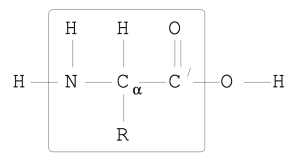
### Protein Docking

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### What Are Amino Acids?

Figure: Illustration of an amino acid. [Neumaier, 2006]



- amino acids polymerize in a specific sequence to a chain
- ▶ the repeating N,  $C_{\alpha}$ , C-pattern is called the protein's backbone

#### Structures

### **Primary Structure**

amino acid sequence

### Secondary Structure

spatial arrangement of amino acid residues that are nearby in the sequence

### Tertiary Structure

its three-dimensional structure, as defined by the atomic coordinates

#### Quaternary Structure

spatial arrangement of multiple folded proteins and the nature of their interactions [Berg, Tymoczko, Stryer, 2002]

### What Is Protein Docking And Why Is It Important?

#### The Problem

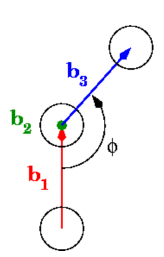
- given tertiary structures, find the most likely quaternary structure
- evaluate its affinity

### **Examples For Quaternary Structures**

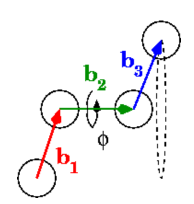
- hemoglobin
- DNA polymerase
- ion channels

Understanding how proteins interact  $\rightarrow$  drug design

▶ bond angle



- ▶ bond angle
- ► dihedral angle



- ▶ bond angle
- ▶ dihedral angle
- internal coordinates

set of bond lenghts, bond angles and dihedral angles; usually only some dihedral angles are allowed to vary though; protein can be seen as a tree-like graph structure

useful to alter a molecule in a chemically meaningful way

- bond angle
- ▶ dihedral angle
- internal coordinates
- absolute coordinates

set of cartesian coordinates of every single atom in a molecule

useful to calculate potentials

### Different Approaches

The problem can be casted as a minimization problem:  $\operatorname{argmin}_X U(X)$ . One can differentiate between several approaches.

#### Classes of Potential Functions

- ▶ **Force fields:** Typically, the potential *U* is a sum over bond, angle, dihedral angle, electrostatic and van der Waals energies.
- ► Knowledge-based/empirical methods: Compare segments with experimentally determined data.

### Choosing a Set of Parameters

- ▶ **Rigid docking:** The parameters X consist of translation and rotation of the smaller protein in  $\mathbb{R}^3$ .
- ▶ Flexible docking: In addition to the 6 parameters of rigid docking, internal parameters (mostly dihedral angles, as bond angles and lengths are relatively stable) are used.

#### The Potential Function In Use

- ▶ let  $X = x, y, z, \alpha, \beta, \gamma, \theta_1, \dots, \theta_n$  free parameters,
- ▶  $M^1(X) = M^1$  be the flexible and  $M^2$  the fixed molecule,
- ▶  $LJ_{M_i,M_j}$  the Lennard-Jones potential, modelling pairwise van der Waals forces, depending on the types of the atoms  $M_i, M_j$  and
- ▶  $||M_i M_j||$  the euclidian distance of their cartesian coordinates.

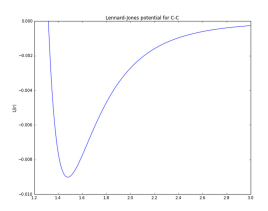
$$U(X) = \underbrace{\sum_{i=1}^{|M^1|} \sum_{j>i}^{|M^1|} LJ_{M_i^1,M_j^1}(\|M_i^1 - M_j^1\|)}_{\text{internal energy of } M^1} + \underbrace{\sum_{i=1}^{|M^1|} \sum_{j=1}^{|M^2|} LJ_{M_i^1,M_j^2}(\|M_i^1 - M_j^2\|)}_{\text{interaction of } M^1 \text{ and } M^2}$$

### Physical Interpretation of LJ

Setting

$$LJ_{M_i,M_j}(r) = \frac{A_{M_i,M_j}}{r^{12}} - \frac{B_{M_i,M_j}}{r^6}$$

for  $A_{M_i,M_j}, B_{M_i,M_j} \in \mathbb{R}^+$  yields the following with a strong divergence at r=0.



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for  $A_{M_i,M_i}, B_{M_i,M_i} \in \mathbb{R}^+$  leads to the following interpretation:

- ▶ A corresponds to the strength of the Pauli-repulsion.
- B corresponds to the attractive long-range term.
- $\varepsilon := \min_{r} LJ(r) = \frac{B^2}{4A}$  is the depth of the potential well.
- $ightharpoonup r_m := \underset{r}{\operatorname{argmin}} LJ(r) = \sqrt[6]{2\frac{A}{B}}$  determines the equilibrium distance of the two elements  $M_i, M_j$ .

#### cons

 Various other physical properties are neglected, e.g. dipole-dipole interactions leading to secondary structures like the α-helix.

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- ▶ All models are simply fitted mathematical functions  $\rightarrow$  treating  $C, C_{\alpha}, O, N, \ldots$  as different elements might account for other physical properties to some extent.

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- ▶ All models are simply fitted mathematical functions  $\rightarrow$  treating  $C, C_{\alpha}, O, N, \ldots$  as different elements might account for other physical properties to some extent.
- It works to a certain extent, i.e.  $\nabla_X U(X) \approx 0$  if X describes the initial configuration which should be an equilibrium state.

### Finding Optimal Parameters

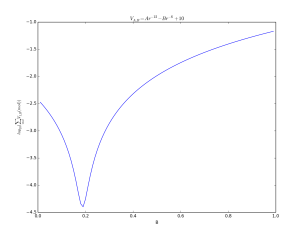
Observe that "M is in equilibrium"  $\Leftrightarrow \nabla_X U_{A,B}(X_M,M)=0$ . Thus, it is natural to choose  $A,B\in (R^+)^{n\times n}$ , where n is the number of different substances one wants to differentiate between, as

$$\underset{A,B}{\operatorname{argmin}} \sum_{M^i \in \operatorname{training set}} \|\nabla_X \mathit{U}_{A,B}(X_{M^i}, M^i)\|.$$

However, this is an overdetermined system  $\to$  fix  $A_{11}=const$  and express all other parameters in terms of  $A_{11}$ , i.e.:  $\tilde{A}_{ij}=A_{11}A_{ij}$ ,  $\tilde{B}_{ij}=A_{11}B_{ij}$ .

### Finding Optimal Parameters

Figure: Projection of the optimization problem in n(n+1)-1 dimensions onto a 1-dimensional subspace ( $B_{11}$  is the only non-constant parameter,  $A_{ij}=1, B_{ij}=B_{11}$ )



### **Numerical Challenges**

#### Conversion Between Internal And Absolute Coordinates

While we want to write the potential as a function of the internal coordinates, it can only be calculated in terms of absolute coordinates  $\rightarrow$  conversion is done often and should thus be implemented efficiently, i.e. minimize rounding errors and duration of computation

### Computation of the Gradient

The potential is not simply a sum over rational functions as the distance between two atoms depends on the internal coordinates  $\rightarrow$  the above conversion has to be included as well; as a result, computing the gradient explicitly is not only a tedious, but also an unnecessary task because of accumulating rounding errors  $\rightarrow$  automatic backwards differentiation

# The End

### References

Arnold Neumaier (2006)

Molecular Modeling of Proteins

