

MLCM-24 schedule

mlcm-24.github.io [updated 05/05/24]

	5/20 MONDAY	5/21 TUESDAY	5/22 WEDNESDAY	5/23 THURSDAY
8:45 - 9:00 AM	Marianne Francois <i>T-division, LANL</i> opening remarks			
9:00 - 9:30 AM	<i>plenary</i> Anatole von Lilienfeld <i>University Of Toronto Vector Institute</i> Quantum Machine Learning	<i>plenary</i> Gabor Csanyi <i>University of Cambridge</i> Foundation models for materials chemistry	<i>plenary</i> Alexandre Tkatchenko <i>University of Luxembourg</i> Navigating Chemical Compound Space with Machine Learning	<i>plenary</i> Markus Reiher <i>ETH Zurich</i> Lifelong Machine Learning Potentials
9:30 - 10:00 AM	<i>plenary</i> Oleg Prezhdo <i>University of Southern California</i> Nonadiabatic Molecular Dynamics with Machine Learning	<i>plenary</i> Mario Barbatti <i>Aix Marseille University, CNRS, ICR</i> Machine Learning Nonadiabatic Dynamics	<i>plenary</i> Roberto Car <i>Princeton University</i> Deep potential models for equilibrium and near equilibrium processes	<i>plenary</i> Olexandr Isayev <i>Carnegie Mellon University</i> AIMNet2: Robust neural network potential for organic, element-organic molecules and chemical reactions
10:00 - 10:20 AM	David Balcells <i>University of Oslo</i> Generative Machine Learning for Transition Metal Chemistry	Reinhard Maurer <i>University of Warwick</i> Machine learning of electronic structure for molecular design	Nong Artrith <i>Debye Institute for Nanomaterials Science</i> ML & XAS for Amorphous Materials	Johannes Margraf <i>University of Bayreuth</i> Materials Discovery with Foundation Models
10:20 - 10:40 AM	Volker Deringer <i>University of Oxford</i> Data-driven interatomic potentials for inorganic materials chemistry	Karel Berka <i>Palacky University Olomouc</i> MolMeDB - free database of molecules on membranes	Zsuzsanna Koczor-Benda <i>University of Warwick</i> Machine learning-based molecular design for plasmonic nanosystems	Matthias Rupp <i>Luxembourg Institute of Science and Technology (LIST)</i> Thermal transport via machine-learning potentials
10:40 - 11:00 AM	Eva Zurek <i>University at Buffalo, SUNY</i> Machine Learned Interatomic Potentials for Binary Carbides Trained on the AFLOW Database	Alexander Shapeev <i>Skolkovo Institute of Science and Technology</i> From quantum mechanics to phase diagrams with machine learning	<i>interactive session</i> Jason Hattrick-Simpers <i>University of Toronto</i> Tutorial on the use of LLMs in Science	Tom Penfold <i>Newcastle Univeristy</i> Deep Neural Networks for X-ray Spectroscopy: Hero or Zero?
11:00 - 11:20 AM	Rohit Goswami <i>University of Iceland / Quansight Labs</i> Throwaway Gaussian Processes for Saddle Searches	Julien Lam <i>CNRS</i> Exploiting constrained linear models for machine-learning interaction potentials		Aurora Clark <i>University of Utah</i> ML in Chemical Separations - Connecting Reaction Stoichiometry with Solution Structure
11:20 - 11:40 AM	Matthew Carbone <i>Brookhaven National Laboratory</i> TBD	Dmitry Zubarev <i>IBM Research</i> Foundational Models for Chemical Discovery		Connor Coley <i>MIT</i> Molecular design and the intersection with synthesis
11:40 am - 1:00 PM	LUNCH BREAK			
1:00 - 1:30 PM	<i>plenary</i> Richard Hennig <i>University of Florida</i> Accelerating Materials Discovery Through Deep Learning and Ultra-Fast Potentials	<i>plenary</i> Kieron Burke <i>University of California, Irvine</i> Machine Learning Density Functionals	<i>plenary</i> Alan Aspuru-Guzik <i>University of Toronto</i> TBA	<i>plenary</i> Rebecca Lindsey <i>University of Michigan, Ann Arbor</i> Explaining Performance of Physics-Informed Machine-Learned Interatomic Models
1:30 - 1:50 PM	Wissam Saidi <i>NETL</i> Materials Modeling and Machine Learning	Y Z <i>University of Michigan</i> Unusual Dynamics of Tetrahedral Liquids Caused by the Competition between Dynamic Heterogeneity and Structural Heterogeneity	Pavlo Dral <i>Xiamen University</i> From fast potentials for dynamics to learning dynamics	Noa Marom <i>Carnegie Mellon University</i> Applications of Machine Learning in First Principles Materials Simulations
1:50 - 2:10 PM	Avanish Mishra <i>LANL</i> Learning from 'Small' Data Using Physics-Based Descriptors	Joshua Rackers <i>Genentech Prescient Design</i> Building Physics into AI for Drug Discovery	Aditya Nandy <i>UCLA University of Chicago</i> Leveraging Community Knowledge to Forge a Path Forward for Transition Metal Complex and Metal-Organic Framework Design	Marivi Fernandez-Serra <i>Stony Brook University</i> Learning the exchange and correlation functional in DFT
2:10 - 2:30PM	Fang Liu <i>Emory University</i> Machine learning aided chemical discovery in the solution phase	Arthur Mar <i>University of Alberta</i> Discovery of Inorganic Solids with Desired Structure Motifs Guided by Machine Learning	Richard Messerly <i>LANL</i> Nanoreactor active learning: Discovering chemistry with a general reactive machine learning potential	David Yaron <i>Carnegie Mellon University</i> Quantum chemical Hamiltonians as flexible and interpretable model forms for machine learning
2:30 - 2:50 PM	Chenru Duan <i>Microsoft</i> Diffusion models on sampling rare events in chemistry	Sakib Matin <i>LANL</i> Machine Learning Potentials with long-range Coulomb interaction: Atomistic simulations of water	Ankur Chaterjee <i>Nicolaus Copernicus University</i> Spatio-temporal dependancies of in-plane and cross-plane thermal transport proerties by hybrid architecture of CNN and RNN	Jenna Pope <i>Pacific Northwest National Laboratory</i> Accelerating Atomic-scale Simulations of Molecules and Materials with Neural Network Potentials
2:50 - 3:10 PM	Rose Cersonsky <i>University of Wisconsin-Madison</i> Data-driven approaches to chemical and materials science:the impact of data selection, representation, and interpretability	Chiho Kim <i>GATech Matmerize, Inc.</i> Machine Learning-Aided Design of Biodegradable Polymers	Bakhtiyor Rasulev <i>North Dakota State University</i> Application of Mixture-type Descriptors in Machine Learning Modeling of Materials	Vignesh Kumar <i>Nicolaus Copernicus University</i> The critical role of XC potential in DFT: Our Big data analysis into DFT
3:10 - 3:30 PM	Johannes Hachmann <i>University at Buffalo</i> TBA	Marina Meila <i>University of Washington</i> TBA	Emily Shinkle <i>LANL</i> Thermodynamic Transferability in Coarse-Grained Force Fields using Graph Neural Networks	Puck van Gerwen <i>EPFL</i> EquiReact: Equivariant Neural Networks for Chemical Reactions
3:30 - 3:50 PM	Will Bricker <i>University of New Mexico</i> Machine-learned electron densities of nucleic acids	Daniel Schwalbe-Koda <i>UCLA</i> Unifying Views on the Extrapolation Power of Machine Learning Potentials and Materials Thermodynamics	Maria Chan <i>Argonne National Laboratory</i> Theory-informed ML for Microscopy, Spectroscopy, and Scattering	Jing Huang <i>Westlake University</i> DP/MM: a hybrid force field model for zinc-protein dynamics