.19** Principle of operation of STMs. A finely sharpened electrically conductive tip is first positioned within about 1 nm of the sample by mechanical translation stages (not shown) and the piezoelectric scanner. At this small separation, electrons tunnel through the gap between tip and sample (Fig. 7.20). The tunneling current depends on the applied bias voltage between tip and sample, the distance, the tip shape, and the chemical compositions of sample and tip. The feedback loop ensures constant height or constant current. Tunneling current and feedback voltage are a measure of surface morphology and composition (Binnig et al., 1982a, 1982b, 1983; Binnig and Rohrer, 1987)



**Fig. 7.20** When the distance between conductive tip and conductive sample is lowered to a few Å, electrons can traverse the gap with some transmission probability. The STM measures not purely distance like the AFM, but the local density of electronic states



**Fig. 7.21** Different possible current paths in STM measurements of extended biological structures lower the resolution and complicate the interpretation of data

Similar as in atomic force microscopy, the quality of the tip is crucial for a high resolution (Figs. 7.7-7.11, 7.22). Commonly tips are micromachined and/or electrochemically etched and have apex radii of 5-100 nm.

STM images of a 1:1 mixture of K344C cytochrome  $P450_{cam}$  / putidaredoxin adsorbed on gold (111) showed a regular array of pairs of the two proteins (Djuricic et al., 2002). I21C/E25C plastocyanin essentially maintains its native redox properties upon immobilization onto a gold substrate as shown by the full potentiostatic control of the electron transfer reaction during STM imaging (Andolfi et al., 2002). Scanning tunneling microscopy demonstrated that the otherwise nearly linear mammalian metallothionein-2 molecule bends by about 20 degrees at its central hinge region between the domains in the presence of ATP (Maret et al., 2002). Electrochemical scanning tunneling microscopy on thiol-derivatized DNA immobilized on a gold (111) single crystal surface showed



Fig. 7.22 An etched tungsten tip for STM (see also Figs. 7.7–7.11)

potential-dependent orientation changes of the DNA in the potential range from 200 to 600 mV (Zhang et al., 2002b). A STM study of morphology and electron transport features in cytochrome c offers evidence for sequential discrete electron-tunneling effects (Khomutov et al., 2002). Tunneling in proteins adsorbed onto a conductive substrate may depend on the applied potential (Facci et al., 2001). The resistance of a single octanedithiol molecule is 900 M $\Omega$  (Cui et al., 2001). The ability to site-specifically introduce cysteine residues and to engineer tags, such as histidine tags and biotin-acceptor peptides, allow the creation of ordered immobilized protein structures that can be characterized both electrochemically and topographically by using scanning probe microscopy and cyclic voltammetry (Gilardi et al., 2001).

### 7.3 Scanning nearfield optical microscope (SNOM)

#### 7.3.1 Overcoming the classical limits of optics

SNOMs (Figs. 7.23–7.25), also known as NSOMs, utilize a light source with a diameter smaller than the wavelength of the light (Synge, 1928; Ash and Nicholls, 1972; Pohl et al., 1984; Betzig et al., 1986, 1991, 1992; Toledo-Crow et al., 1992; Williamson et al., 1998; Egawa et al., 1999; Heimel et al., 2001). By means of this technological innovation they achieve a resolution which may be well beyond the resolution limit, d, of classical Abbe-Fourier optics (see also Sect. 6.1.2):

$$d = \frac{\lambda}{n\sin(\alpha)},\tag{7.1}$$

where  $\lambda$ , *n*, and  $\alpha$  are the vacuum wavelength, refractive index of the medium between sample and objective lens, and half angle of aperture, respectively. For visible light with  $\lambda = 500$  nm, n = 1.6, and  $\alpha$  near 90°, we obtain a resolution limit of classical optics of about 300 nm. Using UV light and image processing can yield improvement beyond this, but it is clear from Eq. 7.1, that classical optics can hardly penetrate the 100-nm resolution barrier. SNOMs have been the first optical microscopes that significantly overcame the limit of Eq. 7.1.



**Fig. 7.23** A design of a SNOM. An optical probe emits light from points above the 2D sample. The subwavelength probe tip takes advantage of the physical effect of optical tunneling. In this SNOM it is made using a metal coated tapered glass fiber: a first taper for the probe was manufactured by melt-drawing, and a second taper at the very end of the probe was etched. The damping of the horizontally vibrating SNOM probe caused by shear forces is taken as a measure for the distance from the sample. A shear force feedback loop which involves an interferometric measurement of the horizontal position of the probe tip keeps it at constant height (Betzig et al., 1992). Near-field optical properties of the sample surface are mapped out by scanning each point within a certain area


Fig. 7.24 Design of an interferometer used for the detection of shear force of the SNOM tip. The principle of operation is based on the extinction of light at the position of the detector due to the interference of the two incident light beams of equal amplitude when the probe tip (movable mirror) is in position 1. A small perturbation of the position of the tip causes a small deviation from  $\lambda/2$  of the phase shift of the two beams inciding on the detector which leads to a non-negligible detector signal. The light filter in the fixed path ensures an equal amplitudes of the two interfering beams

Generally the working principle of SNOMs is as follows (Figs. 7.23–7.25): The subwavelength light source is positioned a few  $\mu$ m above the specimen surface with the help of mechanical coarse translation stages and a piezoelectric fine translation device, e.g., a piezoelectric scan tube. Transmission is measured below the specimen. An image of the specimen surface is obtained by moving the light source (or the sample in some designs) with the help of the piezoelectric fine translation device in horizontal direction. To avoid crashes with the sample, in many SNOMs the light source is oscillated over the specimen surface and damping of oscillations due to source-sample interactions detected. The piezoelectric fine translation device somewhat lifts the subwavelength light source (or lowers the sample) when damping increases. Near-field optical excitation of the sample can be seen as a dipole-dipole energy transfer (Sekatskii and Dietler, 1999).



Fig. 7.25 A design of a SNOM with a xyz-coarse positioning and a xyz-fine positioning

There is virtually no resolution limit of SNOMs as long as one can manufacture light sources of sufficient small size (see next section) and detect very small intensity differences of light passing through thin sample layers.

#### 7.3.2 Design of the subwavelength aperture

The most common methods to manufacture subwavelength apertures are (a) adiabatic pulling of an optical fiber during heating (Betzig et al., 1991; Williamson and Miles, 1996; Figs. 7.26a and 7.27), (b) etching (Muramatsu et al., 1999), in particular tube etching (Turner, 1983; Stöckle et al., 1999a; Figs. 7.26b

and 7.28), and (c) microfabrication (Schurmann et al., 2000; Mitsuoka et al., 2001), e.g., by ion beam milling (e.g., Veerman et al., 1998).



**Fig. 7.26** (a) Melt-drawn probe: in the upper part of the probe, the light is reflected from the walls by total internal reflection (TIR). TIR is the phenomenon involving the reflection of all the incident light off a boundary when both (i) the light is traveling in the more optically dense medium and approaching the less optically dense medium and (ii) the angle of incidence is greater than the so-called critical angle. (b) Etched probe. A desirable high brightness is obtained by a large cone angle. An optical aperture is formed by rotational evaporation of the etched fiber with an opaque metal, e.g., aluminum



**Fig. 7.27** Fabrication of a probe by adiabatic pulling of an optical fiber during heating (melt-drawing; see, e.g., Williamson and Miles, 1996)



**Fig. 7.28** Tube-etching of a quartz glass fiber with hydrofluoric acid (Turner, 1983; Stöckle et al., 1999a, 1999b): the tip forms due to concentration gradients of acid and desolved fiber. The tube serves also for suppression of convection of the acid. These tube-etched tips are inexpensive and have large cone angles permitting high light throughputs



**Fig. 7.29** Propagation of a light wave in a SNOM probe tip. Up to the cut-off diameter (about equal to the wavelength), the light travels with only little attenuation. Beyond this critical diameter, the light intensity very rapidly falls off

Usually the aforementioned probe tips are coated with a metal. Since the light transmission of the tip dramatically depends on the distance from the cut-off region to the aperture (Fig. 7.29), the optical throughput of etched tips with cone angles around  $30^{\circ}$  is, in general, 2-3 orders of magnitude better than that of heat-pulled fiber probes. Even brighter sources can be microfabricated (Fig. 7.30).



Fig. 7.30 SNOM tip made from silicon nitride. It was manufactured by using photolithography, potassium hydroxide etching, and electron beam nanolithography (Zhou et al., 1998, 1999)



**Fig. 7.31** Optical far-field and near-field in the vicinity of a small optical tip for a SNOM. Since the light can transmit through thin layers of metal, the diameter of the light beam in this design cannot be less than a few 10 nm



**Fig. 7.32** A 10-nm sized light source for SNOMs made from fluorescent material. UV light excites fluorescence of the 10-nm sized bead at the end of the probe. The transmission of the fluorescence light through the sample is sensed by a detector which is covered with an UV-absorbent layer. Antioxidants may be added to the fluorescent material to enhance stability and life. The resolution for thin-layered samples enabled by this type of light sources depends on the size of the fluorescent bead and the detectability of small absorbance differences of the fluorescent light

In the aforementioned methods of source fabrication, a limitation for the source size is the transmission of light through thin layers of metal (Fig. 7.31). Sources of 10 nm diameter could hardly be made just by machining an opening in a metal plate or in an optical fiber since light would significantly shine through the walls of the opening. Smaller light sources involve the excitation of small fluorescent particles (Fig. 7.32).

#### 7.3.3 Examples of SNOM applications

An important applicability is seen in cell biology, microbiology (Meixner and Kneppe, 1998), and proteomics (Gao et al., 2001). SNOM imaging visualized domains of photosystem II core complex bound to lipid monolayers (Trudel et al., 2001). Topographic, friction, fluorescence, and surface potential distributions for a Langmuier-Blodgett film can simultaneously be observed using a SNOM-AFM with a thin step-etched optical fiber probe (Horiuchi et al., 1999). Fig. 7.33 demonstrates a 50-nm resolution.



**Fig. 7.33** Absorption of latex beads with approximately 100 nm diameter taken with a SNOM at a resolution of about 50 nm (OMICRON, Taunusstein, Germany)

## 7.4 Scanning ion conductance microscope, scanning thermal microscope and further scanning probe microscopes

Important scanning probe microscopes are also the magnetic force microscope, scanning Hall probe microscope (Chang et al., 1992a, 1992b), friction force microscope (Fig. 7.34; Howald et al., 1995), scanning ion conductance micro-



**Fig. 7.34** Example of a friction force microscope: the detector has sectors in both vertical and horizontal direction so that the torsion of the cantilever can be estimated and a frictional (lateral) force be calculated (see, e.g., Howald et al., 1995)

scope (Fig. 7.35; Hansma et al., 1989; Korchev et al., 1997, 2000a, 2000b; Stachelberger, 2001; Bruckbauer et al., 2002a), and scanning thermal microscope (Fig. 7.36; Mills et al., 1998, 1999). There are two common types of thermal probes for scanning thermal microscopes: thermocouples and thermal resistors. The thermocouple probe in Fig. 7.36 involves two dissimilar metal wires which bisect over the top of a blunt silicon nitride pyramid (Mills et al., 1998). A potential biophysical application is the elucidation of local heating effects in biological tissue due to cell metabolism.



**Fig. 7.35** Detection of membrane pores by a scanning ion conductance microscope. The probe is a nanopipette filled with electrolyte solution. The current between pipette and ion reservoir starts increasing when the pipette tip approaches a pore. The technique permits 100-nm resolution characterization of distribution and sizes of pores



**Fig. 7.36** Design of the tip of a scanning thermal microscope (Mills et al., 1998, 1999). A submicrometer-sized Cu/Ni thermocouple at the end of the cantilever detects the thermal microenvironment



**Fig. 7.37** STM with simultaneous AFM capability: AFM and STM sense different physical properties. The combined information provides greater insight into the chemical nature of the sample ant its physico-chemical properties

The STM with simultaneous AFM capability (Fig. 7.37) provides simultaneously information about surface relief and chemical composition of the specimen.

#### 8 Biophysical nanotechnology

#### 8.1 Force measurements in single protein molecules

Atomic force microscope (AFM)-related techniques can induce and monitor the unfolding of single protein molecules. Experiments on the protein titin, which is a main component of skeletal muscles (Figs. 8.1-8.3), revealed that the force for unfolding of its individual domains with cross sections of less than 5 nm<sup>2</sup> is of the order of 100-300 pN and dependent on the pulling speed (Rief et al., 1997; Gaub and Fernandez, 1998; Carrion-Vasquez et al., 1999). A similar investigation on bacteriorhodopsin showed that its helices are anchored to the bacterial membrane with 100-200 pN (Fig. 8.4; Oesterhelt et al., 2000). Similarly, single-molecule force spectroscopy on spider dragline silk protein molecules revealed that the molecule unfolds through a number of rupture events, indicating a modular structure within single silk protein molecules (Oroudjev et al., 2002). The minimal unfolding module size of 14 nm indicates that the modules are composed of 38 amino acid residues (Oroudjev et al., 2002). Adhesion between two adjacent cell surfaces of the eukaryote Dictyostelium discoideum involves discrete interactions characterized by an unbinding force of about 23 pN. This force probably originates from interactions of individual "contact site A" (csA) glycoprotein molecules (Fig. 8.5; Benoit et al., 2000).



**Fig. 8.1** Molecular architecture of skeletal muscle fibers. AFM-related techniques contributed to the understanding of the role of individual titin molecules in such fibers: some skeletal muscle proteins can withstand drags of 600 kp cm<sup>-2</sup> (see Figs. 8.2 and 8.3; Rief et al., 1997; Gaub and Fernandez, 1998; Carrion-Vasquez et al., 1999)



**Fig. 8.2** Unfolding of a titin fragment with the help of an AFM (Gaub and Fernandez, 1998; Carrion-Vasquez et al., 1999). The unfolding force for the protein, anchored with a cysteine (Cys) to a gold surface, ranged from about 100 to 300 pN (see Fig. 8.3)



**Fig. 8.3** Sketch of a force-distance curve for the unfolding of a titin fragment with the help of an AFM (see Fig. 8.2). The saw tooth-shaped force-extension curve reflects the unfolding of individual titin domains according to an all-or-non mechanism (Gaub and Fernandez, 1998; Carrion-Vasquez et al., 1999)



**Fig. 8.4** Unfolding of individual bacteriorhodopsins. A force of 100-200 pN is required to remove a bacteriorhodopsin helix from the bacterial membrane (Oesterhelt et al., 2000)

Recoverin, a calcium-myristoyl switch protein, binds to a phospholipid bilayer in the presence of  $Ca^{2+}$  with an adhesion force of  $48 \pm 5$  pN (Desmeules et al., 2002). Single molecules of *holo*-calmodulin (i.e., the calcium-loaded form) require a significantly larger force of unfolding by an AFM tip than single molecules of the *apo*-form (Hertadi and Ikai, 2002). Single molecules of the giant filamentous protein titin exhibit mechanical fatigue when exposed to repeated stretch and release cycles (Kellermayer et al., 2001). For further AFM studies on single protein molecules see also Sects. 7.1, 8.2, and 8.3.



Fig. 8.5 Measurement of discrete interactions in cell adhesion (Benoit et al., 2000)

### 8.2 Force measurements in a single polymerase-DNA complex

DNA polymerases catalyze DNA replication. The replication reaction requires single-stranded DNA (ssDNA) as a template. In the course of the reaction, a complementary strand of ssDNA is synthesized to the original ssDNA. Already during the polymerization reaction, both strands coil around each other, leading to a shortening of the end-to-end distance of the DNA. Exerting strain on the DNA strand during polymerization can stop and even revert the extension reaction



**Fig. 8.6** Optical tweezers for the measurement of the effect of template tension on T7 polymerase activity (Fig. 8.7; Smith et al., 1996; Wuite et al., 2000). Regarding the method of optical tweezers see also the groundbreaking studies of single-molecule mechanics by Florin et al. (Florin et al., 1997; Jeney et al., 2001; Pralle and Florin, 2002) and Smith et al. (Bustamante et al., 2000; Liphardt et al., 2001; Smith et al., 2001)



**Fig. 8.7** Measurement of the effect of template tension on T7 polymerase activity (Smith et al., 1996; Wuite et al., 2000): the polymerase which catalyses DNA replication can work against a maximum force of about 34 pN. Exonuclease activity increases about 100-fold above 40 pN template tension.

(Figs. 8.6 and 8.7; Smith et al., 1996; Wuite et al., 2000). For the measurement of the small forces in the single DNA-protein complex, an optical trap was used: a small bead with DNA attached to it is held into position and moved by an intense laser beam. The main mechanism of the action of such optical tweezers is commonly as follows: The bending of light rays through the refractive sphere is connected with a change of momentum of the light which exerts a force back on the sphere. When the sphere is out of focus of the light beam, these light deflection forces pull the sphere back into focus. Another mechanism is as follows: When an isotropically scattering bead moves out of focus, the momentum of the photons scattered in the direction of the movement increases due to the Doppler effect which decelerates the bead. The larger the light intensity the larger is the deceleration of the movement. Thus, the Brownian motion out of focus is energetically unfavorable relative to the motion into focus. Choosing a wavelength just below an absorption maximum of the bead increases the trapping force since then movement causes increased absorption due to the Doppler-shift of the wavelength and thus an additional momentum slowing down the bead.

#### 8.3 Molecular recognition

AFM-related techniques allow the direct measurement of individual intermolecular interactions (Figs. 8.8–8.11; Florin et al., 1994; Dammer et al., 1995, 1996; Merkel et al., 1999; Strunz et al., 1999; De Paris et al., 2000; Fritz et al., 2000; Schwesinger et al., 2000; Zocchi, 2001; Prechtel et al., 2002; see also Sect. 7.1). Virtually any intermolecular interaction forces, e.g., of antibody-antigen interactions, are measurable with the technique illustrated in Fig. 8.8.



**Fig. 8.8** Direct measurement of intermolecular interactions by an AFM-related technique. One of the interacting molecules is immobilized on the surface of the support, the other is connected to the AFM tip by a linker. The tip is approached to the surface so that a specific interaction can take place. Retracting the cantilever ruptures the biophysical interaction. The strength of the interaction is determined from the retract force distance curves (De Paris et al., 2000; Schwesinger et al., 2000; see also Sect. 7.1)



**Fig. 8.9** Sensor of biological agents using recognition between single DNA molecules (Park et al., 2002; Service, 2002; see also Demers et al., 2002). (a) Two electrodes and single-stranded capture DNA strands are attached to a glass substrate. The capture DNA is complementary to one end of the target DNA of the agent. (b) Target DNA and probe strand DNA was added. The probe strand DNA has a gold nanoparticle attached and is complementary to the other end of the target DNA. When all three strands of DNA hybridize together, the gold nanoparticle gets held between the two electrodes. This is detected by an increase of current

Fig. 8.9 illustrates a new type of DNA sensor with potential application for detection of biological contaminants (Park et al., 2002; Service, 2002). It is based on the change of electrical conductivity when gold particles attached to DNA bind



**Fig. 8.10** Force measurement on single molecular contacts through evanescent wave microscopy (Zocchi, 2001). The motion of the bead attached to the wall of the flow channel through a single streptavidin-biotin complex is tracked by detecting the evanescent wave as a force is exerted through a flow. This technique allows the direct measurement of the bond rupture force of the molecular complex



**Fig. 8.11** Nanobiosensor using the cantilever of an AFM (Pereira, 2001). Entering or exiting of specific molecules, including medications, from living cells is observed in real-time and the force of interaction measured

to the targeted sample DNA immobilized between two electrodes. A problem of this detection method might be the difficulty to find pieces of DNA that are unique for the organism of interest. In particular, some genetically engineered bacteria and viruses might remain undetected.

#### 8.4 Protein nanoarrays and protein engineering

Lee et al. (2002b) manufactured protein nanoarrays by means of dip-pen nanolithography (Figs. 8.12 and 8.13): A gold thin-film substrate was patterned, by using an AFM, with a protein-binding chemical linker in the form of dots or grids. Non-patterned surface was inactivated and then protein bound to the linker patterns. Another method for the manufacture of protein nanoarrays is to use germanium pyramids as a support (Fig. 8.14). Calvo et al. (2002) report on the molecular wiring efficiency of glucose oxidase in organized self-assembled nanostructures. Wired protein molecules may be important for future nanotechnological tools (Figs. 8.15 and 8.16). Also the design of non-native macromolecular assemblies, e.g., hexameric helical barrels (Ghirlanda et al., 2002) is an endeavor with implications for nanotechnology (Fig. 8.17). Finally, Fig. 8.18 displays the manufacture of ordered inorganic nanocrystals on top of an array of genetically engineered viruses (Lee et al., 2002c).



**Fig. 8.12** Manufacture of protein nanoarrays for the investigation of molecular interaction and other recognition processes (Lee et al., 2002b; see also Hodneland et al., 2002)



**Fig. 8.13** Example of an engineered protein nanoarray (Lee et al., 2002b). Protein arrays with 100- to 350-nm features were fabricated with dip-pen nanolithography (see Fig. 8.12)



**Fig. 8.14** Protein nanoarray manufactured by self-assembly on nanometer-sized germanium pyramids (Riedel et al., 2001). Dynamic contact angle measurements with water droplets revealed that the germanium substrate is highly hydrophilic, and thus should be suitable for adsorption of hydrophilic proteins. Some protein inactivation was observed, however



**Fig. 8.15** Wired protein molecules might be components in future nanobiotechnological devices (see, e.g., Service, 2001; Calvo et al., 2002; Seeman and Belcher, 2002). Suitable wires are made, e.g., from chemical compounds, peptides, or carbon nanotubes. Proteins may act in such bioelectronic structures, e.g., as redox relays (Calvo et al., 2002)



**Fig. 8.16** Network of single-walled carbon nanotubes and genetically engineered proteins in a future nanobiotechnological device. Carbon nanotubes display an exceptionally low resistance and are seen as especially useful for the production of bridging nanowires and other nanostructures (see, e.g., Fagas et al., 2002; Odom et al., 2002)



**Fig. 8.17** Engineered hexameric helical barrels (Ghirlanda et al., 2002). A dimeric threehelix bundle was designed from first principles. In order to probe the requirements for stabilizing the hexamer, Ghirlanda et al. systematically varied polarity and steric bulk of the residues in the supercore of the hexamer. Formation of the hexameric assembly was best stabilized by changing three polar residues per three-helix bundle to hydrophobic residues (two phenylalanines and one tryptophan)



**Fig. 8.18** *Left:* ordering of inorganic nanocrystals using genetically engineered phages (Lee et al., 2002c). The bacteriophages form the basis of the self-ordering system. Genetic engineering enables the phages to specifically bind to nanocrystals. *Right:* structure of a phage

#### 8.5 Study and manipulation of protein crystal growth

The fabrication of properly diffracting protein and virus crystals is often the main obstacle for the high resolution of proteins using X-ray methods (see Sect. 4.1).



**Fig. 8.19** AFM image of the surface of a turnip yellow mosaic virus crystal (Malkin et al., 1995, 2002; Kuznetsov et al., 2000; McPherson et al., 2000, 2001). AFM investigation revealed the sources of crystal disorder and mechanisms of their formation

AFM is exquisitely useful for the study and manipulation of crystal growth (Fig. 8.19; Durbin and Carlson, 1992; Durbin et al., 1993; Malkin et al., 1995; McPherson et al., 2000; Mollica et al., 2001; Biscarini et al., 2002). Most protein and virus crystals grow, through a process of two-dimensional nucleation, by formation of new crystal layers (McPherson et al., 2000). Scratching the surface of a lysozyme crystal which was completely covered by an impurity stopping crystal growth resulted in resumption of crystal growth (McPherson et al., 2000).

# 8.6 Nanopipettes, molecular diodes, self-assembled nanotransistors, nanoparticle-mediated transfection and further biophysical nanotechnologies

Bone cells respond to stretching by an AFM tip with activation of stretch-activated ion channels (Charras and Horton, 2002). Dissecting bacterial surface layers with an AFM tip provided a better understanding of the high stability of this protective bacterial surface coat (Fig. 8.20; Scheuring et al., 2002). Micronized salbutamol particles stick to glass stronger than to polytetrafluoroethylene (Eve et al., 2002). For further nanobiotechnological innovations see Figs. 8.21–8.27 and Sect. 7.1.



**Fig. 8.20** Unzipping a double layer of lipids adsorbed to mica (not shown), with an AFM tip (Scheuring et al., 2002). Using the AFM stylus as a nanodissector, native bacterial surface layers were separated and their mechanical and protective properties against hostile environments examined



**Fig. 8.21** Conduction between nanoelectrodes through a single molecule (Reimers et al., 2002) or a few organic molecules forming a nanocrystal (Rinaldi et al., 2002). In some cases, rectifying behavior is observed, e.g., for deoxyguanosine nanocrystals (Rinaldi et al., 2002)



**Fig. 8.22** Programmable delivery of DNA through a nanopipette (Ying et al., 2002; Bruckbauer et al., 2002b). The conical geometry of the pipette causes most of the electrical potential drop to occur in the tip region. Pulsatile delivery of DNA molecules is achieved by controlling the applied voltage

The nanopipette in Fig. 8.22 was designed for controlled delivery of macromolecules into living cells (Ying et al., 2002). A voltage applied to the nanopipette moves the DNA molecules slowly towards the tip of the pipette. Since, because of the conical geometry of the nanopipette, most of the potential

drop occurs in the tip region, the DNA molecules are rapidly delivered to the cell once they have reached the tip region. In combination with single molecule detection, individual DNA molecules can be delivered to the living cell in a controlled manner.



(b) Nanoparticle-mediated transfection of cells

**Fig. 8.23** Gold nanoparticle-mediated transfection of cells (Sandhu et al., 2002). (a) Mixed monolayer protected gold cluster functionalized with quaternary ammonium chains binding to DNA. (b) Schematic of the transfection process



**Fig. 8.24** Electrospray deposition of dry proteins for the fabrication of microarrays and nanoarrays (Avseenko et al., 2001, 2002). These arrays can detect antibodies in plasma samples from mice immunized with the proteins used for the arrays



**Fig. 8.25** Continuous-flow preparation of 100-nm nanoparticles for drug delivery, protein delivery, and gene therapy (Prokop et al., 2001; Davda and Labhasetwar, 2002; Igartua et al., 2002; Konan et al., 2002; Haas and Lehr, 2002)



**Fig. 8.26** De-novo designed virus-mimicking particle for drug delivery, protein delivery, and gene therapy (Xu et al., 2002). In contrast to the protein envelope of most viruses, this particle has an envelope made from lipids

Fig. 8.25 illustrates a technique for the fast production of protein-polymer nanoparticles: A solution of polymer and protein is gently mixed with a salt solution. The salting-out effect causes the self-organization of nanoparticles that have different surface properties than the protein in the core (Prokop et al., 2001; Davda and Labhasetwar, 2002; Igartua et al., 2002; Konan et al., 2002; Haas and Lehr, 2002). In a similar way, also DNA can be camouflaged with lipids (Fig. 8.26; Xu et al., 2002). Some of these particles can cross the blood-brain barrier and are seen as promising candidates for future gene therapy and drug delivery.



**Fig. 8.27** Self-assembled monolayer organic field-effect transistor (Collet and Vuillaume, 1998, Collet et al., 2000; Fujita et al., 2003). The figure depicts an example of the principle of operation of a field-effect transistor: The source-drain conductivity is controlled through the gate which is electrically insulated from the device itself: the width of a conducting channel between source and drain is varied by adjusting the voltage between gate and base. (a) Open channel between source and drain: current can flow between source and drain. (b) The source-drain channel is closed by application of a voltage between gate and base: now the source is isolated from the drain

A field effect transistor was manufactured by using self-assembly of organic molecules (Collet and Vuillaume, 1998, Collet et al., 2000). For an example of the principle of operation of field-effect transistors see Fig. 8.27.

## 9 Proteomics: high throughput protein functional analysis

Currently, there is a major effort, on a genome-wide scale, to map protein-drug interactions and to discover drug targets (Sect. 9.1), to map protein-protein interactions (Sect. 9.2), to discover chemical activity of proteins (Sect. 9.3), and to resolve protein structures (Sect. 9.5). This effort, called proteomics, provides significant knowledge of the biology of organisms far beyond the level of sequence information (see, e.g., Adam et al., 2002b; Burbaum and Tobal; Edwards et al., 2000, 2002; Christendat et al., 2000; Figeys, 2002a, 2002c; Gallardo et al., 2002; Hubbard, 2002; Kersten et al., 2002; Koshland and Hamadani, 2002; Lin and Cornish, 2002; Liu et al., 2002; Morrison et al., 2002; Natsume et al., 2002; Yarmush and Jayaraman). The system-wide study of proteins and as well non-proteinaceous interaction partners largely employs protein microarray technology (see, e.g., MacBeath, 2002; Gera et al., 2002; Kukar et al., 2002; Talapatra et al., 2002) and bioinformatic methods (see, e.g., Bork, 2002).

Proteomics-based approaches for the study of organ-specific regulatory and signaling cascades are seen as a key for a better understanding and therapeutical management of diseases (e.g., Jäger et al., 2002). Proteomics has provided new vaccine candidate antigens (Klade, 2002; Nilsson, 2002; Vytvytska et al., 2002). The identification of individual proteins abnormally expressed in tumors may have an important relevance for making diagnosis, prognosis, and treatment (e.g., Celis et al., 2002; Dwek and Rawlings, 2002; Jain, 2002; Michener et al. 2002; Zheng et al., 2003). Proteomics analysis of the neurodegeneration in the brain of transgenic mice discovered 34 proteins with significantly changed intensity (Tilleman et al., 2002). A proteomics approach was used to identify the translation products of squid optic lobe synaptosomes (Jimenez et al., 2002). A central nervous system (CNS) proteome database derived from human tissues is expected to significantly accelerate the development of more specific diagnostic and prognostic disease markers as well as new selective therapeutics for CNS disorders (Rohlff and Southan, 2002). Proteomics provides an extremely powerful tool for the study of variations in protein expression between different ages and for the understanding the changes that occur in individuals as they become older (Cobon et al., 2002).

Innovations towards higher throughput and cost cutting include mass spectrometry advances (Sects. 9.1 and 9.2), DNA microchips (Sect. 9.1), protein microchips (Sect. 9.2), genetic hybrid systems (Sect. 9.2), and lab-on-a-chip technology (Sect. 9.4).

#### 9.1 Target discovery

Two-dimensional electrophoresis and mass spectrometry (Fig. 9.1) are widely used for the study of protein composition and protein changes in humans, animals, and plants. Important applications are (a) the identification of biomarkers specific for certain cell types, disease states, or aging processes, and (b) the study of protein composition changes as a response to drug treatment.

Also, high throughput microarray-based assays hold tremendous promise for the discovery of proteins connected with diseases (Fig. 9.2).



**Fig. 9.1** Discovery of proteins relevant to a certain disease by two-dimensional polyacrylamide gel electrophoresis and mass spectrometry (see, e.g., Edwards et al., 2000; Blomberg, 2002; Kersten et al., 2002; Man et al., 2002; Mo and Karger, 2002; Rohlff and Southan, 2002)



**Fig. 9.2** Discovery of proteins relevant to a certain disease, e.g., cancer markers, by detection of changes in the abundance of mRNA by means of cDNA (complementary DNA) microarray technology (supplied, e.g., by SuperArray, Inc., Bethesda, MD). cDNA chips with spot sizes of  $10-500 \mu m$  are commonly fabricated by high speed robotics or ink-jet printing on glass or nylon substrates. Every spot contains a different, 100-10,000 bases long, immobilized probe cDNA fragment which is complementary to targeted cDNA. The targeted, radioactively labeled cDNA is synthesized by reverse transcriptase from mRNA of the sample cells. Single stranded target cDNA is hybridized with complementary cDNA of the array and non-binding cDNA rinsed off with buffer. Detection of the pattern of radioactivity of the array then shows which mRNA was present in the sample cells, and thus which proteins were expressed

#### 9.2 Interaction proteomics

Analysis of several 100,000 protein-protein interactions using microarray technology (Fig. 9.3) and the yeast two-hybrid system (Fig. 9.4) has led to dozens of



(b) Measurement of protein-protein interactions with the spot of the array



(c) Manufacture of the protein microarray

**Fig. 9.3** (a) Manufacture of a spot of a protein microarray for the assay of protein-protein interactions (see, e.g., Grayhack and Phizicky, 2001; MicroSurfaces, Inc., Minneapolis, MN). For a better maintenance of structural integrity, protein immobilization is carried out on a matrix or layer of a chemical linkers that provide a native-like environment for embedded proteins. (b) Sample protein molecules (target molecules) interact with a probe molecule of a spot of the array. Non-binding sample proteins are simply washed off with buffer. Sample molecules are radioactively or fluorescence labeled so that binding of target protein molecules with probe protein molecules can be detected. (c) Manufacture of an array with 30–100,000 individual probe proteins immobilized on a single slide by chemical treatment of the surface of a quartz glass slide and ink-jet or contact printing of the protein spots



**Fig. 9.4** Discovery of protein-protein interactions by means of the yeast two-hybrid system (McCraith et al., 2000; Ito et al., 2002; Stagljar and Fields, 2002): Two vectors are constructed so that (i) one contains the code for a protein (protein A: e.g., a predicted open reading frame) followed by the code for a DNA-binding protein, and (ii) the other contains the code for a nother protein (protein B: e.g., another predicted open reading frame) followed by the code for a gene expression activator. Expression of both vectors in the cell yields a DNA-binding fusion protein and an expression-activating fusion protein. Attraction of both fusion proteins brings the expression activator into vicinity to the DNA. This leads to reporter gene activation. Diploids expressing the two-hybrid reporter gene in the host cells are then identified

novel findings of important intermolecular interaction (see, e.g., McCraith et al., 2000; Ito et al., 2002; Stagljar and Fields, 2002).

Figs. 9.5 and 9.6 present a mass-spectrometric method for the analysis of a large number of protein-protein and protein-drug interactions, respectively, without need of 2D-chromatography or electrophoresis. Possible ambiguities in the assignment of mass peaks may be resolved by the technique of ion fragmentation (see, e.g., Fig. 3.27). The discovery of new drug targets is highly important for developing new drugs (e.g., Pillutla et al., 2002; Whitelegge and le Coutre, 2002).



**Fig. 9.5** Analysis of millions of protein-protein interactions in one organism without 2Dchromatography or electrophoresis: Target proteins immobilized on the microarray interact simultaneously with all proteins expressed in a certain type of cells. After rinsing off the non-binding molecules, a high resolution mass spectrogram is recorded for each spot of the array. Each mass spectrogram shows a peak corresponding to the target protein and possibly further peaks corresponding to binding proteins. The binding proteins are then identified by their masses



**Fig. 9.6** Large-scale analysis of interactions of chemical compounds with proteins without 2D-chromatography or electrophoresis: The protein array is dipped into the mixture of chemicals. After some time, non-binding chemicals are washed away with buffer. The spots are then mass-spectrometrically analyzed. Since the masses of the involved proteins and chemicals were previously measured, binding chemicals can easily be identified



**Fig. 9.7** Direct determination of strong protein-protein interactions in a cell extract without need for the manufacture of arrays. First the mixture of monomeric proteins and protein-protein complexes is separated according to size by gel chromatography. Conditions for the chromatography are chosen in a way that strong complexes do not completely dissociate. Then mass spectrometry is performed for each chromatographic fraction which identifies the dimeric complexes and their two interacting macromolecules in the fraction: During ionization in the mass spectrometer, the complexes dissociate causing two peaks in the spectrum. These twin peaks are identified by their total mass which is roughly equal to the mass of the non-binding macromolecules. This method may analogously be applied to map out other strong macromolecular interactions

High resolution mass spectrometry even allows the simultaneous mapping-out of a large number of protein-protein and protein-drug interactions without use of microarrays (Fig. 9.7). A fast size exclusion chromatography separates the complicated mixture of interacting molecules into fractions. The parameters for chromatography were chosen such that strongly interacting molecule complexes remain together. Chromatographic fractions are then mass-spectrometrically analyzed. Strongly interacting dimers of molecules appear in the spectrogram as pairs with a total mass of about that of the monomers in the fraction.

#### 9.3 Chemical proteomics

Chemical proteomics assigns molecular and cellular functions to thousands of identified or predicted gene products. Assaying activities of large pools of constructed strains with subsequent deconvolution of active pools is an efficient method to discover new functions of genes (see, e.g., Fig. 9.8; Martzen et al., 1999; Grayhack and Phizicky, 2001; Adam et al., 2002a; Phizicky et al., 2002).



**Fig. 9.8** Example of chemical proteomics (Martzen et al., 1999; Grayhack and Phizicky, 2001; Phizicky et al., 2002). A library of 6144 yeast strains was constructed. Each strain expresses a unique yeast open reading frame (ORF) as a GST-ORF fusion (GST, glutathione S-transferase). Each 96 strains were pooled and biochemical activity of the pools was assayed. Active pools were deconvoluted using the library of strains to identify the GST-ORF responsible for activity. Several previously unknown biochemically active gene products were discovered

### 9.4 Lab-on-a-chip technology and mass-spectrometric array scanners

Protein and DNA microarrays are increasingly often processed with the lab-on-achip technology (Figs. 9.9 and 9.10): tiny channels etched into a glass slide, microswitches, micromixers, and other small devices act as small chemical factories.



**Fig. 9.9** Simplified example for lab-on-a-chip technology (see, e.g., Swedberg et al., 1996; Swedberg and Brennen, 2001; Cheng et al., 2002; Figeys 2002b, Laurell and Marko-Varga, 2002). Tiny channels are microfabricated or etched into the support. The top is then sealed with another plate (not shown). A single chip may contain all the channels, switches and reservoirs necessary for complicated multi-stage chemical reactions



Fig. 9.10 Example of lab-on-a-chip technology. Channels are etched into the glass support

Scanning of protein and DNA arrays with fluorescence detectors usually requires special labeling of the sample and may be prone to errors due to limitations of sensitivity and due to unspecific binding. More importantly, mixtures of signals can often not be resolved. Automatic mass-spectrometric



**Fig. 9.11** Mass-spectrometric array scanner for automatic mass-spectrometric measurement of protein and DNA microarrays. Step motor controlled translation stages rapidly move the microarray into position. A mass spectrum for each of the spots is automatically taken and analyzed. The scanner can be used, e.g., in the methods outlined in Figs. 9.5 and 9.6

detection of the molecules in the spots of the array can greatly enhance the information yield (Fig. 9.11).

#### 9.5 Structural proteomics

Proteomics is driving a substantial effort towards large-scale protein structure prediction (see, e.g., Renfrey and Featherstone, 2002; Schmid, 2002) and determination. High resolution structure determination still relies on X-ray crystallography and NMR (nuclear magnetic resonance). Since both methods are expensive and time-consuming, further optimization of the methods is being in progress. NMR peaks can now automatically be assigned to the corresponding amino acid residues (Nilges et al., 1997; Heinemann et al., 2001). The Berlin Protein Structure Factory develops and applies large-scale NMR and crystallographic methods (Heinemann et al., 2000, 2001; Boettner et al., 2002).

On the other hand, structure computer simulation using simplifications of the conformational space of proteins is rapidly progressing (see Sect. 1.4), and so it can be hoped that comparably inexpensive computational methods will make an increasing contribution to structural proteomics in the near future.
# 10 Ion mobility spectrometry

## 10.1 General design of spectrometers

Ion mobility spectrometry was developed for the simple and cheap detection and characterization of organic compounds (Cohen and Karasek, 1970; Karasek, 1970; Caroll et al., 1971; Caroll, 1972; Cohen et al., 1972; Cohen and Crowe, 1973; Vora et al., 1987; St. Louis and Hill, 1990; Campbell et al., 1991; Burke, 1992; Eiceman and Karas, 1994; Taylor, 1996; Baumbach and Stach, 1998; Baumbach and Eiceman, 1999; Saurina and Hernandez-Cassou, 1999; Asbury and Hill, 2000; Purves et al., 2000; Wu et al., 2000; Beegle et al., 2001; Eiceman et al., 2001; Matz and Hill, 2001; Stone et al., 2001). The ion mobility spectrum reflects the ion mobilities which correlate well with the size-to-charge ratios of the sample compounds.

In the ion mobility spectrometer (IMS), (a) sample molecules in the vapor phase are ionized, (b) the charged sample molecules (ions) are accelerated by an electric field, and (c) their time of flight in the gaseous medium of the drift channel is measured and recorded (Figs. 10.1–10.6). These simple spectrometers can detect and analyze astonishing tiny traces of small and as well large molecules and clusters.



**Fig. 10.1** Principle of operation of ion mobility spectrometers: Molecular ions are generated and accumulated in the reaction area. A gating pulse transfers the molecular ions to the drift channel where said ions are accelerated by an electric field. Ions with different mobilities in the gaseous medium of the drift channel arrive at different times at the detector



**Fig. 10.2** Design of a sample inlet for an IMS. The sampling pump draws air through the semi-permeable membrane which attenuates the influx of large dust particles and other interferents (see also Spangler, 1982). A suitable membrane is, e.g., a  $5-50 \mu m$  polytetrafluoroethylene (PTFE) foil or silicone rubber membrane (Spangler and Carrico, 1983; Kotiaho et al., 1995). For IMS for detection of biological agents, a metal grid instead of a membrane may be more appropriate because of its better transparency for high molecular-weight compounds



(b) Negative ion mode

**Fig. 10.3** Reactions of the sample in the reaction area. Sample molecules are ionized, either directly by dissociation or indirectly by clustering with other ions. In the positive ion mode, positive ions are repelled from the repeller plate and accumulated in front of the shutter grid. In the negative ion mode, negative ions are analogously accumulated. Many IMS operate alternatingly in the positive and negative ion mode



Fig. 10.4 Principle of operation of an ion mobility spectrometer (IMS) (Cohen and Karasek, 1970; Karasek, 1970; Caroll et al., 1971; Keller, 1975; Eiceman and Karas, 1994). Sample molecules are injected into the reaction area (ionization chamber) and ionized. A thin membrane or determines the identity of the sample molecules by matching the spectra to reference signatures. In order to reduce the noise, the IMS is noise to the signal of the Faraday plate, the voltage supply for the guard rings is highly stabilized. In contrast to mass spectrometers, this device needs no hot filaments with a limited lifetime, electron multipliers, energy-consuming vacuum pumps, or expensive vacuum tubes, and its sensitivity can be several orders of magnitude higher than that of a mass spectrometer. Drift times of macromolecules are usually milliseconds a wire mesh grid separates reaction area and drift channel (drift tube, drift region). An electrical pulse applied to the shutter grid (gate, grid, gating grid) transfers the ions into the drift channel where the ions are further accelerated by the electric field which is generated by guard rings see Figs. 10.5 and 10.6). The time of flight of the ions in the gaseous phase is measured with the help of a Faraday plate (collector plate, Faraday cup) or collector grid. The mobility of a gas phase ion is a measure of its collision cross section which in turn depends on its size and ons have different mobilities in the gas of the drift tube, they can result in distinct peaks in the ion mobility spectrum. A computer (not shown) enclosed in a grounded copper foil (not shown). The collector is directly connected with a  $10^{10}$ -V/A preamplifier via a cable of only a few mm ength. The feedback resistor of the preamplifier is selected for a low noise level. Since fluctuations of the electric field of the drift channel add o seconds in 3-20 cm drift tubes with drift voltage gradients of about 100-1000 V/cm. For most IMS, a few seconds are required after each structure. Small ions with compact structures have smaller cross sections than large ions with open structures. Consequently, since different measurement to purge the drift channel



Fig. 10.5 Design of an IMS (Cohen and Karasek, 1970; Karasek, 1970; Caroll et al., 1971; Eiceman and Karas, 1994; Matz and Schröder, 1996, 1997; IUT Institute for Environmental Technologies, Berlin, Germany; Bruker Daltonik, Bremen, Germany; Graseby Dynamics, London, U.K.; Barringer Instruments, Warren, NJ). The ionization source ionizes sample molecules. Guard rings generate an electric field which accelerates the sample ions towards the shutter grid where they accumulate. After a few seconds, a voltage pulse is applied to the shutter grid causing the release of the sample ions into the drift channel. Now the electric field can further move the sample ions towards the ion detector (Faraday plate). Different ions interact differently with the drift gas molecules in the drift channel. This causes the ions to spread out according to their different mobilities. The recorded ion mobility spectrum corresponds to differences in the time of flight of the sample ions. Typically, the drift channel has a length of a few cm, and the electric field strength of in the drift channel is 100-1000 V cm<sup>-1</sup>. At this field strength and length, small organic compounds which have mobilities of a few cm<sup>2</sup> V s<sup>-1</sup> need about 1-100 milliseconds to reach the detector. Biomacromolecules typically travel about 1-3 orders of magnitude slower. Drift channel and reaction area are enclosed in a thermal isolation with heating elements. The operating temperature for the detection of biological agents is typically 100-150 °C

The IMS is comprised of (a) a thermally isolating housing, a source of ionization, reaction area, shutter grid, drift channel with guard rings, possibly an aperture grid, collector, (b) a source of clean gas or a gas filter, (c) a shutter controller, (d) a high voltage supply, (e) an electrometer, (f) temperature control instrumentation, (g) a computer-aided data collection and processing unit.

The major advantage of IMS is the extreme sensitivity (see Sect. 10.2). Another important advantage of IMS, which do not need vacuum parts, is the lower cost and lower energy consumption relative to most mass spectrometers. This makes it particularly suitable for large-scale field applications.



**Fig. 10.6** *Top:* design of an IMS containing 18 stacked copper guard rings that are separated by insulating ceramics spacers and connected by resistors. These resistors are selected for low noise. The voltage supply for the guard rings is stabilized to better than 0.1% rms. In order to ensure a homogenous electric field, the guard rings have sufficiently narrow separations. In this example the IMS is operated in the positive ion mode. *Bottom left:* example of the shutter voltage: Most of the time, a positive shutter voltage prevents positively charged ions from entering the drift channel. After a period of time during which positive ions accumulate in front of the shutter grid, a negative shutter voltage pulse is applied. Now positive ions can enter the drift channel where they are further accelerated. *Bottom right:* recorded spectrum

A suitable inert gas for the detection of many organic compounds is nitrogen, but clean air can also serve for this purpose. Cleaning the drift gas from water vapor and sample residues is frequently performed with molecular sieves (Fig. 10.7). These molecular sieves are made from zeolites which are certain alumino-silicates. Zeolites can adsorb water molecules and other small organic compounds. After a few weeks or month of use in an IMS, the zeolite is saturated. It can be reconstituted by heating it in an oven and be re-used.



**Fig. 10.7** Drying the inert gas with a molecular sieve (Carnahan and Tarassov, 1998; Taylor and Turner, 1999). Suitable dimensions for a length of the drift channel of 5-20 cm, a diameter of the drift channel of 2 - 3 cm, and a height of the ionization chamber of 0.5-1 cm are: flow rate of pump 1: 5-50 ml min<sup>-1</sup>; flow rate of pump 2: 50-500 ml min<sup>-1</sup>

## 10.2 Resolution and sensitivity

The resolution, *R*, of an IMS is defined as

$$R = 0.5 \cdot t_{\rm d} \, \tau^{-1} \,, \tag{10.1}$$

where  $t_d$  is the drift time of the peak,  $\tau$  and is its temporal width at half peak height (St. Louis and Hill, 1990).

Important factors affecting the resolution are: (a) initial ion pulse width and shape (shutter pulse width), (b) broadening by Coulomb repulsion between the ions in both the reaction region and drift channel, (c) ion-molecule and ion-ion reactions in the reaction region, (d) gate depletion, (e) ion-molecule reactions in the drift region, (f) spatial broadening by diffusion of the ion packet during the drift, (g) temperature and pressure inhomogeneities within the spectrometer, (h) capacitive coupling between aperture grid and collector, and (i) the response time of the preamplifier, amplifier, and analog-to-digital converter. Broadening by Coulomb repulsion is particularly severe in narrow designs of reaction region and drift channel.

*R* strongly depends on the design of the spectrometer, but can theoretically exceed 1000 (Figs. 10.8-10.10). A high resolution requires a relatively bulky design with a drift channel of sufficient diameter and large length and a high drift voltage. Most commercial IMS have resolutions of only 20-150 since they are mainly optimized for low weight and portability.



**Fig. 10.8** Theoretical resolution of an IMS at different drift lengths. At short drift lengths, increasing the drift voltage above an optimum does not further improve the resolution. The parameters in this example are: temperature, 130 °C; ion mobility,  $10^{-5}$  m<sup>2</sup> V<sup>-1</sup> s<sup>-1</sup>; initial ion pulse width, 1 ms; shutter pulse voltage, 1000 V; length of the reaction area, 3 mm; distance between aperture grid and Faraday plate, 1 mm; voltage at the aperture grid, 500 V



**Fig. 10.9** Theoretical resolution of an IMS at different drift lengths and ion mobilities: long shutter pulses enhance the sensitivity, but limit the resolution for small ions. The parameters in this example are: temperature, 130 °C; drift voltage, 80 kV; initial ion pulse width, 1 ms; shutter pulse voltage, 1000 V; length of the reaction area, 3 mm; distance between aperture grid and Faraday plate, 1 mm; voltage at the aperture grid, 500 V

*R* is approximately given by Eq. 10.2 (St. Louis and Hill, 1990; Hill et al., 1990; Leonhardt et al., 2001):

$$R = \frac{1}{\sqrt{16\ln 2\frac{k_BT}{qV_d} + \frac{K^2 V_d^2}{l_d^4} \left(t_{Palse} - \frac{s^2}{KV_{Palse}}\right)^2 + \left(\frac{a^2 V_d}{l_d^2 V_{ap}}\right)^2 + \frac{t_{Palse}^2 K^2 V_d^2}{l_d^4}}}{I_d^4}}$$
(10.2)

where  $k_{\rm B}$ , Boltzmann constant (1.3807 × 10<sup>-23</sup> J K<sup>-1</sup>); *T*, absolute temperature; *q*, ion charge;  $V_{\rm d}$ , drift voltage;  $l_{\rm d}$ , drift length; *K*, ion mobility;  $t_{\rm Pulse}$ , initial ion pulse width;  $V_{\rm Pulse}$ , shutter pulse voltage; *s*, distance between space charge in the reaction area and shutter grid (roughly half the length of the reaction area); *a*, distance between aperture grid and Faraday plate;  $V_{\rm ap}$ , voltage at the aperture grid.



Fig. 10.10 Theoretical resolution of an IMS at different drift lengths and ion mobilities. Short shutter pulses and long drift lengths allow the high resolution of both medium-sized and as well large molecules. The parameters in this example are: temperature, 130 °C; drift voltage, 80 kV; initial ion pulse width, 100  $\mu$ s; shutter pulse voltage, 1000 V; length of the reaction area, 3 mm; distance between aperture grid and Faraday plate, 1 mm; voltage at the aperture grid, 500 V

The sensitivity of an IMS can be 1000 times grater than that of a good mass spectrometer: with conventional electronics, theoretically down to 1000 ions are still detectable corresponding to  $3 \times 10^{-19}$  g TNT or  $2.5 \times 10^{-16}$  g botulinum toxin ( $M_w = 150$  kDa). For comparison, the fingerprint of a person likely contains many billion times more residue of organic compounds. Practical detection limits are

typically on the order of 0.1 to 100 ppbv. One reason for the often superior sensitivity relative to mass spectrometers is the simple and large sample inlet. Many IMS contain a thin large area-polymer membrane inlet or a meal grid inlet with cause only little sample losses (Fig. 10.3; Spangler, 1982; Spangler and Carrico, 1983; Kotiaho et al., 1995). In some IMS, the sample inlet solely consists of a tube with a filter which removes dust particles and in some cases water from the sample prior to sample injection into the IMS.

Short gating pulses and high humidity tend to lower the sensitivity. All means of noise reduction, such as (a) small collector capacity, (b) a highly stabilized voltage of the guard rings, (c) an electric shielding of the whole IMS, and (d) low-noise resistors in the electronics, tend to improve the sensitivity. Further, for a high sensitivity and low memory effects, the IMS must be built from materials which adsorb extremely little if any sample molecules, e.g., special ceramics.

Sensitivity and resolution are also affected by the presence or absence of an aperture grid: The function of the aperture grid is to capacitively decouple the collector (Faraday plate or collector grid) from the approaching ion cloud. Without the aperture grid, the collector senses the approaching ion cloud several millimeter prior to its arrival, resulting in line broadening (St. Louis and Hill, 1990). On the other hand, the aperture grid neutralizes ions which decreases the ion current and sensitivity by a factor of roughly 3.

#### 10.3 IMS-based "sniffers"

IMS sniffers (Figs. 10.11 and 10.12) are hand-carried IMS-based devices, mainly manufactured for the detection of dangerous substances. They contain a small IMS and a database of the signatures of substances of interests and interferents. After a few seconds of measurement, the processing unit identifies and signalizes detected agents. The sniffers weighting about 2-10 kg are roughly 25-40 cm long. False alarm rates for the detection of chemicals are quite often 0.01-1%.



**Fig. 10.11** Hand-carried "sniffers" comprising a chemical preconcentrator and an IMS (made, e.g., by Sandia Corporation, Albuquerque, NM, and Barringer Instruments, Inc., Warren, NJ)



**Fig. 10.12** Hand-carried sniffling detector which further comprises a fast gas chromatography (see, e.g., Snyder et al., 1993). The combination with GC improves the false alarm rate for the detection of dangerous substances by typically one order of magnitude

## 10.4 Design details

Figs. 10.11 and 10.12 show a portable IMS and GC/IMS detector with a preconcentrator, respectively. GC in combination with IMS significantly reduces the false alarm rate for the detection of hazardous compounds. Preconcentrators (Fig. 10.13) can increase the sample concentration and the sensitivity of the method by a factor of more than 1000, and can also reduce the false alarm rate.



**Fig. 10.13** Two design variants of preconcentrators for ion mobility spectrometry. It draws a large volume of air and collects biological and heavy chemical organic compounds from the air onto the filter. The filter is made from zeolites – a material which is commonly used in molecular sieves (see Fig. 10.7). After several minutes of sample collection, the heater vaporizes the organic material into a small parcel of air which is delivered to the IMS. Theses preconcentrators increase the sample concentration by a factor of typically 10-1000 (see, e.g., Spangler, 1992a)



interaction with gel molecules; typically optical detection

**Fig. 10.14** The typical set-up for electrophoresis shows an analogy to jon mobility spectrometry: in both methods charged molecules are accelerated by an electric field and slowed down by interaction with molecules of a stationary phase. In electrophoresis, the stationary phase is generally a gel, and the movement of the sample molecules is often optically detected. A gas serves as stationary phase in ion mobility spectrometry, and the movement of the sample ions is electrically detected. A small size and high charge of ions correlates with a large speed

Ion mobility spectrometry has some similarities to electrophoresis (Fig. 10.14). Because of the similarities to common chromatography, originally ion mobility spectrometry was called "plasma chromatography". However, one has to keep in mind that in contrast to common chromatography, ion mobility spectrometers have a narrow linear range due to space charge effects (Bird and Keller, 1976; Blanchard and Bacon, 1989; Spangler, 1992b), and show serious matrix interferences and prolonged memory effects.

Table 10.1 Common methods of sample ionization in ion mobility spectrometry (Lubman and Kronick, 1982, 1983; Baim et al., 1983; Leasure et al., 1986; Eiceman et al., 1988; Shumate and Hill, 1989; Begley et al., 1991; Phillips and Gormally, 1992; Davies, 1994; Spangler et al., 1994; Carnahan and Tarassov, 1995; Leonhardt, 1996; Lee et al., 1998; Wu et al., 1998a, 1998b, 2000; Budovich et al., 1999; Döring et al., 1999; Borsdorf et al., 2000; Megerle and Cohn, 2000; Schnurpfeil and Klepel, 2000; Borsdorf and Rudolph, 2001)

| Method of ionization          | Example   |
|-------------------------------|---|
| Radioactive isotopes          | <sup>3</sup> H, <sup>241</sup> Am foil, or <sup>63</sup> Ni foil; Fig. 10.15  |
| Photoionization               | UV and VUV light from a 30-W krypton or hydrogen<br>lamp with a $MgF_2$ -window, or perpendicular to the drift<br>channel from a frequency-quadrupled Nd:YAG laser at<br>266 nm (Fig. 10.15). A VUV-absorbing compound may<br>be added for an increased degree of ionization. |
| Electrospray                  | Fig. 3.11 in Chap. 3  |
| Laser desorption              | Fig. 10.15  |
| Electrical (corona) discharge | Fig. 10.15  |



(a) Laser desorption and ionization of a solid sample (b) Ionization by an electrical discharge



Fig. 10.15 Examples of ionization methods (see also Table 10.1)

Since most of the interesting chemical and biological substances are not charged, it is necessary to ionize them prior to the drift. Table 10.1 and Figs. 10.15 and 10.16 show methods for ionization in ion mobility spectrometry. Radioactive isotopes and photoionization are the most common methods. When using a <sup>63</sup>Ni foil as the source of ionization and air as the drift gas, the primary ions are mainly short-living N<sub>2</sub><sup>+</sup>, NO<sup>+</sup>, and O<sub>2</sub><sup>-</sup>. These primary ions rapidly react with traces of water in the drift gas to form clusters of the types N<sub>2</sub><sup>+</sup>(H<sub>2</sub>O)<sub>n</sub>, NO<sup>+</sup>(H<sub>2</sub>O)<sub>m</sub>, and O<sub>2</sub><sup>-</sup>(H<sub>2</sub>O)<sub>k</sub>. Photoionization with hydrogen plasma discharge lamps and krypton plasma discharge lamps requires a photon flux of about 10<sup>12</sup> cm<sup>-2</sup> s<sup>-1</sup>. The geometry of the shutter grid is chosen so that most photons cannot enter the drift channel since this would reduce the resolution. Common radioactive sources in IMS have the advantage of relatively long half-lives of several years. For immobilization of <sup>3</sup>H, it is gettered in a thin titanium layer.

Ion mobility spectrometry has gained significant importance in the context of the detection of ultra-trace chemical and biological contaminants (Snyder et al., 1991a, 1991b, 1996a, 1996b, 1999, 2000; Ogden and Strachan, 1993; Strachan et

al., 1995; Dworzanski et al., 1997; Smith et al., 1997), explosives (e.g., Fetterolf and Clark, 1993; Steinfeld and Wormhoudt, 1998; Fig. 10.17), illicit drugs (e.g., Miki et al., 1997, 1998; Keller et al., 1998), pesticides, the detection of animals and animal activity in jungles, and other environmental monitoring.



Fig. 10.16 Gas inlet for electrical-discharge (corona discharge) ionization



Fig. 10.17 Ion mobility spectrogram of TNT (trinitrotoluene). From data supplied by the Institute for Environmental Technologies Ltd., Berlin



Fig. 10.18 A high-resolution two-channel ion mobility spectrometer. Here the sample is simultaneously analyzed in two different ways reducing false identifications of agents. In one of the columns, the sample is chemically and/or physically modified by a chemical addition

Fig. 10.18 outlines an IMS-based twochannel detector which was designed within a feasibility study. In this design the differentiation between various

biological and also some chemical substances is improved by utilizing physical and chemical modifications of the sample in one of the two columns, e.g., by adding an acidizing gas: Most proteins display a strong pH-dependency of the charge (see, e.g., Chap. 2 in Nölting, 2005). Thus, by changing the pH, the charge state of the protein-containing biological material is altered and consequently its speed of diffusion in the drift channel is changed. Further, the added gas can cause chemical changes of some chemical and biological agents. This causes specific changes of the IMS spectra and thus contributes to a further improvement of the correct identification of the agents. Multichannel designs are a further option to reduce false detection rates of IMS. The channels may be operated in the same way speeding up the measurement of slowly drifting substances. Alternatively the sample may simultaneously be distributed over different channels which are operated in various different modes which can improve the resolution of the method.

IMS with an oscillating electric field allow the application of large field strengths without need for a very high voltage (Fig. 10.19). Since very large macromolecules have low mobilities, their fast detection with high resolution requires a high electric field strength in the drift channel. In the common design this may cause safety problems and increases the price of the IMS.



**Fig. 10.19** IMS with an oscillating electric field of the guard rings instead of a constant electrostatic field. Only ions of which the movement is in phase with the oscillating electric field can pass through the drift channel and reach the collector. For further details on frequency-domain IMS see, e.g., Martin et al., 1998



**Fig. 10.20** Modified IMS for the detection of chemicals contained in solid samples and for the identification of different solid materials, e.g., wood (Lawrence et al., 1991; Matz and Schröder, 1997; Schröder et al., 1998)



**Fig. 10.21** Noise reduction with a smaller collector grid: The collector and the guard ring in front of the collector act together as an electrostatic lens. The smaller collector causes less noise and, thus, a higher sensitivity

Measurements on solid samples require special sample inlets with a heating and a temperature-resistant porous membrane (see, e.g., Fig. 10.20) and possibly a higher temperature of the IMS.

An electrostatic lens focusing the ions towards the collector can allow the reduction of the collector size and capacity (Fig. 10.21). This decreases noise and can improve the sensitivity.



**Fig. 10.22** Generating the high voltage for an IMS (Goebel and Breit, 2000). (a) A set of n capacitors is charged with the voltage V. (b) The capacitors are connected together in series for generating the high voltage  $n \cdot V$ 



Fig. 10.23 Improved mechanical stability of the drift channel. For a similar design see also (Karl, 1994)

A technique of high voltage generation which does not necessarily require much weight is illustrated in Fig. 10.22: a set of capacitors is charged in parallel and then connected in series.

Parameters which are important for a high reproducibility of IMS spectra are a constant degree of humidity, constant electric field strength, constant source of ionization, efficient removal of previous samples, mechanical stability. The latter can be improved by a special shape of the guard rings (Fig. 10.23). Figs. 10.24–10.26 display some further important innovations in ion mobility spectrometry.



**Fig. 10.24** Improved purgeability of the reaction area (see, e.g., Snyder et al., 1993) and improved homogeneity of the electric field the ions experiencing in the drift channel: the sample outlet is located close to the shutter grid, and the diameter of the reaction area is smaller than that of the drift channel



Fig. 10.25 Injection of the output of a gas chromatograph or of the vapor from a solid sample into the IMS with the help of a gentle stream of carrier gas



**Fig. 10.26** A multichannel ion mobility spectrometer. For an 8 times higher sampling rate than a single-channel spectrometer, all channels are operated in the same way. For decreased rate of false identifications, one sample may be distributed over different channels that are operated in different modes (see, e.g., Turner, 1993)

## 10.5 Detection of biological agents

Unfortunately, many biological agents are too large to be detected directly: The velocity, v, of a large spherical particle depends on its charge, z, its radius, r, the



**Fig. 10.27** Setup for IMS-detection of biological agents (see, e.g., Snyder et al., 2000). The virtual impactor selects a certain size range of particles, e.g.,  $1-10 \mu m$ , and transfers the selected particles into the pyrolysis tube. Within a few seconds, particles of biological origin are partially decomposed in the pyrolysis reaction at, e.g.,  $350 \,^{\circ}$ C. In the subsequent analysis of the pyrolysis reaction products, a short gas chromatography and an IMS are combined for enhanced resolution. The operating temperature of the GC/IMS is typically  $80-150 \,^{\circ}$ C. For the principle of operation of the virtual impactor see also Fig. 3.20. Due to the highly dispersed 2D-spectra, Py-GC/IMS can potentially much safer unambiguously identify traces of biological agents than a measurement of particle size distribution alone



**Fig. 10.28** Example of a particle of a dangerous biological contaminant. Light and fluffy composites of bacterial spores or viruses to dust-forming particles of about  $1-5 \ \mu m$  diameter can drift in dry air for 100 miles, and can be sucked into the deepest sacs of the lung (Preston, 1998)



Fig. 10.29 Typical sensitivity of IMS detection of compounds with different boiling points and vapor pressures

electric field strength, *E*, and the viscosity of the medium,  $\eta$ :

$$v = zE \ (6\pi\eta r)^{-1} \tag{10.3}$$

Small chemical compounds typically travel in an IMS with several m s<sup>-1</sup>, but a single-charged dust particle ( $z = 1.6 \times 10^{-19}$ C) with a radius of 1 µm travels in air ( $\eta_{air} = 1.8 \times 10^{-5}$  N s m<sup>-2</sup>) at a field strength of E = 300 V mm<sup>-1</sup> only with about 0.14 mm s<sup>-1</sup>. At higher gating frequencies of the IMS, successive spectra of slowly moving agents would superimpose. Also, just average size and size distri-

bution of biological agents offers little information about the precise nature of the agent. That is why biological agents are pyrolyzed prior to analysis in the IMS (Fig. 10.27). Pyrolysis (see also Sect. 3.2) decomposes and vaporizes biological agents and can be applied on bacteria and viruses (Fig. 10.28). The low vapor pressure of most biological compounds requires an operation of the IMS at a sufficiently high temperature (Fig. 10.29). The set-up virtual impactor / pyrolyzer / GC / IMS (Fig. 10.27) is capable to detect a few bacterial spores in a volume of several 1000 liters.

## 11 $\Phi$ -Value analysis

In Chap. 1 some inter-residue contact maps of protein transition states were presented. Here, the method of  $\Phi$ -value analysis underlying such maps and some of its high-resolution applications are presented in more detail: the correlation of inter-residue contacts with  $\Phi$ -values (see Fig. 1.9) is the currently available method with the highest resolution for protein folding transition states (Nölting, 1998, 1999a, b; Nölting and Andert, 2000).

The transition state corresponds to the state with the highest free energy in the course of the reaction. Since it is only extremely short-living, at present its structural resolution can not be carried out with NMR or X-ray crystallographic analysis.  $\Phi$ -Value analysis uses mutants as structural reporters and a combination of equilibrium thermodynamics and kinetics methods (Nölting, 2005).

#### 11.1 The method

Fig. 11.1 shows the free energy changes in the folding reaction for a wild-type protein and a mutant of this protein. One can see that in the course of the reaction



**Fig. 11.1** Energy landscape along the reaction coordinate for the folding reaction of a wild-type protein (top curve) and a mutant of this protein (bottom curve)



**Fig. 11.2** Built-up of an energy difference,  $\Delta\Delta G$ , between wild-type protein and a mutant in the course of the folding reaction.  $\Delta\Delta G_{F-U}$  is the energy difference between wild-type and mutant protein in the folded state

an energy difference,  $\Delta\Delta G$ , builds up between wild-type and mutant protein. This build-up of  $\Delta\Delta G$  corresponds to the build-up of structure in the molecule. In particular, the time point in the reaction at which  $\Delta\Delta G$  becomes significant depends on the time at which the interactions probed by the mutation build up in the molecule: If the interactions altered by the mutagenesis form early in the folding reaction (left curve in Fig. 11.2), one usually observes an early increase of  $|\Delta\Delta G|$ . In contrast, if the interactions probed by mutagenesis are formed late in the folding reaction, there is usually no significant  $\Delta\Delta G$  till late in the reaction (right curve in Fig. 11.2). So, by measuring  $\Delta\Delta G$  at the different stages of the folding reaction one can find out when certain interactions in the molecule are becoming formed. For the methods of measurement of  $\Delta\Delta G$  see Nölting (2005).

The formation of stable interactions in the molecule is usually expressed by the  $\Phi$ -value which is a measure of the structure consolidation at the position of the mutation on a scale from 0 to 1.  $\Phi$  is defined as  $\Phi = \Delta \Delta G / \Delta \Delta G_{F-U}$ , where  $\Delta \Delta G_{F-U}$  is the  $\Delta \Delta G$  in the folded state (Nölting, 2005). A  $\Phi$  of 0 at a certain stage of the folding reaction suggest the absence of stable structure at the position of the mutation at this time. If structure is completely formed at the position of the mutation at this stage of the reaction, one would expect a  $\Phi$ -value of 1. Possible sources of error in this analysis, e.g., the effect of non-native interactions, can be decreased by using several mutants for the same part of the molecule.

So, in order to obtain information on the structure of a transition state one simply needs to measure  $\Phi$  of the transition state for many mutants and correlate the data with the inter-residue contacts in the molecule (Nölting, 1998, 1999a, b).

## 11.2 High resolution of six protein folding transition states

This section presents the structural characteristics of the main transition states of six proteins obtained by correlation of inter-residue contacts with  $\Phi$ -values (see also Sect. 11.1; Chap. 1; Nölting and Andert, 2000). The first four proteins, barstar, barnase, chymotrypsin inhibitor 2 (CI2), and src SH3 domain are monomeric in the native state. Arc repressor is dimeric, and p53 is a tetramer with the structure of a dimer of dimers. In the first five main transition states (Figs. 11.3-11.7) one can see a very non-uniform consolidation of structure. It has been shown that for all five proteins the most consolidated clusters (highlighted as ribbons in Figs. 11.3-11.7; unconsolidated structure is displayed as wires) contain a relatively higher content of residues which belong to secondary structure elements than the non-consolidated parts of the molecules. On the other hand, all six transition states with the exception of the src SH3 domain (Figs. 11.3–11.5, 11.7, 11.8) contain on average a similar content of secondary and tertiary structure interactions (Nölting and Andert, 2000). This high resolution of folding transition states led to an understanding of the mechanism of protein folding and of its astonishing efficiency: folding of many proteins proceeds similar as the growth of a crystal - largely driven by the propensity of secondary structure formation, but also by the hydrophobic effect and other forces, a folding nucleus forms early in the folding reaction. This nucleus then restricts the number of possible conformations and enables further structure growth around it (Nölting and Andert, 2000; Nölting et al., 2003; Nölting, 2005).



Fig. 11.3 Main transition state structure of barstar (Nölting and Andert, 2000). Consolidated structure is highlighted as ribbons; unconsolidated parts of the molecule are shown as wires. Amino acid residues with high  $\Phi$ -values are highlighted as spheres. This figure and the following figures in this section were prepared using MOLMOL, Koradi et al., 1996



**Fig. 11.4** Main transition state structure of barnase (Nölting and Andert, 2000). Consolidated structure is highlighted as ribbons. For further explanation see Fig. 11.3



**Fig. 11.5** Transition state structure of CI2 (Nölting and Andert, 2000). Consolidated structure is highlighted as ribbons. For further explanation see Fig. 11.3



**Fig. 11.6** Transition state structure of the src SH3 domain (Nölting and Andert, 2000). Consolidated structure is highlighted as ribbons. For further explanation see Fig. 11.3



**Fig. 11.7** Transition state structure of Arc repressor (Nölting and Andert, 2000). Consolidated structure is highlighted as ribbons. For further explanation see Fig. 11.3



Fig. 11.8 Main transition state structure of p53 (Nölting and Andert, 2000). In this transition state essentially all parts of the molecule are highly consolidated. Here the residues with low  $\Phi$ -values are highlighted as small spheres

# 12 Evolutionary computer programming

## 12.1 Reasons for the necessity of self-evolving computer programs

Nature offers a tremendous amount of extremely complicated problems which cannot easily be rationalized and resolved. Probably one has to accept that there are scientific and technological problems too complex to be directly rationalized by humans. A well-known example is the non-periodical movement of many gravitationally interacting bodies in space. Since we are not able to imagine their motions with a sufficient degree of perfection, we call it "chaos". Unfortunately, humans obviously have significant intellectual difficulties to find theoretical descriptions or models for phenomena which they do not comprehend. Similarly, as an ape cannot write a mathematical equation beyond its intellectual abilities, humans are not directly able to establish mathematical structures beyond their intellectual limits. However, further progression of science and technology urgently requires to overcome such limits.

One well-known example for a complicated problem is the so-called folding paradox (see, e.g., Nölting, 2005): how can a protein find its unique native conformation among the  $\sim 10^{30}-10^{200}$  possible conformations of an average small protein in the unfolded state? One of the major difficulties of complex phenomena like protein folding is often the lack of an efficient and solvable mathematical description. Often one is able to write down some equations which describe the physics of the system while not being able to solve them.

This chapter describes a method which can potentially provide solutions beyond current human intelligence: in order to overcome the limits of rational design, one lets a computer program evolve itself. The method is exemplarily applied on protein folding and structure predictions (Nölting et al., 2004) and the optimization of optical effects of nanoparticle arrays. In Sect. 12.5 the much wider scope of this method is discussed.

## 12.2 General features of the method

Figs. 12.1 and 12.3 show the principle of operation of the method of self-evolving programs. A so-called wild-type computer program is evolved towards higher efficiency by mutagenesis and selection in a similar way as species evolve in

nature: First a number of mutants of the wild-type program is created. The performance of the mutants is then tested and the best performing program serves



**Fig. 12.1** Scheme of the method of evolutionary computer programming. The best performing program at a certain stage of the evolution is shown as a filled rectangle. Open rectangles indicate less-performing mutants which are usually sorted out. Essentially, the method is based on the mutation and selection of computer programs similarly as for species in the biological evolution. In this way, an initial program which contains mutatable parts becomes highly optimized in the course of successive rounds of evolution



**Fig. 12.2** A suitable structure of a program for self-evolution. The genes are the mutatable parts of the program. The genes improve the start solution of the given task



Initial computer program (wild-type)

## First evolution step

20 mutant programs as a result of the first evolution step

The best-performing program is used as the new wild-type for the second evolution step

## Second evolution step

20 new mutant programs as a result of the second evolution step

The best-performing program is used as the new wild-type for the third evolution step

## Third evolution step

20 new mutant programs as a result of the third evolution step

The best-performing program is used as the new wild-type for the next evolution step

## Further evolution steps ...

Fig. 12.3 Example of evolutionary computer programming

as a template for further mutagenesis (the next evolution step). The pathway of evolution is highlighted by filled rectangles in Fig. 12.1. Open rectangles correspond to dead ends of the evolution, e.g., mutants which did not perform best

at a certain stage of evolution. In order to overcome high energy barriers in the evolution, also less well performing mutants may be tried in further evolution steps (not shown).

Fig. 12.2 gives an example for a suitable program structure. The genes are the parts of the program changed in the course of the evolution. Mutation is performed, e.g., by modification of the genes, deletion, addition, or exchange of genes. As nature teaches us, gene shuffling is an efficient method to improve the genome of the program. The specific changes of the genes in the mutagenesis depend on the technological task the program is intended to fulfil (see the next sections). Important in this method is the significant change of the program code in the course of the evolution. Despite significant flexibility, the program stays slim. Completely different, nonlinear program structures are feasible.

#### 12.3 Protein folding and structure simulations

Evolution computer methods as described in the previous section were developed to demonstrate a new way of protein folding and structure predictions (Nölting et al., 2004). Folding of proteins has been shown to be an extremely complex, but also surprisingly efficient process (see Chap. 11). For a long time, the origin of the extreme efficiency of protein folding remained unknown. Only recently, the development of new methods for the high resolution of transition states of proteins



**Fig. 12.4** Evolution of the program for protein structure predictions (Nölting et al., 2004; Nölting, 2005)

provided some insight into the mechanism of folding (see Chap. 11): several small proteins were shown to fold via a nucleation–condensation mechanism where structure grows similar to a crystallization process (Nölting and Andert, 2000; Chap. 11).

Fig. 12.4 shows the improvement of the program in the course of the evolution (Nölting et al., 2004; Nölting, 2005). With the program at evolution step 20 in Fig. 12.4, the loop in chymotrypsin inhibitor 2 (CI2) was correctly predicted (Nölting et al., 2004). Results with the program at step 24 (Fig. 12.4) show that it can predict essential features of the structure of several small proteins within less than 1 hour run on a PC (Nölting, 2005).

# 12.4 Evolution of nanooptical devices made from nanoparticles

A computer method for the creation and optimization of nanooptical devices is introduced. In this method a computer program evolves itself towards a higher efficiency. In contrast to most simple self-learning programs, in the course of the evolution, the program code changes significantly (see Sect. 12.2). The highly evolved program is then used to calculate optimized arrangements of nanooptical elements with the desired optical properties. In applications on complex arrays of nanoparticles and other complex scientific and technological problems, such selfevolving computer programs may be many orders of magnitude more efficient than programs developed solely by rational design.

#### 12.4.1 Materials and methods

*Evolution of the computer program.* Figs. 12.1 and 12.3 show the principle of operation of the evolution method (Nölting et al., 2004). The method is based on the mutation and selection of computer programs in successive rounds of evolution (see Sects. 12.2, 12.3 and Nölting et al., 2004). Each evolution step of the computer program on the way from the initial wild-type to the finally evolved mutant program proceeds as follows: a) create different mutants of the computer program, b) test the performance of the mutants, c) take the best-performing program within the set of mutants and wild type-program as a template for further mutations. Mutations are, e.g., omissions, additions, or changes of program lines, or changes of the sequence of lines. Only after a few evolution steps, a program suitably designed for this mutagenesis approach can usually perform more than ten times better than the original wild-type program (Nölting et al., 2004).

In this case, the self-evolving program was written in C++ and run on a PC. Because of the aim to achieve a high speed of the program, the time for testing each mutant was always only about 1 minute.



Fig. 12.5 The optical setup the evolution method is applied on (see the text)

*Optical setup:* The specific optical arrangement investigated with this evolution method is shown in Fig. 12.5. A light beam with the shape of a plane wave with a wavelength of 500 nm is passed through an array of nanoparticles. The diffraction pattern is measured on a screen at a distance of 100 nm.

*Calculation of the diffraction pattern.* The diffraction pattern, F(S), of an arrangement of nanoparticles with the diffraction (scattering) power,  $\rho(r) = 1$ , is given by (see Eq. 4.3):

$$F(S) = \int_{-\infty}^{\infty} e^{-2\pi i r S} dr, \qquad (12.1)$$

where *r* is the position of a scattering particle, and  $S = (s - s_0)/\lambda$  is the scattering vector with  $s_0$  = vector (direction) of the incident plane wave, s = vector (direction) of the scattered wave,  $\lambda$  = wavelength of the light.

#### 12.4.2 Results and discussion

Fig. 12.6 presents the success of the evolution process. RMSD (root mean square deviation) is a measure of the quality of the match between the desired optical pattern and the optical pattern caused by the nanoparticles after optimization of their position by the program. One can see that there is a rapid improvement in the course of the program evolution. Fig. 12.7 shows a specific example of an optical effect aimed at. The evolved program started with a random distribution of 29 nanoparticles (Fig. 12.7a). It then moved the nanoparticles and compared the generated diffraction pattern with the diffraction pattern aimed at (Fig. 12.7c). Successful moves were accepted, the others were rejected. Figs. 12.7d and 12.7e show the final diffraction pattern and positions of the nanoparticles, respectively. The evolved program found this arrangement of particles within 1 minute on a PC.



**Fig. 12.6** Evolution of the program for the calculation of complex patterns of nanoparticles with given optical properties. RMSD is the root mean square deviation of the optical pattern generated by the nanoparticle array compared with the demanded pattern. Each evolution step corresponds to one round of mutagenesis and testing



**Fig. 12.7** Optimization of 29 nanoparticles placed on a  $16 \times 16$  pixel array. The size of each pixel is  $100 \times 100$  nm<sup>2</sup>, the wavelength of the light is 500 nm, and the distance between the nanoparticles and screen is 100 nm. (a) initial positions of the nanoparticles; (b) initial diffraction pattern resulting from the arrangement of these particles; (c) diffraction pattern aimed at; (d) diffraction pattern of the optimized arrangement of the nanoparticles; (e) optimized arrangement of the nanoparticles. The shown arrangement of nanoparticles was found by the evolved program on a PC within 1 minute among ~ $10^{38}$  possibilities

These arrays seem simple, but the number, N, of possible arrangements of the 29 nanoparticles on the 256 places of the array is:

$$N = {\binom{256}{29}} = {\binom{256!}{29! \times 227!}} \approx 10^{38} .$$
 (12.2)

At the beginning, the evolution progresses quite rapidly and appears to slow down after a certain number of evolution steps (Fig. 12.6). However, even after a relatively long period of evolution, further jumps are possible: see, e.g., the significant progress from evolution step 20 to 21. The progress of evolution appears to depend largely on the degree of freedom which is allowed for the system. Strong restrictions may cause the evolution to end prematurely. On the other hand, very fanciful and severe mutations are often negative mutations, but, if positive, have a higher potential to enable a sustained progress over many evolution steps.

One might argue that the geometry of the scattering array may more easily be found by Fourier-transforming the demanded pattern. This is because the diffraction pattern, F(S), has the form of the Fourier transform and so the optical arrangement could be calculated with the inverse Fourier transform of the diffraction pattern (see Chap. 4). However, when recording the diffraction pattern on a screen, the phase information gets lost and so the inverse Fourier transform is not easily done. More importantly, however, none of the solutions might be compatible with a fixed number or density of nanoparticles. Furthermore, some optical effects are difficult to calculate by Fourier transform. For example, the Fourier transform is not easily applicable if one tries to find an arrangement with near-invisibility of the nanoparticle array within a certain spectral region. Also, with this approach one might find completely new approximate solutions.

Here, only a short evolution of a simple program is presented. Significant improvement was achieved by changing the positions of the nanoparticles in the third dimension, i.e., the height above the support layer. Even though there are  $\sim 10^{73}$  possible arrangements of 29 particles in a  $16 \times 16 \times 16$  3D array of nanoparticles, the evolved program provided better results (not shown).

It is expected that the following means will lead to further improvement: (a) the program starts with more different random arrangements of the nanoparticles (start solutions in Fig. 12.2) and searches for the one which yields the best results, (b) the program is passed through further rounds of evolution cycles, (c) the program is given more time for the optimization of the arrays.

#### 12.5 Further potential applications

Self-evolving systems have a much wider applicability. In principle such systems should be a powerful way out when the objects of investigation are too compli-

cated and too complex to be easily rationalized (see, e.g., Back, 1996). Adaptive optics (Fig. 12.8), electrical circuits (Fig. 12.9), robots and nanomechanics (Fig. 12.10) are suitable targets.



**Fig. 12.8** Adaptive systems are used in optics, e.g., to compensate for bending of the support structures of large mirrors due to its own weight and thermal fluctuations and for the distortion of the image by density fluctuations of the atmosphere



Fig. 12.9 Macroscopic circuits and nanocircuits made, e.g., from nanoparticles and nanotubes can have an exponential rise of possibilities with the number of structural elements



**Fig. 12.10** The calculation of movements of sophisticated robots can be connected with an exponential growth of possibilities with the degree of freedom of the system and accordingly such movements may be difficult to optimize
Self-evolving programs should also be a powerful tool to study the evolution of species (Fig. 12.11): why have species evolved in the way they did. Which parameters have an effect on the evolution? Which evolutionary mechanisms are encoded in the DNA? What causes dead ends in the evolution? What will, in this regard, be the fate of human kind and how can it be affected? These are important yet largely unanswered questions.



Fig. 12.11 Scheme of the evolution of species

## **13 Conclusions**

Biophysical methods are extremely important for the further understanding of biological processes. Structurally highly resolving methods such as X-ray crystal-lography and scanning probe microscopy in combination with kinetic methods (see, e.g., Nölting, 2005) give us a true understanding of biological processes at a molecular and cellular level. Proteome maps of healthy and ill individuals are compared for identification of up- or down-regulation in disease states and for individual, highly efficient drug targeting. Biophysical nanotechnology and mass spectrometry open new fascinating ways of studying and influencing complex biological systems. Biophysical nanotechnology takes novel approaches to assemble protein nanoarrays, nanoparticles, and nanowires to well-functioning structures. Mass spectrometry and ion mobility spectrometry have significantly advanced towards the detection of ultra-traces not only of chemical, but also of biological agents.

The biophysical understanding of the living world is crucial to develop rational strategies to influence, in a responsible manner, pathological and non-pathological limitations, stress situations, and disease states. In this way, biophysics can contribute to the understanding of the factors that affect the prosperity and evolution of the human society.

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