


KAPITAŁ LUDZKI
 NARODOWA STRATEGIA SPÓJNOŚCI

 Projekt współfinansowany przez
 Unię Europejską w ramach
 Europejskiego Funduszu
 Społecznego

UNIA EUROPEJSKA
 EUROPEJSKI
 FUNDUSZ SPOŁECZNY


Course title		ECTS code	
Monographic lecture - Introduction into quantum computer chemistry		13.3.1034	
Name of unit administrating study			
Faculty of Chemistry			
Studies			
faculty	field of study	type	drugiego stopnia
Wydział Chemii	Biznes chemiczny	form	stacjonarne
		specjalty	wszystkie
		specialization	wszystkie
Teaching staff			
prof. dr hab. Janusz Rak			
Forms of classes, the realization and number of hours		ECTS credits	
Forms of classes		3	
Lecture		classes 30 h	
The realization of activities		tutorial classes 10 h	
classroom instruction		student's own work 35 h	
Number of hours		TOTAL: 75 h - 3 ECTS	
Lecture: 30 hours			
The academic cycle			
2023/2024 summer semester			
Type of course		Language of instruction	
obligatory		polish	
Teaching methods		Form and method of assessment and basic criteria for evaluation or examination requirements	
multimedia-based lecture		Final evaluation	
		Graded credit	
		Assessment methods	
		(mid-term / end-term) test	
		The basic criteria for evaluation	
		Passing with no less than 51% of the maximum score. Those who do not reach the required threshold take an oral examination.	
Method of verifying required learning outcomes			
Required courses and introductory requirements			
A. Formal requirements			
none			
B. Prerequisites			
none			
Aims of education			
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Acquisition of the ability to:			
- choose an appropriate computational chemistry method to a given chemical problem,			
- design a computational algorithm assuring possibly swift solution,			
- judge the accuracy of numerical data.			
Course contents			
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Born-Oppenheimer approximation, time independent Schrödinger equation, one-electron approximation, Slater determinant, Hartree-Fock (HF) and			

Hartree-Fock-Roothan (HFR) methods, semiempirical schemes of the HFR method: CNDO, INDO, NDDO; modified NDDO methods: MNDO, AM1, P<3, M5, RM1, PM6, MNDO/d, SAM1, SAM1d. Basis sets. Electron correlation: configuration interaction, Møller–Plesset perturbation theory (MPn), coupled cluster method (CC). Density functional theory (DFT). Application of HFR and electron correlation methods: choice of the basis set, optimization of molecular geometry, determining reaction enthalpy, harmonic vibrational modes (IR spectrum), NMR shifts and electronic absorption spectrum for a molecular system.

Bibliography of literature

Bibliography of literature

Literature required to pass the course

Lucjan Piela „Idee chemii kwantowej”, PWN 2003.

Frank Jensen „Introduction to Computational Chemistry”, Wiley, 2006.

Christopher J. Cramer „Essentials of Computational Chemistry: Theories and Models”, Wiley, 2004

Extracurricular readings

Attila Szabo, Neil S. Ostlund „Modern Quantum Chemistry: Introduction to Advanced Electronic Structure Theory”, Dover Publications, 1996.

The learning outcomes (for the field of study and specialization)

Knowledge

Skills

Social competence

Contact

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