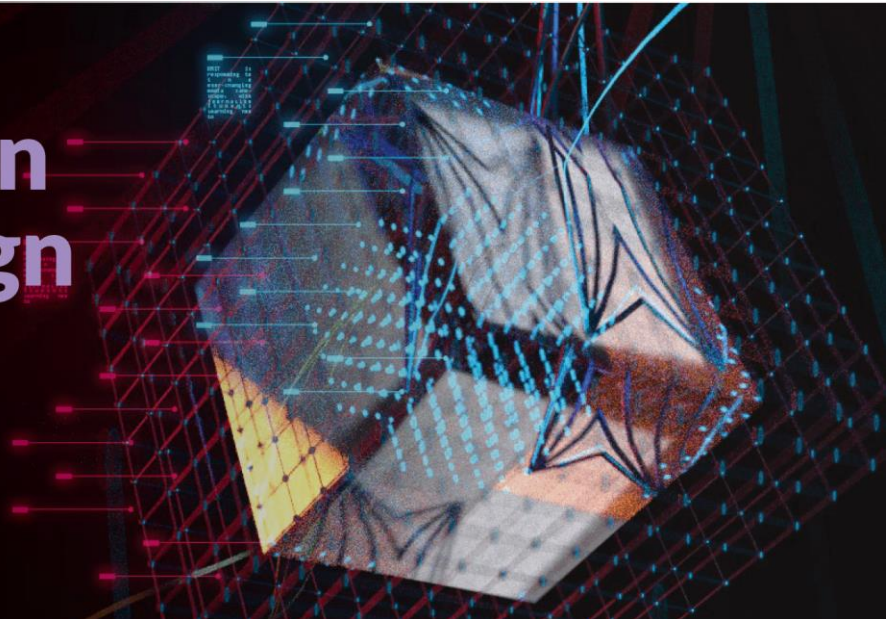


HPC and Data in Materials Design and Discovery

Syllabus



Week	Lecture	Topic	Lecturer
W1	Introduction		
1 Sep	Lecture 1	Overview of HPCD in Materials Design and Discovery <ul style="list-style-type: none"> • General introduction • Summary of course outline • Any additional info re the tutorials 	Prof Sean Smith (ANU)
3 Sep	Lecture 2	Overview of Australian HPC & Data Landscape <ul style="list-style-type: none"> • Australia's capabilities in HPCD research • How we are positioned in the larger international environment • Tier1 Facility - NCI Gadi • Tier1 Facility - Pawsey Setonix 	Overview - Prof Sean Smith (ANU) - 30 min Gadi - Dr Jingbo Wang (NCI) - 15 min Setonix - Dr Maciej Cytowski (Pawsey) - 15 min
W2	Electronic Structure of Finite Molecular Systems - Basics		
8 Sep	Lecture 3	Overview of Methods Used in Electronic Structure Calculations I <ul style="list-style-type: none"> • Intro to semiempirical methods • Intro to density functional theory • Intro to ab initio methods • Intro to force fields and QM/MM methods • Overview of potential energy surfaces 	A/Prof Amir Karton (UWA)

		<ul style="list-style-type: none"> • Overview of molecular properties 	
10 Sep	Lecture 4	<p>Overview of Methods Used in Electronic Structure Calculations II</p> <ul style="list-style-type: none"> • Intro to theory of Intermolecular interactions • Electronic structure methods for calculating intermolecular interactions • Solvation effects in computational chemistry 	Prof Katya Pas (Monash)
W3	Electronic Structure of Finite Molecular Systems – Tutorials on Gadi		
15 Sep	Tut 1	<ul style="list-style-type: none"> • Calculate a potential energy surface for a simple reaction 	A/Prof Karton + NCI Training
17 Sep	Tut 2	<ul style="list-style-type: none"> • Examples of SAPT with Psi4 to extract four fundamental forces in intermolecular complexes • Examples of BSSE calculations with Psi4 & G16 with both MP2 & DFT • Examples of CCSD(T)/CBS with Orca (through DLPNO-CCSD(T)) • Examples of implicit solvation calculations in G16 	Prof Pas + NCI Training
W4	Electronic Structure of Solids – Basics		
22 Sep	Lecture 5	<p>Introduction to the Electronic Structure of Solids</p> <ul style="list-style-type: none"> • Bloch's theorem • The Brillouin zone • Band structure • Density of states • Integrating across the Brillouin 	A/Prof Alister Page (Newcastle)
24 Sep	Lecture 6	<p>Introduction to Practical Electronic Structure Calculations for Solids</p> <ul style="list-style-type: none"> • Choice of basis set • The pseudopotential approximation • Choice of functionals for solids • Car-Parrinello and adiabatic methods • How to choose the most effective code for your problem 	Prof Julian Gale (Curtin)
One week break			

W5		Electronic Structure of Solids – Tutorials on Gadi	
6 Oct	Tut 3	Introduction to Electronic Structure Calculations for Solids	Prof Gale +
8 Oct	Tut 4	Introduction to Electronic Structure Calculations for Solids	NCI Training
W6		Molecular Dynamics Simulations	
13 Oct	Lecture 7	Introduction to MD Simulations I <ul style="list-style-type: none"> • Equations of motion • Periodic boundary conditions • Thermostatting • Applications: material properties using MD simulations 	Prof Debra Bernhardt (UQ)
15 Oct	Lecture 8	Introduction to MD Simulations II <ul style="list-style-type: none"> • Force fields • Advanced sampling techniques • Applications: use of force fields and advanced sampling in MD simulations 	Prof Ben Corry (ANU)
W7		Molecular Dynamics Simulations – Tutorials on Gadi	
20 Oct	Tut 5	TBC	Prof Bernhardt + NCI Training
22 Oct	Tut 6	TBC	Prof Corry + NCI Training
W8		Monte Carlo & Kinetic Monte Carlo Methods	
27 Oct	Lecture 9	Monte Carlo Simulations <ul style="list-style-type: none"> • MC method (overview & history) • MC simulation of chemical systems (Markov chains, importance sampling, detailed balance) • Sampling at constant NVT (Metropolis algorithm, use of non-physical moves) • Sampling at constant NPT (dealing with additional degrees of freedom) • Sampling at constant μVT (absorption in MOFs, systems with a single interface) • Gibbs ensemble technique (application to phase coexistence) 	Dr Asaph Widmer-Cooper (USyd)

		<ul style="list-style-type: none"> • MC code packages 	
29 Oct	Lecture 10	Kinetic Monte Carlo and Lattice Dynamics <ul style="list-style-type: none"> • Introducing time into Monte Carlo simulation - KMC • Using molecular dynamics to obtain rate constants for KMC • Lattice vs molecular dynamics • Optimization methods for location of local minima • Static calculation of reaction barriers • Transition state theory for estimation of rate constants 	Prof Julian Gale (Curtin)
W9	AI / ML for Materials Design & Discovery		
3 Nov	Lecture 11	Automating Electronic Structure Calculations and Large Materials Data <ul style="list-style-type: none"> • Tools to automate electronic structure calculation • Materials property databases • The Materials Project and its API 	Prof Shyue Ping Ong (UC San Diego)
5 Nov	Lecture 12	Machine Learning for Materials Design at Scale <ul style="list-style-type: none"> • Introduction to machine learning (ML) and its role in materials science • Developing ML models • Challenges in ML for materials science 	
W10	AI / ML for Materials Design & Discovery Tutorial on Gadi		
10 Nov	Tut 7	TBC	Dr Sherif Abdulkader Tawfik (Deakin) +
12 Nov	Tut 8	TBC	NCI Staff
W11	Domain Application: Biomaterials		
17 Nov	Lecture 13	Molecular Simulations of Bio-interfaces: Fundamentals and Challenges <ul style="list-style-type: none"> • Introduction to bio-interfaces & typical "simulation appropriate" research questions • Importance of interfacial structural models and understanding their limitations • Describing inter-atomic interactions across the interface 	Prof Tiffany Walsh (Deakin)

		<ul style="list-style-type: none"> • Conformational sampling at bio-interfaces • Example applications and pitfalls 	
19 Nov	Lecture 14	<p>Applications and Examples of Molecular Simulations for Biomaterials</p> <ul style="list-style-type: none"> • Emphasis on connections between the purpose(s) of the investigation and the simulation techniques applied • Highlights of good practise and hazards to avoid in simulation approaches applied to biomaterials • Examples taken from sensing, health, and biotechnology applications 	
W12	Domain Application: Batteries and Hydrogen		
24 Nov	Lecture 15	<p>Understanding and Design of Advanced Battery Electrolytes through Computational Modelling</p> <p>Batteries and the energy transition</p> <ul style="list-style-type: none"> • Examples of large-scale deployment and battery chemistry types • Is Lithium ion the best chemistry for stationary storage? • Beyond Lithium ion battery chemistries • Computational design of new electrolyte systems with properties prediction • Designing electrolyte compositions, structures, interphases to enhance battery performance 	Prof Maria Forsyth (Deakin)
26 Nov	Lecture 16	<p>The Key Role of Computational Modelling for Advancing Hydrogen Technologies</p> <ul style="list-style-type: none"> • Renewable energy storage: the gigantic challenge • Australian climate change response: economic, political and global trade implications • Why hydrogen? • Key technological challenges and why HPC modelling is critical to advancement • Examples of significant Australian contributions 	Prof Sean Smith (ANU)

~ End ~

