C7790 Introduction to Molecular Modelling TSM Modelling Molecular Structures

Lesson 14 Potential Energy Surface II

PS/2020 Distant Form of Teaching: Rev1

Petr Kulhánek

kulhanek@chemi.muni.cz

National Centre for Biomolecular Research, Faculty of Science Masaryk University, Kamenice 5, CZ-62500 Brno

C7790 Introduction to Molecular Modelling -1-

Context

macroworld microworld

states

(thermodynamic properties, G, T,…)

phenomenological thermodynamics

equilibrium (equilibrium constant) kinetics (rate constant)

free energy (Gibbs/Helmholtz)

Description levels (model chemistry):

- quantum mechanics
	- semiempirical methods
	- ab initio methods
	- post-HF methods
	- DFT methods
- molecular mechanics
- coarse-grained mechanics

Simulations:

partition function

statistical thermodynamics

microstates

(mechanical properties, E)

• molecular dynamics

- Monte Carlo simulations
- docking
- …

microstate ≠ microworld

PES

Potential Energy Surface Properties Visualization Important Points (Stationary States)

Configuration space

 $E(\mathbf{R})$ R = point in 3N dimensional space (N is the number of atoms)

The individual points form a configuration space. **Every point** in the configuration space then represents **unique structure** of the system.

The calculation of the potential energy E(R) is possible by:

- **E** approximate solution of the Schrödinger equation (quantum mechanics, QM)
- using empirical force fields (molecular mechanics, MM)
- hybrid QM/MM approach
- using coarse grained models

The calculation of the potential energy E(R) is possible by:

- **T** approximate solution of the Schrödinger equation (quantum mechanics, QM)
- using empirical force fields (molecular mechanics, MM)
- **hybrid QM/MM approach**
- **using coarse grained models**

overview of method categories

The calculation of the potential energy E(R) is possible by:

- approximate solution of the Schrödinger equation (quantum mechanics, QM)
	- HF method
	- post HF methods (MPn, CI, CC)
	- **DFT methods (various functionals)**
- using empirical force fields (molecular mechanics, MM)
	- **forms and parameters of force fields**
- hybrid QM/MM approach
	- interface, type of QM-MM interaction, link atoms, ...
- using coarse grained models

hundreds of methods differing in the approximations used

they do not affect general laws/properties of E(R)

QM (Quantum mechanics) MM (Molecular mechanics) *CGM (Coarse-grained* **mechanics)**

R - position of atom nuclei R - position of atoms R - position of beads

Potential energy surface can be calculated by various method (model chemistry)!

Graphical representation of functions

Graphical representation of functions

Volumetric representation of functions

Ensing, B.; Klein, M. L. Perspective on the Reactions between F– and CH3CH2F: The Free Energy Landscape of the E2 and SN2 Reaction Channels. *PNAS* **2005**, *102* (19), 6755–6759. <https://doi.org/10.1073/pnas.0408094102>.

Displaying E(R)

1 atom

$$
E(x_1, y_1, z_1)
$$

only volumetrically

2 atoms

$$
E(x_1, y_1, z_1, x_2, y_2, z_2)
$$

N atoms

$$
E(x_1, y_1, z_1, x_2, y_2, z_2, \ldots, x_N, y_N, z_N) \quad | \quad
$$

cannot be displayed

Example:

enzyme BsoBI has \sim 10000 atoms => 30000 degrees of freedom, it would be necessary to use 30000 + 1 dimensional space for visualization

Property of E(R)

The potential energy is invariant to:

- **displacement** (translation) of entire system **displacement** (translation) of entire system
• **rotation** (rotation) of entire system
• **rotation** (rotation) of entire system
-

without the action of external force fields

(e.g., electrostatic, magnetic, etc.)

Invariance to displacement

$$
R_1 + T = R_2 \qquad T = \{x_T, y_T, z_T, x_T, y_T, z_T, \dots\}
$$

translation vector

!!! Does not apply to shift in a force field !!!

 $E(R_1) \neq E(R_2)$

Invariance to rotation

 $R_1 = R_2$ rotation matrix

!!! Does not apply to rotation in a force field !!!

 $E(R_1) \neq E(R_2)$

Diatomic molecule

 $E(x_1, y_1, z_1, x_2, y_2, z_2)$

hydrogen molecule

- three degrees of translational freedom
- two rotational degrees of freedom (molecule is linear)

Diatomic molecule

Thriatomic molecule

 $E(x_1, y_1, z_1, x_2, y_2, z_2, x_3, y_3, z_3)$ water

water molecule

- three degrees of translational freedom
- three rotational degrees of freedom

Triatomic molecule

 $E(x_1, y_1, z_1, x_2, y_2, z_2, x_3, y_3, z_3)$ water water molecule three degrees of translational freedom three rotational degrees of freedom $E(r_1, r_2, \theta)$ 9-6=3 r_1 r_2 θ Internal coordinates r_1 , r_2 , θ displayable only volumetrically

Graphical representation of E(R)

- \triangleright The E(R) function is not graphically representable due to its high dimensionality.
- \triangleright Thus, only its relevant part that best captures the problem being studied is displayed in a two- or three-dimensional space

Graphical representation of E(R)

- \triangleright The E(R) function is not graphically representable due to its high dimensionality.
- ➢ Thus, only its relevant part that best captures the problem being studied is displayed in a two- or three-dimensional space

C7790 Introduction to Molecular Modelling -21-

Illustrative example

 x_1 , x_2 and x_3 are stationary points (local extrema of the function). The properties of the quantum states of the system are derived from them.

$$
E_k = E(x_1) + E_{VRT,l}
$$

Function slope is given by **gradient of function** (i.e., first derivative of the function)

$$
\tan(\alpha) = \frac{\partial E(x)}{\partial x}
$$

Condition necessary for a stationary point

$$
\frac{\partial E(x)}{\partial x} = 0
$$

C7790 Introduction to Molecular Modelling -24-

Function slope is given by **gradient of function** (i.e., first derivative of the function)

$$
\tan(\alpha) = \frac{\partial E(x)}{\partial x}
$$

Condition necessary for a stationary point

$$
\frac{\partial E(x)}{\partial x} = 0
$$

C7790 Introduction to Molecular Modelling -25-

Types of stationary points

Determining stationary point type

Taylor series:

$$
E(x + \Delta x) = E(x) + \frac{\partial E(x)}{\partial x} \Delta x + \frac{1}{2} \frac{\partial^2 E(x)}{\partial x^2} (\Delta x)^2 + \dots
$$

Determining stationary point type

Determining stationary point type

Taylor series:
\n
$$
E(x + \Delta x) = E(x) + \frac{\partial E(x)}{\partial x} \Delta x + \frac{1}{2} \frac{\partial^2 E(x)}{\partial x^2} (\Delta x)^2 + ...
$$
\n
$$
E(x + \Delta x) = E(x) + \frac{1}{2} \frac{\partial^2 E(x)}{\partial x^2} (\Delta x)^2 + ...
$$
\ngradient is zero
\ngra
\n
$$
E(x + \Delta x) = E(x) + \frac{1}{2} \frac{\partial^2 E(x)}{\partial x^2} (\Delta x)^2 + ...
$$
\nguare of the equation is

viation is always positive

 (x) 0

 $E(x)$ \qquad

 ∂x^2

 x^2

2 \sim \sim

 $\partial^2 E(x)$

 $2 E(x)$

The value of function increases when deviating from a stationary point, if any **second derivative at a given point has a positive value**. The stationary point is then **local minimum**.

The value of function decreases when deviating from a stationary point, if any **second derivative at a given point has a negative value**. The stationary point is then **local maximum**.

0

0

 > 0

Local minimum:

Local maximum:

!!! required condition !!!

Two-dimensional case

C790 Introduction to Molecular Modelling and Canadian Control of the Canadia

Two-dimensional case

C7790 Introduction to Molecular Modelling -32-

Two-dimensional case

C7790 Introduction to Molecular Modelling -33-

Generalization for E(R)

Stationary point:

 0 of t (R) = $\partial \mathbf{R}$ $\partial E(\mathbf{R})$ **R** $E(\mathbf{R})$ red

required condition, each component of the gradient must be zero

gradient has 3N components

Stationary point type:

Character of the stationary point is determined by **Hessian**, which is a matrix of second derivatives of potential energy.

Not to be confused with Hamiltonian!!!

N number of atoms

Properties of Hessian

Diagonalization of Hessian is a method for finding eigenvalues and eigenvectors.

eigenvector

- 6 (5) eigenvalues are zero it corresponds to the translation and rotation of the system
- remaining eigenvalues:
	- **all positive local minimum**
	- **one negative, other positive first order saddle point**
	- two negative, other positive saddle point of the second order
	- \bullet
	- all negative local maximum

N number of ator

Properties of Hessian

Diagonalization of Hessian is a method for finding eigenvalues and eigenvectors.

eigenvector

- 6 (5) eigenvalues are zero it corresponds to the translation and rotation of the system
- remaining eigenvalues:
	- **all positive local minimum**
	- **one negative, other positive first order saddle point**

chemically significant stationary points

• two negative, other positive - saddle point of the second order

 \bullet

• all negative - local maximum

N number of atom

Why saddle points and local minima only?

C7790 Introduction to Molecular Modelling -37-

Diagonalization of Hessian

C7790 Introduction to Molecular Modelling -38-

Energy, Gradient, Hessian

The calculation of the potential energy E(R) is possible by:

- approximate solution of the Schrödinger equation (quantum mechanics, QM)
	- HF method
	- post HF methods (MPn, CI, CC)
	- **DFT methods (various functionals)**
- using empirical force fields (molecular mechanics, MM)
	- **forms and parameters of force fields**
- hybrid QM/MM approach
	- interface, type of QM-MM interaction, link atoms, ...
- using coarse grained models

hundreds of methods differing in the approximations used

Energy gradient calculation

Energy gradient:

The gradient calculation can be performed:

- **analytically**
- **numerically**

Analytical/Numerical gradient

Analytical gradient calculation is the **preferred method of calculation** in cases where the expression and subsequent calculation of energy derivatives are easy.

computationally more demanding than energy calculation

Numerical gradient calculation is used when the analytical gradient is not available, for example due to the complexity of the implementation of the algorithm for its calculation.

Either **forward differences** (FD) or **central differences** (CD) method can be used to calculate the numerical gradient . In rare cases, it is also possible to use multipoint methods.

The CD method is more accurate than the FD method and therefore the preferred method of gradient calculation.

> *FD requires 2*3*N energy calculations computationally more demanding than analytical gradient calculation*

Calculation of Hessian

Hessian of energy:

it is a matrix; number of components is 3Nx3N

Calculation of Hessian can be implemented:

- **analytically** (**memory and computationally intensive**)
- **numerically** (by the method of central differences)
	- from energies (3 x N x 3 x N x 2 energy calculations)
	- from gradients (3 x N x 2 gradient calculations)

Summary

- \triangleright Quantum states (thermodynamic microstates) from solution of SE are characterized by stationary points on PES.
- \triangleright Stationary point has zero gradient.

Energy and its gradient MUST be calculated at the same level of theory (model chemistry)!

 \triangleright Type of stationary point can be determined from PES curvature at the stationary point (Hessian eigenvalue analysis).

> **Energy, its gradient, and Hessian MUST be calculated at the same level of theory (model chemistry)!**

➢ The most important stationary points are **local minima** (stable states such as reactants, products, intermediates) and **first order saddle points** (transition states).