#### C7790 Introduction to Molecular Modelling TSM Modelling Molecular Structures

#### Lesson 23 Classification of Interactions

**PS/2021** Present Form of Teaching: Rev2

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**C7790 Introduction to Molecular Modelling** 

#### Context



## **Revision: Schrödinger Equation**

Schrödinger equation by its essence provide ultimate description of (bio)chemical systems:

#### $\widehat{H}\psi_k(\mathbf{r},\mathbf{R}) = E_k(\mathbf{R})\psi_k(\mathbf{r},\mathbf{R})$

- > Solution of SR is the **potential energy**  $E_k$  and **wavefunction**  $\psi_k$ .
  - the potential energy quantify strength of inter-atomic interactions
  - the wavefunction provides further information

Remember: use of SE has some dark sides:

- one-electron approximation (correlation energy)
- basis set effects
- Iong-tails of some interactions (dispersion energy in HF and DFT calculations)
- size consistency

By analyzing  $E_k$  and  $\psi_k$  one can classify interactions between atoms to better understand origin of forces that keep them together. Two major categories of interactions between atoms are:

- covalent bonding
- non-covalent interactions

#### **Molecular Mechanics**

Schrodinger equation - quantum mechanical description

$$\hat{H}\psi_{k}^{\mathbf{R}}(\mathbf{r}_{e}) = E_{k}(\mathbf{R})\psi_{k}^{\mathbf{R}}(\mathbf{r}_{e})$$
approximation
electron motions is omitted
(electron motions is implicitly included in empirical parameters)
$$E_{k}(\mathbf{R}) = E_{bonds} + E_{angles} + E_{torsions} + E_{ele} + E_{vdw} + \dots$$
bonded contributions
non-bonded contributions
**Classical physics** - mechanical description
**Molecular Mechanics**

# **Covalent Bonding**

## **Covalent Bonding**

A **covalent bond** is a chemical bond that involves the **sharing of electron pairs** between atoms.

#### Notable types of covalent bonds:

- $\succ$   $\sigma$ -bonds (each atom formally contributes by one electron)
- $\succ \pi$ -bonds (each atom formally contributes by one electron)
- coordinate covalent bonds (one atom contributes two electrons (a lone pair) and the second atom provides a vacant (empty) orbital)
- aromatic bonding
- metallic bonding
- three-center two-electron bond (see boron chemistry)
- ▶ ....

#### https://en.wikipedia.org/wiki/Covalent\_bond

## **Covalent Bonding, cont.**

#### $\sigma$ and $\pi$ covalent bonds:





Bond dissociation energies:

~85 kcal/mol





triple C-C bond (1  $\sigma$ -bond + 2  $\pi$ -bonds)

120.3 pm

106.0 pm

~145 kcal/mol

~200 kcal/mol

#### Conjugated $\pi$ -bonds (aromaticity):



**Coordinate covalent bonds:** 



https://en.wikipedia.org

#### **C7790 Introduction to Molecular Modelling**

### **Molecular Deformations**

- Some molecular deformations can be described by vibrational motions.
- > There is 3N-6(5) unique molecular vibrations, which are called **normal modes**.

Model of harmonic oscillator:

- A normal vibrational mode is a molecular motion, in which ALL atoms oscillate at the same frequency and phase.
- However, some vibrations are more "localized". Meaning that such vibrations exhibit larger amplitudes only on a few atoms, while the rest of the molecule is almost restful.
- > The other vibrations represent skeletal molecular deformations.



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**Bond stretching** 

### **Molecular Deformations, cont.**

Some types of molecular vibrations



down

"no" motion

### **Molecular Deformations, cont.**

In chemistry, conformational isomerism is a form of stereoisomerism in which the isomers can be interconverted just by rotations about formally single bonds (refer to figure on single bond rotation).



# Non-covalent Interactions

#### **Non-covalent Interactions**

Contribution	Additive?	Sign	Comment
Long-range ( $E(R) \sim R^{-n}$ )			
Electrostatic	Yes	+/-	Strong orientation dependence
Induction	No	-	
Dispersion	Approx.	-	Always present
Resonance	No	+/-	Degenerate states only
Magnetic	Yes	+/-	Very small
Short-range ( $E(R) \sim e^{-\alpha R}$ )			
Exchange-repulsion	Approx.	+	Dominates at very short range
Exchange-induction	Approx.	+	
Exchange-dispersion	Approx.	+	
Charge transfer	No	-	Donor-acceptor interactions

Stone, A. J.; Oxford University Press. *The Theory of Intermolecular Forces*; Oxford University Press: Oxford, 2016.

### **HW: Recommended Readings**

**Read Introduction and Overview of given interaction type:** 

Rackers, J. A.; Wang, Q.; Liu, C.; Piquemal, J.-P.; Ren, P.; Ponder, J. W. An Optimized Charge Penetration Model for Use with the AMOEBA Force Field. *Phys. Chem. Chem. Phys.* **2016**, *19* (1), 276–291. <u>https://doi.org/10.1039/C6CP06017J</u>.

Rackers, J. A.; Liu, C.; Ren, P.; Ponder, J. W. A Physically Grounded Damped Dispersion Model with Particle Mesh Ewald Summation. *J Chem Phys* **2018**, *149* (8), 084115. <u>https://doi.org/10.1063/1.5030434</u>.

Rackers, J. A.; Ponder, J. W. Classical Pauli Repulsion: An Anisotropic, Atomic Multipole Model. *J Chem Phys* **2019**, *150* (8), 084104. <u>https://doi.org/10.1063/1.5081060</u>.