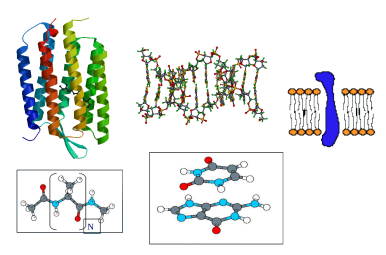
Biomolecular Modeling with Molecular Dynamics Introduction

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2019, April 25

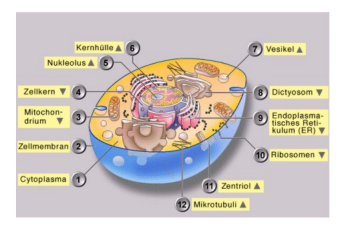
Biomolecular structure

Structural elements of life Biomolecules – proteins, nucleic acids, lipids, carbohydrates ...



Biomolecular structure

Biomolecules \to biomolecular complexes \to aggregates $\to \dots \to$ organelles \to a cell



Biomolecular function

Biophysical processes

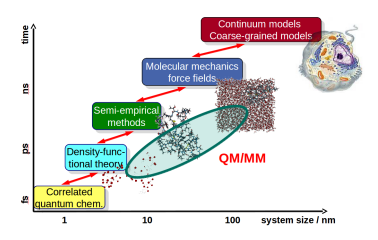
- Bioenergetics reception, transformation and utilization of energy
- Catalysis synthesis of substances, metabolism
- Transport exchange of small molecules, ions up to proteins with the surroundings
- Sensing detection / recognition / binding in presence of a stimulus

Theoretical / computational biophysics

Molecular modeling

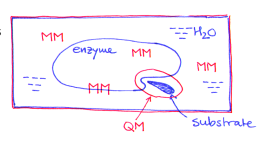
- Focus: understanding on the atomic scale
- The structure and dynamics determine the properties and function of biological molecules
- Prediction of experimentally relevant data
- Molecular design of materials with desired properties
- The way to the goal in general: solution of quantum mechanical many-body problem
- QM MB approach would be too inefficient
 - \rightarrow apply more approximative methods

Methods



Methods

- Quantum chemistry (QM)
 - bonds created/broken
 - computionally costly
 - DFT or ab initio, up to 100 atoms
 - semi-empirical, up to 1000 atoms
- Molecular mechanics (MM)
 - efficient for up to 100,000 atoms
 - generally structural properties
- Hybrid QM/MM
 - chemical reactions etc.



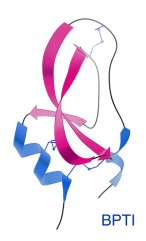
Timeline

- 1687 equations of motion (Newton)
- similar harmonic spring (Hooke)
-
- 1946 molecular mechanics
- 1950s useful computers
- 1959 molecular dynamics of a fluid (Alder & Wainwright)
- 1975 molecular dynamics of a protein (Levitt & Warshel, Gelint & Karplus)
- 1976 QM/MM proposed (Levitt & Warshel)
- 1990 significant QM/MM work (Karplus)

Timeline

1st simulation of protein dynamics

- BPTI, 58 AAs, in vacuo, 9.2 ps
- McCammon, Gelin & Karplus, Nature 1977
- starting point crystal structure



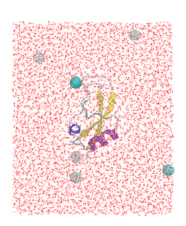
Timeline

simulation of protein in aqueous solution

- BPTI + water, 210 ps
- Levitt & Sharon, PNAS 1988

today's standard

-100,000 atoms, 100 ns



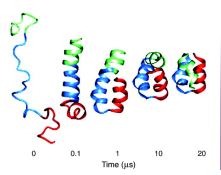
Contact with experimental reality

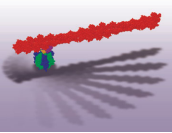
- X-ray and neutron diffraction
- electron cryomiscroscopy (cryo-EM)
- STM/AFM imaging
- electronic / optical spectra
- vibrational / IR spectra
- electronic and nuclear magnetic resonance
- thermodynamic measurements DSC, ITC

Applications

Structure and dynamics of complex biomolecular systems

- protein folding
- protein-ligand interaction
- proteins as nanomachines

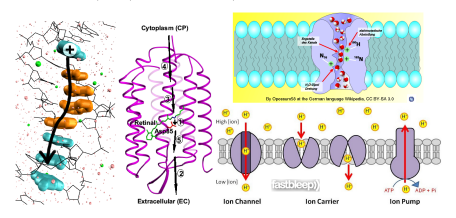




Applications

Transport

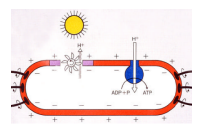
- electrons
- protons across membrane
- water, ions, small molecules, ...

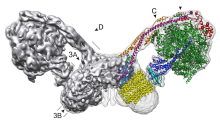


Applications

Enzymes

- catalyzed chemical reactions
- energy conversion (light, chem., mech., gradients)
- bioenergetics





Challenges

- system size limited to ca. 100,000 atoms
- time scales limited to few microseconds
- accuracy of description
 - bonded interaction (vibration, rotation)
 - non-bonded interaction (charge-transfer, polarizability)
- excited electronic states
- quantum character of movement of nuclei

Some can be tackled with

- further development of available methods
- combination of various methods
- optimization of algorithms parallelization, O(N) linear scaling

Nobel prizes for computational chemistry

The Nobel Prize in Chemistry 1998



Walter Kohn Prize share: 1/2



John A. Pople Prize share: 1/2

The Nobel Prize in Chemistry 1998 was divided equally between Walter Kohn "for his development of the density-functional theory" and John A. Pople "for his development of computational methods in quantum chemistry".

Nobel prizes for computational chemistry

The Nobel Prize in Chemistry 2013



© Harvard University Martin Karplus



Photo: © S. Fisch Michael Levitt



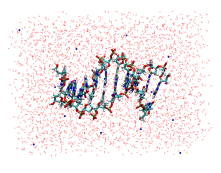
Photo: Wikimedia Commons Arieh Warshel

The Nobel Prize in Chemistry 2013 was awarded jointly to Martin Karplus, Michael Levitt and Arieh Warshel "for the development of multiscale models for complex chemical systems".

Biomolecular simulation

Elementary body - atom

Usually – one molecule/complex of interest (e.g. protein, NA)



Simulation vs. reality

One molecule instead of many Tiny volume of $\approx 10^{-21}$ L instead of $\approx 10^{-5}$ L Dynamics – short time scale of max. $\approx 10^{-5}$ s

Biomolecular simulation

Each atom -x, y, z coordinates "A protein is a set of coordinates." (Gromacs, A. P. Heiner)

```
Peptide in lipid+water
48609
    1LYS
                        4.360
                                4.040
                                         8.207
                                                0.2882
                                                         0.4041 -0.5575
    1LYS
             H1
                        4.416
                                4.119
                                         8.178
                                                0.4151
                                                         0.4652 -0.1555
    1LYS
             H2
                        4.340
                                4.037
                                         8.306
                                                0.8750 -1.7473 -0.4570
    1LYS
             НЗ
                        4.407
                                3.954
                                         8.185
                                                0.3515
                                                        0.2061
                                                                 0.2987
    1LYS
             CA
                        4.231
                                4.037
                                         8.136
                                                0.0777
                                                        0.2898 -0.1753
    1LYS
             HA
                        4.162
                                4.112
                                        8.174 -0.6060 -0.8009
                                                                0.7924
    1LYS
             CB
                        4.262
                                4.069
                                        7.990 -0.3012
                                                        0.3768 -0.2373
    1LYS
            HB1
                        4.360
                                4.025
                                        7.969
                                                0.6025
                                                        1.3517
                                                                 1.6736
                                        7.998
    1LYS
            HB2
                        4.300
                                4.171
                                                0.1892
                                                        0.2300 -0.6695
    1LYS
             CG
                  10
                        4.161
                                4.049
                                         7.877 -0.5841
                                                         0.0286
                                                                 0.0762
    1LYS
            HG1
                  11
                        4.056
                                4.067
                                        7.900 -0.9362 -1.1078 -0.5669
    1LYS
            HG<sub>2</sub>
                  12
                        4.148
                                3.942
                                         7.863 -3.1278
                                                        0.1078
                                                                1.2506
    1LYS
                                4.123
             CD
                  13
                        4.196
                                         7.749
                                                0.0459 -1.0686 -0.3967
    1LYS
            HD1
                  14
                        4.298
                                4.095
                                         7.721 -0.3753 -3.6647
                                                                 0.4016
    1LYS
            HD<sub>2</sub>
                  15
                        4.205
                                4.228
                                         7.778
                                                3.4358 -1.4671
                                                                 0.3786
    1LYS
             CE
                  16
                        4.088
                                4.101
                                         7.644 -0.3622
                                                        0.1377 -0.2469
    1LYS
            HE1
                  17
                        3.992
                                4.138
                                         7.679 -1.1725 -0.1480 -2.0367
    1LYS
            HE2
                  18
                        4.073
                                3.994
                                         7.628 -3.0282
                                                        0.2507
                                                                 0.9872
    1LYS
             NZ
                  19
                        4.124
                                4.174
                                         7.521 -0.0992
                                                        0.0204 -0.2407
                                4.156
    1LYS
            HZ1
                  20
                        4.056
                                         7.449 -2.5018 1.3804
                                                                1.5513
    1LYS
            HZ2
                  21
                        4.118
                                4.275
                                        7.528 -1.2171 -0.0196 -0.4614
```

Simulation vs. reality

Why should we want to simulate molecular systems?

Experiment – the molecule has its genuine properties

Simulation – we need a model to describe the interactions of atoms

– the quality of the model is decisive

Advantage of simulation - structure on atomic level defined

Structure \rightarrow function

Combination of experiment and simulation – added value

Typical hardware



Typical hardware



Cutting-edge hardware





Dror RO, et al. 2012. Annu. Rev. Biophys. 41:429–52