

Subject card

Subject name and code	Materials Science - classical particle approach, PG_00052034							
Field of study	Nanotechnology							
Date of commencement of studies	October 2023		Academic year of realisation of subject		2024/2025			
Education level	second-cycle studies		Subject group		Optional subject group Subject group related to scientific research in the field of study			
Mode of study	Full-time studies		Mode of delivery		at the	at the university		
Year of study	2		Language of instruction		Polish	Polish		
Semester of study	3		ECTS credits		6.0			
Learning profile	general academic profile		Assessment form		assessment			
Conducting unit	Institute of Nanotechnology and Materials Engineering -> Faculty of Applied Physics and Mathematics							
Name and surname of lecturer (lecturers)	Subject supervisor		dr inż. Szymon Winczewski					
	Teachers		Francis Oseko					
			dr inż. Szymon Winczewski					
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							
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 molecular dynamics potentials, boundary advanced concepts of thermostats, barosta	method in theo conditions, sta of MD (selected	ry (integration rting a simulat I topis e.g. ri	of equations of ion, neighbourh	motion) ood, cut	and in off rad	practice (cor lius). Brief tou	nmonly used ur of more

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Extended and well-ordered knowledge pertaining to theoretical and simulation-based classical approaches to the study of annoscale systems. The student is able to employ molecular dynamics to obtain the properties and physical properties straightforward systems. K7_W05	Learning outcomes	Course outcome	Subject outcome	Method of verification		
understanding of particle methods (molecular dynamics) and is aware of the methods (initiations.) (Siyle can position classical and quantum-based methods in the landscape of computational in the landscape of computational manoscale. K7_W01 The student is familiar with the concepts of ordering, phase extended and well-ordered knowledge pertaining to the simulations with classical particle methods in the field of materials science. K7_U06 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 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 (ISU) Assessment of ability the manoscale of the parameters of the simulation (ISU) Assessment of ability the methods and to interper the basic results. K7_U03 The student is able to design and perform a molecular dynamics simulation (ISU) Assessment of ability the parameters of the simulation (ISU) Assessment of ability the parameters of the simulation of the parameters of the parameters of the simulation of the parameters of the		K7_W02	extended and well-ordered knowledge pertaining to theoretical and simulation-based classical approaches to the study of nanoscale systems. The student is able to employ molecular dynamics to obtain the properties and physical properties	[SW1] Assessment of factual knowledge		
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 cannot be supported in the field of materials of the support o		K7_W05	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	[SW1] Assessment of factual knowledge		
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. K7_U03 The student is able to independently perform a molecular dynamics simulation using the LAMMPS package and to independently perform a molecular dynamics simulation using the LAMMPS package and to interpret the basic results. Subject contents Lecture 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 advanta and limitations. Conservation of energy in Newtonian mechanics. Phase space and trajectories. Period mixed boundary conditions, minimum image convention, quasiinfinity, limitations of PBCs. Cut-off radi and its consequences. Hockneys linked cells and Verlet neighbour list. Initializing an MD simulation (positions, velocities), equilibration. Integration of the equations of motion. Visualization in MD, calcular macroscopic observables (energy, temperature, virial, pressure, specific heat, RDF, ADF, S(k), MSD, Potential and its relationship with force. General and particular forms of potentials. Selected potentials: soft- and hard-sphere, Born-Mayer, harmonic, Morse, Stillinger-Weber, Sutton-Chen, GAFF). Polarizat 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 thermo ESM and CSM thermo- and barostats. Hybrid (QM/MM) methods. Computer lab Work on laboratory exercises/example problems, including practical use of simulation methods and techniques discussed in the lecture. Familiarization with scientific software (programs: VMD, gnuplot, LAMMPS, Ovito).		K7_W01	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	[SW3] Assessment of knowledge contained in written work and		
Independently perform a molecular dynamics simulation using the LAMMPS package and to interpret the basic results. Lecture 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 advanta and limitations. Conservation of energy in Newtonian mechanics. Phase space and trajectories. Period mixed boundary conditions, minimum image convention, quasiinifinity, limitations of PBCs. Cut-off radi 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 predict corrector methods. Sources of error in integrating the equations of motion. Visualization in MD, calcula macroscopic observables (energy, temperature, virial, pressure, specific heat, RDF, ADF, SkJM, MSD, I Potential and its relationship with force. General and particular forms of potentials. Selected potentials: soft- and hard-sphere, Born-Mayer, harmonic, Morse, Stillinger-Webers, Sutton-Chen, GAFF), Polarizat 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 thermo ESM and CSM thermo- and barostats. Hybrid (QM/MM) methods. Computer lab Work on laboratory exercises/example problems, including practical use of simulation methods and techniques discussed in the lecture. Familiarization with scientific software (programs: VMD, gnuplot, LAMMPS, Ovito).		K7_U06	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	fulfilment [SU2] Assessment of ability to analyse information [SU4] Assessment of ability to use methods and tools [SU5] Assessment of ability to		
Main questions of modelling at the nanoscale. What is a particle? Dynamical equation.Classical and quantum-based methods, scaling of computational effort. The molecular dynamics method, its advanta and limitations. Conservation of energy in Newtonian mechanics. Phase space and trajectories. Period mixed boundary conditions, minimum image convention, quasiinfinity, limitations of PBCs. Cut-off radii 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 predicts corrector methods. Sources of error in integrating the equations of motion. Visualization in MD, calcular macroscopic observables (energy, temperature, virial, pressure, specific heat, RDF, ADF, S(k), MSD, I Potential and its relationship with force. General and particular forms of potentials. Selected potentials: soft- and hard-sphere, Born-Mayer, harmonic, Morse, Stillinger-Weber, Sutton-Chen, GAFF). Polarizat 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 thermo ESM and CSM thermo- and barostats. Hybrid (QM/MM) methods. Computer lab Work on laboratory exercises/example problems, including practical use of simulation methods and techniques discussed in the lecture. Familiarization with scientific software (programs: VMD, gnuplot, LAMMPS, Ovito).		K7_U03	independently perform a molecular dynamics simulation using the LAMMPS package and	fulfilment [SU5] Assessment of ability to		
Prerequisites		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 and 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). 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.				
	Prerequisites					
and co-requisites The student is acquainted with Newtonian mechanics. The student knows the basics of organization of matter. The student knows the basics of calculus and algebra.	and co-requisites			s the basics of organization of		

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Assessment methods	Subject passing criteria	Passing threshold	Percentage of the final grade		
and criteria	final test	50.0%	40.0%		
	hands-on computer lab	50.0%	60.0%		
Recommended reading	Basic literature	D.C. Rapaport, The Art of Molecular Dynamics Simulation, Cambridge University Press, 2004.			
	Supplementary literature	1. D. Frenkel, Understanding Molecular Simulation, Academic Press, 2001. 2. M.P. Allen, D.J. Tildesley, Computer Simulation of Liquids, Clarendon Press, 1989. 3. V. Bulatov, W. Cai, Computer simulations of dislocations, Oxford University Press, 2006. 4. E.B. Tadmor, R.F. Miller, Modeling Materials, Cambridge University Press, 2011.			
	eResources addresses	Adresy na platformie eNauczanie: Materials Science - Classical Particle Approach 2024/2025 - Moc ID: 41274 https://enauczanie.pg.edu.pl/moodle/course/view.php?id=41274			
Example issues/ example questions/ tasks being completed					
	performed? Why arent continuum m 2. What are the main differences be of nanoscale systems? 3. Briefly explain the principle of ope 4. When is total energy conserved ii 5. What are the main limitations of ti 6. Discuss the notion of periodic boo limitations and what difficulties are a 7. Briefly describe the approaches fi 8. What is the potential cut-off radiu entail? 9. Describe Hockneys linked-cell me are the advantages and disadvantat 10. How would you generate starting 11. How would you generate starting 12. What skew start and how is it re 13. What is equilibration in MD simulation. What are predictor-corrector app 16. What is the pair correlation funct 17. Sketch typical shapes of pair congriphenave as r increases? 18. What additional information on a How can we do that? 19. What is the mean-square displainformation does it hold? 20. What is the angular distribution of 21. How can we calculate the self-decay what is a potential in an MD simplemeral form of a potential used in periodic 12. What is a potential in an MD simplemeral form of a potential used in periodic 12. What is a potential in an MD simplemeral form of a potential used in periodic 12. What is a potential in an MD simplemeral form of a potential used in periodic 12. What is a potential in an MD simplemeral form of a potential used in periodic 12. What is a potential in an MD simplemeral form of a potential used in periodic 12. How can we calculate the self-decay what is a potential in an MD simplemeral form of a potential used in periodic 12. How is a potential in an MD simplemeral form of a potential used in periodic 12. Briefly characterize the GAFF periodic 12. Briefly describe the formal approximation is pressure calculated in M 29. Describe the Ewald approach in 30. What is constrained dynamics? 31. Briefly describe the Berendsen a 33. Describe the three general classimulations.	iffy explain the principle of operation of the molecular dynamics method. en is total energy conserved in an MD simulation and when is it not? at are the main limitations of the molecular dynamics method? cuss the notion of periodic boundary conditions what are they, why are they used, what are their ions and what difficulties are associated with their use? iffy describe the approaches for making MD simulations faster. at is the potential cut-off radius? Why is it used? What difficulties does using a finite cut-off radius? or inche Hockneys linked-cell method and Verlet neighbour list. What do these techniques allow? What e advantages and disadvantages of both? would you generate starting positions and velocities for simulating a liquid with MD? would you generate starting positions and velocities for simulating a crystalline system with MD? hat skew start and how is it realized? hat is equilibration in MD simulations? What are the rules of thumb for performing equilibration? where the Verlet integrator. hat is the pair correlation function? How can it be calculated? tetch typical shapes of pair correlation functions for a crystalline solid, a liquid and a gas. How does shave as r increases? hat additional information on a system can we gather having calculated its pair correlation function? and we do that? hat is the mean-square displacement (MSD) in an MD simulation? How can we calculate it? What ation does it hold? hat is the mean-square displacement (MSD) in an MD simulation? What information does it hold? hat is the pail and MD simulation? How is it related to the force acting on atom i? What is the all form of a potential used in practice? aw the typical shape of the Lennard-Jones potential. What is its functional form? Describe the terms //mbols used. What kinds of systems does this potential describe well? we the formula for the Lennard-Jones potential. What is its functional form? Describe the terms //mbols used. What kinds of systems does this potential describe selected three see. Proper the still			
Work placement	Not applicable				
Work placement					

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