

Subject card

Subject name and code	, PG_00035160								
Field of study	Engineering and Technologies of Energy Carriers								
Date of commencement of studies	February 2025		Academic year of realisation of subject			2025/2026			
Education level	second-cycle studies		Subject group			Optional subject group Subject group related to practical vocational preparation			
Mode of study	Full-time studies		Mode of delivery			at the university			
Year of study	1		Language of instruction			Polish			
Semester of study	2		ECTS credits			3.0			
Learning profile	practical profile		Assessment form			assessment			
Conducting unit	Department of Physical Chemistry -> Faculty of Chemistry								
Name and surname	Subject supervisor		dr hab. inż. M	dr hab. inż. Maciej Śmiechowski					
of lecturer (lecturers)	Teachers								
Lesson types and methods	Lesson type	Lecture	Tutorial	Laboratory	Project	t	Seminar	SUM	
of instruction	Number of study hours	15.0	0.0	30.0	0.0		0.0	45	
	E-learning hours inclu	uded: 0.0		.					
Learning activity and number of study hours	Learning activity Participation in classes include plan				Self-study SUM				
	Number of study hours	45		3.0		27.0		75	
Subject objectives	The aim of the subject is to familiarize the student with the issue of molecular modeling of catalytic processes with a particular consideration of practical aspects of running computer simulations.								
Learning outcomes	Course out	come	Subject outcome			Method of verification			
	K7_U06		Student selects a computational method appropriate for a posited simulation problem considering the computational complexity of the problem and the time consumption of the required calculations.			[SU4] Assessment of ability to use methods and tools [SU2] Assessment of ability to analyse information			
	K7_U07		Student analyzes studied catalytic processes with respect to molecular modifications of reactants ensuring the achievement of better efficiency or selectivity of the process.			[SU4] Assessment of ability to use methods and tools [SU1] Assessment of task fulfilment			
	K7_U05		Student can relate an experimental catalytic process with its molecular mechanism and proposes a simulation-based approach to verify its course.			[SU3] Assessment of ability to use knowledge gained from the subject [SU2] Assessment of ability to analyse information			
	K7_W09		Student knows and understands the molecular mechanism of basic catalytic processes used in chemical technology, especially in the field of heterogeneous catalysis.			[SW1] Assessment of factual knowledge			

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Subject contents	 Repetition in chemical kinetics: reaction order, reaction rate, rate constant and its determination, determination of reaction order, composite reactions, stationary state approximation, temperature dependence of rate constant, Arrhenius equation, activation energy, collision theory, transition state theory and calculation of rate constant, reactions in solutions Repetition in surface science: physical and chemical adsorption, adsorption isotherms, Gibbs isotherm, adsorption from solutions, dispersed phases Repetition in catalysis: catalysts and active centers, homogeneous catalysis, autocatalysis, heterogeneous catalysis (structure of catalysts, metallic catalysts, semiconductor catalysts, stages of a catalyzed reaction) Overview of computational chemistry: molecular mechanics, molecular dynamics, Hartree-Fock method, Kohn-Sham equations, density functional theory, semi-empirical methods, force fields, QM/MM and ONIOM hybrid methods Thermochemical calculations: geometry optimization, vibrational analysis, thermochemical properties, thermodynamics of chemical reactions, optimization to transition state, calculation of rate constant Surface modeling methods: Bravais and reciprocal lattice, periodic boundary conditions, Bloch waves, Brillouin zone, band structure, Fermi level and band gap, density of states, computationa problems (GW method, Hubbard model) Solvation effects: polarizable continuum model (PCM), COSMO model, cluster-continuum models, molecular descriptors in modeling: population analysis, electrostatic potential, localized orbitals, bond order analysis, chemical reactivity indexes (chemical potential, hardness, electrophilicity index, Fukui functions) Reaction path modeling: definition of reaction coordinate, intrinsic reaction coordinate (IRC method), potential energy hypersurface, rare events dynamics, exploration of free energy hypersurface (perturbation methods, thermodynamic integration, umbr						
Prerequisites	Knowledge of material from mathematics and physics in the range of basic academic course in 1st level						
and co-requisites	studies. Basic knowledge of chemical kinetics. Suggested: introductory information in quantum chemistry or physics.						
Assessment methods	Subject passing criteria	Passing threshold	Percentage of the final grade				
and criteria	<u>'</u>	50.0%	50.0%				
	laboratory reports	50.0%	50.0%				
Recommended reading	Basic literature	 P. W. Atkins, Chemia fizyczna, PWN, Warszawa 2001. K. Pigoń, Z. Ruziewicz, Chemia fizyczna 1. Podstawy fenomenologiczne, PWN, Warszawa 2005. K. Pigoń, Z. Ruziewicz, Chemia fizyczna 2. Fizykochemia molekularna, PWN, Warszawa 2005. 					
	Supplementary literature	 Warszawa 2001. E. T. Dutkiewicz, Fizykochemia 1998. L. Piela, Idee chemii kwantowe R. F. Nalewajski, Podstawy i m PWN, Warszawa 2001. A. Kaczmarek-Kędziera, M. Zie 	E. T. Dutkiewicz, Fizykochemia powierzchni, WN-T, Warszawa 1998. L. Piela, Idee chemii kwantowej, PWN, Warszawa 2003 R. F. Nalewajski, Podstawy i metody chemii kwantowej. Wykłady, PWN, Warszawa 2001. A. Kaczmarek-Kędziera, M. Ziegler-Borowska, D. Kędziera, Chemia obliczeniowa w laboratorium organicznym, Wydawnictwo				
	eResources addresses	Adresy na platformie eNauczanie:					
Example issues/ example questions/ tasks being completed	 Define the continuous Fukui function and its three discrete approximations. Relate the reactivity of a molecule towards different types of substitutions to the appropriate discrete functions. The optimization of the geometry of a quantum chemical system has led to a stationary point. How can we distinguish whether it is a local energy minimum or a transition state? How can we predict the direction of deformation of the transition state to a stable structure? Briefly describe the applications of the polarizable continuum model (PCM) in modeling molecules in solution. Define the Fermi level for a solid. Based on its position, divide solids into conductors/semiconductors/insulators. Describe the idea and applications of the internal reaction coordinate (IRC) method. List and briefly characterize three methods of population analysis. Discuss the individual rungs of the "Jacob's ladder" of electron density functionals. Discuss the application of metadynamics in the exploration of the potential energy surface of a system undergoing a chemical reaction. Give the text of Hohenberg-Kohn theorem II. Discuss the basic elements of a typical flexible force field used in molecular mechanics. 						
Work placement	Not applicable						

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