# Digital Biology 

L. Ridgway Scott<br>University of Chicago

Ariel Fernández<br>Rice University

## Release 0.3 DO NOT DISTRIBUTE

February 28, 2008

Draft: February 28, 2008, do not distribute

## Contents

1 How is biology digital? ..... 1
2 Challenges of protein models ..... 3
2.1 Digital nature of molecules ..... 3
2.2 Digital nature of proteins ..... 4
2.3 Eternal Strugole ..... 5
2.4 Biological ambivalence ..... 6
2.5 Pchemomics ..... 7
2.5.1 A new tool? ..... 8
2.5.2 Data mining definition ..... 9
2.5.3 Data mining lens ..... 9
2.5.4 Hydrogen bonds are orientation-dependent ..... 10
2.5.5 What is an answer? ..... 10
2.6 Multiscale models ..... 11
2.7 Exercises ..... 13
3 Electronic forces ..... 15
3.1 Direct bonds ..... 15
3.1.1 Covalent bonds ..... 15
3.1.2 Ionic bonds ..... 16
3.1.3 Hvdrogen bonds ..... 17
3.1.4 Cation- $\pi$ interactions ..... 17
3.2 Charge-force relationship ..... 17
3.3 Interactions involving dipoles ..... 18
3.3.1 Single-file dipole-dipole interactions ..... 18
3.3.2 Parallel dipole-dipole interactions ..... 20
3.3.3 Dipole stability ..... 21
3.3.4 Different dipoles ..... 23
3.4 van der Waals forces ..... 23
3.5 Induced dipoles ..... 24
3.5.1 Debve forces ..... 24
3.5.2 London dispersion forces ..... 25
3.5.3 Lennard-Jones potentials ..... 27
3.6 Hydrophobic interactions ..... 27
3.6.1 Solvent mediation of electric forces ..... 28
3.6.2 Dehydrons ..... 28
3.6.3 Dynamics of dehydrons ..... 29
3.6.4 Simulated dynamics ..... 31
3.6.5 Stickiness of dehydrons ..... 32
3.7 Exercises ..... 32
4 Protein basics ..... 35
4.1 Chains of amino acid residues ..... 35
4.1.1 Hydrogen bonds and secondary structure ..... 37
4.1.2 Taxonomies of amino acids ..... 37
4.1.3 Wrapping of hydrogen bonds ..... 39
4.2 Special bonds ..... 41
4.2.1 Salt Bridges ..... 41
4.2.2 Disulfide bonds ..... 42
4.3 Post-translational modifications ..... 42
4.4 Mechanical properties of proteins ..... 43
4.4.1 Conformational geometry of proteins ..... 43
4.4.2 $\phi . \psi$ versus $\psi . \phi$ : the role of $\theta$ ..... 44
4.4.3 Sidechain rotamers ..... 44
4.4.4 Volume of side chains ..... 45
4.5 Special side chains ..... 46
4.5.1 Glycine (and Alanine) ..... 46
4.5.2 Proline ..... 46
4.5.3 Arginine ..... 46
4.5.4 Cysteine ..... 47
4.5.5 Aromatic sidechains ..... 47
4.5.6 Remembering code names ..... 47
4.6 Sidechain ionization ..... 48
4.7 Exercises ..... 48
5 Hydrogen bonds ..... 51
5.1 Hydrogen bond theory ..... 51
5.2 Types of hydrogen bonds ..... 52
5.3 Identification of hydrogen positions ..... 54
5.4 Geometric criteria for hydrogen bonds ..... 56
5.5 Carboxyl-carboxylate hydrogen bonds ..... 57
5.6 Exercises ..... 58
6 Determinants of protein-protein interfaces ..... 59
6.1 Amino acids at protein-protein interfaces ..... 60
6.2 Interface propensity ..... 61
6.3 Amino acid pairs at interfaces ..... 64
6.4 Pair frequencies ..... 65
6.5 Comparisons and caveats ..... 68
6.6 Conclusions ..... 69
6.7 Exercises ..... 69
7 Wrapping electrostatic bonds ..... 71
7.1 Assessing polarity ..... 72
7.1.1 Electronegativity scale ..... 72
7.1.2 Polarity of groups ..... 73
7.2 Counting residues ..... 75
7.2.1 Desolvation domain ..... 76
7.2.2 Predicting aggregation ..... 78
7.3 Counting nonpolar groups ..... 79
7.3.1 Distribution of wrapping for an antibody complex ..... 79
7.4 Residues versus polar groups ..... 81
7.5 Defining dehydrons via geometric requirements ..... 81
7.6 Exercises ..... 85
8 Stickiness of dehydrons ..... 87
8.1 Surface adherence force ..... 87
8.1.1 Biological surfaces ..... 87
8.1.2 Soluble proteins on a surface ..... 88
8.2 A two-zone model ..... 88
8.2.1 Boundary zone model ..... 89
8.2.2 Diffusion zone model ..... 89
8.2.3 Model validity ..... 90
8.3 Direct force measurement ..... 90
8.4 Membrane morphology ..... 92
8.4.1 Protein adsorption ..... 92
8.4.2 Densitv of invaginations ..... 92
8.5 Kinetic model of morphology ..... 93
8.6 Exercises ..... 94
9 Electronic force details ..... 95
9.1 Global accumulation of electric force ..... 95
9.2 Modeling interactions among polar and charged residues ..... 96
9.2.1 Dipole-dipole interactions ..... 96
9.2.2 Charge-dipole interactions ..... 102
9.2.3 Charge-charge interactions ..... 103
Draft: February 28, 2008, do not distribute ..... iii
9.3 General form of a charge group ..... 105
9.3.1 Asvmptotics of general potentials ..... 105
9.3.2 Application of (9.40) ..... 107
9.3.3 Multipole summary ..... 108
9.4 Quadrupole potential ..... 108
9.4.1 Opposing dipoles ..... 108
9.4.2 Four-corner quadrupole ..... 110
9.4.3 Quadrupole example ..... 111
9.4.4 Water: dipole or quadrupole? ..... 112
9.5 Further results ..... 113
9.5.1 Dipole induction by dipoles ..... 113
9.5.2 Modified dipole interaction ..... 114
9.5.3 Hvdrogen placement for Ser and Thr ..... 115
9.6 Exercises ..... 116
10 Units ..... 119
10.1 Basic units vs. derived units ..... 119
10.1.1 SI units ..... 120
10.2 Biochemical units ..... 120
10.2.1 Molecular length and mass units ..... 120
10.2.2 Molecular time units ..... 121
10.2.3 Charge units ..... 121
10.2.4 Conversion constants ..... 123
10.3 Quantum chemistry units ..... 123
10.4 Laboratorv units ..... 124
10.5 Mathematical units ..... 124
10.6 Evolutionarv units ..... 125
10.7 Other phvsical properties ..... 126
10.7.1 The pH scale ..... 126
10.7.2 Polarity and polarization ..... 126
10.7.3 Water densitv ..... 127
10.7.4 Fluid viscositv and diffusion ..... 127
10.7.5 Kinematic viscosity ..... 127
10.7.6 Diffusion ..... 128
10.8 Exercises ..... 128
11 Case studies ..... 129
11.1 Basic cases ..... 129
11.1.1 A singular case ..... 129
11.1.2 Forming structures ..... 129
11.2 Enzvmatic activitv ..... 131
11.3 Neurophysin/vasopressin binding ..... 132
Draft: February 28, 2008, do not distribute ..... iv
11.4 Variations in proteomic interactivity ..... 133
11.5 Sheets of dehydrons ..... 136
11.6 Exercises ..... 136
12 Sidechain-mainchain hydrogen bonds ..... 137
12.1 Counting the bonds ..... 137
12.2 Proline-like configurations ..... 138
12.2.1 Nearest neighbor connections ..... 139
12.2.2 Further neighbors ..... 144
12.3 All sidechain hydrogen bonds ..... 144
12.4 Unusual hydrogen bonds ..... 150
12.4.1 Hvdrophobic pairs ..... 150
12.4.2 Unusual trios ..... 150
12.5 Exercises ..... 151
13 Aromatic interactions ..... 153
13.1 Cation- $\pi$ interactions ..... 153
13.2 Aromatic-polar interactions ..... 154
13.3 Aromatic-aromatic interactions ..... 154
13.4 Exercises ..... 159
14 Peptide bond rotation ..... 161
14.1 Peptide resonance states ..... 161
14.2 Measuring variations in $\omega$ ..... 162
14.3 Predicting the electric field ..... 163
14.4 Implications for protein folding ..... 166
14.5 Exercises ..... 167
15 Tree representations of data ..... 169
15.1 Distance metrics ..... 169
15.1.1 Triangle inequalitv ..... 170
15.1.2 Non-metric measurements ..... 170
15.2 Sequence distance ..... 172
15.2.1 Two-letter alphabets ..... 172
15.2.2 General alphabets ..... 173
15.2.3 Distance versus score ..... 173
15.3 Feature distance ..... 173
15.4 Tree representation of metrics ..... 175
15.4.1 Three point metrics ..... 176
15.4.2 Four point condition ..... 176
15.4.3 A reduction algorithm ..... 177
15.4.4 Obstruction to reduction ..... 178
15.4.5 The ABC theorem ..... 180
15.5 What UPGMA does ..... 182
15.6 Exercises ..... 182
16 Quantum models ..... 185
16.1 Why quantum models? ..... 185
16.2 The Schrödinger equation ..... 185
16.2.1 Particle spin ..... 186
16.2.2 Interpretation of $\psi$ ..... 186
16.3 The eigenvalue problem ..... 187
16.4 The Born-Oppenheimer approximation ..... 189
16.5 External fields ..... 191
16.5.1 Polarization ..... 192
16.5.2 Induced fields ..... 193
16.5.3 Hartree approximation ..... 197
16.6 Comparisons and conclusions ..... 200
16.7 Repulsive molecular forces ..... 201
16.8 The Madelung equation ..... 203
16.9 Exercises ..... 204
17 Continuum equations for electrostatics ..... 205
17.1 Understanding dielectrics ..... 205
17.2 Dielectric materials ..... 205
17.3 Polarization field ..... 207
17.4 Frequency dependence ..... 208
17.5 Spatial frequency dependence ..... 209
17.5.1 Poisson-Debve equation: bulk case ..... 210
17.5.2 Response to a point charge ..... 211
17.5.3 Non-local relationship between $\mathbf{p}$ and $\mathbf{e}$ ..... 213
17.6 The Poisson-Debve equation: general case ..... 213
17.7 Solving the Poisson-Debve equation ..... 215
17.7.1 Numerical methods for FIO's ..... 215
17.8 Modeling DNA ..... 215
18 Water structure ..... 217
18.1 Understanding dielectrics ..... 217
18.2 Tetrahedral structure of waten ..... 217
18.3 Structural water in proteins ..... 217
18.4 Polarizable water models ..... 219
18.5 A two-D water model ..... 219
19 Hydrophobic effect ..... 221
20 Disorder in Protein Structure ..... 223
Draft: February 28, 2008, do not distribute ..... vi
21 Geckos' feet ..... 225
22 Notes ..... 227
23 Glossary ..... 229

## Chapter 1

## How is biology digital?

Biology can be viewed as an information system. As a simple example, we are biological entities communicating via this book. More to the point, many types of signaling in biological systems involve interactions between proteins and ligands. A type of physical baton-passing is used to communicate requirements. But there are too many examples of information processing in biology to stop here to enumerate them. What is of interest here is to understand how certain biological systems (involving proteins) function as digital information systems despite the fact that the underlying processes are analog in nature.

We primarily study proteins and their interactions. These are often involved in signaling and function in a discrete (or digital, or quantized) way. In addition, proteins are discrete building blocks of larger systems, such as viruses and cells. How they bind together (e.g., in a virus capsid) is also deterministic (repeatable) and precise. But the chemical/physical mechanisms used are fundamentally continuous.

Digital circuits on computer chips are also based on continuous mechanisms, namely electrical currents in wires and electronic components. The analogy with our topic is hopefully apparent. A book by Mead and Conway [167] written at the end of the 1970's transformed computer architecture by emphasizing design rules that simplified the task of converting a fundamentally analog behavior into one that was digital and predictable. We seek to do something analogous here, but we are not in a position to define rules for nature to follow. Rather, we seek to understand how some of the predictable, discrete behaviors of proteins can be explained as if certain methodologies were being used.

The benefits of finding simple rules to explain complicated chemical properties are profound. The octet rule for electron shell completion allowed rapid prediction of molecule formulation by simple counting [192]. Resonance theory (Section 14.1) describes general bonding patterns as a combination of simple bonds (e.g, single and double bonds) [191]. The discrete behavior of DNA elucidated by Crick, Franklin, Watson, Wilkins and others [247, 250, 90] initiated the molecular biology revolution. Our objective here is to provide an introduction to some basic properties of protein-ligand interactions with the hope of stimulating further study of the discrete nature of molecular interactions in biology.

The only force of interest in biochemistry is the electric force. Electrical gradients in proteins
are among the largest known in nature. Moreover, we are primarily interested in proteins operating in an aqueous, and thus dielectric, environment. The dielectric properties of water are among the strongest in nature, and indeed water can be viewed as hostile to proteins. This leads to an interesting contention that we address in more detail in Section [2.3,

Not only is the dielectric coefficient of water remarkably large, but it is also capable of being strongly modulated in ways that are still being unveiled. In particular, hydrophobic effects modulate the dielectric properties of water [53]. Proteins are an amazing assembly of hydrophobic, hydrophilic and amphiphilic side chains. Moreover, the charge variation on proteins is so large that it is hard to make an analogy on larger scales, and the variation in hydrophobicity is equally extreme. Hydrophobic mediation of the dielectric properties of water appears to have significant impact on protein function. Thus we are faced with a series of counterbalancing and extreme properties that must be comprehended in order to see how proteins are functioning at a biophysical level.

Our take home message is that the modulation of the dielectric properties of water by the hydrophobic parts of proteins is an essential aspect of molecular chemistry that needs to be considered carefully. Typical representations of proteins show only physical location, basic bonds and individual charges. Adding a way of viewing the modulation of the dielectric environment is of course complex. We review one effective technique that utilizes a representation which signals the effect of the dielectric modulation on hydrogen bonds. Similar techniques can be applied to other bonds as well. But this is an area where further innovation will be needed.

This is not a summary of finished work. Rather it is intended to stimulate study of the detailed mechanisms of protein interactions. We expect this to require many hands. Our intention here is to help stimulate in particular study of some more mathematical questions, many of which we leave open. To quote Mead and Conway [167], "And thus the period of exploration begins."

## Chapter 2

## Challenges of protein models

We present here a sketch of some of the main ideas that the book will cover. This is not an outline but rather is a narrative that introduces the main goals and challenges to be addressed, and gives a glimpse of some of the major advances.

We describe some challenging features of modeling the interactions of proteins in biological systems as well as opportunities to be addressed in the future. This is meant to provide some orientation, but it is also meant to be a disclaimer. That is, we disclose what we see as limitations of standard approaches which have forced us to adopt new strategies. There may well be other approaches that will be even more successful in the future.

### 2.1 Digital nature of molecules

We begin by illustrating what we mean by digital, or discrete, behavior in analog, or continuous, systems. This gives us an opportunity to review some basic concepts from chemistry. The building blocks of chemistry are atoms. They can be characterized by the number of electrons, protons and neutrons of which they are composed. The atoms of primary interest in protein biochemistry are listed in Table 2.1.

Some comments are in order about Table [2.1. First of all, the number of neutrons can vary; we have listed what is known as the dominant isotope. Neutrons add mass but not charge. Other isotopes are important in various contexts; a hydrogen atom with an extra neutron is called deuterium. Atoms occur naturally in different isotopic forms, and the atomic 'weight' (properly, the mass) reflects this natural variation. Otherwise, the atomic mass would be essentially the sum of the numbers of protons and neutrons, with a small correction for the electronic mass; the rest mass of an electron is less that 0.00055 atomic mass units. In the column 'variation' we give the difference between the atomic 'weight' (the mass in atomic units) and the mass of the standard isotope's protons and neutrons. For chlorine, the atoms with 18 and 19 neutrons are nearly equally common. The given atomic masses are themselves only averages, and any particular set of atoms will vary in composition slightly; see the Periodic Table in [192].

Several rules are encoded in Table 2.1. The first rule is used to reduce the number of columns: the number of protons always equals the number of electrons (the net charge is zero). A second rule

| Atom | Symbol | $+/-$ | neutrons | outer | lacking | variation | radius |
| :---: | :---: | :---: | :---: | :---: | :---: | :--- | :--- |
| Hydrogen | H | 1 | 1 | 1 | 1 | +0.0008 | 1.20 |
| Carbon | C | 6 | 6 | 4 | 4 | +0.01 | 1.70 |
| Nitrogen | N | 7 | 7 | 5 | 3 | +0.007 | 1.55 |
| Oxygen | O | 8 | 8 | 6 | 2 | -0.0006 | 1.52 |
| Fluorine | F | 9 | 10 | 7 | 1 | -0.002 | 1.47 |
| Sodium | Na | 11 | 12 | 1 | 7 | -0.01 | 2.27 |
| Magnesium | Mg | 12 | 12 | 2 | 6 | +0.31 | 1.73 |
| Phosphorus | P | 15 | 16 | 5 | 3 | -0.03 | 1.80 |
| Sulfur | S | 16 | 16 | 6 | 2 | +0.06 | 1.80 |
| Chlorine | Cl | 17 | 18 | 7 | 1 | +0.45 | 1.75 |
| Potassium | K | 19 | 20 | 1 | 7 | +0.10 | 2.75 |

Table 2.1: Subset of the periodic table. The column ' $+/-$ ' denotes the number of protons and electrons in the atom. The column 'outer' is the number of electrons in the outer shell. The column 'lacking' is the number of electrons needed to complete the outer shell. The column 'variation' give the difference between the observed atomic 'weight' (the mass in atomic units) of the naturally occurring isotopic distribution and the mass of the 'standard' isotope's protons and neutrons. The column 'radius' lists the 'mean' van der Waals radius [26].
is that the typical number of neutrons in the dominant isotope is nearly the same as the number of protons. But the most important rule is the octet rule: the number of the electrons in the outer shell plus the number (listed in the 'lacking' column) of electrons contributed by atoms covalently bonded to it is always eight (except for hydrogen). This simple rule facilitates the determination of molecular bond formation.

The digital description of an atom is to be contrasted with the analog description of the Schrödinger equation (see Chapter (16). This equation describes the electron distribution, which is the key determinant of atomic interaction. We are forced to consider effects on this level in many cases, but operating at the atomic level has clear advantages.

There are other simple rules in chemistry that clarify bond formation, such as the resonance principle (Section 14.1). This rule states that observed states of molecular bonds are often a simple convex combination of two elementary states. For example, a benzene ring can be thought of as being made of alternating single and double bonds, whereas in reality each bond is closely approximated by a convex combination of these two bonds. The resonance principle may be thought of as a Galerkin approximation to solutions of the Schrödinger equation (see Chapter 16).

We seek to illuminate rules like these in proteins; see Chapter 4 for an introduction to proteins.

### 2.2 Digital nature of proteins

The digital and deterministic nature of protein function is implied by the fact that their structure is encoded by a discrete mechanism, DNA. There are post-translational events (Section 4.3) which
modify proteins and make their behavior more complex, but it is clear that nature works hard to make proteins in the same way every time.

What is striking about the fact that proteins act in quantized ways is the observation that hydrophobic effects (Section 3.6) are involved in most protein-ligand interactions. Such interactions account not only for the formation of protein complexes, but also for signaling and enzymatic processes. But the hydrophobic effect is essentially nonspecific. Thus its role in a discrete system is intriguing.

We will see that it is possible to quantify the effect of hydrophobicity in discrete ways. The concept of wrapping (see Chapter 7) yields such a description, and we show that this can effect many important phenomena, including protein binding (Chapter 6) and the flexibility of the peptide bond (Chapter 14).

### 2.3 Eternal Struggle

The life of a protein in water is largely a struggle for the survival of its hydrogen bonds. The hydrogen bond (cf. Chapter (5) is the primary determinant of the structure of proteins. But water molecules are readily available to replace the structural hydrogen bonds with hydrogen bonds to themselves; indeed this is a significant part of how proteins are broken down and recycled. We certainly cannot live without water, but proteins must struggle to live with it [147].

Proteins are the fabric of life, playing diverse roles as building blocks, messengers, molecular machines, energy-providers, antagonists, and more. Proteins are initiated as a sequence of amino acids, forming a linear structure. They coil into a three-dimensional structure largely by forming hydrogen bonds. Without these bonds, there would be no structure, and there would be no function. The linear structure of amino acid sequences is entropically more favorable than the bound state, but the hydrogen bonds make the three-dimensional structure energetically favorable.

Water, often called the matrix of life [91, is one of the best makers of hydrogen bonds in nature. Each water molecule can form hydrogen bonds with four other molecules and frequently does so. Surprisingly, the exact bonding structure of liquid water is still under discussion [1, 228, 248, but it is clear that water molecules can form complex bond structures with other water molecules. For example, water ice can take the form of a perfect lattice with all possible hydrogen bonds satisfied.

But water is equally happy to bind to available sites on proteins instead of bonding with other water molecules. The ends of certain side chains of amino acids look very much like water to a water molecule. But more importantly the protein backbone hydrogen bonds can be replaced by hydrogen bonds with water, and this can disrupt the protein structure. This can easily lead to the break-up of a protein if water is allowed to attack enough of the protein's hydrogen bonds.

The primary strategy for protecting hydrogen bonds is to bury them in the core of a protein. But this goes only so far, and inevitably there are hydrogen bonds formed at the surface of a protein. And our understanding of the role of proteins with extensive non-core regions is growing rapidly. The exposed hydrogen bonds are more potentially interactive with water. These are the ones that are most vulnerable to water attack.

Amino acids differ widely in the hydrophobic composition of their side chains (Section 4.1.2). Simply counting carbonaceous groups (e.g., $\mathrm{CH}_{n}$ for $n=1,2$ or 3 ) in the side chains shows a striking
range, from zero (glycine) to nine (tryptophan). Most of the carbonaceous groups are non-polar and thus hydrophobic. Having the right amino acid side chains surrounding, or wrapping, an exposed hydrogen bond can lead to the exclusion of water, and having the wrong ones can make the bond very vulnerable. The concept of wrapping an electrostatic bond by nonpolar groups is analogous to wrapping live electrical wires by non-conducting tape.

We refer to the under-protected hydrogen bonds which are under-wrapped by carbonaceous groups as dehydrons (Section 3.6.2) to simplify terminology. The name derives from the fact that these hydrogen bonds benefit energetically from being dehydrated.

### 2.4 Biological ambivalence

One could imagine a world in which all hydrogen bonds were fully protected. However, this would be a very rigid world. Biology appears to prefer to live at the edge of stability. Moreover, it has been recently observed that exposed hydrogen bonds appear to be sites of protein-protein interactions [79]. Thus what at first appears to be a weakness in proteins is in fact an opportunity.

One could define an epidiorthotric force as one that is associated with the repair of defects. The grain of sand in an oyster that leads to a pearl can be described as an epidiorthotric stimulant. Such forces also have analogies in personal, social and political interactions where forces based on detrimental circumstances cause a beneficial outcome. The defect of an under-protected hydrogen bond gives rise to just such an epidiorthotric force. The action of this force is indirect, so it takes some explaining.

An under-protected hydrogen bond would be much stronger if water were removed from its vicinity. The benefit can be understood first by saying that it is the result of removing a threat of attack (or the intermittent encounter of water forming hydrogen bonds with it). But there is an even more subtle (but mathematically quantifiable) effect due to the change in dielectric environment when water is removed, or even just structured, in the neighborhood. The dielectric constant of water is about eighty times that of the vacuum, or even non-polar materials. Changing the dielectric environment near an under-protected hydrogen bond makes it substantially stronger.

If the removal of water from an under-protected hydrogen bond is energetically favorable, then this means there is a force associated with attracting something that would exclude water. Indeed, one can measure such a force, and it agrees with what would be predicted by calculating the change energy due to the change in dielectric (Section 8.1). You can think of this force as being somewhat like the way that adhesive tape works. Part of the force results from the removal of air between the tape and the surface, leaving atmospheric pressure holding it on. However, the analogy only goes so far in that there is an enhancement of electrical energy associated with the removal of water. For sticky tape, this would correspond to increasing the mass of the air molecules in the vicinity of the tape, by a factor of 80 , without increasing their volume!

Thus the epidiorthotric force associated with water-removal from an under-protected hydrogen bond provides a mechanism to bind proteins together. This is a particular type of hydrophobic effect, because wrapping the bond with hydrophobic groups provide protection from water. It is intriguing that it arises from a defect which provides an opportunity to interact.

Draft: February 28, 2008, do not distribute


Figure 2.1: Number of dehydrons [shown in square brackets] in the protein myoglobin found in various species [79], which are presented in an evolutionary tree determined by sequence alignment distances.

### 2.5 Pchemomics

The term "omics" refers to the use of biological data-bases to extract new knowledge by large-scale statistical surveys. The term "cheminformatics" is an accepted moniker for the interaction of informatics and chemistry, so there is some precedent for combining terms like pchem (a.k.a., physical chemistry) with a term like 'omics.' We do not suggest the adoption of the (unpronounceable) term pchemomics, but it serves to suggest the particular techniques being combined in a unique way. An example of pchemomics is the early study of the hydrogen bond [142]. Indeed, the original study of the structure of the peptide bond (see section 8-4 of [191]) used such an approach. But pchemomics involves a two way interaction with data. In addition to providing a way to learn new properties in physical chemistry, it also involves using physical chemistry to look at standard data in new ways.

The Protein Data-Base (PDB) provides three-dimensional structures that yield continuing opportunities for proteomics discoveries. Using the perspective of physical chemistry in datamining in the PDB, some simple laws about protein families were determined by studying patterns of underwrapped hydrogen bonds [72]. We examine just one such result in Section 2.5.4 many other results in physical chemistry can be likewise explored.

A simple view of the PDB only gives a representation suitable for Lagrangian mechanics (or perhaps just statics). If we keep in mind which atom groups are charged, we begin to see an electrostatic view of proteins, and standard protein viewers will highlight the differently charged groups. But the dielectric effect of the solvent is left to the imagination. And the crucial role of the modulation of the dielectric effect by hydrophobic groups is also missing. Adding such views of proteins involve a type of physical chemistry lens.

When you do look at proteins by considering the effect of wrapping by hydrophobic groups, you see many new things that may be interpreted in ways that are common in bioinformatics. One striking observation is that there is a simple correlation between the number of under-wrapped hydrogen bonds and evolutionary trends. Figure 2.1 depicts the number of dehydrons found in the protein myoglobin (or its analog) in various species [79]. Similar trends are seen with other proteins in Table 11.2

The number of under-wrapped hydrogen bonds appears to be evolving (increasingly), providing increasing opportunities for interaction in advanced species. This provides additional understanding of how higher species may have differentiated function without dramatically increasing the number
of genes which code for proteins.
It is also significant that under-wrapped hydrogen bonds appear to be conserved more than other parts of proteins. But since the number of under-wrapped hydrogen bonds is growing, we should say that once they appear they tend to be conserved [79.

Given our understanding of what it means to be under-wrapped, it is not surprising that underwrapped hydrogen bonds would appear more often in regions of proteins that are themselves not well structured. NORS (NO Recognizable Structure) regions [117] in proteins are large (at least seventy consecutive amino acids) sections which form neither $\alpha$-helices or $\beta$-sheets. These appear more frequently among interactive proteins. Correspondingly, studies [82] have shown a strong correlation between the number of under-wrapped hydrogen bonds and interactivity.

A full understanding of wrapping and the related force associated with under-wrapping requires tools from physical chemistry. Interactions between physical chemistry and "omics" will offer further insights into biological systems. Indeed, precise modeling of water even by explicit solvent methods is still a challenge. Only recently have models begun to predict the temperature behavior of the density of liquid water [149. This means that for very subtle issues one must still be careful about even all-atom simulations. The mysteries of water continue to confront us. But its role in biology will always be central.

### 2.5.1 A new tool?

Since we are seeking to answer new types of research questions, it may be comforting to know that there is a powerful tool that is being used. The combination of data mining and physical chemistry is not new, but its usefulness is far from exhausted. Moreover, it is not so common to see these utilized in conjunction with more conventional techniques of applied mathematics, as we do here. Thus we take a moment to reflect on the foundations of the basic concepts that make up what we refer to as pchemomics.

Typical datamining in bioinformatics uses more discrete information, whereas the PDB uses continuous variables to encode chemical properties. The need for physical chemistry in biology has long been recognized. In the book [235], the following quote is featured:

The exact and definite determination of life phenomena which are common to plants and animals is only one side of the physiological problem of today. The other side is the construction of a mental picture of the constitution of living matter from these general qualities. In the portion of our work we need the aid of physical chemistry.

The emphasis at the end was added as an aid to the eye. These words were written by Jacques Loeb in "The biological problems of today: physiology" which appeared in the journal Science in volume 7 , pages $154-156$, in 1897 . So our theme is not so new, but the domain of physical chemistry has advanced substantially in the last century, so there continues to be an important role for it to play in modern biology.

Draft: February 28, 2008, do not distribute

### 2.5.2 Data mining definition

It is useful to reflect on the nature of data mining, since this is a relatively new term. It is a term from the information age, so it is suitable to look for a definition on the Web. According to WHATIS.COM,

Data mining is sorting through data to identify patterns and establish relationships. Data mining parameters include:

- Association - looking for patterns where one event is connected to another event
- Sequence or path analysis - looking for patterns where one event leads to another later event
- Classification - looking for new patterns (May result in a change in the way the data is organized but that's ok)
- Clustering - finding and visually documenting groups of facts not previously known

Our conclusion? Data mining involves looking at data. If data mining is looking at data then what type of lens do we use?

### 2.5.3 Data mining lens

There are many ways to look at the same biological data. In the field of data mining, this might be called using different filters on the data. However, it is not common to look at the same data with many different filters, so we prefer the different metaphor of a lens. It could be a telescope, a microscope, polarized sunglasses, or just a good pair of reading glasses.

All proteins have chemical representations, e.g., the protein

$$
C_{400} H_{620} N_{100} O_{120} P_{1} S_{1} .
$$

In the early research on proteins [235], discovering such formulæ was a major step. But a much bigger step came with the realization that proteins are composed of sequences of amino acids. This allowed proteins to be described by alphabetic sequences, and they come in different forms: DNA, RNA, amino acid sequences. One can think of these from a linguistic perspective, and indeed this has been a productive approach (153.

The function of DNA is largely to store sequence information, but proteins operate as threedimensional widgets. All proteins have a three-dimensional representation, even if it is not one that forms into a stable, biologically viable, structure. The PDB is a curated database of such structures which provides a starting point to study protein function from a physical chemistry perspective.

But structure alone does not explain how proteins function. Physical chemistry can both simplify our picture of a protein and also allow function to be more easily interpreted. In particular, we will emphasize the role of interpreting the modulation of the dielectric environment by hydrophobic effects. We describe a simple way this can be done to illustrate the effect on individual electronic entities, such as bonds. But there is need for better lenses to look at such complex effects.

Draft: February 28, 2008, do not distribute

### 2.5.4 Hydrogen bonds are orientation-dependent

The hydrogen bond provides a good starting example of the use of "pchem" data mining to reveal its properties. Figure 6 of [142] shows clearly both the radial and the angular dependence of the hydrogen bond. Figure 8 of [257] shows a similar relationship between the angle of the hydrogen bond and its distance, derived using protein data. The data in that figure is consistent with a conical restriction on the region of influence of the bond.

More recently, the orientation dependence of the hydrogen bond has been revisited. An orientationdependent hydrogen bonding potential improves prediction of specificity and structure for proteins and protein-protein complexes [173]. Close agreement between the orientation dependence of hydrogen bonds observed in protein structures and quantum mechanical calculations has also been reported [138].

An alternative method for modeling hydrogen bonds is to study their energetics via quantum mechanical calculations and to interpolate the resulting energy surfaces [223, 179].

Due to the primary importance of the hydrogen bond in protein structure, we will review what is known and not known in Chapter 5

### 2.5.5 What is an answer?

Before we begin to ask questions in earnest, we need to talk about what sort of answers we might expect. In high-school algebra, an answer takes the form of a number, or a small set of numbers. In calculus, the answer is often a function. Here, we will often find that the answer is statistical in nature. There appear to be few absolutes in biology, so a probability distribution of what to expect is the best we can hope for.

A probability distribution provides a way to give answers that combine the types of answers you get with high-school algebra and those you get with calculus. An answer that is a number is a Dirac $\delta$-function, whereas a function corresponds to a measure that is absolutely continuous. This added level of sophistication is especially helpful in a subject where it seems almost anything can happen with some degree of probability.

Mathematics tells us that it is a good idea to have metrics for the space of answers that we expect. Metrics on probability distributions are not commonly discussed. We will not make significant use of such metrics, but we review in Section ?? some possible approaches.

In classical physics, problems were often considered solved only when names for the functions involved could be determined. This paradigm is extremely robust and useful. When the names are familiar, they suggest general properties (exponential versus sinusoidal), and they provide a simple algorithm to general specific values for particular instances. The programming language Fortran was designed specifically to facilitate the evaluation of expressions such as

$$
\begin{equation*}
\sin \left(\log \left(\tan \left(\cos \left(J_{1}\left(e^{x}+\sqrt{\pi x}\right)\right)\right)\right)\right) \tag{2.1}
\end{equation*}
$$

Unfortunately, the classical paradigm is limited by our ability to absorb new names. While the names in (2.1) are familiar to many who have studied Calculus, the list required in practice includes less well known Bessel functions, Hankel functions, elliptic functions, theta functions, zeta functions,


Figure 2.2: Spatial and temporal scales of biomolecular models. See the text for more details.
and so on. Moreover, it may be that each new problem requires a new name, in which case the paradigm fails; it is only successful if it provides an abstraction that allows the simplification of the answer. Moreover, strict adherence to this paradigm causes an unnecessary impediment from a computational point of view. All that we may care about is the asymptotic form of a function, or particular values in a certain range, i.e., a plot, or just the point at which it has a minimum.

The newer computational paradigm is not to associate names to solutions, but rather to associate standard algorithms to problems that can be used to provide the information required to understand the mechanism being studied. For example, we may be content if we can specify a well-posed differential equation to be solved to determine numerical values of a function. Thus we might say that the equation $u^{\prime}=u$ is a sufficient description of the exponential function. When we discuss quantum mechanics, we will adopt this point of view.

### 2.6 Multiscale models

But why don't we just write down a mathematical model and use it to simulate protein dynamics? This is a reasonable question, and we attempt here to show why such an approach at the moment would not be productive. The difficulty is the particular multiscale aspect of the problem: the temporal scales are huge but the spatial scales overlap, as depicted in Figure [2.2. Of course, existing models are useful in limited contexts. However, we will explain limitations in two such models that must be addressed in order to use them on more challenging simulations.

Models for many systems have components which operate at different scales [115]. Scale separation often simplifies the interactions among the different scales. The differences often occur in both physical and temporal scales. Scale separation often simplifies the study of complex systems by allowing each scale to be studied independently, with only weak interactions among the different scales. However, when there is a lack of scale separation, interactions among the scales become

Draft: February 28, 2008, do not distribute
more difficult to model.
There are three models of importance in protein biochemistry. The different spatial and temporal scales for these models are depicted in Figure 2.2. The smallest and fastest scale is that of quantum chemistry (Chapter (16). The model involves continuous variables, partial differential equations and functions as solutions.

The molecular scale is more discrete, described only by the positions of different atoms in space, perhaps as a function of time. The time scale of molecular dynamics is much longer than the quantum scale. But the length scale is comparable with the quantum scale. For example, the Ångstrom can be used effectively to describe both without invoking very large or very small numbers.

Finally, the electric properties of proteins are mediated by the dielectric behavior of water in a way that is suitable for a continuum model [50, 217. But again the length scale is not much bigger than the molecular scale. Many solvated systems are accurately represented using a system in which the size of the solvation layer is the same as the protein dimension. On the other hand, dielectric models are inherently time independent, representing a 'mean field' approximation. Thus there is no natural time scale for the continuum dielectric model, but we have depicted in Figure 2.2 the time scale for so-called Brownian dynamics models which are based on a continuum dielectric model [121].

The lack of dramatic physical scale separation, linked with the extreme time separation, in biological systems is the root of some of the key challenges in modeling them. Note that the temporal scales in Figure 2.2 cover fifteen orders of magnitude whereas the spatial scales cover only three or four orders of magnitude. Many biological effects take place over a time scale measured in seconds, but there may be key ingredients which are determined at a quantum level. This makes it imperative to develop simplified rules of engagement to help sort out behaviors, as we attempt to do here.

We do not give a complete introduction to quantum models, but we do include some material so that we can discuss some relevant issues of interest. For example, molecular-level models utilize force fields that can be determined from quantum models, and this is an area where we can predict significant developments in the future. The hydration structure around certain amino acid residues is complex and something that begs further study. But this may require water models which are currently under development, and these models may require further examination at the quantum level.

Multi-scale models are most interesting and challenging when there is significant information flow between levels. One of the most intriguing examples is the effect of the electric field on the flexibility of the peptide bond [66]. The electric field is governed by the largest-scale model and causes a change in the smallest-scale model, forcing a re-structuring of the molecular model (Chapter 14).

The Schrödinger equation is a well-accepted model for quantum chemistry. However, it is too detailed for use as a numerical model for large systems. Molecular dynamics models are used routinely to simulate protein dynamics, but there are two drawbacks. On the one hand, there are some limitations in the basic theoretical foundations of the model, such as the proper force fields to be used, so the predictions may not be fully accurate (cf. Section 14.4). On the other hand, they

Draft: February 28, 2008, do not distribute 12
are still complicated enough that sufficiently long-time simulations, required for biological accuracy, are often prohibitive [7]. Electrostatic models hope to capture the expected impact of dielectric solvation, but there are limitations here as well. The dielectric coefficient of water is orders of magnitude larger than what would be found inside a large protein. This is a very large jump in a coefficient in a continuum model, and it is prudent to be cautious about any model with such large changes. It is clear that in the neighborhood of the jump in the coefficient, a more complex model might be required [217].

### 2.7 Exercises

Exercise 2.1 Download a PDB file for a protein and compute the distance distribution between sequential $C_{\alpha}$ carbons. What is the mean of the distribution? Compare this with the data in the figure at the top of page 282 in [191].

Exercise 2.2 Download a PDB file for a protein and compute the distance distribution between $C_{\alpha}$ carbons separated in sequence by $k$. That is, the sequential neighbors have $k=1$. How does the mean distance vary as a function of $k$ ? Compare the distributions for $k=3$ and $k=4$; which has $C_{\alpha}$ carbons closer together?

Exercise 2.3 Download a PDB file for a protein and compute the N-O distance distribution between all pairs of carbonyl and amide groups in the peptide bonds (cf. Figure 4.1). What is the part of the distribution that corresponds to ones forming a hydrogen bond? (Hint: exclude the N's and O's that are near neighbors in the peptide bond backbone.)

Exercise 2.4 Acquire a pair of polarized sunglasses and observe objects just below the surface of a body of water both with and without the sunglasses. Do these observations while facing the sun, when it is at a low angle with respect to the water surface. You should observe that the 'glare' is greatly reduced by the polarizing lenses. Also make the same observations when the sun is overhead, and when looking in a direction away from the sun when it is at a low angle.

Exercise 2.5 Quantum-mechanical computations suffer from the 'curse of dimensionality' because each additional electron adds another three dimensions to the problem. Thus a problem with $k$ electrons requires the solution of a partial differential equation in $\mathbb{R}^{3 k}$. If we require a discretization with $m$ degrees of freedom per dimension, then the resulting problem requires $m^{3 k}$ words of memory to store the discrete representation. Compare this with the number of atoms in the observable universe. Assuming we could somehow make a computer using all of these atoms with each atom providing storage for one of the $m^{3 k}$ words of memory required for the discrete representation, determine how large a value of $k$ could be used. Try values of $m=3$ and $m=10$.

## Chapter 3

## Electronic forces

The only force of significance in biochemistry is the electric force. However, it appears in many guises, often modulated by indirection, or induction. Chemistry has classified different regimes of electronic forces by cataloging bonds between different atoms. In terrestrial biology, water plays a dominant role as a dielectric that modulates different types of electronic interactions. Some bonds are more easily affected by water than others.

Here we briefly outline the main types of electronic forces as they relate to biology, and especially to proteins and other molecular structures. There are so many books that could be used as a reference that it is hard to play favorites. But the books by Pauling [191, 192] are still natural references.

The order of forces, or bonds, that we consider is significant. First of all, they are presented in order of strength, starting with the strongest. This order also correlates directly with the directness of interaction of the electrons and protons, from the intertwining of covalent bonds to indirect, induced interactions. Finally, the order is also reflective of the effect of solvent interaction to some extent, in that the dielectric effect of solvent is increasingly important for the weaker bonds.

### 3.1 Direct bonds

The strongest bonds can be viewed as the direct interactions of positive and negative charges, or at least distributions of charge.

### 3.1.1 Covalent bonds

These are the strong bonds of chemistry, and they play a role in proteins, DNA, RNA and other molecules of interest. However, their role in biology is generally static; they rarely break. They form the backbones of proteins, DNA, and RNA and support the essential linear structure of these macromolecules. Typical examples are shown in Figures 4.4 4.7 for aminoacid sidechains and Figure 14.1 for the peptide bond. Single lines represent single bonds and double (parallel) lines represent double bonds. One covalent bond of significant note that is not involved in defining the backbone is the disulfide bond (or disulfide bridge) between two cysteine sidechains (Section 4.2.2)
in proteins. Covalent bonds involve the direct sharing of electrons from two different atoms, as required by the octet rule mentioned in Section 2.1] Such bonds are not easily broken, and they typically survive immersion in water. The octet rule [192, 191] allows the prediction of covalent bond formation through counting of electrons in the outer-most shell of each atom. Explaining further such simple rules for other types of bonds is one of the major goals of this work.

Although covalent bonds are not easily broken, their character can be modified by external influences. The most important covalent bond in proteins is the peptide bond (Figure 14.1) formed between amino acids as they polymerize. This bond involves several atoms that are typically planar in the common form of the peptide bond. But if the external electrical environment changes, as it can if the amide and carbonyl groups lose hydrogen bond partners, the bond can bend. We review this effect in Chapter 14.

### 3.1.2 Ionic bonds

Ionic bonds occur in many situations of biological interest, but it is of particular interest due to its role in what is called a salt bridge (Section 4.2.1). Such an ionic bond occurs between oppositely charged side chains in a protein. Ionic bonds involve the direct attraction of electrons in one atom to the positive charge of another.

The potential for the electrostatic interaction between two charged molecules, separated by a distance $r$, is (see Section 3.2)

$$
\begin{equation*}
V(r)=z_{1} z_{2} r^{-1} \tag{3.1}
\end{equation*}
$$

where $z_{i}$ is the charge on the $i$-th molecule. For two molecules with equal but opposite charges, say, $z_{1}=1$ and $z_{2}=-1$, the potential is $-r^{-1}$.

We will see that different bonds are characterized by the exponent of $r$ in their interaction potential. For potentials of the form $r^{-n}$, we can say that the bonds with smaller $n$ are more long range, since $r^{-n} \gg r^{-m}$ for $n<m$ and $r$ large. The ionic bond is thus the one with the longest range of influence.

In addition to being long range, ionic bonds are often stronger as well. For all bonds of attraction which are of the form $r^{-n}$, there would be infinite attraction at $r=0$. However, there is always some other (electrostatic) force of repulsion that keeps the entities from coalescing. We address the form of such a force of repulsion in Section 16.7. Thus the form of the attractive force is not sufficient to tell us the strength of the bond. However, ionic bonds are often quite strong as well as being long range, second only to covalent bonds in strength.

Although ionic bonds are relatively strong and have a long-range influence, they are also easily disrupted by water, as a simple experiment with table salt introduced into a glass of water will easily show. Salt forms a stable crystal when dry, but when wet it happily dissolves into a sea of separated ions. The source of attraction between the sodium and chloride ions in salt is the ionic bond.

Draft: February 28, 2008, do not distribute

### 3.1.3 Hydrogen bonds

Although weaker than covalent and ionic bonds, hydrogen bonds play a central role in biology. They bind complementary DNA and RNA strands in a duplex structure, and they secure the threedimensional structure of proteins. However, they are also easily disrupted by water, which is the best hydrogen bond maker in nature.

First suggested in 1920, hydrogen bonds were not fully accepted until after 1944 [235]. The detailed structure of hydrogen bonds in biology is still being investigated [138, 173, 228, 248. Most of the hydrogen bonds of interest to us involve a hydrogen that is covalently bonded to a heavy atom X and is noncovalently bonded to a nearby heavy atom Y. Typically the heavy atoms X and Y are N, O, or S in protein systems, e.g., NH - O or $\mathrm{OH}-\mathrm{S}$, etc.; see Table 5.2 for a list. The bond $\mathrm{OH}-\mathrm{O}$ describes the hydrogen bond between two water molecules.

The special nature of the hydrogen bond stems in part from the mismatch in size and charge compared to the other so-called 'heavy' atoms. Carbon is the next smallest atom of major biological interest, with six times as many electrons and protons. The mismatch with nitrogen and oxygen is even greater. Hydrogen bonds will be discussed in more detail in Chapter 5

### 3.1.4 Cation- $\pi$ interactions

Aromatic residues (phenylalanine, tyrosine and tryptophan: see Section 4.5.5) are generally described as hydrophobic, due to the nonpolar quality of the carbon groups making up their large rings. But their carbon rings have a secondary aspect which is polar, in that there is a small negative charge distribution on each side of the plane formed by the rings [95, 258, 48]. This large distribution of negative charge can directly attract the positive charges of cations (e.g., arginine and lysine).

Cation- $\pi$ interactions will be discussed in more detail in Chapter 13 ,

### 3.2 Charge-force relationship

We want to talk about the interaction energy (and force) between two charged groups. The units of charge and energy are not the same, and so we need to introduce a conversion factor to allow this.

Suppose we have a charge $z$ at the origin in space. This induces an electric field $\mathbf{e}$ in all of space, and the relationship between the two is

$$
\begin{equation*}
\varepsilon \nabla \cdot \mathbf{e}=z \delta, \tag{3.2}
\end{equation*}
$$

where $\varepsilon$ is the permittivity and $\delta$ denotes the Dirac delta-function. When the medium is a vacuum, this is the permittivity of free space, $\varepsilon_{0}$. In other media (e.g., water) the value of $\varepsilon$ is much larger.

The exact value for $\varepsilon_{0}$ depends on the units (Chapter (10) chosen for charge, space, time, etc. The electric field e does not have units of force. If there is no other charge in the field, no force will be felt. The resulting force on a second charge $z^{\prime}$ is proportional to the amount of that charge:

Draft: February 28, 2008, do not distribute
$z^{\prime} \mathbf{e}$. So the electric field $\mathbf{e}$ has units of force per unit of charge, whereas $z$ has units of charge. The coefficient $\varepsilon$ provides the change of units required by the relation (3.2), cf. (10.2).

The electric field $\mathbf{e}$ can be written as (minus) the gradient of a potential

$$
\begin{equation*}
\mathbf{e}=-\nabla V \tag{3.3}
\end{equation*}
$$

and therefore the potential is related to the charge by

$$
\begin{equation*}
-\varepsilon \Delta V=z \delta \tag{3.4}
\end{equation*}
$$

It is not too difficult to verify that a solution to (3.4) is

$$
\begin{equation*}
V(r)=\frac{z}{4 \pi \varepsilon r} . \tag{3.5}
\end{equation*}
$$

Note that the units of $V$ are energy per unit charge. We can have a simple representation of the relationship between charge $z$ and its electric potential

$$
\begin{equation*}
V(r)=\frac{z}{r} \tag{3.6}
\end{equation*}
$$

provided we choose the units (Chapter (10)) appropriately so that $\varepsilon=1 / 4 \pi$. The resulting potential energy of a pair of charges $z_{1}$ and $z_{2}$ is thus given by (3.1).

We will make this simplification in much of our discussion, but it should be remembered that there is an implicit constant proportional to the permittivity in the denominator. In particular, we see that a larger permittivity leads to a smaller potential and related force.

### 3.3 Interactions involving dipoles

Many interactions can be modeled as dipole-dipole interactions, e.g., between water molecules. More generally, the use of partial charges (cf. Table 13.1) represents many interactions as dipoledipole interactions. Forces between molecules with fixed dipoles are often called Keesom forces 93]. For simplicity, we consider dipoles consisting of the same charges of opposite signs, separated by a distance $2 \epsilon$. If the charges have unit value, then the dipole strength $\mu=2 \epsilon$. Interacting dipoles have two orientations which produce no torque on each other.

### 3.3.1 Single-file dipole-dipole interactions

In the single-file orientation, the base dipole has a positive charge at $(\epsilon, 0,0)$ and a negative charge at $(-\epsilon, 0,0)$; the other dipole is displaced on the $x$-axis at a distance $r$ : a positive charge at $(r+\epsilon, 0,0)$ and a negative charge at $(r-\epsilon, 0,0)$ (cf. Figure 3.1). The potential due to the base dipole at a distance $r \gg \epsilon$ along the $x$-axis is

$$
\begin{align*}
V(r) & =\frac{1}{r-\epsilon}-\frac{1}{r+\epsilon}=\frac{(r+\epsilon)-(r-\epsilon)}{(r-\epsilon)(r+\epsilon)}  \tag{3.7}\\
& =\frac{2 \epsilon}{(r-\epsilon)(r+\epsilon)}=\frac{2 \epsilon}{r^{2}-\epsilon^{2}} \approx 2 \epsilon r^{-2}=\mu r^{-2}
\end{align*}
$$

Draft: February 28, 2008, do not distribute 18


Figure 3.1: Single-file dipole-dipole configuration.
where $\mu=2 \epsilon$ is the dipole strength.
We use the expression $f(r) \approx g(r)$ to mean that the expression $f(r)$ is a good approximation to $g(r)$. More precisely, in this case we mean that the two expressions are asymptotically equal for large $r$, that is, that

$$
\begin{equation*}
\lim _{r \rightarrow \infty} g(r) / f(r)=1 \tag{3.8}
\end{equation*}
$$

In (3.7), $f(r)=1 /\left(r^{2}-\epsilon^{2}\right.$ ) and $g(r)=r^{-2}$, so that $g(r) / f(r)=1-\epsilon^{2} / r^{2}$, and thus (3.8) follows. Moreover, we can get a quantitative sense of the approximation: the approximation in (3.7) is $99 \%$ accurate for $r \geq 10 \epsilon$, and even $75 \%$ accurate for $r \geq 2 \epsilon$.

In the field of the dipole (3.7), the potential energy of a single charge on the $x$-axis at a distance $r$ is thus $\mu r^{-2}$, for a charge of +1 , and $-\mu r^{-2}$, for a charge of -1 . In particular, we see that the charge-dipole interaction has a potential one order lower $\left(r^{-2}\right)$ than a charge-charge interaction $\left(r^{-1}\right)$. The charge-dipole interaction is very important, but we defer a full discussion of it until Section 9.2.2,

The combined potential energy of two opposite charges in the field generated by a dipole is given by the difference of terms of the form (3.7). In this way, we derive the potential energy of a dipole, e.g., a positive charge at $(r+\epsilon, 0,0)$ and a negative charge at $(r-\epsilon, 0,0)$, as the sum of the potential energies of the two charges in the field of the other dipole:

$$
\begin{equation*}
\frac{\mu}{(r+\epsilon)^{2}}-\frac{\mu}{(r-\epsilon)^{2}} . \tag{3.9}
\end{equation*}
$$

Considering two such charges as a combined unit allows us to estimate the potential energy of two dipoles as

$$
\begin{align*}
\frac{\mu}{(r+\epsilon)^{2}}-\frac{\mu}{(r-\epsilon)^{2}} & =-\mu \frac{(r+\epsilon)^{2}-(r-\epsilon)^{2}}{(r+\epsilon)^{2}(r-\epsilon)^{2}}  \tag{3.10}\\
& =-\mu \frac{4 r \epsilon}{(r+\epsilon)^{2}(r-\epsilon)^{2}} \approx-4 \mu \epsilon r^{-3}=-2 \mu^{2} r^{-3} .
\end{align*}
$$

The negative sign indicates that there is an attraction between the two dipoles in the configuration Figure 3.1 .

The electric force field $\mathbf{F}$ is the gradient of the potential $\nabla V$. For $V$ defined by (3.7), only the $x$-component of $\nabla V$ is non-zero along the $x$-axis, by symmetry. Differentiating (3.7), we find that for $r \gg \epsilon$ along the $x$-axis,

$$
\begin{align*}
F_{x}(r, 0,0) & =-(r-\epsilon)^{-2}+(r+\epsilon)^{-2} \\
& =\frac{-(r+\epsilon)^{2}+(r-\epsilon)^{2}}{(r-\epsilon)^{2}(r+\epsilon)^{2}}=\frac{-4 \epsilon r}{(r-\epsilon)^{2}(r+\epsilon)^{2}}  \tag{3.11}\\
& \approx-4 \epsilon r^{-3}=-2 \mu r^{-3}
\end{align*}
$$

Draft: February 28, 2008, do not distribute


Figure 3.2: Parallel dipole-dipole configuration.

The attractive force experienced by a dipole displaced on the $x$-axis at a distance $r$, with a positive charge at $(r+\epsilon, 0,0)$ and a negative charge at $(r-\epsilon, 0,0)$, is thus (asymptotically)

$$
\begin{align*}
-\frac{2 \mu}{(r+\epsilon)^{3}}+\frac{2 \mu}{(r-\epsilon)^{3}} & =2 \mu \frac{(r+\epsilon)^{3}-(r-\epsilon)^{3}}{(r+\epsilon)^{3}(r-\epsilon)^{3}}  \tag{3.12}\\
& =2 \mu \frac{6 r^{2} \epsilon+2 \epsilon^{3}}{(r+\epsilon)^{3}(r-\epsilon)^{3}} \approx 6 \mu^{2} r^{-4},
\end{align*}
$$

which is equal to the derivative of the potential (3.10) as we would expect.

### 3.3.2 Parallel dipole-dipole interactions

In the parallel orientation, the base dipole has a positive charge at $(0, \epsilon, 0)$ and a negative charge at $(0,-\epsilon, 0)$; the other dipole is displaced on the $x$-axis at a distance $r$ : a positive charge at $(r,-\epsilon, 0,0)$ and a negative charge at $(r,+\epsilon, 0,0)$ (cf. Figure 3.2).

The potential in the $(x, y)$-plane due to the base dipole at a distance $r$ along the $x$-axis is

$$
\begin{equation*}
V(x, y)=\frac{1}{\sqrt{(y-\epsilon)^{2}+x^{2}}}-\frac{1}{\sqrt{(y+\epsilon)^{2}+x^{2}}} \tag{3.13}
\end{equation*}
$$

The potential energy of a dipole displaced on the $x$-axis at a distance $r$, with a positive charge at $(r,-\epsilon, 0)$ and a negative charge at $(r, \epsilon, 0)$, is thus

$$
\begin{align*}
\left(\frac{1}{\sqrt{(2 \epsilon)^{2}+r^{2}}}-\frac{1}{r}\right) & -\left(\frac{1}{r}-\frac{1}{\sqrt{(2 \epsilon)^{2}+r^{2}}}\right)=-2\left(\frac{1}{r}-\frac{1}{\sqrt{(2 \epsilon)^{2}+r^{2}}}\right) \\
& =-2 \frac{\sqrt{(2 \epsilon)^{2}+r^{2}}-r}{r \sqrt{(2 \epsilon)^{2}+r^{2}}}=-2 \frac{\sqrt{(2 \epsilon / r)^{2}+1}-1}{r \sqrt{(2 \epsilon / r)^{2}+1}}  \tag{3.14}\\
& \approx-\frac{(2 \epsilon / r)^{2}}{r}=-\mu^{2} r^{-3} .
\end{align*}
$$

Thus the potential energy of the parallel orientation is only half of the single-file orientation.
The potential $V(x, y)$ in (3.13) vanishes when $y=0$. Therefore, its derivative along the $x$-axis also vanishes: $\frac{\partial V}{\partial x}(r, 0)=0$. However, this does not mean that there is no attractive force between


Figure 3.3: General $\theta$-dependent dipole-dipole configuration.
the dipoles, since (by symmetry) $\frac{\partial V}{\partial x}(r, \pm \epsilon)= \pm f(\epsilon, r)$. Thus the attractive force is equal to $2 f(\epsilon, r)$. For completeness, we compute the expression $f(\epsilon, r)$ :

$$
\begin{equation*}
\frac{\partial V}{\partial x}(x, y)=\frac{-x}{\left((y-\epsilon)^{2}+x^{2}\right)^{3 / 2}}+\frac{x}{\left((y+\epsilon)^{2}+x^{2}\right)^{3 / 2}} \tag{3.15}
\end{equation*}
$$

for general $y$. Choosing $y= \pm \epsilon$, (3.15) simplifies to

$$
\begin{align*}
\frac{\partial V}{\partial x}(r, \pm \epsilon) & =\mp r^{-2} \pm \frac{r}{\left((2 \epsilon)^{2}+r^{2}\right)^{3 / 2}}=\mp r^{-2}\left(1-\frac{1}{\left((\mu / r)^{2}+1\right)^{3 / 2}}\right) \\
& =\mp \frac{\left((\mu / r)^{2}+1\right)^{3 / 2}-1}{r^{2}\left((\mu / r)^{2}+1\right)^{3 / 2}} \approx \mp \frac{3 \mu^{2}}{2 r^{4}} \tag{3.16}
\end{align*}
$$

for large $r / \epsilon$. The net force of the field (3.16) on the two oppositely charged particles on the right side of Figure 3.2 is thus $3 \mu^{2} r^{-4}$, consistent with what we would find by differentiating (3.14) with respect to $r$.

The electric force field in the direction of the second dipole (that is, the $y$-axis) is

$$
\begin{equation*}
\frac{\partial V}{\partial y}(r, y)=\frac{\epsilon-y}{\left((y-\epsilon)^{2}+r^{2}\right)^{3 / 2}}+\frac{\epsilon+y}{\left((y+\epsilon)^{2}+r^{2}\right)^{3 / 2}} \tag{3.17}
\end{equation*}
$$

At a distance $r \gg \epsilon$ along the $x$-axis, this simplifies to

$$
\begin{equation*}
\frac{\partial V}{\partial y}(r, \pm \epsilon)=\frac{\mu}{\left(\mu^{2}+r^{2}\right)^{3 / 2}} \approx \mu r^{-3} \tag{3.18}
\end{equation*}
$$

for large $r / \epsilon$. Although this appears to be a force in the direction of the dipole, the opposite charges on the dipole on the right side of Figure 3.2 cancel this effect. So there is no net force on the dipole in the direction of the $y$-axis.

### 3.3.3 Dipole stability

Only the single-file dipole orientation is stable with respect to perturbations. This can be seen as follows. Suppose the dipoles are arranged along the $x$-axis as above but that they are both tilted away from the $x$-axis at an angle $\theta$, as shown in Figure 3.3. Define $\theta$ so that $\theta=0$ (and $\theta=\pi$ ) is


Figure 3.4: Potential energy variation $v(\rho, \theta)$ as defined in (3.22) (vertical axis) of dipoles as a function of $\theta$ (horizontal axis) for the configurations shown in Figure 3.3 for $\rho=0.02$ (top), $0.05,0.1,0.2$ (bottom), where $\rho$ is defined in (3.21).
the single-file dipole configuration and $\theta=\pi / 2$ is the parallel configuration. Thus one dipole has a positive charge at $\epsilon(\cos \theta, \sin \theta, 0)$ and a negative charge at $-\epsilon(\cos \theta, \sin \theta, 0)$. The other dipole is displaced on the $x$-axis at a distance $r$ : a positive charge at $(r+\epsilon \cos \theta,-\epsilon \sin \theta, 0)$ and a negative charge at $(r-\epsilon \cos \theta, \epsilon \sin \theta, 0)$.

The potential at the point $(x, y, 0)$ due to the rotated base dipole is

$$
\begin{equation*}
V(x, y)=\frac{1}{\sqrt{(x-\epsilon \cos \theta)^{2}+(y-\epsilon \sin \theta)^{2}}}-\frac{1}{\sqrt{(x+\epsilon \cos \theta)^{2}+(y+\epsilon \sin \theta)^{2}}} \tag{3.19}
\end{equation*}
$$

Therefore the potential energy of the second rotated dipole, with a positive charge at $(r+\epsilon \cos \theta,-\epsilon \sin \theta, 0)$ and and a negative charge at $(r-\epsilon \cos \theta, \epsilon \sin \theta, 0)$, is thus

$$
\begin{align*}
& V(r, \theta)=\frac{1}{\sqrt{r^{2}+(2 \epsilon \sin \theta)^{2}}}-\frac{1}{r+2 \epsilon \cos \theta}-\left(\frac{1}{r-2 \epsilon \cos \theta}-\frac{1}{\sqrt{r^{2}+(2 \epsilon \sin \theta)^{2}}}\right) \\
&=\frac{2}{\sqrt{r^{2}+(2 \epsilon \sin \theta)^{2}}}-\frac{1}{r+2 \epsilon \cos \theta}-\frac{1}{r-2 \epsilon \cos \theta} \\
&=\frac{2}{\sqrt{r^{2}+(2 \epsilon \sin \theta)^{2}}}-\frac{2 r}{r^{2}-(2 \epsilon \cos \theta)^{2}}  \tag{3.20}\\
&=\frac{2}{r}\left(\frac{1}{\sqrt{1+\rho \sin ^{2} \theta}}-\frac{1}{1-\rho \cos ^{2} \theta}\right):=\frac{2}{r} v(\rho, \theta),
\end{align*}
$$

where the (nondimensional) parameter $\rho$ is defined by

$$
\begin{equation*}
\rho=(2 \epsilon / r)^{2} . \tag{3.21}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute


Figure 3.5: Single-file dipole-dipole configuration with different dipole strengths.

This expression

$$
\begin{equation*}
v(\rho, \theta)=\frac{1}{\sqrt{1+\rho \sin ^{2} \theta}}-\frac{1}{1-\rho \cos ^{2} \theta} \tag{3.22}
\end{equation*}
$$

in (3.20) has a maximum when $\theta=0$ and a minimum when $\theta=\pi / 2$. A plot of $v$ in (3.22) is shown in Figure 3.4 for various values of $\rho$. When $\rho$ is small, the expression (3.22) tends to the limit

$$
\begin{align*}
v(\rho, \theta) & \approx \frac{1}{1+\frac{1}{2} \rho \sin ^{2} \theta}-\frac{1}{1-\rho \cos ^{2} \theta}  \tag{3.23}\\
& \approx\left(1-\frac{1}{2} \rho \sin ^{2} \theta\right)-\left(1+\rho \cos ^{2} \theta\right)=-\frac{1}{2} \rho\left(1+\cos ^{2} \theta\right)
\end{align*}
$$

Of course, what we have presented is only an indication of the stability and energy minimum of the single-file dipole configuration. We leave a complete proof as Exercise 3.8.

### 3.3.4 Different dipoles

So far, we considered dipoles with identical charges and charge distributions (separations). Here we consider a single-file configuration as in Figure 3.1, but with the dipole on the right consisting of charges $\pm q$ separated by a distance $\delta$, as depicted in Figure 3.5. We consider the potential energy of the right-hand dipole in the potential field (3.7) of the left dipole. Similar to (3.10), we find

$$
\begin{equation*}
\frac{\mu q}{(r+\delta)^{2}}-\frac{\mu q}{(r-\delta)^{2}}=-\mu q \frac{4 r \delta}{(r+\delta)^{2}(r-\delta)^{2}} \approx-4 \mu q \delta r^{-3}=-2 \mu \nu r^{-3} \tag{3.24}
\end{equation*}
$$

where $\nu=2 q \delta$ is the strength of the dipole on the right. Notice that the expression (3.24) is symmetric in the two dipole strengths $\mu$ and $\nu$.

## 3.4 van der Waals forces

Many of the electric forces we consider are induced rather than direct. The best known of these are called van der Waals forces, although this term covers a range of forces known by other names. Keesom forces, which we covered in Section ??, are often included in this group, but we will see that there is a qualitative difference in behavior. One prominent web site went as far as to say "all intermolecular attractions are known collectively as Van der Waals forces" but this seems a bit extreme.

Debye forces and London dispersion forces [93] involve induced dipole-dipole interactions, which we will study using the results derived in Section ??. The most significant example is the London
(a)

(b)


Figure 3.6: A dipole (left) inducing a dipole in a polarizable molecule (right). The upper configuration (a) shows the dipole and polarizable molecule well separated, and the lower configuration (b) shows them closer, with the molecule on the right now polarized.
dispersion force 93 which results from both dipoles being induced. This often takes the form of a symmetry breaking, and we give a derivation of the dependence of the magnitude of the induced dipole on distance in Section 16.5.2.

The Lennard-Jones model

$$
\begin{equation*}
V(r)=c r^{-12}-c^{\prime} r^{-6} \tag{3.25}
\end{equation*}
$$

is commonly used to model van der Waals interactions. The attractive potential $r^{-6}$ is a precise result of the interaction of a fixed dipole and an induced dipole, which we derive in Section 3.5. In Section 3.5.2, we consider a second, but defer to Section 16.5 .2 a detailed discussion. The repulsive term $r^{-12}$ is a convenient model, whereas other terms are more accurate [46] (cf. Section 16.7).

We cover van der Waals forces in detail here to clarify that they are electrostatic in nature, and not some new or different type of force. As such they are susceptible to modulation by solvent dielectric behavior.

### 3.5 Induced dipoles

Dipoles can be induced in two ways. Fixed dipoles, such as water molecules, induce a dipole in any polarizable material. Such interactions give rise to what are frequently called Debye forces 93]. More subtly, two polarizable molecules can induce dipoles in each other, via what are called London dispersion forces 93].

### 3.5.1 Debye forces

If a polarizable molecule is subjected to an electric field of strength $\mathbf{F}$, then it is reasonable to expect that an induced dipole $\mu_{i}$ will result, given by

$$
\begin{equation*}
\mu_{i} \approx \alpha \mathbf{F} \tag{3.26}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
for small $\mathbf{F}$, where $\alpha$ is the polarizability. This is depicted visually in Figure 3.6, where the upper configuration (a) shows the dipole and polarizable molecule well separated, and the lower configuration (b) shows them closer, with the molecule on the right now polarized.

In general, the electric field $\mathbf{F}$ is a vector and the polarization $\alpha$ is a tensor (or matrix). Also, note that a dipole is a vector quantity: it has a magnitude and direction. In our previous discussion, we considered only the magnitude, but the direction was implicit (the line connecting the two charges). For simplicity, we assume here that $\alpha$ can be represented as a scalar (times the identity matrix), that is, that the polarizability is isotropic. The behavior in (3.26) will be deduced by a perturbation technique for small $\mathbf{F}$ from the concepts in Section 16.5

We can approximate a polarized molecule as a simple dipole with positive and negative charges $\pm q$ displaced by a distance $\delta$, as depicted in Figure 3.5. This takes some justification, but it will be addressed in Chapter 9 , There is ambiguity in the representation in that only the product $q \delta$ matters: $\mu=q \delta$.

We derived in (3.11) that the electric force field due to a fixed dipole $\mu_{f}$ has magnitude

$$
\begin{equation*}
F_{x}=2 \mu_{f} r^{-3} \tag{3.27}
\end{equation*}
$$

where the $x$-axis connects the two charges of the fixed dipole. We assume that the molecule whose dipole is being induced also lies on this axis. By combining (3.26) and (3.27), we conclude that the strength of the induced dipole is

$$
\begin{equation*}
\mu_{i} \approx 2 \alpha \mu_{f} r^{-3} \tag{3.28}
\end{equation*}
$$

From (3.24), we know that the potential energy of the two dipoles is

$$
\begin{equation*}
V(r) \approx-2 \mu_{f} \mu_{i} r^{-3} \approx-4 \alpha \mu_{f}^{2} r^{-6} \tag{3.29}
\end{equation*}
$$

in agreement with the Lennard-Jones model in (3.25).

### 3.5.2 London dispersion forces

Suppose now that we start with two nonpolar, but polarizable, molecules that are well separated. Due to the long range interaction (correlation) of the electron distributions of the two molecules (to be explained in Section (16.5), they can become polarized. To get a sense of what might happen, suppose one of them polarizes first so that it becomes the dipole depicted on the left in Figure 3.6. Then as it approaches the other molecule, it induces a dipole in it. But what if the molecules are identical? Then the induced dipole is the same as the 'fixed' dipole that was in the case of the Debye force: $\mu_{i}=\mu_{f}$. Thus there is only one $\mu$ in the discussion now.

The dipole $\mu$ is induced by the electric field of the other dipole, so that again $\mu \approx \alpha \mathbf{F}$ where $\mathbf{F}$ is the electric field strength and $\alpha$ is the polarizability. The electric strength of the field $\mathbf{F}$ is again given by (3.27): $F_{x}=2 \mu r^{-3}$. But now the electric field strength and the dipole strength are coupled in a new way, and it is not simple to solve this system.

The expression (3.24) remains valid for the potential energy of the induced dipoles:

$$
\begin{equation*}
V \approx-2 \mu^{2} r^{-3} \tag{3.30}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

But how big is the induced dipole $\mu$ in expression (3.30)? We saw in (3.28) that the dipole induced by a fixed dipole has a magnitude that is asymptotic to $r^{-3}$. If such an asymptotic behavior were to hold in the case of doubly induced dipoles, it would lead to an expression for the potential energy of the induced dipoles of the form

$$
\begin{equation*}
V \approx c r^{-9} \tag{3.31}
\end{equation*}
$$

which is quite different from the Lennard-Jones model.
Let us review the arguments used to estimate the magnitude of the dipole induced by a fixed dipole to see where it fails for doubly induced dipoles. It is reasonable to assume that the dipole strength is a monotone function of the induced field $\mathbf{F}$; we used the ansazt that $\mu \approx \alpha \mathbf{F}$ for small $\mathbf{F}$ in the derivation of the $r^{-6}$ dependence of $V$ for a dipole induced by a fixed dipole. But since $\mathbf{F}$ depends on $r$, so must $\mu$ depend on $r$, and this would mean that our expression (3.30) would not be a complete description of the asymptotic behavior of $V$ as a function of $r$, and it would imply that the behavior of $\mathbf{F}=\nabla V$ would go to zero faster than $r^{-3}$. This would imply that $\mu \approx \alpha \mathbf{F}$ would be even smaller. In fact, if we iterate the argument, we would never converge on a finite power of $r$. Let us analyze the argument in more detail.

We used two key equations, namely (3.26) and (3.27), in deriving the expression for $V$ for a dipole induced by a fixed dipole. If we now assume that $\mu=\mu_{i}=\mu_{f}$, the two equations $\mu=\alpha \mathbf{F}$ (in scalar form, $\mu=\alpha F_{x}$ ) and $F_{x}=2 \mu r^{-3}$ can be solved to find

$$
\begin{equation*}
r=\sqrt[3]{2 \alpha} \tag{3.32}
\end{equation*}
$$

Note that this is dimensionally correct, since the units of the polarizability $\alpha$ are the same as volume. Thus using the two equations (3.26) and (3.27) together with the simplification $\mu=\mu_{i}=\mu_{f}$ determines a particular value of $r$, in contradiction to our derivation of an expression valid for various values of $r$. We will see in Section 16.5.3 that this value of $r$ can be interpreted in a mathematical, if not physical, sense.

We will derive in (16.70) a result that confirms the basic dependence of the dipole strength on $r$, namely

$$
\begin{equation*}
\mu \approx c r^{-3} \tag{3.33}
\end{equation*}
$$

where an expression for the constant $c$ will be made explicit. Thus, at least for $r$ sufficiently large, the expression (3.30) appears to be the correct asymptotic behavior.

The above arguments could be interpreted in the following way. For very large $r$, the potential of the induced dipole is very small, comparable to $r^{-9}$. As the induced field $\mathbf{F}$ increases, the polarizability will saturate, and $\mu$ will tend to a limit. For example, this might take a form similar to

$$
\begin{equation*}
\mu(r) \approx \frac{c_{1}}{1+c_{2} r^{-3}} . \tag{3.34}
\end{equation*}
$$

Thus the potential energy varies from (3.31) for large $r$ to $\mathrm{Cr}^{-3}$ for smaller values of $r$. For this reason, using the intermediate exponent of 6 , approximating the behavior of the potential energy by $r^{-6}$, may be a reasonable compromise.

Draft: February 28, 2008, do not distribute

| atom | $\rho$ | $\epsilon$ | $V(\rho / 2)$ | $D$ | $\kappa$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| C (aliphatic) | 1.85 | 0.12 | 476 | 1.54 | 83.1 |
| O | 1.60 | 0.20 | 794 | 1.48 | 33.2 |
| H | 1.00 | 0.02 | 79 | 0.74 | 104.2 |
| N | 1.75 | 0.16 | 635 | 1.45 | 38.4 |
| P | 2.10 | 0.20 | 794 | 1.87 | 51.3 |
| S | 2.00 | 0.20 | 794 | 1.81 | 50.9 |

Table 3.1: Lennard-Jones parameters from AMBER for various atoms involving the van der Waals radius $\rho$ measured in Ångstroms and energy (well depth) $\epsilon$ in kcal/mol. For comparison, covalent bond lengths $D$ and strengths [191] $\kappa$ are given in kcal/mol, together with the repulsion potential energy $V(\rho)$ at the van der Waals radius $\rho$.

### 3.5.3 Lennard-Jones potentials

The van der Waals interactions are often modeled via the Lennard-Jones potential

$$
\begin{equation*}
V(r):=\epsilon\left(\left(\frac{\rho}{r}\right)^{12}-2\left(\frac{\rho}{r}\right)^{6}\right) . \tag{3.35}
\end{equation*}
$$

The minimum of $V$ is at $r=\rho$, with $V(\rho)=-\epsilon$, so we can think of the well depth $\epsilon$ as giving the energy scale. The parameter $\rho$ is called the van der Waals radius, and can be defined as the separation distance at which the force of attraction and repulsion cancel [26]. Typical values for these parameters, from the AMBER force field, are shown in Table 3.1. Note that $V(\rho / 1.2) \approx-3 V(\rho)$, and $V(\rho / 2)=-3968 V(\rho)$, so the repulsion is quite strong in this model.

### 3.6 Hydrophobic interactions

There is another force that is crucial in biology, sometimes said to be more important than even the hydrogen bond force [130]. It is called the hydrophobic force [22], which derives from the hydrophobic effect [234]. This effect is one of the central topics of our study. However, the hydrophobic effect has many manifestations in protein behavior.

There is a simple view of how hydrophobic forces work. There are certain molecules that are hydrophobic (cf. Section 3.6 .2 and Chapter 7), meaning that they repel water. Regions of proteins that have many such molecules, e.g., a protein with a large number of hydrophobic residues on a part of its surface, would tend to prefer association with another such surface to reduce the frustration of having two water-hydrophobe interfaces. It is this simple effect that makes cooking oil form a single blob in water even after it has been dispersed by vigorous stirring.

More precisely, the argument is that the elimination of two hydrophobic surfaces with a water interface is energetically favorable. One could also argue by considering volume changes (cf. Section 4.4.4) since hydrophobic side chains take up more volume in water. Recent results show how a hydrophobic force can arise through a complex interaction between polarizable (e.g., hydrophobic)

Draft: February 28, 2008, do not distribute
molecules and (polar) water molecules [53, 54]. These arguments are compelling, but they suggest a nonspecific interaction. Indeed, hydrophobic attraction leads to nonspecific binding [81].

But there are other kinds of hydrophobic effects as well. We will show that hydrophobicity plays a central role in a number of electrostatic forces by modulating the dielectric effect of water. In addition, water removal can affect the local polar environment, which can modify the nature of covalent bonds.

### 3.6.1 Solvent mediation of electric forces

Some bonds become substantially altered in the presence of water. We have already noted that certain ionic bonds (in table salt) are easily disrupted by water. The main bond holding proteins together is the hydrogen bond, and this bond is extremely susceptible to alteration by water interaction since water molecules can each make four hydrogen bonds themselves. So protein survival depends on keeping the hydrogen bond dry in water 67. More generally, solvent mediation can alter any electrostatic force via dielectric effects (Chapter 17).

One type of solvent effect that is expressed on the quantum level is the rigidity of the peptide bond (Chapter (14) which requires an external field to select one of two resonant states. Such a field can be due to hydrogen bonds with the amide or carbonyl groups, either with other backbone or sidechain groups, or with water. In some situtations, water removal can cause a switch in the resonance state to a flexible mode [66].

Another example of a change of electrical properties resulting from differences in the water environment involves a more gross change. Proteins which penetrate a cell membrane go from a fully solvated environment to one that is largely solvent-free (inside the membrane). We will see that this can be related to a large-scale change in the secondary structure of the protein conformation that has implications for drug delivery 81].

Changes in dielectric properties of the environment can have a substantial impact on any electrical property. But rather than try to address this by a general model, we prefer to introduce the concept by example. We thus begin by looking at one particular example of hydrophobic modulation of the dielectric behavior of water around hydrogen bonds.

### 3.6.2 Dehydrons

In [79, a quantifiable structural motif, called dehydron, was shown to be central to protein-ligand interactions. A dehydron is a defectively 'wrapped' hydrogen bond in a molecular structure whose electrostatic energy is highly sensitive to water exclusion by a third party. Such pre-formed, but underprotected, hydrogen bonds are effectively adhesive, since water removal from their vicinity contributes to their strength and stability, and thus they attract partners that make them more viable.

A review of protein structure and the role of hydrogen bonds will be presented in Chapter 4 , The concept of 'wrapping' of a hydrogen bond is based on the hydrophobic effect [22, 234]. At the simplest level, wrapping occurs when sufficient nonpolar groups $\left(\mathrm{CH}_{n}, n=1,2,3\right)$ are clustered in the vicinity of intramolecular hydrogen bonds, protecting them by excluding surrounding water

Draft: February 28, 2008, do not distribute 28


Figure 3.7: (a) Well wrapped hydrogen bond (b) Underwrapped hydrogen bond.
[77. The concept of wrapping of a hydrogen bond is depicted informally in Figure 3.7. A well wrapped hydrogen bond (Figure 3.7(a)) is surrounded by $\mathrm{CH}_{n}$ groups on all sides, and water is kept away from the hydrogen bond formed between the C-O group of one peptide and the $\mathrm{N}-\mathrm{H}$ group of another peptide (Section 4.1). An underwrapped hydrogen bond (Figure 3.7(b)) allows a closer approach by water to the hydrogen bond, and this tends to disrupt the bond, allowing the distance between the groups to increase and the bond to weaken.

It is possible to identify dehydrons as under wrapped hydrogen bonds (UWHB) by simply counting the number of hydrophobic side chains in the vicinity of a hydrogen bond. This approach is reviewed in Section 7.2, More accurately, a count of all (nonpolar) carbonaceous groups gives a more refined estimate (Section [7.3). However, it is possible to go further and quantify a force associated with dehydrons which provides a more refined measure of the effect geometry [79] of the wrappers (Section 7.5).

We have already seen in Figure 2.1] that dehydrons are a sensitive measure of protein differences. At the structural level, a significant correlation can be established between dehydrons and sites for protein complexation (Chapter (7). The HIV-1 capsid protein P24 complexed with antibody FAB25.3 provides a dramatic example [79].

### 3.6.3 Dynamics of dehydrons

The extent of wrapping changes the nature of hydrogen bond. Hydrogen bonds that are not protected from water do not persist [49]. Figure 5 of [49] shows the striking difference of water residence times for well wrapped and underwrapped hydrogen bonds. Private communication with the authors of 49 have confirmed that there is a marked difference as well in the fluctuations of the hydrogen bonds themselves. Under wrapped hydrogen bond lengths are larger (on average) than well wrapped hydrogen bonds. More strikingly, the distributions of bond lengths as shown in Fig-


Figure 3.8: Dynamics of water near hydrogen bonds, reproduced from Fig. 5 in 49]. (A) Hydrogen bond (H3) is well wrapped. (B) Hydrogen bond (H1) is underwrapped.
ure 3.9 are quite different, confirming our prediction based on Figure 3.7 that the coupling of the hydrogen bond characteristics with the water environment would be different.

The H-bond R208-E212 depicted in Fig. 5(A) 49 is well wrapped whereas V189-T193 depicted in Fig. 5(B) is a dehydron (see Fig 3a in [74] page 6448). Well-wrapped hydrogen bonds are visited by fewer water molecules but have longer-lasting water interactions (due to the structuring effect of the hydrophobes), whereas the behavior of dehydrons is more like that of bulk water: frequent re-bonding with different water molecules [49].

The long residence time of waters around a well-wrapped hydrogen bond would seem to have two contributing factors. On the one hand, the water environment is structured by the hydrophobic barrier, so the waters have reduced options for mobility: once trapped they tend to stay. But also, the polar effect of the hydrogen bond which attracts the water is more stable, thus making the attraction of water more stable. With a dehydron, both of these effects go in the opposite direction. First of all, water is more free to move in the direction of the hydrogen bond. Secondly, the fluctuation of the amide and carbonyls comprising the hydrogen bond contribute to a fluctuating electrostatic environment. The bond can switch from the state depicted in Figure 3.7(b) when water is near, to one more like that depicted in Figure 3.7(a) if water molecules move temporarily away. More precisely, the interaction of the bond strength and the local water environment becomes a strongly coupled system for an underwrapped hydrogen bond, leading to increased fluctuations. For a well wrapped hydrogen bonds, the bond strength and water environment are less strongly coupled.


Figure 3.9: Distribution of bond lengths for two hydrogen bonds formed in a structure of the sheep prion [49]. The horizontal axis is measured in nanometers, whereas the vertical axis represents numbers of occurrences taken from a simulation with 20,000 data points with bin widths of 0.1 Ångstrom. The distribution for the well-wrapped hydrogen bond (H3) is depicted with a solid line, whereas the distribution for the underwrapped hydrogen bond (H1) is depicted with a dashed line.

The distance distribution for under-wrapped hydrogen bonds can be interpreted as reflecting a strong coupling with the thermal fluctuations of the solvent. Thus we see a Boltzmann-type distribution for the under-wrapped hydrogen bond distances in Figure 3.9, It is natural to expect the mean distances in this case to be larger than the mean distances for the underwrapped case, but the tails of the distribution are at first more confusing. The distribution in the underwrapped case exhibit a Gaussian-like tail (that is, exponential of the distance squared), whereas the wellwrapped case decays more slowly, like a simple exponential. Thus the well-wrapped hydrogen bond is sustaining much larger deviations, even though the typical deviation is much smaller than in the underwrapped case. To explain how this might occur, we turn to a simulation with a simple model.

### 3.6.4 Simulated dynamics

The data in Figure 3.9 can be interpreted via a simulation which is depicted in Figure 3.10. This figure records the distribution of positions for a random walk subject to a restoring force defined by

$$
\begin{equation*}
x_{i+1}=x_{i}+\Delta t\left(f_{i}+\phi\left(x_{i}\right)\right) \tag{3.36}
\end{equation*}
$$

with $f_{i}$ drawn randomly from a uniform distribution on $[-0.5,0.5]$, and with $\phi$ being a standard Lennard-Jones potential

$$
\begin{equation*}
\phi(x)=(0.1 / x)^{12}-(0.1 / x)^{6} . \tag{3.37}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute


Figure 3.10: Simulation of a random walk with a restoring force. Shown is the distribution of values $x_{i}$ defined in (3.36) for $10^{5}$ time steps $i$, starting with $x_{1}=0.1$, scaled by a factor of $10^{-3}$. Also shown is a graph of $\phi+0.03$ where $\phi$ is the potential (3.36). The dot-dashed horizontal line provides a reference axis to facilitate seeing where $\phi$ is positive and negative. The + 's indicate the part of the distribution exhibiting an exponential decay; the dashed line is a least-squares fit to the logarithm of these distribution values. The distribution has been scaled by a factor of $10^{-3}$ so that it fits on the same plot with $\phi$.

The particular time step used in Figure 3.10 is $\Delta t=0.02$; the simulation was initiated with $x_{1}=0.1$ and carried out for $10^{5}$ steps.

The simulation (3.36) represents a system that is forced randomly with a restoring force back to the stationary point $x=0.1$, quantified by the potential $\phi$ in (3.37). Such a system exhibits a distribution with an exponential decay, as verified in Figure 3.10 by comparison with a least-squares fit of the logarithm of the data to a straight line.

### 3.6.5 Stickiness of dehydrons

Desolvation of an underwrapped hydrogen bond can occur when a ligand binds nearby, as depicted in Figure 3.11. The removal of water lowers the dielectric and correspondingly strengthens the hydrogen bond. The resulting change in energy due to the binding effectively means that there is a force of attraction for a dehydron. This is explained in more detail in Chapter 8

### 3.7 Exercises

Exercise 3.1 Show that the approximation in (3.7) is $96 \%$ accurate for $r \geq 5 \epsilon$.


Figure 3.11: Cartoon showing dehydration due to ligand binding and the resulting strengthening of an underwrapped hydrogen bond.

Exercise 3.2 Pour salt into a glass of water and watch what happens to the salt. Take a small amount out and put it under a microscope to see if the picture stays the same.

Exercise 3.3 Prove that (3.10) is still correct if we use the exact form in (3.7) instead of the approximation $V(r) \approx \mu r^{-2}$.

Exercise 3.4 Prove that (3.12) is still correct if we use the exact form in (3.11), $F_{x}(r, 0,0)=$ $-4 \epsilon r(r-\epsilon)^{-2}(r+\epsilon)^{-2}$, instead of the approximation $F_{x}(r, 0,0) \approx-2 \mu r^{-3}$.

Exercise 3.5 Pour cooking oil into a glass of water and stir it vigorously until the oil is well dispersed. Now wait and watch as the oil droplets coalesce. Do the individual droplets retain any sort of discrete form? Or does the hydrophobic force just create a blob in the end?

Exercise 3.6 Consider the expression in (3.20). Prove that, for any $\rho<1$, it has a maximum when $\theta=0$ and a minimum when $\theta=\pi / 2$.

Exercise 3.7 Pour salt into a glass of water and stir it until it dissolves. Now also add some oil to the water and stir it until small droplets form. Look at the surface of the oil droplets and see if you can see salt crystals that have reformed due to the change in electrostatic environment there. This might best be done on a slide beneath a microscope objective.

Exercise 3.8 Prove that the single-file dipole configuration is stable and an energy minimum. (Hint: derive a formula for the general orientation of two dipoles in three dimensions, cf. Figure 2.2 in [118]. This can be done with one distance parameter and three angular parameters.)

Exercise 3.9 Describe the orientation of the dipoles that corresponds to $\theta=2 \pi$ in Figure 3.4.

## Chapter 4

## Protein basics

Proteins are sequences of amino acids which are covalently bonded along a "backbone." Proteins of biological significance fold into a three-dimensional structure by adding hydrogen bonds between carbonyl and amide groups on the backbone of different amino acids. In addition, other bonds, such as a salt bridge (Section 4.2.1) or a disulfide bond (Section 4.2.2), can form between particular amino acids (Cysteine has sulfur atoms in its sidechain). However, the hydrogen bond is the primary mode of structure formation in proteins.

It is not our intention to provide a complete introduction to the structure of proteins. Instead, we suggest consulting texts [47, 196] for further information. Moreover, we suggest acquiring a molecular modeling set so that accurate three-dimensional models can be constructed. In addition, it will be useful to become familiar with a graphical viewer for PDB files (even the venerable 'rasmol' would be useful). We present some essential information and emphasize concepts needed later or ones that may be novel.

### 4.1 Chains of amino acid residues

Proteins are chains of amino acid residues whose basic unit can be considered to be the peptide group shown in Figure 4.1. The trans form (a) of the peptide bond is the most common state, but the cis form (b) has a small but significant occurrence [107, 186].

The chain is repeated many times, up to many hundreds of backbone $C_{\alpha}$ carbons. The residues $\mathrm{R}^{i}$ come from the different amino acids and will be described in more detail in Section 4.1.2, See Figure 4.4 for some of the smaller residues. A cartoon of a peptide sequence is depicted in Figure 4.2,

The peptide chain is joined at the double bond indicated between the N and the O in Figure 4.1 , Thus we refer to the coordinates of the nitrogen and hydrogen as $N^{i+1}$ and $H^{i+1}$ and to the coordinates of the oxygen and carbon as $O^{i}$ and $C^{i}$.

At the ends of the chain, things are different. The N-terminus, or $\mathbf{N}$-terminal end, has an $\mathrm{NH}_{2}$ group instead of just N , and nothing else attached. In the standard numbering scheme, this is the beginning of the chain. The C-terminus, or $\mathbf{C}$-terminal end, has a $\mathrm{CO}_{2} \mathrm{H}$ group instead of just CO, and nothing else attached. In the standard numbering scheme, this is the end of the chain.

(a)

(b)

Figure 4.1: The rigid state of the peptide bond: (a) trans form, (b) cis form. The double bond between the central carbon and nitrogen keeps the peptide bond planar. Compare Figure 14.1 .


Figure 4.2: Cartoon of a peptide sequence where all of the peptides are in the trans form (cf. Figure 4.1). The small boxes represent the C-alpha carbons, the arrow heads represent the amide groups NH, the arrow tails represent the carbonyl groups CO, and the thin rectangular boxes are the double bond between the backbone C and N . The different residues are indicated by R's. The numbering scheme is increasing from left to right, so that the arrow formed by the carbonyl-amide pair points in the direction of increasing residue number. The three-dimensional nature of the protein is left to the imagination, but in particular where the arrow heads appear to be close in the plane of the figure they would be separated in the direction perpendicular to the page.
(a)

(b)

(c)


Figure 4.3: The hydrogen bond (dashed line) configuration in (a) $\alpha$-helix, (b) antiparallel $\beta$-sheet, and (c) parallel $\beta$-sheet. The amide ( $\mathrm{N}-\mathrm{H}$ ) groups are depicted by arrow heads and the carbonyl (O-C) groups are depicted by arrow tails.

### 4.1.1 Hydrogen bonds and secondary structure

The representation of proteins as a linear sequence of amino acid residues depicted in Figure 4.2 is called the primary structure. Proteins have a hierarchy of structure, the next being secondary structure consisting of two primary types: alpha-helices and beta-sheets (a.k.a., $\alpha$-helices and $\beta$-sheets).

Alpha helices are helical arrangements of the subsequent peptide complexes with a distinctive hydrogen bond arrangement between the amide (NH) and carbonyl (OC) groups in peptides separated by $k$ steps in the sequence, where primarily $k=4$ but with $k=3$ and $k=5$ also occurring less frequently. The hydrogen bond arrangement is depicted in Figure 4.3(a) between two such peptide groups.

Beta sheets represent different hydrogen bond arrangements, as depicted in Figure 4.3 (b) is the anti-parallel arrangement and (c) is the parallel. Both structures are essentially flat, in contrast to the helical structure in (a).

### 4.1.2 Taxonomies of amino acids

There are many ways that one can categorize the amino acid sidechains of proteins. We are mainly interested in protein interactions, so we will focus initially on a scale that is based on interactivity. We postpone until Chapter [6 a full explanation of the rankings, but suffice it to say that we rank amino acid sidechains based on their likelihood to be found in a part of the protein surface that is involved in an interaction.

In the following, we will use the standard terminology for the common twenty amino acids ${ }^{1}$ In Table 4.1 we recall the naming conventions and the RNA codes for each residue. Complete descriptions of the sidechains for the amino acids can be found in Figures 4.4 -4.7.

In Table 4.2, we present some elements of a taxonomy of sidechains. We give just two descriptors of sidechains, but even these are not completely independent. For example, all the hydrophilic

[^0]Draft: February 28, 2008, do not distribute


Figure 4.4: Periodic table of amino acid sidechains (residues). Not shown is the $C_{\alpha}$ carbon (see Figure (4.1), located at the top of the residue where the name appears. The smallest, and most likely to be involved in protein-ligand interactions, ordered from the left (asparagine).

| Full name of <br> amino acid | three <br> letter | single <br> letter | The various RNA codes <br> for this amino acid |
| :---: | :---: | :---: | :--- |
| alanine | Ala | A | GCU, GCC, GCA, GCG |
| arginine | Arg | R | CGU, CGC, CGA, CGG, AGA, AGG |
| asparagine | Asn | N | AAU, AAC |
| aspartate | Asp | D | GAU, GAC |
| cysteine | Cys | C | UGU, UGC |
| glutamine | Gln | Q | CAA, CAG |
| glutamate | Glu | E | GAA, GAG |
| glycine | Gly | G | GGU, GGC, GGA, GGG |
| histidine | His | H | CAU, CAC |
| isoleucine | Ile | I | AUU, AUC, AUA |
| leucine | Leu | L | UUA, UUG, CUU, CUC, CUA, CUG |
| lysine | Lys | K | AAA, AAG |
| methionine | Met | M | AUG |
| phenylalanine | Phe | F | UUU, UUC |
| proline | Pro | P | CCU, CCC, CCA, CCG |
| serine | Ser | S | UCU, UCC, UCA, UCG, AGU, AGC |
| threonine | Thr | T | ACU, ACC, ACA, ACG |
| tryptophan | Trp | W | UGG |
| tyrosine | Tyr | Y | UAU, UAC |
| valine | Val | V | GUU, GUC, GUA, GUG |
| stop codons |  |  | UAA, UAG, UGA |

Table 4.1: Amino acids, their (three-letter and one-letter) abbreviations and the RNA codes for them. For completenes, the "stop" codons are listed on the last line.

| Full name <br> of residue | three <br> letter | single <br> letter | water <br> preference | sidechain <br> type | nonpolar <br> $C H_{n}$ groups | intrinsic <br> $p K_{a}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Alanine | Ala | A | phobic | small | 1 | NA |
| Arginine | Arg | R | amphi | positive | 2 | 12 |
| Asparagine | Asn | N | philic | polar | 1 | NA |
| Aspartate | Asp | D | philic | negative | 1 | $3.9-4.0$ |
| Cysteine | Cys | C | philic | polar | 1 | $9.0-9.5$ |
| Glutamine | Gln | Q | amphi | polar | 2 | NA |
| Glutamate | Glu | E | amphi | negative | 2 | $4.3-4.5$ |
| Glycine | Gly | G | NA | tiny | 0 | NA |
| Histidine | His | H | philic | positive | 1 | $6.0-7.0$ |
| Isoleucine | Ile | I | phobic | aliphatic | 4 | NA |
| Leucine | Leu | L | phobic | aliphatic | 4 | NA |
| Lysine | Lys | K | amphi | positive | 3 | $10.4-11.1$ |
| Methionine | Met | M | amphi | polar | 3 | NA |
| Phenylalanine | Phe | F | phobic | aromatic | 7 | NA |
| Proline | Pro | P | phobic | cyclic | 2 | NA |
| Serine | Ser | S | philic | polar | 0 | NA |
| Threonine | Thr | T | amphi | polar | 1 | NA |
| Tryptophan | Trp | W | amphi | aromatic | 7 | NA |
| Tyrosine | Tyr | Y | amphi | aromatic | 6 | $10.0-10.3$ |
| Valine | Val | V | phobic | aliphatic | 3 | NA |

Table 4.2: A taxonomy of amino acids. The code for water interaction is: phobic, hydrophobic; philic, hydrophilic; amphi, amphiphilic. Values of $p K_{a}$ for ionizable residues are taken from Table 1.2 of [47].
residues are either charged or polar. But the converse of this relationship (that is, hydrophobic implies not charged or polar) is false, and there are few such general correlations. For example, the aromatic residues are among the most hydrophobic even though they are polar, cf. Section 4.5.5.

We focus here on the properties of individual sidechains, but these properties alone do not determine protein structure: the context is essential. Studying pairs of sidechains that are interacting in some way (e.g., ones that appear sequentially) gives a first approximation of context.

### 4.1.3 Wrapping of hydrogen bonds

A key element of protein structure is the protection of hydrogen bonds from water attack. A different taxonomy amino acids can be based on their role in the protection of hydrogen bonds. We will see in Chapter 6 that this correlates quite closely with the propensity to be at an interface.

Some hydrogen bonds are simply buried in the interior of a protein. Others are near the surface and potentially subject to water attack. These can only be protected by the sidechains of other


Figure 4.5: Periodic table of amino acid sidechains (sidechains only shown). Not shown is the $C_{\alpha}$ carbon (see Figure 4.1), located at the top of the residue where the name appears. The middle ground in terms of interactivity.
nearby amino acids. Such protection is provided by the hydrophobic effect. We discuss in Chapter 19 some details regarding the hydrophobic effect [22, [234], but suffice it to say that a key element has to do with the fact that certain non-polar groups, such as the carbonaceous groups $\mathrm{CH}_{n}$ ( $n=1,2,3$ ), tend to repel polar molecules like water. They are non-polar and thus do not attract water strongly, and moreover, they are polarizable and thus damp nearby water fluctuations. Such carbonyl groups are common in amino acid sidechains; Val, Leu, Ile, Pro, and Phe have only such carbonaceous groups. We refer to the protection that such sidechains offer as the wrapping of hydrogen bonds. For reference, the number of nonpolar $\mathrm{CH}_{n}$ groups for each residue is listed in Table 4.2

The standard thinking about sidechains has been to characterize them as being hydrophobic or hydrophilic or somewhere in between. Clearly a sidechain that is hydrophobic will repel water and thus protect anything around it from water attack. Conversely, a sidechain that is hydrophilic will attract water and thus might be complicit in compromising an exposed hydrogen bond. In some taxonomies [196], Arg, Lys, His, Gln, and Glu are listed as hydrophilic. However, we will see that they are indeed good wrappers. On the other hand, Ala is listed as hydrophobic and Gly, Ser, Thr, Cys and others are often listed as "in between" hydrophobic and hydrophilic. And we will see that they are among the most likely to be near underwrapped hydrogen bonds. This is not surprising since they are both polar (see Section 4.5.1) and have a small number of carbonaceous groups.

What is wrong with a simple philic/phobic dichotomy of amino acids is that the "call" of philic versus phobic is made primarily based on the final group in the sidechain (the bottom in Figures 4.4 4.7). For example, Lys is decreed to be hydrophilic when the bulk of its sidechain is a set of four carbonaceous groups. What is needed is a more complete picture of the role of all the groups in the entire sidechain. This requires a detailed understanding of this role, and in a sense that is a major object of this monograph. Thus it will require some in depth analysis and comparison


Figure 4.6: Periodic table of amino acid sidechains (sidechains only shown). Not shown is the $C_{\alpha}$ carbon (see Figure 4.1), located at the top of the residue where the name appears. The less likely to be interactive.
with data to complete the story. However in the subsequent chapters this will be done, and it will appear that one can provide at least a broad classification, if not a linear ordering, of amino acid sidechains based on either their ability or propensity to wrap (or not) exposed hydrogen bonds or other electronic bonds.

The ordering of the most interactive proteins is based on a statistical analysis which is described in more detail in Chapter 6. We will also see there that these are likely to be associated with underwrapped hydrogen bonds. On the other hand, it is relatively easy to predict the order for good wrappers based on counting the number of carbonaceous groups. There is not a strict correlation between interactivity and bad wrapping, but a significant trend exists.

### 4.2 Special bonds

In addition to the covalent bonds of the backbone and the ubiquitous hydrogen bonds in proteins, there are two other bonds that are significant.

### 4.2.1 Salt Bridges

Certain sidechains are charged, as indicated in Table 4.2. Depending on the pH level, His may or may not be positively charge, but both Arg and Lys can be considered positively charged in most biological environments. Similarly, Asp and Glu are typically negatively charged. When sidechains of opposite charge form an ionic bond (Section 3.1.2) in a protein, it is called a salt bridge. Thus there are four (or six, depending on His) possible salt bridges.

Unmatched charged residues are often found on the surface of a protein, but inside a protein core they would not likely prevail as they could instead be solvated by several water molecules.

Draft: February 28, 2008, do not distribute


Figure 4.7: Periodic table of amino acid sidechains. The amino acids (sidechains only shown) least likely to be involved in interactions. Not shown is the $C_{\alpha}$ carbon (see Figure 4.1), located at the top of the residue where the name appears.

### 4.2.2 Disulfide bonds

Proteins are also held together by disulfide bonds or disulfide bridges which are bonds which form between two sulfurs on cysteines. Specifically, the hydrogens attached to the sulfur atom on the two Cys sidechains are liberated, and a covalent bond forms between the two sulfur atoms. This is a much stronger bond than a hydrogen bond, but it is also much more specialized. It appears frequently in neurotoxins [99, 183]. These proteins would be highly disordered without the disulfide bridges.

Disulfide bonds can also form between two separate proteins to form a larger system. This occurs in insulin and in antibodies.

### 4.3 Post-translational modifications

Proteins are not quite so simple as the protein sequence might imply. The term post-translational modification means changes that occur after the basic sequence has been set. Modifications (glycosylation, phosphorylation, etc.) add groups to sidechains and change the function of the resulting protein. A change in pH can cause the ends of some sidechains to be modified, as we discuss in Section 4.6.

Phosphorylation occurs by liberating the hydrogen atom in the OH group of Serine, Threonine and Tyrosine, and adding a complex of phosphate groups (see Section 13.1 for illustrations).

Phosphorylation can be inhibited by the presence of wrappers. Serine phosphorylates ten times more often than Tyrosine, even though the benzene ring presents the OH group further from the backbone.

Phosphorylation is expressed in PDB files by using a non-standard amino acid code, e.g., PTR for
phosphotyrosine (phosphorylated tyrosine) and TPO for phosphothreonine (phosphorylated threonine).

### 4.4 Mechanical properties of proteins

The Protein Data Bank (PDB) supports a simple mechanical view of proteins. The positions of the backbone and sidechain atoms are specified, together with the positions of some observed water molecules and other atoms. This basic information allows the derivation of extensive additional information, as we will explain subsequently. But for the moment, we simply recall some information on the static description of proteins.

### 4.4.1 Conformational geometry of proteins

We recall the basic ingredients of the peptide group from Figure 4.1. If $x$ is a given residue, then $N(x), H(x), C(x)$ and $O(x)$ denote the position vectors of the corresponding atoms in the peptide group. For the remaining atoms, the standard notation from the PDB is as follows:

$$
C_{\alpha}(x), C_{\beta}(x), C_{\gamma}(x), C_{\delta}(x), C_{\epsilon}(x), C_{\eta}(x)
$$

are the $\alpha, \beta, \gamma, \delta, \epsilon, \eta$ carbons (denoted in plain text in the PDB by $\mathrm{CA}, \mathrm{CB}, \mathrm{CG}, \mathrm{CD}, \mathrm{CE}, \mathrm{CH}$ ) in the sidechain structure of residue $x$. Most of these can also appear with subscripts, e.g., $C_{\gamma^{i}}$ for $i=1,2$ in Ile and Val. Correspondingly, $N_{\delta^{i}}(x), N_{\epsilon^{i}}(x), N_{\eta^{i}}(x)$ are the $i$-th $\delta, \epsilon, \eta$ nitrogens, denoted in plain text in the PDB by NDi, NE $i$, NH $i$ for $i=1,2$. Notation for oxygens is similar. Unfortunately, the plain text descriptor OH for $O_{\eta}$ in Tyr is a bit confusing, since this oxygen has an attached hydrogen.

We can view $C_{\alpha}(x), N_{\delta^{i}}(x)$, etc., as three-dimensional vectors, using the corresponding coordinates from the PDB. For amino acids $x_{i}, x_{i+1}$ which are adjacent in the protein sequence, the backbone vector is defined as

$$
\begin{equation*}
\mathcal{B}=C_{\alpha}\left(x_{i+1}\right)-C_{\alpha}\left(x_{i}\right) \tag{4.1}
\end{equation*}
$$

The sidechain vector $\mathcal{S}(x)$ for a given amino acid $x$, defined by

$$
\begin{equation*}
\mathcal{S}(x)=C_{\beta}(x)-C_{\alpha}(x) \tag{4.2}
\end{equation*}
$$

will be used to measure sidechain orientation. $\mathcal{S}$ involves the direction of only the initial segment in the sidechain, but we will see that it is a significant indicator of sidechain conformation. For $x=G l y$, we can substitute the location of the sole hydrogen atom in the residue in place of $C_{\beta}$. For each neighboring residue pair $x_{i}, x_{i+1}$, the sidechain angle $\theta\left(x_{i}, x_{i+1}\right)$ is defined by

$$
\begin{equation*}
\cos \theta\left(x_{i}, x_{i+1}\right)=\frac{\mathcal{S}\left(x_{i}\right) \cdot \mathcal{S}\left(x_{i+1}\right)}{\left|\mathcal{S}\left(x_{i}\right)\right|\left|\mathcal{S}\left(x_{i+1}\right)\right|} \tag{4.3}
\end{equation*}
$$

where $\mathcal{B}$ is defined in (4.1), and $A \cdot B$ denotes the vector dot-product.
Draft: February 28, 2008, do not distribute

It is not common to characterize the secondary structures (helix and sheet) by $\theta$, but $\theta$ is strongly correlated with secondary structure [151], and it gives a simple interpretation. Values $70 \leq \theta \leq 120$ are typical of $\alpha$-helices, since each subsequent residue turns about 90 degrees in order to achieve a complete ( 360 degree) turn in four steps (or 72 degrees for five steps, or 120 degrees for three steps). Similarly $140 \leq \theta \leq 180$ is typical of $\beta$-sheets, so that the sidechains are parallel but alternate in direction, with one exception. Some $\beta$-sheets have occasional 'spacers' in which $\theta$ is small [151], in keeping with the planar nature of sheets.

The distribution of the $\theta$ angle peaks roughly at 44, 82 and 167 degrees [151. The peptide bond makes it difficult for $\theta$ to be much less than 50 degrees, thus the smaller peak corresponds to a motif where the side chains align as closely as possible. A small number of these occur in beta sheets, but the majority of them constitute an independent motif whose properties deserve further study.

The different structural motifs have characteristic sidechain compositions 8, 151. For the larger values of $\theta$, hydrophobic residues are found in most pairs; $\beta$-sheets have alternating hydrophobic and hydrophilic pairs [151]. By contrast [151], the most common pairs involve predominantly polar or charged residues for $\theta \leq 50$. The ends (or caps) of $\alpha$-helices necessarily must be different from the middle to terminate the structure [8].

We also recall the standard main-chain dihedral angles. Given a sequence of four main chain atoms $a_{i}$, let $\left[a_{1}, a_{2}, a_{3}, a_{4}\right]$ denote the dihedral (or torsion) angle between the planes defined by the points $a_{1}, a_{2}, a_{3}$ and $a_{2}, a_{3}, a_{4}$. Then the $\psi, \omega$ and $\phi$ angles are defined by

$$
\begin{align*}
\psi\left(x_{i}\right) & =\left[N\left(x_{i}\right), C_{\alpha}\left(x_{i}\right), C\left(x_{i}\right), N\left(x_{i+1}\right)\right] \\
\omega\left(x_{i+1}\right) & =\left[C_{\alpha}\left(x_{i}\right), C\left(x_{i}\right), N\left(x_{i+1}\right), C_{\alpha}\left(x_{i+1}\right]\right.  \tag{4.4}\\
\phi\left(x_{i+1}\right) & =\left[C\left(x_{i}\right), N\left(x_{i+1}\right), C_{\alpha}\left(x_{i+1}\right), C\left(x_{i+1}\right)\right] .
\end{align*}
$$

In Chapter 14 we study the effect of a polar environment on the flexibility of $\omega$.

### 4.4.2 $\phi, \psi$ versus $\psi, \phi$ : the role of $\theta$

The pair of angles $\phi_{i}, \psi_{i}$ captures the rotation of the peptide chain around the $i$-th $C_{\alpha}$ carbon atom. The $\theta$ angle measures the rotation that corresponds with comparing angles $\psi_{i}, \phi_{i+1}$ in successive peptides (cf. Exercise 4.3). This correlation has recently been observed to have significant predictive power 87 .

The conformations of $\phi_{i}, \psi_{i}$ are typical of different secondary structures, such as $\alpha$-helix or $\beta$-sheet. The Ramachandran plot depicts the distributions of angles that are commonly adopted (cf. Exercise 4.7).

### 4.4.3 Sidechain rotamers

The sidechains are not rigid, so the geometric description of a sidechain requires more information than $\phi, \psi$ and so forth. Libraries of angular orientations of the different segments have been developed [156]. The possible orientations are not uniformly distributed in many cases, but rather show a strong bias for a few discrete orientations. For example, carbon chains typically orient so that the hydrogen atoms are in complementary positions. In Figure 4.8, the three primary

Draft: February 28, 2008, do not distribute


Figure 4.8: The primary sidechain rotamer conformations (a) gauche+, (b) gauche-, and (c) trans, corresponding to $\chi_{1}$ values of (a) -60 degrees, (b) +60 degrees and (c) 180 degrees. The view is oriented so that the $C_{\alpha}$ and $C_{\beta}$ atoms are aligned perpendicular to the plane of the page. The closer (and therefore larger) atoms are indicated with dashed lines and bold letters. The hydrogens for $C_{\beta}$ are indicated. The atom marked ' XG ' corresponds to either a $C_{\gamma}$ or an $O_{\gamma}$ atom.
conformations are shown for side chains with $C_{\beta}$ and $C_{\gamma}$ constituents. These conformations are known as gauche + , gauche-, and trans, corresponding to mean $\chi_{1}$ values of -60 degrees, +60 degrees and 180 degrees, respectively.

However, the distributions can change depending on local neighbor context [152].

### 4.4.4 Volume of side chains

The sizes of amino acid side chains varies significantly. But there is also a more subtle size issue that is solvent dependent. In 1975, Chothia initiated a study of the size of sidechains and the change in size in the core of proteins. This was an early use of datamining in the PDB. That study was later revisited [108, and subsequent studies have further refined estimates of sidechain volume, including sizes of individual atom groups [237].

One of the significant conclusions [108] is that hydrophobic residues occupy less volume inside the core of a protein than they do in bulk water. Similarly, hydrophilic residues occupy more volume inside the core of a protein than they do in bulk water. Given our general understanding of the hydrophobic effect, this is not surprising. However, it gives a clear understanding of an important packing effect.

In typical proteins, the increase in volume due to burying hydrophilic residues is compensated by the decrease in volume due to burying hydrophobic residues. That is, the net volume change upon folding is typically quite small. However, for other systems, such a balance does not seem to be so close. For example, cell membranes are made of lipid layers which are composed substantially of hydrophobic chains. Thus simple pressure tends to keep such cell membranes intact. To break apart, the cell membranes constituents would have to undergo a substantial increase in volume and thus induce a significant increase in pressure.

The volumetric cost of burying hydrophilic residues makes one wonder why they appear in proteins at all. It may be that their role in the electrostatic balance of the protein is their reason for existence. On the other hand, if proteins had only hydrophobic cores, they would be harder to
unfold. Both of these effects may contribute to the reason for having charged and polar residues in protein cores.

### 4.5 Special side chains

There are many ways that sidechains can be classified, according to polarity, hydrophobicity and so on. When all such designations are taken into account, each sidechain becomes essentially unique. Indeed, it is advisable to study more complete descriptions of the unique properties of individual sidechains [47. But there are some special properties of sidechains that deserve special mention here for emphasis.

### 4.5.1 Glycine (and Alanine)

Glycine is special because it has essentially no sidechain. More precisely, it is the only aminoacid without a $C_{\beta}$ carbon. As a result, it is appropriate to think of Gly as polar, since the polarity of the backbone itself has a significant impact on the environment near the sidechain. In this regard, alanine can also be viewed to be somewhat polar. Alanine has a $C_{\beta}$ carbon, but no other heavy atoms in its sidechain, a feature unique to Ala.

### 4.5.2 Proline

Proline is unique because it connects twice to the backbone. This causes a special rigidity not found with other residues. There is a special conformation of protein structures called PP2 (a.k.a. PPII or PII) which refers to the type of structure that a polyproline strand adopts 855, 221].

### 4.5.3 Arginine

The uniqueness of arginine is highlighted by the fact that its residue is the guanidinium ion. Guanidinium, like urea (a.k.a. carbamide), has the property that it can denature proteins, that is, cause them to unfold. How this is achieved, for either denaturant, is not fully understood. One feature of the arginine residue is that the positive charge at the end of the residue is distributed quite broadly among the atoms at the end of the residue (see Table [7.5). How or why this might have a special effect is not clear.

It is very difficult to form natural water structures around an arginine [163]. There are two NH groups (in the two NH2 groups at the end) in which the N-H vectors are nearly parallel. Model building shows that it is very hard for waters attached to these hydrogens to cohabitate. There is a similar difficulty with the NH and NH2 groups, where there are N-H vectors nearly parallel. One can think of the planar structure of the terminal CN3H5 group as like a knife blade that cuts through water structures.

One property of arginine is that polyarginine is the polypeptide most able to cross a cell membrane without the help of a transporter molecule [169], and compounds rich in Arg have similar behavior [266].

Draft: February 28, 2008, do not distribute

### 4.5.4 Cysteine

What makes cysteine special is the ability of the sidechain to bond with another cysteine sidechain, making a disulfide bridge (Section 4.2.2). This is the only sidechain that forms a covalent bond with another sidechain.

### 4.5.5 Aromatic sidechains

Three sidechains (Tyr, Trp, Phe) have benzene rings as a significant part of their structure. At first, these appear simply hydrophobic, but the electron structure of aromatic rings is complex [55]. There is a doughnut of positive charge centered in the plane of the carbon and hydrogen atoms, and the hole of the doughnut contains disks of negative charge on either side of the main positive ring (see Figure 2A of [55]). This makes these side chains polar. Tyrosine is also polar in a more conventional way at the end of the sidechain due to the OH group there.

Tryptophan deserves special mention for various reasons, not just because of its pop-culture notoriety for sleep induction [178] and other behavioral impact 198. It is the largest and most complex sidechain, involving two types of rings, the indole ring in addition to the benzene ring ${ }^{2}$ Tryptophan is also the least common and most conserved (least likely to be mutated in homologous proteins) sidechain.

### 4.5.6 Remembering code names

Many of the single letter codes for sidechains are obvious (Alanine, Glycine, Histidine, Isoleucine, Leucine, Methionine, Proline, Serine, Threonine, Valine), but others require some method to remember. We propose here some non-serious mnemonic devices that may aid in retaining their assignments.

Asp and Glu are the negatively charged residues, and the alphabetic order also corresponds with the size order (Asp is smaller than Glu). The code names are also alphabetical (D and E); the choice of E corresponds to the charge $e$ of the extra electron.

Two of the positive sidechains also have special codes. To remember the R for arginine is to think of it as the pirate's favorite sidechain. To "lyse" means to destroy or disorganize, so we can think of lysine as the Killer sidechain.

The biggest sidechains (the aromatic ones) also have letter codes which need special treatment. A way to remember the single letter code for Phe is to misspell it with the Ph changed to F . A way to remember the single letter code for $\operatorname{Trp}$ is that it is the Widest sidechain. A way to remember the single letter code for Tyr is to focus on the second letter Y in the name.

The two remaining proteins are comparable to Asp and Glu, but with nitrogen groups replacing one of the oxygens: asparagiNe and Qlutamine. The emphasis on Nitrogen is clear in Asn, since it is the third letter of the code. The letter $G$ is one the most overloaded among first letters in the sidechain names, but Q is a close match for a poorly written G .

[^1]Draft: February 28, 2008, do not distribute

### 4.6 Sidechain ionization

We will not consider extensively pH effects, although these clearly involve a type of modulation of electrical forces. There is significant pH variation in different parts of cells, and thus it has a potential role in affecting protein-ligand interactions.

The effects of pH are both localized and dynamic in nature, since the number of ions that can be involved in protein-ligand interactions is not large. For example, a well solvated large biomolecule [251] can be modeled dynamically with just over $10^{5}$ atoms, and significantly less than $10^{5}$ water molecules. But at pH 7 , there is just one hydronium molecule per $5.5508 \times 10^{8}$ water molecules (cf. Section 10.7.1). The number of water molecules in the simulation in 251] used fewer than 55,508 water molecules, and thus would not have included a hydronium ion until the pH was less than three. On the other hand, ions cluster around proteins since they have charged and polar residues, so a more complex model is required to account for their effects.

The ends of some sidechains can vary depending on the ionic composition of the solvent 47. The pH value (Section 10.7.1) relevant for ionization is out of the range of biological interest in most cases, with the exception of His. We list the intrinsic $p K_{a}$ values [47] in Table 4.2 for reference. This value is the pH at which half of the residues would be in each of the two protonation states. For example, below pH four, Asp would be more likely to be protonated, so that one of the terminal oxygens would instead be an OH group. In this case, it would be appropriate to refer to the residue as aspartic acid. Similarly, for pH below 4.4, Glu would more likely have an OH terminal group, and be called glutamic acid. For simplicity, we refer to the residues in their form that is most common at physiological values of pH .

### 4.7 Exercises

Exercise 4.1 Draw all the atoms in the tri-peptide GAG, including the $C$-terminal and $N$-terminal ends.

Exercise 4.2 Determine how long each sidechain is by scanning the PDB. That is, determine the distribution of distances from the $C_{\alpha}$ carbon to the terminal (heavy) atoms for each residue (amino acid) type.

Exercise 4.3 In typical peptide bonds, the $\omega$ angle is constrained to so that the peptide bond is planar (cf. Figure 14.1). In this case, there is a relationship imposed between $\theta, \phi$ and $\psi$. Determine what this relationship is.

Exercise 4.4 Proteins are oriented: there is a C-terminal end and an $N$-terminal end. Determine whether there is a bias in $\alpha$-helices in proteins with regard to their macrodipole $\mu$ which is defined as follows. Suppose that a helix consists of the sequence $p_{i}, p_{i+1}, \ldots, p_{i+\ell}$ where each $p_{j}$ denotes an amino-acid sidechain. Let $\mathcal{C}(p)$ denote the charge of the sidechain $p$, that is, $\mathcal{C}(D)=\mathcal{C}(E)=-1$ and $\mathcal{C}(K)=\mathcal{C}(R)=\mathcal{C}(H)=+1$, with $\mathcal{C}(p)=0$ for all other $p$. Define

$$
\begin{equation*}
\mu\left(p_{i}, p_{i+1}, \ldots, p_{i_{\ell}}\right)=\sum_{j=0}^{\ell} \mathcal{C}\left(p_{i+j}\right)\left(j-\frac{1}{2} \ell\right) \tag{4.5}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

Plot the distribution of $\mu$ over a set of proteins. Compare with the peptide dipole, which can be modeled as a charge of +0.5 at the $N$-terminus of the helix and a charge of -0.5 at the $C$-terminus of the helix. How does this differ for left-handed helices versus right-handed helices? (Hint: the $P D B$ identifies helical regions of protein sequences. The peptide dipole in our simplification is just $\ell$, so $\mu / \ell$ provides a direct comparison.)

Exercise 4.5 Consider the definition of macrodipole introduced in Exercise 4.4. Explain why the $\alpha$-helical polypeptide Glu $_{20}$ Ala $a_{20}$ would be more stable than Ala $_{20}$ Glu $_{20}$.

Exercise 4.6 Determine the Ramachandran plot for a set of proteins. That is, plot the $\phi_{i}$ and $\psi_{i}$ angles for all peptides in the set. Use a different symbol or color for the cases where the $i$-th peptide is said to be a helix, sheet or turn in the PDB file.

Exercise 4.7 Determine the Ramachandran plot for a set of proteins. That is, plot the $\phi_{i}$ and $\psi_{i}$ angles for all peptides in the set. Instead of using the designation in the PDB file (as helix, sheet or turn), use the software DSSPcont [4] and use a different symbol or color for the different classes.

## Chapter 5

## Hydrogen bonds

Hydrogen bonds are the most important bond in biochemistry, so we need to understand them in some depth. Unfortunately, there are several challenges. First of all, although hydrogen bonds in proteins have been studied extensively [17, 142], they are not yet fully understood and are still actively studied [122, 123. Secondly, in most PDB files, hydrogens are not listed at all, due to the difficulty of locating them by typical imaging techniques. We describe how their locations can be inferred starting in Section 5.3. We begin by reviewing some of the main results.

The general hydrogen bond is of the form XH - - Y where X and Y are 'heavy atoms' such as F, N, O, S or even C in some cases. The X atom is called the donor of the bond, and the Y atom is called the acceptor of the bond.

### 5.1 Hydrogen bond theory

Hydrogen bonds differ based on the heavy atoms that are involved. The variation in bond distance and strength is illustrated in Table 5.1 which has been extracted from [134. What is clear from this data is that the donor type (the side of the bond that includes the hydrogen) is the primary determinant of the hydrogen bond strength (and length) in these cases. This is interpreted to mean that the charge dipole of the donor is the determining factor [134]. In some sources (including Wikipedia), the electronegativity of the constituents is given as the key factor. But according to [134], "the ability of proton donors and acceptors to form hydrogen bonds (X-H. . . Y) is more closely related to their respective acidity or basicity than to the electronegativities of X and Y ."

There is a strong angular dependence for the energy of the hydrogen bond [179]. One might hope that modeling the hydrogen bond as a simple dipole-dipole interaction (Section 9.2.1) would be sufficient to capture the angular dependence. But a purely partial-charge (i.e., dipole-dipole) model of hydrogen bonds is not sufficient to capture the angular dependence of the energy: "At the distances where H bonding occurs, the dipole moment approximation is a poor one and higher multipoles must be considered" [134], as we confirm in Section 9.2.1.

A model based only on atom distances has been proposed [179], in which the dominant term appears to be a strong repulsion term between the like-charged atoms. Such a model is simple to implement because it uses exactly the same data as a dipole model, but with a more complex form

| Donor | Acceptor | System | $R(\AA)$ | $\Delta E($ kcal $)$ |
| :---: | :---: | :---: | :---: | :---: |
| $\mathrm{NH}_{3}$ | HF | $\mathrm{HF}-\mathrm{HNH}_{2}$ | 3.45 | 1.3 |
| $\mathrm{NH}_{3}$ | $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{H}_{2} \mathrm{O}-\mathrm{HNH}_{2}$ | 3.41 | 2.3 |
| $\mathrm{NH}_{3}$ | $\mathrm{H}_{3} \mathrm{~N}$ | $\mathrm{H}_{3} \mathrm{~N}-\mathrm{HNH}_{2}$ | 3.49 | 2.7 |
| $\mathrm{H}_{2} \mathrm{O}$ | HF | $\mathrm{HF}-\mathrm{HOH}$ | 3.08 | 3.0 |
| $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{H}_{2} \mathrm{O}-\mathrm{HOH}$ | 3.00 | 5.3 |
| $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{H}_{3} \mathrm{~N}$ | $\mathrm{H}_{3} \mathrm{~N}-\mathrm{HOH}$ | 3.12 | 5.8 |
| HF | HF | $\mathrm{HF}-\mathrm{HF}$ | 2.72 | 9.4 |
| HF | $\mathrm{H}_{2} \mathrm{O}$ | $\mathrm{H}_{2} \mathrm{O}-\mathrm{HF}$ | 2.75 | 11.7 |
| HF | $\mathrm{H}_{3} \mathrm{~N}$ | $\mathrm{H}_{3} \mathrm{~N}-\mathrm{HF}$ | 2.88 | 4.6 |

Table 5.1: $R$ is the distance (in Ångstroms) between the donor and acceptor (heavy) atoms. The energy $\Delta E$ of the hydrogen bond is given in kcal/mole. Note that $R$ "is primarily a function of the degree of positive charge on the hydrogen in the H bond" [134].
and with additional data derived from ab initio quantum chemistry calculations.
The accurate computation of the most basic hydrogen bond, the water dimer, has been of recent interest [133], even though this computation has been carried out for several decades 134]. The fact that this simple interaction is still studied is an indicator of the difficulty of determining information about general hydrogen bonds.

One question to ask about hydrogen bonds is whether the hydrogens take on a symmetric position between the donor and acceptor, or whether it favors one side (donor) over the other. The answer is: yes and no [199. Both situations arise in nature, and there is an intriguing bifurcation between the two states, as depicted by the caricature in Figure 5.1. Depicted is a curve that was fit [199] to extensive data on bond lengths of OH - - O hydrogen bonds. The horizontal axis is the distance between the oxygen centers, and the vertical coordinate is the (larger) distance between oxygen and hydrogen. The upper-left segment, where the O-H distance is exactly half of the O-O distance, is the symmetric arrangement. The dashed parts of the curves indicate where data has been found in both states. But what is striking is the void in the O-H distance region between $1.1 \AA$ and nearly $1.2 \AA$. Thinking in bifurcation terms, one can stretch the O-O distance in the symmetric configuration, but at a certain point it loses stability and has to jump to the asymmetric one in which the hydrogen has a preferred partner. Moreover, as the O-O distance continues to increase, the (smaller) O-H distance decreases, as the influence of the other oxygen decreases with increasing distance. Note that the O-O distance (for waters) reported in Table 5.1 is $3.0 \AA$, thus clearly in the asymmetric regime (actually off the chart in Figure 5.1]).

### 5.2 Types of hydrogen bonds

As indicated in Table 5.1, hydrogen bonds vary in character depending on the donor and acceptor. In proteins, there are two classes of donors and acceptors, mainchain (or backbone) and sidechain.

Draft: February 28, 2008, do not distribute


Figure 5.1: Cartoon of the bifurcation of O-H..O hydrogen bonds from a symmetric arrangement to an asymmetric arrangement, based on Figure 4 of 199. The horizontal axis is the O-O distance and the vertical coordinate is the O-H distance (both in Angstroms). The upper-left segment is the symmetric arrangement.

All backbone nitrogens (with the exception of proline, unless it is N-terminal) can act as donors of hydrogen bonds, and all backbone oxygens can be acceptors of hydrogen bonds. In addition, many of the standard sidechains can act as donors or acceptors, as listed in Table 5.2. Note that certain atoms can be both donors and acceptors.

Given two classes of contributors, mainchain (M) and sidechain (S), there are four classes of bond pairs: M-M, M-S, S-M, and S-S. We have differentiated between S-M and M-S depending on whether the donor or acceptor is M or S , but in some cases these two classes are lumped into one class.

Given the rigidity of the backbone and the flexibility of the sidechains, it would be reasonable to assume that S-S bonds were the most common and M-M the least. Curiously, it is just the opposite. In Chapter 12, we will see that mainchain-mainchain are much more common. By simple counts in a database of 1547 nonredundant structures, the number of M-M bonds is nearly four times the number of mainchain-sidechain (M-S and S-M) bonds combined, and it is seven times the number of sidechain-sidechain bonds. On the other hand, one finds a significant number of potential sidechainwater hydrogen bonds in many PDB files. These include apparent water bridges [197, [257]. It is not clear how fully waters in PDB files are reported, but their importance to protein structure is significant.

Typical hydrogen donors would make only one hydrogen bond, whereas typical oxygen acceptors can make two hydrogen bonds. However, more complex patterns are possible; see the figures on page 139 of [123].

| Full name of <br> amino acid | three <br> letter | single <br> letter | Donors <br> (PDB name) | Acceptors <br> (PDB name) |
| :---: | :---: | :---: | :---: | :---: |
| Arginine | Arg | R | NE, NH1, NH2 | - |
| Asparagine | Asn | N | ND2 | OD1 |
| Aspartate | Asp | D | - | OD1, OD2 |
| Cysteine | Cys | C | SG* | SG |
| Glutamine | Gln | Q | NE2 | OE1 |
| Glutamate | Glu | E | - | OE1, OE2 |
| Histidine | His | H | ND1, NE2 | ND1, NE2 |
| Lysine | Lys | K | NZ | - |
| Methionine | Met | M | - | SD |
| Serine | Ser | S | OG | OG |
| Threonine | Thr | T | OG1 | OG1 |
| Tryptophan | Trp | W | NE1 | - |
| Tyrosine | Tyr | Y | OH | OH |

Table 5.2: Donors and acceptors for sidechain hydrogen bonds. ${ }^{*}$ If a Cys is involved in a disulfide bridge, it cannot be a hydrogen bond donor.

### 5.3 Identification of hydrogen positions

Most PDB files do not include locations of hydrogens. Only the heavier atoms are seen accurately in the typical imaging technologies. However, in many cases, the positions of the missing hydrogens can be inferred according to simple rules. For example, the position of the hydrogen that is attached to the mainchain nitrogen (see Figure 4.1) can be estimated by a simple formula. The C-O vector and the $\mathrm{N}-\mathrm{H}$ vector are very nearly parallel, so one can simply take

$$
\begin{equation*}
H=N+|C-O|^{-1}(C-O) \tag{5.1}
\end{equation*}
$$

since the N-H distance is approximately one Angstrom. We leave as an exercise (Exercise 5.1) to make the small correction suggested by the figure on page 282 in [191].

As another simple example, the position of the hydrogens that are attached to the terminal nitrogen in Asn and Gln can also be estimated by a simple formula. The terminal O-C-NH2 group of atoms are all coplanar, and the angles formed by the hydrogens around the nitrogen are all 120 degrees, as depicted in Figure 5.2. The angle between the C-N and the C-O vectors is very close to 120 degrees [171, so the C-O vector and one of the N-H vectors are very nearly parallel. So one can again take

$$
\begin{equation*}
H^{1}=N+|C-O|^{-1}(C-O) \tag{5.2}
\end{equation*}
$$

as the location for one of the hydrogens attached to N , since again the $\mathrm{N}-\mathrm{H}$ distance is approximately one $\AA$ Angstrom. For the other hydrogen bond, the direction we want is the bisector of the C-O and C-N directions. Thus the second hydrogen position can be defined as

$$
\begin{equation*}
H^{2}=N+\frac{1}{2}\left(|O-C|^{-1}(O-C)+|N-C|^{-1}(N-C)\right) \tag{5.3}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

| Full name of non-standard residue or molecule | PDB three letters | Donors | Acceptors |
| :---: | :---: | :---: | :---: |
| Acetyl group | ACE |  | O |
| Glycerol | GOL | O1, O2, O3 | O1, O2, O3 |
| Nitrate Ion | NO3 |  | O1, O2, O3 |
| Phosphotyrosine | PTR | $\mathrm{N}, ~ \mathrm{O} 2 \mathrm{P}^{\ddagger}, ~$ O3 $\mathrm{P}^{\ddagger}$ | O, OH, O1P, O2P, O3P |
| Pyroglutamic acid | PCA | $\mathrm{N}^{\dagger}$ | O, OE |
| Phosphono group | PHS |  | O1P, O2P, O3P |
| Phosphate Ion | PO4 |  | O1, O2, O3, O4 |
| Sulphate Ion | SO4 |  | O1, O2, O3, O4 |

Table 5.3: PDB codes for donor and acceptor atoms in some nonstandard residues and molecules. Key: $\dagger$ Only N-terminus. $\ddagger$ In case that the hydrogens PHO2, PHO3 exist in the PDB files.


Figure 5.2: Hydrogen placement for Asn and Gln. Shown is the terminal group of atoms for the sidechains. The atom marked C' denotes the preceding carbon in the sidechain, viz., CB for Asn and CG for Gln.

We leave as an exercise (Exercise 5.2) to make the small corrections suggested by the Figure 13 in [171].

The position of hydrogens can be modeled by the bond lengths and angles given in [171]. A program called HBPLUS [165] was developed based on this information to provide hydrogen positions in a PDB format.

Most hydrogens can be located uniquely. In particular, the Appendix in 171 depicts the locations of such hydrogens, as well as providing precise numerical coordinates for their locations. However, other hydrogens are not uniquely determined. For example, the hydrogen attached to the terminal oxygen in the tyrosine sidechain has two potential positions. The hydrogen must be in the plane of the aromatic ring, but there are two positions that it can take. This is depicted in Figure 5.3. The one which makes the stronger H-bond with an acceptor is presumably the one that is adopted.

Draft: February 28, 2008, do not distribute

### 5.4. GEOMETRIC CRITERIA FOR HYDROGEN BONDEHAPTER 5. HYDROGEN BONDS



Figure 5.3: Hydrogen placement for Tyr. Both positions are possible for the terminal hydrogen.


Figure 5.4: Hydrogen placement for Ser and Thr: anywhere on the dotted circle. A Cys sidechain not in a disulfide bond would be similar, with O replaced by S .

The terminal OH groups in serine and threonine are even less determined, in that the hydrogen can be in any position in a circle indicated in Figure 5.4. A Cys sidechain that is not engaged in a disulfide bond would be similar, with the oxygen in Figure 5.4 replaced by a sulfur.

An interesting example of the ambiguity of the assignment of the hydrogen location for serines and threonines occurs in the PDB file 1C08. In chain B, Thr30 and Ser28 form a sidechain-sidechain hydrogen bond involving the terminal OH groups. But which is the donor and which is the acceptor cannot be differentiated by the data in the PDB file in a simple way. Model building shows that both are possible, and indeed there could be a resonance (Section 14.1) between the two states. One state may be forced by the local environment, but without further determining factors both states are possible. It is possible to critique the detailed geometry by considering the quality of the corresponding dipole-dipole interaction (see Section 9.2.1). According to this metric, Thr30 is the preferred donor.

### 5.4 Geometric criteria for hydrogen bonds

One approach to approximating the angular dependence of the hydrogen bond is to use angular limits, as well as distance limits, in the definition. Each hydrogen bond can be defined by the geometric criteria (Figure 5.6) based on those used in [165], as we now enumerate:


Figure 5.5: Ambiguous hydrogen placement for serine-28 (lower right) - threonine-30 (upper left) sidechain-sidechain hydrogen bond involving the terminal O-H groups; from the B chain in the PDB file 1C08. The sidechain of isoleucine- 29 has been omitted but the backbone atoms are shown connecting the two residues. Only the oxygen atoms in the terminal O-H groups are shown.


Figure 5.6: Geometric model for hydrogen bonds: D is the donor atom, H the hydrogen, A the acceptor, B acceptor antecedent (i.e. an atom one covalent bond away from the acceptor).

1. Distance between donor and acceptor $|D-A|<3.5 \AA$
2. Distance between hydrogen and acceptor $|H-A|<2.5 \AA$
3. Angle of donor-hydrogen-acceptor $\angle D H A>90^{\circ}$
4. Angle of donor-acceptor-acceptor antecedent $\angle D A B>90^{\circ}$
5. Angle of hydrogen-acceptor-acceptor antecedent $\angle H A B>90^{\circ}$

### 5.5 Carboxyl-carboxylate hydrogen bonds

Under suitable conditions, the terminal groups of Asp and Glu can be come protonated. The resulting OH group can then form hydrogen bonds with oxygens, including the ones in the terminal
groups of other Asp and Glu residues [253]. These are referred to as carboxyl-carboxylate hydrogen bonds. Although these bonds would be expected in low pH environments [213], they have been found to be critical elements of ion channels [174. In typical PDB structures, the hydrogen in a carboxyl-carboxylate hydrogen bond would not be visible. Thus it could be associated with either oxygen unless further information is available to reveal the association.

### 5.6 Exercises

Exercise 5.1 Refine the formula (5.1) to give a more precise location for the hydrogen attached to the nitrogen in the peptide bond, e.g., following the figure on page 282 in [191].

Exercise 5.2 Refine the formulas (5.2) and (5.3) to give a more precise location for the hydrogens attached to the terminal nitrogen in the residues Asn and Gln, using the data in Figure 13 in [171.

Exercise 5.3 Use the improved model for the energy of a hydrogen bond in [179] to estimate the strength of hydrogen bonds. Apply this to antibody-antigen interfaces to investigate the evolution of the intermolecular hydrogen bonds at the interfaces.

Exercise 5.4 Hydrogen positions can be inferred using neutron diffraction data, because hydrogen is a strong neutron scatterer. There are over a hundred PDB files including neutron diffraction data. Use this data to critique the models for hydrogen locations presented in this chapter.

Exercise 5.5 Helical secondary structure is formed by amide-carbonyl hydrogen bonding between peptides $i$ and $j$ where $3 \leq|i-j| \leq 5$. Determine how frequent it is to have $i-j=k$ for the different possible values of $k=-5,-4,-3,3,4,5$. Are there instances where amide-carbonyl hydrogen bonding between peptides $i$ and $j$ where $|i-j|=2$ or $|i-j|=6$ ?

Exercise 5.6 The $C$ - $O$ (carbonyl) groups in the peptide backbone can make two hydrogen bonds (typically), whereas the $N$-H (amide) group usually forms only one hydrogen bond. How common is it for carbonyl groups to make two bonds in helical secondary structures? In $\beta$-sheet structures? How often are the bonds mainchain-mainchain bonds, versus sidechain-mainchain bonds? As a first step, you can define helical carbonyls to be ones where there is bonding between peptides $i$ and $j$ where $3 \leq|i-j| \leq 5$, but determine how many double bonds there are for each value of $k=i-j$.

Exercise 5.7 Determine the angular dependence of the mainchain-mainchain hydrogen bond. What is the distribution of $O-H$ distances and $C-O, N-H$ angles? Consider the different classes of bonds separately: those in (1) parallel and (2) anti-parallel sheets, and those in helices of separation $k= \pm 3, \pm 4, \pm 5$ (cases 3-8). How does the bond distance and angle correlate? What is the mean distance and angle in each case?

## Chapter 6

## Determinants of protein-protein interfaces

We now turn to a key question: what factors are most influential in protein-ligand binding? We review attempts to answer this question both to give a sense for the historical development and also to emphasize key aspects of the datamining techniques used. Later in the book we will clarify the role of dehydrons in this process, but for now we proceed navely to get a sense of how the ideas developed.

Protein associations are at the core of biological processes, but their physical basis, often attributed to favorable pairwise interactions, remains an active topic of research [205, 43, 25, 125, 155, 245, 92, 170. Hydrophobic-polar mismatches at protein-protein interfaces are all too common and difficult to properly account for. The prediction and rationalization of binding sites for soluble proteins require that we quantify pairwise energy contributions, and concurrently, the extent to which surrounding water is immobilized or excluded from the interactive residue pairs. as proteins associate, their local solvent environments become modified in ways that can dramatically affect the intramolecular energy [245, 84, 177, 73, 12, 70, 64].

It is well known that water removal from hydrophobic patches on the protein surface results in a high thermodynamic benefit [205, 43, 25, 125, 155, 245, 92, 170], due to an entropic gain by the solvent. Thus, hydrophobic patches might become suitable binding regions provided a geometric match on the binding partner is obtained. However, such patches are rare: most protein surfaces have the expected high ratios (typically $7: 1$ to 10:1) of hydrophilic to hydrophobic residues [205, 43, 25, 125, 155, 245, 92, 170]. Furthermore, even if overexposed hydrophobic patches become involved in associations, the resulting interface often presents hydrophobic-polar mismatches [229].

At the simplest level, one would expect the sort of bonds that help proteins form their basic structure would also be involved in joining two different proteins together. Both hydrogen bonds and salt bridges play a significant role at protein interfaces [257]. The density of hydrogen bonds between two different proteins at an interface is about one per two square nanometers. If you think of a checkerboard with nanometer sized squares, then it is like having one hydrogen bond on each of the red squares. The average number of hydrogen bonds per interface is about ten. On the other hand, the average number of salt bridges per interface is only two. Disulfide bonds play a more

### 6.1. AMINO ACIDSCAIA PRERIGINIDHRCEIFENIXNANTERREFEROTEIN-PROTEIN INTERFACES

limited and specialized role.
It might be that the story of protein-protein interactions ends here, with the intermolecular hydrogen bonds and salt bridges being the whole story. However, three of the 54 high-resolution structures studied in [257] have no hydrogen bonds or salt bridges, and another dozen have no salt bridges and five or fewer hydrogen bonds. Not surprisingly, we will begin to see indications of the role of intramolecular hydrogen bonds that become enhanced upon binding, as we depicted in Figure 3.11

One factor that complicates the picture of protein-protein interactions is the appearance of water molecules which appear to play a structural role, as opposed to simply mediating interactions via dielectric effects. In the protein interfaces studied in [257], polar atom pairs bridged by water across the interface with hydrogen bonds were more numerous than direct hydrogen bond pairs, with each water molecule connecting 3.8 cross-chain atom pairs on average.

### 6.1 Amino acids at protein-protein interfaces

We begin with a simple use of datamining applied to the understanding of amino acid tendencies at interfaces. There are different questions that one can ask, and of course it is natural that amino acids get ranked in different orders accordingly. For simplicity, we contrast just two, but we also review others in Section 6.5. The data here is drawn primarily from [30, 83, (97].

The site specificity of protein-protein interactions has been widely studied due to its central biological significance [51, 97, 116, 125, 126, 127. Hydrophobic residues such as Leu and Val are more abundant at protein-ligand interfaces. As a result, the removal of water surrounding hydrophobic residues on the protein surface has been assumed to be a driving force for association [77, 227]. But it is also true that such residues are more abundant over-all (see Table 6.2).

The first question [83] we consider is about the amino acid composition of protein-protein interfaces. This can be done by simply counting, once an identification has been made regarding which amino acids are at an interface. However, simple frequencies are misleading: Leu is the most common residue at interfaces, but it is also overwhelmingly the most common residue in most proteins. Thus one has to normalize by the natural frequencies of amino acids in proteins [30].

The second question [97] is about the amino acid composition for pairs of amino acids at interfaces that are interacting. There are many ways to define interaction, but proximity [97] is a natural metric. That is, two residues are defined [97] as interacting if their $C_{\beta}$ coordinates differ by at most $6 \AA$ (with a similar scheme to include Gly). This notion is simplistic in that the $C_{\beta}$ atom is only the first in the sequence, but it is notable that the same sort of simple measure based on the initial segment is successful in other contexts [152].

Let us compare and contrast the two questions. The first question seeks to determine clues for protein-protein association by investigating all residues, suitably normalized. The second question assumes that proximity of sidechain pairs is a significant factor in protein-protein association, and thus looks for consequences of restricting to such pairs. Not surprisingly, each question returns different answers regarding the relative significance of different residues. In Figure 6.1, we depict the difference between the two data sets. We allow for the fact that being 'at the interface' may be differently defined in each case, leading to the possibility that neither set contains the other.


Figure 6.1: Cartoon showing possible relationship between two datasets.
The distribution of amino acid composition in proteins displays evolutionary trends [30], and this can require extra care to reveal subtle relationships. Here we limit our investigations to fairly strong trends for simplicity. However, the precise numerical data presented would differ if different databases were chosen for the primary data being used.

### 6.2 Interface propensity

The common belief is that hydrophobic residues on the surface of proteins are likely candidates to support interfaces in protein-protein association. In Section 6.3, we present evidence that supports this case with suitable clarifications. However, 83] presents data with a distinctively different conclusion, by focusing on all residues found at an interface and normalizing the relative abundance of residues at the interface by their over-all abundances. The residues with the highest relative propensity [83] to be at interfaces are, in decreasing order of frequency, Asn, Thr, Gly, Ser, Asp, Ala, and Cys, the group depicted in Figure 4.4. None of these residues is distinctively hydrophobic. This is quite a surprising result, and it demands an explanation.

To begin with, let us clarify the basic notions. If we have a dataset with $N$ different types of characteristics (e.g., $N=20$ and the characteristics are the different amino acids), then the frequency $f_{i}$ of the $i$-th characteristic is defined by

$$
\begin{equation*}
f_{i}=\frac{o_{i}}{\sum_{j=1}^{N} o_{j}} \tag{6.1}
\end{equation*}
$$

where $o_{j}$ is the number of occurrences of the $j$-th characteristic in the dataset. In some cases, frequencies are represented as percentages, in which case we simply multiply by 100 in (6.1).

If we have two datasets with the same characteristics, with frequencies $f_{i}$ and $g_{i}$, respectively, then one can define a relative frequency

$$
\begin{equation*}
r_{i}=f_{i} / g_{i} \tag{6.2}
\end{equation*}
$$

of the characteristics between the two datasets. There are some problems with this measure of occurrence. First of all, it might happen that $g_{k}=0$ for some $k$, making the interpretation difficult.

Related to this is the need for normalization in order to be able to compare two different comparisons. In [83], the following approach was taken.

Define a normalized relative propensity via

$$
\begin{equation*}
R_{i}=\frac{r_{i}}{\sum_{j=1}^{N} r_{j}} . \tag{6.3}
\end{equation*}
$$

These relative propensities sum to one, so we can think of them like ordinary frequencies. Similarly, we multiply by 100 in (6.1) to convert to percentages as the unit of "frequency."

If we apply this approach to datasets of proteins, and the characteristics are the different amino acid constituents, then we obtain the scheme used in 83]. In this case, the sum of the relative propensities (in percentage units) is one hundred, so the mean is five. In Table 6.1, data from 83 is presented in terms of the deviation of these relative propensities from the mean of five. That is, the data represent $100 R_{i}-5$.

The unusual ranking of residues in Table 6.1 was explained in 83] by noting that it correlates closely with the propensity to be engaged in under-wrapped backbone hydrogen bonds, among amino acids acting as either proton donors or acceptors for main-chain hydrogen bonds. These data are presented in the fifth column in Table 6.1, and the correlation is striking. Such bonds, in turn, are determinants of protein-protein associations, as discussed subsequently.

Since we expect a significant number of intermolecular hydrogen bonds (and some salt bridges) at interfaces, we might expect residues capable of making them (cf. Table 5.2) to be more likely at interfaces. But these residues are uniformly distributed in Table [6.1] not clustered near the top. If anything, the charged residues are clustered near the bottom. This implies that another factor determines the propensity to be at an interface, as suggested in 83, namely, the amount of wrapping a residue can provide.

As noted in [83], the seven residues in Figure 4.4, with the highest propensity for being engaged in under-desolvated hydrogen bonds, also have at most one torsional degree of freedom in their side chain. Thus, the entropic loss resulting from the conformational hindrance of the sidechains upon protein association is minimal with these sidechains, so that the energetic benefit of intermolecular protection of pre-formed hydrogen bonds is most beneficial. The only purely hydrophobic residue that has an appreciable propensity to be in an interface is Val (cf. Figure 4.5), with only one sidechain rotameric degree of freedom. Therefore, its conformational hindrance upon binding also entails minimal loss in conformational entropy.

Considering the residues ranked at the bottom of Table 6.1 demonstrates that hydrophobic residues on the protein surface are infrequent relative to their over-all abundance. This implies that they are negatively selected to be part of binding regions, and thus they must play a secondary role in terms of binding.

Note that the polar residues (Asn, Asp, Ser, Cys and Thr) with a minimal distance from their polar groups to the backbone are likely to be engaged in dehydrons, according to Table 6.1. It is presumed [83] that this arises not only because they have minimal nonpolar carbonaceous groups, but also because the relative proximity of their polar groups to a backbone hydrogen bond may limit further clustering of hydrophobic groups around the bond. Gly is itself the greatest under-wrapper and can even be thought of as polar due to the fact that the polar environment of the peptide bond

Draft: February 28, 2008, do not distribute

| 3-letter <br> code | 1-letter <br> code | Nonpolar <br> Carbons | Interface <br> Rel. Prop. | Dehydron <br> Rel. Prop. | Hydro- <br> pathy |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Asn | N | 1 | +1.28 | +1.63 | -3.5 |
| Thr | T | 1 | +1.10 | +1.41 | -0.7 |
| Gly | G | 0 | +0.99 | +1.42 | -0.4 |
| Ser | S | 0 | +0.60 | +0.80 | -0.8 |
| Asp | D | 1 | +0.34 | +0.76 | -3.5 |
| Ala | A | 1 | +0.29 | +0.6 | 1.8 |
| Cys | C | 1 | +0.25 | +0.24 | 2.5 |
| Val | V | 3 | +0.20 | -0.31 | 4.2 |
| Met | M | 3 | +0.10 | +0.10 | 1.9 |
| Tyr | Y | 6 | +0.10 | +0.10 | -1.3 |
| His | H | 1 | -0.25 | -0.25 | -3.2 |
| Pro | P | 3 | -0.25 | -0.25 | -1.6 |
| Trp | W | 7 | -0.33 | -0.4 | -0.9 |
| Arg | R | 2 | -0.35 | -0.4 | -4.5 |
| Leu | L | 4 | -0.35 | -1.10 | 3.8 |
| Phe | F | 7 | -0.40 | -0.40 | 2.8 |
| Lys | K | 3 | -0.42 | -0.38 | -3.9 |
| Glu | E | 2 | -0.50 | -0.11 | -3.5 |
| Gln | Q | 2 | -0.62 | -0.6 | -3.5 |
| Ile | I | 4 | -0.70 | -0.92 | 4.5 |

Table 6.1: Amino acids ranked according to their likelihood of being found at protein-protein interfaces. The second column indicates the number of carbon groups in the side chain. Interface and dehydron relative propensity (Rel. Prop.) is given as $R_{i}-5$ as in (6.3). Dehydron Propensity is also presented as frequency $f-5 ; 5 \%$ is the average propensity to be at interface or engaged in a dehydron. The hydropathy scale of Kyte et al. [143] is included for reference.

| Res. <br> Code | Pairing <br> Rel. Prop. | Pairing <br> Rel. Freq. | Pairing <br> Freq. $[97]$ | Total Abun- <br> dance [30] | Interface <br> Rel. Prop. | Rim/Core <br> freq. [40] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Cys | 5.4 | 2.40 | 1.87 | 0.78 | +0.25 | 0.45 |
| Trp | 1.9 | 1.60 | 1.63 | 1.02 | -0.33 | 0.32 |
| Pro | 1.7 | 1.55 | 6.74 | 4.35 | -0.25 | 1.24 |
| Ser | 1.5 | 1.50 | 7.01 | 4.66 | +0.60 | 1.04 |
| Asn | 1.3 | 1.46 | 4.90 | 3.36 | +1.28 | 1.19 |
| Thr | 1.1 | 1.41 | 6.87 | 4.87 | +1.10 | 1.19 |
| His | 0.76 | 1.33 | 2.56 | 1.92 | -0.25 | 0.52 |
| Tyr | 0.32 | 1.23 | 3.70 | 3.00 | +0.10 | 0.67 |
| Gly | 0.11 | 1.18 | 8.59 | 7.30 | +0.99 | 1.16 |
| Ala | 0.11 | 1.18 | 9.18 | 7.77 | +0.29 | 0.95 |
| Phe | -0.15 | 1.12 | 4.02 | 3.61 | -0.40 | 0.33 |
| Gln | -0.33 | 1.08 | 3.41 | 3.15 | -0.62 | 1.03 |
| Met | -0.72 | 0.99 | 2.38 | 2.41 | +0.10 | 0.54 |
| Asp | -0.98 | 0.93 | 5.06 | 5.42 | +0.34 | 1.48 |
| Val | -1.2 | 0.87 | 7.12 | 8.17 | +0.20 | 1.09 |
| Leu | -1.6 | 0.79 | 7.05 | 8.91 | -0.35 | 0.82 |
| Ile | -1.8 | 0.75 | 5.00 | 6.66 | -0.70 | 0.76 |
| Arg | -1.9 | 0.71 | 4.46 | 6.27 | -0.35 | 1.19 |
| Glu | -2.6 | 0.55 | 4.71 | 8.59 | -0.50 | 1.87 |
| Lys | -2.9 | 0.48 | 3.73 | 7.76 | -3.9 | 2.16 |

Table 6.2: Amino acids which occur in pairs at interfaces and their relative abundances. Primary data is taken from the indicated references. Relative Propensity is defined in (6.3) and Relative Frequency is defined in (6.2). Interface Relative Propensity from Table 6.1 is included for comparison.
is exposed; Ala is the penultimate under-wrapper and may also exhibit some of the polar qualities of Gly (cf. Section 4.5.1).

### 6.3 Amino acid pairs at interfaces

We now return to the second question raised at the beginning of the chapter regarding the amino acid composition for interacting pairs of amino acids at interfaces. We review the results in 97] which use proximity as an interaction metric in which two residues are defined as interacting if their $C_{\beta}$ coordinates differ by at most $6 \AA$. In this setting, some dominant residues are indeed hydrophobic, although it is pointed out in 97] that they "occurred more often in large contact surfaces, while polar residues prevailed in small surfaces," anticipating the subsequent discussion regarding "core" versus "rim" residues. We present in Table 6.2 the residues and their relative propensities, as defined in (6.3), in decreasing order.

Two of the residues in Table 6.2 with greatest relative propensity, namely Trp and Pro, are distinctively hydrophobic, as we might expect. However, these are also two of the most unique residues, as discussed in Section 4.5. Moreover, other high-ranking residues are as found in Table 6.1, The differences between this table and Table 6.1 reflect the fact that we are now asking about residues which are in proximity of a specific residue and thus may be interacting in some direct way.

Since Table 6.2 does not provide relative abundances directly, we need to say how these have been derived. The fundamental data in Table 6.2 is Table II on page 93 in 97, which lists the "contact" matrix $C_{i j}$. This is a matrix that counts the number of times that residue $i$ contacts (is within the proximity radius of) residue $j$. Summing a column (or row) of $C_{i j}$ and normalizing appropriately gives the total frequency $F_{i}$ of the $i$-th amino acid involved in such pairings. More precisely, to report frequencies as a percentage, define

$$
\begin{equation*}
F_{i}=100 \frac{\sum_{j=1}^{20} C_{i j}}{\sum_{i, j=1}^{20} C_{i j}} \tag{6.4}
\end{equation*}
$$

to be the amino acid pairing frequency, shown in the column entitled 'Pairing Freq. 97]' in Table 6.2.
The abundance of each amino acid in such pairings needs to be normalized by an appropriate measure. Here we have taken for simplicity the abundances published in [30] which are reproduced in the column entitled 'Total Abundance [30]' in Table 6.2. We do not claim that this provides the optimal reference to measure relative abundance in this setting, but it certainly is a plausible data set to use. The data shown in the column entitled 'Pairing Rel. Freq.' in Table 6.2 represents the ratio of $F_{i}$, defined in (6.4), to the abundances reported in (30].

The fact that Cys appears to have the highest relative abundance in pairs at interfaces reflects the simple fact that when Cys appears paired with another residue, it is unusually frequently paired with another Cys to form a disulfide bond (Section 4.2.2), as confirmed in 97].

### 6.4 Pair frequencies

In addition to looking at the frequencies of individual residues, one can also look at the frequencies of pairings. A standard tool for doing this is the odds ratio. Suppose that $f_{i}$ is the frequency of the $i$-th amino acid in some dataset, and suppose that $C_{i j}$ is the frequency of the pairing of the $i$-th amino acid with the $j$-th amino acid. Then the odds ratio $O_{i j}$ is defined as

$$
\begin{equation*}
O_{i j}=\frac{C_{i j}}{f_{i} f_{j}} \tag{6.5}
\end{equation*}
$$

and has the following simple interpretation. If the pairing of the $i$-th amino acid with the $j$-th amino acid were random and uncorrelated, then we would have $C_{i j}=f_{i} f_{j}$, and thus $O_{i j}=1$. Therefore an odds ratio bigger than one implies that the pairing is more common than would be expected for a random pairing, and conversely if it is less than one.

The log odds ratio is often defined by simply taking the logarithm of the odds ratio. This has the benefit of making the more likely pairings positive and the less likely pairings negative. In 97], a quantity $G_{i j}$ is defined by multiplying the log odds ratio by a numerical factor of ten.

Draft: February 28, 2008, do not distribute

It is noteworthy that the odds ratios indicated in Table III of [30 are all between one half and two. That is, there are no pairs which occur even as much as twice as frequently as would be expected randomly (or half as frequently).

The pair with the highest odds ratio (1.87) is Cys-Cys, a disulfide bridge. Although Cys is uncommon, when it does appear we can expect it to be involved in a disulfide bridge. The next highest odds ratio pair is Trp-Pro (1.42), which pairs two of the most unique sidechains (Sections 4.5.2 and 4.5.5). The lack of rotational freedom in proline may be significant since there is no entropic loss in the pairing, but the story is likely more complex, e.g., Trp-Pro can be involved in a sandwich [201].

The following four pairs with the next highest odds ratios involve charged residues: Asp-His (1.25), Arg-Trp (1.23), Asp-Ser (1.22) and Asp-Thr (1.21). The first of these is a salt-bridge, and the second is a charge-polar interaction known as a cation- $\pi$ interaction [95, 258, 48] (see Section (13.1) based on the special polarity of aromatic residues (Section 4.5.5). The latter two pairs are charged and polar residues as well. The next four pairs in ranking of odds ratio are Cys-Ser (1.20), Asp-Arg (1.19), Met-Met (1.16) and Cys-His (1.15). These show a similar mix of polar interactions.

There is no absolute scale on which to measure odds ratios, and the significance of any deviation from one is context dependent. But it is notable that the pair frequencies reported in [97] are much smaller than found for alpha helices or beta sheets [152]. The top thirty values for the odds ratios for amino acid pairs with $\theta<50$ (Section 4.4.1) are all greater than two, with the highest being 3.75 [152]. Moreover, the top fifteen values for the odds ratios for amino acid pairs with $\theta>155$, that is pairs in $\beta$ sheets, are all greater than two [152]. We interpret that to mean that the hydrophobic pairs involved in interfaces are more nearly random, none of which occur with very high odds ratios.

When we add the further analysis in 40 which differentiated the prevalence of core versus rim residues in protein interfaces, the picture is clarified. In [40], interface topology was characterized in detail, and it was found that interfaces could typically be described in terms of discrete patches of about $1600 \AA^{2}$ in area. For each patch, the boundary (rim) residues were identified versus the interior (core) residues. The statistics for amino acid preferences for the rim versus the core are reproduced in Table 6.2. There is a strong correlation between being charged or polar and preferring the rim, as indicated in Table 6.3.

Similarly, it is noteworthy that the variance in relative propensities is much greater for pairs of interacting residues at interfaces (Table 6.2) than it is for all (unrestricted) residues at interfaces (Table 6.1). This is not surprising because we have selected for a particular subset of pairs (instead of including all pairs). Combining the previous two observations, we can say that interacting pairs at the core of interfaces are more likely to involve a hydrophobic residue, but the pair compositions involving hydrophobes are nearly random.

In [97], the typical configuration of Arg-Trp is pictured, and similar polar pairings are highlighted, such as Lys-Lys (odds ratio 0.81).

Draft: February 28, 2008, do not distribute

| Res. <br> Code | Rel. <br> Prop. | Rel. <br> Freq. | Pair <br> Freq. 97$]$ | Total Abun- <br> dance [30] | Rim/Core <br> freq. 40] | Homodimer <br> Rim/Core [15] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Lys | -2.9 | 0.48 | 3.73 | 7.76 | 2.16 | 2.19 |
| Glu | -2.6 | 0.55 | 4.71 | 8.59 | 1.87 | 1.48 |
| Asp | -0.98 | 0.93 | 5.06 | 5.42 | 1.48 | 1.61 |
| Pro | 1.7 | 1.55 | 6.74 | 4.35 | 1.24 | 1.51 |
| Asn | 1.3 | 1.46 | 4.90 | 3.36 | 1.19 | 1.49 |
| Thr | 1.1 | 1.41 | 6.87 | 4.87 | 1.19 | 1.16 |
| Gly | 0.11 | 1.18 | 8.59 | 7.30 | 1.16 | 1.38 |
| Arg | -1.9 | 0.71 | 4.46 | 6.27 | 1.19 | 0.85 |
| Val | -1.2 | 0.87 | 7.12 | 8.17 | 1.09 | 0.83 |
| Ser | 1.5 | 1.50 | 7.01 | 4.66 | 1.04 | 1.15 |
| Gln | -0.33 | 1.08 | 3.41 | 3.15 | 1.03 | 1.22 |
| Ala | 0.11 | 1.18 | 9.18 | 7.77 | 0.95 | 0.93 |
| Leu | -1.6 | 0.79 | 7.05 | 8.91 | 0.82 | 0.61 |
| Ile | -1.8 | 0.75 | 5.00 | 6.66 | 0.76 | 0.55 |
| Tyr | 0.32 | 1.23 | 3.70 | 3.00 | 0.67 | 0.58 |
| Met | -0.72 | 0.99 | 2.38 | 2.41 | 0.54 | 0.68 |
| His | 0.76 | 1.33 | 2.56 | 1.92 | 0.52 | 0.85 |
| Cys | 5.4 | 2.40 | 1.87 | 0.78 | 0.45 | 0.81 |
| Phe | -0.15 | 1.12 | 4.02 | 3.61 | 0.33 | 0.40 |
| Trp | 1.9 | 1.60 | 1.63 | 1.02 | 0.32 | 0.60 |

Table 6.3: Amino acids which occur in pairs at interfaces and their relative abundances. Primary data is taken from the indicated references.

### 6.5 Comparisons and caveats

We have made several observations based on analyzing existing data sets. These conclusions should be viewed as preliminary since these data sets must be viewed as incomplete. Our primary intent was to introduce a methodology for exploring such data sets and to indicate the type of results that can be obtained.

Our basic analysis of pairwise interaction data was taken from 97. However, the methodology is quite similar to that of the earlier paper [236], although there are differences in the way the interior (and non-interior) sidechains in the interaction zone are defined. That is, the classification of rim and core residues in the interface [97] is different in definition from exposed and interior residues in the interface in [236], although similar in spirit. Figure 3B of [236] shows how the residues that are interacting (proximate) in an interface are very similar in composition to ones in the interior of proteins.

To illustrate the sensitivity of results depending on the database chosen, we review the results in [15] which is very similar in spirit to [40, the difference being the use of homodimers for the study of interfaces. In Table 6.3 we present this data, with the residues reordered to give the rim/core preferences in order for the data in [40] to facilitate comparison with the data in [15]. What we see is the same general trend, namely that charged and polar residues prefer the rim, but with changes in the particular rankings among the different groups. However, there is a significant reversal in the roles of arginine and valine [15].

The dissection trilogy is completed in [16] in which an attempt is made to determine aminoacid distributions for "nonspecific" interactions. This is intended to be a proxy for any surfaces which might bind however briefly to other protein surfaces. The dataset is determined by looking at crystal contact surfaces in the PDB. We leave as an exercise to compare the data for these surfaces with the other data presented here. See [16] for a comparison with the data in (40] and [15.

Protein-ligand interfaces differ in function, and interfaces with different function can have different composition. In [125, basic differences between protein-antibody and enzyme-inhibitor pairs, as well as others, are explored. Using more extensive datasets available more recently, this approach has been refined to allow classification of interface type based on aminoacid composition [182].

In [24], an attempt is made to identify so-called "hot spots" on protein surfaces. They report on the results of an experimental technique called alanine scanning in which residues are replaced by alanine and compared with the original protein by some activity assay. What they discover is that the most common sidechains at hot spots are the ones that are bulkiest, Trp, Tyr and Arg. This is not surprising since the replacement by Ala has the greatest change in geometry for these residues. However, such substitutions might be extremely rare. What might be a better test of importance would be other mutations, e.g., ones which do not change the volume or geometry of the side chain. Systematic replacement of all amino acids by all other amino acids is clearly an order of magnitude more work than just replacing by a fixed side chain. Having a better model of what governs protein-protein interactions could lead to a more directed study of sidechain mutation effects.

The aromatic sidechains do play a special role in protein interfaces through what is called a cation- $\pi$ interaction 95] (see Section 13.1). The special polar nature of the aromatic residues
(Section 4.5.5) provides the opportunity for interaction with positively charged (cation) residues (Lys, Arg, His). The cation- $\pi$ motifs play a special role in protein interfaces [48, 258]. The cation- $\pi$ interaction also has a significant role in $\alpha$-helix stabilization [220].

A study of the role of evolution on protein interface composition can be found in [36]. In [106, 158], interacting amino acids across interfaces are studied and compared with regard to conservation and hot spots.

Protein-protein interactions can be classified in different ways, e.g., by how transient they are, and studies have been done to examine differences in size of interaction zones and sidechain propensities [180, 181].

Identification of individual sidechains that may play the role of 'anchors' in protein-ligand recognition is studied in [202] via molecular dynamics simulations. Individual residues are identified that appear to fit into geometric features on paired protein surfaces both in crystal structures and in the dynamic simulations.

It is possible to refine the concept of sidechain interactions to one involving the interactions of individual atoms in structures. This approach has been suggested 44] as a way to discriminate between correct structures and incorrect ones. In [44], this concept was proposed as a way to critique structures being determined based on experimental imaging techniques, but the same concept could be applied to discriminate between native and decoy structures that are proposed via computational techniques.

### 6.6 Conclusions

Two main conclusions were obtained. The first is that residue hydrophobicity is not the primary variable that determines proximity of a residue to interaction sites. Instead, there is a different 'interactivity' order that governs the likelihood of an amino acid residue being in an active zone. This interactivity scale is related strongly to the number of nonpolar constituents of sidechains, which governs the local dielectric environment. Thus the likelihood of a residue being at an interface is to some extent inversely proportional to its hydrophobicity.

On the other hand, pairwise interactions with hydrophobic residues do play a secondary role in protein-protein interactions, especially in the interior, or core, regions of interaction domains. Moreover, their interactions tend to be less specific than might be the case in other pairings, such as in alpha helices and beta sheets. The role of hydrophobic sidechains in such interactions is not revealed by such an analysis. In particular, the definition of 'interaction' has been taken to be simple proximity, so it is misleading to infer that there is any identified form of interaction.

### 6.7 Exercises

Exercise 6.1 Compare the data for the surfaces in [15, 16, 40] by constructing a table analogous to Table 6.3.

Exercise 6.2 The aminoacid frequencies for different datasets constitute probability distributions on the set of aminoacids. Different datasets have different distributions. In [16], the distributions for

### 6.7. EXERCISES CHAPTER 6. DETERMINANTS OF PROTEIN-PROTEIN INTERFACES

nonspecific interaction surfaces are compared with the distributions for other surfaces 40, 15]. The comparison metric is the $L^{2}$ norm. Consider the effect of using the KL-divergence, Jensen-Shannon metric, and the earth-moving metric Section??.

Exercise 6.3 The frequency of location at interfaces provides a linear ranking (Table 6.1) of residues that can be useful in making predictions based on techniques from learning theory. As an example, consider using this to identify under-wrapped hydrogen bonds in $\alpha$-helices directly from sequence data. For an $\alpha$-helix, there will be hydrogen bonds formed between residues at a distance of 3, 4, or 5 residues. Generate data from a protein sequence by computing the product of the product of interface ranks of two neighbors. That is, for a sequence abcd define $x=\operatorname{rank}(a) \operatorname{rank}(b)$ and $y=\operatorname{rank}(c) \operatorname{rank}(d)$. Thus for every four letter sequence, we assign a pair of numbers $(x, y)$ in the unit square. If there is a dehydron associated with abcd then we expect ( $x, y$ ) near zero. Using data from the $P D B$, construct a support-vector machine to separate dehydrons from wrapped hydrogen bonds. Then use this machine to predict dehydrons in sequences for which the sequence is not known.

## Chapter 7

## Wrapping electrostatic bonds

For a protein structure to persist in water, its electrostatic bonds must be shielded from water attack [77, 84, 195, 238. This can be achieved through wrapping by nonpolar groups (such as $\left.\mathrm{CH}_{n}, n=1,2,3\right)$ in the vicinity of electrostatic bonds to exclude surrounding water [77]. Such desolvation enhances the electrostatic contribution and stabilizes backbone hydrogen bonds [18]. In a nonbonded state, exposed polar amide and carbonyl groups which are well wrapped are hindered from being hydrated and more easily return to the bonded state [49, as depicted in Figure 3.7.

The thermodynamic benefit associated with water removal from pre-formed structure makes under-wrapped proteins adhesive [70, 78, 80]. As shown in [77, under-wrapped hydrogen bonds (UWHB's) are determinants of protein associations. In Section 8.1. we describe the average adhesive force exerted by an under-wrapped hydrogen bond on a test hydrophobe.

The dielectric environment of a chemical bond can be enhanced in different ways, but wrapping is a common factor. There are different ways to quantify wrapping. Here we explore two that involve simple counting. One way of assessing a local environment around a hydrogen bond involves just counting the number of 'hydrophobic' residues in the vicinity of a hydrogen bond. This approach is limited for two reasons.

The first difficulty of this approach relates to the taxonomy of residues being used. The concept of 'hydrophobic residue' appears to be ambiguous for several residues. In some taxonomies, Arg, Lys, Gln, and Glu are listed as hydrophilic. However, we will see that they contribute substantially to a hydrophobic environment. On the other hand, Gly, Ala, Ser, Thr, Cys and others are often listed variously as hydrophobic or hydrophilic or amphiphilic. We have identified these five residues in Chapter 4 as among the most likely to be neighbors of underwrapped hydrogen bonds, as will be discussed at more length in Chapter 6. As noted in Section 4.5.1] glycine, and to a lesser extent alanine, can be viewed as polar, and hence hydrophilic, but alanine has only a nonpolar group in its sidechain representation and thus would often be viewed as hydrophobic.

A second weakness of the residue-counting method is that it is based solely on the residue level and does not account for more subtle, 'sub-residue' features. We will see that these limitations can be overcome to a certain extent with the right taxonomy of residues. However, we will also consider (Section [7.3) a measure of wrapping that looks into the sub-residue structure by counting all neighboring non-polar groups. The residue-counting method is included both for historical and

| atomic symbol | H | C | N | O | F | Na | Mg | P | S |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| electronegativity | 2.59 | 2.75 | 3.19 | 3.66 | 4.0 | 0.56 | 1.32 | 2.52 | 2.96 |
| nuclear charge | 1 | 6 | 7 | 8 | 9 | 11 | 12 | 15 | 16 |
| outer electrons | 1 | 4 | 5 | 6 | 7 | 1 | 2 | 5 | 6 |

Table 7.1: Electronegativity scale [192, 210] of principal atoms in biology. The 'outer electrons' row lists the number of electrons needed to complete the outer shell.
pedagogical reasons, although we would not recommend using it in general.
In our first measure of wrapping, we define precisely two classes of residues relevant to wrapping. This avoids potential confusion caused by using taxonomies of residues based on standard concepts. In Section [7.2.2, we show that this definition is sufficient to give some insight into protein aggregation and make predictions about protein behaviors.

However, it is also possible to provide a more refined measure that looks below the level of the residue abstraction and instead counts all non-polar groups, independent of what type of sidechain they inhabit. We present this more detailed approach in Section 7.3. We will show in Section 8.1 that there is a measurable force associated with an UWHB that can be identified by the second definition. Later we will define this force rigorously and use that as part of the definition of dehydron in Section 7.5. In Section 7.5, we will review a more sophisticated technique that incorporates the geometry of nonpolar groups as well as their number to assess the extent of protection via dielectric modulation.

### 7.1 Assessing polarity

The key to understanding hydrophobicity is polarity. Nonpolar groups repel water molecules (or at least do not attract them strongly) and polar groups attract them. We have already discussed the concept of polarity, e.g., in the case of dipoles (Section 3.3). Similarly, we have noted that certain sidechains, such as glutamine, are polar, even though there is no apparent charge difference in relevant molecules. Here we explain how such polarity can arise due to more subtle differences in charge distribution.

### 7.1.1 Electronegativity scale

The key to understanding the polarity of certain molecules is the electronegativity scale 192, [210], part of which is reproduced in Table 7.1. Atoms with similar electronegativity tend to form nonpolar groups, such as $C H_{n}$ and $C-S$. Atomic pairs with differences in electronegativity tend to form polar groups, such as $C-O$ and $N-H$. The scaling of the electronegativity values is arbitrary, and the value for fluorine has been taken to be exactly four.

Let us show how the electronegativity scale can be used to predict polarity. In a C-O group, the O is more electronegative, so it will pull charge from C , yielding a pair with a negative charge associated with the O side of the group, and a positive charge associated with the C side of the pair.

Draft: February 28, 2008, do not distribute

Similarly, in an N-H group, the N is more electronegative, so it pulls charge from the H , leaving a net negative charge near the N and a net positive charge near the H. In Section 7.1.2, we will see that molecular dynamics codes assign such partial charges. The electronegativity difference for $\mathrm{C}-\mathrm{O}$ is 0.91 , and for $\mathrm{N}-\mathrm{H}$ it is 0.6 . Thus, it would be expected to find larger partial charges for $\mathrm{C}-\mathrm{O}$ than for $\mathrm{N}-\mathrm{H}$, as we will see. Of course, the net charge for both $\mathrm{C}-\mathrm{O}$ and $\mathrm{N}-\mathrm{H}$ must be zero.

Only the differences in electronegativity have any chemical significance. But these differences can be used to predict the polarity of atomic groups, as we now illustrate for the carbonyl and amide groups. For any atom $X$, let $\mathcal{E}(X)$ denote the electronegativity of $X$. Since $\mathcal{E}(O)>\mathcal{E}(C)$, we conclude that the dipole of the carbonyl group $C-O$ can be represented by a positive charge on the carbon and a negative charge on the oxygen. Similarly, because $\mathcal{E}(N)>\mathcal{E}(H)$, the dipole of the amide group $N-H$ can be represented by a positive charge on the hydrogen and a negative charge on the nitrogen. A more detailed comparison of the electronegativities of $C, O, N$, and $H$ gives

$$
\begin{equation*}
\mathcal{E}(O)-\mathcal{E}(C)=3.66-2.75=0.91>0.60=3.19-2.59=\mathcal{E}(N)-\mathcal{E}(H) \tag{7.1}
\end{equation*}
$$

Thus we conclude that the charge difference in the dipole representation of the carbonyl group $(C-O)$ is larger than the charge difference in the dipole representation of the amide $(N-H)$ group.

It is beyond our scope to explain electronegativity here, but there is a simple way to comprehend the data. Electronegativity represents the power of an atom to attract electrons in a covalent bond 192. Thus a stronger positive charge in the nucleus would lead to a stronger attraction of electrons, which is reflected in the correlation between nuclear charge and electronegativity shown in Table 7.1. More precisely, there is a nearly linear relationship between the electronegativity scale and the number of electrons in the outer shell. The value for hydrogen can be explained by realizing that the outer shell is half full, as it is for carbon.

The atoms with a complete outer shell (helium, neon, argon, etc.) are not part of the electronegativity scale, since they have no room to put electrons that might be attracted to them. Similarly, atoms with just a few electrons in the outer shell seem to be more likely to donate electrons than acquire them, so their electronegativity is quite small, such as sodium and magnesium. Hydrogen and carbon are in the middle of the scale, not surprisingly, since they are halfway from being full and empty of electrons.

### 7.1.2 Polarity of groups

Using the electronegativity scale, we can now estimate the polarity of groups of atoms. For example, the near match of electronegativity of carbon and hydrogen leads to the correct conclusion that the carbonaceous groups $\mathrm{CH}_{n}, n=1,2,3$ are not polar, at least in appropriate contexts. The typically symmetric arrangement of hydrogens also decreases the polarity of a carbonaceous group, at least when the remaining $4-n$ atoms bonded to it are other carbons or atoms of similar electronegativity.

If a carbon is not covalently attached exclusively to carbon or hydrogen then it is likely polarized and carries a partial charge. Thus, $C_{\alpha}$ carbons in the peptide bonds of all residues are polar. Sidechain carbons are polar if they are covalently attached to heteroatoms such as N or O. Sulfur $(S)$ is a closer electronegative match with carbon and polarizes carbon to a lesser extent.

Draft: February 28, 2008, do not distribute

| Full name of <br> amino acid | three <br> letter | single <br> letter | The various PDB codes for the <br> nonpolar carbonaceous groups |
| :---: | :---: | :---: | :--- |
| Alanine | Ala | A | CB |
| Arginine | Arg | R | CB, CG |
| Asparagine | Asn | N | CB |
| Aspartate | Asp | D | CB |
| Cysteine | Cys | C | CB |
| Glutamine | Gln | Q | CB, CG |
| Glutamate | Glu | E | CB, CG |
| Glycine | Gly | G | NA |
| Histidine | His | H | CB |
| Isoleucine | Ile | I | CB1, CB2, CG, CD1 |
| Leucine | Leu | L | CB, CG, CD1, CD2 |
| Lysine | Lys | K | CB, CG, CD |
| Methionine | Met | M | CB (CG, CE) |
| Phenylalanine | Phe | F | CB, CG, CD1, CD2, CE1, CE2, CZ |
| Proline | Pro | P | CB, CG |
| Serine | Ser | S | NA |
| Threonine | Thr | T | CG2 |
| Tryptophan | Trp | W | CB, CG, CD2, CE1, CE2, CZ3, CH2 |
| Tyrosine | Tyr | Y | CB, CG, CD1, CD2, CE1, CE2 |
| Valine | Val | V | CB, CG1, CG2 |

Table 7.2: PDB codes for nonpolar carbonaceous groups. The carbonaceous groups surrounding the sulfer in Met may be considered polar.

| Full name of <br> compound | PDB <br> code | The various PDB codes for the <br> nonpolar carbonaceous groups |
| :---: | :---: | :--- |
| pyroglutamic acid | PCA | CB, CG |
| phosphorylated tyrosine | PTR | CB, CG, CD1, CD2, CE1, CE2 |
| staurosporine | STU | $\mathrm{C} i, i=1, \ldots, 7 ; i=11, \ldots, 16 ; \mathrm{C} 24, \mathrm{C} 26$ |

Table 7.3: Sample PDB codes and nonpolar carbonaceous groups for some nonstandard amino acids and other compounds.

| Residues | atom type | PDB codes | charge |
| :--- | :---: | :---: | :---: |
| ASP (GLU) | C | CG (CD) | 0.27 |
|  | OM | OD $i(\mathrm{OE} i) i=1,2$ | -0.635 |
| ASN (GLN) | NT | ND2 (NE2) | -0.83 |
|  | H | HD2i (HE2i), $i=1,2$ | 0.415 |
|  | C | CG (CD) | 0.38 |
|  | O | OD1 (OE1) | -0.38 |
| CYS | S | SG | -0.064 |
|  | H | HG | 0.064 |
| THR | CH1 | CB | 0.15 |
|  | OA | OG1 | -0.548 |
|  | H | HG1 | 0.398 |
| SER | CH2 | CB | 0.15 |
|  | OA | OG | -0.548 |
|  | H | HG | 0.398 |

Table 7.4: Partial charges from the Gromos force field for polar and negatively charged amino acids.

The case $\mathrm{CH}_{n}$ with $n=0$ is not encountered in biology unless the carbon is attached to at least one heteroatom.

To illustrate the polarity of the atoms not listed in Table 7.2 we present the partial charges of the remaining atoms as utilized in the Gromos code in Table 7.4 and Table 7.5. In Table 13.1 , partial charges for aromatic sidechains are listed.

In addition to the the charges shown for the individual sidechain atoms, the backbone is assigned partial charges as follows: the charges of the amide group are $\pm 0.28$ and the carbonyl group are $\pm 0.38$. That is, in the amide $(N-H)$ group, the $N$ is given a partial charge of -0.28 and the $H$ is given a partial charge of +0.28 . Similarly, in the carbonyl $(C-O)$ group, the $O$ is given a partial charge of -0.38 and the $C$ is given a partial charge of +0.38 . Note that the partial charges for $C-O$ are larger than the partial charges for $N-H$, in accord with our prediction using the electronegativity scale in (7.1).

The N-terminal and C-terminal groups also have appropriate modifications. The C-terminal oxygens have a charge of -0.635 , and the attached carbon has a charge of 0.27 . The N-terminal nitrogen has a charge of 0.129 , and the attached three hydrogens have a charge of 0.248 . All of the groups listed in Table 7.2 have zero partial charge.

### 7.2 Counting residues

In [74], the definition of 'well-wrapped' was based on the proximity of certain residues and defined in relation to the observed distribution of rapping among a large sample set of proteins. The extent of hydrogen-bond desolvation was defined by the number of residues $\rho_{R}$ with at least two nonpolar carbonaceous groups $\left(\mathrm{CH}_{n}, n=1,2,3\right)$ whose $\beta$-carbon is contained in a specific desolvation domain,

| Residue | atom type | PDB codes | charge |
| :--- | :---: | :---: | :---: |
| ARG | CH 2 | CD | 0.09 |
|  | NE | NE | -0.11 |
|  | C | CZ | 0.34 |
|  | NZ | $\mathrm{NH} i, i=1,2$ | -0.26 |
|  | H | $\mathrm{HE}, \mathrm{HH} i j, i, j=1,2$ | 0.24 |
| LYS | CH 2 | CE | 0.127 |
|  | NL | NZ | 0.129 |
|  | H | $\mathrm{HZ} i, i=1,3$ | 0.248 |
| HIS (A/B) | C | $\mathrm{CD} 2 / \mathrm{CG}$ | 0.13 |
|  | NR | $\mathrm{NE} 2 / \mathrm{ND} 1$ | -0.58 |
|  | CR 1 | CE 1 | 0.26 |
|  | H | $\mathrm{HD} 1 / \mathrm{HE} 2$ | 0.19 |

Table 7.5: Partial charges from the Gromos force field for positively charged amino acids. The partial charges for His represent two possible ionized states which carry neutral charge.
as depicted in Figure 7.1. In Section 7.1.2, we explained how to determine the polarity of groups using the electronegativity scale. The nonpolar carbonaceous groups are listed in Table 7.2 ,

The $C_{\alpha}$ carbons in all residues are covalently bonded to a nitrogen atom. The mismatch in electronegativity between carbon and nitrogen (Table 7.11) implies that the $C_{\alpha}$ carbons are polar and thus do not contribute to repelling water. Sidechain carbons are counted only if they are not covalently attached to heteroatoms such as N or O . The CH groups in serine and threonine are attached to an oxygen, which renders them polar. However, the CH groups in methionine attached to a sulfur are not polar. Similarly, a lone carbon that is attached to oxygens is also polar. Thus the seven residues listed in Figure 4.4 are eliminated from the group of wrappers.

### 7.2.1 Desolvation domain

The desolvation domain was chosen in [74] to be the union of two (intersecting) $7 \AA$-radius spheres centered at the $C_{\alpha}$-carbons of the residues paired by the hydrogen bond, as shown in Figure 7.1. The choice of the $C_{\alpha}$ carbons as the centers of the desolvation spheres is justified in Figure 7.2, These figures show that the center of the line joining the centers of the desolvation spheres is often the center of the hydrogen bonds in typical secondary structures. In the case of a parallel $\beta$-sheet, the desolvation domain is the same for two parallel hydrogen bonds. The radius represents a typical cutoff distance to evaluate interactions between nearby residues. $C_{\alpha}$-carbons which are neighboring in protein sequence are about $3.8 \AA$ apart (cf. Exercise 2.1). The distance between other $C_{\alpha}$-carbons is easily determined by datamining in the PDB (cf. Exercise 2.2).

An amide-carbonyl hydrogen bond was defined in 74 by an N-O (heavy-atom) distance within the range 2.6-3.4 (typical extreme bond lengths) and a 60 -degree latitude in the N-H-O angle (cf. Section 5.4). As a scale of reference, at maximum density, water occupies a volume that


Figure 7.1: Caricature showing desolvation spheres with various side chains. The open circles denote the nonpolar carbonaceous groups, and the solid circles represent the $C_{\alpha}$ carbons. The hydrogen bond between the amide ( $\mathrm{N}-\mathrm{H}$ ) and carbonyl ( $\mathrm{O}-\mathrm{C}$ ) groups is shown with a dashed line. Glycines appear without anything attached to the $C_{\alpha}$ carbon. There are 22 nonpolar carbonaceous groups in the union of the desolvation spheres and six sidechains with two or more carbonaceous groups whose $C_{\beta}$ carbon lie in the spheres.
(a)

(b)

(c)


Figure 7.2: The hydrogen bond (dashed line) configuration in (a) $\alpha$-helix, (b) antiparallel $\beta$-sheet, and (c) parallel $\beta$-sheet. A dotted line connects the $C_{\alpha}$ carbons (squares) that provide the centers of the spheres forming the desolvation domains in Figure 7.1. The amide ( $\mathrm{N}-\mathrm{H}$ ) groups are depicted by arrow heads and the carbonyl (O-C) groups are depicted by arrow tails.
corresponds to a cube of dimension just over $3.1 \AA$ on a side (cf. Section 10.7.3).
The average extent of desolvation, $\rho_{R}$, over all backbone hydrogen bonds of a monomeric structure can be computed from any set of structures. In [74], a nonredundant sample of 2811 PDBstructures was examined. The average $\rho_{R}$ over the entire sample set was found to be 6.6 [74]. For any given structure, the dispersion (standard deviation) $\sigma$ from the mean value of $\rho_{R}$ for that structure can be computed. The dispersion averaged over all sampled structures was found to be $\sigma=1.46$ [74]. These statistics suggested a way to identify the extreme of the wrapping distribution as containing three or fewer wrapping residues in their desolvation domains. This can be interpreted as defining underwrapped as $\rho_{R}$ values that are more than two standard deviations from the mean.

The distribution of the selected proteins as a function of their average wrapping as measured by $\rho_{R}$ is shown in Fig. 5 in [74]. The probability distribution has a distinct inflection point at $\rho=6.2$. Over $90 \%$ of the proteins studied have $\rho_{R}>6.2$, and none of these are yet known to yield amyloid aggregation under physiological conditions. In addition, individual sites with low wrapping on selected proteins were examined and found to correlate with known binding sites.

In Section 7.2.2, we will see that the known disease-related amyloidogenic proteins are found in the relatively under-populated $3.5<\rho_{R}<6.2$ range of the distribution, with the cellular prion proteins located at the extreme of the spectrum $\left(3.5<\rho_{R}<3.75\right)$. We discuss there the implications regarding a propensity for organized aggregation. Approximately $60 \%$ of the proteins in the critical region $3.5<\rho_{R}<6.2$ which are not known to be amyloidogenic are toxins whose structures are stabilized mostly by disulfide bonds.

To further assess the virtues of the residue-based assessment of wrapping, we review additional results and predictions of [74].

### 7.2.2 Predicting aggregation

Prediction of protein aggregation can be based on locating regions of the protein surface with high density of defects which may act as aggregation sites [114, 139, 166]. Figure 3a of [74] depicts the (many) UWHB's for the human cellular prion protein (PDB file 1QM0) [200, 204, 260]. Over half of the hydrogen bonds are UWHB's, indicating that many parts of the structure must be open to water attack. For example, $\alpha$-helix 1 has the highest concentration of UWHB's, and therefore may be prone to structural rearrangement.

In helix 1 (residues 143 to 156), all of the hydrogen bonds are UWHB's, and this helix has been identified as undergoing an $\alpha$-helix to $\beta$-strand transition [200, 204, 260]. Furthermore, helix 3 (residues 199 to 228) contains a significant concentration of UWHB's at the C-terminus, a region assumed to define the epitope for protein-X binding [200]. The remaining UWHB's occur at the helix-loop junctures and may contribute to flexibility required for rearrangement.

The average underwrapping of hydrogen bonds in an isolated protein may be a significant indicator of aggregation, but it is not likely to be sufficient to determine amyloidogenic propensity. For instance, protein L (PDB file 2PTL) is not known to aggregate even though its $\rho_{R}=5.06$ value is outside the standard range of sufficient wrapping. Similarly, trp-repressor (PDB file 2WRP) has $\rho_{R}=5.29$, and the factor for inversion stimulation (PDB file 3FIS) has $\rho_{R}=4.96$. Many neurotoxins (e.g., PDB file 1CXO with $\rho_{R}=3.96$ ) are in this range as well.

The existence of short fragments endowed with fibrillogenic potential [14, 52, 63, 101, 114, [166, 139] suggests a localization or concentration of amyloid-related structural defects. In view of this, a local wrapping parameter, the maximum density $\delta_{\max }$ of UWHB's on the protein surface was introduced [74]. A statistical analysis involving $\delta_{\max }$ [74] established that a threshold $\delta_{\max }>$ $0.38 / \mathrm{nm}^{2}$ distinguishes known disease-related amyloidogenic proteins from other proteins with a low extent of hydrogen bond wrapping. On the basis of a combined assessment, identifying both low average wrapping and high maximum density of underwrapping, it was predicted [74] that six proteins might posses amyloidogenic propensity. Three of them,

- angiogenin (cf. PDB files 1B1E and 2ANG),
- meizothrombin (cf. PDB file 1A0H), and
- plasminogen (cf. PDB file 1B2I),
are involved in some form of blood clotting or wound healing, and not something related to disease.
Not all protein aggregation is related to disease. Angiogenesis refers to the growth of new capillaries from an existing capillary network, and many processes involve this, including wound healing. Angiogenin is only one of many proteins involved in the angiogenesis process, but it appears to have certain unique properties [146. Meizothrombin is formed during prothrombin activation, and is known to be involved in blood clotting [129] and is able to bind to procoagulant phospholipid membranes [194]. Plasminogen has been identified as being a significant factor in wound healing [207].


### 7.3 Counting nonpolar groups

A more refined measure of hydrogen-bond protection has been proposed based on the number of vicinal nonpolar groups [70, 77]. The desolvation domain for a backbone hydrogen bond is defined again as the union of two intersecting spheres centered at the $\alpha$-carbons of the residues paired by the hydrogen bond, as depicted in Figure [7.1] In this case, all of the dark circles are counted, whether or not the base of the sidechain lies within the desolvation domain. The extent of intramolecular desolvation of a hydrogen bond, $\rho_{P G}$, is defined by the number of sidechain nonpolar groups $\left(\mathrm{CH}_{n}\right.$, $n=1,2,3$ ) in the desolvation domain (see Table 7.2).

The distribution of wrapping for a large sample of non-redundant proteins is given in Figure 12.1 for a radius of $6 \AA$ for the definition of the desolvation domain. In [78], an UWHB was defined by the inequality $\rho_{P G}<12$ for this value of the radius. Statistical inferences involving this definition of $\rho_{P G}$ were found to be robust to variations in the range $6.4 \pm 0.6 \AA$ for the choice of desolvation radius [77, 84]. In Figure 7.3 the distribution of wrapping is presented for a particular PDB file.

### 7.3.1 Distribution of wrapping for an antibody complex

It is instructive to consider wrapping of hydrogen bonds from a more detailed statistical point of view. In Figure 7.3 the distribution of wrapping is presented for the antibody complex whose

Draft: February 28, 2008, do not distribute


Figure 7.3: Distribution of wrapping for PDB file 1P2C. There are three chains: light, heavy chains of the antibody, and the antigen (HEL) chain. The desolvation radius is $6.0 \AA$. Smooth curves (7.2) are added as a guide to the eye.
structure is recorded PDB file 1P2C. There are three chains, two in the antibody (the light and heavy chains), and one in the antigen, hen egg-white lysozyme (HEL).

What is striking about the distributions is that they are bi-modal, and roughly comparable for all three chains. We have added a smooth curve representing the distributions

$$
\begin{equation*}
d_{i}(r)=a_{i}\left|r-r_{0}\right| e^{-\left|r-r_{0}\right| / w_{i}} \tag{7.2}
\end{equation*}
$$

to interpolate the actual distributions. More precisely, $d_{1}$ represents the distribution for $r<r_{0}$, and $d_{2}$ represents the distribution for $r>r_{0}$. The coefficients chosen were $w_{1}=2.2$ and $w_{2}=3.3$. The amplitude coefficients were $a_{1}=12$ and $a_{2}=9$, and the offset $r_{0}=18$ for both distributions. In this example, there seems to be a line of demarcation at $\rho=18$ between hydrogen bonds that are well wrapped and those that are underwrapped.

The distributions in Figure 7.3 were computed with a desolvation radius of $6.0 \AA$. Larger desolvation radii were also used, and the distributions are qualitatively similar. However the sharp gap at $\rho=18$ becomes blurred for larger values of the desolvation radius.

Draft: February 28, 2008, do not distribute

### 7.4 Residues versus polar groups

The two measures considered here for determining UWHB's share some important key features. Both count sidechain indicators which fall inside of desolvation domains that are centered at the $C_{\alpha}$ backbone carbons. The residue-based method counts the number of residues (of a restricted type) whose $C_{\beta}$ carbons fall inside the desolvation domain. The group-based method counts the number of carbonaceous groups that are found inside the desolvation domain.

We observed that the average measure of wrapping based on counting residues was $\rho_{R}=6.6$, whereas the average measure of wrapping based on counting non-polar groups is $\rho_{P G}=15.9$. The residues in the former count represent at least two non-polar groups, so we would expect that $\rho_{P G}>2 \rho_{R}$. We see that this holds, and that the excess corresponds to the fact that some residues have three or more non-polar groups. Note that these averages were obtained with different desolvation radii, $6.0 \AA$ for $\rho_{P G}$ and $7.0 \AA$ for $\rho_{R}$. Adjusting for this difference would make $\rho_{P G}$ even larger, indicating an even greater discrepancy between the two measures. This implies that $\rho_{P G}$ provides a much finer estimation of local hydrophobicity.

The structural analysis in [74] identified site mutations which might stabilize the part of the cellular prion protein (PDB file 1QM0) believed to nucleate the cellular-to-scrapie transition. The (Met134, Asn159)-hydrogen bond has a residue wrapping factor of only $\rho_{R}=3$ and is only protected by Val161 and Arg136 locally, which contribute only a minimal number (five) of non-polar carbonaceous groups. Therefore it is very sensitive to mutations that alter the large-scale context preventing water attack. It was postulated in [74] that a factor that triggers the prion disease is the stabilization of the (Met134, Asn159) $\beta$-sheet hydrogen bond by mutations that foster its desolvation beyond wild-type levels.

In the wild type, the only nonadjacent residue in the desolvation domain of hydrogen bond (Met134, Asn159) is Val210, thus conferring marginal stability with $\rho_{R}=3$. Two of the three known pathogenic mutations (Val210Ile and Gln217Val) would increase the number of non-polar carbonaceous groups wrapping the hydrogen bond (Met134, Asn159), even though the number of wrapping residues would not change. Thus we see a clearer distinction in the wrapping environment based on counting non-polar carbonaceous groups instead of just residues.

The third known pathogenic mutation, Thr183Ala, may also improve the wrapping of the hydrogen bond (Met134, Asn159) even though our simple counting method will not show this, as both Thr and Ala contribute only one nonpolar carbonaceous group for desolvation. However, Ala is four positions below Thr in Table 6.1 and is less polar than Thr. Table 6.1 reflects a more refined notion of wrapping for different sidechains, but we do not pursue this here.

### 7.5 Defining dehydrons via geometric requirements

The enhancement of backbone hydrogen-bond strength and stability depends on the partial structuring, immobilization or removal of surrounding water. In this section we review an attempt [79] to quantify this effect using a continuous representation of the local solvent environment surrounding backbone hydrogen bonds [33, 70, 77, 84, 113, 184, 244. The aim is to estimate the changes in the permittivity (or dielectric coefficient) of such environments and the sensitivity of
the Coulomb energy to local environmental perturbations caused by protein interactions [70, 84]. However, induced-fit distortions of monomeric structures are beyond the scope of these techniques.

The new ingredient is a sensitivity parameter $M_{k}$ assessing the net decrease in the Coulomb energy contribution of the $k$-th hydrogen bond which would result from an exogenous immobilization, structuring or removal of water due to the approach by a hydrophobic group. This perturbation causes a net decrease in the permittivity of the surrounding environment which becomes more or less pronounced, depending on the pre-existing configuration of surrounding hydrophobes in the monomeric state of the protein. In general, nearby hydrophobic groups induce a structuring of the solvent needed to create a cavity around them and the net effect of this structuring is a localized reduction in the solvent polarizability with respect to reference bulk levels. This structuring of the solvent environment should be reflected in a decrease of the local dielectric coefficient $\epsilon$. This effect has been quantified in recent work which delineated the role of hydrophobic clustering in the enhancement of dielectric-dependent intramolecular interactions [70, 84].

We now describe an attempt to estimate $\epsilon$ as a function of the fixed positions $\left\{r_{j}: j=1, \ldots, n_{k}\right\}$ of surrounding nonpolar hydrophobic groups $\left(\mathrm{CH}_{n}\right.$, with $n=1,2,3$, listed in Table 7.2). The simpler estimates of wrapping considered so far could fail to predict an adhesive site when it is produced by an uneven distribution of desolvators around a hydrogen bond, rather than an insufficient number of such desolvators. Based on the fixed atomic framework for the monomeric structure, we now identify Coulomb energy contributions from intramolecular hydrogen bonds that are most sensitive to local environmental perturbations by subsuming the effect of the perturbations as changes in $\epsilon$.

Suppose that the carbonyl oxygen atom is at $\mathbf{r}_{O}$ and that the partner hydrogen net charge is at $\mathbf{r}_{H}$. The electrostatic energy contribution $E_{\text {CouL }}(k, \mathbf{r})$ for this hydrogen bond in a dielectric medium with dielectric permittivity $\epsilon(\mathbf{r})$ is approximated (see Chapter 17) by

$$
\begin{equation*}
E_{\mathrm{COUL}}(\mathbf{r})=\frac{-1}{4 \pi \epsilon(\mathbf{r})} \frac{q q^{\prime}}{\left|\mathbf{r}_{O}-\mathbf{r}_{H}\right|} \tag{7.3}
\end{equation*}
$$

where $q, q^{\prime}$ are the net charges involved and where $|\cdot|$ denotes the Euclidean norm.
Now suppose that some agent enters in a way to alter the dielectric field, e.g., a hydrophobe that moves toward the hydrogen bond and disrupts the water that forms the dielectric material. This movement will alter the Coulombic energy as it modifies $\epsilon$, and we can use equation (7.3) to determine an equation for the change in $\epsilon$ in terms of the change in $E_{\text {Coul }}$. Such a change in $E_{\text {Coul }}$ can be interpreted as a force (cf. Chapter (3). We can compute the resulting effect as a derivative with respect to the position $R$ of the hydrophobe:

$$
\begin{equation*}
\nabla_{R}(1 / \epsilon(\mathbf{r}))=\frac{4 \pi\left|\mathbf{r}_{O}-\mathbf{r}_{H}\right|}{q q^{\prime}}\left(-\nabla_{R} E_{\mathrm{COUL}}(\mathbf{r})\right)=\frac{4 \pi\left|\mathbf{r}_{O}-\mathbf{r}_{H}\right|}{q q^{\prime}} F(\mathbf{r}) \tag{7.4}
\end{equation*}
$$

where $F(\mathbf{r})=-\nabla_{R} E_{\text {COUL }}(\mathbf{r})$ is a net force exerted on the hydrophobe by the fixed pre-formed hydrogen bond. This force represents a net 3-body effect [70, involving the bond, the dielectric material (water) and the hydrophobe. If $E_{\text {Coul }}$ is decreased in this process, the hydrophobe is attracted to the hydrogen bond because in so doing, it decreases the value of $E_{\text {CouL }}(\mathbf{r})$.

To identify the 'opportune spots' for water exclusion on the surface of native structures we need to first cast the problem within the continuous approach, taking into account that $1 / \epsilon$ is the factor
in the electrostatic energy that subsumes the influence of the environment. Thus to identify the dehydrons, we need to determine for which Coulombic contributions the exclusion or structuring of surrounding water due to the proximity of a hydrophobic 'test' group produces the most dramatic increase in $1 / \epsilon$. The quantity $M_{k}$ was introduced [79] to quantify the sensitivity of the Coulombic energy for the $k$-th backbone hydrogen bond to variations in the dielectric. For the $k$-th backbone hydrogen bond, this sensitivity is quantified as follows.

Define a desolvation domain $D_{k}$ with border $\partial D_{k}$ circumscribing the local environment around the $k$-th backbone hydrogen bond, as depicted in Figure 7.1. In 79, a radius of $7 \AA$ was used. The set of vector positions of the $n_{k}$ hydrophobic groups surrounding the hydrogen bond is extended from $\left\{\mathbf{r}_{j}: j=1,2, \ldots, n_{k}\right\}$ to $\left\{\mathbf{r}_{j}: j=1,2, \ldots, n_{k} ; R\right\}$ by adding the test hydrophobe at position $R$. Now compute the gradient $\left.\nabla_{R}(1 / \epsilon)\right|_{R=R_{o}}$, taken with respect to a perpendicular approach by the test hydrophobe to the center of the hydrogen bond at the point $R=R_{o}$ located on the circle consisting of the intersection $C$ of the plane perpendicular to the hydrogen bond with the boundary $\partial D_{k}$ of the desolvation domain. Finally, determine the number

$$
\begin{equation*}
M_{k}=\max \left\{\left|\nabla_{R}(1 / \epsilon)\right|_{R=R_{o}} \mid: R_{o} \in C\right\} . \tag{7.5}
\end{equation*}
$$

The number $M_{k}$ quantifies the maximum alteration in the local permittivity due to the approach of the test hydrophobe in the plane perpendicular to the hydrogen bond, centered in the middle of the bond, at the surface of the desolvation domain.

The quantity $M_{k}$ may be interpreted in physical terms as a measure of the maximum possible attractive force exerted on the test hydrophobic group by the pre-formed hydrogen bond. The only difficulty in estimating $M_{k}$ is that it requires a suitable model of the dielectric permittivity $\epsilon$ as a function of the geometry of surrounding hydrophobic groups. We will consider the behavior of the dielectric permittivity more carefully in Chapter [17, but for now we consider a heuristic model used in (79].

The model in [79] for the dielectric may be written

$$
\begin{equation*}
\epsilon^{-1}=\left(\epsilon_{o}^{-1}-\epsilon_{w}^{-1}\right) \Omega\left(\left\{\mathbf{r}_{j}\right\}\right) \Phi\left(\mathbf{r}_{H}-\mathbf{r}_{O}\right)+\epsilon_{w}^{-1} \tag{7.6}
\end{equation*}
$$

where $\epsilon_{w}$ and $\epsilon_{o}$ are the permittivity coefficients of bulk water and vacuum, respectively, and

$$
\begin{equation*}
\Omega\left(\left\{\mathbf{r}_{j}\right\}\right)=\prod_{j=1, \ldots, n_{k}}\left(1+e^{-\left|\mathbf{r}_{O}-\mathbf{r}_{j}\right| / \Lambda}\right)\left(1+e^{-\left|\mathbf{r}_{H}-\mathbf{r}_{j}\right| / \Lambda}\right) \tag{7.7}
\end{equation*}
$$

provides an estimate of the change in permittivity due to the hydrophobic effects of the carbonaceous groups. In [79], a value of $\Lambda=1.8 \AA$ was chosen to represent the characteristic length associated with the water-structuring effect induced by the solvent organization around the hydrophobic groups. Further, a cut-off function

$$
\begin{equation*}
\Phi(\mathbf{r})=(1+|\mathbf{r}| / \xi) e^{-|\mathbf{r}| / \xi} \tag{7.8}
\end{equation*}
$$

where $\xi=5 \AA$ is a water dipole-dipole correlation length, approximates the effect of hydrogen bond length on its strength [79].

Draft: February 28, 2008, do not distribute

### 7.5. DEFINING DEHYDRONS VIA GEHAIRTITRHCTREORAARPINEANESECTROSTATIC BONDS



Figure 7.4: The function $\omega(x, y)$ plotted as a function of the distance along the $x$-axis connecting $r_{H}$ and $r_{O}$, for three different values of the distance $y$ from that axis: $y=1$ (solid line), $y=2$ (dashed line), $y=3$ (dotted line). The coordinates have been scaled by $\Lambda$ and the value of $\left|\mathbf{r}_{O}-\mathbf{r}_{H}\right|=1$ was assumed.

We can write the key expression $\Omega$ in (7.7) as

$$
\begin{equation*}
\Omega\left(\left\{\mathbf{r}_{j}\right\}\right)=\prod_{j=1, \ldots, n_{k}} \omega\left(\mathbf{r}_{j}\right), \tag{7.9}
\end{equation*}
$$

where the function $\omega$ is defined by

$$
\begin{equation*}
\omega(\mathbf{r})=\left(1+e^{-\left|\mathbf{r}_{O}-\mathbf{r}\right| / \Lambda}\right)\left(1+e^{-\left|\mathbf{r}_{H}-\mathbf{r}\right| / \Lambda}\right) . \tag{7.10}
\end{equation*}
$$

The function $\omega$ is never smaller than one, and it is maximal in the plane perpendicular to the line connecting $r_{H}$ and $r_{O}$. Moreover, it is cylindrically symmetric around this axis. The values of $\omega$ are plotted in Figure 7.4 as a function of the distance from the perpendicular bissector of the axis connecting $\mathbf{r}_{H}$ and $\mathbf{r}_{O}$, for three different values of the distance $y$ from the line connecting $\mathbf{r}_{H}$ and $\mathbf{r}_{O}$.

We see that the deviation in $\omega$ provides a strong spatial dependence on the dielectric coefficient in this model. Thus hydrophobes close to the plane bissecting the line connecting $r_{H}$ and $r_{O}$ are counted more strongly than those away from that plane, for a given distance from the axis, and those closer to the line connecting $r_{H}$ and $r_{O}$ are counted more strongly than those further away. When the product $\Phi \Omega=1$, we get $\epsilon=\epsilon_{o}$ reflecting the maximal amount of water exclusion possible. Correspondingly, if $\Phi \Omega=0, \epsilon=\epsilon_{w}$ indicating a dielectric similar to bulk-water. Thus bigger values of $\Omega$ correspond to the effect of wrapping.

The definition (7.6) of the dielectric has not been scaled in a way that assures a limiting value of $\epsilon=\epsilon_{o}$. However, since we are only interested in comparing relative dielectric strength, this scaling is inessential. What matters is that larger values of $\Omega$ correspond to a lower dielectric and thus stronger bonds.

Draft: February 28, 2008, do not distribute

The computation of $M_{k}$ involves computing the gradient of

$$
\begin{equation*}
\Omega\left(\left\{\mathbf{r}_{1}, \ldots, \mathbf{r}_{n_{k}}, R\right\}\right)=\omega(R) \Omega\left(\left\{\mathbf{r}_{1}, \ldots, \mathbf{r}_{n_{k}}\right\}\right) \tag{7.11}
\end{equation*}
$$

with respect to $R$. Due to the cylindrical symmetry of $\omega,\left|\nabla_{R} \omega\right|_{R=R_{o}}$ is a constant depending only on the desolvation radius $\left|R_{o}\right|$ and the hydrogen bond length $\left|\mathbf{r}_{O}-\mathbf{r}_{H}\right|$ for all $R_{o} \in C$. Thus, for a fixed desolvation radius $\left|R_{o}\right|, M_{k}$ may be written as a function of $\left|\mathbf{r}_{O}-\mathbf{r}_{H}\right|$ times $\Omega\left(\left\{\mathbf{r}_{1}, \ldots, \mathbf{r}_{n_{k}}\right\}\right)$ when using the model (7.6).

A sensitivity threshold for hydrogen bonds was established in [79] by statistical analysis on a sample of native structures for soluble proteins. Only $8 \%$ of backbone hydrogen bonds from a sample of 702 proteins, of moderate sizes $(52<N<110)$ and free from sequence redundancies [112], were found to be highly sensitive in the sense that

$$
\begin{equation*}
M_{k}>\lambda / 10 \tag{7.12}
\end{equation*}
$$

where $\lambda$ was defined to be

$$
\begin{equation*}
\lambda=\frac{\epsilon_{o}^{-1}-\epsilon_{w}^{-1}}{2 \AA} \tag{7.13}
\end{equation*}
$$

On the other hand, $91.6 \%$ of backbone hydrogen bonds were found to be relatively insensitive to water removal, namely,

$$
\begin{equation*}
0<M_{k}<\lambda / 100 \tag{7.14}
\end{equation*}
$$

This remarkable separation in the (nearly bimodal) distribution of sensitivities led [79] to the definition of a dehydron as a backbone hydrogen bond satisfying (7.12).

### 7.6 Exercises

Exercise 7.1 It was predicted [74] that the three proteins

- anti-oncogene $A(P D B$ file $1 A 1 U)$;
- $R A D R$ zinc finger peptide (PDB file $1 A 1 K$ ) and
- rubredoxin (PDB file 1B20).
might have amyloidogenic tendencies. Investigate these three proteins to see why this might be the case.

Exercise 7.2 In Section 4.4.3, we noted that sidechains have different conformations. Determine the number of different rotameric states possible for each sidechain (hint: read [156]). Compare the number of rotameric degrees of freedom for the seven residues listed in Figure 4.4 with the remaining group of thirteen sidechains.

Exercise 7.3 In Figure 7.2(a), it appears that the dotted line joining the two $C_{\alpha}$ cabons intersects the dashed line joining the amide and carbonyl groups. By searching the PDB, determine the distribution of distances between the midpoints of these two lines for $\alpha$-helices.

Draft: February 28, 2008, do not distribute

Exercise 7.4 Explain whether you would expect methyl flouride to have a polar carbon, based on the electronegativity scale.

Exercise 7.5 Determine the extent to which the rapping in a given protein is bimodal, as depicted in Figure 7.3. Determine a way to measure consistently the degree to which a distribution is bimodal, and survey a large set of PDB structures with this measure.

## Chapter 8

## Stickiness of dehydrons

We have explained why under-wrapped hydrogen bonds benefit from the removal of water. This makes them susceptible to interaction with molecules that can replace water molecules in the vicinity of the hydrogen bond. Conceptually, this implies that under-wrapped hydrogen bonds attract entities that can dehydrate them. Thus they must be sticky. If so, it must be possible to observe this experimentally. Here we review several papers that substantiate this conclusion. One of them involves a mesoscopic measurement of the force associated with a dehydron [78]. A second presents data on the direct measurement of the dehydronic force using atomic force microscopy 68. Another paper examines the effect of such a force on a deformable surface 80 .

### 8.1 Surface adherence force

We defined the notion of an under-wrapped hydrogen bond by a simple counting method in Chapter 7 and have asserted that there is a force associated with UHWB's. Here we describe measurements of the adhesion of an under-wrapped hydrogen bond by analyzing the flow-rate dependence of the adsorption uptake of soluble proteins onto a phospholipid bilayer.

### 8.1.1 Biological surfaces

The principal biological surface of interest is the cell membrane. This is a complex system, but a key component is what is called a phospholipid bilayer. The term lipid refers to a type of molecule that is a long carbonaceous polymer with a polar (phospho) group at the 'head.' This it is hydrophobic at one end and hydrophilic at the other. These molecules align to form a complex that could be described as a bundle of pencils, with the hydrophilic head group (the eraser) at one side of the surface and the hydrophobic 'tail' on the other side. These bundles can grow to form a surface when enough pencils are added. A second surface can form in the opposite orientation, with the two hydrophobic surfaces in close proximity. This results in a membrane that is hydrophilic on both sides, and thus can persist in an aqueous environment.

One might wonder what holds together a lipid bilayer. We have noted that there is a significant volume change when a hydrophobic molecule gets removed from water contact in Section 4.4.4.

The volume change causes self-assembly of lipids and provides a substantial pressure that holds the surface together. The architecture of a lipid bilayer is extremely adaptive. For example, a curved surface can be formed simply by allocating more lipid to one side than the other. Moreover, it easily allows insertion of other molecules of complex shape but with other composition. Much of a cell membrane is lipid, but there are also proteins with various functions as well as other molecules such as cholesterol. However, a simple lipid bilayer provides a useful model biological surface.

### 8.1.2 Soluble proteins on a surface

One natural experiment to perform is to release soluble proteins in solution near a lipid bilayer and to see to what extent they attach to the bilayer. Such an experiment [71] indicated a significant correlation between the under-wrapping of hydrogen bonds and bilayer attachment. The results were explained by assuming that the probability of successful landing on the liquid-solid interface is proportional to the ratio of UWHB's to all hydrogen bonds on the protein surface. Here, the number of surface hydrogen bonds is taken simply as a measure of the surface area. Thus the ratio can be thought of as an estimate of the fraction of the surface of the protein that is under-wrapped. The experiments in [71] indicated that more dehydrons lead to more attachments, strongly suggesting that dehydrons are sticky. However, such indications were only qualitative.

A more refined analysis of lipid bilayer experiments was able to quantify a force of attachment [78]. The average magnitude of the attractive force exerted by an UWHB on a surface was assessed based on measuring the dependence of the adsorption uptake on the flow rate of the ambient fluid above the surface. The adhesive force was measured via the decrease in attachment as the flow rate was increased.

Six proteins were investigated in 78, as shown in Table 8.1, together with their numbers of well-wrapped hydrogen bonds as well as dehydrons. The UWHB's for three of these are shown in Fig. 1a-c in 78. The particular surface was a Langmuir-Blodgett bilayer made of the lipid DLPC (1,2 dilauroyl-sn-glycero-3 phosphatidylcholine) [206]. We now review the model used in [78] to interpret the data.

### 8.2 A two-zone model

In [78], a two-zone model of surface adhesion was developed. The first zone deals with the experimental geometry and predicts the number of proteins that are likely to reach a fluid boundary layer close to the lipid bylayer. The probability $\Pi$ of arrival is dependent on the particular experiment, so we only summarize the model results from [78]. The second zone is the fluid boundary layer close to the lipid bilayer, where binding can occur. In this layer, the probability $P$ of binding is determined by the thermal oscillations of the molecules and the solvent as well as the energy of binding.

The number $M$ of adsorbed molecules is given by

$$
\begin{equation*}
M=\Pi P\left(n_{U W}, n_{W}, T\right) N \tag{8.1}
\end{equation*}
$$

where $\Pi$ is the fraction of molecules that reach the (immobile) bottom layer of the fluid, $P\left(n_{U W}, n_{W}, T\right)$ is the conditional probability of a successful attachment at temperature $T$ given that the bottom

Draft: February 28, 2008, do not distribute
layer has been reached, and $N$ is the average number of protein molecules in solution in the cell. The quantities $n_{U W}$ and $n_{W}$ are the numbers of underwrapped and well-wrapped hydrogen bonds on the surface of the protein, respectively. These will be used to estimate the relative amount of protein surface area related to dehydrons. The fraction $\Pi$ depends on details of the experimental design, so we focus initially on on the second term $P$.

### 8.2.1 Boundary zone model

Suppose that $\Delta U$ is the average decrease in Coulombic energy associated with the desolvation of a dehydron upon adhesion. It is the value of $\Delta U$ that we are seeking to determine. Let $\Delta V$ be the Coulombic energy decrease upon binding at any other site. Let $f$ be the fraction of the surface covered by dehydrons. As a simplified approximation, we assume that

$$
\begin{equation*}
f \approx \frac{n_{U W}}{n_{U W}+n_{W}} \tag{8.2}
\end{equation*}
$$

Then the probability of attachment at a dehydron is predicted by thermodynamics as

$$
\begin{equation*}
P\left(n_{U W}, n_{W}, T\right)=\frac{f e^{\Delta U / k_{B} T}}{(1-f) e^{\Delta V / k_{B} T}+f e^{\Delta U / k_{B} T}} \approx \frac{n_{U W} e^{\Delta U / k_{B} T}}{n_{W} e^{\Delta V / k_{B} T}+n_{U W} e^{\Delta U / k_{B} T}} \tag{8.3}
\end{equation*}
$$

with $k_{B}=$ Boltzmann's constant. In [78], $\Delta V$ was assumed to be zero. In this case, (8.3) simplifies to

$$
\begin{equation*}
P\left(n_{U W}, n_{W}, T\right)=\frac{f e^{\Delta U / k_{B} T}}{(1-f)+f e^{\Delta U / k_{B} T}} \approx \frac{n_{U W} e^{\Delta U / k_{B} T}}{n_{W}+n_{U W} e^{\Delta U / k_{B} T}} \tag{8.4}
\end{equation*}
$$

(cf. equation (2) of [78]). Note that this probability is lower if $\Delta V>0$.

### 8.2.2 Diffusion zone model

The probability $\Pi$ in (8.1) of penetrating the bottom layer of the fluid is estimated in [78] by a model for diffusion via Brownian motion in the plane orthogonal to the flow direction. This depends on the solvent bulk viscosity $\mu$, and the molecular mass, $m$, and the hydrodynamic radius [211] or Stokes radius [109] of the protein. This radius $R$ associates with each protein an equivalent sphere that has approximately the same flow characteristics at low Reynolds numbers. This particular instance of a 'spherical cow' approximation [56, 140] is very accurate, since the variation in flow characteristics due to shape variation is quite small [211]. The drag on a sphere of radius $R$, at low Reynolds numbers, is $F=6 \pi R \mu v$ where $v$ is the velocity. The drag is a force that acts on the sphere through a viscous interaction. The coefficient

$$
\begin{equation*}
\xi=6 \pi R \mu / m=F / m v \tag{8.5}
\end{equation*}
$$

where $m$ is the molecular mass, is a temporal frequency (units: inverse time) that characterizes Brownian motion of a protein. The main non-dimensional factor that appears in the model is

$$
\begin{equation*}
\alpha=\frac{m \xi^{2} L^{2}}{2 k_{B} T}=\frac{L^{2}(6 \pi R \mu)^{2} / m}{2 k_{B} T} \tag{8.6}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
which has units of energy in numerator and denominator. We have [2]

$$
\begin{align*}
\Pi(v, R, m) & =\int_{\Lambda} \int_{\Omega \backslash \Lambda} \int_{[0, \tau]} \frac{\alpha L^{-2}}{\pi \Gamma(t)} e^{-\alpha L^{-2}\left|\mathbf{r}-\mathbf{r}_{0}\right|^{2} / \Gamma(t)} d t d \mathbf{r}_{0} d \mathbf{r} \\
& =\int_{\tilde{\Lambda}} \int_{\tilde{\Omega} \backslash \tilde{\Lambda}} \int_{[0, L / v]} \frac{\alpha}{\pi \Gamma(t)} e^{-\alpha\left|\tilde{\mathbf{r}}-\tilde{\mathbf{r}}_{0}\right|^{2} / \Gamma(t)} d t d \tilde{\mathbf{r}}_{0} d \tilde{\mathbf{r}} \tag{8.7}
\end{align*}
$$

where $\mathbf{r}$ is the two-dimensional position vector representing the cell cross-section $\Omega,|\mathbf{r}|$ denotes the Euclidean norm of $\mathbf{r}, \Lambda$ is the $6 \AA \times 10^{8} \AA$ cross-section of the bottom layer, and $\Gamma(t)=2 \xi t-$ $3+4 e^{-\xi t}-e^{-2 \xi t}$. The domains $\tilde{\Lambda}$ and $\tilde{\Omega}$ represents domains scaled by the length $L$, and thus the variables $\tilde{\mathbf{r}}$ and $\tilde{\mathbf{r}}_{0}$ are non-dimensional. In particular, the length of $\tilde{\Lambda}$ and $\tilde{\Omega}$ is one in the horizontal coordinate. Note that $\Gamma(t)=\frac{2}{3}(\xi t)^{3}+\mathcal{O}\left((\xi t)^{4}\right)$ for $\xi t$ small. Also, since the mass $m$ of a protein tends to grow with the radius cubed, $\alpha$ actually decreases like $1 / R$ as the Stokes radius increases.

### 8.2.3 Model validity

The validity of the model represented by equations (8.1 8.7) was established by data fitting. The only parameter in the model, $\Delta U$, was varied, and a value was found that consistently fits within the confidence band for the adsorption data for the six proteins (see Fig. 3 of [78]) across the entire range of flow velocities $v$. This value is

$$
\begin{equation*}
\Delta U=3.91 \pm 0.67 \mathrm{~kJ} / \mathrm{mole}=\Delta U=0.934 \pm 0.16 \mathrm{kcal} / \mathrm{mole} \tag{8.8}
\end{equation*}
$$

This value is within the range of energies associated with typical hydrogen bonds. Thus we can think of a dehydron as a hydrogen bond that gets turned 'on' by the removal of water due to the binding of a ligand.

Using the estimate (8.8) of the binding energy for a dehydron, an estimate was made [78] of the force

$$
\begin{equation*}
|F|=7.78 \pm 1.5 \mathrm{pN} \tag{8.9}
\end{equation*}
$$

exerted by the surface on a single protein molecule at a $6 \AA$ distance from the dehydron.

### 8.3 Direct force measurement

The experimental techniques reviewed in the previous section suggest that the density of dehydrons correlates with protein stickiness. However, the techniques are based on measuring the aggregate behavior of a large number of proteins. One might ask for more targeted experiments seeking to isolate the force of a dehydron, or at least a small group of dehydrons. Such experiments were reported in [68] based on atomic force microscopy (AFM).

We will not give the details of the experimental setup, but just describe the main points. The main concept was to attach hydrophobic groups to the tip of an atomic force microscope. These were then lowered onto a surface capable of forming arrays of dehydrons. This surface was formed by

| protein name | PDB code | residues | WWHB | dehydrons |
| :---: | :---: | :---: | :---: | :---: |
| apolipoprotein A-I | 1AV1 | 201 | 121 | 66 |
| $\beta$ lactoglobulin | 1BEB | 150 | 106 | 3 |
| hen egg-white lysozyme | 133 L | 130 | 34 | 13 |
| human apomyoglobin | 2 HBC | 146 | 34 | 3 |
| monomeric human insulin | 6INS | 50 | 30 | 14 |
| human $\beta_{2}$-microglobulin | 1I4F | 100 | 17 | 9 |

Table 8.1: Six proteins and their hydrogen bond distributions. WWHB=well-wrapped hydrogen bonds.
a self-assembling monolayer of the molecules $\mathrm{SH}-\left(\mathrm{CH}_{2}\right)_{11}-\mathrm{OH}$. The OH "head" groups are capable of making $\mathrm{OH}-\mathrm{OH}$ hydrogen bonds, but these will be exposed to solvent and not well protected.

The data obtained by lowering a hydrophobic probe on such a monolayer are complex to interpret. However, they become easier when they are compared with a similar monolayer not containing dehydrons. In [68], the molecule $\mathrm{SH}-\left(\mathrm{CH}_{2}\right)_{11}-\mathrm{Cl}$ was chosen.

The force-displacement curve provided by the AFM have similarities for both monolayers 68]. For large displacements, there is no force, and for very small displacements the force grows substantially as the tip is driven into the monolayer. However, in between, the characteristics are quite different.

For the OH-headed monolayer, as the displacement is decreased to the point where the hydrophobic group on the tip begins to interact with the monolayer, the force on the tip decreases, indicating a force of attraction. Near the same point of displacement, the force on the tip increases for the chlorine-headed monolayer. Thus we see the action of the dehydronic force in attracting the hydrophobes to the dehydron-rich OH-headed layer. On the other hand, there is a resistance at the similar displacement as the hydrophobic tip begins to dehydrate the chlorine-headed monolayer. Ultimately, the force of resistance reaches a maximum, and then the force actually decreases to a slightly negative (attractive) value as the monolayer becomes fully dehydrated. It is significant that the displacement for the force minimum is approximately the same for both monolayers, indicating that they both correspond to a fully dehydrated state.

The force-displacement curves when the tip is removed from the surface also provide important data on the dehydronic force. The force is negative for rather large displacements, indicating the delay due to the requirements of rehydration. Breaking the hydrophobic bond formed by the hydrophobic groups on the tip and the monolayer requires enough force to be accumulated to completely rehydrate the monlayer. This effect is similar to the force that is required to remove sticky tape, in which one must reintroduce air between the tape and the surface to which it was attached. For the chlorine-headed monolayer, there is little change in force as the displacement is increased by four $\AA$ Angstroms from the point where the force is minimal. Once the threshold is reached then the force returns abruptly to zero, over a distance of about one Ångstrom. For the OH-headed monolayer, the threshold is delayed by another two Angstroms, indicating the additional effect of the dehydronic force.

Draft: February 28, 2008, do not distribute

The estimation of the dehydronic force is complicated by the fact that one must estimate the number of dehydrons that will be dehydrated by the hydrophobic groups on the tip. But the geometry of AFM tips is well characterized, and the resulting estimate [68] of

$$
\begin{equation*}
|F|=5.9 \pm 1.2 \mathrm{pN} \tag{8.10}
\end{equation*}
$$

at a distance of $5 \AA$ is in close agreement with the estimate (8.9) of $7.78 \pm 1.5 \mathrm{pN}$ at a distance of $6 \AA$ in [78]. Part of the discrepancy could be explained by the fact that in [78] no energy of binding was attributed to the attachment to areas of a protein lacking dehydrons. If there were such an energy decrease, due e.g. to the formation of intermolecular interactions, the estimate of the force obtained in [78] would be reduced.

### 8.4 Membrane morphology

Since dehydrons have an attractive force that causes them to bind to a membrane, then the equal and opposite force must pull on the membrane. Since membranes are flexible, then this will cause the membrane to deform.

The possibility of significant morphological effect of dehydrons on membranes was suggested by the diversity of morphologies [218] of the inner membranes of cellular or subcellular compartments containing soluble proteins [80]. These vary from simple bag-like membranes 60] (e.g., erythrocytes, a.k.a. red blood cells) to highly invaginated membranes [241] (e.g., mitochondrial inner membranes). This raises the question of what might be causing the difference in membrane structure [136, 148, 175, 243.

Some evidence [80 suggests that dehydrons might play a role: hemoglobin subunits (which comprise the bulk of erythrocyte contents) are generally well wrapped, whereas two mitochondrial proteins, cytochrome c and pyruvate dehydrogenase, are less well wrapped. The correlation between the wrapping difference and the morphology difference provided motivation to measure the effect experimentally [80].

### 8.4.1 Protein adsorption

Morphology induction was tested in fluid phospholipid (DLPC) bilayers (Section 8.1) coating an optical waveguide [80. The density of bilayer invaginations was measured by a technology called evanescent field spectroscopy which allowed measurement of both the thickness and refractive index of the adlayer [203, 231]. DLPC was added as needed for membrane expansion, with the portion remaining attached to the waveguide serving as a nucleus for further bilayer formation. Stable invaginations in the lipid bilayer formed after 60 -hour incubation at $\mathrm{T}=318 \mathrm{~K}$.

### 8.4.2 Density of invaginations

The density of invaginations correlates with the extent of wrapping, $\rho$, of the soluble protein structure (Fig. 1, 2a in [80]). Greater surface area increase corresponds with lack of wrapping of backbone
hydrogen bonds. The density of invaginations as a function of concentration (Figure 2 b in [80]) shows that protein aggregation is a competing effect in the protection of solvent-exposed hydrogen bonds (77, 70, 71, 65, [84]): for each protein there appears to be a concentration limit beyond which aggregation becomes more dominant.

### 8.5 Kinetic model of morphology

The kinetics of morphology development suggest a simple morphological instability similar to the development of moguls on a steep ski run. When proteins attach to the surface, there is a force that binds the protein to the surface. This force pulls upward on the surface (and downward on the protein) and will increase the curvature in proportion to the local density of proteins adsorbed on the surface [71. The rate of change of curvature $\frac{\partial g}{\partial t}$ is an increasing function of the force $f$ :

$$
\begin{equation*}
\frac{\partial g}{\partial t}=\phi(f) \tag{8.11}
\end{equation*}
$$

for some increasing function $\phi$. Note that

$$
\begin{equation*}
\phi(0)=0 ; \tag{8.12}
\end{equation*}
$$

if there is no force, there will be no change. The function $\phi$ represents a material property of the surface.

The probability $p$ of further attachment increases as a function of the curvature at that point since there is more area for attachment where the curvature is higher. That is, $p(g)$ is also an increasing function.

Of course, attachment also reduces surface area, but we assume this effect is small initially. However, as attachment grows, this neglected term leads to a 'saturation' effect. There is a point at which further reduction of surface area becomes the dominating effect, quenching further growth in curvature. But for the moment, we want to capture the initial growth of curvature in a simple model. We leave as Exercise 8.2 the development of a more complete model.

Assuming equilibrium is attained rapidly, we can assert that the force $f$ is proportional to $p(g)$ : $f=c p(g)$ at least up to some saturation limit, which we discuss subsequently. If we wish to be conservative, we can assert only that

$$
\begin{equation*}
f=\psi(p(g)) \tag{8.13}
\end{equation*}
$$

with $\psi$ increasing. In any case, we conclude that $f$ may be regarded as an increasing function of the curvature $g$, say

$$
\begin{equation*}
f=F(g):=\phi(\psi(p(g))) . \tag{8.14}
\end{equation*}
$$

To normalize forces, we should have no force for a flat surface. That is, we should assume that $p(0)=0$. This implies, together with the condition $\phi(0)=0$, that

$$
\begin{equation*}
F(0)=0 . \tag{8.15}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

The greater attachment that occurs locally causes the force to be higher there and thus the curvature to increase even more, creating an exponential runaway (Fig. 4 in [80]). The repeated interactions of these two reinforcing effects causes the curvature to increase in an autocatalytic manner until some other process forces it to stabilize.

The description above can be captured in a semiempirical differential equation for the curvature $g$ at a fixed point on the bilayer. It takes the form

$$
\begin{equation*}
\frac{\partial g}{\partial t}=F(g) \tag{8.16}
\end{equation*}
$$

where $F$ is the function in (8.14) that quantifies the relationships between curvature, probability of attachment and local density of protein described in the previous paragraph. Abstractly, we know that $F$ is increasing because it is the composition of increasing functions. Hence $F$ has a positive slope $s$ at $g=0$. Moreover, it is plausible that $F(0)=0$ using our assumptions made previously.

Thus the curvature should grow exponentially at first with rate $s$. In the initial stages of interface development, $F$ may be linearly approximated by virtue of the mean value theorem, yielding the autocatalytic equation:

$$
\begin{equation*}
\frac{\partial g}{\partial t}=s g \tag{8.17}
\end{equation*}
$$

Figure 4 in [80] indicates that the number of invaginations appears to grow exponentially at first, and then saturates.

We have observed that there is a maximum amount of protein that can be utilized to cause morphology (Figure 2b in [80]) beyond which aggregation becomes a significantly competitive process. Thus, a 'crowding problem' at the surface causes the curvature to stop increasing once the number of adsorbed proteins gets too high at a location of high curvature.

### 8.6 Exercises

Exercise 8.1 Determine the minimal distance between a hydrophobe and a backbone hydrogen bond in protein structures. That is, determine the number of wrappers as a function of the desolvation radius, and determine when, on average, this tends to zero.

Exercise 8.2 Derive a more refined model of morphological instability accounting for the reduction of surface area upon binding. Give properties of a function $F$ as in 8.14) that incorporate the effect of decreasing surface area, and show how it would lead to a model like (8.16) which would saturate (rather than grow exponentially forever), reflecting the crowding effect of the molecules on the lipid surface.

## Chapter 9

## Electronic force details

In Section 3.3, we introduced the basic electronic interactions. Here we look at them in more detail. The basic electronic entities are groups of charges that are constrained to be together, such as dipoles. In Section 9.2 .1 we study dipole-dipole interactions. In Section 9.2.2 we consider chargedipole interactions such as arise in cation-pi pairs such as Arg-Tyr or Lys-Phe. We also consider like-charge repulsion such as occurs with Arg-His or Asp-Glu pairs in Section 9.2.3.

There is a natural hierarchy of charged groups. These can be ranked by the rate of decay of their potentials, and thus by their globality. At the highest level is the single charge, with a potential $r^{-1}$. The dipole is a combination of opposite charges at nearby locations, with a potential $r^{-2}$. The quadrupole is a collection of four charges arranged in appropriate positions with a potential $r^{-3}$. Some important entities, such as water, involve four charges at positions with substantial symmetry, and it is important to know whether they constitute quadruples or just dipoles. This determines the global accumulation of charge and thus has significant implications as we now discuss. We subsequently return to the question of whether water is a dipole or quadrupole.

### 9.1 Global accumulation of electric force

The reason that we need to know the order of decay of the potential, or the associated force, for various types of charged groups is quite simple to explain. Suppose that we have a material made of an assembly of electrostatic entities, such as water. We would like to understand the locality of forces exerted by the entities on each other. In particular, are they local, or do global contributions have a significant effect?

To quantify this question, suppose we try to estimate the force on a particular entity by all the others, and suppose this force is proportional to $r^{-n}$ for some $n$. Summing over all space, we determine the total force. We can estimate this sum by computing sums over expanding spherical shell sets $\left\{\mathbf{r} \in \mathbb{R}^{3}: R-1 \leq|\mathbf{r}|<R\right\}$ for $R=1,2,3, \ldots$. In each spherical shell region, the sum of all forces, ignoring possible cancellations, would be approximately $c R^{2-n}$ since all values of $\mathbf{r}$ in the set would be comparable to $R$, and there would be approximately $c R^{2}$ of them (assuming as we


Figure 9.1: Dipole-dipole interaction: rotated in-line configuration.
do that they are uniformly distributed). Then the total force would be proportional to

$$
\begin{equation*}
\sum_{R=1}^{R_{\mathrm{mac}}} R^{2-n} \tag{9.1}
\end{equation*}
$$

which is divergent (as $R_{\text {mac }}$ increases) for $n \leq 3$.
Note that $R_{\text {mac }}$ is the size of a macroscopic system in microscopic units, so it is related to Avogadro's number, hence should be viewed as nearly infinite. The borderline case $n=3$, for which the divergence is only logarithmic, corresponds to the electric force in the charge-dipole interaction. For the dipole-dipole interaction, $n=4$, the first exponent where the forces can be said to be local, but the convergence rate is rather slow: $\mathcal{O}\left(1 / R_{\text {cut }}\right)$ if we take $R_{\text {cut }}$ to be a cut-off radius beyond which we ignore external effects. This explains to some extent why molecular dynamics simulations have to expend so much computational effort to compute electrostatic interactions in order to represent the forces accurately.

### 9.2 Modeling interactions among polar and charged residues

We have seen that certain bonds can be modeled by simple interactions by charge groups. For example, polar groups can be modeled simply by placing partial charges appropriately at atom centers, as described in Section 7.1.2 Here we investigate in detail the angular dependence of these models.

### 9.2.1 Dipole-dipole interactions

Let us consider the effect of angular orientation on the strength of interaction of two dipoles. Since the possible set of configurations has a high dimension, we break down into special cases.

## In-line interaction configuration

Suppose we have two dipoles as indicated in Figure 9.1. The exact positions of the charges are as follows. The position of the positive charge on the right we take as the origin, and we assume the separation distance of the charges is one. The separation of the positive charge on the right and the negative charge on the left is $R$. Thus the charge centers of the dipole on the left are at


Figure 9.2: Dipole-dipole (in-line) interaction energy, scaled by $R^{3}$, for $R=2,4,10,1000$. Horizontal $\phi$-axis measured in radians. The flattest curve corresponds to $R=2$.
$(-R-1,0)$ (positive charge) and $(-R, 0)$ (negative charge). The negative charge on the right is at $(\cos \phi, \sin \phi)$.

The distances between the various charges are easy to compute. The distance between the negative charge on the left and the positive charge on the right is $R$, and the distance between the two positive charges is $R+1$. The distance between the two negative charges is

$$
\begin{align*}
|(\cos \phi, \sin \phi)-(-R, 0)| & =\sqrt{(R+\cos \phi)^{2}+\sin ^{2} \phi}  \tag{9.2}\\
& =\sqrt{1+R^{2}+2 R \cos \phi},
\end{align*}
$$

and the distance between the positive charge on the left and the negative charge on the right is

$$
\begin{align*}
|(\cos \phi, \sin \phi)-(-R-1,0)| & =\sqrt{(1+R+\cos \phi)^{2}+\sin ^{2} \phi}  \tag{9.3}\\
& =\sqrt{1+(R+1)^{2}+2(R+1) \cos \phi}
\end{align*}
$$

Thus the interaction energy for the dipole pair (assuming unit charges) is

$$
\begin{equation*}
\frac{1}{R+1}-\frac{1}{R}+\frac{1}{\sqrt{1+R^{2}+2 R \cos \phi}}-\frac{1}{\sqrt{1+(R+1)^{2}+2(R+1) \cos \phi}} \tag{9.4}
\end{equation*}
$$

A plot of the interaction energy (9.4) is given in Figure 9.2 as a function of $\phi$ for various values of $R$. Since we know (cf. (3.10)) that the interaction energy will decay like $R^{-3}$, we have scaled the energy in Figure 9.2 by $R^{3}$ to keep the plots on the same scale. The value of $R=1000$ indicates the asymptotic behavior; see Exercise 9.1 for the analytical expression. Indeed, there is little difference between $R=100$ (not shown) and $R=1000$. The flatter curve is the smallest value of $R(=2)$ and shows only limited angular dependence. Thus modeling a hydrogen bond using a simple dipole-dipole interaction does not yield a very strong angular dependence.

Draft: February 28, 2008, do not distribute


Figure 9.3: Dipole-dipole interaction: rotated parallel configuration.

## Parallel interaction configuration

Let us consider the effect of a different angular orientation on the strength of interaction of two dipoles. Suppose we have two dipoles as indicated in Figure 9.3 . Here the dipoles stay parallel, but the one on the right is displaced by an angle $\phi$ from the axis through the dipole on the left. The exact positions of the charges are as follows.

The position of the negative charge on the left we take as the origin, and we assume the separation distance of the charges is one. The separation of the positive charge on the right and the negative charge on the left is $R$. Thus the charge centers of the dipole on the right are at $R(\cos \phi, \sin \phi)$ (positive charge) and ( $1+R \cos \phi, R \sin \phi$ ) (negative charge).

The distance between the positive charges is the same as the distance between the negative charges because the dipoles are parallel:

$$
\begin{equation*}
|(1+R \cos \phi, R \sin \phi)|=\sqrt{1+R^{2}+2 R \cos \phi} \tag{9.5}
\end{equation*}
$$

Similarly, the distance between the positive charge on the left and the negative charge on the right is

$$
\begin{equation*}
|(2+R \cos \phi, R \sin \phi)|=\sqrt{4+R^{2}+4 R \cos \phi} \tag{9.6}
\end{equation*}
$$

Thus the interaction energy for the dipole pair (assuming unit charges) is

$$
\begin{equation*}
-\frac{1}{R}+\frac{2}{\sqrt{1+R^{2}+2 R \cos \phi}}-\frac{1}{\sqrt{4+R^{2}+4 R \cos \phi}} \tag{9.7}
\end{equation*}
$$

A plot of the interaction energy (9.7) is given in Figure 9.4 as a function of $\phi$ for various values of $R$. Since we know (cf. (3.10)) that the interaction energy will decay like $R^{-3}$, we have scaled the energy in Figure 9.2 by $R^{3}$ to keep the plots on the same scale. The value of $R=1000$ indicates the asymptotic behavior; see Exercise 9.2 for the analytical expression. Again, there is little difference between $R=100$ (not shown) and $R=1000$. The flatter curve is the smallest value of $R(=2)$ and shows only limited angular dependence.

## Two-parameter interaction configuration

Now we consider the effect of a dual angular orientation on the strength of interaction of two dipoles. Suppose we have two dipoles as indicated in Figure 9.5. The exact positions of the charges are as follows. The position of the negative charge on the left we take as the origin, and we assume the


Figure 9.4: Dipole-dipole (parallel) interaction energy, scaled by $R^{3}$, for $R=2,4,10,1000$. Horizontal $\phi$-axis measured in radians.
separation distance of the charges is one. The separation of the positive charge on the right and the negative charge on the left is $R$. Thus the charge centers of the dipole on the right are at $R(\cos \theta, \sin \theta)$ (positive charge) and $R(\cos \theta, \sin \theta)+(\cos \phi, \sin \phi)$ (negative charge).

The distance between the negative charge on the left and the positive charge on the right is $R$, and the separation between the positive charge on the right and the positive charge on the left is

$$
\begin{equation*}
|(1+R \cos \theta, R \sin \theta)|=\sqrt{1+R^{2}+2 R \cos \theta} \tag{9.8}
\end{equation*}
$$

The separation between the positive charge on the right and the negative charge on the left is

$$
\begin{align*}
|R(\cos \theta, \sin \theta)+(\cos \phi, \sin \phi)| & =\sqrt{(R \cos \theta+\cos \phi)^{2}+(R \sin \theta+\sin \phi)^{2}} \mid \\
& =\sqrt{\left(R^{2}+1+2 R(\cos \theta \cos \phi+\sin \theta \sin \phi)\right.} \mid \tag{9.9}
\end{align*} .
$$

Finally, the distance (squared) between the positive charge on the left and the negative charge on the right is

$$
\begin{align*}
\mid R(\cos \theta, \sin \theta) & +(\cos \phi, \sin \phi)-\left.(-1,0)\right|^{2}=|(1+R \cos \theta+\cos \phi, R \sin \theta+\sin \phi)|^{2} \\
& =(1+\cos \phi)^{2}+2 R \cos \theta(1+\cos \phi)+R^{2}+2 R \sin \theta \sin \phi+\sin ^{2} \phi  \tag{9.10}\\
& =2(1+\cos \phi)+2 R \cos \theta(1+\cos \phi)+R^{2}+2 R \sin \theta \sin \phi \\
& =2(1+\cos \phi)(1+R \cos \theta)+R^{2}+2 R \sin \theta \sin \phi
\end{align*}
$$

Thus the interaction energy for the dipole pair (assuming unit charges) is

$$
\begin{align*}
& -\frac{1}{R}+\frac{1}{\sqrt{1+R^{2}+2 R \cos \theta}} \\
& -\frac{1}{\sqrt{R^{2}+2(1+\cos \phi)(1+R \cos \theta)+2 R \sin \theta \sin \phi}}  \tag{9.11}\\
& +\frac{1}{\sqrt{1+R^{2}+2 R(\cos \theta \cos \phi+\sin \theta \sin \phi)}}
\end{align*}
$$

Draft: February 28, 2008, do not distribute


Figure 9.5: Dipole-dipole (two-angle) interaction configuration.


Figure 9.6: Dipole-dipole (two-angle) interaction energy, scaled by $R^{3}$, for $R=3$, as a function of $\phi$ for various fixed values of $\theta=0,0.2,0.4,0.6,0.8$. Approximate minimum values of the energy are indicated by circles at the points $\phi=1.7 \theta$. The horizontal $\phi$-axis is measured in radians.

## Minimum energy configuration

Since there are now two angles to vary, it is not so clear how to display the energy in a useful way. But one question we may ask is: what is the minimum energy configuration if we allow $\phi$ to vary for a given $\theta$ ? We might think that the dipole on the right would always point at the negative charge at the left. This would correspond to having the minimum energy configuration at $\phi=\theta$. This is clearly true at $\theta=0$, but say at $\theta=\pi / 2$, we might expect the minimum energy configuration to occur when the dipole on the right is flipped, that is at $\phi=\pi=2 \theta$. We plot the energy as a function of $\phi$ for various values of $\theta$ in Figure 9.6. As an aid to the eye, we plot a circle at a point close the minimum in energy, as a way to see how the optimum $\phi$ varies as a function of $\theta$. In particular, we have plotted the point not at $\phi=\theta$, nor at $\phi=2 \theta$, but rather $\phi=1.7 \theta$. This is convincing evidence that the relationship between the optimum value of $\phi$ for a fixed value of $\theta$ is complex.


Figure 9.7: Optimal $\phi$ angle (minimum interaction energy) as a function of $\theta$ for the dipole-dipole (two-angle) interaction, for $R=3,5,10,1000$ (the left-most curve corresponds to $R=3$, and they move to the right with increasing $R$ ). Horizontal $\theta$-axis and vertical $\phi$-axis are measured in radians. The line $\phi=2 \theta$ has been added as a guide.

In the case that $\theta=\pi / 2$, the expression (9.11) simplifies to

$$
\begin{equation*}
-\frac{1}{R}+\frac{1}{\sqrt{1+R^{2}}}-\frac{1}{\sqrt{R^{2}+2(1+\cos \phi)+2 R \sin \phi}}+\frac{1}{\sqrt{1+R^{2}+2 R \sin \phi}} . \tag{9.12}
\end{equation*}
$$

Then if $\phi=\pi$, this further simplifies to $-2 R^{-1}+2\left(1+R^{2}\right)^{-1 / 2}$ as we would expect. However, the minimum of the expression (9.12) does not occur at $\phi=\pi$, due to the asymmetry of the expression around this value. We leave as Exercise 9.4 to plot (9.12) as a function of $\phi$ for various values of $R$ to see the behavior.

When $R$ is large, we might expect that $\phi_{\mathrm{opt}} \approx \theta$, since the dipole should point in the general direction of the other dipole. However, this is not the case; rather there is a limiting behavior that is different. In Figure 9.7, the optimal $\phi$ is plotted as a function of $\theta$, and we note that it is very nearly equal to $2 \theta$, but not exactly. For $\theta$ small, it behaves more nearly like $\phi \approx 1.7 \theta$, but for larger values of $\theta$ the optimal $\phi$ increases to, and then exceeds, $2 \theta$, before returning to the value of $2 \theta$ near $\theta=\pi$.

The minimum $\phi$ has been determined by computing the energies for discrete values of $\phi$ and then interpolating the data by a quadratic around the discrete minimum. Necessary adjustments at the ends of the computational domain are evident. Limited resolution in the computations contributes to the visible jaggedness of the curves in the plot. We leave as an exercise to produce smoother plots, as well as to explore the asymptotic behavior as $R \rightarrow \infty$.

The energy, again scaled by $R^{3}$, at the optimal value of $\phi$ is plotted as a function of $\theta$ in Figure 9.8. Since the curves in this figure are not horizontal, the dipole system has a torque that would tend to move them to the $\theta=0$ position if $\theta$ were not fixed (as we assume it is, due to some external geometric constraint).

Draft: February 28, 2008, do not distribute


Figure 9.8: Dipole-dipole (two-angle) interaction energy minimum, scaled by $R^{3}$, for $R=$ $3,5,10,1000$ (top to bottom), as a function of $\theta$. Horizontal $\theta$-axis measured in radians. Plotted is the energy at the optimal value of $\phi$ that minimizes the energy as a function of $\phi$ for fixed $\theta$.


Figure 9.9: Charge-dipole interaction configuration.

### 9.2.2 Charge-dipole interactions

Charge-dipole interactions are simpler to analyze, and we have already anticipated their asymptotic strength in (3.7). On the other hand, this forms a very important class of interactions. Although mainchain-mainchain (hydrogen bond) interactions do not involve such pairs, all of the three other interactions among sidechains and mainchains can involve charge-dipole interactions. In addition, more complex interactions, such as cation- $\pi$ interactions (Section 13.1) are of this form. Thus we develop the basics of charge-dipole interactions in some detail.

By choosing coordinates appropriately, we can assume that the positive and negative sites of the dipole align on the $x$-axis, and that the charge is located in the $x, y$ plane, as depicted in Figure 9.9 , Assume that the negative charge of the dipole is at the origin and that the isolated charge is positive, located at $(r \cos \theta, r \sin \theta, 0)$. We choose scales such that the charges of the dipole are of unit size and the positive charge of the dipole is at $(-1,0,0)$. If $a$ is the charge of the isolated charge, then the interaction energy of the system is

$$
\begin{equation*}
V(r, \theta)=-\frac{a}{r}+\frac{a}{\sqrt{(1+r \cos \theta)^{2}+r^{2} \sin ^{2} \theta}} . \tag{9.13}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

We leave as Exercise 9.8 to show that

$$
\begin{equation*}
V(r, \theta) \approx-\frac{a \cos \theta}{r^{2}} \tag{9.14}
\end{equation*}
$$

for large $r$ and fixed $\theta$.
We will be interested in the force field that the dipole exerts on the charge as well. It is easier to compute the gradient of $V$ in Cartesian coordinates (note that we can ignore the $z$ direction in our computations):

$$
\begin{equation*}
V(x, y)=-\frac{a}{\sqrt{x^{2}+y^{2}}}+\frac{a}{\sqrt{(1+x)^{2}+y^{2}}} \tag{9.15}
\end{equation*}
$$

To improve readability, we will use the notation $[x, y]$ to denote the vector with components $x$ and $y$. Similarly, we will use $r=\sqrt{x^{2}+y^{2}}$ to reduce bookkeeping. Thus we find

$$
\begin{align*}
\nabla V(x, y) & =\frac{a[x, y]}{\left(x^{2}+y^{2}\right)^{3 / 2}}-\frac{a[1+x, y]}{\left((1+x)^{2}+y^{2}\right)^{3 / 2}} \\
& =\frac{a[x, y]\left(\left(1+2 x+r^{2}\right)^{3 / 2}-r^{3}\right)}{r^{3}\left(1+2 x+r^{2}\right)^{3 / 2}}-\frac{a[1,0]}{\left(1+2 x+r^{2}\right)^{3 / 2}}  \tag{9.16}\\
& =a \frac{[x, y]\left(\left(r^{-2}(1+2 x)+1\right)^{3 / 2}-1\right)-[1,0]}{\left(1+2 x+r^{2}\right)^{3 / 2}}
\end{align*}
$$

This expression can be used to evaluate the force field on a charge in a dipole. For example, for $r=2$ we find

$$
\begin{equation*}
\nabla V_{r=2}(\theta)=a \frac{[\cos \theta, \sin \theta]\left(\left(\frac{5}{4}+\cos \theta\right)^{3 / 2}-1\right)-\left[\frac{1}{2}, 0\right]}{4\left(\frac{5}{4}+\cos \theta\right)^{3 / 2}} \tag{9.17}
\end{equation*}
$$

We can approximate (9.16) for large $r$ as

$$
\begin{align*}
\nabla V(x, y) & \approx a \frac{[x, y]\left(r^{-2}\left(\frac{3}{2}+3 x\right)\right)-[1,0]}{\left(1+2 x+r^{2}\right)^{3 / 2}}=a \frac{[x, y]\left(\frac{3}{2}+3 x\right)-\left[r^{2}, 0\right]}{r^{2}\left(1+2 x+r^{2}\right)^{3 / 2}} \\
& =a \frac{\left[\frac{3}{2} x+2 x^{2}-y^{2}, \frac{3}{2} y+3 x y\right]}{r^{2}\left(1+2 x+r^{2}\right)^{3 / 2}} \approx a \frac{\left[2 x^{2}-y^{2}, 3 x y\right]}{r^{5}}  \tag{9.18}\\
& =a r^{-3}\left[2 \cos ^{2} \theta-\sin ^{2} \theta, 3 \cos \theta \sin \theta\right]=\frac{1}{2} a r^{-3}[1+3 \cos 2 \theta, 3 \sin 2 \theta]
\end{align*}
$$

Finally, we recall that these calculations are fully valid in three dimensions, so we have derived expressions valid for all $z$ as well. In all cases, the $z$-component of $\nabla V$ is zero.

### 9.2.3 Charge-charge interactions

We now consider the preferred angular orientation for two like charged groups as one finds in residues such as Asp and Glu. Suppose we have two charge groups as indicated in Figure 9.10. The exact positions of the charges are as follows. We assume the separation distance of the charges is two,

Draft: February 28, 2008, do not distribute

### 9.2. MODELING INTERACTIONS AMONG CHARFERND (HARCABORIICSHU日ESE DETAILS



Figure 9.10: Charge-charge interaction configuration similar to what is found in an interaction between Asp and Glu.
and we assume that the origin is the center of the two negative charges on the right. Thus there are negative charges at $(\cos \phi, \sin \phi)$ and $(-\cos \phi,-\sin \phi)$. The separation between the charge groups is $R$; the negative charges on the left are fixed at $(R \pm 1,0)$. Thus the interaction energy for the charged pairs (assuming unit charges) depend on the distances

$$
\begin{align*}
& r_{++}=|+(\cos \phi, \sin \phi)-(R+1,0)|, \\
& r_{-+}=|-(\cos \phi, \sin \phi)-(R+1,0)|, \\
& r_{+-}=|+(\cos \phi, \sin \phi)-(R-1,0)|, \text { and }  \tag{9.19}\\
& r_{--}=|-(\cos \phi, \sin \phi)-(R-1,0)| .
\end{align*}
$$

With the denotations $C=\cos \phi, S=\sin \phi$, we find

$$
\begin{align*}
r_{++}^{2} & =(C-R-1)^{2}+S^{2}=2+R^{2}-2 \cos \phi(R+1)+2 R, \\
r_{-+}^{2} & =(C+R+1)^{2}+S^{2}=2+R^{2}+2 \cos \phi(R+1)+2 R, \\
r_{+-}^{2} & =(C-R+1)^{2}+S^{2}=2+R^{2}-2 \cos \phi(R-1)-2 R, \quad \text { and }  \tag{9.20}\\
r_{--}^{2} & =(C+R-1)^{2}+S^{2}=2+R^{2}+2 \cos \phi(R-1)-2 R .
\end{align*}
$$

We can write these expressions succinctly as

$$
\begin{equation*}
r_{ \pm_{1} \pm_{2}}=\sqrt{2+R^{2} \pm_{1} 2\left(\cos \phi\left(R \pm_{2} 1\right)\right) \pm_{2} 2 R} \tag{9.21}
\end{equation*}
$$

The energy (of repulsion) for the charge groups is

$$
\begin{equation*}
\frac{1}{r_{++}}+\frac{1}{r_{-+}}+\frac{1}{r_{+-}}+\frac{1}{r_{--}} \tag{9.22}
\end{equation*}
$$

and we seek to find the value of $\phi$ that minimizes it. We leave as Exercise 9.5 to plot the expression in (9.22) which is symmetric around $\phi=\pi / 2$ and has a simple minimum there.

A more realistic model of the charge group for Asp and Glu is depicted in Figure 9.11. The configuration is now three-dimensional, with the carbon joining the oxygens below the plane. We assume a positive charge on the left at $(-R,-1,-z)$ and on the right at $(\sin \phi,-\cos \phi,-z)$. The negative charges are at $(\cos \phi, \sin \phi, 0),(-\cos \phi,-\sin \phi, 0)$, and $(R \pm 1,0,0)$. We leave it as an exercise to investigate the minimum energy configuration.

Draft: February 28, 2008, do not distribute

## CHAPTER 9. ELECTRONIC FORCE DETAOL GENERAL FORM OF A CHARGE GROUP



Figure 9.11: Charge-charge interaction configuration similar to what is found in an interaction between Asp and Glu, but with a more refined model.

### 9.3 General form of a charge group

The general form of a potential for a charged system can be written as a sum of point charge potentials

$$
\begin{equation*}
V(\mathbf{r})=\sum_{k=1}^{K} \frac{q_{k}}{\left|\mathbf{r}-\mathbf{r}_{k}\right|}, \tag{9.23}
\end{equation*}
$$

where the charges $q_{k}$ are at $\mathbf{r}_{k}$. When the net charge of the system is zero, we can interpret $V$ as being defined by a difference operator applied to the fundamental charge potential

$$
\begin{equation*}
W(\mathbf{r})=1 /|\mathbf{r}| \tag{9.24}
\end{equation*}
$$

as follows. Define a translation operator $T_{\mathbf{x}}$ by

$$
\begin{equation*}
\left(T_{\mathbf{x}} f\right)(\mathbf{r})=f(\mathbf{r}-\mathbf{x}) \tag{9.25}
\end{equation*}
$$

for any function $f$. Then we can interpret the expression (9.23) as

$$
\begin{equation*}
V=\sum_{k=1}^{K} q_{k} T_{\mathbf{r}_{k}} W \tag{9.26}
\end{equation*}
$$

In view of (9.26), we define the operator

$$
\begin{equation*}
\mathcal{D}=\sum_{k=1}^{K} q_{k} T_{\mathbf{r}_{k}} . \tag{9.27}
\end{equation*}
$$

We will see that this corresponds to a difference operator when the net charge of the system is zero.

### 9.3.1 Asymptotics of general potentials

The decay of $V(\mathbf{r})$ for simple dipoles can be determined by algebraic manipulations as in Section 3.3, However, for more complex arrangements, determining the rate is quite complicated. Multipole

Draft: February 28, 2008, do not distribute
expansions such as in Section 16.5 .1 become algebraically complex as the order increases. Here we offer an alternative calculus to determine asymptotic behavior of general potentials. We begin with some more precise notation.

Let as assume that there is a small parameter $\epsilon$ that defines the distance scale between the charge locations. That is, we define

$$
\begin{equation*}
V_{\epsilon}(\mathbf{r})=\sum_{k=1}^{K} \frac{q_{k}}{\left|\mathbf{r}-\epsilon \mathbf{r}_{k}\right|} \tag{9.28}
\end{equation*}
$$

There is a dual relationship between the asymptotics of $V_{\epsilon}(\mathbf{r})$ as $\mathbf{r} \rightarrow \infty$ and $\epsilon \rightarrow 0$, as follows:

$$
\begin{equation*}
V_{\epsilon}(\mathbf{r})=|\mathbf{r}|^{-1} V_{\epsilon /|\mathbf{r}|}\left(|\mathbf{r}|^{-1} \mathbf{r}\right) . \tag{9.29}
\end{equation*}
$$

The proof just requires changing variables in (9.28):

$$
\begin{align*}
V_{\epsilon}(\mathbf{r}) & =\frac{1}{|\mathbf{r}|} \sum_{k=1}^{K} \frac{|\mathbf{r}| q_{k}}{\left|\mathbf{r}-\epsilon \mathbf{r}_{k}\right|}=\frac{1}{|\mathbf{r}|} \sum_{k=1}^{K} \frac{q_{k}}{|\mathbf{r}|^{-1}\left|\mathbf{r}-\epsilon \mathbf{r}_{k}\right|}  \tag{9.30}\\
& =\frac{1}{|\mathbf{r}|} \sum_{k=1}^{K} \frac{q_{k}}{\left|\left(|\mathbf{r}|^{-1} \mathbf{r}\right)-(\epsilon /|\mathbf{r}|) \mathbf{r}_{k}\right|}=|\mathbf{r}|^{-1} V_{\epsilon /|\mathbf{r}|}\left(|\mathbf{r}|^{-1} \mathbf{r}\right) .
\end{align*}
$$

Given a general potential $V$ of the form (9.23), we can think of this as having $\epsilon=1$, that is, $V=V_{1}$. Using (9.30), we can derive the asymptotic form

$$
\begin{equation*}
V(\mathbf{r})=V_{1}(\mathbf{r})=\left.|\mathbf{r}|^{-1} V_{\mid \mathbf{r}}\right|^{-1}\left(|\mathbf{r}|^{-1} \mathbf{r}\right)=\epsilon V_{\epsilon}(\omega) \tag{9.31}
\end{equation*}
$$

where we now define $\epsilon=|\mathbf{r}|^{-1}$ and $\omega=|\mathbf{r}|^{-1} \mathbf{r}$ satisfies $|\omega|=1$. This says that we can determine asymptotics of $V$ as $\mathbf{r} \rightarrow \infty$ by considering instead the behavior of $V_{\epsilon}$ on bounded sets (e.g., $\omega$ with $|\omega|=1)$ as $\epsilon \rightarrow 0$.

The reason that $V_{\epsilon}$ is useful is that we can write it in terms of a difference operator applied to $W$. Recalling (9.27), we define

$$
\begin{equation*}
\mathcal{D}_{\epsilon}=\sum_{k=1}^{K} q_{k} T_{\epsilon \mathbf{r}_{k}} \tag{9.32}
\end{equation*}
$$

and observe from (9.23) and (9.24) that

$$
\begin{equation*}
V_{\epsilon}=\mathcal{D}_{\epsilon} W \tag{9.33}
\end{equation*}
$$

We will see in typical cases that, for some $k \geq 0$,

$$
\begin{equation*}
\lim _{\epsilon \rightarrow 0} \epsilon^{-k} \mathcal{D}_{\epsilon}=\mathcal{D}_{0} \tag{9.34}
\end{equation*}
$$

where $\mathcal{D}_{0}$ is a differential operator of order $k$. The convergence in (9.34) is (at least) weak convergence, in the sense that for any smooth function $f$ in a region $\Omega \subset \mathbb{R}^{3}$,

$$
\begin{equation*}
\lim _{\epsilon \rightarrow 0} \epsilon^{-k} \mathcal{D}_{\epsilon} f(\mathbf{x})=\mathcal{D}_{0} f(\mathbf{x}) \tag{9.35}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

## CHAPTER 9. ELECTRONIC FORCE DETA川Ls GENERAL FORM OF A CHARGE GROUP

uniformly for $\mathbf{x} \in \Omega$. In particular, we will be mainly interested in sets $\Omega$ that exclude the origin, where the potentials are singular. Thus we conclude that

$$
\begin{equation*}
\lim _{\epsilon \rightarrow 0} \epsilon^{-k} V_{\epsilon}=V_{0} \tag{9.36}
\end{equation*}
$$

where the limiting potential is defined by

$$
\begin{equation*}
V_{0}(\mathbf{r})=\mathcal{D}_{0} W(\mathbf{r}) . \tag{9.37}
\end{equation*}
$$

Applying (9.31), (9.36), and (9.37), we conclude that

$$
\begin{equation*}
V(\mathbf{r}) \approx \frac{1}{|\mathbf{r}|^{k+1}} \mathcal{D}_{0} W\left(|\mathbf{r}|^{-1} \mathbf{r}\right) \tag{9.38}
\end{equation*}
$$

for large $\mathbf{r}$. More precisely, we will typically show that

$$
\begin{equation*}
\epsilon^{-k} \mathcal{D}_{\epsilon} \phi(\mathbf{r})=\mathcal{D}_{0} \phi(\mathbf{r})+\mathcal{O}(\epsilon) \tag{9.39}
\end{equation*}
$$

in which case we can assert that

$$
\begin{equation*}
V(\mathbf{r})=\frac{1}{|\mathbf{r}|^{k+1}} \mathcal{D}_{0} W\left(|\mathbf{r}|^{-1} \mathbf{r}\right)+\mathcal{O}\left(|\mathbf{r}|^{-k-2}\right) \tag{9.40}
\end{equation*}
$$

### 9.3.2 Application of (9.40)

Let us show how (9.40) can be used in practice by considering a known situation, that of a dipole. Thus take $\mathbf{r}_{1}=\left(\frac{1}{2}, 0,0\right)$ and $\mathbf{r}_{2}=\left(-\frac{1}{2}, 0,0\right)$. We can compute the action of $\mathcal{D} \epsilon$ on smooth functions via

$$
\begin{equation*}
\mathcal{D}_{\epsilon} \phi(x, y, z)=\phi\left(x+\frac{1}{2} \epsilon, y, z\right)-\phi\left(x-\frac{1}{2} \epsilon, y, z\right) . \tag{9.41}
\end{equation*}
$$

By Taylor's theorem, we can expand a function $\psi$ to show that

$$
\begin{equation*}
\psi(x+\xi)-\psi(x-\xi)=2 \xi \psi^{\prime}(x)+\frac{1}{3} \xi^{3} \psi^{(3)}(x)+\mathcal{O}\left(\xi^{5}\right) \tag{9.42}
\end{equation*}
$$

Applying (9.42) to $\psi(x)=\phi(x, y, z)$, we have

$$
\begin{equation*}
\mathcal{D}_{\epsilon} \phi(x, y, z)=\epsilon \frac{\partial}{\partial x} \phi(x, y, z)+\mathcal{O}\left(\epsilon^{3}\right) \tag{9.43}
\end{equation*}
$$

Taking limits, we see that

$$
\begin{equation*}
\epsilon^{-1} \mathcal{D}_{\epsilon} \rightarrow \frac{\partial}{\partial x} \tag{9.44}
\end{equation*}
$$

as $\epsilon \rightarrow 0$. Thus we conclude that the potential for a dipole is $\mathcal{O}\left(|\mathbf{r}|^{-2}\right)$ for large $\mathbf{r}$, in keeping with the derivation in Section 3.3. More precisely, applying (9.40) we have

$$
\begin{equation*}
V(\mathbf{r})=|\mathbf{r}|^{-2} \frac{\partial}{\partial x} W\left(|\mathbf{r}|^{-1} \mathbf{r}\right)+\mathcal{O}\left(|\mathbf{r}|^{-3}\right) \tag{9.45}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

| charge group | nonzero net charge | dipole | quadrapole | octapole |
| :--- | :---: | :---: | :---: | :---: |
| decay rate | $r^{-1}$ | $r^{-2}$ | $r^{-3}$ | $r^{-4}$ |

Table 9.1: Asymptotic decay rates for potentials of various charge groups.

| charge group | nonzero | dipole | quadrapole | octapole |
| :--- | :---: | :---: | :---: | :---: |
| nonzero | $r^{-1}$ | $r^{-2}$ | $r^{-3}$ | $r^{-4}$ |
| dipole | $r^{-2}$ | $r^{-3}$ | $r^{-4}$ | $r^{-5}$ |
| quadrapole | $r^{-3}$ | $r^{-4}$ | $r^{-5}$ | $r^{-6}$ |
| octapole | $r^{-4}$ | $r^{-5}$ | $r^{-6}$ | $r^{-7}$ |

Table 9.2: Asymptotic decay rates for interaction energies of various charge groups.

### 9.3.3 Multipole summary

Let us summarize the asymptotic behavior of the various potentials that can arise. We have seen in (9.40) that the order of decay of the potential can be determined by the arrangement of the charges (9.32). When the net charge is non-zero, we have $k=0$, but when the net charge is zero then $k \geq 1$. The dipole is the case $k=1$, and $k=2$ is called the quadrapole. Similarly, $k=3$ is an octapole, and so on. We summarize in Table 9.1 the different powers for the potential of these different charge groups.

The interaction energy between different charge groups has been worked out in specific cases. We summarize the general case in Table 9.2. We leave as Exercise 9.11 to verify the additional cases not already covered.

### 9.4 Quadrupole potential

The most important potential after the dipole is the quadrupole. As the name implies, it typically involves four charges. For this reason, the geometry can be quite complex. This provides an opportunity to apply the techniques developed in Section 9.3 . We begin with a simple case.

### 9.4.1 Opposing dipoles

Two opposing dipoles tend to cancel each other out, but the result is not zero, rather it is a quadrupole. For example, suppose there unit negative charges at $( \pm a, 0,0)$, where $a$ is some (positive) distance parameter, with unit positive charges at $(a+1,0,0)$ and ( $-a-1,0,0$ ). These four charges can be arranged as two dipoles, one centered at $a+\frac{1}{2}$ and the other centered at $-a-\frac{1}{2}$. Thus the separation distance $S$ between the two dipoles is

$$
\begin{equation*}
S=2 a+1 \tag{9.46}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute


Figure 9.12: Comparison of single charge, dipole and quadrupole potentials. Dipole separation (a) two units and (b) four units. The locations of the negative charges are indicated by circles and the locations of the positive charges are indicated by squares. The upper solid line is the potential for a single positive charge indicated by the right-most square. The middle, short-dashed line is the potential for the dipole corresponding to the right-most dipole. The lower, longer-dashed line is the potential for the dipole corresponding to the quadrupole formed by the pair of dipoles.

The partial charges for a benzene ring as modeled in Table 13.1 consist of three sets of such paired dipoles, arranged in a hexagonal fashion.

The potential for such a charge group can be estimated by algebraic means, as we did in Chapter 3, or we can utilize the technology of Section 9.3. We define

$$
\begin{equation*}
\mathcal{D}_{\epsilon}=T_{\epsilon(a+1,0,0)}-T_{\epsilon(a, 0,0)}+T_{\epsilon(-a-1,0,0)}-T_{\epsilon(-a, 0,0)} \tag{9.47}
\end{equation*}
$$

In evaluating $\mathcal{D}_{\epsilon} \phi$, we may as well assume that $\phi$ is only a function of $x$. Applying (9.42) to $\phi$ and $\phi^{\prime}$ we find that

$$
\begin{align*}
\mathcal{D}_{\epsilon} \phi(x) & =\phi(x-\epsilon(a+1))-\phi(x-\epsilon a)+\phi(x+\epsilon(a+1))-\phi(x+\epsilon a) \\
& =\epsilon \phi^{\prime}\left(x-\epsilon\left(a+\frac{1}{2}\right)\right)-\epsilon \phi^{\prime}\left(x+\epsilon\left(a+\frac{1}{2}\right)\right)+\mathcal{O}\left(\epsilon^{3}\right) \\
& =\epsilon^{2}(2 a+1) \phi^{\prime \prime}(x)+\mathcal{O}\left(\epsilon^{3}\right)  \tag{9.48}\\
& =\epsilon^{2} S \phi^{\prime \prime}(x)+\mathcal{O}\left(\epsilon^{3}\right),
\end{align*}
$$

where $S$ is the separation distance between the dipoles. Thus

$$
\begin{equation*}
\lim _{\epsilon \rightarrow 0} \epsilon^{-2} \mathcal{D}_{\epsilon}=(2 a+1) \frac{\partial^{2}}{\partial x^{2}}=S \frac{\partial^{2}}{\partial x^{2}} \tag{9.49}
\end{equation*}
$$

where $S=2 a+1$ is the separation distance (9.46) between the dipoles. Applying (9.40), we find

$$
\begin{equation*}
V(\mathbf{r})=|\mathbf{r}|^{-3} S \frac{\partial^{2}}{\partial x^{2}} W\left(|\mathbf{r}|^{-1} \mathbf{r}\right)+\mathcal{O}\left(|\mathbf{r}|^{-4}\right) \tag{9.50}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute


Figure 9.13: Four corner quadrupole potential. (a) The potential is plotted as a function of distance $s$ along the line $(x(x), y(s))=((2+s / \sqrt{2}, 1-s / \sqrt{2})$ which emanates from the lower-right corner of the quadrupole. The locations of the negative charges are indicated by circles and the locations of the positive charges are indicated by squares. (b) Schematic representation. The line used for the plot in (a) is indicated as a dashed line. The potential vanishes, by symmetry, on the dotted lines.
for large $\mathbf{r}$, where $W(=1 / r)$ is defined in (9.24).
The potential for opposing dipoles is depicted in Figure 9.12 for two separation distances, $S=2$ (a) and $S=4(\mathrm{~b})$. For the larger value of the separation, there is little difference between the dipole and quadrupole potentials near the right-most charge. There is a much greater difference between the potentials for a single charge and that of a dipole. Thus the separation distance affects substantially the cancellation of the second dipole, at least locally. If the distance units are interpreted as Angstroms, then the separation $S=4(\mathrm{~b})$ is roughly comparable to the partial charge model of a benzene ring (cf. Table 13.1) consisting of three sets of such paired dipoles.

### 9.4.2 Four-corner quadrupole

The four-corner arrangement provides a two-dimensional arrangement of opposing dipoles. This quadrupole system has positive charges $q_{1}=q_{2}=1$ at $\mathbf{r}_{1}=(-1,1,0)$ and $\mathbf{r}_{2}=(1,-1,0)$ and negative charges $q_{3}=q_{4}=-1$ at $\mathbf{r}_{3}=(1,1,0)$ and $\mathbf{r}_{4}=(-1,-1,0)$. A plot of the potential along a diagonal where it is maximal is given in Figure 9.13 . Note that it dies off a bit more rapidly than the potential for the opposing dipoles (cf. Figure 9.12). Defining

$$
\begin{equation*}
\mathcal{D}_{\epsilon}=\sum_{k=1}^{K} q_{k} T_{\epsilon r_{k}} \tag{9.51}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute


Figure 9.14: A near quadrupole found in the PDB file 1 I 4 M of the human prion. (a) The four charged groups are nearly aligned on the right side of the figure. Shown is the residue sequence DRYYRE. (b) Detail of the charged groups indicating the alignment of the opposing dipoles.
and applying (9.42) twice, we see that

$$
\begin{align*}
\mathcal{D}_{\epsilon} \phi(\mathbf{r}) & =\sum_{k=1}^{K} q_{k} \phi\left(\mathbf{r}-\epsilon \mathbf{r}_{k}\right)  \tag{9.52}\\
& =4 \frac{\partial}{\partial x} \frac{\partial}{\partial y} \phi(0) \epsilon^{2}+\mathcal{O}\left(\epsilon^{3}\right)
\end{align*}
$$

Thus

$$
\begin{equation*}
V(\mathbf{r})=|\mathbf{r}|^{-3} 4 \frac{\partial}{\partial x} \frac{\partial}{\partial y} W\left(|\mathbf{r}|^{-1} \mathbf{r}\right)+\mathcal{O}\left(|\mathbf{r}|^{-4}\right) \tag{9.53}
\end{equation*}
$$

It is not hard to generalize these results to the case where the opposing charges form the four corners of any parallelogram.

### 9.4.3 Quadrupole example

An example of a (near) quadrupole is found in the human prion (PDB file 1 I 4 M ) in the motif DRYYRE. This is shown in Figure 9.14 The charges closely approximate the 'four corner' arrangement for a suitable parallelogram. The DRYYRE residue group forms a helical structure. Note that the four charged sidechains are nearly planar, with the tyrosines transverse to this plane. The detail Figure 9.14(b) shows the skewness of the two opposing dipoles.


Figure 9.15: Configuration of charges in water model. Open circles indicate negative charge locations; shaded circles indicate locations of positive charge.

### 9.4.4 Water: dipole or quadrupole?

Water can be written as a combination of two dipoles, following the general pattern of Section 9.3 , So is water a quadrupole or just a dipole? The answer is crucial to determine the locality or globality of water-water interaction.

We can write water as system with positive charges

$$
\begin{equation*}
q_{1}=q_{2}=a \text { at } \mathbf{r}_{1,2}=( \pm c,-1,0) \tag{9.54}
\end{equation*}
$$

and negative charges

$$
\begin{equation*}
q_{3}=q_{4}=-b \text { at } \mathbf{r}_{3.4}=\left(0, y^{0}, \pm d\right), \tag{9.55}
\end{equation*}
$$

where $y^{0}>0$ denotes the position above the $x$-axis of the lone-pair oxygen charges. Note that we have chosen the spatial unit so that the hydrogens are exactly one unit below the $x$-axis (and the charge center is the origin), but otherwise all positions are arbitrary. This is exactly the model of water that is used by Tip5P [160], with $a=b$. We would like to show that this system is a dipole; by that, we mean two things, one of which is that it is not a quadrupole.

To discover the exact multipole nature of our water model encoded in (9.54) and (9.55), we modify it to form a quadrupole. We extend the system (9.54 9.55) to involve two more charges:

$$
\begin{align*}
q_{5}=-2 a & \text { at } \mathbf{r}_{5}
\end{align*}=(0,-1,0) \text { and }, ~=2 b \text { at } \mathbf{r}_{6}=\left(0, y_{0}, 0\right) .
$$

The configuration of charges is depicted in Figure 9.15

Draft: February 28, 2008, do not distribute

The extended system is a quadrupole due to the cancellations leading to an expression such as (9.52). More precisely, note that the charges at locations 1, 2 and 5 correspond to a second difference stencil centered at point 5 for approximating

$$
\begin{equation*}
\frac{\partial^{2} \phi}{\partial x^{2}}(0,-1,0) \tag{9.57}
\end{equation*}
$$

(with suitable scaling). Similarly, the charges at locations 3, 4 and 6 correspond to a second difference stencil centered at point 6 for approximating

$$
\begin{equation*}
\frac{\partial^{2} \phi}{\partial z^{2}}\left(0, y^{0}, 0\right) \tag{9.58}
\end{equation*}
$$

(with suitable scaling). Therefore

$$
\begin{align*}
\mathcal{D}_{\epsilon} \phi(0) & =\sum_{k=1}^{6} q_{k} \phi\left(\epsilon \mathbf{r}_{k}\right)  \tag{9.59}\\
& =a c^{2} \epsilon^{2} \frac{\partial^{2} \phi}{\partial x^{2}}(0,-1,0)-b d^{2} \epsilon^{2} \frac{\partial^{2} \phi}{\partial z^{2}}\left(0, y^{0}, 0\right) \epsilon^{2}+\mathcal{O}\left(\epsilon^{4}\right),
\end{align*}
$$

and a similar result would hold when expanding about any point $\mathbf{r}$.
Let $V^{D}$ denote the potential of the system with charges as indicated in (9.56). We leave as Exercise 9.10 to show that this is a dipole provided $a=b$. Let $V_{Q}$ denote the quadrupole potential associated with (9.59), and let $V^{W}$ be the water potential using the model (9.54-9.55). Thus we have written the water potential as

$$
\begin{equation*}
V^{W}=V^{D}+V^{Q} \tag{9.60}
\end{equation*}
$$

for an explicit dipole potential $V^{D}$, with charges at $\mathbf{r}_{5}$ and $\mathbf{r}_{6}$, and a quadrupole. Thus the water model 9.540 .55$)$ is asymptotically a dipole, and not a quadrupole. Moreover, we see that the axis of the dipole is the $y$-axis, the bisector of the angle $\angle H O H$.

### 9.5 Further results

We collect here some further results about electrostatic interactions.

### 9.5.1 Dipole induction by dipoles

Water has both a fixed dipole and an inducible dipole. That is, water is both polar and polarizable. The dipole strength of water in the gas phase $\mu \approx 0.5 \mathrm{e}-\AA$ (cf. Section 10.7.2), and the polarizability $\alpha \approx 1.2 \AA^{3}$. Thus an electric field strength of only one tenth of an electron per square Ångstrom $\left(0.1 \mathrm{e}-\AA^{-2}\right)$ could make a substantial modification to the polarity of water, since the change in polarity is approximated by the product of the polarizability and the electric field strength (see (??)).

Draft: February 28, 2008, do not distribute


Figure 9.16: Dipole-dipole (in-line) interaction energy, scaled by $R^{3}$, for $R=2$ for the two models. Horizontal $\phi$-axis measured in radians. The flattest (dashed) curve corresponds to $\alpha=0.8, \beta=$ $1.0, \gamma=0.8, \delta=0.2$, whereas the solid curve corresponds to $\alpha=0.8, \beta=1.0, \gamma=0.8, \delta=0.2$,

### 9.5.2 Modified dipole interaction

Since we found that the dipole-dipole interaction does not reproduce the sort of angular dependence we expect for certain bonds, e.g., hydrogen bonds, it is reasonable to try to modify the model. We ask the the question: if the hydrogen charge density is represented in a more complex way, will a stronger angular dependence appear? To address this question, we introduce a negative charge to represent the electron density beyond the hydrogen. The exact positions of the charges are as follows. The position of the negative charge on the right we take as the origin, and we assume the separation distance of the charges is one. The separation of the positive charge on the left and the negative charge on the right is $R$. Thus the charge centers of the multipole on the left are at $(-R-1,0)$ (negative charge $-\alpha),(-R, 0)$ (positive charge $+\beta$ ) and $(-R+\delta, 0)$ (negative charge $-\gamma)$. The positive charge on the right is at $(\cos \phi, \sin \phi)$.

The distances between the various charges are easy to compute. The distance between the positive charge on the left and the negative charge on the right is $R$, and the distance between the main $(\alpha)$ negative charge on the left and the negative charge on the right is $R+1$. The distance between the minor $(\gamma)$ negative charge on the left and the negative charge on the right is $R-\delta$.

The distance between the positive charge on the right and the minor $(\gamma)$ negative charge on the left is

$$
\begin{align*}
|(\cos \phi, \sin \phi)-(-R+\delta, 0)| & =\sqrt{(R-\delta+\cos \phi)^{2}+\sin ^{2} \phi}  \tag{9.61}\\
& =\sqrt{1+(R-\delta)^{2}+2(R-\delta) \cos \phi}
\end{align*}
$$

The distance between the two positive charges is

$$
\begin{align*}
|(\cos \phi, \sin \phi)-(-R, 0)| & =\sqrt{(R+\cos \phi)^{2}+\sin ^{2} \phi}  \tag{9.62}\\
& =\sqrt{1+R^{2}+2 R \cos \phi}
\end{align*}
$$

Draft: February 28, 2008, do not distribute
and the distance between the main $(\alpha)$ negative charge on the left and the positive charge on the right is

$$
\begin{align*}
|(\cos \phi, \sin \phi)-(-R-1,0)| & =\sqrt{(1+R+\cos \phi)^{2}+\sin ^{2} \phi}  \tag{9.63}\\
& =\sqrt{1+(R+1)^{2}+2(R+1) \cos \phi}
\end{align*}
$$

Thus the interaction energy for the dipole pair (assuming unit charges) is

$$
\begin{align*}
& \frac{\alpha}{R+1}-\frac{\beta}{R}+\frac{\gamma}{R-\delta}-\frac{\gamma}{\sqrt{1+(R-\delta)^{2}+2(R-\delta) \cos \phi}} \\
+ & \frac{\beta}{\sqrt{1+R^{2}+2 R \cos \phi}}-\frac{\alpha}{\sqrt{1+(R+1)^{2}+2(R+1) \cos \phi}} \tag{9.64}
\end{align*}
$$

A plot of the interaction energy (9.64) is given in Figure 9.16 as a function of $\phi$ for $R=2$, scaled by $R^{-3}=8$. The flatter curve corresponds to the new model with a more complex dipole. Thus we see that this does not produce an improved model of the angular dependence of a hydrogen bond.

### 9.5.3 Hydrogen placement for Ser and Thr

Let us consider the problem of determining the angular orientation of the hydrogen in serine and threonine, depicted in Figure 5.4. We choose coordinates so that the $x, y$ plane contains the terminal carbon and oxygen from the sidechain of Ser/Thr and the negative site of the partial charge of the moitie that is forming the hydrogen bond, as depicted in Figure 9.17. In the special case that the positive charge in the dipole forming the hydrogen acceptor is also in this plane, then we can argue by symmetry that the hydrogen must lie in this plane as well, at one of the solid dots indicated at the intersection of the circle with the plane of the page.

But in general, we must assume that the location of the positive partial charge is outside of this plane.

In Figure 9.17 (b), we indicate the view from the plane defined by the positions of the oxygen and the negative and positive partial charges of the dipole. The circle of possible locations for the hydrogen (see Figure 5.4) is now clearly visible, and the intersection points with the plane of the page are again indicated by black dots. Now we see it is not obvious what the optimal position for the hydrogen would be.

To determine the optimal hydrogen position, let us assume that the coordinates are as in Figure 9.17 , with the origin chosen to be at the center of the circle. Thus, the plane of the page is the $x, y$ plane, and the coordinates of the circle are $(0, \cos \theta, \sin \theta)$. The position of the negative partial charge is then $\left(x_{0}, y_{0}, 0\right)$ and the positive partial charge is $\left(x_{1}, y_{1}, z_{1}\right)$. The interaction potential between the dipole and the hydrogen is thus

$$
\begin{equation*}
\frac{-1}{\sqrt{x_{0}^{2}+\left(y_{0}-\cos \theta\right)^{2}+\sin ^{2} \theta}}+\frac{1}{\sqrt{x_{1}^{2}+\left(y_{1}-\cos \theta\right)^{2}+\left(z_{1}-\sin \theta\right)^{2}}} \tag{9.65}
\end{equation*}
$$

For given $x_{0}, y_{0}, x_{1}, y_{1}, z_{1}$, this expression can be minimized to find the optimal $\theta$.

Draft: February 28, 2008, do not distribute
(a)



(b)


Figure 9.17: Configuration for the placement of hydrogen at the end of the sidechain of serine or threonine in response to a nearby dipole. The dashed line indicates the circle of possible hydrogen placements. (a) The plane of the circle is orthogonal to the plane of the page. (b) The plane of the circle is skew to the plane of the page.

We can also use the expression (9.16) to find the optimum $\theta$. In coordinates determined so that the hydrogen and the dipole lie in a plane, the interaction field (9.16) has a zero component orthogonal to the plane. For the hydrogen position on the circle to be correct, the tangent to the circle must be orthogonal to the gradient of the interaction potential at that point. Suppose that we write the circle as $(x(\phi), y(\phi), z(\phi))$ in these coordinates. Then a necessary condition is that

$$
\begin{equation*}
\nabla V(x(\phi), y(\phi), z(\phi)) \cdot\left(x^{\prime}(\phi), y^{\prime}(\phi), z^{\prime}(\phi)\right)=0 . \tag{9.66}
\end{equation*}
$$

### 9.6 Exercises

Exercise 9.1 Show that the interaction energy (9.4) tends to the asymptotic form

$$
\begin{equation*}
\frac{-2 \cos \phi}{R^{3}} \tag{9.67}
\end{equation*}
$$

Exercise 9.2 Show that the interaction energy 9.7) tends to the asymptotic form

$$
\begin{equation*}
\frac{-\frac{1}{2}-\frac{3}{2} \cos \phi}{R^{3}} \tag{9.68}
\end{equation*}
$$

Exercise 9.3 Verify that the second term in the energy expression in (9.11) is indeed the same as (9.5). Also verify that the fourth term in the energy expression in (9.11) is correct.

Exercise 9.4 Plot the expression in (9.12) and verify that it is not symmetric around $\phi=\pi$ for finite $R$. Determine the limiting expression as $R \rightarrow \infty$ after scaling by $R^{3}$.

Exercise 9.5 Plot the expression in (9.22) and verify that it is symmetric around $\phi=\pi / 2$ and has a simple minimum there.

Draft: February 28, 2008, do not distribute

Exercise 9.6 Carry out the calculations leading to the expression in 9.22) in the case that the charge group has a positive charge as well as the negative charges, as shown in Figure 0.11. Take the charges to be appropriate for Asp or Glu. Investigate the minimum energy configuration. Also consider three-dimensional configurations in which the positive charge is located below the negative charges.

Exercise 9.7 Investigate the optimal (minimum energy) configuration for charge-dipole pairs in which the charge is fixed at a distance $r$ from the center of the dipole, which is free to rotate by and angle $\phi$. Determine the value of $\phi$ at the minimum.

Exercise 9.8 Prove that the asymptotic expression 9.14) is valid for fixed $\theta$ and large $r$. (Hint: show that

$$
\begin{equation*}
V(r, \theta)=\frac{a}{r}\left(\frac{1-\sqrt{1+2 r^{-1} \cos \theta+r^{-2}}}{\sqrt{1+2 r^{-1} \cos \theta+r^{-2}}}\right) \tag{9.69}
\end{equation*}
$$

and expand the expression in the numerator. Is this asymptotic approximation uniformly valid for all $\theta$ ?)

Exercise 9.9 Determine the percentage error in the approximation (9.14) when $\theta=\pi / 4$ and $r=3$.
Exercise 9.10 Show that a charge system with only the charges as indicated in (9.56) forms a dipole provided $a=b$ and examine its asymptotic behavior.

Exercise 9.11 Verify the interaction energies listed in Table 9.2 for the cases not already covered. (Hint: develop technology similar to Section 9.3.1. The interaction introduces an additional difference operator that multiplies the one associated with the potential. The order of the product of the two limiting differential operators is equal to the sum of the orders of the individual operators.)

## Chapter 10

## Units

It is helpful to pick the right set of units in order to reason easily about a physical subject. In different contexts, different units are appropriate. Small boat enthusiasts will recognize the need to determine whether depth on a chart is labeled in feet or fathoms. It is common in the United States coastal waters to label the depths in feet where mostly small boats will be expected to be found. But commercial vessels might prefer to think in fathoms (a fathom is six feet) since their depth requirements will be some number of fathoms (and thus a much larger number of feet). The phrase "mark twain" was used by riverboats for whom twelve feet of water provided safe passage.

In astronomy, we may measure distances in light-years. But this is the wrong unit for our discussion. Just like the choice between fathoms for commercial vessels and feet for small pleasure boats, we need to find the right size for our mental models.

### 10.1 Basic units vs. derived units

We encounter units for many things: length, time, mass, charge, viscosity, energy, and so forth. There are only so many of these that are independent. Once we choose a set of units, others must be derived from them. In Table 10.1, we give a simple example of a particular choice of basic and derived units.

As a simple example, energy $(E)$ is measured in units of mass $(m)$ times velocity $(v)$ squared, and velocity has units length $(\ell)$ over time $(t)$ :

$$
\begin{equation*}
E=m v^{2}=m(\ell / t)^{2} \tag{10.1}
\end{equation*}
$$

However, there is no canonical definition of which units are basic and which are derived.

| basic units | length, time, mass, charge, temperature |
| :--- | :--- |
| derived units | energy, viscosity, kinematic viscosity, permittivity, speed of light |

Table 10.1: An example of basic versus derived units.

| basic units | meter, second, kilogram, .... |
| :--- | :--- |
| derived units | joule (energy), newton (force), .... |

Table 10.2: The SI system of basic and derived units.

We might want certain units to have a prescribed value, and thus we take those units to be basic. For example, we might want the permittivity of free space to be one, as we discuss in Section 10.2.3,

### 10.1.1 SI units

One standard set of units is the SI system. Mass is measured in (kilo)grams, distance in meters, time in seconds. There are other basic units as well, but let us stop here as it allows us to define the standard units of energy and force.

The SI standard unit of energy is the joule. Energy has units mass times velocity squared, as we know from Einstein's famous relation. A joule is a newton-meter, the work related to applying the force of a newton for a distance of a meter. A newton is one kilogram-meter/second ${ }^{2}$, so a joule as one kilogram-(meter/second) ${ }^{2}$.

These quantities are familiar macroscopic measures. A kilogram is the weight of a good book, and a meter per second is understandable as a walking speed: 3.6 kilometers per hour. Thus a joule is the energy required to get a book up to walking speed. In many cases, the older unit calorie is used, which differs from the joule by a small factor: one calorie is 4.1868 joules.

### 10.2 Biochemical units

There are natural units associated with biochemical phenomena which relate more to the nanoscale. For example, the frequently used unit for energy is kcal/mole. This of course refers to one-thousand calories per mole of particles, or per $6.022 \times 10^{23}$ particles, which is Avogadro's number. That is a big number, but we can squash it down with the right word: it is 0.6022 yotta-particles (yotta is a prefix which means $10^{24}$, just as kilo means $10^{3}$ or nano means $10^{-9}$ ). The kcal is 4.1868 kilojoules, or 3.9683 Btu (British thermal units, a unit used in describing the power of both residential and commercial heating and cooling systems).

### 10.2.1 Molecular length and mass units

At the molecular scale, the typical units of mass (e.g., the gram) are much too large to be meaningful. Moreover, the units such as the meter and gram are based on macro-scale quantities. It would be much more reasonable to pick scales more appropriate for the atomic scale [233] such as the Bohr radius $a_{0}=0.529189 \AA$, which is based on properties of the electron distribution for the hydrogen atom. Similarly, a more natural mass unit would be based on an atomic mass, e.g., the dalton (or $\mathbf{D a}$ ) which is essentially the mass of the hydrogen atom. More precisely, it is one-twelfth of

Draft: February 28, 2008, do not distribute

| basic units | Ångstrom, svedberg $\left(10^{-13} \mathrm{~s}\right)$, dalton (mass) |
| :--- | :--- |
| derived units | $($ energy $)$ dalton- $(\AA / \text { svedberg })^{2}=0.239 \mathrm{kcal} / \mathrm{mole}$ |

Table 10.3: Some units relevant for biochemistry.
the mass of carbon twelve. The dalton is almost identical to the previous standard known as the atomic mass unit (or $\mathbf{a m u}$ ) The dalton mass unit $=1.6605310^{-27}$ kilograms.

### 10.2.2 Molecular time units

A natural time scale for biochemistry is the femtosecond ( $10^{-15}$ second) range. This is the temporal scale to observe the dynamics of molecules above the quantum level. For example, time-stepping schemes for molecular dynamics simulation are often a few femtoseconds, although some systems (e.g., liquid argon) appear to be stable for timesteps up to 100 femtoseconds. The svedberg is a time unit equal to 100 femtoseconds ( $10^{-13}$ second).

This svedberg provides a time scale that resolves molecular motion, but does not over resolve it: it is a scale at which to see details evolving the way a mechanical system would evolve in our everyday experience. We perceive things happening in a fraction of a second and are aware of motions that take place over many seconds. Runners and other athletes are timed to hundredths of a second, so we can think of that as a timestep for our perception. Thus our typical perception of motion covers $10^{4}$ or $10^{5}$ of our perceptual timesteps. By this reckoning, there are about $2 \times 10^{11}$ timesteps in a typical human lifetime. Biological events, such as protein folding, take up to $10^{11}$ svedbergs, and even more. Note that a typical human height is about $2 \times 10^{10}$ Ångstroms.

There is a natural length scale associated with any temporal scale when electromagnetic waves will be of interest. Just like the light-year, it is natural to consider the distance light travels in the natural time unit here, the svedberg, about $2.9979 \times 10^{-5}$ meters, or 30 micometers. This may seem odd. You might have expected a spatial unit on the order of an atomic unit such as the Angstrom, but this is 0.03 millimeters, a scale we can almost resolve with a magnifying glass. This means that light is still very fast at these molecular scales. We hesitate to give this length a name, but it is clearly a light-svedberg.

If we pick the svedberg as time unit and the Angstrom as spatial unit, then the natural velocity scale is the $\AA$ ngstrom per svedberg, which is equal to $10^{-10+13}=1000$ meters per second, about three times the speed of sound in air at sea level. With the dalton as mass unit, the natural energy unit in these units is one dalton- $(\text { Ångstrom per svedberg })^{2}$. One dalton- $(\AA / \text { svedberg })^{2}=$ $1.66 \times 10^{-21}$ joules $=0.239 \mathrm{kcal} / \mathrm{mole}$. Thus the chosen units of mass, length and time lead to a nearly unit value for the commonly used unit of energy, kcal/mole.

### 10.2.3 Charge units

The natural unit of charge for protein chemistry is the charge of the electron, $q_{e}$. When we look at macromolecules, we can resolve individual units and their charges. The coulomb is an aggregate

| basic units | kcal/mole (energy), $q_{e}$ (charge), $\varepsilon$ (permittivity) |
| :--- | :--- |
| derived units | (length) 417 nanometers |

Table 10.4: The units implied by Debye's assumptions.
charge constant defined so that $q_{e}=1.602 \times 10^{-19} \mathrm{C}$. That is, $\mathrm{C}=6.242 \times 10^{18} q_{e}$. The actual definition of a coulomb is the charge associated with an ampere flowing for a second. Thus a hundred amp-hour battery has 360,000 coulombs of charge, or about $2.25 \times 10^{24} q_{e}$, which corresponds to 3.7 moles of electrons.

Permittivity has units charge-squared per energy-length:

$$
\begin{equation*}
\text { permittivity }=\frac{\text { charge }^{2}}{\text { energy-length }}=\frac{\text { charge }^{2} \text { time }^{2}}{\text { mass-length }^{3}} . \tag{10.2}
\end{equation*}
$$

Thus it is possible to have the permittivity, charge and energy be one in any units by varying the spatial (length) unit:

$$
\begin{equation*}
\text { length }=\frac{\text { charge }^{2}}{\text { energy-permittivity }} \tag{10.3}
\end{equation*}
$$

However, it would not be possible to specify length, permittivity, energy, and charge independently.
The permittivity of free space $\varepsilon_{0}$ is $8.8542 \times 10^{-12} \mathrm{~F} \mathrm{~m}^{-1}$ (farads per meter). A farad is a coulomb squared per newton-meter. That is, we also have

$$
\begin{align*}
\varepsilon_{0} & =8.8542 \times 10^{-12} \mathrm{C}^{2} \mathrm{~N}^{-1} \mathrm{~m}^{-2} \\
& =3.450 \times 10^{26} q_{e}^{2} \mathrm{~N}^{-1} \mathrm{~m}^{-2}=3.450 \times 10^{26} q_{e}^{2} \mathrm{~J}^{-1} \mathrm{~m}^{-1} \\
& =1.444 \times 10^{27} q_{e}^{2} \mathrm{cal}^{-1} \mathrm{~m}^{-1}=1.444 \times 10^{30} q_{e}^{2} \mathrm{kcal}^{-1} \mathrm{~m}^{-1}  \tag{10.4}\\
& =2.40 \times 10^{6} q_{e}^{2}(\mathrm{kcal} / \mathrm{mole})^{-1} \mathrm{~m}^{-1}=2.40 \times q_{e}^{2}(\mathrm{kcal} / \mathrm{mole})^{-1} \mu \mathrm{~m}^{-1} \\
& =0.72 q_{e}^{2}(\mathrm{kcal} / \mathrm{mole})^{-1} \mathrm{lfs}^{-1}
\end{align*}
$$

where 'lfs' stands for light-femtosecond, the distance travelled by light (in a vacuum) in a femtosecond. Thus we see that in the units in which energy is measured in kcal/mol, charge is measured in units of the charge of the electron, $q_{e}$, and length is the light-femtosecond (lfs), we find the permittivity of free space to be on the order of unity. It is noteworthy that Debye [50] used units so that $\varepsilon_{0}=1$, together with energy measured in kcal/mol and charge measured in units of $q_{e}$. This means that the implied spatial unit is 1.39 lfs , or about 417 nanometers, or just under half a micron, a length related to the Debye screening length in water [110. If this is the chosen spatial unit, then $\varepsilon_{0}=1$ in these units. For reference, very large viruses [265] are between one and two tenths of a micron in diameter.

Draft: February 28, 2008, do not distribute

| basic units | $\varepsilon_{0}$ (permittivity), $a_{0}$ (length), $m_{e}$ (mass), $10^{-3}$ svedberg (time) |
| :--- | :--- |
| derived units | hartree $=\hbar^{2}$ (energy) |

Table 10.5: Quantum chemistry units.

### 10.2.4 Conversion constants

Boltzmann's constant, $k_{B}=1.380 \times 10^{-23}$ joules per degree Kelvin, relates energy to temperature. This seems really small, so let us convert it to the "kcal/mole" energy unit. We get

$$
\begin{align*}
k_{B} & =1.380 \times 10^{-23} \mathrm{~J} / \mathrm{K} \\
& =\frac{1.380 \times 6.022}{4.1868} \mathrm{cal} / \mathrm{mole}-\mathrm{K}  \tag{10.5}\\
& =1.984 \mathrm{cal} / \mathrm{mole}-\mathrm{K}
\end{align*}
$$

For example, at a temperature of $T=303 \mathrm{~K}$, we have $k_{B} T=0.601 \mathrm{kcal} / \mathrm{mole}$.
If temperature is in degrees Kelvin, velocities are measured in Ångstroms per picosecond (around 224 miles per hour), and masses in daltons, then $k_{B} \approx 0.831$.

Planck's constant,

$$
\begin{equation*}
h=6.626068 \times 10^{-34} \mathrm{~m}^{2}-\mathrm{kg} / \mathrm{s}=39.90165 \text { Ångstroms }{ }^{2} \text {-dalton/picosecond, } \tag{10.6}
\end{equation*}
$$

has units energy-time, which is a unit of action. The other Planck constant $\hbar=h / 2 \pi$ is then $\hbar=6.35055$ Ångstroms ${ }^{2}$-dalton/picosecond.

The ratio of Planck's constant to Boltzmann's constant has an interesting interpretation. It is $h / k_{B}=4.80 \times 10^{-11}$ seconds per degree Kelvin, or 48 picoseconds per degree Kelvin.

### 10.3 Quantum chemistry units

The Schrödinger equation has three terms which must have the same units in order to be dimensionally correct. If we divide (16.1) by $\hbar$, then the diffusion term is multiplied by the constant $\hbar / 2 \mathrm{~m}$. Fortunately, $\hbar / m$ has units of length-square over time, as required. In the Schrödinger equation (16.1) we have implicitly assumed that the permittivity of free space $\varepsilon_{0}=1 / 4 \pi$. We can do this, as noted above, but we need to choose the right spatial and energy units to make it all work out. Unfortunately, if the energy unit is kcal/mole, the natural scale for biochemistry, then the spatial unit is quite large, four orders of magnitude larger than the typcial scale of interest.

A more typical choice of spatial unit at the quantum scale [233] would be to use the Bohr radius $a_{0}=0.529189 \AA$. This scale only differes by a factor of about two from what we have been considering so far. But the natural unit of mass is the mass of the electron $m_{e}=9.10938 \times 10^{-31}$ $\mathrm{kg}=5.48579 \times 10^{-4}$ dalton. In these units, Planck's constant is

$$
\begin{equation*}
h=39.90165 \text { Ångstroms }^{2} \text {-dalton/picosecond }=137.45 a_{0}^{2}-m_{e} / \text { femtosecond. } \tag{10.7}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

If we also adopt the hartre 1 [222] $E_{h}$ for the unit of energy, and we adopt the mass of the electron $m_{e}$ as the unit of mass, then things are better. By definition, the Hartree $E_{h}$ is

$$
\begin{equation*}
E_{h}=m_{e} c^{2} \alpha^{2}=4.356 \times 10^{-18} \text { joules }=1.040 \times 10^{-21} \mathrm{kcal}=626.5 \mathrm{kcal} / \mathrm{mole} \tag{10.8}
\end{equation*}
$$

(cf. Table 3.1) where $\alpha \approx .007$ is a dimensionless number known as the fine structure constant, cf. Exercise 10.7. Moreover, we also have the coefficient of the potential in (16.1) equal to $E_{h}$; that is, $e^{2} /\left(4 \pi \varepsilon_{0} a_{0}\right)=E_{h}$.

The time-derivative term in (16.1) is multiplied by $\hbar$, which fortunately has units of energy times time. Planck's constant $h=6.626068 \times 10^{-34}$ joule-seconds $=1.521 \times 10^{-16}$ Hartree-seconds $=0.1521$ Hartree-femtoseconds. Dividing by $2 \pi$, we find that Planck's constant $\hbar=0.02421$ Hartree-femtoseconds. That is, if we take the time unit to be femtoseconds, then the coefficient of the time derivative term is $=0.02421$, or about one over forty. This is a small term. It implies that changes can happen on the scale of a few tens of attoseconds, whereas on the scale of a few femtoseconds (the typical time step of molecular dynamics simulations), the time-derivative term in (16.1) can plausibly be ignored, or rather time-averaged. To cast this in terms of the units suggested for biochemistry, the natural timescale for quantum chemistry is about $10^{-3}$ smaller, about a milli-svedberg.

In thinking of the Schrödinger equation in classical terms as describing the probability of an electron's position as it flies around the nucleus, it is interesting to think about the time scale for such a motion. At the speed of light, it takes an attosecond to go 3 Ångstroms. The time-scale of the Schrödinger equation is 24 attoseconds, and in this time anything moving at the speed of light would go 72 Ångstroms. If the Schrödinger equation represents the average behavior of electrons moving around the nucleus at anything approaching the speed of light, then they can make many circuits in this basic time unit of the Schrödinger equation. So it is plausible that it represents such an average of dynamic behavior.

### 10.4 Laboratory units

In a laboratory, it would be confusing to use units appropriate at the molecular scale. A typical mass unit would be a milligram (mg), and a typical volume unit would be a milliliter (mL). Inside cells, protein concentrations can exceed 100 mg per mL . The term 'micromolar' is used to express the fractional concentration of one substance in another, e.g., water.

### 10.5 Mathematical units

There is a natural set of units that might be called mathematical units. They are based on the observation that many named constants are really just conversion factors. For example, Boltzmann's constant really just converts temperature to energy. Thus with the right temperature scale, Boltzmann's constant is one (cf. Exercise 10.4). Similarly, Planck's constant has units energy times time,

[^2]Draft: February 28, 2008, do not distribute

| basic units | speed of light, Planck's constant, Boltzmann's constant |
| :--- | :--- |
| derived units | time unit $=0.7$ yoctoseconds, length unit $=0.2$ femtometers |

Table 10.6: Some units to simplify mathematical equations.
and it will be one with the right relationship between energy and time. This places a constraint on the relationship between mass, length, and time. A natural mass unit is the dalton, since it is roughly the mass of the smallest atom. With the dalton as the mass unit, the largest masses in the Schrödinger equation are of order one, although the smallest (i.e., the electrons) have a tiny mass in this unit. It is natural to take the speed of light to be one, so this sets a relationship between length and time.

If we divide Planck's constant by the speed of light we get $\hbar / c=0.212 \times 10^{-15}$ dalton-meters. If we want $\hbar=1$ and $c=1$ [189], then we need to have the length unit to be $0.212 \times 10^{-15}$ meters $=$ 0.212 femtometers. The diameter of a proton is approximately one femtometer.

If we divide Planck's constant by the speed of light squared we get $\hbar / c^{2}=0.7066 \times 10^{-24}$ daltonsecond. If we want $\hbar=1$ and $c=1$, then we need to have the time unit to be $0.7066 \times 10^{-24}$ seconds $=0.7066$ yoctoseconds. If these independent calculations are correct, we would find that the speed of light is about 0.3 femtometers per yoctosecond. A femtometer per yoctosecond is $10^{9}$ meters per second, so we have agreement.

To summarize, if we take length to be measured in multiples of $\mathrm{L}=0.212$ femtometers, time to be measured in multiples of $\mathrm{t}=0.7066$ yoctoseconds, and mass in daltons, then $c=\hbar=1$. See Exercise 10.5 for the similar case where the unit of mass is the mass of the electron. As noted above, a joule in these units is $6.7006 \times 10^{9}$ dalton- $(\mathrm{L} / \mathrm{t})^{2}$. Similarly, in these units $k_{B}=9.2468 \times 10^{-14} \mathrm{~K}^{-1}$.

### 10.6 Evolutionary units

There are also other time scales of interest in biology, and geology. The molecular clock refers to the time it takes for a single point mutation to occur in DNA. This is measured by comparing divergent genomes of related species, for which the time of divergence is estimated from the fossil record. The unit of measure for sequence divergence is percentage divergence, which refers to the fraction of times each individual sequence entry is expected to have been modified in a given time unit. Typically, this is a very small number, so the time unit is often taken to be large. Thus a $2 \%$ sequence divergence per million years means that the probability of mutation of each individual sequence entry is only 0.02 in a million years. However, in a hundred million years, we would expect each entry to be modified twice.

Estimates for molecular clocks vary on the order of one percent divergence per $10^{6}$ years, although turtle mitochondrial DNA mutation tends to be slower [13], possibly due to their slower internet connections. Fortunately, this is a slow scale from a human perspective. However, over geologic time, it is significant.

It is interesting to note that using typical estimates of the age of the earth 35], there has not been enough time for this type of mutation to cause a complete change to a typical chromosome.

Draft: February 28, 2008, do not distribute

Time is usually measured in units of Mya (millions of years ago), or bya (billions of years ago); the latter unit is often replaced by the shorter Ga (giga-annum). The age of the earth is estimated to be at least 4.5 Ga [35].

We can put these two pieces of data together to estimate how many times a complete genome might have been modified. If the variation is occuring at a rate of one percent per $10^{6}$ years, then in $10^{8}$ years it could become completely modified. But given the estimated age of the earth, no genome would be expected to have been fully modified more than about forty five times via single point mutations.

From the point of view of a dynamical system, a genome that has been modified 45 times might well have reached a stable equilibrium (if there is one). On the other hand, more recently evolved species would not yet be at a point where many cylces have taken place, so their genomes might be still far from an equilibrium point. Species that are older that $10^{6}$ years would have experienced substantial modification (more than one base-pair in a hundred modified). But species much younger than $10^{8}$ years might exhibit little limiting behvior, still undergoing substantial modification due to random mutations.

### 10.7 Other physical properties

We now consider some other physical properties that are measured in much the same way that basic units are. Some of them could themselves be treated as basic units, such as viscosity. Others deal with more complex issues, such as the pH scale which describes a mixture of materials.

### 10.7.1 The pH scale

At a pH of $k$, there are $10^{-k}$ moles of hydronium ions (and hydroxyl ions) per liter of water. A mole of water weighs 18.0153 grams. At 4 degress Centigrade, where water has its maximum density, one gram of water occupies one cubic centimeter, or one milliliter. Thus a mole of water occupies 0.0180153 liters (at $4^{\circ} \mathrm{C}$ ), so a liter of water has 55.508 moles of water. Thus the ratio of hydronium ions to water molecules at a pH of $k$ is roughly one hydronium ion per $5.5508 \times 10^{k+1}$ water molecules. Humans seem happiest at pH seven, which corresponds to a ratio of approximately one hydronium ion per half billion water molecules. However, the pH in cells can be much lower.

### 10.7.2 Polarity and polarization

The Debye is the standard unit for dipole moment, and is $3.338 \times 10^{-30}$ coulomb-meters. A more useful unit would be a $q_{e}$-Ångstrom, where $q_{e}$ is the charge of an electron, and this turns out to be about 4.8 Debye. Recall that a coulomb is $6.242 \times 10^{18} q_{e}$. Thus, a Debye is $0.2084 q_{e}$ - $\AA$ ngstrom. The dipole moment of water ranges from about 1.9 Debye to 3.5 Debye depending on the enviroment [100, 89].

Polarization is the effect of an external field to change the strength of a dipole. An interesting feature is that the polarization coefficient has units of volume (i.e., length cubed). Thus there is a natural motif that can be used to illustrate the polarizability of an object: the volume of its
representation. For example, if we are representing atoms as spheres, the volume of the sphere could be taken to be its polarization coefficient.

Polarization is a tensor, and it need not be isotropic. However, in many cases, a scalar approximation is appropriate. The polarizability of water is $\alpha \approx 1.2 \AA^{3}$.

### 10.7.3 Water density

Water is a molecule with a complex shape, but it is possible to estimate the volume that an individual molecule occupies. A mole of water, $6.022 \times 10^{23}$ water molecules, weighs 18.0153 grams. At 4 degress Centigrade, where water has its maximum density, one gram of water occupies one cubic centimeter, or $10^{24} \AA^{3}$. Thus a mole of water occupies $180.153 \times 10^{23} \AA^{3}$ (at $4^{\circ} \mathrm{C}$ ), so a single molecule of water occupies about $29.92 \AA^{3}$. This corresponds to a cube of just over $3.1 \AA$ Ångtroms on a side. It is interesting to compare this distance with the typical O-O distance in water (about $2.75 \AA$ ).

### 10.7.4 Fluid viscosity and diffusion

Fluids display an aggregate behavior known as viscosity. Fluid dynamicists [] call the viscosity $\mu$ and physicists [] call it $\eta$. The units of the coefficient of viscosity (often called dynamic viscosity) are mass per length-time. A standard unit of viscosity is the poise 2 which is one gram per centimeter-second. One poise is 0.1 Pascal-second, where a Pascal is a unit of pressure or stress. One pascal is one newton per meter-squared, where we recall that a newton (one kilogram-meter/second-squared) is a measure of force.

The viscosity of water at 293 degrees Kelvin (20 degrees Centigrade) is about one centipoise, or about 0.001 Pascal-second. The viscosity of olive oil is about 80 times larger, so the ratio of viscosities of olive oil and water is roughly the ratio the dielectric of water and vacuum. The viscosity of air is 0.0018 centipoise, over a factor of five-hundred smaller.

### 10.7.5 Kinematic viscosity

Another scaling factor is significant in fluid flow, namely the fluid density. The ratio of viscosity (or dynamic viscosity) and density is called kinematic viscosity, usually labelled $\nu$. This has units length-squared per time, since density has units of mass per length-cubed. Thus kinematic viscosity has the same units as a spatial diffusion constant. The stoke is one centimeter-squared per second. The kinematic viscosity of water is about one millimeter-squared per second, or one centistoke, whereas the kinematic viscosity of air is rougly two times larger. That is, air is more viscous than water! The viscosity of fluids varies significantly with temperature, but we have provided values at roughly the same temperature ( 293 K ) for comparison.

[^3]Draft: February 28, 2008, do not distribute

Viscous drag is the effective force of viscosity in opposing motion. It provides a retarding force in the direction oppositve to the motion. The drag coefficient has the units of force divided by velocity, or mass per time unit.

### 10.7.6 Diffusion

### 10.8 Exercises

Exercise 10.1 Using the Bohr radius $a_{0}=0.5291772 \AA$ as the basic unit of length, the svedberg as time unit, and the amu as mass unit, compute the unit of energy, one amu-(bohradius/svedberg) ${ }^{2}$ in terms of the unit kcal/mole.
Exercise 10.2 Determine a basic unit of length $L$ such that, with the svedberg as time unit, and the dalton as mass unit, then the unit of energy of one dalton-(L/svedberg $)^{2}$ is exactly one kcal/mole. Compare $L$ with the van der Waals radius of different atoms (which are closest?).
Exercise 10.3 Determine a three-dimensional volume which can be used to tile space and fits a water molecule better than a cubic box. Use this volume to estimate the density of water.
Exercise 10.4 Suppose we take the dalton as the mass unit and that we choose space and time units so that the speed of light and Planck's constant are both one. What is the temperature scale that makes Boltzmann's constant equal to one?

Exercise 10.5 Suppose we take the mass of the electron as the mass unit, and that we want units so that the speed of light and Planck's constant are both one. What are the corresponding time and length scales?

Exercise 10.6 Suppose temperature is in degrees Kelvin and mass is in daltons. Determine a velocity scale such that $k_{B}=1$.
Exercise 10.7 The fine structure constant is

$$
\begin{equation*}
\alpha=\frac{q_{e}^{2}}{2 h \varepsilon_{0} c}, \tag{10.9}
\end{equation*}
$$

where $m_{e}$ and $q_{e}$ are the mass and charge of the electron, respectively, $\varepsilon_{0}$ is the permittivity of free space, $h$ is Planck's constant, and $c$ is the speed of light in a vacuum. Prove that $\alpha$ is dimensionless. Determine other combinations of various physical constants that are also dimensionless.
Exercise 10.8 The Rydberg constant $R_{\infty}$ is

$$
\begin{equation*}
R_{\infty}=\frac{m_{e} q_{e}^{4}}{8 \varepsilon_{0}^{2} h^{3} c} \tag{10.10}
\end{equation*}
$$

where $m_{e}$ and $q_{e}$ are the mass and charge of the electron, respectively, $\varepsilon_{0}$ is the permittivity of free space, $h$ is Planck's constant, and $c$ is the speed of light in a vacuum. Prove that the hartree $E_{h}=2 R$.

Exercise 10.9 We have two equations for the Hartree $E_{h}$, namely, $E_{h}=m_{e} c^{2} \alpha^{2}$ and $q_{e}^{2} /\left(4 \pi \varepsilon_{0} a_{0}\right)=$ $E_{h}$. Show that these are compatible, that is, $4 \pi \varepsilon_{0}=q_{e}^{2} / a_{0} m_{e} c^{2} \alpha^{2}$.

## Chapter 11

## Case studies

We have established that dehydrons are sticky binding sites for protein ligands. Here we examine several cases where these play an important biological role.

### 11.1 Basic cases

Several examples of the role of dehydrons in protein associations were given in [79]. We review them here briefly.

### 11.1.1 A singular case

One striking example of a dehydron involved in protein-protein association is the binding of the light chain of antibody FAB25.3 with the HIV-1 capsid protein P24. In the monomeric state, the capsid protein has two dehydrons, one of which appears in a helix. The binding interface for this complex is quite small and yet the single helical dehydron of the HIV-1 capsid protein lies precisely at the interface with the extended finger of the light chain.

### 11.1.2 Forming structures

Dehydrons also appear more broadly in structural roles 79. In some cases, they guide formation of quarternary structues, such as dimers. For example, dehydrons (G49, G52), (G78, T80), and (T91, G94) in the HIV-1 protease guide the formation of the dimer structure [76]. In other cases, the structures can be quite complex.

Virus capsids provide a valuable model for protein-protein interaction [262, 263, 264]. The formation of the capsid in picornaviruses [94, 141] was shown to be essentially determined by the distribution of dehydrons in the individual virus-peptide (VP) subunits [79]. This distribution concentrates at the symmetry centers of the capsid and edge-to-edge subunit positioning.

There are three individual VP (virus peptide) subunits which assemble into a virus unit for the foot-and-mouth disease virus (FMDV) [94. The atomic coordinates correspond to pdb entry 1BBT. Regions of high concentration of dehydrons are found at the symmetry centers of the capsid,
and those dehydrons become desensitized upon unit associations (Figure 5a in [79). Approximately $67 \%$ of the dehydrons involved in domain-swapping and unit assembly become desensitized upon formation of the unit. However, the unit has four remaining regions with a high concentration of dehydrons.

A comparison between Figures 5a and 5b of [94] reveals that the four regions in the unit with a high dehydron density can be readily associated either with the centers of symmetry of the capsid or to the inter-unit edge-to-edge assembly of VP2 and VP3 94. Thus, the FMDV unit presents two loop regions highly sensitive to water removal on VP1 and VP3 which are directly engaged in nucleating other units at the pentamer and hexamer (with three-fold symmetry) centers of the capsid. Furthermore, a helical region on VP2 is severely under-wrapped and occupies the central dimeric center (shown in Fig. 5b of [94]).

The structurally defective 'handle' region on FMDV-VP2, an under-wrapped $\beta$-hairpin near the N-terminus (12-27 region) is also present in the VP2 subunits of all the picornaviruses. Because of its high density of dehydrons, this region is a strong organizing center in the capsid.

Similar roles for dehydrons are seen in the human rhinovirus (PDB code 1R1A) [131. While most dehydrons can be attributed to the unit assembly, there are three particular sites with high dehydron density which do not become well wrapped either after the formation of the unit or after the assembly of the whole capsid. Two of them correspond to antibody binding sites, but the other site lies under the so-called canyon of the VP1 structure and has been known to be the target region for the drug WIN 51711 used to treat common cold [131.

The Mengo encephalomyocarditis virus (PDB code 2MEV) [141] has a large number of crystal contacts. These are points at which each individual unit makes contact with another in the formation of a crystal used in X-ray imaging. Thus, they may be thought of as a type of artifact of the protein-protein interactions related to the imaging process. The viral unit has only two very strong dehydron centers on its rim: the pentamer center located in VP1, and the ubiquitous VP2VP3 edge-to-edge region located in VP2. The remaining 15 dehydrons are listed in Table 11.1 are not involved in the organization of the capsid. Of the 60 residues known to be engaged in crystal contacts for this virus [141, 54 of them have sidechain carbonaceous groups in the desolvation domains of the 15 dehydrons marked in Table 11.1. Thus, the dehydons not involved in capsid organization correlate with the crystal packing.

The number of dehydrons of a given protein in different organisms can vary substantially [79] as we have depicted in Figure 2.1] for myoglobin. Six other groups of proteins were analyzed in [79], and shown to display similar trends. Species with a rich interactive complexity for a particular protein are have a higher number of dehydrons, whereas in more archaic species, the homologous proteins are shown to have far fewer dehydrons. A clear illustration of this trend is provided in Fig. 3 in [79] where three different versions of myoglobin corresponding to aplysia limacina (gastropode, mollusc), whale and human are displayed and their respective distribution of dehydrons is highlighted. The myoglobin from aplysia limacina is one of the best wrappers of hydrogen bonds in the entire PDB 79.

Draft: February 28, 2008, do not distribute

| A84-LEU-(H)-A81-TYR-(O) |
| :--- |
| A98-GLU-(H)-A96-GLY-(O) |
| A96-GLY-(H)-A102-GLU-(O) |
| B162-LYS-(H)-A96-GLY-(O) |
| A101-SER-(H)-A98-GLU-(O) |
| A205-HIS-(H)-B208-GLN-(O) |
| A210-ASN-(H)-A207-ARG-(O) |
| A212-GLY-(H)-A210-ASN-(O) |
| B252-VAL-(H)-B101-ARG-(O) |
| B161-ARG-(H)-B159-THR-(O) |
| B228-ALA-(H)-B225-SER-(O) |
| C72-VAL-(H)-C70-THR-(O) |
| C141-GLN-(H)-C138-SER-(O) |
| C180-ILE-(H)-C178-ALA-(O) |
| C204-SER-(H)-C202-PRO-(O) |

Table 11.1: List of dehydrons for the protein unit of Mengo encephalomyocarditis virus (PDB code 2MEV) 141 engaged in known crystal contacts (their wrapping hydrophobic groups belong to the side chains of residues known to form crystal contacts [131). The proton donor residue is marked as $(\mathrm{H})$ and the electron-donor residue supplying the carbonyl group is marked $(\mathrm{O})$.

### 11.2 Enzymatic activity

Enzymes are proteins that catalyze (i.e., enhance) chemical reactions. We can think of them as machines, since they start with an input resource, a molecule called the substrate, and convert the substrate into a different molecule, the product. The process is called catalysis. The chemical reaction that is facilitated by an enzyme would occur naturally without a catalyst, but at a much slower rate.

Enzymes can be quite large proteins [252, 251], but the 'active site' in which catalysis takes place is localized. There are thousands of enzymes that play a role in metabolic pathways [214]. Many drugs are designed to be inhibitors of enzymes [76], disrupting the role of an enzyme in the metabolic system.

The active site of an enzyme will contain water until the substrate enters. Thus water removal is critical to the enzymatic process [251, 76]. The process of water removal can be enhanced energetically by the presence of dehydrons near the active site [76. Although most drugs were not designed to facilitate water removal, we can see retrospectively that particular drugs play a significant role in wrapping dehydrons 76.

Draft: February 28, 2008, do not distribute


Figure 11.1: Wrapping of hydrogen bond Asn48N-Gln45O in PDB file 2BN2. The hydrogen (in white) attached to Asn 48 N has been added to indicate the bond, which is in the middle of the figure. The ligand pair Tyr99 and Phe98 is in the upper left of the figure. Residues 45 to 50 in the A chain are shown to indicate the bulk of the wrapping.

### 11.3 Neurophysin/vasopressin binding

Aromatic rings can in principle interact with any polar moiety (cf. Section 13.2), and this type of interaction has been suggested as an explanation for the binding of neurophysin and vasopressin. We analyze this interaction here and exhibit some additional candidates to explain the particular details of the interaction zone.

In [29], a polar/aromatic interaction is suggested for the residue Tyr99 of the hormone vasopressin, a ligand of the bovine protein neurophysin-II, a transporter of hormones along axons. Part of the reason for this suggestion is the large number of disulfide bonds in neurophysin [212]. There are seven such bonds in a protein with only 79 residues; nearly a fifth of all of the residues are involved in one. Moreover, there is a disulfide bond in the ligand vasopressin.

However, it is also the case that Tyr99 wraps an under-wrapped mainchain-mainchain hydrogen bond between Asn48-N and Gln45-O in PDB file 2BN2. It is wrapped by 13 nonpolar groups in chain A: PRO24-CB; GLN45-CB,CG; GLU 46-CB,CG; GLU 47-CB,CG; ASN48-CB; TYR49CB,CG,CD1CE1,CB. However, 12 of these come from the surrounding helix and leave a large gap. The active site, in which Tyr99 inserts, leaves a large water-exposed area. Indeed, both Tyr99 (CD2,CE2) and Phe98 (CB,CG) provide wrappers for the Asn48N-Gln45O hydrogen bond. It is clear that they provide wrapping in a sector that is otherwise exposed to water attack.

The terminal O-H group in Tyr99 also forms a sidechain-mainchain hydrogen bond with Cys44-
O. This presumably accounts for the additional stability that this ligand displays compared to the Phe98-Phe99 variant [29], the corresponding residue pair at the binding site for the ligand phenypressin, a hormone in Australian macropods.

The role of Phe98 is also of interest. It is also a wrapper of the sidechain-sidechain hydrogen bond ARG8-NE-SER52-OG. But it further makes mainchain-mainchain intermolecular hydrogen bonds, with CYS54-N and either LEU50-O or SER52-O. Apparently, only peptides with Phe in the first position bind with detectable affinity, and the fact that Phe98 wraps two dehydrons, as well as making intermolecular hydrogen bonds, contributes to its uniqueness. However, it is known that the change from Phe to Leu in position one still preserves significant affinity [29]. If the position of Leu is the same as that of Phe, it would preserve both its mainchain hydrogen bonds and its contribution to wrapping two dehydrons.

On the other hand, the combination Phe-Leu has significantly reduced affinity. This could be due to two different reasons. One could be a polar-aromatic interaction of some sort, but it could also be due to the factors discussed here. Compared with Tyr, Leu lacks the sidechain-mainchain bond with Cys44. Moreover, it is positions CD and CE on residue 99 that provide the wrapping. These positions would be near the end of the Leu sidechain, so its ability to wrap, compared with Phe, would be reduced. Similarly, two other second-position sidechains (Met and His) show minimal affinity, consistent with their lack of wrappers at the end of the sidechain and lack of ability to form a hydrogen bond at the end with Cys44.

The role of residue 48 in chain A was explored in [29] by a natural 'mutation' arising due to the fact that bovine (Asn48) and ovine (Ile48) differ at this location. The ovine affinity is slightly higher, but of the same order. The 'mutation' Asn $48 \rightarrow$ Ile48 is isosteric, so it is plausible that the mainchain-mainchain hydrogen bond to Gln45-O is maintained.

A similar picture emerges in the related PDB structure 1JK4 bound to a larger fragment of vasopressin [255]. The individual bonds are different, but the general picture is the same. One new ingredient is the dehydron in the vasopressin fragment CYS6-N-TYR2-O, which becomes wrapped by GLU47-CB and ASP76-CB.

### 11.4 Variations in proteomic interactivity

We now review how dehydrons provide a novel indicator of proteomic interactivity [82]. We have seen in Section 2.3 that dehydrons in the protein myoglobin vary among different species. This trend is further confirmed in Table 11.2 with other proteins. A larger number of dehydrons implies a larger number of possible interactions, due to the fact that they are sticky (Chapter [8). Thus it is reasonable to ask if these observation can be used to relate classes of proteins using their potential interactivity, as measured by the number of dehydrons, as an indicator.

Relations among proteins can be determined by various means. The notion of 'fold' or 'domain' characterizes the basic unit of tertiary structure of proteins [256, 254]. Although most proteins consist of a single domain, a significant number contain multiple domains. Thus one way to form a relation among proteins is to define the vertices of a graph to be the protein domains and the edges of the graph to be pairs of domains found in a single protein [256]. The resulting graph of protein domain connectivity has many nodes with low degree with just a few highly connected
nodes, commonly referred to as a scale-free (or small world) graph, in agreement with a power-law distribution [256].

Although relating protein folds by their inclusion in a single protein reveals some type of fold interactivity, it is possible to choose more direct measures of protein interaction. Interactivity of proteins can be determined by various means, and this is documented in various databases [28], such as DIP, the Database of Interacting Proteins [209]. The objective of DIP is "to integrate the diverse body of experimental evidence on protein-protein interactions" [209].

In [82], a subset of DIP was selected containing domains from the yeast proteome whose interactions were determined by a single class of experiments. For the proteins in this subset having PDB structures, the average amount of wrapping of all hydrogen bonds was determined using a desolvation-sphere radius of $6.4 \AA$ together with the number of dehydrons using a strict requirement of $\rho_{P G}<12$ (cf. Section [7.3) imposed in the definition of dehydron [82]. A strong correlation was found between the number of dehydrons and the number of interacting partners as reported by DIP. Thus there is a strong indication that the potential interactions related to dehydrons correlate with actual protein interactions.

By examining conserved folds across species, significant differences in the number and distribution of dehydrons were found. Within a conserved domain fold, the number of dehydrons in higher eukaryotes is consistently greater than in species (e.g., bacteria) of lower complexity.

Known data on protein interactivity can be used to represent a network, where each protein domain is represented as a node. Nodes with few connections are common, and highly connected nodes are rare, in agreement with a power-law distribution [256]. Connectivity of different domains in this representation is proportional to the average number of dehydrons in the family [82]. Furthermore, the dehydron patterns associated with structural domains for a given species consistently define a characteristic scale-free interactive network. the defects in the packing of structural domains within a given species consistently determines a characteristic exponent that describes the node distribution within a scale-free interaction network [21, 6].

The greater proteomic interactivity of higher eukaryotes does not result from their genome sizes or even estimated transcriptome sizes [135, 256, 240]. Even the combinatorial multiplicity due to domain shuffling cannot account for qualitative differences in the power-law description of proteomic connectivity [135].

A similar analysis was carried out in [82] using a different definition of interactivity, based on the Structural Classification of Proteins (or SCOP) superfamilies [176, 154] as vertices in a graph. Interactivity among the superfamilies was defined using a variant of method described initially [256] utilizing protein domains with identified interactions (complexation or intramolecular interaction) in PDB files from different superfamilies (cf. [188, 120, 124, 3, 57]) to define graph edges. To account for differences in protein length, an average $r_{d / H B}$ of dehydrons per hundred hydrogen bonds was determined [82]. Again, a strong correlation emerged between interactivity and $r_{d / H B}$.

With $r_{d / H B}$ established as an indicator of protein interaction, it is then possible to look for differences among species. Define [82] data points $f_{r}$ to be the percentage of protein (SCOP) superfamilies having $r_{d / H B}=r$, where the values of $r$ consist the set of $r_{d / H B}$ values for all the superfamilies. Thus $f_{0}=100$ and $f$ forms a type of decreasing relation. Since two families can have the same $r_{d / H B}$ value, $f$ cannot be extended to be a function of a continuous parameter $r$, only a

Draft: February 28, 2008, do not distribute

| Protein | Species (common name) | PDB code | $N$ | $H B$ | $\rho$ | $d$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Cytochrome c | Chlamydomonas reinhardtii (algae) | 1 cyi | 89 | 52 | 19.74 | 6 |
| Cytochrome c | Rhodophila globiformis (bacteria) | 1 hro | 105 | 50 | 17.52 | 7 |
| Cytochrome c | Oryza sativa (Asian rice) | 1 ccr | 111 | 55 | 14.94 | 11 |
| Cytochrome c | Katsuwonus pelamis (skipjack tuna) | 1 cyc | 103 | 41 | 14.03 | 12 |
| Cytochrome c | Thunnus alalunga (albacore tuna) | 5 cyt | 103 | 53 | 14.25 | 13 |
| Cytochrome c | Equus caballus (horse) | 1 giw | 104 | 44 | 14.01 | 14 |
| Hemoglobin | Vitreoscilla stercoraria (bacteria) | 2 vhb | 136 | 102 | 23.50 | 0 |
| Hemoglobin | Lupinus luteus (yellow lupin, plant) | 1 gdj | 153 | 109 | 23.43 | 0 |
| Hemoglobin | Paramecium caudatum (protozoa) | 1 dlw | 116 | 77 | 22.02 | 0 |
| Hemoglobin | (Nonsymbiotic) Oryza sativa (rice) | 1 d 8 u | 165 | 106 | 23.58 | 2 |
| Hemoglobin | Equus caballus (horse) | 1 gob | 146 | 101 | 21.45 | 2 |
| Hemoglobin | H. sapiens | $1 \mathrm{bz0}$ | 146 | 103 | 21.45 | 3 |
| Myoglobin | Aplysia limacina (mollusc) | 1 mba | 146 | 106 | 23.42 | 0 |
| Myoglobin | Chironomus thummi thummi (insect) | 1 eca | 136 | 101 | 21.31 | 3 |
| Myoglobin | Thunnus albacares (yellow-fin tuna) | 1 myt | 146 | 110 | 21.15 | 8 |
| Myoglobin | Caretta caretta (sea turtle) | 11 ht | 153 | 110 | 21.09 | 11 |
| Myoglobin | Physeter catodon (sperm whale) | $1 \mathrm{bz6}$ | 153 | 113 | 20.98 | 11 |
| Myoglobin | Sus scrofa (wild boar) | 1 mwc | 153 | 113 | 19.95 | 12 |
| Myoglobin | Equus caballus (horse) | 1 dwr | 152 | 112 | 18.90 | 14 |
| Myoglobin | Elephas maximus (Asian elephant) | 1 emy | 153 | 115 | 18.90 | 15 |
| Myoglobin | Phoca vitulina (seal) | 1 mbs | 153 | 109 | 18.84 | 16 |
| Myoglobin | H. sapiens | 2 hbc | 146 | 102 | 18.80 | 16 |

Table 11.2: Variations in numbers of dehydrons in homologous proteins in different species. $N$ is the number of residues in the protein, $H B$ denotes the number of hydrogen bonds, $\rho$ is the average wrapping of the hydrogen bonds in the protein, and $d$ denotes the number of dehydrons in the protein. The quantity $r_{d / H B}$ in [82] is $100 d / H B$.
relation. The set of values $\left(r, f_{r}\right)$ were plotted in [82] on a log-log scale and a power-law behavior

$$
\begin{equation*}
f(r) \approx c r^{-\gamma} \tag{11.1}
\end{equation*}
$$

emerged for each species (E. coli, mouse and human). Not surprisingly, in view of Table 11.2, the values of $\gamma$ were significantly different for different species (2.1, 1.49, 1.44, respectively).

The results reviewed here present two different views of the correlation of dehydrons and interactivity. On the one hand, interactivity of broad classes of protein with similar folds (including many different species) tends to increase as a function of the average number of dehydrons in that class. Thus for example, we see in Table 11.2 that hemoglobin has far fewer dehydrons than myoglobin, when averaging over all the species represented. However, it is also possible to subdivide the distribution of dehydrons across all proteins according to species, and a differentiation appears in the form (11.1). Although these two statements may seem contradictory at first, they are rather complementary.

### 11.5 Sheets of dehydrons

It is possible for protein systems to form using only indirect dehydration forces. One example is given by associations of $\beta$-sheets [69].

### 11.6 Exercises

Exercise 11.1 Determine the residues most likely to be involved in catalytic activity in the active site of an enzyme.

Exercise 11.2 The residue Asp is often involved in catalytic activity in the active site of an enzyme. It is often found to make local sidechain-mainchain bonds (cf. Chapter [12) in an underwrapped environment. Explore the possible correlation of these two observations.

## Chapter 12

## Sidechain-mainchain hydrogen bonds

It is remarkable that there are more mainchain-mainchain hydrogen bonds than ones involving sidechains. Data taken from a subset of the PDB Select database consisting of 1547 proteins complexes (PDB files) has only 68,994 hydrogen bonds between sidechains and mainchains, contrasted with 233,879 mainchain-mainchain hydrogen bonds. In addition, there are 33,021 sidechainsidechain hydrogen bonds, 152 of which involve terminal oxygens. If we classify bonds according to whether the sidechain is the donor (S-M) or acceptor (M-S), then we find that 30,640 of the total hydrogen bonds between sidechains and mainchains have the sidechain as acceptor (M-S), not including another 75 which involve terminal oxygens. Correspondingly, the remainder $(38,277)$ are S-M bonds.

### 12.1 Counting the bonds

In thinking about the likelihood of finding one type of bond versus another in proteins, there are two ways of looking at the question. Above, we have taken the view that is suitable for the following question: when looking at a protein, are we more likely to see mainchain-only hydrogen bonds than ones involving sidechains? This is a useful question to ask, since it says something about the contributors to the energy of binding contributed by the various types of hydrogen bonds. But there is a different point of view we could take. We might instead be interested in the likelihood of a particular donor or acceptor being involved in a hydrogen bond. This is a different question because the numbers of donors and acceptors are different for the mainchain versus the sidechains. Exploring this question reveals typical issues that have to be dealt with when datasets are examined from different perspectives.

Some of the differences in numbers of mainchain-mainchain hydrogen versus bonds involving sidechains can be explained by the differences in the number of hydrogen bond donors and acceptors. Every mainchain unit can form hydrogen bonds, but not all sidechains can. In Table 5.2, we see that there are about one-third fewer (thirteen out of twenty) donors and acceptors for sidechains than mainchains. Note that the counts are simplifed because there are both thirteen donors and thirteen acceptors for sidechains. If the numbers were different, it would be more difficult to compare them with the mainchain donors and acceptors.

Making a correction for the differences in numbers of donors/acceptors narrows the gap somewhat, but the observed difference is still greater. To account for the deficit in donors and acceptors, we can multiply the number of M-S and S-M bonds by $20 / 13 \approx 1.54$, the ratio of potential donors/acceptors for M-S or S-M versus M-M bonds. The resulting number corresponds to 'virtual' bonds that would exist if the numbers of donors and acceptors were the same, and these numbers can be directly compared with the number of M-M bonds. Combining the number of M-S and S-M bonds $(68,917)$ and multiplying by the factor $20 / 13$, we get about 106 K 'virtual' bonds. For the sidechain-sidechain bonds, we need to multiply by $(20 / 13)^{2} \approx 2.37$, yielding about 78 K 'virtual' bonds, for a total of 186 K 'virtual' bonds. Thus the likelihood of forming mainchain-mainchain bonds could be viewed as about a quarter more frequent than formation of a sidechain bond. Nevertheless, in terms of energy budget, the mainchain-mainchain bonds remain dominant.

The likelihood of finding a sidechain involved in a hydrogen bond depends on the likelihood of finding that sidechain in a protein. Thus a more careful analysis would involve the frequencies of individual residues in proteins (cf. Table 6.2). We leave this task as Exercise 12.2 ,

### 12.2 Proline-like configurations

Proline is the unique residue that turns back and attaches a second time to the backbone. This forms a very rigid configuration which thus has special properties. However, other residue configurations have somewhat the same character, based on hydrogen bonds formed by the sidechains with the nearby amide or carbonyls on the backbone. We review this interesting behavior here and note the relationship to wrapping.

One feature of hydrogen bonds between mainchains and sidechains is that a large portion of them involve a bond between the sidechain and the amide or carbonyl of their own peptide, or of a neighboring peptide in the sequence. In this way, these sidechains form a structure that is similar to that of proline, but with the covalent bond in proline replaced by a weaker hydrogen bond.

One thing that characterizes such local attachments is that they tend to be underwrapped compared to hydrogen bonds between sidechains and mainchains that are more distant in sequence. The distribution of wrapping of all hydrogen bonds in this set of protein structures is depicted in Figure 12.1. First of all, we need to say how the wrapping of bonds involving sidechains was computed. To keep within the framework used for estimating wrapping of mainchain-mainchain bonds (Chapter (7), we used again a desolvation domain consisting of spheres centered at the two $C_{\alpha}$ carbons of the corresponding peptides. However, it should be noted that hydrogen bonds involving sidechains are quite different, and it is likely that a different metric would be more appropriate (and accurate).

We see that the wrapping of all hydrogen bonds involving sidechains is less than that for mainchain-mainchain bonds, with the deviation greatest for the M-S bonds for which the sidechain forms the acceptor of the bond, the donor being a mainchain amide group. An underwrapped mainchain amide or carbonyl would be a likely target for water attachment. Thus the structural defect associated with underwrapping appears to be corrected by certain sidechains making hydrogen bonds with the exposed backbone amides or carbonyls. Moreover, the formation of the hydrogen bond also removes the sidechain from water exposure as well. The major contributors to

Draft: February 28, 2008, do not distribute 138


Figure 12.1: Distribution of wrapping for hydrogen bonds in a subset of the PDB Select structures. Bonds involving self-attachment, which have a significantly smaller desolvation domain, have been excluded. Solid line: M-M; large-dash line: M-S; small-dash line: S-M; dotted line: S-S. Desolvation radius $6 \AA$.
this motif are Thr, Ser and Asp, which have relatively little sidechain flexibility, so only a limited amount of entropy is lost in these associations.

We should note that we are only considering sidechain bonds that are formed within the same chain. That is, the intermolecular bonds formed between different chains are not counted here.

### 12.2.1 Nearest neighbor connections

Some sidechains can form a hydrogen bond with their own mainchain peptide group. For example, Asp and Glu can form hydrogen bonds between their terminal oxygens and the NH group on the backbone. An example of this is found in the PDB file 1NDM in the bond between the NH and OE1 of B-GLU306. In this bond, the N-O distance is only $2.78 \AA$ (cf. Table 5.1, second row), and the angle between the NH and the CO is quite favorable. This is depicted in Figure 12.2 (a). A similar bond is formed by A-GLU81 in 1NDM, and the N-O distance is only $2.37 \AA$. In the homologous structure in 1NDG, we find the second of these motifs (A-GLU81) repeated, as well as two more: A-GLU123 and B-GLU301. But the simple motif involving B-GLU306 becomes a complex of two Glu's (B-GLU 306 and B-GLU 405) with symmetric mainchain-sidechain bonds (N-H bonded to OE's) between the two as well as a self-bond (B-GLU 405), as shown in Figure 12.2 (b).

The analogous type of bond can be formed with Gln, as in 1MPA (H-GLN113) and 1WEJ (H-GLN109). Although the turn is tighter, this motif also occurs with Asp: in 2H1P (H-ASP480) and in 2BSR (A-ASP106). And similarly, the motif occurs with Asn: in 1CU4 (L-ASN138), 1E4W (P-ASN4) and 1JRH (I-ASN53).

A similar type of motif can occur with Asn, in which the terminal amide group bonds with the


Figure 12.2: Ball and stick representation of the main atoms. Oxygens in red, nitrogen in blue, carbons in light grey, and hydrogen in white. Hydrogen positions have been estimated. (a) BGLU306 sidechain in the PDB file 1NDM, including the C-O group from peptide 305. (b) B-GLU306 (lower right) and B-GLU 405 (upper left) complex in the PDB file 1NDG, including the C-O groups from sidechains 305 and 404.
backbone oxygen of the same residue, such as in 1IC4 (Y-ASN37), 1JRH (I-ASN62) and 1MPA (H-ASN3). With Thr, the terminal OH group can bond with the backbone oxygen of the same residue as well, as in 1NDM (B-THR431) and 1DQM (H-THR132). With Ser, the terminal OH group can bond with the backbone oxygen of the same residue as well, as in 2H1P (L-SER32) and 1IGC (L-SER202).

The terminal $\mathrm{NH}_{3}$ group on lysine can bond with its own backbone oxygen, as occurs in 1WEJ (H-LYS136). A related type of motif can occur with Gln, in which the terminal amide group bonds with the oxygen of the preceding residue, such as (A-GLN89 NE2 - A-GLN90 O) in both PDB files 1NDG and 1NDM. This can also happen with Ser, with the terminal oxygen bonding with the next backbone amide group, as in 1DQJ (C-SER86 N-C-SER85 OG).

In Table 12.1, we tabulate the occurrences all of the observed local bonds where the sidechain bonds to its own backbone. This data is taken from a subset of the PDB Select database consisting of 1547 proteins complexes (PDB files) having 68,994 hydrogen bonds between sidechains and mainchains. (This is to be contrasted with a total of 233,879 mainchain-mainchain hydrogen bonds, and 33,217 sidechain-sidechain hydrogen bonds, 152 of which involve terminal oxygens.) If we classify bonds according to whether the sidechain is the donor (S-M) or acceptor (M-S), then we find that 30,642 of the total hydrogen bonds between sidechains and mainchains have the sidechain as acceptor (M-S), not including another 75 which involve terminal oxygens. Correspondingly, the remainder $(38,277)$ are S-M bonds. Thus the special bonds tabulated in Table 12.1 represent 2,729 (2,078 M-S and 651 S-M) hydrogen bonds, or about $4 \%$ of the total hydrogen bonds between sidechains and mainchains ( $6.7 \%$ of M-S and $1.7 \%$ of S-M).

The hydrogen bonds in Table 12.1 appear at first to be extremely underwrapped, but it must be remembered that in the case of a self-bond, the definition of the desolvation domain would involve only one sphere. Thus the desolvation domain is about $40 \%$ smaller than a desolvation domain

| offset | residue | type | donor | acceptor | frequency | wrapping | corrected |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 0 | ASN | M-S | N | OD1 | 116 | 8.5 | 14.4 |
| 0 | HIS | M-S | N | ND1 | 236 | 9.0 | 15.3 |
| 0 | GLN | M-S | N | OE1 | 267 | 10.2 | 17.3 |
| 0 | ASP | M-S | N | OD1 | 478 | 8.7 | 14.7 |
| 0 | GLU | M-S | N | OE1/2 | 981 | 10.5 | 17.8 |
| 0 | CYS | S-M | SG | O | 24 | 8.0 | 13.6 |
| 0 | LYS | S-M | NZ | O | 55 | 9.6 | 16.3 |
| 0 | SER | S-M | OG | O | 67 | 6.8 | 11.5 |
| 0 | THR | S-M | OG1 | O | 79 | 8.3 | 14.1 |
| 0 | GLN | S-M | NE2 | O | 83 | 11.0 | 18.7 |
| 0 | ASN | S-M | ND2 | O | 119 | 7.7 | 13.0 |
| 0 | ARG | S-M | Nx | O | 224 | 10.0 | 17.0 |
| 0 | ARG | S-M | NH2 | O | 19 | 9.7 | 16.4 |
| 0 | ARG | S-M | NE | O | 65 | 9.7 | 16.4 |
| 0 | ARG | S-M | NH1 | O | 140 | 10.1 | 17.1 |

Table 12.1: Observed local hydrogen bonds between a residue and its own mainchain amide or carbonyl groups. The bonds made by arginine are further subdivided according to the specific hydrogen bond donor group. The corrected wrapping values in the final column are simply the wrapping value times 1.7, to account for the difference in size of desolvation domain (desolvation radius $6 \AA$ ). See Figure 12.3 parts (a) and (b).
(a)



(d)


Figure 12.3: Hydrogen bond-forming sidechain configurations in the case of (a) zero-offset, mainchain-sidechain (M-S) bonds; (b) zero-offset, sidechain-mainchain (S-M) bonds; (c) +1 offset, mainchain-sidechain (M-S) bonds; and (d) -1 offset, sidechain-mainchain (S-M) bonds. See Table 12.1 for (a) and (b) and Table 12.2 for (c) and (d).
consisting of two spheres centered on $C_{\alpha}$ 's separated by $6 \AA$ (for a sphere radius of $6 \AA$ ). Thus we might increase the wrapping numbers by about $70 \%$ in order to get a realistic comparison. With this correction (listed in the last column in Table 12.1), the bonds still appear slightly underwrapped.

The average amount of wrapping found in the same set of protein complexes was 17.8 for M-S hydrogen bonds and 19.9 for S-M hydrogen bonds (for a desolvation sphere radius of $6 \AA$ ), with the bonds removed from the calculation that involve the same residue, due to the smaller desolvation domain in that case. The distribution is shown in Figure 12.1. Adjusting these for a $40 \%$ decrease in volume of the desolvation domain would suggest expected values of 10.7 (M-S) and 11.9 (S-M), respectively. For reference, the mean wrapping of mainchain-mainchain hydrogen bonds for this data set is 21.4 and 20.5 for sidechain-sidechain hydrogen bonds. Thus we see that, in general, sidechain hydrogen bonds are less well wrapped than mainchain-mainchain bonds, with M-S hydrogen bonds significantly less well wrapped.

In particular, we see that both Glu and Gln are about as well wrapped when they make M-S hydrogen bonds to their own backbone amide groups as M-S hydrogen bonds are in general. On the other hand, Asp and Asn tend to be significantly less well wrapped when they make M-S hydrogen bonds to their own backbone amide groups compared with M-S hydrogen bonds are in general. The residues of Asp and Asn themselves contribute one less wrapper to the desolvation domain, but even adding one to the corrected wrapping values (to account for the additional intrinsic wrapper in the sidechains of Glu/Gln versus Asp/Asn) leaves their mean wrapping values more than two lower than the average of M-S hydrogen bonds in general.

On the other hand, all of the sidechains that form S-M bonds to their own carbonyl groups are, on average, significantly underwrapped compared to the average wrapping (19.9) of S-M bonds. The only exception to this is Gln, whose average wrapping in this configuration is only one less than the average.

The configuration of the sidechains relating to the data in Table 12.1 is depicted in Figure 12.3 (ab). We can see that these are the two closest possible mainchain locations for hydrogen bonding by the sidechain. But we also realize that other locations are also quite close, involving nearest sequence

| offset | residue | type | donor | acceptor | frequency | wrapping |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| +1 | HIS | M-S | N | ND1 | 38 | 12.0 |
| +1 | GLN | M-S | N | OE1 | 69 | 14.1 |
| +1 | THR | M-S | N | OG1 | 95 | 13.5 |
| +1 | SER | M-S | N | OG | 105 | 13.3 |
| +1 | GLU | M-S | N | OE1/2 | 138 | 14.9 |
| +1 | ASN | M-S | N | OD1 | 249 | 14.1 |
| +1 | ASP | M-S | N | OD1/2 | 627 | 15.0 |
| -1 | ASN | S-M | ND2 | O | 19 | 16.2 |
| -1 | CYS | S-M | SG | O | 38 | 14.8 |
| -1 | GLN | S-M | NE2 | O | 42 | 15.7 |
| -1 | LYS | S-M | NZ | O | 120 | 15.6 |
| -1 | ARG | S-M | Nx | O | 179 | 15.0 |
| -1 | ARG | S-M | NH2 | O | 42 | 15.3 |
| -1 | ARG | S-M | NE | O | 54 | 15.2 |
| -1 | ARG | S-M | NH1 | O | 83 | 14.7 |
| -1 | THR | S-M | OG1 | O | 236 | 14.7 |
| -1 | SER | S-M | OG | O | 286 | 14.7 |

Table 12.2: Observed local hydrogen bonds between the residue and the nearest mainchain amide or carbonyl groups of its sequence neighbors. The bonds made by arginine are further subdivided according to the specific hydrogen bond donor group. Nx refers to the collection of the three NH groups; the frequency is the sum of the frequencies and the wrapping is the average. See Figure 12.3 parts (c) and (d).
neighbors. These possible hydrogen bonds are depicted in Figure 12.3 (c-d). In Table 12.2, we list all of the observed local bonds that can occur where the sidechain bonds to the nearest position on the backbone of its sequence neighbor. The special bonds tabulated in Table 12.2 represent 1,321 M-S and 920 S-M hydrogen bonds, or about $4.3 \%$ of M-S and $2.4 \%$ of S-M hydrogen bonds. Thus the combined M-S bonds involving sidechain hydrogen bonds with the amide group on either the same peptide or the subsequent peptide constitute $11 \%$ of all M-S hydrogen bonds.

The desolvation domains for sequence-neighbor residues will also be slightly smaller in size. The mean separation of $C_{\alpha}$ 's in sequence neighbors is about $3.84 \AA$, and thus the desolvation domain (with radius $6 \AA$ ) is about $13 \%$ smaller in volume that a desolvation domain where the $C_{\alpha}$ 's are a typical $6 \AA$ apart. However, even with this correction, the amount of wrapping depicted is still significantly depressed from the expected averages (17.8 for M-S and 19.9 for S-M hydrogen bonds). Indeed, the volume of the desolvation domain depends on the distance between $C_{\alpha}$ atoms used in its definition, but taking a $6 \AA$ separation as typical, we see that when the separation varies from 5 $\AA$ to $7 \AA$, the corresponding volume variation is no more that $6 \%$ (for a desolvation radius of $6 \AA$ ), cf. Exercise 12.1 .

The next closest positions for a residue to make hydrogen bonds with the mainchain of its


Figure 12.4: Configuration of the (a) +1 offset sidechains for the sidechain-mainchain ( $\mathrm{S}-\mathrm{M}$ ) configuration, and the (b) -1 offset sidechains for the mainchain-sidechain (M-S) configuration. See Table 12.3
sequence neighbor are depicted in Figure 12.4. In Table 12.3, we list all of the observed local bonds that can occur where the sidechain bonds to the nearest position on the backbone of its sequence neighbor. The special bonds tabulated in Table 12.2 represent $64 \mathrm{M}-\mathrm{S}$ and $842 \mathrm{~S}-\mathrm{M}$ hydrogen bonds, or about $0.2 \%$ of M-S and $2.2 \%$ of S-M hydrogen bonds.

There are a few examples of bonds that occur only rarely in the dataset considered here. These are collected in Table 12.4

### 12.2.2 Further neighbors

By contrast, once we look beyond nearest sequence neighbors, the picture changes dramatically. There are $7,969 \mathrm{M}-\mathrm{S}$ hydrogen bonds between the sidechain of residue $k$ and the amide group on peptide $k+2$. This represents over a quarter of all M-S bonds in this dataset. Three fifths of them involved either Asp or Asn. However, the mean wrapping for these hydrogen bonds is still low: 14.2 for Asp and 14.9 for Asn. In Table 12.5, we compare the data on these configurations with the data on Asp and Asn that bond with closer neighbors. In these configurations, Glu and Gln are relatively less frequent, occurring only 153 and 48 times, respectively. Ser (1530) and Thr (1052) are more strongly represented in this group, while His (279) and Cys (117) are less other commonly occurring sidechains found in $k+2$ M-S bonds. The number of S-M bonds in this configuration is much smaller, having only 1180 occurrences. Similarly there are only 128 M-S, and 708 S-M, hydrogen bonds between the sidechain of residue $k$ and peptide $k-2$.

### 12.3 All sidechain hydrogen bonds

To amplify the assessment of attachments of sidechains to mainchain amides and carbonyls on peptides nearby in sequence, we now analyze sidechain hydrogen bonds with mainchains in general. In Table [12.6, we collect some pertinent statistics, again drawn from the same subset of 1547

| offset | residue | type | donor | acceptor | frequency | wrapping |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| +1 | CYS | S-M | SG | O | 21 | 14.0 |
| +1 | GLN | S-M | NE2 | O | 22 | 13.8 |
| +1 | ASN | S-M | ND2 | O | 48 | 13.6 |
| +1 | LYS | S-M | NZ | O | 62 | 13.7 |
| +1 | SER | S-M | OG | O | 177 | 11.1 |
| +1 | THR | S-M | OG1 | O | 217 | 13.2 |
| +1 | ARG | S-M | Nx | O | 295 | 16.3 |
| +1 | ARG | S-M | NH2 | O | 35 | 15.9 |
| +1 | ARG | S-M | NE | O | 45 | 15.9 |
| +1 | ARG | S-M | NH1 | O | 215 | 16.5 |
| -1 | ASN | M-S | N | OD1 | 6 | 11.0 |
| -1 | GLN | M-S | N | OE1 | 8 | 10.1 |
| -1 | GLU | M-S | N | OE1/2 | 50 | 11.6 |

Table 12.3: Observed local hydrogen bonds between the residue and the next nearest mainchain amide or carbonyl groups of its sequence neighbors. The bonds made by arginine are further subdivided according to the specific hydrogen bond donor group. Nx refers to the collection of the three NH groups; the frequency is the sum of the frequencies and the wrapping is the average. See Figure 12.4

| offset | residue | type | donor | acceptor | frequency | wrapping |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 0 | THR | M-S | N | OG1 | 1 | 13.0 |
| -1 | HIS | S-M | NE2 | O | 1 | 8.0 |
| -1 | TRP | S-M | NE1 | O | 1 | 12.0 |
| -1 | HIS | M-S | N | ND1 | 2 | 9.5 |
| -1 | THR | M-S | N | OG1 | 2 | 9.0 |
| -1 | ASP | M-S | N | OD1 | 3 | 15.7 |

Table 12.4: Rarely observed local hydrogen bonds between the residue and various nearby mainchain amide or carbonyl groups of its sequence neighbors.

| offset | residue | type | donor | acceptor | frequency | wrapping |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 0 | ASN | M-S | N | OD1 | 116 | $(14.4)$ |
| 0 | ASP | M-S | N | OD1 | 478 | $(14.7)$ |
| +1 | ASN | M-S | N | OD1 | 249 | 14.1 |
| +1 | ASP | M-S | N | OD1/2 | 627 | 15.0 |
| +2 | ASN | M-S | N | OD1 | 1613 | 14.9 |
| +2 | ASP | M-S | N | OD1 | 3175 | 14.2 |

Table 12.5: Comparison of wrapping for Asp and Asn M-S bonds in different contexts. Wrapping values for bonds to their own peptide are the corrected values, shown in parentheses, from Table 12.1.
proteins complexes from the PDB Select database.
Some explanations are required for the data in Table 12.6. The column 'type' indicates whether the sidechain forms a bond with an amide (M-S) or a carbonyl (S-M). That is, the first letter indicates the hydrogen bond donor and the second letter indicates the hydrogen bond acceptor. The column 'count' gives the total number of residues of this type in the PDB Select subset studied. The column 'bonds' indicates the number of hydrogen bonds of the specified within the total dataset.

The mode ' $M$ ' refers to the distribution of sequence distances between a sidechain and mainchain making a hydrogen bond. More precisely, we form the frequency distribution of sidechains with sequence number $i$ that are bonded to the mainchain amide or carbonyl of peptide $i+j$ as a function of $j$. These distributions are highly peaked (cf. Figure 12.7), and the mode provides a useful statistic to characterize them. The mode $M$ of this distribution of sequence distances is the number such that the largest number of sidechains $i$ makes bonds with peptide $i+M$. For example, $M=0$ means that the majority of the sidechains are bonded to their own mainchain.

The distribution of all sequence distances for all types of sidechains is shown in Figure 12.5 , We see that it is heavily concentrated on small distances, and Figure 12.6 provides a view of the distribution in this region. From this, we conclude that a substantial fraction of the distribution is concentrated for distances of magnitude ten or less, and that the character of the distribution changes outside this region. There is a significant difference in the distributions for M-S versus S-M bonds, and these differences are contrasted in Figure 12.6 as well.

The subsequent column (\%-loc) helps to characterize further the distribution of sequence distances between a sidechain and mainchain making a hydrogen bond. It gives the percentage of sidechains with sequence number $i$ that are bonded to the mainchain amide or carbonyl of peptide $i+M+j$ for $|j| \leq 2$. When this percentage is large, it indicates how peaked the distribution is around its mode $M$. The smaller percentages indicate distributions that are more spread out. Typical distributions are highly peaked around the mode, as indicated in Figure 12.7 for the M-S bonds for Thr, Ser and Asp. The distributions for the S-M bonds for Thr and Ser are similar but just shifted to the right by two units, corresponding to having $M=4$.

The column NNW gives the average wrapping for sidechains $i$ bonded to mainchains $i \pm 1$. If there are no such hydrogen bonds, then an average cannot be formed, and this is indicated in the table by NA. This special group is singled out due to the fact that the desolvation domain is about

| res. | type | count | bonds | $M$ | \%-loc | NNW | in-W | out-W | $\%$-out |
| :---: | :---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| ARG | S-M | 63565 | 8913 | 5 | 14.49 | 15.87 | 17.61 | 20.00 | 64.31 |
| ASN | M-S | 52997 | 4571 | 2 | 67.11 | 14.66 | 15.78 | 18.63 | 25.86 |
| ASN | S-M |  | 1688 | 4 | 19.98 | 14.37 | 17.90 | 19.08 | 58.41 |
| ASP | M-S | 68906 | 10185 | 2 | 61.04 | 14.91 | 15.44 | 18.09 | 25.51 |
| CYS | M-S | 29731 | 285 | 2 | 66.20 | NA | 14.83 | 15.97 | 24.91 |
| CYS | S-M |  | 1052 | 4 | 69.36 | 14.43 | 21.26 | 20.24 | 16.83 |
| GLN | M-S | 49222 | 1745 | -3 | 28.27 | 13.53 | 17.69 | 20.26 | 42.92 |
| GLN | S-M |  | 1365 | 2 | 31.16 | 15.05 | 17.47 | 20.12 | 47.91 |
| GLU | M-S | 84047 | 4555 | 0 | 31.40 | 14.07 | 16.70 | 19.89 | 36.55 |
| HIS | M-S | 27377 | 784 | 2 | 77.14 | 11.90 | 16.88 | 19.23 | 16.20 |
| HIS | S-M |  | 763 | 3 | 8.14 | 8.00 | 19.24 | 19.77 | 79.03 |
| LYS | S-M | 85292 | 4191 | 3 | 21.26 | 14.96 | 18.13 | 20.17 | 61.66 |
| SER | M-S | 71354 | 4496 | 2 | 69.12 | 13.42 | 15.45 | 17.94 | 21.57 |
| SER | S-M |  | 8298 | 4 | 59.46 | 13.31 | 16.60 | 18.45 | 24.57 |
| THR | M-S | 65455 | 3272 | 2 | 69.58 | 13.48 | 15.68 | 19.08 | 22.68 |
| THR | S-M |  | 9127 | 4 | 64.09 | 14.14 | 17.61 | 19.11 | 21.96 |
| TRP | S-M | 16203 | 1119 | 5 | 15.30 | 12.00 | 25.08 | 26.98 | 70.78 |
| TYR | M-S | 39743 | 747 | -5 | 13.14 | NA | 24.72 | 26.78 | 74.97 |
| TYR | S-M |  | 1761 | 4 | 10.34 | NA | 25.19 | 27.38 | 80.01 |

Table 12.6: Key: 'type' of bond (see text); 'count' is the number of residues of this type in the PDB Select subset; 'bonds' is the number of hydrogen bonds of the specified type involving this residue; $M$ is the mode of the distribution of sequence distances between a sidechain and mainchain making a hydrogen bond (the largest number of sidechains $i$ are bonded to the mainchain amide or carbonyl of peptide $i+M)$; \%-loc= percentage of hydrogen bonds with sequence distances of the form $M \pm i$ for $|i| \leq 2$; NNW=average wrapping for the nearest neighbors (sidechain $i$ bonded to the mainchain amide or carbonyl of peptide $i \pm 1$; NA indicates that there are no such bonds); in- $\mathrm{W}=$ average wrapping for sidechain $i$ bonded to the mainchain amide or carbonyl of peptide $i+j$ for $1<|j| \leq 10$; out-W=average wrapping for sidechain $i$ bonded to the mainchain amide or carbonyl of peptide $i+j$ for $|j|>10 ; \%$-out=percent of sidechains $i$ bonded to the mainchain amide or carbonyl of peptide $i+j$ for $|j|>10$;


Figure 12.5: Distribution of sequence distances between donors and acceptors in hydrogen bonds between mainchains and sidechains. Both M-S and S-M bonds are included.


Figure 12.6: Distribution of sequence distances between donors and acceptors in hydrogen bonds between mainchains and sidechains. Solid line: S-M bonds; dashed line: M-S bonds.


Figure 12.7: Distribution of sequence distances for M-S hydrogen bonds for Thr (solid line), Ser (dashed line) and Asp (dotted line).
$13 \%$ smaller than is typical. This was done to avoid contamination of the assessment 'in-W' of the wrapping of hydrogen bonds between sidechains with sequence number $i$ that are bonded to the mainchain amide or carbonyl of peptide $i+j$ for $1<|j| \leq 10$. Self-bonding $(i=j)$ was also eliminated in computing the 'in-W' statistics (these data are found in Table 12.1).

In most cases, the ratio of in-W to NNW is about what would be expected due to the slight difference in sizes of the desolvation domains. In the cases where there is a significant difference, the set of sidechains $i$ bonded to mainchains $i \pm 1$ is quite small. The column 'out-W' lists the average wrapping of hydrogen bonds between sidechains with sequence number $i$ that are bonded to the mainchain amide or carbonyl of peptide $i+j$ for $|j|>10$. These are the hydrogen bonds without any bias due to locality in sequence. The distribution of sequence distances extends into the hundreds in each direction. The percentage of such hydrogen bonds is indicated by $\%$-out. In some cases, this is the minority of bonds, but the percentages are still large enough in the important cases to give a good estimate of the average wrapping for non-local sidechain-mainchain bonds of the particular types.

What is most striking about the data in Table 12.6 is that the local average wrapping, as indicated by NNW and in-W, is significantly less than the non-local average wrapping, as indicated by out-W. That is, the indicated sidechain-mainchain hydrogen bonds are far more likely to occur with nearby sidechain-mainchain pairs when there is a local wrapping deficit. These sidechains tend to correct the wrapping defect by forming hydrogen bonds with the mainchain.

A significant fraction of certain residues are devoted to these local bonds formed in underwrapped environments. For example, $19 \%$ of threonine residues form sidechain bonds with the mainchain, as do $18 \%$ of the serines and $15 \%$ of aspartates. The majority of these bonds are made with near sequence neighbors and are deficiently wrapped.

Draft: February 28, 2008, do not distribute


Figure 12.8: Distribution of sequence distances between donors and acceptors in arginine S-M bonds.

The distribution of distances for arginine is somewhat of an exception from the others. It is still quite localized, as indicated in Figure [12.8. However, it is spread more broadly over -10 to +10 , rather than being concentrated in a region $\pm 2$ around its mode $(M=5)$.

### 12.4 Unusual hydrogen bonds

### 12.4.1 Hydrophobic pairs

When Val is paired with Val in a backbone-backbone hydrogen bond it is frequently the case that there is a pair of bonds. For example in 1CU4, there are both H-VAL133N-H-VAL180O and H-VAL180N-H-VAL133O bonds. This type of pair is also found in 1CU4 and 1E4W (133,180 and 194,203), 1F90 (138,193 and 210,219), 1IGC (143,190 and 204,213), 1JRH (210,219), 1MPA (144, 191 and 205,214 ), 1WEJ ( 140,187 and 201,210 ), 2CII $(199,249)$, and 2H1P ( 443,490 and 504,513 ).

A similar Met pair is in 1MPA $(20,81)$, and a His pair is found in $1 \mathrm{G} 6 \mathrm{~V}(94,119)$. The Ile pair $(34,51)$ appears in 1 CU 4 . Tyr pairs appear frequently: the pair $(142,172)$ appears in both 1CU4 and 1 E 4 W , the pair $(7,99)$ appears in $2 \mathrm{BSR}, 2 \mathrm{BSS}, 2 \mathrm{BVO}$, and 2 BVQ , and the pair $(36,87)$ is in 1JRH. Phe pairs are also common: $(208,241)$ in 2 BSR , $2 \mathrm{BSS}, 2 \mathrm{BVO}$, and 2BVQ; $(30,62)$ in 2 BSR and 2CII; and $(36,87)$ in 1 E 4 W .

### 12.4.2 Unusual trios

The trio of arginines $(75,79,83)$ appears in 2BSR, 2 BSS , and 2 BVO . These are involved in a sequence of mainchain hydrogen bonds: A-ARG 83 N - A-ARG 79 O , A-ARG 79 N - A-ARG 75 O. A

Draft: February 28, 2008, do not distribute 150
similar trio of lysines $(9,55,103)$ is in 1 IGC. Such a trio of residues forms an unusual charged structure. A similar type of trio also occurs with Phe: H-PHE32 N - H-PHE29 O , H-PHE29 N - H-PHE27 O is found in 1IGC.

### 12.5 Exercises

Exercise 12.1 Let $V_{\rho, R}$ be the volume of the desolvation domain when the desolvation radius is $\rho$ and the separation between the $C_{\alpha}$ is $R$. When $R \geq 2 \rho, V_{\rho, R}=2 V_{\rho, 0}$. Compute the change in volume in the desolvation domain as a function of the separation of the $C_{\alpha}$ atoms used in its definition. Define a separation parameter $r=R / 2 \rho$, so that $r=0$ is the case when the desolvation spheres coincide, and $r=1$ is the point at which they become completely separated. Verify that the ratio of volumes $\phi(r)=V_{\rho, R} / V_{\rho, 2 \rho}$ may be written as

$$
\begin{equation*}
\phi(r)=\frac{1}{2}+\frac{3}{4}\left(r-\frac{1}{3} r^{3}\right) \tag{12.1}
\end{equation*}
$$

and is in particular independent of $\rho$.
Exercise 12.2 Repeat the analysis in Section 12.1 but count the potential frequency of bond formation by including the relative frequencies of each amino acid based on the data in Table 6.2.

Exercise 12.3 Determine if wrapping correlates (or not) with the formation of mainchain-mainchain hydrogen bonds. Consider the average wrapping of carbonyls and amides that do not form hydrogen bonds and compare this with what is found in Figure 12.1. Note that there is a significant number of amide and carbonyl groups that are well wrapped and yet form no hydrogen bonds; these are the peptides in group Ib in Section 14.3 .

## Chapter 13

## Aromatic interactions

Aromatic sidechains are special for many reasons. Their large size allows for multifaceted roles. On the one hand, the benzene rings in them are largely hydrophobic, and they have very small dipole moments. However, the charge distribution of these rings creates a significant quadrupole moment (Section 9.4).

Partial charges are frequently used to model aromatics, as shown in Table 13.1. However, this model is planar and only approximates an important aspect of aromatic polarity. The center of a surface above each side of the face of the aromatics is negatively charged, and the C-H groups in the ring form positive charge centers [55]. In Figure 13.4 we plot the electrostatic potential corresponding to the partial charges in Table 13.1 in a plane parallel to the plane of the ring but at a distance of $1 \AA$ from the plane of the ring. We do see that the face of the ring has a negative charge, as required.

However, the planar partial charges cause a large polar behavior in the plane of the ring near the locations of the hydrogens. In Figure [13.4, we plot the electrostatic potential corresponding to the partial charges in Table 13.1 in a plane near the plane of the ring (at a distance from the plane of the ring of only $0.1 \AA$ ). In this plane, there is a strong polarity, one that might lead to hydrophilic behavior.

A less crude approximation of the aromatic ring would be to put negative charges at positions near the carbons but in the direction normal to the ring. This improvement would be similar to current models of water, such as Tip5P [160].

### 13.1 Cation- $\pi$ interactions

Among pair interactions at interfaces (Section 6.3), the Arg-Trp interaction has the fourth highest log-odds ratio. This pair is an example of what is known as a cation- $\pi$ pair [95, 258, 48]. It has a strength comparable to that of a hydrogen bond. It is based on an interaction between the negative charge on the face of aromatic residues and positively charged (cation) residues (Lys, Arg, His). The cation- $\pi$ motifs play a special role in protein interfaces [258, 48]. The cation- $\pi$ interaction also has a significant role in $\alpha$-helix stabilization [220].

Cation- $\pi$ interactions can take place with other resides, such as a phosphorylated tyrosine. And

| Residue | atom type | PDB code | charge |
| :--- | :---: | :---: | :---: |
| PHE | C | $\mathrm{CD} i, \mathrm{CE} i, i=1,2, \mathrm{CZ}$ | -0.1 |
|  | HC | $\mathrm{HD} i, \mathrm{HE} i, i=1,2, \mathrm{HZ}$ | 0.1 |
| TYR | C | $\mathrm{CD} i, \mathrm{CE} i, i=1,2$ | -0.1 |
|  | HC | $\mathrm{HD} i, \mathrm{HE} i, i=1,2$ | 0.1 |
|  | C | CZ | 0.15 |
|  | OA | OH | -0.548 |
|  | H | HH | 0.398 |
| TRP | C | CG | -0.14 |
|  | C | $\mathrm{CD} 1, \mathrm{CE} 3, \mathrm{CZ} i, i=2,3, \mathrm{CH} 2$ | -0.1 |
|  | HC | $\mathrm{HD} 1, \mathrm{HE} 3, \mathrm{HZ} i i=2,3, \mathrm{HH} 2$ | 0.1 |
|  | NR | NE 1 | -0.05 |
|  | H | HE 1 | 0.19 |

Table 13.1: Partial charges from the Gromos force field for aromatic amino acids.
the two-sided nature of the polarity of an aromatic ring means that it can interact with two cations at one time, one on each side. This is depicted in Figure 13.1 which shows a phosphorylated tyrosine flanked by an arginine and a lysine in the SH2 domain in the PDB file 1JYR. SH2 domains [150] specifically recognize phosphorylated tyrosines and bind proteins containing them, and the cation- $\pi$ interaction is presumably crucial in the binding process.

Correspondingly, a cation could be sandwiched between two aromatics, as shown in Figure 13.2 , The arginine (A67) interacts with both a phosphorylated tyrosine (PTR-I3) and a phenylalanine (Phe-I4). In addition, Arg-A67 is hydrogen bonded with the backbone oxygen on Phe-I4 and one of the terminal oxygens on PTR-I3.

Finally, we show a cation- $\pi$ grouping involving a complex of two cations and two aromatics, as shown in Figure 13.3. The arginine is interacting with both of the aromatics, whereas the lysine is interacting only with the phosphorylated tyrosine.

### 13.2 Aromatic-polar interactions

Aromatic rings can in principle interact with any polar moiety. In [123], a hydrogen bond with the polar face of an aromatic ring is described (in Figure 6).

### 13.3 Aromatic-aromatic interactions

It is possible for two aromatics to interact via more complex interactions. In Figure 13.6 the relative orientations of two phenylalanines is shown [5] as found in the PDB file 1TLA. A vector along the line formed by the $C_{\gamma}-C_{\zeta}$ carbons in Phe117 (upper) is pointing toward the face of Phe153.

Draft: February 28, 2008, do not distribute


Figure 13.1: Cation- $\pi$ group found in the SH2 domain in the PDB file 1JYR. Shown are a phosphorylated tyrosine (PTR1003) flanked by Arg67 (upper left) and Lys109 (lower right).


Figure 13.2: Cation- $\pi$ group found in the SH2 domain in the PDB file 1BMB. Shown are a phosphorylated tyrosine (PTR-I3, lower right) and a phenylalanine (Phe-I4, upper right) flanking an Arg-A67 (middle left).


Figure 13.3: Cation- $\pi$ group found in the SH2 domain in the PDB file 1TZE. Shown are a phosphorylated tyrosine (PTR-I4, middle left) and a phenylalanine (Phe-I3, upper right) together with Arg-E67 (lower right) and Lys-E109 (upper left).


Figure 13.4: Electric potential of the partial-charge model of a benzene ring in a parallel plane, 0.1 Ångstrom above the ring, using the partial charges in Table 13.1 .


Figure 13.5: Electric potential of the partial-charge model of a benzene ring in a parallel plane, one Angstrom above the ring, using the partial charges in Table 13.1 .


Figure 13.6: Pairwise interactions of two phenylalanines (117 and 153) in a mutant (Ser117 $\rightarrow$ Phe) of T4 lysozyme detailed in the PDB file 1TLA.

The interaction depicted in Figure 13.6 is consistent with a model of Phe in which there is a small positive partial charge, or positive polarity, at the end of the sidechain, that is near the $C_{\zeta}$ carbon. The partial charge model in Table 13.1 is precisely of that type. There is a dipole formed between $C_{\zeta}$ and $H_{\zeta}$, with no opposing dipole related to $C_{\gamma}$. The other dipoles in the ring are always counter-balanced, forming quadrupoles. Thus there is a net dipolar behavior to the carbonaceous ring of Phe in the partial charge model in Table 13.1, consistent with the Phe-Phe interaction in Figure 13.6

### 13.4 Exercises

Exercise 13.1 Investigate the interactions of two phenylalanine residues as depicted in Figure 13.6. Determine the distribution of distances from the terminal carbon of one of the residues to the face of the other. Also record the angle between the two faces.

## Chapter 14

## Peptide bond rotation

We now consider an application of data mining that links the quantum scale with the continuum level electrostatic field. In other cases, we have considered the modulation of dielectric properties and the resulting effect on electronic fields. Here we study the effect of the local electronic field on a particular covalent bond. Wrapping by hydrophobic groups plays a role in our analysis, but we are interested primarily in a secondary effect of the interaction. We consider modulations in the electronic field that cause a significant change in the electron density of the peptide bond and lead to a structural change in the principle bond.

We first describe the results and then consider their implications for protein folding.

### 14.1 Peptide resonance states

The peptide bond is characterized in part by the planarity of the six atoms shown in Figure 14.1 , The angle $\omega$ (see Section 4.4.1) quantifies this planarity. The bond is what is known as a resonance [191] between two states, shown in Figure 14.1. The "keto" state (A) on the left side of Figure 14.1 is actually the preferred state in the absence of external influences [191]. However, an external polarizing field can shift the preference to the "enol" state (B) on the right side of Figure [14.1, as we illustrate in Figure 14.2 .

The resonant state could be influenced by an external field in different ways, but the primary cause is hydrogen bonding. If there is hydrogen bonding to the carbonyl or amide groups in a peptide bond, this induces a significant dipole which forces the peptide bond into the (B) state shown in Figure 14.1. Such hydrogen bonds could be either with water, with sidechains, or with other backbone donor or acceptor groups, and such a configuration is indicated in Figure 14.2, On the right side of this figure, the polar environment is indicated by an arrow from the negative charge of the oxygen partner of the amide group in the peptide bond to the positive charge of the hydrogen partner of the carbonyl group. The strength of this polar environment will be less if only one of the charges is available, but either one (or both) can cause the polar field.

However, if neither type of hydrogen bond is available, then the resonant state moves toward the preferred (A) state in Figure 14.1. The latter state involves only a single bond and allows $\omega$ rotation. Thus the electronic environment of peptides determines whether they are rigid or flexible.

(B)


Figure 14.1: Two resonance states of the peptide bond. The double bond between the central carbon and nitrogen keeps the peptide bond planar in the right state (B). In the left state (A), the single bond can rotate.


Figure 14.2: The resonance state $(\mathrm{B})$ of the peptide bond shown, on the left, with hydrogen bonds which induce a polar field to reduce the preference for state (A). On the right, the (B) state is depicted with an abstraction of the dipole electrical gradient induced by hydrogen bonding indicated by an arrow.

Since any hydrogen bond can enable the (B) state, it will prevail whenever a water molecule or a mainchain or sidechain donor or acceptor is appropriately located. Otherwise said, the (A) state persists only when water is removed and there are no other binding partners.

### 14.2 Measuring variations in $\omega$

There is no simple way to measure the flexibility of the peptide bond from typical structural data. If flexibility were linked with mobility of the $\omega$ bond, then it could potentially be inferred from the 'fuzziness' of the electron densities, e.g., as measured by the B-factors reported in the PDB. But such observations might be attributed to other factors, and there is no reason to believe that flexibility in $\omega$ would necessarily mean that the angle adopted would not be fixed and well resolved. For any given peptide bond, the value of $\omega$ could correspond to the rigid (B) state even if it is fully
in the (A) state. The particular value of $\omega$ depends not only on the flexibility but also on the local forces that are being applied. These could in principle be determined, but it would be complicated to do so. However, by looking at a set of peptide bonds, we would expect to see a range of values of $\omega$ corresponding to a range of local forces. It is reasonable to assume that these local forces would be randomly distributed in some way if the set is large enough. Thus, the dispersion $\Delta_{\omega}$ for a set of peptide bond states can give a clear signal as to the flexibility or rigidity of the set of peptide bonds.

The assessment of the flexibility of the peptide bond requires a model. Suppose that we assume that the bond adopts a configuration that can be approximated as a fraction $C_{B}$ of the rigid (B) state and corresponding the fraction $C_{A}$ of the (A) state. Let us assume that the flexibility of the peptide bond is proportional to $C_{A}$. That is, if we imagine that the peptide bond is subjected to random forces, then the dispersion $\Delta_{\omega}$ in $\omega$ is proportional to $C_{A}$ :

$$
\begin{equation*}
\Delta_{\omega}=\gamma C_{A} \tag{14.1}
\end{equation*}
$$

where $\gamma$ is a constant of proportionality.
We are assuming that the resonant state $\psi$ has the form $\psi=C_{A} \psi_{A}+C_{B} \psi_{B}$ where $\psi_{X}$ is the eigenfunction (see Chapter (16) for state $X(=A$ or $B)$. Since these are eigenstates normalized to have $L^{2}$ norm equal to one, we must have $C_{A}^{2}+C_{B}^{2}=1$, assuming that they are orthogonal. Therefore

$$
\begin{equation*}
\Delta_{\omega}=\gamma C_{A}=\gamma \sqrt{1-C_{B}^{2}} \tag{14.2}
\end{equation*}
$$

We can thus invert this relationship to provide $C_{B}$ as a function of $\Delta_{\omega}$ :

$$
\begin{equation*}
C_{B}=\sqrt{1-\left(\Delta_{\omega} / \gamma\right)^{2}} \tag{14.3}
\end{equation*}
$$

The assumption (14.1) can be viewed as follows. The flexibility of the peptide bond depends on the degree to which the central covalent bond between carbon and nitrogen is a single bond. The (A) state is a pure single bond state and the $(B)$ state is a pure double bond state. Since the resonance state is a linear combination, $\psi=C_{A} \psi_{A}+C_{B} \psi_{B}$, it follows that there is a linear relationship between $C_{A}$ and flexibility provided that the single bond state can be quantified as a linear functional. To prove this relationship, we seek a functional $L_{A}$ such that $L_{A} \psi_{A}=1$ and $L_{A} \psi_{B}=0$. If $\psi_{A}$ and $\psi_{B}$ are orthogonal, this is easy to do. We simply let $L_{A}$ be defined by taking inner-products with $\psi_{A}$, provided (as we assume) that $\psi_{A}$ and $\psi_{B}$ are orthogonal.

### 14.3 Predicting the electric field

Since the preference for state (A) or (B) is determined by the local electronic environment, the easiest way to study the flexibility would be to correlate it with the gradient of the external electric field at the center of the peptide bond. However, this field is difficult to compute precisely due to the need to represent the dielectric effect of the solvent (Chapter 17) and to account for the polarizability of all the molecular groups. A dynamic simulation with an explicit water model and full representation of polarizability might be able to correctly estimate this accurately, but this

Draft: February 28, 2008, do not distribute
is beyond current technological capability. However, it is possible to estimate the likelihood of a significant dipole moment based on the local environment 66.

The major contributor to a local dipole would be the hydrogen bonding indicated in Figure 14.2 , These bonds can arise in two ways, either by backbone hydrogen bonding (or perhaps backbonesidechain bonding, which is more rare) or by contact with water. The presence of backbone hydrogen bonds is indicated by the PDB structure, but the presence of water is not consistently represented. However, there is a a proxy for the probability of contact with water: the wrapping of the local environment. So we can approximate the expected local electric field by analyzing the backbone hydrogen bonding and the wrapping of these amide and carbonyl groups.

In [66], sets of peptide bonds were classified in two ways. First of all, they were separated into two groups, as follows. Group I consisted of peptides forming no backbone hydrogen bonds, that is, ones not involved in either $\alpha$-helices or $\beta$-sheets. Group II consisted of peptides forming at least one backbone hydrogen bond. In each major group, subsets were defined based on the level $\rho$ of wrapping in the vicinity of backbone.

Figure 14.3(a) depicts the resulting observations for group I peptide bonds, using the model (14.3) to convert observed dispersion $\Delta_{\omega}(\rho)$ to values of $C_{B}$, with a constant $\gamma=22^{\circ}$. This value of $\gamma$ fits well with the estimated value $C_{B}=0.4$ [191] for the vacuum state of the peptide bond 191] which is approached as $\rho$ increases. Well wrapped peptide bonds that do not form hydrogen bonds should closely resemble the vacuum state. However, poorly wrapped peptide amide and carbonyl groups would be strongly solvated, and thus strongly polarized, leading to a larger component of (B) as we expect and as Figure 14.3(a) shows.

Using this value of $\gamma$ allows an interesting assessment of the group II peptide bonds, as shown in Figure 14.3 (b). These are bonds that, according to the PDB structures, are capable of participating in backbone hydrogen bonds. We see that these bonds also have a variable resonance structure depending on the amount of wrapping. Poorly wrapped backbone hydrogen bonds will likely be solvated, and thus the group I and group II peptides can be expected to behave similarly for small $\rho$. As with group I peptide bonds, we expect state (B) to be dominant for small $\rho$. Indeed, the two curves in Figure 14.3 (a) and (b) are quite similar for small $\rho$. As dehydration by wrapping improves, the polarity of the environment due to waters decreases, and the proportion of state (B) decreases. But a limit occurs in this case, unlike with group I, due to the fact that wrapping now enhances the strength of the backbone hydrogen bonds, and thus increases the polarity of the environment.

The interplay between the decreasing strength of polarization due to one kind of hydrogen bonding (with water) and the increasing strength of backbone hydrogen bonding is quite striking. As water is removed, hydrogen bonds strengthen and increase polarization of peptide bond. Figure 14.3(b) shows that there is a middle ground in which a little wrapping is not such a good thing. That is, small amounts of wrapping appear to remove enough water to decrease the polar environment. Moreover, with minimal wrapping, the backbone hydrogen bonds are screened, and therefore the resulting external polar environment of the peptide is weaker. But the effect of the hydrogen bonds increase as wrapping is increased.

An alternate way of viewing the data is as follows. We consider four groups:

- Group Ia consists of peptides forming no backbone hydrogen bonds and the amide and carbonyl groups are not well wrapped,

Draft: February 28, 2008, do not distribute


Figure 14.3: From [66]: Fraction of the double-bond (planar) state in the resonance for residues in two different classes (a) Neither amide nor carbonyl group is engaged in a backbone hydrogen bond (group I). As water is removed, so is polarization of peptide bond. (b) At least one of the amide or carbonyl groups is engaged in backbone hydrogen bond (group II).

- Group Ib consists of well wrapped peptides forming no backbone hydrogen bonds,
- Group IIa consists of peptides capable of forming a backbone hydrogen bond but not well wrapped, and
- Group IIb consists of well wrapped peptides forming a backbone hydrogen bond.

The left side of Figure 14.3 corresponds to Group Ia and Group IIa (panels a and b, respectively), and we see that the behavior is the same for the two groups. That is, the underwrapped peptides have a similar dispersion $\Delta_{\omega}$, corresponding to a dominant (B) state, whether or not they appear to be capable of hydrogen bonding. On the other hand, there is a difference between the Ib and IIb groups, as indicated on the right side of Figure 14.3 (panels a and b, respectively). Group Ib prefers the vacuum state (A), whereas IIb tends to the (B) state.

Figure 14.3 presents an even more refined analysis. It involves groups $\mathrm{I} \rho$ and $\mathrm{II} \rho$ for different values of $\rho$.

### 14.4 Implications for protein folding

After the "hydrophobic collapse" [261] a protein is compact enough to exclude most water. At this stage, few hydrogen bonds have fully formed. But most amide and carbonyl groups are protected from water. The data in Figure 14.3(a) therefore implies that many peptide bonds are flexible in final stage of protein folding. This effect is not included in current models of protein folding. This effect buffers the entropic cost of hydrophobic collapse in the process of protein folding.

New models need to allow flexible bonds whose strengths depend on the local electronic environment. Typical molecular dynamics (MD) models would either have peptide bonds fixed in the planar configuration or have a large spring constant for rotation in the $\omega$ angle. Here we need the spring constant to depend on the gradient of the electric field in the vicinity of the peptide bond.

The gradient of the electric field at a point $\mathbf{r}$ is given by

$$
\begin{equation*}
\sum_{k} q_{k} \frac{\mathbf{r}-\rho_{k}}{\left|\mathbf{r}-\rho_{k}\right|^{3}} \tag{14.4}
\end{equation*}
$$

cf. (17.9). In particular, the quantity of interest is the strength of the dot product of the electric field gradient and the vector $\overline{O-H}$ pointing from O to H in the peptide group. If $\mathbf{r}_{0}$ is the centroid of the peptide group, then one would seek a bending strength depending on the quantity

$$
\begin{equation*}
\sum_{k} q_{k} \frac{\left(\mathbf{r}_{0}-\rho_{k}\right) \cdot \overline{O-H}}{\left|\mathbf{r}_{0}-\rho_{k}\right|^{3}} . \tag{14.5}
\end{equation*}
$$

Note that we are invoking a sum over all (charged) atoms in the system, and this type of global term will make the simulation much more costly. Using a cut-off radius to limit the number of charged atoms involved may be practical, but it introduces an approximation into the model whose effect would have to be assessed.

Draft: February 28, 2008, do not distribute

### 14.5 Exercises

Exercise 14.1 The dipole vector is twice as strong if both amide and carbonyl groups are involved in hydrogen bonds. Split group II into two groups, group II1 and II2 depending on the number of hydrogen bonds. How does the preference for the (B) state differ between groups II1 and II2?

Exercise 14.2 Scan some PDB files and form the groups Ia, Ib, IIa, IIb indicated in Section 14.3. Plot the distributions of $\omega$ angles for each group. Do the distributions for Ia and IIa look similar? How are the distributions for $I b$ and IIb different?

Exercise 14.3 Using a model of the dielectric effect (cf. Chapter 17), estimate the dipole vector at each peptide in a set of PDB files. Use this estimate to predict whether the peptide bond is in the $(A)$ state or the (B) state. Compare this with the measured value of $\omega$.

Exercise 14.4 Using a molecular dynamics model with explicit water, estimate the dipole vector at each peptide in a set of PDB files. Use this estimate to predict whether the peptide bond is in the $(A)$ state or the (B) state. Compare this with the measured value of $\omega$.

Exercise 14.5 Using a quantum chemistry model, calculate the flexibility of the $\omega$ bond as a function of an imposed dipole as indicated in Figure 14.2.

## Chapter 15

## Tree representations of data

The process of evolution is naturally expressed via a tree. For this reason, tree representations of data are widely used in biology. We have used a tree representation of relationships among different myoglobin molecules in various species in Figure 2.1. In that representation, the distances along the tree are intended to represent the evolutionary distance, as measured by sequence similarity, between different versions of myoglobin. This sequence distance was used to contrast differences in the numbers of dehydrons in the different versions of myoglobin.

We explain here some of the basic issues with tree representations, mainly that they rarely represent the data exactly. Moreover, different algorithms are commonly used, and they do not give the same trees in many cases. Although we are not able to explain the differences that may arise in general, we do give one hint of how this may be tolerated in practice. We also give various examples of interest as illustration.

### 15.1 Distance metrics

The concept of distance is formalized mathematically in a metric space. Finite metric spaces are quite simple. They are characterized by having objects that can be labelled by integers $i=1, \ldots, n$, and corresponding distances between the points given by a disatnce matrix $\mathcal{D}(i, j)$ where the individual values of the entries of the matrix are the distances in the space between objects $i$ and $j$.

Several of the key properties of distance matrices are both easy to verify by inspection and easy to motivate. By definition, the diagonal of $\mathcal{D}$ is always zero, since the distance from one point to itself should be zero. Also, since they are distances, all of the entries are positive. Similarly, the matrix is assumed to be symmetric, reflecting the presumption that the cost of going from A to $B$ is the same as going from $B$ to $A$. In some cases, this might not be true, so a different kind of mathematics would need to be used.

The most important condition on a distance matrix is the triangle inequality. This condition encapsulates an important geometric condition and is not easily verified by inspection.

### 15.1.1 Triangle inequality

The triangle inequality for a distance matrix $\mathcal{D}$ requires that

$$
\begin{equation*}
\mathcal{D}(i, j) \leq \mathcal{D}(i, k)+\mathcal{D}(k, j) \tag{15.1}
\end{equation*}
$$

for all $i, j, k=1, \ldots, n$. For example, if there are only three objects in the metric space (three proteins, say), then the general form of a distance matrix is

$$
\left(\begin{array}{lll}
0 & a & b  \tag{15.2}\\
a & 0 & c \\
b & c & 0
\end{array}\right)
$$

where $a, b$, and $c$ are positive parameters. Suppose that, without loss of generality, that $c$ is the smallest:

$$
\begin{equation*}
c \leq \min \{a, b\} \tag{15.3}
\end{equation*}
$$

(If not, re-label the names of the elements of the metric space.) Then we leave as Exercise 15.2 that the triangle inequality implies that

$$
\begin{equation*}
|b-a| \leq c \tag{15.4}
\end{equation*}
$$

We leave as Exercise 15.3 to show that any coefficients $a, b$, and $c$ satisfying the bounds (15.3) and (15.4) generate a distance matrix of the form (15.2) satisfying the triangle inequality. Thus the set of possible values for $a, b$, and $c$ occupy a non-convex cone (with non-empty interior) in three dimensions defined by the bounds (15.3) and (15.4) (cf. Exercise 15.4).

### 15.1.2 Non-metric measurements

In many cases in biology, there are values $\widetilde{\mathcal{D}}(i, j)$ that represent relationships between objects $i$ and $j$ that are all positive but do not necessarily satisfy the triangle inequality (15.1). Thus they do not faithfully represent the 'cost' or 'distance' in going from $i$ to $j$. For example, if $\widetilde{\mathcal{D}}(i, j)>\widetilde{\mathcal{D}}(i, k)+\widetilde{\mathcal{D}}(k, j)$, then the path

$$
\begin{equation*}
i \rightarrow k \rightarrow j \tag{15.5}
\end{equation*}
$$

represents a cheaper way (with cost $\widetilde{\mathcal{D}}(i, k)+\widetilde{\mathcal{D}}(k, j)$ ) to get from $i$ to $j$, rather than going directly. Thus $\widetilde{\mathcal{D}}$ no longer satisfies our general notion of a collection of direct distances. However, it is possible to define a metric related to $\widetilde{\mathcal{D}}$ that represents the relations faithfully, as follows.

Using the metaphor of distance, we can think of progressing from $i$ to $j$ via a finite sequence of steps like (15.5), i.e.,

$$
\begin{equation*}
i=k_{0} \rightarrow k_{1} \rightarrow \cdots \rightarrow k_{r}=j \tag{15.6}
\end{equation*}
$$

Then we define $\mathcal{D}(i, j)$ as the minimum cost over all possible paths of the type (15.6), viz.,

$$
\begin{equation*}
\mathcal{D}(i, j)=\min _{i=k_{0}, k_{1}, \ldots, k_{r}=j} \sum_{\ell=1}^{r} \widetilde{\mathcal{D}}\left(k_{\ell-1}, k_{\ell}\right) . \tag{15.7}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

Without loss of generality, we can assume that all paths satisfy $k_{\ell-1} \neq k_{\ell}$, since if they are equal there is no contribution to the sum in (15.7), and the corresponding paths with the repetitions of terms eliminated lead to the same value.

Lemma 15.1 Suppose that the matrix $\widetilde{\mathcal{D}}$ is symmetric, has all off-diagonal entries positive and all diagonal entries zero. Then the matrix $\mathcal{D}$ defined by 15.7) is a metric (in particular, satisfies the triangle inequality) and

$$
\begin{equation*}
\mathcal{D}(i, j) \leq \widetilde{\mathcal{D}}(i, j) \tag{15.8}
\end{equation*}
$$

for all $i$ and $j$.
Proof. Since one of the possible paths is $i=k_{0}, k_{1}=j$, we have $\mathcal{D}(i, j) \leq \widetilde{\mathcal{D}}(i, j)$ for all $i$ and $j$, which proves (15.8).

Let $\epsilon=\min \{\widetilde{\mathcal{D}}(i, j): i, j=1, \ldots n\}$ and $\mu=\max \{\widetilde{\mathcal{D}}(i, j): i, j=1, \ldots n\}$. Then

$$
\begin{equation*}
\sum_{\ell=1}^{r} \widetilde{\mathcal{D}}\left(k_{\ell-1}, k_{\ell}\right) \geq \epsilon r \tag{15.9}
\end{equation*}
$$

In view of (15.8), we can restrict to paths such that

$$
\begin{equation*}
\sum_{\ell=1}^{r} \widetilde{\mathcal{D}}\left(k_{\ell-1}, k_{\ell}\right) \leq \mu \tag{15.10}
\end{equation*}
$$

because paths with a larger sum will not change the value of the minimum in (15.7). So we can assume that $r \leq \mu / \epsilon$. Thus the number of paths can be assumed to be finite and the minimum in (15.7) is thus positive for all $i$ and $j$. Moreover, we can assume that the minimum is attained by a particular path in (15.7):

$$
\begin{equation*}
\mathcal{D}(i, j)=\sum_{\ell=1}^{r} \widetilde{\mathcal{D}}\left(k_{\ell-1}, k_{\ell}\right) \tag{15.11}
\end{equation*}
$$

for some path of the form (15.6).
The matrix $\mathcal{D}$ defined by (15.7) is symmetric because any path from $i$ to $j$ given via steps $i=k_{0}, k_{1}, \ldots, k_{r}=j$ gives a path from $j$ to $i$ given by $j=k_{r}, k_{r-1}, \ldots, k_{1}=i$, and conversely. And the inequality (15.8) guarantees that $\mathcal{D}$ is zero on the diagonal.

Now let us confirm that the triangle inequality (15.1) holds for the matrix $\mathcal{D}$ defined by (15.7). Let $i, j$, and $m$ be arbitrary. Let $i=k_{0}, k_{1}, \ldots, k_{r}=m$ be a path such that

$$
\begin{equation*}
\mathcal{D}(i, m)=\sum_{\ell=1}^{r} \widetilde{\mathcal{D}}\left(k_{\ell-1}, k_{\ell}\right) . \tag{15.12}
\end{equation*}
$$

Similarly, let $m=k_{0}^{\prime}, k_{1}^{\prime}, \ldots, k_{r^{\prime}}^{\prime}=j$ be a path such that

$$
\begin{equation*}
\mathcal{D}(m, j)=\sum_{\ell=1}^{r^{\prime}} \widetilde{\mathcal{D}}\left(k_{\ell-1}^{\prime}, k_{\ell}^{\prime}\right) \tag{15.13}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

Using the path $i=k_{0}, k_{1}, \ldots, k_{r}=m=k_{0}^{\prime}, k_{1}^{\prime}, \ldots, k_{r^{\prime}}^{\prime}=j$ as a possible path in (15.7) proves that

$$
\begin{equation*}
\mathcal{D}(i, j) \leq \sum_{\ell=1}^{r} \widetilde{\mathcal{D}}\left(k_{\ell-1}, k_{\ell}\right)+\sum_{\ell=1}^{r^{\prime}} \widetilde{\mathcal{D}}\left(k_{\ell-1}^{\prime}, k_{\ell}^{\prime}\right)=\mathcal{D}(i, m)+\mathcal{D}(m, j) \tag{15.14}
\end{equation*}
$$

## QED

### 15.2 Sequence distance

The simplest form of sequence comparison is based on editing strings and counting the number of edits required to get from one sequence to the other. The formulation of string-edit distance $d_{e}$ balances two different types of edits. The simplest is replacement of a single letter by another letter. To start with, we need a metric on the set of letters in the alphabet $\Sigma$ for our sets of sequences. Let $D_{\Sigma}$ be a metric on the alphabet $\Sigma$. Then $D_{\Sigma}(\xi, \eta)$ measures the difference between the two letters $\xi$ and $\eta$ in $\Sigma$. Now we show how to extend this to a metric $d_{e}$ on the set of all finite strings $\Sigma^{*}$.

If two strings $x$ and $y$ differ only in the $k$-th position, then we set $d_{e}(x, y)=D_{\Sigma}\left(x_{k}, y_{k}\right)$. In general, when there are multiple replacements, string edit distance is based on just summing the effects. However, string-edit distance also allows a different kind of change as well: insertion and deletion. For example, we can define $x_{\hat{k}}$ to mean the string $x$ with the $k$-th entry removed. It might be that $x_{\hat{k}}$ agrees perfectly with the string $y$, and so we assign $d(x, y)=\delta$ where $\delta$ is the deletion penalty. Similarly, insertions of characters are allowed to determine edit distance. Clearly, if $y=x_{\hat{k}}$, then adding $x_{k}$ to $y$ at the $k$-th position yields $x$. Again, the effect of multiple insertions/deletions is additive, and this allows strings of different lengths to be compared.

The use of both replacements and instertion/deletions to determine edit distance means that an edit path from $x$ to $y$ is not unique. Edit distance is therefore defined by taking the minimum over all possible representations, as we define formally in (15.18). But this will not in general define a metric unless appropriate conditions on $\delta$ and $D_{\Sigma}$ are satisfied. These conditions can be defined by extending the alphabet $\Sigma$ and metric $D_{\Sigma}$ to include a "gap" as a character, say "." (let $\widetilde{\Sigma}$ denote the extended alphabet), and by assigning a distance $D_{\widetilde{\Sigma}}\left(x,,_{-}\right)$for each character $x$ in the original alphabet.

Theorem 9.4 of [246] says that $d_{e}$ is a metric on strings of letters in $\Sigma$ whenever $D_{\widetilde{\Sigma}}$ is a metric on the extended alphabet.

### 15.2.1 Two-letter alphabets

The simplest non-trivial example is an alphabet with two letters, say $x$ and $y$, when there is only one distance $D_{\Sigma}(x, y)$ that is non-zero. The requirement that the triangle inequality hold for $D_{\widetilde{\Sigma}}$ reduces to three inequalities that can be expressed as

$$
\begin{equation*}
\left|D_{\widetilde{\Sigma}}(x,-)-D_{\widetilde{\Sigma}}(y,-)\right| \leq D_{\Sigma}(x, y) \leq D_{\widetilde{\Sigma}}(x,-)+D_{\widetilde{\Sigma}}(y,-) \tag{15.15}
\end{equation*}
$$

Together with the condition that all distances be non-negative, we see that (15.15) characterizes completely the requirement for $D_{\widetilde{\Sigma}}$ to be a metric in the case of a two-letter alphabet $\Sigma$.

Draft: February 28, 2008, do not distribute

### 15.2.2 General alphabets

For a general alphabet $\Sigma$, if

$$
\begin{equation*}
\alpha \leq D_{\Sigma}(x, y) \leq 2 \alpha \tag{15.16}
\end{equation*}
$$

for all $x \neq y$ (including _) for some $\alpha>0$, then $D_{\Sigma}$ is a metric (that is, the triangle inequality holds). This is because

$$
\begin{equation*}
D_{\Sigma}(x, y) \leq 2 \alpha \leq D_{\Sigma}(x, z)+D_{\Sigma}(z, y) \tag{15.17}
\end{equation*}
$$

for any $z \in \Sigma$. One simple choice for a metric on letters is to choose $D_{\Sigma}(x, y)=1$ for all $x \neq y$, and then to take $D_{\widetilde{\Sigma}}(x,-)=2$; the resulting $D_{\widetilde{\Sigma}}$ satisfies (15.16) for $\widetilde{\Sigma}$. However, condition (15.16) is far from optimal as the example (15.15) shows.

The edit distance $d_{e}$ is derived from the extended alphabet distance $D_{\widetilde{\Sigma}}$ as follows. We introduce the notion of alignment $\mathcal{A}$ of sequences $\left(x^{*}, y^{*}\right)=\mathcal{A}(x, y)$ where $x *$ has the letters of $x$ in the same order but possibly with gaps _ inserted, and similarly for $y^{*}$. We suppose that $x *$ and $y^{*}$ have the same length even if $x$ and $y$ did not, which can always be achieved by adding gaps at one end or the other. Then

$$
\begin{equation*}
d_{e}(x, y)=\min _{\mathcal{A}} \sum_{i} D_{\widetilde{\Sigma}}\left(x_{i}^{*}, y_{i}^{*}\right) \tag{15.18}
\end{equation*}
$$

for any $z \in \Sigma$. The minimum is over all alignments $\mathcal{A}$ and the sum extends over the length of the sequences. Fortunately, string-edit distance $d_{e}$, and even more complex metrics involving more complex gap penalties, can be computed efficiently by the dynamic programming algorithm [246].

The simple string-edit distance $d_{e}$ described here is useful in many contexts. However, more complex metrics would be required in other applications.

### 15.2.3 Distance versus score

Typically biologists prefer to work with a "score" that is large when two sequences are close as opposed to a distance which is small in such a case. The dynamic programming algorithm can equivalently be used to mininmize the distance or maximize a score. There is a formal correspondence that can be made between scores and distances, as follows [246].

### 15.3 Feature distance

Another way to define metrics for biological entities is by comparing feature differences. Different features can indicate a propensity for protein-protein interaction, such as surface curvature, wrapping (dehydrons), 'hot spots,' etc. These features can be compared on different proteins to see if they are similar or not. Different features may not be conserved across homologs, and thus a distance metric can encapsulate such differences.

One feature that can be compared is the hydrogen bond location for homologous proteins. This can be done in a variety of ways, but one way is to first form a standard structural alignment (http://www.ncbi.nlm.nih.gov/structure/CN3D/cn3d.shtml) and then to trim the protein sequences by restricting to those residues that have been structurally aligned in a one-to-one correspondence.

Draft: February 28, 2008, do not distribute

|  | Ack | Kit | Abl | Lck | Chk1 | Pdk1 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| EGFR | 2 | 2 | 3 | 3 | 4 | 4 |
| Ack |  | 1 | 4 | 4 | 5 | 5 |
| Kit |  |  | 4 | 4 | 5 | 5 |
| Abl |  |  |  | 1 | 4 | 4 |
| Lck |  |  |  |  | 4 | 4 |
| Chk1 |  |  |  |  |  | 1 |

Table 15.1: Dehydron distances for seven homologous tyrosine kinase proteins. The PDB files corrsponding to the abbreviations in the table are: Abl (1FPU), Ack1 (1U54), Chk1 (1IA8), C-kit (1T45), EGFR (1M17), Lck (3LCK), Pdk1 (1UVR).

An indicator matrix for each protein is constructed that is indexed by all pairs of residues. It consists only of zeros unless two residues are paired by a hydrogen bond, in which case the corresponding entry is one. Then, a Hamming-type distance [75] can be defined based on the number of disagreements between two such indicator matrices. More precisely, a matrix $H_{i j}$ is constructed by choosing $H_{i j}=1$ if residues $i$ and $j$ are paired by a hydrogen bond and $H_{i j}=0$, otherwise. Then, a Hamming-type distance [75]

$$
\begin{equation*}
\mathcal{D}(m, n)=\sum_{i<j}\left|H_{i j}(m)-H_{i j}(n)\right| \tag{15.19}
\end{equation*}
$$

is used, where $m$ and $n$ are the indices of different proteins.
Metrics based on comparison of hydrogen bond structures can be refined by limiting to certain classes of hydrogen bonds. A very successful use of this idea is to restrict to the underwrapped hydrogen bonds (the dehydrons) 81. Since this metric reflects defects in the packing of the hydrogen bonds by hydrophobic groups, it is called the 'packing distance.' The packing distance for seven homolgous tyrosine kinase proteins is depicted in Table 15.1

Another example of restricting to special hydrogen bonds is depicted in Table 15.2 Here the distance is based on the intermolecular hydrogen bonds formed between the antibody and the antigen in homologous antibody-antigen structures.

A distance based on nonpolar regions on proteins (a.k.a., hydrophobic patches) can similarly be defined [41]. A pharmacological distance matrix can also be constructed based on affinity profiling against specified drugs [75, 62].

Many other features could potentially be used to define useful metrics. For example, geometric [19] features such as surface curvature can be measured by

$$
\begin{equation*}
\mathcal{D}(m, n)=\sum_{i}\left|K_{i}(m)-K_{i}(n)\right| \tag{15.20}
\end{equation*}
$$

where $K_{i}$ is the curvature of the protein-ligand interface at the $i$-th position of proteins $m, n$.
Such distance matrices can be used to construct dendrograms (or trees) and are implicitly behind the dendrograms commonly seen [62, 88]. Such trees are easily constructed by using standard

Draft: February 28, 2008, do not distribute


Figure 15.1: Packing similarity tree (PST, bottom in black) for the seven structurally aligned paralogs of Bcr-Abl, based on the dehydron distances in Table 15.1. The PST restricted to the alignments of the Gleevec wrapped region in Bcr-Abl is shown (top) with blue dashed lines. The paralogs in red have the most similar packing in the region that aligns with the Gleevec wrapped region in $\mathrm{Bcr}-\mathrm{Abl}$ and are also primary targets of this inhibitor 62.
algorithms for phylogenetic analysis [102]. Distance in the tree reflects distance as encoded in the distance matrix, with the proviso that certain restrictions must apply to the distance matrix to get an exact representation [102. For example, with dehydrons, we refer to this tree as a 'packing' similarity tree (PST). The resulting tree gives a visual indication of 'closeness' for the various proteins, cf. Figure 15.1 which depicts two such trees. The PST allows the assessment of possible effects of targeting given features in drug design to determine potential specificity. Nearby proteins (Abl and Lck, for example) have similar features in our simple example.

Although visual inspection and comparison of trees is useful, the lack of uniqueness of trees is a cause for concern. A direct comparison of distance matrices provides a more rigorous comparison of measured properties, such as a comparison of pharmacological distance and packing distance [75]. In general, if a distance matrix is to be represented faithfully by a tree, it must satisfy a 'four-point condition' [102] that includes the familiar triangle inequality but is substantially more restrictive. This requirement implies that typical biological trees will not uniquely represent a given biological distance matrix. Thus direct comparison of distance matrices is a more reliable technique [75].

### 15.4 Tree representation of metrics

A tree naturally defines a metric space for the leaves of the tree, where the distance between leaves is the shortest path in the tree. We now invert this observation and ask which metrics can be represented by trees.

### 15.4.1 Three point metrics

A distance matrix $\mathcal{D}(i, j)$ for three points can always be represented by a tree. There are only three unique values in the distance matrix: $\mathcal{D}(1,2), \mathcal{D}(2,3)$ and $\mathcal{D}(1,3)$. A tree that connects the points 1,2 , and 3 via a central node (call it 0 ) has only three distances to define it: the distance $\delta_{i}$ from 0 to node $i$ for $i=1,2,3$. The relatinship between the two representations of distance is $\mathcal{D}(i, j)=\delta_{i}+\delta_{j}$. It is easy to see that this $3 \times 3$ system of equations is invertible.

However, in general a distance matrix $\mathcal{D}(i, j)$ for $k>3$ points cannot be exactly represented by the distances in a tree. We will see this explicitly in the case $k=4$, but it is easy to see why there is a difficulty in this case. The tree of interest will take the form in Figure 15.3, There are only five internal distances available in this tree. But there are six different distinct values in a distance matrix $\mathcal{D}(i, j)$ for $k=4$. Even if these six values are chosen to satisfy the triangle inequality, there is one too many of them to be able to be matched by five parameters.

Different algorithms are used in practice, and they have the property that given a distance matrix $\mathcal{D}(i, j)$ they will produce an answer. It is not known in general what the relattionship is between the products of different popular algorithms. However, for $k=4$ we can at least give an interpretation of the extent of ambiguity.

### 15.4.2 Four point condition

In general, if a distance matrix is to be represented faithfully by a tree, it must satisfy the following four-point condition 102

$$
\begin{equation*}
\mathcal{D}(i, j)+\mathcal{D}(m, n) \leq \max \{\mathcal{D}(i, m)+\mathcal{D}(j, n), \mathcal{D}(j, m)+\mathcal{D}(i, n)\} \tag{15.21}
\end{equation*}
$$

for all $i, j, m$, and $n$. The four-point condition generalizes the triangle inequality (take $m=n$ ). For a distance matrix satisfying the triangle inequality, it suffices to take $i, j, m$, and $n$ distinct. We will see that generically most distance matrices do not satisfy the four-point condition. This implies that typical biological trees will not rigorously represent a given biological distance matrix.

Definition 15.1 A matrix that satisfies the four point condition is called additive.
The following theorem may be found in [102].
Theorem 15.1 A distance matrix can be represented by distances in a tree if and only if it satisfies the four point condition (i.e., it is additive).

There is a simple interpretation of the four point condition, as follows. Given distinct values of $i, j, m$, and $n$, define parameters $A, B$, and $C$ by

$$
\begin{equation*}
A=\mathcal{D}(i, j)+\mathcal{D}(m, n), \quad B=\mathcal{D}(i, m)+\mathcal{D}(j, n), \quad \text { and } \quad C=\mathcal{D}(j, m)+\mathcal{D}(i, n) \tag{15.22}
\end{equation*}
$$

When we have three values $A, B$, and $C$, it is natural to expect that there are three distinct values given by

$$
\begin{equation*}
\min \{A, B, C\} \leq \operatorname{mid}\{A, B, C\} \leq \max \{A, B, C\} \tag{15.23}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

## CHAPTER 15. TREE REPRESENTATIONS GF4DAПRAEE REPRESENTATION OF METRICS



Figure 15.2: Original tree (left) and reduced tree (right) where nodes $i$ and $j$ have been removed and replaced by node $k$. The corresponding distances are defined via (15.26).

Here, the 'mid' function picks out the value between the min and the max. The four-point condition is the requirement that

$$
\begin{equation*}
\operatorname{mid}\{A, B, C\}=\max \{A, B, C\} \tag{15.24}
\end{equation*}
$$

Otherwise said, of the three values $A, B$, and $C$, the two largest must be equal for the four-point condition to hold.

As an example, consider the intermolecular hydrogen bond distance data presented in Table 15.2 , For a metric space with only four points, there is only one choice for the distinct values of $i, j, m$, and $n$, although relabelling is possible. For the data in Table 15.2, we find that

$$
\begin{equation*}
\{A, B, C\}=\{8,12,12\} \tag{15.25}
\end{equation*}
$$

so Table 15.2 is an additive matrix.
It is possible to motivate the four-point condition by considering a simple algorithm for reducing the size of the metric space. This algorithm is only suggestive, but it does make clear why the condition arises, and it also motivates one of the widely used algorithms for determining trees from data.

### 15.4.3 A reduction algorithm

Consider the following algorithm for constructing a tree from a distance matrix. Loop over all pairs $i, j$. Suppose there is a graph rerpresentation with nodes $i$ and $j$ appearing as leaves of a parent node as depicted on the left-hand side of Figure 15.2. Call the parent node $k$, where $k$ is an index not being used in the current indexing scheme. Then define

$$
\begin{equation*}
\mathcal{D}(m, k)=\mathcal{D}(k, m)=\frac{1}{2}(\mathcal{D}(i, m)+\mathcal{D}(j, m)-\mathcal{D}(i, j)) \tag{15.26}
\end{equation*}
$$

Create a new discrete space by eliminating $i$ and $j$ and adding $k$; in terms of the distance matrix, we eliminate the $i$ and $j$ rows and add the new information defined by (15.26). By the triangle inequality, this new matrix is non-negative. Moreover, if we find $\mathcal{D}(k, m)=0$ we can take $k=m$ and avoid the addition to the discrete space, so we can assume that this new matrix is non-degenerate.

This new matrix satisfies the triangle inequality, that is $\mathcal{D}(k, m) \leq \mathcal{D}(k, n)+\mathcal{D}(n, m)$ for all $n$. This is equivalent to

$$
\begin{equation*}
(\mathcal{D}(i, m)+\mathcal{D}(j, m)-\mathcal{D}(i, j)) \leq(\mathcal{D}(i, n)+\mathcal{D}(j, n)-\mathcal{D}(i, j))+2 \mathcal{D}(n, m) \tag{15.27}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
or equivalently

$$
\begin{equation*}
\mathcal{D}(i, m)+\mathcal{D}(j, m) \leq \mathcal{D}(i, n)+\mathcal{D}(j, n)+2 \mathcal{D}(n, m) \tag{15.28}
\end{equation*}
$$

But the triangle inequality for the original matrix implies this: we just add

$$
\begin{equation*}
\mathcal{D}(i, m) \leq \mathcal{D}(i, n)+\mathcal{D}(n, m) \tag{15.29}
\end{equation*}
$$

to

$$
\begin{equation*}
\mathcal{D}(j, m) \leq \mathcal{D}(j, n)+\mathcal{D}(n, m) \tag{15.30}
\end{equation*}
$$

to prove (15.28).
The reduction to a smaller metric space just described motivates the popular UPGMA algorithm. The algorithm proceeds by clustering nodes $i$ and $j$ for which $\mathcal{D}(i, j)$ is smallest. The heuristic is that smaller $\mathcal{D}(i, j)$ values should mean that $i$ and $j$ are closer in the tree (which is correct) and (which is not correct) the closest points must be children of the same parent node in the tree. We give an example to show how this fails in Section 15.5

### 15.4.4 Obstruction to reduction

The difficulty arises in the assignment of the distances between the new point and the deleted points. If all were well, we would have

$$
\begin{equation*}
\mathcal{D}(i, k)=\mathcal{D}(i, m)-\mathcal{D}(m, k)=\frac{1}{2}(\mathcal{D}(i, m)-\mathcal{D}(j, m)+\mathcal{D}(i, j)) \tag{15.31}
\end{equation*}
$$

for any $m$, and similarly for $\mathcal{D}(j, k)$ :

$$
\begin{equation*}
\mathcal{D}(j, k)=\mathcal{D}(j, m)-\mathcal{D}(m, k)=\frac{1}{2}(\mathcal{D}(j, m)-\mathcal{D}(i, m)+\mathcal{D}(i, j)) \tag{15.32}
\end{equation*}
$$

Since $m$ is arbitrary in (15.31), it must hold as well for any other node $n$ replacing $m$. Since the left-hand side of (15.31) remains unchanged, we must have

$$
\begin{equation*}
\mathcal{D}(i, m)-\mathcal{D}(j, m)+\mathcal{D}(i, j)=\mathcal{D}(i, n)-\mathcal{D}(j, n)+\mathcal{D}(i, j) \tag{15.33}
\end{equation*}
$$

for any $m$ and $n$, which is the same as saying

$$
\begin{equation*}
\mathcal{D}(i, m)+\mathcal{D}(j, n)=\mathcal{D}(i, n)+\mathcal{D}(j, m) \tag{15.34}
\end{equation*}
$$

which we will see is equivalent to the ' $\mathrm{mid}=\mathrm{max}$ ' interpretation of the four-point condition.
To complete the derivation of the four-point condition from (15.34), we note that the common value in (15.34) may be written as (see Exercise 15.8)

$$
\begin{equation*}
\mathcal{D}(i, m)+\mathcal{D}(j, n)=\mathcal{D}(i, n)+\mathcal{D}(j, m)=\mathcal{D}(i, j)+\mathcal{D}(k, m)+\mathcal{D}(k, n) \tag{15.35}
\end{equation*}
$$

By the triangle inequality,

$$
\begin{equation*}
\mathcal{D}(i, j)+\mathcal{D}(m, n) \leq \mathcal{D}(i, j)+\mathcal{D}(m, k)+\mathcal{D}(k, n) \tag{15.36}
\end{equation*}
$$

Combining (15.36) and (15.35) completes the derivation of the four-point condition. Moreover, it proves that $\mathcal{D}(i, j)+\mathcal{D}(m, n)$ has to be the 'min' value of these ABC values, for all $m$ and $n$. This appears at first to be stronger than the four point condition, so we state it formally.

Draft: February 28, 2008, do not distribute

|  | 1DQJ | 1C08 | 1NDG |
| :---: | :---: | :---: | :---: |
| 1NDM | 3 | 7 | 6 |
| 1DQJ |  | 6 | 5 |
| 1C08 |  |  | 5 |

Table 15.2: Distance matrix for intermolecular hydrogen bond interactions between antibody and antigen in the PDB complexes: $a=1 \mathrm{NDM}, \mathrm{b}=1 \mathrm{DQJ}, \mathrm{c}=1 \mathrm{C} 08, \mathrm{~d}=1 \mathrm{NDG}$. Zeros on the diagonal and the redundant lower triangular part of the matrix have been omitted.

Lemma 15.2 Suppose that the matrix $\mathcal{D}$ can be represented as a tree. Then there are indices $i$ and $j$ such that

$$
\begin{equation*}
\mathcal{D}(i, j)+\mathcal{D}(m, n) \leq \min \{\mathcal{D}(i, m)+\mathcal{D}(j, n), \mathcal{D}(i, n)+\mathcal{D}(j, m)\} \tag{15.37}
\end{equation*}
$$

for all $m$ and $n$.

The problem of identifying indices $i$ and $j$ such that the reduction algorithm can be applied is complex. Although discovered some time ago [208], the condition has received a great deal of attention [32, 96] and continues to be of interest [59, 61, [168]. The required condition [32] is to minimize the expression

$$
\begin{align*}
Q_{i j} & =(n-2) \mathcal{D}(i, j)-\sum_{k=1}^{n} \mathcal{D}(i, k)-\sum_{k=1}^{n} \mathcal{D}(j, k)  \tag{15.38}\\
& =(n-4) \mathcal{D}(i, j)-\sum_{k \neq i, j ; k=1}^{n} \mathcal{D}(i, k)+\mathcal{D}(j, k) .
\end{align*}
$$

The leaves $i$ and $j$ identified in this way not only allow the reduction algorithm to construct a tree for an additive metric, they also are good choices in the case that a metric is not additive [32, 59, 61, 96, 168].

The algorithm we have described for constructing a tree by recursively reducing the size of the metric space is called neighbor joining. The neighbors being joined are $i$ and $j$. The choice given by minimizing (15.38) is widely used, but not universal. The UPGMA algorithm uses the far simpler heuristic of minimizing $\mathcal{D}(i, j)$. We explain why this approach leads to the wrong answer in Section 15.5 even for an additive metric.

Although we will see that there are many interesting biological metrics that are additive, in general this is not the case. Thus it is of interest to understand to what extent we can approximate a general distance matrix via an additive metric [20]. Neighbor joining, especially using the criterion of minimizing (15.38) [32, 59, 61, 96, 168]. gives an effective approximation. In many cases, what is of most interest is not the lengths of the edges in the tree, but rather just the topology of the tree. We now consider one simple case in which we can understand this approximation problem in some detail.

Draft: February 28, 2008, do not distribute


Figure 15.3: Tree representation of the distance matrix in Table 15.2

### 15.4.5 The ABC theorem

It is possible to show that the topology of the tree representation is essentially unique under very mild conditions, as follows. Consider the three independent quantities that figure in the four point condition:

$$
\begin{align*}
& A=\mathcal{D}(i, m)+\mathcal{D}(j, n) \\
& B=\mathcal{D}(i, n)+\mathcal{D}(j, m)  \tag{15.39}\\
& C=\mathcal{D}(i, j)+\mathcal{D}(n, m)
\end{align*}
$$

based on the three ways to partition the index set $\{i, j, m, n\}$ into distinct pairs. These quantities determine the topology of the tree representations, as follows. There are four distinct cases. Three of them involve two internal nodes and one internal edge, and are categorized by the following three distinct possibilities for additive matrices: $A=B>C, B=C>A$, and $C=A>B$. The fourth tree corresponds to $A=B=C$. Note that when $A=B=C$, the tree representing the distance matrix is a star. That is, there is one internal node $k$, and four edges joining the four indices to $k$.

We will show that even in the case that a matrix is not additive, a unique assignment of one of these topology classes is possible in most cases.

Suppose $\mathcal{D}$ is a general distance matrix that is not necessarily additive. Without loss of generality, by renaming the indices if necessary, we can assume that the terms are ordered:

$$
\begin{equation*}
A \geq B \geq C \tag{15.40}
\end{equation*}
$$

The four-point condition can now be stated simply: $A=B$. In this case, the distance matrix can be represented exactly by a tree. Now we consider the other case, that $A>B$. First, we define the $\ell^{1}$-norm for distance matrices:

$$
\begin{equation*}
\|\mathcal{D}\|_{\ell^{1}}=\sum_{i<j}|\mathcal{D}(i, j)| \tag{15.41}
\end{equation*}
$$

Note that we allow for negative entries, as we intend to apply the norm to differences of distance matrices.

Theorem 15.2 Suppose that $A>B$. Then

$$
\begin{equation*}
\inf \left\{\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}: \mathcal{D}^{\prime} \text { satisfies the four-point condition }\right\}=A-B \tag{15.42}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

## CHAPTER 15. TREE REPRESENTATIONS GF4DADREEE REPRESENTATION OF METRICS

Moreover, if $B>C$, then all additive distance matrices $\mathcal{D}^{\prime}$ which satisfy

$$
\begin{equation*}
\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}=A-B \tag{15.43}
\end{equation*}
$$

have trees with the same topology.
When $B=C$, there is an ambiguity in representing $\mathcal{D}$ since there are additive matrices $\mathcal{D}^{\prime}$ all equally close in $\ell^{1}$ norm with different topology types. We leave as an exercise that there is a matrix $\mathcal{D}^{\prime}$ with $A^{\prime}=B^{\prime}=C^{\prime}=B=C$, as well as two others: one with $A^{\prime}=B^{\prime}=A$ and $C^{\prime}=B=C$ and the other with with $A^{\prime}=C^{\prime}=A$ and $B^{\prime}=B=C$.
Proof. To prove these assertions, we first show that

$$
\begin{equation*}
\inf \left\{\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}: \mathcal{D}^{\prime} \text { satisfies four-point condition }\right\} \leq A-B \tag{15.44}
\end{equation*}
$$

To so so, we simply need to exhibit a $\mathcal{D}^{\prime}$ which satisfies the four-point condition and $\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}=$ $A-B$. We can do this if we keep $A^{\prime}=A$ and increase $B^{\prime}$ to be equal to $A$. For example, we can set

$$
\begin{equation*}
\mathcal{D}_{i n}^{\prime}=\mathcal{D}_{i n}+A-B, \tag{15.45}
\end{equation*}
$$

leaving all other entries of $\mathcal{D}^{\prime}$ the same as for $\mathcal{D}$. Thus by explicit construction, we have $\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}=$ $A-B$. Similarly, since we also have $A^{\prime}=A=B^{\prime}, \mathcal{D}^{\prime}$ satisfies the four point condition.

Now we demonstrate the other inequality. We can clearly write

$$
\begin{equation*}
\left|A-A^{\prime}\right|+\left|B-B^{\prime}\right|+\left|C-C^{\prime}\right| \leq\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}} \tag{15.46}
\end{equation*}
$$

for any distance matrices. Now suppose it were the case that for some any $\mathcal{D}^{\prime}$ we have $\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}<$ $A-B$. Then by (15.46) we have

$$
\begin{align*}
B^{\prime}-A^{\prime}+A-B & =A-A^{\prime}+B^{\prime}-B \\
& \leq\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}<A-B \tag{15.47}
\end{align*}
$$

from which we conclude that $B^{\prime}<A^{\prime}$, so $\mathcal{D}^{\prime}$ cannot satisfy the four point condition. This completes the proof of the equality (15.42).

Now suppose that $\mathcal{D}^{\prime}$ is additive and satisfies (15.43). Then we want to show that

$$
\begin{equation*}
A \geq A^{\prime}=B^{\prime} \geq B \tag{15.48}
\end{equation*}
$$

Suppose that $A<B^{\prime}$. Then $B^{\prime}-B>A-B$ and applying (15.46) we find that $\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}>A-B$. On the other hand, if $A^{\prime}<B$ then $A-A^{\prime}>A-B$, and again (15.46) yields a contradiction. Thus (15.48) has to hold.

Applying (15.48) in (15.46), we get

$$
\begin{align*}
A-B & =A-A^{\prime}+B^{\prime}-B \\
& =\left|A-A^{\prime}\right|+\left|B-B^{\prime}\right| \\
& \leq\left|A-A^{\prime}\right|+\left|B-B^{\prime}\right|+\left|C-C^{\prime}\right|  \tag{15.49}\\
& \leq\left\|\mathcal{D}-\mathcal{D}^{\prime}\right\|_{\ell^{1}}=A-B
\end{align*}
$$

Draft: February 28, 2008, do not distribute

|  | b | c | d |
| :--- | :--- | :--- | :--- |
| a | 3 | 5 | 4 |
| b |  | 4 | 5 |
| c |  |  | 7 |

Table 15.3: Additive distance matrix for which UPGMA gives the wrong tree.


Figure 15.4: UPGMA tree (left) and additive tree for distance matrix in Table 15.3
which means that equality holds throughout the experession (15.49), so we must have $C=C^{\prime}$. In particular, we conclude that $A^{\prime}=B^{\prime} \geq C^{\prime}$. If $B>C$, then we also have $B^{\prime}>C^{\prime}$.

Now it is easy to show that all additive matrices with $A^{\prime}=B^{\prime}>C^{\prime}$ have the same topology. QED

### 15.5 What UPGMA does

The UPGMA algorithm [102 applied to a distance matrix will coallesce the closest points, that is, the two for which the entry in the distance matrix is smallests, say $\mathcal{D}(i, j)$. One might hope that UPGMA would find the correct tree for an additive matrix.

The nearest neighbors in the tree for an additive matrix are the indices that combine to form the term $C$ that is the smallest of the three terms involved in the four point condition: $C<B=$ $A$. However, even though it may hold that $\mathcal{D}(i, j)$ is the smallest entry in the distance matrix, $C=\mathcal{D}(i, j)+\mathcal{D}(m, n)$ is not smaller than $A$ or $B$. In this case, UPGMA finds the wrong tree. For example, consider the distance matrix in Table 15.3 with $A=5+5=10, B=4+4=8$, and $C=3+7=10$. In Figure 15.4 we see both the tree that UPGMA will generate (left) for the data together with the tree (right) that precisely represents this additive matrix.

### 15.6 Exercises

Exercise 15.1 Show that the distance matrices in Table 15.1, Table 15.2, and Table 15.3 satisfy the triangle inequality.

Exercise 15.2 Prove that any distance matrix (15.2) must satisfy $|b-a| \leq c$ provided that $c \leq$ $\min \{a, b\}$. (Hint: apply the triangle inequality (15.1).)

Exercise 15.3 Suppose that a distance matrix (15.2) satisfies (15.3) and (15.4). Prove that it satisfies the triangle inequality (15.1). (Hint: reverse the derivation in Exercise 15.4.)

Exercise 15.4 Suppose that a distance matrix (15.2) satisfies $c \leq \min \{a, b\}$. Sketch the cone of values of $a, b$, and $c$ that satisfy the triangle inequality (15.1). (Hint: apply Exercise 15.2)

Exercise 15.5 Determine the set of the (six) allowable values for a distance matrix for a fourelement metric space (cf. Exercise 15.4).

Exercise 15.6 Does that the matrix

$$
\left(\begin{array}{lll}
0 & 4 & 2  \tag{15.50}\\
4 & 0 & 1 \\
2 & 1 & 0
\end{array}\right)
$$

satisfy the triangle inequality (15.1)? (Hint: see Exercise 15.2.)
Exercise 15.7 PAM and Blossum matrices attempt to encode evolutionary distance between sequence elements. Examine these to see if they satisfy the triangle inequality (15.1).

Exercise 15.8 Prove (15.35).

## Chapter 16

## Quantum models

Computational models for approximating the quantum mechanics of chemical systems 45] represent one of the most computationally intense areas of computational science. There has been recent interest in computational methods for quantum chemistry that involve algorithms with improved computational complexity [216, 230]. Recent books [37, 38] present the subject from a more mathematical point of view. Here we review of the major ideas as a guide to understanding the basis for molecular force fields.

Our objective is to clarify the notions of polarizability and induced fields. We present a novel derivation of the asymptotic behavior of the London dispersion force which is based on the induced dipole. We confirm the well known $R^{-6}$ dependence of the force due to an induced dipole, but our techniques do not rely on the standard eigenvalue (second-order) perturbation approach [193]. Instead, our approach performs a perturbation of the wavefunction and identifies a boundary value problem for the asymptotic form of the induced dipole, which incidentally provides a way to compute the coefficient of $R^{-6}$ for any two systems.

### 16.1 Why quantum models?

One might ask why we include a discussion of quantum models in a discussion of protein interactions. Molecular-level models utilize force fields that can be determined from quantum models, and this is an area where we can predict significant developments in the future. The hydration structure around certain amino acid residues is complex and something that begs further study [232, 187, 119, [187, 105, [259]. But this may require water models which are currently considered quite expensive, and these models may require further examination at the quantum level.

### 16.2 The Schrödinger equation

The Schrödinger equation is taken as the fundamental model for the molecular systems considered here. It is a time-dependent partial differential equation for the wave function $\psi$ for the system as a function of the positions in space of the nuclei and electrons in a molecule.

For one particle of mass $m$ moving in a force field having potential $V$, it can be written

$$
\begin{equation*}
\iota \hbar \frac{\partial \psi}{\partial t}=\mathcal{H} \psi \tag{16.1}
\end{equation*}
$$

where $\hbar$ is Planck's constant divided by $2 \pi, \iota$ is the imaginary unit and

$$
\begin{equation*}
\mathcal{H} \phi(\mathbf{r}):=-\frac{\hbar^{2}}{2 m} \nabla^{2} \phi(\mathbf{r})+V(\mathbf{r}) \phi(\mathbf{r}) \tag{16.2}
\end{equation*}
$$

For a system of $k$ particles, each of mass $m_{i}$, the equation (16.1) remains valid with

$$
\begin{equation*}
\mathcal{H}_{k} \phi\left(r_{1}, \ldots, r_{k}\right):=-\sum_{i=1}^{k} \frac{\hbar^{2}}{2 m_{i}} \nabla_{i}^{2} \phi\left(r_{1}, \ldots, r_{k}\right)+V_{k}\left(r_{1}, \ldots, r_{k}\right) \phi\left(r_{1}, \ldots, r_{k}\right) \tag{16.3}
\end{equation*}
$$

and $\nabla_{i}$ denotes the gradient with respect to $r_{i}$.
The general form of the potential $V_{k}$ is

$$
\begin{equation*}
V_{k}\left(r_{1}, \ldots, r_{k}\right):=-\sum_{i=1}^{k} \sum_{j=i+1}^{k} \frac{z_{i} z_{j}}{\left|r_{i}-r_{j}\right|} \tag{16.4}
\end{equation*}
$$

where $z_{i}$ denotes the charge of particle $i$.
The single particle Hamiltonian $\mathcal{H}_{1}$ will appear in subsequent approximate models for multiparticle systems.

### 16.2.1 Particle spin

In actuality, there are more independent variables than just position and time. The spin $\sigma$ of the particles is a discrete variable which does not effect the form of the equation (16.1) but will just enter as a parameter [98, 190]. Thus the correct functional form of the wave function is $\psi\left(t, r_{1}, \ldots, r_{k}, \sigma_{1}, \ldots, \sigma_{k}\right)$. Pauli [190] postulated the antisymmetry of the wave function:

$$
\begin{equation*}
\psi\left(t, r_{1}, \ldots, r_{k}, \sigma_{1}, \ldots, \sigma_{k}\right)=(-1)^{|\mu|} \psi\left(t, r_{\mu(1)}, \ldots, r_{\mu(k)}, \sigma_{\mu(1)}, \ldots, \sigma_{\mu(k)}\right) \tag{16.5}
\end{equation*}
$$

for any permutation $\mu$ of the integers $\{1, \ldots, k\}$, where $|\mu|$ is the signature of the permutation (zero for even permutations, one for odd permutations).

This mathematical expression implies the Pauli exclusion principle, that there is zero probability of having two particles with the same spin and the same position. Note however, two particles of different spin can occupy the same position with nonzero probability.

### 16.2.2 Interpretation of $\psi$

The solution $\psi\left(t, r_{1}, \ldots, r_{k}\right)$ of (16.1) is called the time-dependent wave function, and $\left|\psi\left(t, r_{1}, \ldots, r_{k}\right)\right|^{2}$ is the probability density of finding particles at positions $r_{1}, \ldots, r_{k}$ at time $t$.

Draft: February 28, 2008, do not distribute

The electric moment may thus be defined in terms of $\psi$. Let $m_{i}$ denote the moment

$$
\begin{equation*}
m_{i}:=\int r_{i}\left|\psi\left(t, r_{1}, \ldots, r_{k}\right)\right|^{2} d r_{1} \cdots d r_{k} \tag{16.6}
\end{equation*}
$$

which is a vector in $R^{3}$. Then the electric moment $m$ of the entire system is

$$
\begin{equation*}
m:=\sum_{i=1}^{k} m_{i} z_{i}=\int \sum_{i=1}^{k} z_{i} r_{i}\left|\psi\left(t, r_{1}, \ldots, r_{k}\right)\right|^{2} d r_{1} \cdots d r_{k} \tag{16.7}
\end{equation*}
$$

where $z_{i}$ is the charge of the $i$-th particle.

### 16.3 The eigenvalue problem

The eigenvalue problem for the Schrödinger equation provides sufficient information to model many chemical interactions. It is a time-independent partial-differential eigenvalue problem of the form

$$
\begin{equation*}
\mathcal{H} \Psi=\lambda \Psi \tag{16.8}
\end{equation*}
$$

where $\mathcal{H}$ is as in Section 16.2
The motivation for the eigenvalue problem is that $\psi=e^{i \lambda t} \Psi$ will be a typical time-dependent solution of (16.1). This assumption requires some thought, given the extreme complexity of the system. In particular, it represents only one mode of oscillation of the system. The related equation of Madelung [242] displays the potential for more complex behavior. However, we will assume that it is sufficient to consider only the fundamental mode of oscillation of the Schrödinger equation here.

In the case of a nucleus of charge $Z$ and a single electron (the hydrogen atom if $Z=1$ ) equation (16.8) can be written

$$
\begin{equation*}
-\frac{\hbar^{2}}{2 m} \nabla^{2} \phi(r)-\frac{Z}{|r|} \phi=\lambda \phi \tag{16.9}
\end{equation*}
$$

by taking the nucleus fixed at the origin. The solutions to this equation can be determined in closed form since the angular and radial variables can be separated, that is,

$$
\begin{equation*}
\phi(r)=\phi(|r| \theta)=u(|r|) v(\theta) \tag{16.10}
\end{equation*}
$$

where $v$ is a spherical harmonic and $u$ decays exponentially at infinity.
In the case of two protons and two electrons (the helium atom), equation (16.8) can be written

$$
\begin{equation*}
-\frac{\hbar^{2}}{2 m} \nabla_{x}^{2} \phi-\frac{\hbar^{2}}{2 m} \nabla_{y}^{2} \phi-Z \frac{1}{|x|} \phi-Z \frac{1}{|y|} \phi+\frac{1}{|x-y|} \phi=\lambda \phi \tag{16.11}
\end{equation*}
$$

by taking the nucleus again fixed at the origin. The solutions $\phi(x, y)$ to this equation apparently can not be separated into a product $\phi_{1}(x) \phi_{2}(y)$ of functions of the the $x$ and $y$ variables, respectively, due to the coupling term $\frac{1}{|x-y|}$.

Draft: February 28, 2008, do not distribute

It is interesting to note that the operator $\mathcal{H}_{k}$ defined in equation (16.3) has certain symmetry properties. We will say that a function $f$ of the $k$ variables $r_{1}, \ldots, r_{k}$ is symmetric if $f\left(r_{1}, \ldots, r_{k}\right)=$ $f\left(x_{\mu(1)}, \ldots, r_{\mu(k)}\right)$ for any permutation $\mu$ of the integers $\{1, \ldots, k\}$. Then it is easy to see that $\mathcal{H}_{k} f$ is symmetric as long as $f$ is symmetric. Similarly, we will say that a function $f$ of the $k$ variables $r_{1}, \ldots, r_{k}$ is antisymmetric if $f\left(r_{1}, \ldots, r_{k}\right)=-f\left(x_{\mu(1)}, \ldots, r_{\mu(k)}\right)$ for any permutation $\mu$ of the integers $\{1, \ldots, k\}$. Then $\mathcal{H}_{k} f$ is antisymmetric as long as $f$ is antisymmetric.

In the case of $k=2$ (as for the helium atom), any function can be decomposed into the sum of a symmetric and an antisymmetric part.

We recall that the Pauli exclusion principle is derived from the inclusion of spin variables $\sigma$ and the assertion that only antisymmetric eigenfunctions are of physical interest. In the case of electrons, the only possible spin values are $\pm \frac{1}{2}$, however for $k$ electrons there are then $2^{k}$ possible spin combinations.

Note that the inclusion of spin in the independent variables does not change the form of the eigenproblem. More precisely, first observe that solutions to the time dependent problem (16.1) which satisfy the condition of antisymmetry (16.5) can be expressed in terms of the eigenmodes of the problem (16.8) restricted to a space of antisymmetric functions satisfying

$$
\begin{equation*}
\phi\left(r_{1}, \ldots, r_{k}, \sigma_{1}, \ldots, \sigma_{k}\right)=(-1)^{|\mu|} \phi\left(r_{\mu(1)}, \ldots, r_{\mu(k)}, \sigma_{\mu(1)}, \ldots, \sigma_{\mu(k)}\right) \tag{16.12}
\end{equation*}
$$

for any permutation $\mu$ of the integers $\{1, \ldots, k\}$, where $|\mu|$ is the signature of the permutation (zero for even permutations, one for odd permutations).

Secondly, one can show that the eigenvalue problem in a space of antisymmetric functions of space and spin (i.e. $r$ and $\sigma$ ) is equivalent to the corresponding eigenvalue problem in a space of antisymmetric functions of space alone. If

$$
\begin{equation*}
\mathcal{H} \psi(r, \sigma)=\lambda_{\sigma} \psi(r, \sigma) \tag{16.13}
\end{equation*}
$$

then how do we know that the function

$$
\begin{equation*}
\tilde{\psi}(r)=\sum_{\sigma} \psi(r, \sigma) \tag{16.14}
\end{equation*}
$$

(where the summation is over all spin values) satisfies the eigenproblem

$$
\begin{equation*}
\mathcal{H} \psi(r)=\lambda \psi(r) \tag{16.15}
\end{equation*}
$$

???
On the other hand, if $\psi(r)$ is any antisymmetric eigenfunction of (16.15) then defining $\psi(r, \sigma)=$ $\psi(r)$ for all $\sigma$ yields an eigenfunctions of (16.13).

Space of antisymmetric functions can be generated by linear combinations of determinants, as observed by Slater [225, [226] and later proved by Löwdin [157]. More precisely, suppose that a given space $\mathcal{F}$ of functions of $k$ variables can be written as a $k$-fold tensor product of a space $F$ of functions of one variable. Here, let us presume that "written as" means equality as topological vector spaces (e.g., Banach spaces). This means that $f\left(r_{1}, \ldots, r_{k}\right) \in \mathcal{F}$ can be written as a limit of linear combinations of products of $k$ functions $f_{i}\left(r_{i}\right) \in F$ of a single variable. Then the antisymmetric functions in $\mathcal{F}$ can be expressed as limits of linear combinations of determinants of matrix expressions $f_{i}\left(r_{j}\right)$.

### 16.4 The Born-Oppenheimer approximation

The Born-Oppenheimer approximation makes the approximation that the movement of electrons does not strongly affect the position of the nuclei. In this case, we write the force field resulting from the interactions of $k$ electrons and $n$ nuclei in the form

$$
\begin{align*}
& V_{k}^{\mathrm{BO}}\left(x_{1}, \ldots, x_{k} ; \xi_{1}, \ldots, \xi_{n}\right):=\sum_{i=1}^{k} \sum_{j=i+1}^{k}\left|x_{i}-x_{j}\right|^{-1}  \tag{16.16}\\
& -\sum_{i=1}^{k} \sum_{\nu=1}^{n} Z_{\nu}\left|x_{i}-\xi_{\nu}\right|^{-1}-\sum_{\mu=1}^{n} \sum_{\nu=\mu+1}^{n} Z_{\mu} Z_{\nu}\left|\xi_{\mu}-\xi_{\nu}\right|^{-1}
\end{align*}
$$

where $Z_{\nu}$ is the charge of the $\nu$-th nucleus, $\left\{x_{i}: i=1, \ldots, k\right\}$ are the locations of the electrons and $\left\{\xi_{\nu}: \nu=1 \ldots, n\right\}$ are the locations of the nuclei. We assume that the charge units have been arranged so that the charge of the electrons is -1 .

Of course, with the appropriate renaming of variables, the Born-Oppenheimer potential $V_{k}^{\mathrm{BO}}$ is the same as the full potential $V_{n+k}$ defined in equation (16.4). However, we choose to think of $V_{k}^{\mathrm{BO}}$ as a function of $3 k$ variables with $3 n$ parameters as opposed to a function of $3(k+n)$ variables. With this in mind, we write

$$
\begin{equation*}
V_{k}^{\mathrm{BO}}\left(x_{1}, \ldots, x_{k} ; \xi_{1}, \ldots, \xi_{n}\right)=V_{k}\left(x_{1}, \ldots, x_{k}\right)+V_{k}^{\mathrm{EF}}\left(x_{1}, \ldots, x_{k} ; \xi_{1}, \ldots, \xi_{n}\right) \tag{16.17}
\end{equation*}
$$

where $V_{k}\left(x_{1}, \ldots, x_{k}\right)=\sum_{i=1}^{k} \sum_{j=i+1}^{k}\left|x_{i}-x_{j}\right|^{-1}$ represents the direct interaction of the $k$ electrons, and $V_{k}^{\mathrm{EF}}$ represents the remaining two terms in (16.16) which we can think of producing an external field.

The full Hamiltonian can similarly be written in the form

$$
\begin{equation*}
\mathcal{H}_{k}=-\frac{\hbar^{2}}{2} \sum_{i=1}^{n} \frac{1}{M_{i}} \Delta_{\xi_{i}}-\frac{\hbar^{2}}{2 m} \Delta_{\mathbf{x}}+V_{k}^{\mathrm{BO}}(\mathbf{x}, \xi) \tag{16.18}
\end{equation*}
$$

where $m$ is the electron mass. Here, $\Delta_{\mathbf{x}}=\sum_{i=1}^{k} \Delta_{x_{k}}$. We will assume that

$$
\begin{equation*}
M:=\min M_{i} \gg m \tag{16.19}
\end{equation*}
$$

Note that $H$ acts on $L^{2}\left(\mathbb{R}^{(3 k+3 n)}\right)$. The Born-Oppenheimer approximation is essentially a perturbation scheme in the small parameter $m / M$, but as is clear from (16.18), this is a very singular perturbation.

The Born-Oppenheimer approximation consists of first studying the spectral problem for the "electronic Hamiltonian,"

$$
\begin{equation*}
\mathcal{H}_{k}^{\mathrm{BO}}(\xi)=-\frac{\hbar^{2}}{2 m} \Delta_{\mathbf{x}}+V_{k}^{\mathrm{BO}}(\mathbf{x}, \xi) \tag{16.20}
\end{equation*}
$$

acting on $L^{2}\left(\mathbb{R}^{3 k}\right)$ (here $\left.\mathbf{x} \in \mathbb{R}^{3 k}\right)$, and treating the nuclear positions as parameters. Thus, we will obtain a sequence of eigenvalues $\lambda_{1}(\xi)<\lambda_{2}(\xi) \leq \lambda_{3}(\xi) \leq \ldots$, for each choice of the nuclear
positions $\xi$. Born and Oppenheimer then argued that the eigenvalues of the full problem are given (approximately) by the sequence of simplified "nuclear Hamiltonians"

$$
\begin{equation*}
\mathcal{H}_{j}^{\mathrm{N}}=-\frac{\hbar^{2}}{2 M} \Delta_{\underset{\sim}{ }}+\lambda_{j}(\xi) \tag{16.21}
\end{equation*}
$$

which act on $L^{2}\left(\mathbb{R}^{3 n}\right)\left(\right.$ here $\left.\xi \in \mathbb{R}^{3 n}\right)$.
Some insight into the relation between the spectrum of (16.20) and (16.21) comes from remarking that if $u_{1}(\cdot ; \xi)$ is the eigenfunction of (16.20), with eigenvalue $\lambda_{1}(\xi)$, and if $\psi(\cdot)$ is an eigenfunction of the nuclear Hamiltonian $H_{1}^{N}$, then $\psi(\xi) u_{1}(\mathbf{x}, \xi)$ is an eigenfunction of the full Hamiltonian (16.18). There is, however, no reason for all of the eigenfunctions of the full problem to factor in this way, and a better way of understanding the relationship between $H^{B O}$ and $H_{j}^{N}$ is through a reduction method first introduced by Feshbach in the context of nuclear physics [86]. Suppose that we expand an arbitrary function $\psi(\mathbf{x}, \xi)=\sum_{n=0}^{\infty} \phi_{n}(\mathbf{x}, \xi) \psi_{n}(\xi)$, where for each $\xi$, we choose a complete, orthonormal set of eigenfunctions $\left\{\phi_{n}(\cdot, \xi)\right\}$ for $H^{B O}(\xi)$. (One can choose these eigenfunctions to be at least $C^{2}$ with respect to $\xi$ [162.) If one then substitutes this expansion into the "full" eigenvalue problem:

$$
\begin{equation*}
H \psi(\mathbf{x}, \xi)=\lambda \psi(\mathbf{x}, \xi) \tag{16.22}
\end{equation*}
$$

and then projects onto the eigenfunctions $\phi_{j}$, one obtains the coupled system of eigenvalue problems

$$
\begin{equation*}
H_{j}^{N} \psi_{j}+\sum_{k} \frac{\hbar^{2}}{2 M}\left\langle\phi_{j}, \Delta_{\underset{\xi}{\xi}} \phi_{k}\right\rangle_{x} \psi_{k}=\lambda \psi_{j} \quad j=0,1,2,3 \ldots \tag{16.23}
\end{equation*}
$$

Here, $\langle\cdot, \cdot\rangle_{x}$ means that we take the inner product only with respect to the electron coordinates.
Feshbach then notes that if one defines projection operators $P_{0}$, which projects onto the $\psi_{0}$ direction, and $\hat{P}_{0}$, the orthogonal complement of $P_{0}$, then one can rewrite the $j=0$ equation in (16.23) as an uncoupled equation for $\psi_{0}$. If one applies $P_{0}$ and $\hat{P}_{0}$ to (16.22), one obtains:

$$
\begin{gather*}
P_{0} H P_{0} \psi+P_{0} H \hat{P}_{0} \psi=\lambda P_{0} \psi  \tag{16.24}\\
\hat{P}_{0} \psi=\left(\hat{P}_{0}(J-\lambda) \hat{P}_{0}\right)^{-1} \hat{P}_{0} H \hat{P} \psi \tag{16.25}
\end{gather*}
$$

where $\left(\hat{P}_{0}(J-\lambda) \hat{P}_{0}\right)^{-1}$ exists provided there is a gap between the lowest eigenvalue of $H^{B O}$ and the remainder of its spectrum - i.e. provided $\delta=\inf _{\xi} \operatorname{dist}\left(\lambda_{0}(\xi), \operatorname{spec}\left(H^{B O}(\xi) \backslash \lambda_{0}(\xi)\right)>0\right.$. Inserting the first of these equations into the second gives

$$
\begin{equation*}
P_{0} H P_{0}+P_{0} H\left(\hat{P}_{0}(J-\lambda) \hat{P}_{0}\right)^{-1} \hat{P}_{0} H P_{0} \psi=\lambda P_{0} \psi \tag{16.26}
\end{equation*}
$$

Recalling that $H=-\frac{\hbar^{2}}{2 M} \Delta_{\mathcal{\xi}}+H^{B O}(\xi)$, and that $P_{0}$ and $\hat{P}_{0}$ are spectral projections for $H^{B O}$, one finds that $\hat{P}_{0} H P_{0}=-\frac{\hbar^{2}}{2 M} \hat{P}_{0} \Delta_{\underset{\sim}{\xi}} P_{0}$, while $P_{0} H P_{0}=-\frac{\hbar^{2}}{2 M} P_{0} \Delta_{\underset{\sim}{\xi}} P_{0}+\lambda_{0}(\xi) P_{0}$. Thus, since $\psi_{0}=P_{0} \psi$, (16.24) is equivalent to

$$
\begin{align*}
\left(-\frac{\hbar^{2}}{2 M} \Delta_{\xi}+\lambda_{0}(\xi)\right) \psi_{0} & +\frac{\hbar^{2}}{2 M}\left\langle\phi_{0}, \Delta_{\xi} \phi_{0}\right\rangle_{x} \psi_{0} \\
& +\left(\frac{\hbar^{2}}{2 M}\right)^{2} \hat{P}_{0} \Delta_{\xi} P_{0}\left(\hat{P}_{0}(J-\lambda) \hat{P}_{0}\right)^{-1} \hat{P}_{0} \Delta_{\xi} \psi_{0}=\lambda \psi_{0} \tag{16.27}
\end{align*}
$$

Draft: February 28, 2008, do not distribute

Thus, up to the correction terms, $\frac{\hbar^{2}}{2 M}\left\langle\phi_{0}, \Delta_{\xi} \phi_{0}\right\rangle_{x}$ and $\left(\frac{\hbar^{2}}{2 M}\right)^{2} \hat{P}_{0} \Delta_{\xi} P_{0}\left(\hat{P}_{0}(J-\lambda) \hat{P}_{0}\right)^{-1} \hat{P}_{0} \Delta_{\xi}$, the spectral problem for (16.22) is equivalent to that for $H_{0}^{N}$, at least when $\lambda \approx \lambda_{0}$.

Note further that the perturbations in (16.27) are no longer singular perturbations, since both of the perturbations are relatively bounded with respect to $H_{0}^{N}$. Indeed, while it is technically rather involved, the above approach can be made rigorous, [103], [104], [132], [162] and one finds that there exists an asymptotic expansion of the eigenvalues of (16.22) in powers of $\left(\frac{m}{M}\right)^{1 / 2}$ with the leading order in this expansion given by the eigenvalues of $H_{0}^{N}$.

From now on, we will assume that the Born-Oppenheimer approximation is in force.

### 16.5 External fields

Now let us consider the effect of an external field on solutions of the eigenvalue problem for the Schrödinger equation. This takes the form

$$
\begin{equation*}
\mathcal{H} \Psi+\epsilon B \Psi=\lambda \Psi \tag{16.28}
\end{equation*}
$$

where $\mathcal{H}$ is as in Section 16.2 and $B$ represents the external field. We introduce a small parameter $\epsilon$ to facilitate discussion.

More precisely, we have $\mathcal{H}:=\mathcal{H}_{k}+V_{k}^{\mathrm{BO}}$ and $B=\epsilon^{-1} V_{k}^{\mathrm{EF}}$ where

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(x_{1}, \ldots, x_{k} ; \eta_{1}, \ldots, \eta_{n}\right)=-\sum_{i=1}^{k} \sum_{\nu=1}^{n} z_{i} Z_{\nu}\left|x_{i}-\eta_{\nu}\right|^{-1}-\sum_{\mu=1}^{n} \sum_{\nu=\mu+1}^{n} Z_{\mu} Z_{\nu}\left|\eta_{\mu}-\eta_{\nu}\right|^{-1} \tag{16.29}
\end{equation*}
$$

where the charges $Z_{\nu}$ produce the external field. The second term is independent of $x_{i}$, so we can just treat it as a constant, $C_{Z}$. Suppose that the two systems are separated by a distance $R=\min \left\{\left|x_{i}-\eta_{\nu}\right|\right\}$, so that we can write $\eta_{\nu}=\eta_{\nu}^{\prime}-R \mathbf{E}$ for some unit vector $\mathbf{E}$ and expand the first term in powers of $R$ :

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(x_{i}, \eta_{\nu}\right)-C_{Z}=-\sum_{i=1}^{k} \sum_{\nu=1}^{n} \frac{z_{i} Z_{\nu}}{\left|x_{i}-\eta_{\nu}\right|}=-\frac{1}{R} \sum_{i=1}^{k} \sum_{j=1}^{k} \frac{z_{i} Z_{\nu}}{\left|E+R^{-1}\left(x_{i}-\eta_{\nu}\right)\right|} . \tag{16.30}
\end{equation*}
$$

We then have the approximation

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(x_{i}, \eta_{i}\right)=C_{Z}+R^{-1}\left(A_{0}+A_{1} R^{-1}+A_{2} R^{-2}+\cdots\right) \tag{16.31}
\end{equation*}
$$

where

$$
\begin{equation*}
A_{0}=-\sum_{i=1}^{k} \sum_{\nu=1}^{n} z_{i} Z_{\nu} \tag{16.32}
\end{equation*}
$$

and

$$
\begin{equation*}
A_{1}=\sum_{i=1}^{k} \sum_{\nu=1}^{n} z_{i} Z_{\nu} \mathbf{E} \cdot\left(x_{i}-\eta_{\nu}\right) \tag{16.33}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

If we assume that $\sum_{j=1}^{k} z_{j}=0$, then (16.32) simplifies to $A_{0}=0$. Similarly, $A_{1}$ becomes

$$
\begin{equation*}
A_{1}=\sum_{i=1}^{k} \sum_{\nu=1}^{n} z_{i} Z_{\nu} \mathbf{E} \cdot x_{i}=\left(\sum_{\nu=1}^{n} Z_{\nu}\right) \mathbf{E} \cdot\left(\sum_{i=1}^{k} z_{i} x_{i}\right) \tag{16.34}
\end{equation*}
$$

Thus we can write

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(r_{i}^{1}, r_{i}^{2}\right) \approx C_{Z}+\mathcal{E} \cdot M\left(x_{i}\right) \tag{16.35}
\end{equation*}
$$

where $\mathcal{E}$ denotes the external electric field

$$
\begin{equation*}
\mathcal{E}=\frac{1}{R^{2}}\left(\sum_{\nu=1}^{n} Z_{\nu}\right) \mathbf{E} \tag{16.36}
\end{equation*}
$$

and

$$
\begin{equation*}
M\left(x_{i}\right):=\sum_{i=1}^{k} z_{i} x_{i} \tag{16.37}
\end{equation*}
$$

is the electric moment of the system with charges $z_{i}$ at positions $x_{i}$. With these approximations, our eigenvalue problem becomes

$$
\begin{equation*}
\mathcal{H}_{k} \Psi+\mathcal{E} \cdot M\left(x_{i}\right) \Psi=\left(\lambda-C_{Z}\right) \Psi \tag{16.38}
\end{equation*}
$$

which is of the form (16.28) with $\epsilon B=\mathcal{E} \cdot M\left(x_{i}\right)$, and with the eigenvalue shifted by a constant.
An eigenvalue problem is a nonlinear system, so a perturbation is not simple to analyze, except in the case where the perturbation is small. Let us call the solution to (16.38) $\Psi^{\epsilon}$. Hence $\Psi^{0}$ is the solution to the unperturbed eigenvalue problem (16.8). Let us suppose that we can write $\Psi^{\epsilon} \approx \Psi^{0}+\epsilon \Psi^{\prime}$ for some $\Psi^{\prime}$ at least for small $\epsilon$.

In general [172], the dependence of $\lambda^{\epsilon}$ on $\epsilon$ can be quite singular. However, let us assume that $\lambda^{\epsilon} \approx \lambda^{0}+\epsilon \lambda^{\prime}$ for small $\epsilon$. Then taking difference quotients and letting $\epsilon \rightarrow 0$, we find

$$
\begin{equation*}
\left(\mathcal{H}-\lambda^{0} \mathcal{I}\right) \Psi^{\prime}=-B \Psi^{0}+\lambda^{\prime} \Psi^{0} \tag{16.39}
\end{equation*}
$$

Since $\left\|\Psi^{\epsilon}\right\|=1$ for all $\epsilon$, we must have $\Psi^{\prime} \in\left[\Psi^{0}\right]^{\perp}$, where $\left[\Psi^{0}\right]$ denotes the linear space spanned by $\Psi^{0}$ (Exercise 16.3). If $\lambda^{0}$ is a simple eigenvalue, then this implies that (16.39) is solvable provided $\lambda^{\prime}=\left(\Psi^{0}, B \Psi^{0}\right)$, since this implies that $B \Psi^{0}+\lambda^{\prime} \Psi^{0} \in\left[\Psi^{0}\right]^{\perp}$.

### 16.5.1 Polarization

Polarization of a chemical system is simply the result of the action of an external field. The polarizability $\alpha$ of a material can be defined as the infinitesimal response to a small external field. More precisely, $\alpha$ is the matrix that gives the change in the electric moment (16.7) as a response to the input external field.

The electric moment for a system where $z_{i}$ is the charge of the $i$-th particle is defined in terms of $\Psi$ as in (16.7) via

$$
\begin{equation*}
m:=\sum_{i=1}^{k} z_{i} \int r_{i}\left|\Psi\left(r_{1}, \ldots, r_{k}\right)\right|^{2} d r_{1} \cdots d r_{k} \tag{16.40}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

If $\mathcal{E}=\epsilon \mathbf{e}^{j}$ is our external field, then the resulting infinitesimal change in $i$-th moment is given by

$$
\begin{align*}
& \Delta m_{i}^{j}:  \tag{16.41}\\
&=\lim _{\epsilon \rightarrow 0} \frac{1}{\epsilon} \int r_{i}\left(\left|\Psi^{\mathcal{E}}\left(r_{1}, \ldots, r_{k}\right)\right|^{2}-\left|\Psi^{0}\left(r_{1}, \ldots, r_{k}\right)\right|^{2}\right) d r_{1} \cdots d r_{k} \\
&=2 \int r_{i} \Psi^{0}\left(r_{1}, \ldots, r_{k}\right) \Psi_{j}^{\prime}\left(r_{1}, \ldots, r_{k}\right) d r_{1} \cdots d r_{k}
\end{align*}
$$

where $\Psi_{j}^{\prime} \in\left[\Psi^{0}\right]^{\perp}$ solves the linear system

$$
\begin{equation*}
\left(\mathcal{H}-\lambda^{0} \mathcal{I}\right) \Psi_{j}^{\prime}=-e^{j} \cdot M(r) \Psi^{0}+\lambda_{j}^{\prime} \Psi^{0} \tag{16.42}
\end{equation*}
$$

with

$$
\begin{align*}
\lambda_{j}^{\prime} & =\left(\Psi^{0}, e^{j} \cdot M(r) \Psi^{0}\right) \\
& =\int \sum_{i=1}^{k} z_{i} r_{i}^{j}\left|\Psi^{0}\left(r_{1}, \ldots, r_{k}\right)\right|^{2} d r_{1} \cdots d r_{k} \tag{16.43}
\end{align*}
$$

Thus $\alpha$ is the matrix

$$
\begin{equation*}
\alpha_{j, \ell}=2 \int \sum_{i=1}^{k} z_{i} r_{i}^{\ell} \Psi^{0}\left(r_{1}, \ldots, r_{k}\right) \Psi_{j}^{\prime}\left(r_{1}, \ldots, r_{k}\right) d r_{1} \cdots d r_{k} \tag{16.44}
\end{equation*}
$$

Note that $\lambda_{j}^{\prime}$ is the $j$-th component of the electric moment of the unperturbed system defined in (16.40).

### 16.5.2 Induced fields

Induced dipoles represent an intriguing example of external fields. Each dipole provides the external field that induces the other dipole. This sounds like a circular argument, and so it is deserving of a more detailed examination, which we now present. When two parts of an atomic system are far apart from each other, it is reasonable to assume that the coupling should be quite weak. In particular, let us suppose that the two sub-systems are identical, for simplicity.

Let us return to the basic notation of (16.3), but now we refer to the variables of one system by $r_{i}^{1}$ and the other by $r_{i}^{2}$. The full system operator is thus

$$
\begin{align*}
\mathcal{H}_{k} \phi\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right): & =-\sum_{i=1}^{k} \frac{\hbar^{2}}{2 m_{i}} \nabla_{r_{i}^{1}}^{2} \phi-\sum_{i=1}^{k} \frac{\hbar^{2}}{2 m_{i}} \nabla_{r_{i}^{2}}^{2} \phi \\
+V_{k}\left(r_{1}^{1}, \ldots, r_{k}^{1}\right) \phi & +V_{k}\left(r_{1}^{2}, \ldots, r_{k}^{2}\right) \phi+V_{k}^{\mathrm{EF}}\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right) \phi  \tag{16.45}\\
& =\left(A_{r^{1}}+A_{r^{2}}+V_{k}^{\mathrm{EF}}\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right)\right) \phi
\end{align*}
$$

where $\phi=\phi\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right)$ and we have separated the potential terms as in (16.17). Suppose that the two systems are separated by a distance $R$ along an axis that is parallel to a unit vector $E$, so that by renaming variables by

$$
\begin{equation*}
r_{j}^{2} \rightarrow-r_{j}^{2}+R \mathbf{E} \tag{16.46}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
we can expand $V_{k}^{\mathrm{EF}}$ as

$$
\begin{align*}
V_{k}^{\mathrm{EF}}\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right) & =-\sum_{i=1}^{k} \sum_{j=1}^{k} \frac{z_{i} z_{j}}{\left|r_{i}^{1}-R \mathbf{E}+r_{j}^{2}\right|}  \tag{16.47}\\
& =-\frac{1}{R} \sum_{i=1}^{k} \sum_{j=1}^{k} \frac{z_{i} z_{j}}{\left|E-R^{-1}\left(r_{i}^{1}+r_{j}^{2}\right)\right|}
\end{align*}
$$

We can make the asymptotic expansion

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right)=\frac{a_{0}}{R}+\frac{a_{1}}{R^{2}}+\frac{a_{2}}{R^{3}}+\cdots \tag{16.48}
\end{equation*}
$$

where we can easily verify that

$$
\begin{equation*}
a_{0}=-\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j}=-\sum_{i=1}^{k} z_{i}\left(\sum_{j=1}^{k} z_{j}\right) \tag{16.49}
\end{equation*}
$$

If the net charge on each molecule is zero, then

$$
\begin{equation*}
\sum_{j=1}^{k} z_{j}=0 \tag{16.50}
\end{equation*}
$$

so (16.48) simplifies since $a_{0}=0$. We will show subsequently that

$$
\begin{equation*}
a_{1}=\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j} \mathbf{E} \cdot\left(r_{i}^{1}+r_{j}^{2}\right) \tag{16.51}
\end{equation*}
$$

Expanding we find that

$$
\begin{align*}
a_{1} & =\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j} \mathbf{E} \cdot r_{i}^{1}+\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j} \mathbf{E} \cdot r_{j}^{2}  \tag{16.52}\\
& =\left(\sum_{j=1}^{k} z_{j}\right) \sum_{i=1}^{k} z_{i} \mathbf{E} \cdot r_{i}^{1}+\left(\sum_{i=1}^{k} z_{i}\right) \sum_{j=1}^{k} z_{j} \mathbf{E} \cdot r_{j}^{2}=0
\end{align*}
$$

provided (16.50) holds.
To clarify the computations, let us write $\epsilon=1 / R$ and then

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right)=-\epsilon S(\epsilon), \tag{16.53}
\end{equation*}
$$

where the expression $S$ is given by

$$
\begin{equation*}
S(\epsilon)=\sum_{i=1}^{k} \sum_{j=1}^{k} \frac{z_{i} z_{j}}{\left|\mathbf{E}-\epsilon\left(r_{i}^{1}+r_{j}^{2}\right)\right|} \tag{16.54}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

We have $S(0)=a_{0}$, and computing the derivative of $S$ with respect to $\epsilon$ yields $a_{1}$. Let $S_{i j}$ denote an individual term in the double summation (16.54):

$$
\begin{equation*}
S_{i j}(\epsilon)=\frac{z_{i} z_{j}}{|\mathbf{E}-\epsilon r|}=z_{i} z_{j}\left(|\mathbf{E}|^{2}-2 \epsilon \mathbf{E} \cdot r+\epsilon^{2}|r|^{2}\right)^{-1 / 2} \tag{16.55}
\end{equation*}
$$

where $r=r_{i}^{1}+r_{j}^{2}$. Therefore

$$
\begin{equation*}
S_{i j}^{\prime}(\epsilon)=z_{i} z_{j}\left(\mathbf{E} \cdot r-\epsilon|r|^{2}\right)\left(|\mathbf{E}|^{2}-2 \epsilon \mathbf{E} \cdot r+\epsilon^{2}|r|^{2}\right)^{-3 / 2} \tag{16.56}
\end{equation*}
$$

so that setting $\epsilon=0$ confirms the computation of $a_{1}=S^{\prime}(0)$, since $|\mathbf{E}|=1$. Differentiating (16.56), we find

$$
\begin{align*}
S_{i j}^{\prime \prime}(\epsilon) & =3 z_{i} z_{j}\left(\mathbf{E} \cdot r-\epsilon|r|^{2}\right)^{2}\left(|\mathbf{E}|^{2}-2 \epsilon \mathbf{E} \cdot r+\epsilon^{2}|r|^{2}\right)^{-5 / 2}  \tag{16.57}\\
& -z_{i} z_{j}|r|^{2}\left(|\mathbf{E}|^{2}-2 \epsilon \mathbf{E} \cdot r+\epsilon^{2}|r|^{2}\right)^{-3 / 2}
\end{align*}
$$

and thus $S_{i j}^{\prime \prime}(0)=3 z_{i} z_{j}(\mathbf{E} \cdot r)^{2}-z_{i} z_{j}|r|^{2}$. Therefore

$$
\begin{align*}
S^{\prime \prime}(0) & =\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j}\left(3\left(\mathbf{E} \cdot\left(r_{i}^{1}+r_{j}^{2}\right)\right)^{2}-\left|r_{i}^{1}+r_{j}^{2}\right|^{2}\right) \\
& =\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j}\left(3\left(\left(\mathbf{E} \cdot r_{i}^{1}\right)^{2}+2\left(\mathbf{E} \cdot r_{i}^{1}\right)\left(\mathbf{E} \cdot r_{j}^{2}\right)+\left(\mathbf{E} \cdot r_{j}^{2}\right)^{2}\right)-\left(\left|r_{i}^{1}\right|^{2}+2 r_{i}^{1} \cdot r_{j}^{2}+\left|r_{j}^{2}\right|^{2}\right)\right) \\
& =\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j}\left(6\left(\mathbf{E} \cdot r_{i}^{1}\right)\left(\mathbf{E} \cdot r_{j}^{2}\right)-2\left(r_{i}^{1} \cdot r_{j}^{2}\right)\right) \tag{16.58}
\end{align*}
$$

provided (16.50) holds. Define the electric moment

$$
\begin{equation*}
D\left(r_{1}^{\ell}, \ldots, r_{k}^{\ell}\right)=\sum_{i=1}^{k} z_{i} r_{i}^{\ell}, \quad \ell=1,2 \tag{16.59}
\end{equation*}
$$

Note that $a_{2}=\frac{1}{2} S^{\prime \prime}(0)$. Then if (16.50) holds, (16.58) implies

$$
\begin{align*}
a_{2} & \left.=\sum_{i=1}^{k} \sum_{j=1}^{k} z_{i} z_{j}\left(3\left(\mathbf{E} \cdot r_{i}^{1}\right)\left(\mathbf{E} \cdot r_{j}^{2}\right)\right)-\left(r_{i}^{1} \cdot r_{j}^{2}\right)\right)  \tag{16.60}\\
& =3\left(\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)-D\left(\mathbf{r}^{1}\right) \cdot D\left(\mathbf{r}^{2}\right)\right) .
\end{align*}
$$

where we set $\mathbf{r}^{\ell}=\left(r_{1}^{\ell}, \ldots, r_{k}^{\ell}\right)$. From (16.48), we conclude that

$$
\begin{equation*}
V_{k}^{\mathrm{EF}}\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right) \approx \frac{a_{2}}{R^{3}}=\frac{1}{R^{3}}\left(3\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)-D\left(\mathbf{r}^{1}\right) \cdot D\left(\mathbf{r}^{2}\right)\right) \tag{16.61}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute 195

Now the eigenvalue problem can be written

$$
\begin{equation*}
A f_{\epsilon}+\epsilon B f_{\epsilon}=\lambda_{\epsilon} f_{\epsilon}, \tag{16.62}
\end{equation*}
$$

where $A=A_{\mathbf{r}^{1}}+A_{\mathbf{r}^{2}}, \epsilon=R^{-3}$ and $B$ is the multiplication operator defined by

$$
\begin{equation*}
B\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)=B\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right)=3\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)-D\left(\mathbf{r}^{1}\right) \cdot D\left(\mathbf{r}^{2}\right) \tag{16.63}
\end{equation*}
$$

Suppose that $f_{\epsilon}=f_{0}+\epsilon f^{\prime}+\mathcal{O}\left(\epsilon^{2}\right)$ and that $f_{0}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)=\hat{f}_{0}\left(\mathbf{r}^{1}\right) \hat{f}_{0}\left(\mathbf{r}^{2}\right)$ with the property that $\hat{f}_{0}$ satisfies

$$
\begin{equation*}
\int D(\mathbf{r})\left|\hat{f}_{0}\left(r_{1}, \ldots, r_{k}\right)\right|^{2} d r_{1} \cdots d r_{k}=0 \tag{16.64}
\end{equation*}
$$

Here, the integrand is a vector, so we are saying the integral of all three components is zero. If we take the dot-product with $\mathbf{E}$, we also find

$$
\begin{equation*}
\int \mathbf{E} \cdot D(\mathbf{r})\left|\hat{f}_{0}\left(r_{1}, \ldots, r_{k}\right)\right|^{2} d r_{1} \cdots d r_{k}=0 \tag{16.65}
\end{equation*}
$$

Therefore (16.64) implies that

$$
\begin{align*}
& \int B\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)\left|f_{0}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{1} d \mathbf{r}^{2}= \\
& \int\left(3\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)-D\left(\mathbf{r}^{1}\right) \cdot D\left(\mathbf{r}^{2}\right)\right)\left|\hat{f}_{0}\left(\mathbf{r}^{1}\right) \hat{f}_{0}\left(\mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{1} d \mathbf{r}^{2}= \\
& 3 \int\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)\left|\hat{f}_{0}\left(\mathbf{r}^{1}\right)\right|^{2} d \mathbf{r}^{1} \int\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)\left|\hat{f}_{0}\left(\mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{2}  \tag{16.66}\\
& -\int D\left(\mathbf{r}^{1}\right)\left|\hat{f}_{0}\left(\mathbf{r}^{1}\right)\right|^{2} d \mathbf{r}^{1} \cdot \int D\left(\mathbf{r}^{2}\right)\left|\hat{f}_{0}\left(\mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{2}=0 .
\end{align*}
$$

Note that $f^{\prime}$ satisfies $f^{\prime} \perp f_{0}$ and

$$
\begin{align*}
-\sum_{i=1}^{k} \frac{\hbar^{2}}{2 m_{i}} \nabla_{r_{i}^{1}}^{2} f^{\prime}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)-\sum_{i=1}^{k} \frac{\hbar^{2}}{2 m_{i}} \nabla_{r_{i}^{2}}^{2} f^{\prime}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)+ & \left(V_{k}\left(\mathbf{r}^{1}\right)+V_{k}\left(\mathbf{r}^{2}\right)-\lambda_{0}\right) f^{\prime}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)  \tag{16.67}\\
= & -B\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right) \hat{f}_{0}\left(\mathbf{r}^{1}\right) \hat{f}_{0}\left(\mathbf{r}^{2}\right)
\end{align*}
$$

since $f_{0}$ is orthogonal to $B f_{0}$, by (16.66). Even though $f_{0}$ factors into a product, we cannot solve (16.67) by separation of variables as a product. The coupling of the right-hand side term $D\left(\mathbf{r}^{1}\right) \cdot D\left(\mathbf{r}^{2}\right)$ prevents this. Thus the dipole $\widetilde{D}_{\epsilon}$ induced is

$$
\begin{align*}
\widetilde{D}_{\epsilon} & =\int D\left(\mathbf{r}^{1}\right)\left|f_{\epsilon}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{1} d \mathbf{r}^{2} \\
& \approx \int D\left(\mathbf{r}^{1}\right)\left|\left(f_{0}+\epsilon f^{\prime}\right)\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{1} d \mathbf{r}^{2}+\mathcal{O}\left(\epsilon^{2}\right)  \tag{16.68}\\
& =2 \epsilon \int D\left(\mathbf{r}^{1}\right)\left(f_{0} f^{\prime}\right)\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right) d \mathbf{r}^{1} d \mathbf{r}^{2}+\mathcal{O}\left(\epsilon^{2}\right) \\
& =\frac{2}{R^{3}} \int D\left(\mathbf{r}^{1}\right)\left(f_{0} f^{\prime}\right)\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right) d \mathbf{r}^{1} d \mathbf{r}^{2}+\mathcal{O}\left(R^{-6}\right) .
\end{align*}
$$

Draft: February 28, 2008, do not distribute

Let us give a name to the dipolar coefficient of $R^{-3}$ in (16.68):

$$
\begin{equation*}
\mathcal{D}=\int D\left(\mathbf{r}^{1}\right)\left(f_{0} f^{\prime}\right)\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right) d \mathbf{r}^{1} d \mathbf{r}^{2} \tag{16.69}
\end{equation*}
$$

Then we can state our main result as follows: the induced dipole $\widetilde{D}_{\epsilon}$ is

$$
\begin{equation*}
\widetilde{D}_{\epsilon}=\int D\left(\mathbf{r}^{1}\right)\left|f_{\epsilon}\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{1} d \mathbf{r}^{2}=\frac{2}{R^{3}} \mathcal{D}+\mathcal{O}\left(R^{-6}\right) \tag{16.70}
\end{equation*}
$$

where $\mathcal{D}$ is defined in (16.69).
It is noteworthy that the term $\frac{2}{R^{3}}$ is significant for $R=4$, a distance at which van der Waals forces are often a dominant effect. The error term $\epsilon^{2}=2^{-18}$ is quite small for $R=4$, so we can trust the accuracy of the prediction (16.70) in this range.

In the approach reported in [193], the perturbation analysis is based on the beavhior of $\lambda^{\epsilon}$ instead of the wave function $f_{\epsilon}$ as we have done here. The deviation $\lambda^{\epsilon}-\lambda^{0}$ is a direct measure of the energy of the induced dipole. As we have determined, $\lambda^{\prime}=0$, so it would appear that

$$
\begin{equation*}
\lambda^{\epsilon}-\lambda^{0}=\mathcal{O}\left(\epsilon^{2}\right)=\mathcal{O}\left(R^{-6}\right) \tag{16.71}
\end{equation*}
$$

but we have not identified an experession for the coefficient $\lambda^{\prime \prime}$ required to quantify this asymptotic estimate. The degeneracy caused by having $\lambda^{\prime}=0$ necessitates a so-called second-order perturbation method in 193, and only an approximate version of that is utilized.

By emphasizing the calculation of the wave function, we have derived an expression that is not degenerate in $\epsilon$ and indeed gives the expected behavior of the induced dipole strength as a function of $R$. Of course, evaluating numerically the coefficient of $R^{-3}$ presented in (16.68) requires some work, but conceptually it is clear how to do this by solving the Schrödinger equation together with a closely related auxiliary equation.

### 16.5.3 Hartree approximation

The Hartree approximation can sometimes provide interesting information. It uses the approximation

$$
\begin{equation*}
\phi\left(r_{1}^{1}, \ldots, r_{k}^{1}, r_{1}^{2}, \ldots, r_{k}^{2}\right) \approx f^{1}\left(r_{1}^{1}, \ldots, r_{k}^{1}\right) f^{2}\left(r_{1}^{2}, \ldots, r_{k}^{2}\right) \tag{16.72}
\end{equation*}
$$

We will see that this leads to an interesting interpretation of the induced dipole, with similar asymptotic behvior.

Let us write $\mathcal{H}_{k}=\mathcal{H}_{k}^{1}+\mathcal{H}_{k}^{2}+V_{k}^{\mathrm{EF}}$. Then

$$
\begin{align*}
\mathcal{H}_{k} \phi & \approx f^{2} \mathcal{H}_{k}^{1} f^{1}+f^{1} \mathcal{H}_{k}^{2} f^{2}+V_{k}^{\mathrm{EF}} f^{1} f^{2} \\
& =f^{2}\left(\mathbf{r}^{2}\right)\left(\mathcal{H}_{k}^{1} f^{1}+\frac{1}{2} V_{k}^{\mathrm{EF}}\left(\cdot, \mathbf{r}^{2}\right) f^{1}\right)+f^{1}\left(\mathbf{r}^{1}\right)\left(\mathcal{H}_{k}^{2} f^{2}+\frac{1}{2} V_{k}^{\mathrm{EF}}\left(\mathbf{r}^{1}, \cdot\right) f^{2}\right)  \tag{16.73}\\
& \approx f^{2}\left(\mathbf{r}^{2}\right)\left(\mathcal{H}_{k}^{1} f^{1}+\frac{1}{2} R^{-3} B\left(\cdot, \mathbf{r}^{2}\right) f^{1}\right)+f^{1}\left(\mathbf{r}^{1}\right)\left(\mathcal{H}_{k}^{2} f^{2}+\frac{1}{2} R^{-3} B\left(\mathbf{r}^{1}, \cdot\right) f^{2}\right)
\end{align*}
$$

Draft: February 28, 2008, do not distribute
where we used the approximation (16.61). We further approximate by replacing the dipoles in the definition of $B$ by ones given by their expected positions. That is, let $\hat{r}_{i}^{1}$ denote the expected position of the $i$-th atom,

$$
\begin{equation*}
\hat{r}_{i}^{\ell}:=\int r_{i}^{\ell}\left|f^{\ell}\left(r_{1}^{\ell}, \ldots, r_{k}^{\ell}\right)\right|^{2} d r_{1}^{\ell} \cdots d r_{k}^{\ell} \tag{16.74}
\end{equation*}
$$

and define

$$
\begin{equation*}
\widehat{D}\left(r^{\ell}\right)=\widehat{D}^{\ell}=\sum_{i=1}^{k} z_{i} \hat{r}_{i}^{\ell} \tag{16.75}
\end{equation*}
$$

for $\ell=1,2$. Thus

$$
\begin{equation*}
\widehat{D}^{\ell}:=\sum_{i=1}^{k} z_{i} \int r_{i}^{\ell}\left|f^{\ell}\left(r_{1}^{\ell}, \ldots, r_{k}^{\ell}\right)\right|^{2} d r_{1}^{\ell} \cdots d r_{k}^{\ell} \tag{16.76}
\end{equation*}
$$

We can thus write the expression $B\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right)$ as either

$$
\begin{equation*}
B\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right) \approx 3\left(\mathbf{E} \cdot \widehat{D}^{1}\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)-\widehat{D}^{1} \cdot D\left(\mathbf{r}^{2}\right) \tag{16.77}
\end{equation*}
$$

or

$$
\begin{equation*}
B\left(\mathbf{r}^{1}, \mathbf{r}^{2}\right) \approx 3\left(\mathbf{E} \cdot \widehat{D}^{2}\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)-\widehat{D}^{2} \cdot D\left(\mathbf{r}^{1}\right) \tag{16.78}
\end{equation*}
$$

depending on the context. Then (16.73) becomes

$$
\begin{align*}
\mathcal{H}_{k} \phi & \approx f^{2}\left(\mathbf{r}^{2}\right)\left(\mathcal{H}_{k}^{1} f^{1}+\frac{1}{2} R^{-3}\left(3\left(\mathbf{E} \cdot \widehat{D}^{2}\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{1}\right)\right)-\widehat{D}^{2} \cdot D\left(r^{1}\right) f^{1}\right)\right)  \tag{16.79}\\
& +f^{1}\left(\mathbf{r}^{1}\right)\left(\mathcal{H}_{k}^{2} f^{2}+\frac{1}{2} R^{-3}\left(3\left(\mathbf{E} \cdot \widehat{D}^{1}\right)\left(\mathbf{E} \cdot D\left(\mathbf{r}^{2}\right)\right)-\widehat{D}^{1} \cdot D\left(r^{2}\right) f^{2}\right)\right)
\end{align*}
$$

Let us make the ansatz that $f^{1}=f^{2}$. This reduces the eigenvalue problem to one for a single distribution $f=f^{1}=f^{2}$. Then the eigenvalue problem $\mathcal{H}_{k} \phi=\lambda \phi$ would collapse into a single eigenvalue problem

$$
\begin{equation*}
\mathcal{H}_{k}^{\rho} f+\frac{1}{2} R^{-3}(3(\mathbf{E} \cdot \widehat{D})(\mathbf{E} \cdot D(\rho))-\widehat{D} \cdot D(\rho)) f=\lambda f \tag{16.80}
\end{equation*}
$$

where $\rho$ denotes the spatial variable name, and the induced dipole is

$$
\begin{equation*}
\widehat{D}:=\sum_{i=1}^{k} z_{i} \int \rho_{i}\left|f\left(\rho_{1}, \ldots, \rho_{k}\right)\right|^{2} d \rho_{1} \cdots d \rho_{k} \tag{16.81}
\end{equation*}
$$

Then (16.80) appears to be a single eigenvalue problem that is parameterized by the coefficient $R^{-3} \widehat{D}$, but the problem is that $\widehat{D}$ also depends on $f$ as well. Now we proceed to remove this recursive dependence on $f$.

Suppose that we know by symmetry that $\widehat{D}=d \mathbf{E}$ where $\mathbf{E}$ is the unit vector used in the change of variables (16.46). Then we have

$$
\begin{equation*}
d=\mathbf{E} \cdot \widehat{D}=\sum_{i=1}^{k} z_{i} \int \mathbf{E} \cdot \rho_{i}\left|f\left(\rho_{1}, \ldots, \rho_{k}\right)\right|^{2} d \rho_{1} \cdots d \rho_{k} \tag{16.82}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
and the term multiplying $\frac{1}{2} R^{-3}$ in (16.80) can be simplified as

$$
\begin{equation*}
3(\mathbf{E} \cdot \widehat{D})(\mathbf{E} \cdot D(\rho))-\widehat{D} \cdot D(\rho)=3 d(\mathbf{E} \cdot D(\rho))-d \mathbf{E} \cdot D(\rho)=2 d(\mathbf{E} \cdot D(\rho)) \tag{16.83}
\end{equation*}
$$

Therefore the eigenvalue problem (16.80) can be written

$$
\begin{equation*}
\mathcal{H}_{k}^{\rho} f_{\epsilon}+\epsilon B f_{\epsilon}=\lambda_{\epsilon} f_{\epsilon}, \tag{16.84}
\end{equation*}
$$

where we make the definitions

$$
\begin{equation*}
\epsilon=d R^{-3} \tag{16.85}
\end{equation*}
$$

and

$$
\begin{equation*}
B=\mathbf{E} \cdot D(\rho) \tag{16.86}
\end{equation*}
$$

Written in this way, we have eliminated the recursive dependence of $f_{\epsilon}$ on $f_{\epsilon}$, but at the expense of losing $R$ as an independent variable. But we can recover $R$ as needed from (16.85). Thus we have $R=(d / \epsilon)^{1 / 3}$ so that (16.82) implies

$$
\begin{equation*}
R=R_{\epsilon}=\left(\frac{1}{\epsilon} \sum_{i=1}^{k} z_{i} \int E \cdot \rho_{i}\left|f_{\epsilon}\left(\rho_{1}, \ldots, \rho_{k}\right)\right|^{2} d \rho_{1} \cdots d \rho_{k}\right)^{1 / 3} \tag{16.87}
\end{equation*}
$$

Suppose that $f_{\epsilon}=f_{0}+\epsilon f^{\prime}+\mathcal{O}\left(\epsilon^{2}\right)$ and that

$$
\begin{equation*}
\sum_{i=1}^{k} z_{i} \int E \cdot \rho_{i}\left|f_{0}\left(\rho_{1}, \ldots, \rho_{k}\right)\right|^{2} d \rho_{1} \cdots d \rho_{k}=0 \tag{16.88}
\end{equation*}
$$

Then

$$
\begin{equation*}
R_{\epsilon} \approx\left(2\left(\sum_{i=1}^{k} z_{i} \int E \cdot \rho_{i} f_{0}\left(\rho_{1}, \ldots, \rho_{k}\right) f^{\prime}\left(\rho_{1}, \ldots, \rho_{k}\right) d \rho_{1} \cdots d \rho_{k}\right)+\mathcal{O}(\epsilon)\right)^{1 / 3} \tag{16.89}
\end{equation*}
$$

Note that $f^{\prime}$ satisfies $f^{\prime} \perp f_{0}$ and

$$
\begin{equation*}
\left(\mathcal{H}_{k}^{\rho}-\lambda_{0} I\right) f^{\prime}=-B f_{0}=-\mathbf{E} \cdot D(\rho) f_{0} \tag{16.90}
\end{equation*}
$$

since $f_{0}$ is orthogonal to $B f_{0}$. The system (16.90) can be viewed as a Hartree approximation to (16.67).

Recalling (16.87), we find

$$
\begin{equation*}
R_{\epsilon} \rightarrow R_{0}:=\left(2\left(\sum_{i=1}^{k} z_{i} \int \mathbf{E} \cdot \rho_{i} f_{0}\left(\rho_{1}, \ldots, \rho_{k}\right) f^{\prime}\left(\rho_{1}, \ldots, \rho_{k}\right) d \rho_{1} \cdots d \rho_{k}\right)\right)^{1 / 3} \tag{16.91}
\end{equation*}
$$

Note the similarlity between the expression for $R_{0}$ and the one for $\mathcal{D}$ defined in (16.69); the expression (16.91) can be viewed as a Hartree approximation to (16.69). More precisely, define

$$
\begin{equation*}
\mathcal{D}^{H}=\int D(\rho) f_{0}\left(\rho_{1}, \ldots, \rho_{k}\right) f^{\prime}\left(\rho_{1}, \ldots, \rho_{k}\right) d \rho_{1} \cdots d \rho_{k} \tag{16.92}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

Then we have

$$
\begin{align*}
|\widehat{D}| & =d=\epsilon R_{\epsilon}^{3} \\
& \approx \epsilon R_{0}^{3}+o(\epsilon)  \tag{16.93}\\
& =2 \epsilon E \cdot \mathcal{D}^{H}+o(\epsilon) .
\end{align*}
$$

Suppose that

$$
\begin{equation*}
R_{\epsilon} \approx R_{0}-\epsilon R^{\prime}+o(\epsilon), \tag{16.94}
\end{equation*}
$$

so that $\epsilon=\left(R_{0}-R\right) / R^{\prime}+o(\epsilon)$. Then

$$
\begin{equation*}
d(R) \approx \frac{2\left(R_{0}-R\right)}{R^{\prime}} E \cdot \mathcal{D}^{H}+o\left(R_{0}-R\right) \tag{16.95}
\end{equation*}
$$

We get agreement with (16.70) provided that

$$
\begin{equation*}
\frac{\left(R_{0}-R\right)}{R^{\prime}} E \cdot \mathcal{D}^{H} \approx \frac{2}{R^{3}} E \cdot \mathcal{D} \tag{16.96}
\end{equation*}
$$

These results tell us that the Hartree model yields a bifurcation interpretation of the induced dipole (London dispersion) interaction. For $R>R_{0}$, the two atoms do not interact. In principle, for $R \leq R_{0}$, they might also not interact, but this state is presumably not stable. At $R=R_{0}$, a bifurcation occurs, and the induced dipole grows linearly in the parameter $R_{0}-R$ according to the expression (16.95).

### 16.6 Comparisons and conclusions

We now summarize the main conclusions of our results on the approximation of the induced dipole, a.k.a., London dispersion. We have been able to derive the asymptotic form $\left(R^{-6}\right)$ of the interaction of induced dipoles, by determining a formula for the asymptotic form of the electron distribution polarization due to the induction. This provides a length scale at which the induced dipole becomes significant in size.

The main result was the asympototic formula for the polarization due to an induced dipole, encapsulated in (16.69). We can interpret the asymptotic formula (16.70) as defining a length scale $R_{1}$ at which the induced dipole achieves an order-one effect. It should be remembered that this is dimensionally correct, in that the units of polarization are volume (Section 10.7.2). More precisely, let us define

$$
\begin{equation*}
R_{1}=(E \cdot \mathcal{D})^{1 / 3} \tag{16.97}
\end{equation*}
$$

We now compare the Hartree approximation with the exact derivation in which electron correlations were included.

Comparing (16.97) with (16.91), we can say that $R_{0}$ is the Hartree approximation of $R_{1}$. The latter is the length scale at which the induced dipole appears (according to the asymptotic theory) to develop an order-one influence. Similarly, in the Hartree theory, $R_{0}$ is the scale at which the
induced dipole influence begins. In the Hartree theory, the change in size of the induced dipole is linear in $R$, so it becomes of unit size almost instantaneously.

The Hartree approximation fails to predict the asymptotic form of the decay of the induced dipole. On the other hand, the exact theory only tells us reliable estimates of the induced dipole only for $R \gg R_{1}$. So in a sense, both theories fail to tell us reliable information for $R$ smaller than the distances $\left(R_{0}\right.$ or $\left.R_{1}\right)$ predicted by the asymptotic form of the induced dipole moment. But we can say that the Hartree approximation correctly predicts the length scale at which the induced dipole becomes significant, provided that $R_{0} \approx R_{1}$.

It would be of interest to investigate the form of the electron distribution at distances comparable to, and smaller than, $R_{1}$ (and $R_{0}$ ). This would include the effects of repulsive forces (Section 16.7) as well.

### 16.7 Repulsive molecular forces

In addition to the force of attraction due to London dispersion, it is also of interest to investigate the asymptotic form of the force of repulsion as $R$ decreases. This is often modelled via a Lennard-Jones potential as behaving like $R^{-12}$, although there is no particular physical basis for this. It is often quoted that an exponential form is more realistic, e.g., espression (2.19) in 46] and the discussion in Chapter XII of [193]. However, these are not exponentials in $R^{-1}$ but rather of the form $c_{0} e^{-c_{1} R}$ (for constants $c_{0}$ and $c_{1}$ ) and hence tend to a finite limit as $R \rightarrow 0$. Thus the expressions may only capture the form of the repulsion for large $R$ and not the behavior of the repulsive force as $R \rightarrow 0$. Thus it would be of interest to determine exactly what this is.

We do not attempt a general treatment of the repulsive force here, but instead we review calculations of the potential energey of interaction between two hydrogen atoms as given in equation (43-11) on page 343 of [193]. The dominant term in the energy as the distance $r_{A B}$ between the two hydrogen nuclei goes to zero is proportional to $1 / r_{A B}$ (and not a higher power). The repulsive effects of the electron-electron interactions is mollified by the integral against the kernel $r_{12}^{-1}=\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1}$ which acts as a smoothing operator and remain bounded as $r_{A B} \rightarrow 0$. This statement requires a proof, which we initiate subsequently with the formal definition of the smoothing operator in (16.98).

The asymptotic behavior of the interaction energy as $r_{A B} \rightarrow \infty$ is more complex to describe. It decays like exponentials in $-r_{A B}$ (with various constants) times polynomials in $r_{A B}$ [193]. Thus there is not a simple exponential decay. However, it is possible to establish bounds for decay rates. Again, certain integrals are the key to doing this.

The most complex computation that arises in the analysis of the hydrogen molecule $\left(\mathrm{H}_{2}\right)$ in [193] is an integral of the form

$$
\begin{equation*}
\iint f\left(\mathbf{r}^{1}\right) g\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2} \tag{16.98}
\end{equation*}
$$

that can be interpreted (and computed) as follows. The function

$$
\begin{equation*}
u\left(\mathbf{r}^{2}\right):=\int f\left(\mathbf{r}^{1}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} \tag{16.99}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
is a solution to the differential equation $-\Delta u=f$ together with Dirichlet boundary conditions $(u(\mathbf{r}) \rightarrow 0$ as $|\mathbf{r}| \rightarrow \infty)$ provided that $f$ decays at infinity sufficiently rapidly (which holds, say, for the wave function for the hydrogen atom). Thus

$$
\begin{equation*}
\iint f\left(\mathbf{r}^{1}\right) g\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2}=\int u\left(\mathbf{r}^{2}\right) g\left(\mathbf{r}^{2}\right) d \mathbf{r}^{2} \tag{16.100}
\end{equation*}
$$

In the particular case that $g=f$, this simplifies to

$$
\begin{align*}
\iint f\left(\mathbf{r}^{1}\right) f\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2} & =\int u\left(\mathbf{r}^{2}\right) f\left(\mathbf{r}^{2}\right) d \mathbf{r}^{2} \\
& =\int u\left(\mathbf{r}^{2}\right)\left(-\Delta u\left(\mathbf{r}^{2}\right)\right) d \mathbf{r}^{2}  \tag{16.101}\\
& =\int\left|\nabla u\left(\mathbf{r}^{2}\right)\right|^{2} d \mathbf{r}^{2}
\end{align*}
$$

Since $-\nabla \cdot(\nabla u)=f$, we can think of $\nabla u$ as an anti-derivative of $f$. Hence, the expression in (16.101) may be viewed as a type of Sobolev $H^{-1}$ norm of $f$. That is,

$$
\begin{equation*}
\left(\iint f\left(\mathbf{r}^{1}\right) f\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2}\right)^{1 / 2} \leq \sup _{v \in H_{0}^{1}} \frac{\int v(\mathbf{r}) f(\mathbf{r}) d \mathbf{r}}{\left(\int|\nabla v(\mathbf{r})|^{2} d \mathbf{r}\right)^{1 / 2}} \tag{16.102}
\end{equation*}
$$

This explains why the electron-electron repulsion is less strong than the proton-proton repulsion, as $\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right| \rightarrow 0$.

To prove (16.102), choose $v=u$ and use (16.101) twice to get

$$
\begin{align*}
\left(\iint f\left(\mathbf{r}^{1}\right) f\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2}\right)^{1 / 2} & =\frac{\int u(\mathbf{r}) f(\mathbf{r}) d \mathbf{r}}{\left(\int|\nabla u(\mathbf{r})|^{2} d \mathbf{r}\right)^{1 / 2}}  \tag{16.103}\\
& \leq \sup _{v \in H_{0}^{1}} \frac{\int v(\mathbf{r}) f(\mathbf{r}) d \mathbf{r}}{\left(\int|\nabla v(\mathbf{r})|^{2} d \mathbf{r}\right)^{1 / 2}}
\end{align*}
$$

By Sobolev's inequality, $H_{0}^{1} \subset L^{6}$; that is $\|v\|_{L^{6}} \leq c\|\nabla v\|_{L^{2}}$ (in three dimensions). Therefore Hölder's inequality yields

$$
\begin{align*}
\left(\iint f\left(\mathbf{r}^{1}\right) f\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2}\right)^{1 / 2} & \leq \sup _{v \in H_{0}^{1}} \frac{\int v(\mathbf{r}) f(\mathbf{r}) d \mathbf{r}}{\left(\int|\nabla v(\mathbf{r})|^{2} d \mathbf{r}\right)^{1 / 2}} \\
& \leq \sup _{v \in H_{0}^{1}} \frac{\|v\|_{L^{6}}\|f\|_{L^{6 / 5}}}{\left(\int|\nabla v(\mathbf{r})|^{2} d \mathbf{r}\right)^{1 / 2}}  \tag{16.104}\\
& \leq c\|f\|_{L^{6 / 5}}
\end{align*}
$$

The expression (43-10) on page 343 of 193] is of the form (16.101), with

$$
\begin{equation*}
f(\mathbf{r})=e^{-\left|\mathbf{r}-\mathbf{r}_{A}\right|-\left|\mathbf{r}-\mathbf{r}_{B}\right|} \tag{16.105}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
whereas expression (43-8) on page 342 of [193] is of the more general form (16.100) with

$$
\begin{equation*}
f(\mathbf{r})=e^{-2\left|\mathbf{r}-\mathbf{r}_{A}\right|}, g(\mathbf{r})=e^{-2\left|\mathbf{r}-\mathbf{r}_{B}\right|} \tag{16.106}
\end{equation*}
$$

Applying (16.104) to $f$ as in (16.105), we easily see that

$$
\begin{align*}
\iint f\left(\mathbf{r}^{1}\right) f\left(\mathbf{r}^{2}\right)\left|\mathbf{r}^{1}-\mathbf{r}^{2}\right|^{-1} d \mathbf{r}^{1} d \mathbf{r}^{2} & \leq c^{2}\|f\|_{L^{6 / 5}}^{2} \\
& =c^{2}\left(\int e^{-(6 / 5)\left(\left|\mathbf{r}-\mathbf{r}_{A}\right|+\left|\mathbf{r}-\mathbf{r}_{B}\right|\right)} d \mathbf{r}\right)^{5 / 3}  \tag{16.107}\\
& \leq C_{m} e^{-m\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right|}
\end{align*}
$$

for any $m<2$, where $C_{m}$ is some constant. The latter estimate follows because for all $\mathbf{r}$

$$
\begin{equation*}
e^{-\left(\left|\mathbf{r}-\mathbf{r}_{A}\right|+\left|\mathbf{r}-\mathbf{r}_{B}\right|\right)} \leq e^{-\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right|} \tag{16.108}
\end{equation*}
$$

and because $e^{-\epsilon\left(\left|\mathbf{r}-\mathbf{r}_{A}\right|+\left|\mathbf{r}-\mathbf{r}_{B}\right|\right)}$ is integrable for any $\epsilon>0$. Examining the remaining terms in equation (43-11) on page 343 of [193], we see that

$$
\begin{equation*}
W_{S}-2 W_{H} \leq C_{m} e^{-m\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right|} \text { as }\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right| \rightarrow \infty \tag{16.109}
\end{equation*}
$$

for any $m<2$, where $C_{m}$ is some constant. Here, $W_{S}$ is energy of the symmetric hydrogen pair, $W_{H}$ is the energy of a single hydrogen, so that $W_{S}-2 W_{H}$ represents the energy increase due to the bond.

Note that the calculations in Chapter XII of [193] that we have reviewed in this section do not reflect the effects of the induced dipole. Thus they only provide a rough guide to potential asymptotic effects. Being based on a Galerkin approximation (with only two basis functions), they only provide an upper-bound on the energy. That is why they miss the significant contribution from the induced dipole. This also means that we cannot prove that there is an $\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right|^{-1}$ dependence on the energy as $\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right| \rightarrow 0$, only that it is no worse than this. However, they do give an indication that there are contributions to a repulsive force which grow exponentially from zero as $\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right|$ decreases from infinity, and these can presumably balance the attractive force of the induced dipole, defining the iteratomic distance for a stable configuration. We leave as an exercise (Exercise 16.6) to include basis functions based on the limiting form of the induced dipole to see what new information this approach can provide.

We also note that the calculations in Chapter XII of [193] indicate that there could be a stronger repulsive force, proportional to $\left|\mathbf{r}_{A}-\mathbf{r}_{B}\right|^{-2}$, for two hydrogen atoms whose electrons have incompatible (meaning: the same) spins.

### 16.8 The Madelung equation

The Schrödinger equation can be recast by a simple change of variables. Write $\psi=e^{R+i S}$.
But there is a constraint [242] that must be satisfied by $\mathbf{v}=\nabla S$, namely,

$$
\begin{equation*}
\oint_{L} \mathbf{v} \cdot d \ell=2 \pi j \tag{16.110}
\end{equation*}
$$

for some $j$, for any closed loop $L$.

Draft: February 28, 2008, do not distribute

### 16.9 Exercises

Exercise 16.1 Verify the derivation of (16.39).
Exercise 16.2 Verify the computations in (16.61).
Exercise 16.3 Suppose that $X(s)$ is a smooth function from $[0,1]$ into an inner-product space with the property that $\|X(s)\|=1$ for all $s$. Let $Y$ be the derivative of $X$ at $s=0$. Show that $Y$ and $X(0)$ are orthogonal. (Hint: just expand

$$
\left.1=\|X(s)\|^{2}=\left\|X(0)+s Y+\mathcal{O}\left(s^{2}\right)\right\|=\|X(0)\|^{2}+2 s(X(0), Y)+\mathcal{O}\left(s^{2}\right) .\right)
$$

Exercise 16.4 Derive an expression for the term $R^{\prime}$ in 16.94) in terms of $f^{\prime}$, $f_{0}$, and any other required quantities. Under what conditions can you assert that $R^{\prime} \neq 0$ ?

Exercise 16.5 Carry out the computations indicated in (16.69) and (16.92) in the case of the hydrogen atom (in which case $f_{0}$ is known in closed form [193]). This requires solving (16.67) and (16.90) to determine $f^{\prime}$.

Exercise 16.6 Improve the computations in section $43 a$. in [193] by including as basis function $f^{\prime}$ computed in Exercise 16.5. Note that $f_{0}$ and $f^{\prime}$ are orthogonal.

Exercise 16.7 Carry out a full numerical simulation of the exact problem for the interaction of the hydrogen molecule (cf. Exercise 16.5 and Exercise 16.6). Consider using a Galerkin method in which the basis functions used in Exercise 16.6 are augmented in some way and an adaptive scheme is used for mesh refinement. This may be essential to control the memory requirements for this sixdimensional problem. Also explore dimensional reductions that may be possible due to symmetries of the problem.

## Chapter 17

## Continuum equations for electrostatics

The basic equations of electrostatics for a collection of charges of strength $q_{i}$ at positions $\mathbf{r}_{i}$ can be derived from the simple expression

$$
\begin{equation*}
\nabla \cdot\left(\epsilon_{0} \mathbf{e}\right)=\sum_{i} q_{i} \delta\left(\mathbf{r}-\mathbf{r}_{i}\right) \tag{17.1}
\end{equation*}
$$

where $\epsilon_{0}$ is the permittivity of the vacuum. Here $\mathbf{e}$ is the induced electric field.
We will split the charge distribution $\sum_{i} q_{i} \delta\left(\mathbf{r}-\mathbf{r}_{i}\right)$ into two parts, $\gamma+\rho$, where $\gamma$ is the part of the charge density corresponding to charge groups with net charge zero, and $\rho$ denotes the remainder of the charge density.

### 17.1 Understanding dielectrics

Dipole molecules can moderate charge by aligning with an external field. This effect is small on a small scale, but the effect of many dielectric molecules can coordinate on a large scale to achieve a substantial effect. In Figure 17.1, we present a simple example of a single charge moderated by a string of dipoles along a single line. There is a detectable depression in the average potential due to the dipole alignments.

Water is really a double dipole, with an effective dipole. In Tip5P, there are partial charges $q_{1}=-q_{2}=0.241$, positioned at distances $l_{1}=0.9572$ and $l_{2}=0.70$ from the oxygen center.

### 17.2 Dielectric materials

The dielectric properties of materials are important in many contexts [23, 224, 164]. A dielectric medium [31] is characterized by the fact that the charges are organized in local groups with net charge zero. Specifically we assume that (at least part of) the $\mathbf{r}_{i}$ and $q_{i}$ can be enumerated as $i=(j, k)$, where $j$ is the index for the group and $k$ is the index within each group, with $\mathbf{r}_{i}:=\mathbf{r}_{j}-\mathbf{r}_{j k}$ and $q_{i}:=q_{j k}$ where the $j$-th group of charges $q_{j k}$ sums to zero for all $j$ :

$$
\begin{equation*}
\sum_{k} q_{j k}=0 . \tag{17.2}
\end{equation*}
$$



Figure 17.1: Single charge $(+2)$ at zero and dipoles at $2,3, \ldots, 10$ of width 0.4 and strength 1 .

Then the expression for the charge density can be simplified as

$$
\begin{equation*}
\gamma(\mathbf{r})=\sum_{j} \sum_{k} q_{j k} \delta\left(\mathbf{r}-\mathbf{r}_{j}-\mathbf{r}_{j k}\right) \tag{17.3}
\end{equation*}
$$

The expression $\delta$ in (17.1) can be interpreted in several ways. As a first abstraction, we can take it to be the Dirac delta function, which provides a rigorous model of a point charge [215]. In [111], a mollification of the Dirac delta function is introduced, which makes it possible to reason classically about expressions involving $\delta$. This is a very useful device, and it can also be given a physical interpretation. We can think of $\delta$ representing the actual charge cloud that would be seen at a quantum scale. With this interpretation, there is an assumption being made, namely, that the local charge distribution can be represented by a single function $\delta(\mathbf{r})$, independent of the charge $q$ and independent of the atom in question. This is of course not exact, but it gives a physical interpretation to the mollifier used in [111]. A closer approximation might be obtained by letting $q$ be fractional, with positions $\mathbf{r}_{j k}$ chosen to improve the representation [160].

Let us suppose that the charge groups are homogeneous in the sense that

$$
\begin{equation*}
\mathbf{r}_{j k}=\mathcal{R}\left(\theta_{j}\right) \rho_{k} \tag{17.4}
\end{equation*}
$$

for fixed vectors $\rho_{k}$ and for some angle $\theta_{j} \in S_{2}$ (where $S_{2}$ denotes the unit 2-sphere), and further that $q_{j k}=q_{k}$ independent of $j$. This would be the case for water, for example [160]. Then

$$
\begin{align*}
\sum_{k} q_{j k} \delta\left(\mathbf{r}-\mathbf{r}_{j}-\mathbf{r}_{j k}\right) & =\sum_{k} q_{k} \delta\left(\mathbf{r}-\mathbf{r}_{j}-\mathcal{R}\left(\theta_{j}\right) \rho_{k}\right)  \tag{17.5}\\
& =\mathcal{F}\left(\theta_{j}, \mathbf{r}-\mathbf{r}_{j}\right)
\end{align*}
$$

Draft: February 28, 2008, do not distribute
where $\mathcal{F}$ is defined by

$$
\begin{equation*}
\mathcal{F}(\theta, \mathbf{r})=\sum_{k} q_{k} \delta\left(\mathbf{r}-\mathcal{R}(\theta) \rho_{k}\right) \tag{17.6}
\end{equation*}
$$

Now suppose that $\delta$ is rotationally invariant. Then

$$
\begin{align*}
\mathcal{F}(\theta, \mathcal{R}(\theta) \mathbf{r}) & =\sum_{k} q_{k} \delta\left(\mathcal{R}(\theta) \mathbf{r}-\mathcal{R}(\theta) \rho_{k}\right) \\
& =\sum_{k} q_{k} \delta\left(\mathbf{r}-\rho_{k}\right)  \tag{17.7}\\
& =\nabla \cdot \mathcal{W}(\mathbf{r})
\end{align*}
$$

with $\mathcal{W}(\mathbf{r})=\nabla \psi(\mathbf{r})$ where $\psi$ solves a Poisson equation of the form

$$
\begin{equation*}
\Delta \psi=\sum_{k} q_{k} \delta\left(\mathbf{r}-\rho_{k}\right) \tag{17.8}
\end{equation*}
$$

If $\delta$ is the Dirac $\delta$-function, then $\mathcal{W}$ is a generalized multipole expression

$$
\begin{equation*}
\mathcal{W}(\mathbf{r})=-\sum_{k} q_{k} \frac{\mathbf{r}-\rho_{k}}{\left|\mathbf{r}-\rho_{k}\right|^{3}} \tag{17.9}
\end{equation*}
$$

Then

$$
\begin{align*}
\sum_{k} q_{j k} \delta\left(\mathbf{r}-\mathbf{r}_{j}-\mathbf{r}_{j k}\right) & =\mathcal{F}\left(\theta_{j}, \mathbf{r}-\mathbf{r}_{j}\right)  \tag{17.10}\\
& =\nabla \cdot \mathcal{W}\left(\mathcal{R}\left(\theta_{j}\right)^{t}\left(\mathbf{r}-\mathbf{r}_{j}\right)\right)
\end{align*}
$$

Therefore we have an exact representation of the dielectric field $\gamma$ defined in (17.3), viz.,

$$
\begin{equation*}
\gamma(\mathbf{r})=\sum_{j} \nabla \cdot \mathcal{W}\left(\mathcal{R}\left(\theta_{j}\right)^{t}\left(\mathbf{r}-\mathbf{r}_{j}\right)\right) \tag{17.11}
\end{equation*}
$$

### 17.3 Polarization field

Now, let us suppose that there is an additional charge density $\rho$ that is not related to the dielectric, and thus the equations (17.1) would be written

$$
\begin{equation*}
\nabla \cdot\left(\epsilon_{0} \mathbf{e}\right)=\rho+\sum_{j} \nabla \cdot \mathcal{W}\left(\mathcal{R}\left(\theta_{j}\right)^{t}\left(\mathbf{r}-\mathbf{r}_{j}\right)\right) \tag{17.12}
\end{equation*}
$$

Let us define the polariztion $\mathbf{p}$ by

$$
\begin{equation*}
\mathbf{p}(\mathbf{r})=-\sum_{j} \mathcal{W}\left(\mathcal{R}\left(\theta_{j}\right)^{t}\left(\mathbf{r}-\mathbf{r}_{j}\right)\right) \tag{17.13}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
and set $\mathbf{d}=\epsilon_{0} \mathbf{e}+\mathbf{p}$. Then

$$
\begin{equation*}
\nabla \cdot \mathbf{d}=\epsilon_{0} \nabla \cdot \mathbf{e}+\nabla \cdot \mathbf{p}=\gamma+\nabla \cdot \mathbf{p}=\rho \tag{17.14}
\end{equation*}
$$

The electric field $\mathbf{e}$ thus satisfies $\epsilon_{0} \mathbf{e}=\mathbf{d}-\mathbf{p}$.
The directions $\theta_{j}$ that determine the polarization tend on average to cause $\mathbf{p}$ to line up with the induced field. In fact, in a thermalized system, there will be fluctuations in the angles $\theta_{j}$, and we can only talk about the mean angles. Debye [50] suggested that we can write $\widetilde{\mathbf{p}}=\left(\epsilon-\epsilon_{0}\right) \widetilde{\mathbf{e}}$ where $\epsilon$ denotes an effective permittivity. Here, $\widetilde{\mathbf{p}}$ (resp., $\widetilde{\mathbf{e}}$ ) denotes a temporal average over a timescale that is long with respect to the basic thermal motions. The latter occur on the order of fractions of picoseconds, so we could imaging a time average of the order of picoseconds.

If one is uncomfortable with the Debye ansatz, we can define $\epsilon$ as follows. We decompose $\mathbf{p}$ into one part in the direction $\mathbf{e}$ and the other perpendicular. That is, we write

$$
\begin{equation*}
\mathbf{p}=\left(\epsilon-\epsilon_{0}\right) \mathbf{e}+\zeta \mathbf{e}^{\perp}, \tag{17.15}
\end{equation*}
$$

where $\epsilon$ is defined by

$$
\begin{equation*}
\epsilon=\epsilon_{0}+\frac{\mathbf{p} \cdot \mathbf{e}}{\mathbf{e} \cdot \mathbf{e}} \tag{17.16}
\end{equation*}
$$

with the appropriate optimism that $\mathbf{p}=0$ when $\mathbf{e}=0$. That is, $\epsilon-\epsilon_{0}$ reflects the correlation between $\mathbf{p}$ and $\mathbf{e}$. As defined, $\epsilon$ is a function of $\mathbf{r}$ and $t$, and potentially singular. However, Debye postulated that a suitable average

$$
\begin{equation*}
\tilde{\epsilon}=\epsilon_{0}+\left\langle\frac{\mathbf{p} \cdot \mathbf{e}}{\mathbf{e} \cdot \mathbf{e}}\right\rangle \tag{17.17}
\end{equation*}
$$

should be well behaved. For simplicity, we will drop the tildes and think from now on that everything represents temporal, or spatial, averages.

The expression (17.17) provides an operational definition for a computationally determined dielectric constant. That is, in a molecular dynamics computation, one can define a local dielectric constant by averaging the correlation coefficient

$$
\begin{equation*}
\frac{\mathbf{p} \cdot \mathbf{e}}{\mathbf{e} \cdot \mathbf{e}} \tag{17.18}
\end{equation*}
$$

over space, over certain molecules, and/or time. This correlation coefficient need not be positive, so it is conceivable that $\widetilde{\epsilon}<\epsilon_{0}$, and we could even have $\widetilde{\epsilon}<0$.

### 17.4 Frequency dependence

The relationship proposed by Debye between $\mathbf{p}$ and $\mathbf{e}$ depends on frequency. In particular, we have

$$
\begin{equation*}
\epsilon(\nu)=\epsilon_{0}+\frac{\epsilon_{1}-\epsilon_{0}}{1+\nu^{2} \tau^{2}} \tag{17.19}
\end{equation*}
$$

where $\tau$ is a characteristic time associated with the dielectric material.
This relationship has been verified extensively by experimental data, as indicated in Figure 17.2 , But it is instructive to review the theoretical derivation that Debye gave to justify the behavior.


Figure 17.2: This figure is reproduced from [128] and we intend to get the permission of the authors and journal.

### 17.5 Spatial frequency dependence

We are interested in electric fields which are not time varying (i.e., $\nu=0$ ) but rather spatially varying. It is easy to see that the driving electrical fields $\rho$ generated by proteins have high frequencies. Salt bridges involve charge alternations on the order of a few Angstroms. And polar sidechains correspond to even higher frequencies, although at a smaller amplitude.

In addition, the dielectric coefficient varies by a factor of nearly one hundred from inside the protein to the dielectric bulk away from the surface. This forces a kink in the electric field in the vicinity of the boundary in the direction normal to the surface of the protein. This further engenders high frequency components in the electric field. Thus there are high-frequency components in the electric field in both the direction normal to the "surface" of the protein as well as in directions along the "surface."

On the other hand, it is clear that the dielectric response has to go to zero for high frequencies. If the electric field varies at a spatial frequency whose wavelength is smaller than the size of a water molecule, the water molecule feels a diminished effect of that field component. Therefore, the dielectric coefficient must be a function of spatial wave number and go to zero for high frequencies.

Thus we use the ansatz that the dielectric properties depend on spatial wave number $\xi$ proportional to a factor $\kappa$ where

$$
\begin{equation*}
\kappa(\xi)=\epsilon_{0}+\frac{\epsilon_{1}-\epsilon_{0}}{1+|\lambda \xi|^{2}} \tag{17.20}
\end{equation*}
$$

where $\lambda$ is the length scale of the transition from one value of the dielectric to the other. In general, $\lambda$ could be a matrix, allowing for anisotropy. But for the time being we will think of it as a scalar.

Empirical evidence suggests that the length scale $\lambda$ should be approximately 1.7 Ångstroms [217] when modelling water in a bulk environment.

We have anticipated that the dielecric coefficient may depend on space (and time), so we will be interested in cases where $\lambda$ depends on the spatial variable $\mathbf{r}$, and in some cases the length scale will tend to infinity. For this reason, we introduce $\nu=1 / \lambda$, and write

$$
\begin{equation*}
\kappa(\mathbf{r}, \xi)=\epsilon_{0}+\nu(\mathbf{r}) \frac{\epsilon_{1}-\epsilon_{0}}{\nu(\mathbf{r})+|\xi|^{2}} \tag{17.21}
\end{equation*}
$$

where $\nu(\mathbf{r})$ is a spatial frequency scale. However, for simplicity we assume that the model (17.20) is sufficient for the moment.

### 17.5.1 Poisson-Debye equation: bulk case

We can expand $\mathbf{e}$ and $\mathbf{p}$ in a Fourier series and use the Debye-like relationship (17.20) to relate the resulting coefficients in the series. That is, we have (using the inverse Fourier transform)

$$
\begin{equation*}
\frac{1}{(2 \pi)^{3}} \int_{\boldsymbol{R}^{3}} e^{-i \mathbf{r} \cdot \xi} \kappa(\xi) \widehat{\mathbf{e}}(\xi) d \xi=\mathbf{p}(\mathbf{r}), \tag{17.22}
\end{equation*}
$$

where here and subsequently we use the notation $\widehat{u}$ to denote the Fourier transform of a function $u$ :

$$
\begin{equation*}
\widehat{u}(\xi):=\int_{\mathbb{R}^{3}} e^{i \xi \cdot \mathbf{r}} u(\mathbf{r}) d \mathbf{r} . \tag{17.23}
\end{equation*}
$$

Therefore the basic equation is

$$
\begin{equation*}
\nabla \cdot\left(\frac{1}{(2 \pi)^{3}} \int_{\boldsymbol{R}^{3}} e^{-i \mathbf{r} \cdot \xi} \kappa(\xi) \widehat{\mathbf{e}}(\xi) d \xi\right)=4 \pi \rho(\mathbf{r}) \tag{17.24}
\end{equation*}
$$

We can write $\mathbf{e}=\nabla \phi$ using Maxwell's equations. Therefore $\widehat{\mathbf{e}}(\xi)=i \xi \widehat{\phi}(\xi)$. Therefore, (17.23) becomes

$$
\begin{equation*}
\nabla \cdot\left(\frac{1}{(2 \pi)^{3}} \int_{\mathbb{R}^{3}} e^{-i \mathbf{r} \cdot \xi} \kappa(\xi) i \xi \widehat{\phi}(\xi) d \xi\right)=4 \pi \rho(\mathbf{r}) \tag{17.25}
\end{equation*}
$$

Taking the Fourier transform (17.23) of (17.25) provides the simple relation

$$
\begin{equation*}
\widehat{\phi}(\xi)=\frac{4 \pi \widehat{\rho}(\xi)}{|\xi|^{2} \kappa(\xi)} \tag{17.26}
\end{equation*}
$$

which can be used to compute $\phi$ (and thus e) from $\rho$.
The expression (17.24) can be simplified in certain limits. We have

$$
\begin{equation*}
\frac{1}{(2 \pi)^{3}} \int_{\mathbb{R}^{3}} e^{-i \mathbf{r} \cdot \xi} \kappa(\xi) \widehat{\mathbf{e}}(\xi) d \xi \approx \epsilon_{j} \mathbf{e}(\mathbf{r}) \tag{17.27}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute
where $j=1$ when $\mathbf{e}$ is very smooth and $j=0$ when e consists of only high frequencies. However, for general fields e it is not possible to approximate the Fourier integral in this way. Thus we cannot think of (17.24) as a partial differential equation, except approximately in special cases.

When a non-dielectric material is immersed in a dielectric (e.g., a protein in water), it might be plausible to approximate (17.27) with $j=1$ in the dielectric, switching to $j=0$ at the interface of the non-dielectric material (which introduces high frequencies due to the abrupt change in material). This leads to the standard Poisson equation with a spatially varying permittivity $\epsilon_{j}$ that jumps from $j=1$ in the dielectric to $j=0$ in the non-dielectric material; this is often used to model macromolecular systems in solvent [121. However, it is not clear what to do when very small non-dielectric objects, such as nanotubes [] are introduced into a dielectric. The scale of a nanotube is so small that there would be almost no $\epsilon_{0}$ region in such models, so that any predictions of electrostatics would be essentially the same as if there were pure dielectric. It is possible to introduce a spatially varying permittivity that changes more smoothly between the two extremes [217], but this does not capture accurately the behavior of the wave-number dependence.

The characteristic scale $\lambda$ represents a correlation length relating the way changes in the dielectric influence each other spatially. When the dielectric molecules are constrained, for example, at a material boundary, the characteristic scale $\lambda$ increases. This is because the dielectric molecules lose freedom near a wall, and thus changes propagate further than in bulk. It is also clear that these changes may be anisotropic, with changes parallel to the wall more affected than perpendicular to the wall. Such changes near an interface could cause $\lambda$ to increase effectively to infinity at the surface of the bounding material. Thus it might be reasonable to view the kernel $\kappa$ in this case as continuous across material boundaries.

The equation (17.24) involves a Fourier integral operator [58]. Due to the special form of $\kappa(\nu)$, it is possible to write (17.24) as a fourth-order elliptic partial differential operator for the potential $\phi$ :

$$
\begin{equation*}
\nabla \cdot\left(\left(\epsilon_{1}-\epsilon_{0} \lambda^{2} \Delta\right) \nabla \phi\right)=\left(1-\lambda^{2} \Delta\right) \rho \tag{17.28}
\end{equation*}
$$

provided that $\lambda$ is constant. However, if $\lambda$ is a function of $\mathbf{r}$ this is no longer valid. Also, the limit $\lambda \rightarrow \infty$ is harder to interpret in this setting.

### 17.5.2 Response to a point charge

The first question to ask with the model (17.24) is what the electric field (or potential) looks like for a single charge $\rho=4 \pi \delta$. More precisely, (17.24) with $\rho=4 \pi \delta$ defines a family of potentials $\phi_{\epsilon_{r}, \lambda}$ for any given $\epsilon_{r}$ and $\lambda$, where $\epsilon_{r}=\epsilon_{1} / \epsilon_{0}$. Thus a simple change of variables implies that

$$
\begin{equation*}
\phi(\mathbf{r})=\frac{\epsilon_{0}}{\lambda} \phi_{\epsilon_{r}, 1}(\mathbf{r} / \lambda) \tag{17.29}
\end{equation*}
$$

where $\phi_{\epsilon_{r}, 1}$ is defined using the kernel $\kappa=1+\frac{\epsilon_{r}-1}{1+\xi^{2}}$. In the limit $\lambda \rightarrow \infty$ we find $\phi_{\infty}=c / \epsilon_{0} r$.
The computation of $\phi_{\epsilon_{r}, 1}$ requires us to consider the Fourier transform of a radially symmetric function $u$ (which is itself radially symmetric). The following formula holds:

$$
\begin{equation*}
|\xi| \widehat{u}(\xi):=4 \pi \int_{0}^{\infty} r u(r) \sin (|\xi| r) d r \tag{17.30}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute


Figure 17.3: Comparison of empirical permittivity formula in [219] (dotted line) with the model (17.24) (solid line) where $\kappa$ is defined in (17.20) with $\lambda=1.7$. Plotted are the ratios of the effective permittivity experienced by a dipole to the vacuum permittivity as a function of the separation distance in $\AA$, where the bulk permittivity is that of water.
where $u(r)$ means $u(\mathbf{r})$ with $r=|\mathbf{r}|$.
Thus the expression $\epsilon_{0} \phi_{\infty} / \phi_{\lambda}=c / r \phi_{\lambda}$ is depicted in Figure 17.3. This indicates that $\phi_{\lambda}$ behaves like $\phi_{\infty}$, the vacuum potential for a point charge, near the point charge. At a distance of $20 \AA$ from the charge, the potential has been reduced by a factor of nearly eighty due to the effect of the dielectric. However, note that the dielectric does not take full effect until a distance of over $10 \lambda$ is reached.

There is a physical interpretation for the suppression of the dielectric effect near the point charge. In the immediate vicinity of the charge, water molecules are strongly structured. In particular, the immediate layer of hydrogen bonds between the point charge and the first layer of water molecules would be expected to be nearly constant. The immobilization of such a layer of water immediately surrounding the charge leads to significant water structuring in subsequent layers. Thus only when one reaches a significant distance (e.g., 10 $\lambda$ ) are the water molecules without noticeable effect from the point charge.

It is useful to observe that the permittivity of free space $\epsilon_{0}$ can be written in convenient units as $\epsilon_{0}=0.24 q_{e}^{2}(\mathrm{kcal} / \mathrm{mol})^{-1} \mu \mathrm{~m}^{-1}$ where $-q_{e}$ is the charge of the electron.

### 17.5.3 Non-local relationship between pand e

A model has been used by different people [145, 137, 65, 42] to account for the frequency dependence of the (zero temporal frequency) dielectric relationship. It is often expressed as a non-local dependence of the polarization on the electric field and written in the form

$$
\begin{equation*}
\mathbf{p}(\mathbf{r})=\int K\left(\mathbf{r}, \mathbf{r}^{\prime}\right) \mathbf{e}\left(\mathbf{r}^{\prime}\right) d \mathbf{r}^{\prime} \tag{17.31}
\end{equation*}
$$

where the averaging kernel $K$ satisfies

$$
\begin{equation*}
K\left(\mathbf{r}, \mathbf{r}^{\prime}\right)=K\left(\mathbf{r}-\mathbf{r}^{\prime}\right)=\int e^{i \mathbf{k} \cdot\left(\mathbf{r}-\mathbf{r}^{\prime}\right)} \kappa(\mathbf{k}) d \mathbf{k} \tag{17.32}
\end{equation*}
$$

with the expression $\kappa$ representing the Debye-like frequency dependence (17.20). Taking Fourier transforms, we see that the "non-local" model (17.20) is the same as (17.22). However, it is not possible to represent the mollifier $K$ as an ordinary function. Clearly, $K=\epsilon_{0} \delta+\left(\epsilon_{1}-\epsilon_{0}\right) \widetilde{K}$ where

$$
\begin{equation*}
\widetilde{K}(\mathbf{r})=\int e^{i \mathbf{k} \cdot \mathbf{r}} \tilde{\kappa}(\mathbf{k}) d \mathbf{k} \tag{17.33}
\end{equation*}
$$

and $\tilde{\kappa}$ is defined by

$$
\begin{equation*}
\tilde{\kappa}(\mathbf{k})=\frac{1}{1+|\mathbf{k}|^{2} \lambda^{2}} . \tag{17.34}
\end{equation*}
$$

We easily identify $\tilde{\kappa}$ as the Fourier transform of the fundamental solution of the Laplace operator $1-\lambda^{2} \Delta$, so that

$$
\begin{equation*}
\widetilde{K}(\mathbf{r})=\frac{\lambda e^{-|\mathbf{r}| / \lambda}}{4 \pi|\mathbf{r}|} \tag{17.35}
\end{equation*}
$$

Although this expression appears singular, we realize it is less singular than the Dirac $\delta$-function, which simply evaluates a function at a point instead of averaging. The exponential decay insures that the averaging is fairly local in nature. The kernel for the non-local expression for the polarization can be written formally (in the sense of distributions [215]) as

$$
\begin{equation*}
K(\mathbf{r})=\epsilon_{0} \delta+\frac{\lambda\left(\epsilon_{1}-\epsilon_{0}\right) e^{-|\mathbf{r}| / \lambda}}{4 \pi|\mathbf{r}|} \tag{17.36}
\end{equation*}
$$

More precisely, we have

$$
\begin{equation*}
\mathbf{p}(\mathbf{r})=\epsilon_{0} \mathbf{e}(\mathbf{r})+\frac{\lambda\left(\epsilon_{1}-\epsilon_{0}\right)}{4 \pi} \int \frac{e^{-\left|\mathbf{r}-\mathbf{r}^{\prime}\right| / \lambda}}{\left|\mathbf{r}-\mathbf{r}^{\prime}\right|} \mathbf{e}\left(\mathbf{r}^{\prime}\right) d \mathbf{r}^{\prime} \tag{17.37}
\end{equation*}
$$

### 17.6 The Poisson-Debye equation: general case

A general relationship between $\mathbf{p}$ and $\mathbf{e}$ of the form $\mathbf{p}=\mathcal{P} \mathbf{e}$ leads to an equation for the electric field of the following form:

$$
\begin{equation*}
\nabla \cdot \mathcal{P} \nabla \phi=\rho \tag{17.38}
\end{equation*}
$$

Draft: February 28, 2008, do not distribute

We can write a general relationship between the electric field and the polarization vector using an operator $\mathcal{P}$ (defined component-wise) given by

$$
\begin{equation*}
\mathcal{P} v(\mathbf{r})=\int_{\boldsymbol{R}^{3}} \int_{\mathbb{R}^{3}} e^{i \mathbf{k} \cdot\left(\mathbf{r}-\mathbf{r}^{\prime}\right)} \kappa\left(\mathbf{k}, \mathbf{r}, \mathbf{r}^{\prime}\right) v\left(\mathbf{r}^{\prime}\right) d \mathbf{k} d \mathbf{r}^{\prime} \tag{17.39}
\end{equation*}
$$

where we define the symbol $\kappa$ by the Debye-like relationship

$$
\begin{equation*}
\kappa\left(\mathbf{k}, \mathbf{r}, \mathbf{r}^{\prime}\right)=\epsilon_{0}+\frac{\epsilon_{1}-\epsilon_{0}}{1+\left|\lambda\left(\mathbf{r}, \mathbf{r}^{\prime}\right) \mathbf{k}\right|^{2}} \tag{17.40}
\end{equation*}
$$

where we have assumed that the length scale is allowed to vary as a function of the spatial coordinate.
Our justification for such an approach is based on the fact that models in which $\lambda$ is constant but of different size depending on the context have been successful in modelling dielectric behavior near boundaries [217]. At boundaries, $\lambda$ increases to infinity, so it is perhaps easier to express $\kappa$ in terms of $\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)=1 / \lambda$ :

$$
\begin{equation*}
\kappa\left(\mathbf{k}, \mathbf{r}, \mathbf{r}^{\prime}\right)=\epsilon_{0}+\frac{\left(\epsilon_{1}-\epsilon_{0}\right) \nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)^{2}}{\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)^{2}+|\mathbf{k}|^{2}} \tag{17.41}
\end{equation*}
$$

In this description, $\kappa$ tends to a well defined limit as $\nu \rightarrow 0$. However, note that the limit depends on whether $|\mathbf{k}|=0$ or not. In particular, $\kappa\left(0, \mathbf{r}, \mathbf{r}^{\prime}\right)=\epsilon_{1}$ for all $\nu$. However, for $|\mathbf{k}|>0, \kappa\left(0, \mathbf{r}, \mathbf{r}^{\prime}\right)$ tends to $\epsilon_{0}$ as $\nu$ tends to zero.

The dependence on the length scale $\lambda$ on the distance from a hydrophobic surface was discussed in [217]. In terms of $\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)=1 / \lambda$ it is given by

$$
\begin{equation*}
\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)=\frac{1}{\lambda_{b}} \prod_{j}\left(1-e^{-\left(\left|\mathbf{r}-\mathbf{r}_{j}\right|+\left|\mathbf{r}^{\prime}-\mathbf{r}_{j}\right|\right) / \lambda_{b}}\right) \tag{17.42}
\end{equation*}
$$

where $\lambda_{b}$ denotes the bulk coordination length and hydrophobic entities are located at the points $\mathbf{r}_{j}$.

This can be interpreted in the following ways. First of all, it says that $\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right) \approx 1 / \lambda_{b}$ whenever either $\mathbf{r}$ or $\mathbf{r}^{\prime}$ is far from the hydrophobes. In particular, if there are no hydrophobes at all (the bulk case), we have $\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)=1 / \lambda_{b}$.

When both $\mathbf{r}$ and $\mathbf{r}^{\prime}$ are near a hydrophobe, say $\mathbf{r}_{j}$, then $e^{-\left(\left|\mathbf{r}-\mathbf{r}_{j}\right|+\left|\mathbf{r}^{\prime}-\mathbf{r}_{j}\right|\right) / \lambda_{b}} \approx 1$. This means that $\nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right) \approx 0$ for such such $\mathbf{r}, \mathbf{r}^{\prime}$. In such a case, we find $\kappa\left(\mathbf{k}, \mathbf{r}, \mathbf{r}^{\prime}\right) \approx \epsilon_{0}$.

This approach does not explicitly involve a definition of the interior or boundary of the protein. This could be done using the function $\nu$, for example by defining

$$
\begin{equation*}
\Omega=\{\mathbf{r}: \nu(\mathbf{r}, \mathbf{r})<\tau\} \tag{17.43}
\end{equation*}
$$

for some fixed tolerance $\tau>0$, e.g., $\tau=0.1$. But we do not explicitly need an expression for such a domain, and in particular there is no need to provide a mesh that is senstitive to the boundary $\partial \Omega$. This simplified representation of the effect of the protein in modulating the dielectric may compenstate for the fact that the resulting model is more complex.

Draft: February 28, 2008, do not distribute

### 17.7 Solving the Poisson-Debye equation

The Poisson-Debye equation (17.38) involves the Fourier Integral Operator $\mathcal{P}$ defined by (17.39). Clearly, the symbol $\kappa\left(\mathbf{k}, \mathbf{r}, \mathbf{r}^{\prime}\right) \geq \epsilon_{0}$ for all $\mathbf{r}, \mathbf{r}^{\prime}, \mathbf{k}$ (recall that $\epsilon_{1}>\epsilon_{0}$ ). Similarly, the symbol of $\mathcal{P}$ is bounded above by a fixed constant for all $\mathbf{r}, \mathbf{r}^{\prime}, \mathbf{k}$. Thus a standard variational theory gives existence, uniqueness and stability for the Poisson-Debye equation (17.38).

### 17.7.1 Numerical methods for FIO's

Curvlets have been applied to FIO's [39] but efficiency seems to require smooth coefficients.
In our case,

$$
\begin{equation*}
\kappa\left(\mathbf{k}, \mathbf{r}, \mathbf{r}^{\prime}\right)=\epsilon_{0}+\frac{\epsilon_{1}-\epsilon_{0}}{1+\left(|\mathbf{k}| / \nu\left(\mathbf{r}, \mathbf{r}^{\prime}\right)\right)^{2}} \tag{17.44}
\end{equation*}
$$

is homogeneous in $|\mathbf{k}| / \nu$ and thus singular when $\nu \rightarrow 0$.
Therefore, direct application does not provide a sparse representation via curvelets.

### 17.8 Modeling DNA

The environment of DNA has been modeled as a dielectric cylinder with a given surface charge density immersed in an electrolite [185.

## Chapter 18

## Water structure

Each water molecule can potentially form hydrogen bonds with four other molecules, and typicaly does so when frozen, when water ice takes the form of a perfect lattice with all possible hydrogen bonds satisfied. However, the exact bonding structure of liquid water is still being studied and debated [1, 248, 228.

### 18.1 Understanding dielectrics

Dipole molecules can moderate charge. Induced dipole interactions lead to van der Waals forces.

### 18.2 Tetrahedral structure of water

Water is really a quadrupole, as indicated in Figure 18.3. The two positive charge regions provided by the hydrogens are matched by two negatively chaged regions oriented in a plane perpendicular to the plane of the water molecule. These negative lobes have been referred to variously as rabbit ears [144] and squirrel ears [161]. The deviation from a spherical electron density is actually quite small, yet distinct enough to localize hydrogen bonding.

Recent research has suggested that water is typically involved in only about half of its possible hydrogen bonds [249, 228].

### 18.3 Structural water in proteins

There is growing evidence that water plays some sort of structural role in proteins.
The frequent appearance of like-charged sidechain pairs has been noted previously. [159, 239] The stability of such repellent pairs comes from their interaction with stable clusters of water molecules. 159 Such favorable close approach of charged groups in water, due to polarization of the intervening water, has been demonstrated by ab initio quantum mechanical calculations and by observation of proteins in the PDB. [239] The role of backbone solvation and electrostatics in generating preferred peptide backbone conformations has recently been studied. 10] The effect of


Figure 18.1: Single charge $(+2)$ at zero and dipoles at $2,3, \ldots, 10$ of width 0.4 and strength 1 . Sharp peaks on the graph are located at the positive charge of the dipole, and the valleys are at the negative charge.


Figure 18.2:


Figure 18.3: In Tip5P, see (d), $q_{1}=-q_{2}=0.241, l_{1}=0.9572, l_{2}=0.70, \theta=104.52, \phi=109.47$.
neighboring residues on backbone conformation has been attributed to peptide backbone solvation and shielding from water of peptide groups by adjacent sidechains. 11]

### 18.4 Polarizable water models

Standard water models, such as Tip5P [149, do not allow for the polarizability of water ( $\alpha \approx$ $1.47 \AA^{3}$ ). More complex models incorporate this explicitly [34]. Apparently, the water dipole can be significantly enhanced by the polarization inherent in a condensed phase [100]. For a polarizability of $\alpha=1 \AA^{3}$, a charge density of $0.2084 q_{e}$ per $\AA^{2}$ would result in a change of one Debye. The dipole moment of an isolated water molecule is about 1.9 Debye and increases to 2.5 Debye in the condensed phase [100]. This would be consistent with a typical charge density of $0.1 q_{e}$ per $\AA^{2}$.

### 18.5 A two-D water model






Figure 18.4: A two-D water model.

## Chapter 19

## Hydrophobic effect

The hydrophobic effect [234, 22] was proposed in the 1950's [235] as a major contributor to protein structure. However, it is only recently that the detailed nature of hydrophobic forces have been understood. Indeed, the dehydron can be viewed as a particular type of hydrophobic effect.

In two recent papers, further understanding of hydrophobic forces have been provided [53, 54]. In the second, it is seen that the role of hydrophobic modulation of solvent dielectric is critical to the hydrophobic force [54].

## Chapter 20

## Disorder in Protein Structure

Disorder is a common feature of proteins. It can range from a small amount of flexibility to complete disorder. It has been argued that complete disorder is the common case for the vast majority of possible amino acid sequences []. Thus biologically relevant proteins represent a highly selective sampling.

One example of disorder is exhibited by prions.
The review [27] lists several programs that predict disorder based on protein sequence.

## Chapter 21

## Geckos' feet

It has recently been discovered that the feet of geckos are able to attach to surfaces by a type of van der Waals interaction (9].

## Chapter 22

## Notes

The following notes are intended to complement the discussion in the indicated chapters.

## Chapter 2

We are indebted to the noted photographer Ron Scott for the suggestion of the grain of sand in an oyster to explain epidiorthotric effects. Regarding epiorthotric forces in psychosocial contexts, we have in mind something like a sense of insecurity that often drives highly talented people. Groucho Marx expressed this via the conundrum that he would never join a club that would stoop so low as to have him as a member. A prolific inventor, Larkin Burneal Scott (1917-1991) expressed it in a jest as follows: "I don't have an inferiority complex; I am inferior."

## Chapter 3

It is useful to reflect on the history of the development of the understanding of different types of electrostatic forces and structural features of proteins. In Table 22.1, we present these together with approximate dates of emergence and representative citations. Dates should be considered only approximate and not a definitive statement about priority. Similarly, the references presented are only intended to give a sense of some central contributions.

Chapter 4
Chapter 7
Chapter 6
Chapter 14
Chapter 8
Chapter 17
Chapter ??
Chapter 16

| Force/structure name | emergence date(s) | selected references |
| :---: | :---: | :--- |
| van der Waals | 1893 |  |
| Keesom dipole | 1912 |  |
| covalent bond | 1916 |  |
| Debye induction | 1920 |  |
| hydrogen bond | $1920-33-44$ |  |
| London dispersion | 1930 |  |
| alpha/beta structure | 1936 | 53 |
| hydrophobic force | $1954-$ present | $[55$ |
| cation- $\pi$ interaction | 1996 | $[77,71,79]$ |
| dehydron | 2003 |  |

Table 22.1: A brief history of the development of understanding of the principle bonds that are significant for protein structure and interaction.

## Chapter 18

Chapter 19
Chapter ??
Chapter 9
Chapter 21

## Chapter 23

## Glossary

The following definitions are intended only as an informal description. Other sources should be consulted for definitive meanings. Our objective is simply to provide a rapid, if approximate, way for the reader to return to the main part of the book without needing to resort to another text or an on-line resource. The definitions often depend on other definitions; terms that are used in one definition and are defined separately are inidicated in a separate font like this. Many terms are defined fully in the text, and the index provides pointers to these explications. If a term is not found here, consult the index next.

An aliphatic chain is a linear sequence of carbonaceous $\left(\mathrm{CH}_{n}\right)$ groups.
Allostery is derived from the Greek meaning "other shape." An allosteric effect is one that is induced at one part of a molecule by an effect (e.g., ligand binding) at another.

An amino acid is a molecule that forms the basis of the sidechain of a protein.
An amide group is the $N-H$ pair in a peptide bond as shown in Figure 14.1 .
A protein antagonist $\qquad$
A beta-hairpin (or $\beta$-hairpin) is ....
The backbone is the name for the sequence of $C_{\alpha}$ carbons in a protein chain, that is, the lower left and upper right C's in Figure 14.1 .

A capsid is the outer coat of a virus, typically a protein complex.
A carbonyl group is the $C=O$ pair in a peptide bond as shown in Figure 14.1 .
C. elegans, or more completely Caenorhabditis elegans, is a worm.

A chain is an individual protein in a protein complex
The conformation of a protein is the three-dimensional shape that it adopts. A change in conformation means that a new shape is adopted.

A covalent bond is an electrostatic bond in which the electrons of different atoms become intertwined and can no longer be identified as belonging to a distinct atom.

A crystal is a lattice of objects, such as proteins, that can form under certain conditions. The repeated (periodic) structure in particular allows them to be imaged using X-rays.

A dimer is an object made of two monomers, typically the same or very similar.
A protein domain is the basic unit of tertiary strucutre. A single protein can consist of a single domain or many domains. A protein fold is a synonym for domain.

DNA is the acronym for DioxyriboNucleicAcid.
E. coli, or more completely Escherichia coli, is a bacterium commonly found in food.

Electron density refers to the fact that electrons can not be located exactly, but rather a probabilistic description is used in quantum mechanics to describe where they spend a fraction of their time.

Endocytosis is a process of cell surface folding that ingests a substance.
The term epidiorthotric was introduced in Section [2.4 to refer to an effect that occurs as the result of a repair of a defect.

An epitope is a small region of a protein involved in a binding event, such as the part of an antigen where an antibody binds.

A protein fold is the basic unit of tertiary strucutre. A single protein can consist of a single fold or many folds. A protein domain is a synonym for fold.

Homo sapien (sometimes written H. sapien) is the formal biological name for a human being.
Something is hydrophobic if it repels water.
Hydrophobic packing refers to the placement of carbonaceous groups in the vicinity.
A ligand is anything that binds to something.
A moity is a portion of a whole, usually with a defined property or structure.
A monomer is a single unit, e.g., a peptide, that can join with one or more other monomers of the same type to form a larger complex. The term can be used for something as small as a single molecule or as large as a protein.

A motif is a characteristic feature.
A multimer is something formed from small units, often called monomers, cf. polymer.
Mus musculus is the formal biological name for the common house mouse.
A noncovalent bond, or interaction, is an electrostatic interaction in which the electrons of the atoms in the bond remain sufficiently apart to remain identified with their atoms, even though they may be strongly correlated.

A packing defect is a defficiency in wrapping, that is, a lack of adequate carbonaceous groups in the vicinity.

A partial charge is a model to account for the fact that electrons may be unevenly distributed in a molecule.

A peptide is the basic unit of a protein.
Something is polar if it has a positive partial charge on one side and a negative partial charge on the other.

A polymer is something formed from small units, often called monomers, cf. multimer.
To polymerize is to form a larger system from small units, e.g., a chain such as a protein.
A polypeptide is the result of polymerizing peptides.
A graph has power-law distribution if the number of vertices with degree $k$ is rougly $k^{-\gamma}$ for a fixed $\gamma$.

The primary structure of a protein is the sequence of its amino acids.
A protein complex is a collection of two or more proteins that are bound together.
The quartenary structure of a protein system is the three-dimensional arrangement of the different protein chains of the system.

Draft: February 28, 2008, do not distribute

An amino acid residue is part of the amino acid that remains when it is cleaved to form a sidechain on a protein.

RNA is the acronym for RiboNucleicAcid.
A graph is scale free if obeys a power-law distribution, that is, if the number of verticeswith degree $k$ is rougly $k^{-\gamma}$ for a fixed $\gamma$.

A small world graph is one in which most of the vertices have low degree, such as a graph with a power-law distribution.

The secondary structure of a protein is the set of alpha helices and beta sheets.
A sidechain is another name for residue.
A protein is soluble if it can form a stable and functional form in water.
A steric effect is one that involves the shape of an object (the word steric derives from the Greek word for 'shape;' also see the explanation of allostery).

Protein structure is hierarchical, involving primary, secondary, tertiary, and, for a protein complex, quartenary structure.

A subunit of a protein complex is one of the proteins in the collection.
The tertiary structure of a protein is the three-dimensional shape of the fully folded proteins. A tetramer is an object made of four monomers, typically the same or very similar.
The three-letter code for RNA and DNA is the sequence of three letters that code for a particular amino acid.

A trimer is an object made of three monomers, typically the same or very similar.
UWHB is the acronym for underwrapped hydrogen bond, a.k.a. dehdydron.
Vicinal means 'in the vicinity' or nearby.

## Bibliography

[1] Noam Agmon. Tetrahedral displacement: The molecular mechanism behind the Debye relaxation in water. Journal of Physical Chemistry, 100(1):1072-1080, 1996.
[2] M. P. Allen and D. J. Tildesley. Computer Simulation of Liquids. Oxford Science Publications, Clarendon Press, Oxford, 1989.
[3] Patrick Aloy and Robert B. Russell. Interrogating protein interaction networks through structural biology. Proceedings of the National Academy of Sciences, USA, 99:5896-5901, 2002.
[4] Claus A.F. Andersen, Arthur G. Palmer, Sren Brunak, and Burkhard Rost. Continuum secondary structure captures protein flexibility. Structure, 10:175-184, 2002.
[5] D. E. ANDERSON, J. H. HURLEY, H. NICHOLSON, W. A. BAASE, and B. W. MATTHEWS. Hydrophobic core repacking and aromatic-aromatic interaction in the thermostable mutant of T4 lysozyme Ser $117 \rightarrow$ Phe. Protein Science, 2(8):1285-1290, 1993.
[6] G. Apic, J. Gough, and S. A. Teichmann. An insight into domain combinations. Bioinformatics, 17 (Suppl.1):S83-89, 2001.
[7] Isaiah T. Arkin, Huafeng Xu, Morten O. Jensen, Eyal Arbely, Estelle R. Bennett, Kevin J. Bowers, Edmond Chow, Ron O. Dror, Michael P. Eastwood, Ravenna Flitman-Tene, Brent A. Gregersen, John L. Klepeis, Istvan Kolossvary, Yibing Shan, and David E. Shaw. Mechanism of $\mathrm{Na}+/ \mathrm{H}+$ antiporting. Science, 317(5839):799-803, 2007.
[8] R. Aurora and G. D. Rose. Helix capping. Protein Science, 7(1):21-38, 1998.
[9] Kellar Autumn, Metin Sitti, Yiching A. Liang, Anne M. Peattie, Wendy R. Hansen, Simon Sponberg, Thomas W. Kenny, Ronald Fearing, Jacob N. Israelachvili, and Robert J. Full. Evidence for van der Waals adhesion in gecko setae. Proceedings of the National Academy of Sciences, USA, 99(19):12252-12256, 2002.
[10] Franc Avbelj and Robert L. Baldwin. Role of backbone solvation and electrostatics in generating preferred peptide backbone conformations: distributions of phi. Proceedings of the National Academy of Sciences, USA, 100(10):5742-5747, 2003.

Draft: February 28, 2008, do not distribute
[11] Franc Avbelj and Robert L. Baldwin. Origin of the neighboring residue effect on peptide backbone conformation. Proceedings of the National Academy of Sciences, USA, 101(30):1096710972, 2004.
[12] Franc Avbelj, P. Luo, and Robert L. Baldwin. Energetics of the interaction between water and the helical peptide group and its role in determining helix propensities. Proceedings of the National Academy of Sciences, USA, 97:10786-10791, 2000.
[13] JC Avise, BW Bowen, T Lamb, AB Meylan, and E Bermingham. Mitochondrial DNA evolution at a turtle's pace: evidence for low genetic variability and reduced microevolutionary rate in the Testudines. Mol Biol Evol, 9(3):457-473, 1992.
[14] R. Azriel and E. Gazit. Analysis of the minimal amyloid-forming fragment of the islet amyloid polypeptide. An experimental support for the key role of the phenylalanine residue in amyloid formation. Journal of Biological Chemistry, 276(36):34156-34161, Sep 2001.
[15] Ranjit Prasad Bahadur, Pinak Chakrabarti, Francis Rodier, and Joöl Janin. Dissecting subunit interfaces in homodimeric proteins. Proteins: Str. Func. Gen., 53:708-719, 2003.
[16] Ranjit Prasad Bahadur, Pinak Chakrabarti, Francis Rodier, and Joöl Janin. A dissection of specific and non-specific protein-protein interfaces. Journal of Molecular Biology, 336:943-955, 2004.
[17] E. N. Baker and R. E. Hubbard. Hydrogen bonding in globular proteins. Progress in Biophysics and Molecular Biology, 44:97-179, 1984.
[18] Robert L. Baldwin. Protein folding: Making a network of hydrophobic clusters. Science, 295:1657-1658, 2002.
[19] Y.-E. A. Ban, H. Edelsbrunner, and J. Rudolph. Interface surfaces for protein-protein complexes. In Proceedings of the 8th Annual International Conference on Research in Computational Molecular Biology (RECOMB), pages 205-212, 2004.
[20] Hans-Jurgen Bandelt and Andreas Dress. Reconstructing the shape of a tree from observed dissimilarity data. Advances in Applied Mathematics, 7:309-343, Sep 1986.
[21] A. Barabasi and R. Albert. Emergence of scaling in random networks. Science, 286:509-512, 1999.
[22] Arieh Y. Ben-Naim. Hydrophobic Interactions. Springer, 1980.
[23] Marco Bittelli, Markus Flury, and Kurt Roth. Use of dielectric spectroscopy to estimate ice content in frozen porous media. WATER RESOURCES RESEARCH, 40:W04212, 2004.
[24] A. A. Bogan and K. S. Thorn. Anatomy of hot spots in protein interfaces. Journal of Molecular Biology, 280:1-9, 1998.

Draft: February 28, 2008, do not distribute
[25] A. A. Bogan and K. S. J. Thorn. Anatomy of hot spots in protein interfaces. Journal of Molecular Biology, 280:1-9, 1998.
[26] A. Bondi. van der Waals volumes and radii. Journal of Physical Chemistry, 68(3):441-451, 1964.
[27] Clay Bracken, Lilia M. Iakoucheva, Pedro R. Romero, and A. Keith Dunker. Combining prediction, computation and experiment for the characterization of protein disorder. Curr Opin Struct Biol, 14(5):570-576, 2004.
[28] Bobby-Joe Breitkreutz, Chris Stark, Teresa Reguly, Lorrie Boucher, Ashton Breitkreutz, Michael Livstone, Rose Oughtred, Daniel H. Lackner, Jrg Bhl, Valerie Wood, Kara Dolinsk, and Mike Tyers. The BioGRID Interaction Database: 2008 update. Nucl. Acids Res., 36(database issue):D637-640, 2008.
[29] E Breslow, V Mombouyran, R Deeb, C Zheng, JP Rose, BC Wang, and RH Haschemeyer. Structural basis of neurophysin hormone specificity: Geometry, polarity, and polarizability in aromatic ring interactions. Protein Science, 8(4):820-831, 1999.
[30] Dawn J. Brooks, Jacques R. Fresco, Arthur M. Lesk, and Mona Singh. Evolution of amino acid frequencies in proteins over deep time: Inferred order of introduction of amino acids into the genetic code. Molecular Biology and Evolution, 19:1645-1655, 2002.
[31] William Fuller Brown, Jr. Dielectrics. In Handbuch der Physik, volume 14, pages 1-154. Springer, 1956.
[32] David Bryant. On the uniqueness of the selection criterion in neighbor-joining. Journal of Classification, 22:3-15, Jun 2005. 10.1007/s00357-005-0003-x.
[33] R. G. Bryant. The dynamics of water-protein interactions. Annu. Rev. Biophys. Biomol. Struct., 25:29-53, 1996.
[34] Christian J. Burnham and Sotiris S. Xantheas. Development of transferable interaction models for water. iii. reparametrization of an all-atom polarizable rigid model (ttm2-r) from first principles. Journal of Chemical Physics, 116(4):1500-1510, 2002.
[35] G. Manhès C. J. Allègre and Christa Göpel. The age of the earth. Geochimica et Cosmochimica Acta, 59(8):1445-1456, 1995.
[36] Daniel R. Caffrey, Shyamal Somaroo, Jason D. Hughes, Julian Mintseris, and Enoch S. Huang. Are protein-protein interfaces more conserved in sequence than the rest of the protein surface? Protein Science, 13(1):190-202, 2004.
[37] Eric Cancès, Claude Le Bris, and Yvon Maday. Mèthodes mathèmatiques en chimie quantique. Springer, 2006.

Draft: February 28, 2008, do not distribute
[38] Eric Cancès, M. Defranceschi, W. Kutzelnigg, Claude Le Bris, and Yvon Maday. Computational quantum chemistry: a primer. In Ph. Ciarlet and C. Le Bris, editors, Handbook of numerical analysis. Volume X: special volume: computational chemistry, pages 3-270. Elsevier, 2003.
[39] E. Candes and L. Demanet. The curvelet representation of wave propagators is optimally sparse. Comm. Pure Appl. Math., 58(11):1472-1528, 2005.
[40] Pinak Chakrabarti and Joöl Janin. Dissecting protein-protein recognition sites. Proteins: Str. Func. Gen., 47:334-343, 2002.
[41] Jianping Chen, Xi Zhang, Ridgway Scott, and Ariel Fernández. Molecular basis for promiscuity and specificity in the drugable kinome. Bioinformatics, page 7, 2006.
[42] Dmitry A. Cherepanov. Force oscillations and dielectric overscreening of interfacial water. Phys. Rev. Lett., 93:266104, 2004.
[43] T. Clackson and J. A. Wells. A hot spot of binding energy in a hormone-receptor interface. Science, 267:383-386, 1995.
[44] C. COLOVOS and T. O. YEATES. Verification of protein structures: Patterns of nonbonded atomic interactions. Protein Sci, 2(9):1511-1519, 1993.
[45] David B. Cook, editor. Handbook of Computational Quantum Chemistry. Oxford Univ. Press, 1998.
[46] Christopher J. Cramer. Essentials of Computational Chemistry. Wiley, second edition, 2004.
[47] Thomas E. Creighton. Proteins: Structures and molecular properties. W. H. Freeman, 1993.
[48] Peter B. Crowley and Adel Golovin. Cation- $\pi$ interactions in protein-protein interfaces. PROTEINS: Structure, Function, and Bioinformatics, 59:231-239, 2005.
[49] Alfonso De Simone, Guy G. Dodson, Chandra S. Verma, Adriana Zagari, and Franca Fraternali. Prion and water: Tight and dynamical hydration sites have a key role in structural stability. Proceedings of the National Academy of Sciences, USA, 102:7535-7540, 2005.
[50] P. Debye. Polar molecules. Dover, New York, 1945.
[51] W. L. DeLano, M. H. Ultsch, A. M. de Vos, and J. A. Wells. Convergent solutions to binding at a protein-protein interface. Science, 287:1279-1283, 2000.
[52] A. H. DePace, A. Santoso, P. Hillner, and J. S. Weissman. A critical role for aminoterminal glutamine/asparagine repeats in the formation and propagation of a yeast prion. Cell, 93(7):1241-1252, Jun 1998.
[53] Florin Despa and R. Stephen Berry. The origin of long range attraction between hydrophobes in water. Biophysical Journal, 92:373-378, 2007.

Draft: February 28, 2008, do not distribute
[54] Florin Despa, Ariel Fernández, and R. Stephen Berry. Dielectric modulation of biological water. Phys. Rev. Lett., 93:269901, 2004.
[55] Dennis A. Dougherty. Cation-pi interactions in chemistry and biology: a new view of benzene, Phe, Tyr, and Trp. Science, 271(5246):163-168, 1996.
[56] John Doyle. Beyond the sperical cow. Nature, 411:151-152, May 2001.
[57] Xiaoqun Joyce Duan, Ioannis Xenarios, and David Eisenberg. Describing biological protein interactions in terms of protein states and state transitions. Mol. Cell. Proteomics, 1:104-116, 2002.
[58] Johannes J. Duistermaat. Fourier Integral Operators. Springer, Berlin, 1996.
[59] Kord Eickmeyer, Peter Huggins, Lior Pachter, and Ruriko Yoshida. On the optimality of the neighbor-joining algorithm, 2007.
[60] A. Elgsaeter, B. T. Stokke, A. Mikkelsen, and D. Branton. The molecular basis of erythrocyte shape. Science, 234:1217-1223, 1986.
[61] Isaac Elias and Jens Lagergren. Fast Neighbor Joining, pages 1263-1274. LNCS Volume 3580. Springer, 2005. 10.1007/11523468_102.
[62] Miles A. Fabian, William H. Biggs, Daniel K. Treiber, Corey E. Atteridge, Mihai D. Azimioara, Michael G. Benedetti, Todd A. Carter, Pietro Ciceri, Philip T. Edeen, Mark Floyd, Julia M. Ford, Margaret Galvin, Jay L. Gerlach, Robert M Grotzfeld, Sanna Herrgard, Darren E. Insko, Michael A. Insko, Andiliy G Lai, Jean-Michel Lelias, Shamal A. Mehta, Zdravko V. Milanov, Anne Marie Velasco, Lisa M. Wodicka, Hitesh K. Patel, Patrick P. Zarrinkar, and David J. Lockhart. A small molecule-kinase interaction map for clinical kinase inhibitors. Nature Biotechnology, 23(3):329-336, 2005.
[63] M. Fandrich, M. A. Fletcher, and C. M. Dobson. Amyloid fibrils from muscle myoglobin. Nature, 410(6825):165-166, Mar 2001.
[64] Ariel Fernández. Cooperative walks in a cubic lattice: Protein folding as a many-body problem. J. Chem. Phys., 115:7293-7297, 2001.
[65] Ariel Fernández. Intramolecular modulation of electric fields in folding proteins. Phys. Lett. A, 299:217-220, 2002.
[66] Ariel Fernández. Buffering the entropic cost of hydrophobic collapse in folding proteins. Journal of Chemical Physics, 121:11501-11502, 2004.
[67] Ariel Fernández. Keeping dry and crossing membranes. Nature Biotechnology, 22:1081-1084, 2004.

Draft: February 28, 2008, do not distribute
[68] Ariel Fernández. Direct nanoscale dehydration of hydrogen bonds. Journal of Physics D: Applied Physics, 38:2928-2932, 2005.
[69] Ariel Fernández. What factor drives the fibrillogenic association of beta-sheets? FEBS Letters, 579:6635-6640, 2005.
[70] Ariel Fernández and R. Stephen Berry. Extent of hydrogen-bond protection in folded proteins: a constraint on packing architectures. Biophysical Journal, 83(5):2475-2481, Nov 2002.
[71] Ariel Fernández and R. Stephen Berry. Proteins with H-bond packing defects are highly interactive with lipid bilayers: Implications for amyloidogenesis. Proceedings of the National Academy of Sciences, USA, 100:2391-2396, 2003.
[72] Ariel Fernández and R. Stephen Berry. Molecular dimension explored in evolution to promote proteomic complexity. Proceedings of the National Academy of Sciences, USA, 101:1346013465, 2004.
[73] Ariel Fernández and M. Boland. Solvent environment conducive to protein aggregation. FEBS Letters, 529:298-303, 2002.
[74] Ariel Fernández, Jösef Kardos, L. Ridgway Scott, Yuji Goto, and R. Stephen Berry. Structural defects and the diagnosis of amyloidogenic propensity. Proceedings of the National Academy of Sciences, USA, 100(11):6446-6451, 2003.
[75] Ariel Fernández and Sridhar Maddipati. A priori inference of cross reactivity for drug-targeted kinases. Journal of Medicinal Chemistry, 49(11):3092-3100, 2006.
[76] Ariel Fernández, Kristina Rogale Plazonic, L. Ridgway Scott, and Harold A. Scheraga. Inhibitor design by wrapping packing defects in HIV-1 proteins. Proceedings of the National Academy of Sciences, USA, 101:11640-11645, 2004.
[77] Ariel Fernández and Harold A. Scheraga. Insufficiently dehydrated hydrogen bonds as determinants of protein interactions. Proceedings of the National Academy of Sciences, USA, 100(1):113-118, Jan 2003.
[78] Ariel Fernández and L. Ridgway Scott. Adherence of packing defects in soluble proteins. Phys. Rev. Lett., 91:18102(4), 2003.
[79] Ariel Fernández and L. Ridgway Scott. Dehydron: a structurally encoded signal for protein interaction. Biophysical Journal, 85:1914-1928, 2003.
[80] Ariel Fernández and L. Ridgway Scott. Under-wrapped soluble proteins as signals triggering membrane morphology. Journal of Chemical Physics, 119(13):6911-6915, 2003.
[81] Ariel Fernández and L. Ridgway Scott. Modulating drug impact by wrapping target proteins. Expert Opinion on Drug Discovery, 2:249-259, 2007.

Draft: February 28, 2008, do not distribute
[82] Ariel Fernández, L. Ridgway Scott, and R. Stephen Berry. The nonconserved wrapping of conserved protein folds reveals a trend towards increasing connectivity in proteomic networks. Proceedings of the National Academy of Sciences, USA, 101(9):2823-2827, 2004.
[83] Ariel Fernández, L. Ridgway Scott, and Harold A. Scheraga. Amino-acid residues at proteinprotein interfaces: Why is propensity so different from relative abundance? J. Phys. Chem. B, 107(36):9929-9932, 2003.
[84] Ariel Fernández, Tobin R. Sosnick, and Andres Colubri. Dynamics of hydrogen bond desolvation in protein folding. Journal of Molecular Biology, 321(4):659-675, Aug 2002.
[85] Josephine C. Ferreon and Vincent J. Hilser. The effect of the polyproline II (PPII) conformation on the denatured state entropy. Protein Sci, 12(3):447-457, 2003.
[86] Herman Feshbach. Unified theory of nuclear reactions. Annals of Physics, 5:357-390, 1958.
[87] J.E. Fitzgerald, A.K. Jha, T.R. Sosnick, and K.F. Freed. Polypeptide motions are dominated by peptide group oscillations resulting from dihedral angle correlations between nearest neighbors. Biochemistry, 46(3):669-682, 2007.
[88] Anton F. Fliri, William T. Loging, Peter F. Thadeio, and Robert A. Volkmann. Biological spectra analysis: Linking biological activity profiles to molecular structure. Proceedings of the National Academy of Sciences, USA, 102(2):261-266, 2005.
[89] Rodolphe Vuilleumier Anne Boutin François-Xavier Coudert. Dipole moment, hydrogen bonding and ir spectrum of confined water. ChemPhysChem, 7(12):2464-2467, 2006.
[90] Rosalind Franklin and R. G. Gosling. Evidence for 2-chain helix in crystalline structure of sodium deoxyribonucleate. Nature, 172:156-157, 1953.
[91] Felix Franks. Water: a matrix for life. Royal Society of Chemistry, 2000.
[92] Ernesto Freire. The propagation of binding interactions to remote sites in proteins: Analysis of the binding of the monoclonal antibody D1.3 to lysozyme. Proceedings of the National Academy of Sciences, USA, 96:10118-10122, 1999.
[93] Robert H. French. Origins and applications of London dispersion forces and Hamaker constants in ceramics. Journal of the American Ceramic Society, 83(9):2117-46, 2000.
[94] E. Fry, R. Acharya, and D. Stuart. Methods used in the structure determination of foot-andmouth disease virus. Acta Crystall. A, 49:45-55, 1993.
[95] Justin P. Gallivan and Dennis A. Dougherty. Cation-pi interactions in structural biology. Proceedings of the National Academy of Sciences, USA, 96(17):9459-9464, 1999.
[96] Olivier Gascuel and Mike Steel. Neighbor-joining revealed. Mol Biol Evol, 23(11):1997-2000, 2006.

Draft: February 28, 2008, do not distribute
[97] Fabian Glaser, David M. Steinberg, Ilya A. Vakser, and Nir Ben-Tal. Residue frequencies and pairing preferences at protein-protein interfaces. Proteins: Str. Func. Gen., 43:89-102, 2001.
[98] S. A. Goudschmidt and G. H. Uhlenbeck. Spinning electrons and the structure of spectra. Nature, 117(2938):264-265, 1926.
[99] G.A. Grant, C.W. Luetje, R. Summers, and X.L. Xu. Differential roles for disulfide bonds in the structural integrity and biological activity of $\kappa$-bungarotoxin, a neuronal nicotinic acetylcholine receptor antagonist. Biochemistry, 37(35):12166-12171, 1998.
[100] J. K. Gregory, D. C. Clary, K. Liu, M. G. Brown, and R. J. Saykally. The water dipole moment in water clusters. Science, 275(5301):814-817, 1997.
[101] J. I. Guijarro, M. Sunde, J. A. Jones, I. D. Campbell, and C. M. Dobson. Amyloid fibril formation by an SH3 domain. Proceedings of the National Academy of Sciences, USA, 95(8):4224-4228, Apr 1998.
[102] Dan Gusfield. Algorithms on Strings, Trees, and Sequences: Computer Science and Computational Biology. Cambridge University Press, January 1997.
[103] George Hagedorn. High order corrections to the time-independent Born-Oppenheimer approximation. I. smooth potentials. Ann. Inst. Henri Poincaré, 47:1-16, 1987.
[104] George Hagedorn. High order corrections to the time-independent Born-Oppenheimer approximation. II. diatomic Coulomb systems. Comm. Math. Phys., 116:23-44, 1988.
[105] Bertil Halle. Protein hydration dynamics in solution: a critical survey. Philosophical Transactions of the Royal Society B: Biological Sciences, 359(1448):1207-1224, 2004.
[106] Inbal Halperin, Haim Wolfson, and Ruth Nussinov. Protein-protein interactions; coupling of structurally conserved residues and of hot spots across interfaces. implications for docking. Structure, 12:1027-1038, 2004.
[107] D. Hamelberg and J.A. McCammon. Fast peptidyl cis-trans isomerization within the flexible Gly-rich flaps of HIV-1 protease. Journal of the American Chemical Society, 127(40):1377813779, 2005.
[108] Yehouda Harpaz, Mark Gerstein, and Cyrus Chothia. Volume changes on protein folding. Structure, 2:641-649, 1994.
[109] J. P. Helfrich. Dynamic laser light scattering technology for the molecular weight and hydrodynamic radius characterization of proteins. Pharmaceutical Laboratory, 1:34-40, 1998.
[110] M. Heuberger, T. Drobek, and N. D. Spencer. Interaction forces and morphology of a proteinresistant poly(ethylene glycol) layer. Biophysical Journal, 88(1):495-504, 2005.

Draft: February 28, 2008, do not distribute
[111] Nora E. Hill, Worth E. Vaughan, A. H. Price, and Mansel Davies. Dielectric properties and molecular behaviour. van Nostrand, London, 1969.
[112] U. Hobohm, M. Scharf, R. Schneider, and C. Sander. election of representative protein data sets. Protein Science, 1(3):409-417, 1992.
[113] B. Honig and A. S. Yang. Free energy balance in protein folding. Adv. Protein Chem., 46:27-58, 1995.
[114] M. Hoshino, H. Katou, Y. Hagihara, K. Hasegawa, H. Naiki, and Y. Goto. Mapping the core of the beta(2)-microglobulin amyloid fibril by H/D exchange. Nature Structural Biology, 19:332-336, 2002.
[115] Thomas Hou and Petros Koumoutsakos. Special section on multiscale modeling in materials and life sciences. SIAM J. Multiscale Modeling ES Simulation, 4(1):213-214, 2005.
[116] Zengjian Hu, Buyong Ma, Haim Wolfson, and Ruth Nussinov. Conservation of polar residues as hot spots at protein interfaces. Proteins: Str. Func. Gen., 39:331-342, 2000.
[117] L. M. Iakoucheva and K. A. Dunker. Order, disorder, and flexibility: prediction from protein sequence. Structure, 11:1316-1317, 2003.
[118] Jacob Israelachvili. Intermolecular and surface forces. Academic Press, second edition, 1991.
[119] Jacob Israelachvili and Hakan Wennerstrom. Role of hydration and water structure in biological and colloidal interactions. Nature, 379:219-225, 1996.
[120] Joël Janin, S. Miller, and Cyrus Chothia. Surface, subunit interfaces and interior of oligomeric proteins. Journal of Molecular Biology, 204:155-164, 1988.
[121] J.D.Madura, J.M.Briggs, R.C.Wade, M.E.Davis, B.A.Luty, A. Ilin, J.Antosciewicz, M.K.Gilson, B. Bagheri, L.R.Scott, and J.A.McCammon. Electrostatics and diffusion of molecules in solution: Simulations with the University of Houston Brownian Dynamics program. Comp. Phys. Commun., 91:57-95, Sept. 1995.
[122] George A. Jeffrey. An introduction to hydrogen bonds. Oxford Science Publications, Clarendon Press, Oxford, 1997.
[123] George A. Jeffrey. Hydrogen-bonding: An update. Crystallography Reviews, 9:135-176, Jan 2003. 10.1080/08893110310001621754.
[124] S. Jones, A. Marin, and J. M. Thornton. Protein domain interfaces: characterization and comparison with oligomeric protein interfaces. Protein Eng., 13:77-82, 2000.
[125] Susan Jones and Janet M. Thornton. Principles of protein-protein interactions. Proceedings of the National Academy of Sciences, USA, 93:13-20, 1996.

Draft: February 28, 2008, do not distribute
[126] Susan Jones and Janet M. Thornton. Analysis of protein-protein interaction sites using surface patches. Journal of Molecular Biology, 272:121-132, 1997.
[127] Susan Jones and Janet M. Thornton. Prediction of protein-protein interaction sites using patch analysis. Journal of Molecular Biology, 272:133-143, 1997.
[128] U. Kaatze, R. Behrends, and R. Pottel. Hydrogen network fluctuations and dielectric spectrometry of liquids. Journal of Non-Crystalline Solids, 305(1):19-28, 2002.
[129] R. Kaufmann, U. Junker, M. Schilli-Westermann, C. Klötzer, J. Scheele, and K. Junker. Meizothrombin, an intermediate of prothrombin cleavage potently activates renal carcinoma cells by interaction with par-type thrombin receptors. Oncology Reports, 10:493-496, 2003.
[130] W. Kauzmann. Some factors in the interpretation of protein denaturization. In Advances in Protein Chemistry, volume 14, pages 1-63. Academic Press, 1959.
[131] S. S. Kim, T. J. Smith, M. S. Chapman, M. C. Rossmann, D. C. Pevear, F. J. Dutko, P. J. Felock, G. D. Diana, and M. A. McKinlay. Crystal structure of human rhinovirus serotype 1A (HRV1A). Journal of Molecular Biology, 210:91-111, 1989.
[132] M. Klein, A. Martinez, R. Seiler, and X. P. Wang. Born-Oppenheimer expansion for polyatomic molecules. Comm. Math. Phys, pages 607-639, 1992.
[133] W. Klopper, J. G. C. M. van Duijneveldt-van de Rijdt, and F. B. van Duijneveldt. Computational determination of equilibrium geometry and dissociation energy of the water dimer. Physical Chemistry Chemical Physics, 2:2227- 2234, 2000.
[134] Peter A. Kollman and Leland C. Allen. The theory of the hydrogen bond. Chemical Reviews, 72(3):283-303, 1972.
[135] E. V. Koonin, Y. I. Wolf, and G. P. Karev. The structure of the protein universe and genome evolution. Nature, 420:218-223, 2002.
[136] J. Korlach, P. Schwille, W. W. Webb, and G. W. Feigenson. Characterization of lipid bilayer phases by confocal microscopy and fluorescence correlation spectroscopy. Proceedings of the National Academy of Sciences, USA, 96:8461-8466, 1999.
[137] A. A. Kornyshev and A. Nitzan. Effect of overscreeming on the localization of hydrated electrons. Zeitschrift für Physikalische Chemie, 215(6):701-715, 2001.
[138] Tanja Kortemme, Alexandre V. Morozov, and David Baker. An orientation-dependent hydrogen bonding potential improves prediction of specificity and structure for proteins and protein-protein complexes. Journal of Molecular Biology, 326:1239-1259, 2003.

Draft: February 28, 2008, do not distribute
[139] Gennady V. Kozhukh, Yoshihisa Hagihara, Toru Kawakami, Kazuhiro Hasegawa, Hironobu Naiki, and Yuji. Goto. Investigation of a peptide responsible for amyloid fibril formation of beta 2-microglobulin by achromobacter protease I. Journal of Biological Chemistry, 277(2):1310-1315, Jan 2002.
[140] Lawrence M. Krauss. Fear of Physics. Basic Books, 1994.
[141] S. Krishnaswamy and Michael G. Rossmann. Structural refinement and analysis of mengo virus. Journal of Molecular Biology, 211:803-844, 1990.
[142] J. Kroon, J. A. Kanters, J. G. C. M. van Duijneveldt-van De Rijdt, F. B. van Duijneveldt, and J. A. Vliegenthart. O-H • O hydrogen bonds in molecular crystals a statistical and quantum-chemical analysis. Journal of Molecular Structure, 24:109-129, Jan 1975.
[143] J. Kyte and R. F. Doolittle. A simple method for displaying the hydropathic character of a protein. Journal of Molecular Biology, 157:105-132, 1982.
[144] M. Laing. No rabbit ears on water. J. Chem. Ed., 64:124-128, 1987.
[145] S. Leikin and A. A. Kornyshev. Theory of hydration forces. nonlocal electrostatic interaction of neutral surfaces. J. Chem. Phys., 92(6):6890-6898, 1990.
[146] P.A. Leland, K.E. Staniszewski, C. Park, B.R. Kelemen, and R.T. Raines. The ribonucleolytic activity of angiogenin. Biochemistry, 41(4):1343-1350, 2002.
[147] Yaakov Levy and Jose N. Onuchic. Water and proteins: A love-hate relationship. Proceedings of the National Academy of Sciences, USA, 101(10):3325-3326, 2004.
[148] Laurent Limozin and Erich Sackmann. Polymorphism of cross-linked actin networks in giant vesicles. Phys Rev Lett, 89(16):168103, Oct 2002.
[149] M. Lisal, J. Kolafa, and I. Nezbeda. An examination of the five-site potential (TIP5P) for water. Journal of Chemical Physics, 117:8892-8897, November 2002.
[150] Bernard A. Liu, Karl Jablonowski, Monica Raina, Michael Arcé, Tony Pawson, and Piers D. Nash. The human and mouse complement of SH2 domain proteins - establishing the boundaries of phosphotyrosine signaling. Molecular Cell, 22(6):851-868, 2006.
[151] Jing Liu, L. Ridgway Scott, and Ariel Fernández. Interactions of aligned nearest neighbor protein side chains. Journal of Bioinformatics and Computational Biology, 7:submitted-2006.
[152] Jing Liu, L. Ridgway Scott, and Ariel Fernández. Interactions of aligned nearest neighbor protein side chains. Journal of Bioinformatics and Computational Biology, 7:submitted-2006.

Draft: February 28, 2008, do not distribute
[153] Jing Liu, L. Ridgway Scott, and John Goldsmith. Journal of Bioinformatics and Computational Biology, 7:submitted-, 2006.
[154] Loredana Lo Conte, S. E. Brenner, T. J. Hubbard, Cyrus Chothia, and A. G. Murzin. SCOP database in 2002: refinements accommodate structural genomics. Nucleic Acids Res., 30:264267, 2002.
[155] Loredana Lo Conte, Cyrus Chothia, and Joël Janin. The atomic structure of protein-protein recognition sites. Mol. Biol., 285:2177-2198, 1999.
[156] Simon C. Lovell, J. Michael Word, Jane S. Richardson, and David C. Richardson. The penultimate rotamer library. PROTEINS: Structure, Function, and Genetics, 40:389-408, 2000.
[157] Per-Olov Löwdin. Expansion theorems for the total wave function and extended Hartree-Fock schemes. Rev. Mod. Phys., 32(2):328-334, 1960.
[158] Buyong Ma, Tal Elkayam, Haim Wolfson, and Ruth Nussinov. Protein-protein interactions: Structurally conserved residues distinguish between binding sites and exposed protein surfaces. Proceedings of the National Academy of Sciences, USA, 100(10):5772-5777, 2003.
[159] A. Magalhaes, B. Maigret, J. Hoflack, J. A. N. F. Gomes, and H. A. Scheraga. Contribution of unusual arginine-arginine short-range interactions to stabilization and recognition in proteins. J. Protein Chem., 13(2):195-215, 1994.
[160] Michael W. Mahoney and William L. Jorgensen. A five-site model for liquid water and the reproduction of the density anomaly by rigid, nonpolarizable potential functions. J. Chem. Phys., 112:8910-8922, 2000.
[161] R. B. Martin. Localized and spectroscopic orbitals: Squirrel ears on water. J. Chem. Ed., 65:668-670, 1988.
[162] A. Martinez. Eigenvalues and resonances of polyatomic molecules in the Born-Oppenheimer approximation. In Erik Balslev, editor, Schrödinger Operators, The Quantum Mechanical Many-Body Problem, pages 145-152, Berlin, 1992. Springer-Verlag.
[163] P. E. Mason, G. W. Neilson, C. E. Dempsey, A. C. Barnes, and J. M. Cruickshank. The hydration structure of guanidinium and thiocyanate ions: Implications for protein stability in aqueous solution. Proceedings of the National Academy of Sciences, USA, 100(8):4557-4561, 2003.
[164] C. Matzler and U. Wegmuller. Dielectric properties of freshwater ice at microwave frequencies. Journal of Physics D: Applied Physics, 20(12):1623-1630, 1987.
[165] I.K. McDonald and Janet M. Thornton. Satisfying hydrogen bonding potential in proteins. Journal of Molecular Biology, 238:777-793, 1994.

Draft: February 28, 2008, do not distribute
[166] Victoria J. McParland, Arnout P. Kalverda, Steve W. Homans, and Sheena E. Radford. Structural properties of an amyloid precursor of beta(2)-microglobulin. Nature Structural Biology, 9(5):326-331, May 2002.
[167] Carver Mead and Lynn Conway. Introduction to VLSI Systems. Addison-Wesley, 1979.
[168] Radu Mihaescu, Dan Levy, and Lior Pachter. Why neighbor-joining works. Algorithmica, 2008. 10.1007/s00453-007-9116-4.
$[169]$ D. J. Mitchell, L. Steinman, D. T. Kim, C. G. Fathman, and J. B. Rothbard. Polyarginine enters cells more efficiently than other polycationic homopolymers. Journal of Peptide Research, 56(5):318-325, 2000.
[170] C. Momany, L. C. Kovari, A. J. Prongay, W. Keller, R. K. Gitti, B. M. Lee, A. E. Gorbalenya, L. Tong, J. McClure, L. S. Ehrlich, M. F. Carter, and M. G. Rossmann. Crystal structure of dimeric HIV-1 capsid protein. Nature Structural Biology, 3:763-770, 1996.
[171] F.A. Momany, R.F. McGuire, A.W. Burgess, and H.A. Scheraga. Energy parameters in polypeptides. vii. geometric parameters, partial atomic charges, nonbonded interactions, hydrogen bond interactions, and intrinsic torsional potentials for the naturally occurring amino acids. Journal of Physical Chemistry, 79(22):2361-2381, 1975.
[172] J. Moro, J.V. Burke, and M.L. Overton. On the Lidskii-Lyusternik-Vishik perturbation theory for eigenvalues with arbitrary Jordan structure. SIAM J. Matrix Anal. Appl., 18:793-817, 1997.
[173] Alexandre V. Morozov, Tanja Kortemme, Kiril Tsemekhman, and David Baker. Close agreement between the orientation dependence of hydrogen bonds observed in protein structures and quantum mechanical calculations. Proceedings of the National Academy of Sciences, USA, 101:6946-6951, 2004.
[174] James A. Morrill and Roderick MacKinnon. Isolation of a single carboxyl-carboxylate proton binding site in the pore of a cyclic nucleotide-gated channel. J. Gen. Physiol., 114(1):71-84, 1999.
[175] A. S. Muresan, H. Diamant, and K.-Y. Lee. Effect of temperature and composition on the formation of nanoscale compartments in phospholipid membranes. J. Am. Chem. Soc., 123:69516952, 2001.
[176] A. G. Murzin, S. E. Brenner, T. J. Hubbard, and C. Chothia. SCOP: a structural classification of proteins database for the investigation of sequences and structures. Journal of Molecular Biology, 247:536-540, 1995.
[177] G. Némethy, I. Z. Steinberg, and H. A. Scheraga. Influence of water structure and of hydrophobic interactions on the strength of side-chain hydrogen bonds in proteins. Biopolymers, 1:43-69, 1963.

Draft: February 28, 2008, do not distribute
[178] Alexander Neumeister, Nicole Praschak-Rieder, Barbara Hesselmann, Oliver Vitouch, Manfred Rauh, Arnd Barocka, Johannes Tauscher, and Siegfried Kasper. Effects of tryptophan depletion in drug-free depressed patients who responded to total sleep deprivation. Arch. Gen. Psychiatry, 55(2):167-172, 1998.
[179] Kyoung Tai No, Oh Young Kwon, Su Yeon Kim, Mu Shik Jhon, and Harold A. Scheraga. A simple functional representation of angular-dependent hydrogen-bonded systems. 1. amide, carboxylic acid, and amide-carboxylic acid pairs. Journal of Physical Chemistry, 99:34783486, 1995.
[180] Irene M. A. Nooren and Janet M. Thornton. Diversity of proteinprotein interactions. EMBO Journal, 22:3486-3492, 2003.
[181] Irene M. A. Nooren and Janet M. Thornton. Structural characterisation and functional significance of transient protein-protein interactions. Journal of Molecular Biology, 325(5):991-1018, 2003.
[182] Yanay Ofren and Burkhard Rost. Analysing six types of protein-protein interfaces. Journal of Molecular Biology, 335:377-387, 2003.
[183] Diana O. Omecinsky, Katherine E. Holub, Michael E. Adams, and Michael D. Reily. Threedimensional structure analysis of $\mu$-agatoxins: Further evidence for common motifs among neurotoxins with diverse ion channel specificities. Biochemistry, 35(9):2836-2844, 1996.
[184] T. Ooi. Thermodynamics of protein folding: effects of hydration and electrostatic interactions. Advan. Biophys., 30:105-154, 1994.
[185] D. Ouroushev. Electric field of dielectric cylinder with given surface charge density immersed in electrolite. Journal of Physics A: Mathematical and General, 31(16):3897-3902, 1998.
[186] Doreen Pahlke, Dietmar Leitner, Urs Wiedemann, and Dirk Labudde. COPS-cis/trans peptide bond conformation prediction of amino acids on the basis of secondary structure information. Bioinformatics, 21(5):685-686, 2005.
[187] Samir Kumar Pal, Jorge Peon, and Ahmed H. Zewail. Biological water at the protein surface: Dynamical solvation probed directly with femtosecond resolution. Proceedings of the National Academy of Sciences, USA, 99(4):1763-1768, 2002.
[188] J. Park, M. Lappe, and S. A. Teichmann. Mapping protein family interactions: Intramolecular and intermolecular protein family interaction repertoires in the PDB and yeast. Journal of Molecular Biology, 307:929-938, 2001.
[189] Wolfgang Pauli. The connection between spin and statistics. Phys. Rev., 58(8):716-722, Oct 1940.
[190] Wolfgang Pauli. Exclusion principle and quantum mechanics. In Nobel Lectures, Physics 1942-1962, pages 27-43, Amsterdam, 1964. Elsevier Publishing Company.

Draft: February 28, 2008, do not distribute
[191] Linus Pauling. Nature of the Chemical Bond. Cornell Univ. Press, third edition, 1960.
[192] Linus Pauling. General Chemistry. Dover, 1970.
[193] Linus Pauling and E. Bright Wilson. Introduction to Quantum Mechanics with Applications to Chemistry. Dover, 1985.
[194] Gang Pei, Thomas M. Laue, Ann Aulabaugh, Dana M. Fowlkes, and Barry R. Lentz. Structural comparisons of meizothrombin and its precursor prothrombin in the presence or absence of procoagulant membranes. Biochemistry, 31:6990-6996, 1992.
[195] D. Petrey and B. Honig. Free energy determinants of tertiary structure and the evaluation of protein models. Protein Science, 9(11):2181-2191, Nov 2000.
[196] Gregory A. Petsko and Dagmar Ringe. Protein Stucture and Function. New Science Press, 2004.
[197] MICHAEL PETUKHOV, DAVID CREGUT, Clàudio M. SOARES, and LUIS SERRANO. Local water bridges and protein conformational stability. Protein Science, 8:1982-1989, 1999.
[198] Amira Pierucci-Lagha, Richard Feinn, Vania Modesto-Lowe, Robert Swift, Maggie Nellissery, Jonathan Covault, and Henry R. Kranzler. Effects of rapid tryptophan depletion on mood and urge to drink in patients with co-morbid major depression and alcohol dependence. Psychopharmacology, 171(3):340-348, 2004.
[199] G. C. Pimentel and A. L. McClellan. Hydrogen bonding. Annual Review of Physical Chemistry, 22(1):347-385, 1971.
[200] S. B. Prusiner. Prions. Proceedings of the National Academy of Sciences, USA, 95(23):1336313383, Nov 1998.
[201] Sergei Radaev and Peter Sun. Recognition of immunoglobulins by Fc $\gamma$ receptors. Molecular Immunology, 38(14):1073-1083, 2002.
[202] Deepa Rajamani, Spencer Thiel, Sandor Vajda, and Carlos J. Camacho. Anchor residues in protein-protein interactions. Proceedings of the National Academy of Sciences, USA, 101:11287-11292, 2004.
[203] J. J. Ramsden. Review of optical methods to probe protein adsorption at solid-liquid interfaces. J. Stat. Phys., 73:853-877, 1993.
[204] R. Riek, G. Wider, M. Billeter, S. Hornemann, R. Glockshuber, and K. Wuthrich. Prion protein NMR structure and familial human spongiform encephalopathies. Proceedings of the National Academy of Sciences, USA, 95(20):11667-11672, Sep 1998.
[205] Dagmar Ringe. What makes a binding site a binding site? Curr. Opin. Struct. Biol., 5:825829, 1995.

Draft: February 28, 2008, do not distribute
[206] G. G. Roberts, editor. Langmuir-Blodgett Films. Plenum Press, New York, 1990.
[207] John Ròmer, Thomas H. Bugge, Leif R. Lund, Matthew J. Flick, Jay L. Degen, and Keld Danò. Impaired wound healing in mice with a disrupted plasminogen gene. Nature Medicine, 2(3):287-292, 1996.
[208] N Saitou and M Nei. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol Biol Evol, 4(4):406-425, 1987.
[209] Lukasz Salwinski, Christopher S. Miller, Adam J. Smith, Frank K. Pettit, James U. Bowie, and David Eisenberg. The Database of Interacting Proteins: 2004 update. Nucl. Acids Res., 32(database issue):D449-451, 2004.
[210] R. T. Sanderson. Electronegativity and bond energy. Journal of the American Chemical Society, 105:2259-2261, 1983.
[211] Seppo J. Karrila Sangtae Kim. Microhydrodynamics : Principles and Selected Applications. Dover Publications, 2005.
[212] LEISTER J. SAUNDERS, GREGORY B. YOUNG, and GARY J. PIELAK. Polarity of disulfide bonds. Protein Science, 2(7):1183-1184, 1993.
[213] Lindsay Sawyer and Michael N. G. James. Carboxyl-carboxylate interactions in proteins. Nature, 295:79-80, Jan 1982. 10.1038/295079a0.
[214] Ida Schomburg, Antje Chang, Christian Ebeling, Marion Gremse, Christian Heldt, Gregor Huhn, and Dietmar Schomburg. BRENDA, the enzyme database: updates and major new developments. Nucl. Acids Res., 32(database issue):D431-433, 2004.
[215] Laurent Schwartz. Théorie des distributions. Hermann, Paris, 1966.
[216] Eric Schwegler, Matt Challacombe, and Martin Head-Gordon. Linear scaling of the Fock matrix. II. rigorous bounds on exchange integrals and incremental Fock build. J. Chem. Phys., 106:9708-9717, 1997.
[217] L. Ridgway Scott, Mercedes Boland, Kristina Rogale, and Ariel Fernández. Continuum equations for dielectric response to macro-molecular assemblies at the nano scale. Journal of Physics A: Math. Gen., 37:9791-9803, 2004.
[218] J. Seelig and A. Seelig. Lipid conformation in model membranes and biological membranes. Q. Rev Biophys, 13(1):19-61, Feb 1980.
[219] Min-Yi Shen and Karl F. Freed. Long Time Dynamics of Met-Enkephalin: Comparison of Explicit and Implicit Solvent Models. Biophysical Journal, 82(4):1791-1808, 2002.
[220] Z. Shi, C.A. Olson, and N.R. Kallenbach. Cation- $\pi$ interaction in model $\alpha$-helical peptides. Journal of the American Chemical Society, 124(13):3284-3291, 2002.

Draft: February 28, 2008, do not distribute
[221] Zhengshuang Shi, Kang Chen, Zhigang Liu, Tobin R. Sosnick, and Neville R. Kallenbach. PII structure in the model peptides for unfolded proteins - studies on ubiquitin fragments and several alanine-rich peptides containing qqq, sss, fff, and vvv. Proteins - Structure, Function, and Bioinformatics, 63(2):312-321, 2006.
[222] H. Shull and G. G. Hall. Atomic units. Nature, 184:1559-1560, Nov 1959. 10.1038/1841559a0.
[223] U. C. Singh and Peter A. Kollman. A water dimer potential based on ab initio calculations using Morokuma component analyses. Journal of Chemical Physics, 83(8):4033-4040, 1985.
[224] O. Sipahioglu, S.A. Barringer, I. Taub, , and A.P.P. Yang. Characterization and modeling of dielectric properties of turkey meat. JOURNAL OF FOOD SCIENCE, 68(2):521-527, 2003.
[225] John C. Slater. Solid-State and Molecular Theory: A Scientific Biography. John Wiley \& Sons, New York, 1975.
[226] John C. Slater. The Calculation of Molecular Orbitals. John Wiley \& Sons, New York, 1979.
[227] G. R. Smith and M. J. Sternberg. Prediction of protein-protein interactions by docking methods. Curr Opin Struct Biol, 12:28-35, 2002.
[228] Jared D. Smith, Christopher D. Cappa, Kevin R. Wilson, Benjamin M. Messer, Ronald C. Cohen, and Richard J. Saykally. Energetics of hydrogen bond network rearrangements in liquid water. Science, 306:851-853, 2004.
[229] Peter Sondermann, Robert Huber, Vaughan Oosthuizen, and Uwe Jacob. The 3.2-å crystal structure of the human IgG1 Fc fragment-Fc $\gamma$ RIII complex. Nature, 406:267-273, 2000.
[230] Mattthew C. Strain, Gustavo E. Scuseria, and Michael J. Frisch. Achieving linear scaling for the electronic quantum Coulomb problem. Science, 271:51-53, 1996.
[231] P. A. Suci and G. G. Geesey. Comparison of adsorption behavior of two mytilus edulis foot proteins on three surfaces. Colloids Surf. B Biointerfaces, 22:159-168, 2001.
[232] D. I. Svergun, S. Richard, M. H.J. Koch, Z. Sayers, S. Kuprin, and G. Zaccai. Protein hydration in solution: Experimental observation by x-ray and neutron scattering. Proceedings of the National Academy of Sciences, USA, 95(5):2267-2272, 1998.
[233] Attila Szabo and Neil S. Ostlund, editors. Modern Quantum Chemistry. Dover, 1996.
[234] Charles Tanford. Hydrophobic Effect. Wiley, 1973.
[235] Charles Tanford and Jacqueline Reynolds. Nature's Robots, a history of proteins. Oxford Univ. Press, 2001.
[236] Chung-Jung Tsai, S. L. Lin, Haim J. Wolfson, and Ruth Nussinov. Studies of protein-protein interfaces: A statistical analysis of the hydrophobic effect. Protein Science, 6:53-64, 1997.

Draft: February 28, 2008, do not distribute
[237] Jerry Tsai, Robin Taylor, Cyrus Chothia, and Mark Gerstein. The packing density in proteins: Standard radii and volumes. Journal of Molecular Biology, 290:253-266, 1999.
[238] J. A. Vila, D. R. Ripoll, and H. A. Scheraga. Physical reasons for the unusual alpha-helix stabilization afforded by charged or neutral polar residues in alanine-rich peptides. Proceedings of the National Academy of Sciences, USA, 97(24):13075-13079, Nov 2000.
[239] Jorge A. Vila, Daniel R. Ripoll, Myriam E. Villegas, Yury N. Vorobjev, and Harold A. Scheraga. Role of hydrophobicity and solvent-mediated charge-charge interactions in stabilizing alpha-helices. Biophysical Journal, 75:2637-2646, 1998.
[240] D. Vitkup, E. Melamud, J. Moult, and C. Sander. Completeness in structural genomics. Nature Structural Biology, 8:559-566, 2001.
[241] D. Voet and J. G. Voet. Biochemistry. John Wiley \& Sons, New York, 1990.
[242] Timothy C. Wallstrom. Inequivalence between the Schrödinger equation and the Madelung hydrodynamic equations. Phys. Rev. A, 49:1613-1617, 1994.
[243] Xiaoyuan. Wang, Mikhail. Bogdanov, and William. Dowhan. Topology of polytopic membrane protein subdomains is dictated by membrane phospholipid composition. EMBO J., 21(21):5673-5681, Nov 2002.
[244] A. Warshel and A. Papazyan. Electrostatic effects in macromolecules: fundamental concepts and practical modeling. Curr. Opin. Struct. Biol., 8:211-217, 1998.
[245] Arieh Warshel. Electrostatic origin of the catalytic power of enzymes and the role of preorganized active sites. Journal of Biological Chemistry, 273:27035-27038, 1998.
[246] Michael Waterman. Introduction to Computational Biology. Chapman \& Hall/CRC Press, 1995.
[247] James D. Watson and Francis H. C. Crick. Molecular structure of nucleic acids: A structure for deoxyribose nucleic acid. Nature, 171:737-738, 1953.
[248] Ph. Wernet, D. Nordlund, U. Bergmann, M. Cavalleri, M. Odelius, H. Ogasawara, L. Å. Nåslund, T. K. Hirsch, L. Ojamåe, P. Glatzel, L. G. M. Pettersson, and A. Nilsson. The structure of the first coordination shell in liquid water. Science, 304(5673):995-999, 2004.
[249] Ph. Wernet, D. Nordlund, U. Bergmann, M. Cavalleri, M. Odelius, H. Ogasawara, L.Å. Näslund, T. K. Hirsch, L. Ojamaë, P. Glatzel, L. G. M. Pettersson, and A. Nilsson. The structure of the first coordination shell in liquid water. Science, 304:995-999, 2004.
[250] Maurice Wilkins, A. R. Stokes, and H. R. Wilson. Molecular structure of deoxypentose nucleic acids. Nature, 171:738-740, 1953.

Draft: February 28, 2008, do not distribute
[251] S. T. Wlodek, T. W. Clark, L. R. Scott, and J. A. McCammon. Molecular dynamics of acetylcholinesterase dimer complexed with tacrine. J. Am. Chem. Soc., 119:9513-9522, 1997.
[252] Stanislaw T. Wlodek, Tongye Shen, and J.A.McCammon. Electrostatic steering of substrate to acetylcholinesterase: Analysis of field fluctuations. Biopolymers, 53(3):265-271, 2000.
[253] Gerd Wohlfahrt. Analysis of pH-dependent elements in proteins: Geometry and properties of pairs of hydrogen-bonded carboxylic acid side-chains. Proteins: Structure, Function, and Bioinformatics, 58:396-406, 2005.
[254] Yuri I. Wolf, Steven E. Brenner, Paul A. Bash, and Eugene V. Koonin. Distribution of protein folds in the three superkingdoms of life. Genome Res, 9(1):17-26, Jan 1999.
[255] Chia Kuei Wu, Bing Hu, John P. Rose, Zhi-Jie Liu, Tam L. Nguyen, Changsheng Zheng, Esther Breslow, and Bi-Cheng Wang. Structures of an unliganded neurophysin and its vasopressin complex: Implications for binding and allosteric mechanisms. Protein Science, 10(9):1869-1880, 2001.
[256] Stefan Wuchty. Scale-free behavior in protein domain networks. Molecular Biology and Evolution, 18(9):1694-1702, Sep 2001.
[257] Dong Xu, Chung-Jung Tsai, and Ruth Nussinov. Hydrogen bonds and salt bridges across protein-protein interfaces. Protein Engineering, 10(9):999-1012, 1997.
[258] Wai-Ming Yau, William C. Wimley, Klaus Gawrisc, and Stephen H. White. The preference of tryptophan for membrane interfaces. Biochemistry, 37:14713-14718, 1998.
[259] Tsuyoshi Yokomizo, Masayoshi Nakasako, Toshimasa Yamazaki, Heizaburo Shindo, and Junich Higo. Hydrogen-bond patterns in the hydration structure of a protein. Chemical Physics Letters, 401:332-336, 2005.
[260] R. Zahn, A. Liu, T. Luhrs, R. Riek, C. von Schroetter., F. Lopez Garcia, M. Billeter, L. Calzolai, G. Wider, and K. Wüthrich. NMR solution structure of the human prion protein. Proceedings of the National Academy of Sciences, USA, 97(1):145-150, Jan 2000.
[261] Ruhong Zhou, Xuhui Huang, Claudio J. Margulis, and Bruce J. Berne. Hydrophobic collapse in multidomain protein folding. Science, 305(5690):1605-1609, 2004.
[262] Z. H. Zhou, M. L. Baker, W. Jiang, M. Dougherty, J. Jakana, G. Dong, G. Lu, and W. Chiu. Electron cryomicroscopy and bioinformatics suggest protein fold models for rice dwarf virus. Nature Structural Biology, 8:868-873, 2001.
[263] Z. H. Zhou, W. Chiu, K. Haskell, H. Spears, J. Jakana, F. J. Rixon, and L. R. Scott. Parallel refinement of herpesvirus B-capsid structure. Biophysical Journal, 73:576-588, January 1997.
[264] Z. H. Zhou, M. Dougherty, J. Jakana, J. He, F. J. Rixon, and W. Chiu. Seeing the herpesvirus capsid at 8.5 å. Science, 288:877-880, 2000.

Draft: February 28, 2008, do not distribute
[265] Z. H. Zhou, Sharon J. Macnab, Joanita Jakana, L. R. Scott, W. Chiu, and F. J. Rixon. Identification of the sites of interaction between the scaffold and outer shell in herpes simplex virus-1 capsids by difference electron imaging. Proceedings of the National Academy of Sciences, USA, 95:2778-2783, March 1998.
[266] Matjaz Zorko and Ülo Langel. Cell-penetrating peptides: mechanism and kinetics of cargo delivery. Advanced Drug Delivery Reviews, 57(4):529-545, 2005.


[^0]:    ${ }^{1}$ There are more than twenty biologically related amino acids that have been identified, but we will limit our study to the twenty "classical" amino acids commonly found.

[^1]:    ${ }^{2}$ In this regard tryptophan shares structure similar to the compound psilocybin which is known to fit into the same binding sites as the neurotransmitter serotonin.

[^2]:    ${ }^{1}$ Douglas Hartree (1897-1958) pioneered approximation methods for quantum chemistry calculations.

[^3]:    ${ }^{2}$ The unit of viscosity is named for Jean Louis Marie Poiseuille (1799-1869) who, together with Gotthilf Heinrich Ludwig Hagen (1797-1884) established the basic properties of viscous flow in simple geometries.

