

Subject card

Subject name and code	Materials Science - classical particle approach, PG_00063960								
Field of study	Nanotechnology								
Date of commencement of studies	October 2024		Academic year of realisation of subject			2025/2026			
Education level	second-cycle studies		Subject group			Specialty subject group Subject group related to scientific			
						research in the field of study			
Mode of study	Full-time studies		Mode of delivery			at the university			
Year of study	2		Language of instruction			English			
Semester of study	3		ECTS credits			6.0			
Learning profile	general academic profile		Assessmer	ssessment form		assessment			
Conducting unit	Division of Physics of Disordered Systems -> Institute of Nanotechnology and Materials Engineering -> Faculty of Applied Physics and Mathematics -> Wydziały Politechniki Gdańskiej								
Name and surname	Subject supervisor		dr hab. inż. Jacek Dziedzic						
of lecturer (lecturers)	Teachers		dr inż. Szymon Winczewski						
	dr hab. inż. Jacek Dziedzic								
Lesson types and methods of instruction	Lesson type	Lecture	Tutorial	Laboratory	Projec	t	Seminar	SUM	
	Number of study hours	30.0	0.0	45.0	0.0		0.0	75	
	E-learning hours included: 0.0								
	eNauczanie source address: https://enauczanie.pg.edu.pl/2025/course/view.php?id=1135								
	Moodle ID: 1135 Materials Science - classical particle approach (zima 25/26) https://enauczanie.pg.edu.pl/2025/course/view.php?id=1135								
Learning activity and number of study hours	Learning activity	Participation in didactic classes included in study plan		Participation in consultation hours		Self-study		SUM	
	Number of study hours	75		5.0		70.0		150	
Subject objectives	Discussion of particle methods in the context of modelling of nanoscale systems. Detailed description of the molecular dynamics method in theory (integration of equations of motion) and in practice (commonly used potentials, boundary conditions, starting a simulation, neighbourhood, cut-off radius). Brief tour of more advanced concepts of MD (selected topics e.g. rigid molecules, shell model, constrained dynamics, thermostats, barostats, Ewald method).								

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Learning outcomes	Course outcome	Subject outcome	Method of verification			
	[K7_U03] has enhanced abilities of using advanced specialist software packages	The student is able to independently perform a molecular dynamics simulation using the LAMMPS package and to interpret the basic results.	[SU5] Assessment of ability to present the results of task [SU1] Assessment of task fulfilment			
	[K7_W01] has extended and organized knowledge of materials science.	The student is familiar with the concepts of ordering, phase transition. They have acquired extended and well-ordered knowledge pertaining to simulations with classical particle methods in the field of materials science.	[SW3] Assessment of knowledge contained in written work and projects [SW1] Assessment of factual knowledge			
	[K7_U06] can plan and conduct theoretical and numerical calculations, simulations of phenomena and processes, critically analyze their results, draw conclusions and formulate reasoned conclusions – within their specialization.	The student is able to design and perform a simple simulation using the LAMMPS package, together with a critical analysis of obtained results, visualize the system's trajectory, prepare plots of key parameters of the simulation.	[SU5] Assessment of ability to present the results of task [SU4] Assessment of ability to use methods and tools [SU2] Assessment of ability to analyse information [SU1] Assessment of task fulfilment			
	[K7_W05] has enhanced knowledge of mathematical, numerical, simulation, classical and quantum methods, applied in modeling nanostructures.	The student has in-depth understanding of particle methods (molecular dynamics) and is aware of the methods' limitations. (S)he can position classical and quantum-based methods in the landscape of computational methods suitable for the nanoscale.	[SW1] Assessment of factual knowledge			
Subject contents	Main questions of modeling at the nanoscale. What is a particle? Dynamical equation. Classical and quantum-based methods, scaling of computational effort. The molecular dynamics method, its advantages and limitations. Conservation of energy in Newtonian mechanics. Phase space and trajectories. Periodic ar mixed boundary conditions, minimum image convention, quasiinifinity, limitations of PBCs. Cut-off radius and its consequences. Hockneys linked cells and Verlet neighbour list. Initializing an MD simulation (positions, velocities), equilibration. Integration of the equations of motion. Verlet, leapfrog and predictor-corrector methods. Sources of error in integrating the equations of motion. Visualization in MD, calculating macroscopic observables (energy, temperature, virial, pressure, specific heat, RDF, ADF, S(k), MSD, D(T)) Potential and its relationship with force. General and particular forms of potentials. Selected potentials: LJ, soft- and hard-sphere, Born-Mayer, harmonic, Morse, Stillinger-Weber, Sutton-Chen, GAFF, AMOEBA). Polarizability and shell models (Cochran, Fincham). Constrained dynamics, formal approach, SHAKE, RATTLE, QSHAKE. Rigid molecules in MD simulations, Euler angles, rotation matrix, vector transformations, quaternions. Coulombic interactions in MD, Ewald method. NVT and NPT ensembles, primitive thermostats, ESM and CSM thermo- and barostats. Hybrid (QM/MM) methods.					
Computer lab: The aim is to familiarize students with the practical aspects of modeling at the atomic scale, we carried out in a computer laboratory. The scope includes the following topics: Defining and visualizing atomic systems Introduction to the VMD program Processing and presenting numerical calculation results using gnuplot Introduction to the LAMMPS program Preparation, execution, and analysis of molecular dynamics (MD) simulation results Studying the vibrations of the Ar molecule based on MD simulations Determining the specific heat of an argon single crystal using MD simulations Introduction to the OVITO program Determining the bulk modulus of an argon single crystal using MD simulations						
Prerequisites and co-requisites	The student is acquainted with Newtonian mechanics. The student knows the basics of the organization of matter. The student knows the basics of calculus and algebra.					
Assessment methods	Subject passing criteria	Passing threshold	Percentage of the final grade			
and criteria	written assessment	50.0%	50.0%			
	lab reports	50.0%	50.0%			
Recommended reading	Basic literature 1. D.C. Rapaport, The Art of Molecular Dynamics Simulation Cambridge University Press, 2004.					

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	Supplementary literature	 D. Frenkel, Understanding Molecular Simulation, Academic Press, 2001. M.P. Allen, D.J. Tildesley, Computer Simulation of Liquids, Clarendon Press, 1989. V. Bulatov, W. Cai, Computer simulations of dislocations, Oxford University Press, 2006. E.B. Tadmor, R.F. Miller, Modeling Materials, Cambridge University Press, 2011.
		5. Materials provided by the teacher.
	eResources addresses	
Example issues/ example questions/ tasks being completed		
	Resources addresses 1. What are the typical system sizes and timescales for which molecular dynamics simulations are performed? Why arent continuum methods used for such systems? 2. What are the main differences between classical and quantum-based methods of computational and of nanoscale systems? 3. Briefly explain the principle of operation of the molecular dynamics method. 4. When is total energy conserved in an MD simulation and when is it not? 5. What are the main limitations of the molecular dynamics method? 6. Discuss the notion of periodic boundary conditions what are they, why are they used, what are their limitations and what difficulties are associated with their use? 7. Briefly describe the approaches for making MD simulations faster. 8. What is the potential cut-off radius? Why is it used? What difficulties does using a finite cut-off radius? 9. Describe Hockneys linked-cell method and Verlet neighbour list. What do these techniques allow? are the advantages and disadvantages of both? 10. How would you generate starting positions and velocities for simulating a liquid with MD? 11. How would you generate starting positions and velocities for simulating a reystalline system with M 12. What skew start and how is it realized? 13. What is equilibration in MD simulations? What are the rules of thumb for performing equilibration? 14. Derive the Verlet integrator. 15. What are predictor-corrector approaches? What are their advantages compared to other integrator. 16. What six the pair correlation function? How can it be calculated? 17. Sketch typical shapes of pair correlation functions for a crystalline solid, a liquid and a gas. How d (f) behave as r increases? 18. What additional information on a system can we gather having calculated its pair correlation function? Sketch its typical shape for various systems. 21. How can we calculate the self-diffusion coefficient in an MD simulation? What information does it hold? 22. What is a poential in an MD simulation? How is it related to the	
Work placement	Not applicable	

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