

Artificial Intelligence for Science in Quantum, Atomistic, and Continuum Systems

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Advances in artificial intelligence (AI) are fueling a new paradigm of discoveries in natural sciences. Today, AI has started to advance natural sciences by improving, accelerating, and enabling our understanding of natural phenomena at a wide range of spatial and temporal scales, giving rise to a new area of research known as AI for science (AI4Science). Being an emerging research paradigm, AI4Science is unique in that it is an enormous and highly interdisciplinary area. Thus, a unified and technical treatment of this field is needed yet challenging. This paper aims to provide a technically thorough account of a subarea of AI4Science; namely, AI for quantum, atomistic, and continuum systems. These areas aim at understanding the physical world from the subatomic (wavefunctions and electron density), atomic (molecules, proteins, materials, and interactions), to macro (fluids, climate, and subsurface) scales and form an important subarea of AI4Science. A unique advantage of focusing on these areas is that they largely share a common set of challenges, thereby allowing a unified and foundational treatment. A key common challenge is how to capture physics first principles, especially symmetries, in natural systems by deep learning methods. We provide an in-depth yet intuitive account of techniques to achieve equivariance to symmetry transformations. We also discuss other common technical challenges, including explainability, out-of-distribution generalization, knowledge transfer with foundation and large language models, and uncertainty quantification. To facilitate learning and education, we provide categorized lists of resources that we found to be useful. We strive to be thorough and unified and hope this initial effort may trigger more community interests and efforts to further advance AI4Science.

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1 INTRODUCTION

Decades of artificial intelligence (AI) research has culminated in the renaissance of neural networks [LeCun et al. 1998] under the name of deep learning. Since AlexNet [Krizhevsky et al. 2012], a decade of intensive research has led to many breakthroughs in deep learning, including, for example, ResNet [He et al. 2016], diffusion and score-based models [Ho et al. 2020; Song et al. 2020], attention, transformers [Vaswani et al. 2017], and recently large language models (LLM) and ChatGPT [OpenAI 2023], etc. These developments have led to continuously improved performance for deep models. When coupled with growing computing power and large-scale datasets, deep learning methods are becoming dominant approaches in various fields, such as computer vision and natural language processing. Propelled by these advances, AI has started to advance natural sciences by improving, accelerating, and enabling our understanding of natural phenomena at a wide range of spatial and temporal scales, giving rise to a new area of research, known as AI for science. It is our belief that AI for science opens a door for a new paradigm of scientific discovery and represents one of the most exciting areas of interdisciplinary research and innovation.

Historically, the importance of computing in accelerating discoveries in natural sciences has been noted. Almost one hundred years ago in 1929, the quantum physicist Paul Dirac stated that *“The underlying physical laws necessary for the mathematical theory of a large part of physics and the whole of chemistry are thus completely known, and the difficulty is only that the exact application of these laws leads to equations much too complicated to be soluble.”* In quantum physics, it is known that the Schrödinger’s equation provides precise descriptions of behaviors of quantum systems, but solving such an equation is only possible for very small systems due to its exponential complexity. In fluid mechanics, the Navier-Stokes equations describe spatiotemporal dynamics of fluid flows, but solving these equations of practically useful sizes is highly demanding, especially when computing efficiency is also required. Similar to these two examples, the underlying physics of many natural science problems are known and can be described by a set of mathematical equations. The key difficulty lies in how to solve these equations accurately and efficiently. Recent studies have shown that deep learning methods can accelerate the computing of solutions for these equations. For example, deep learning methods have been used to compute the solutions of Schrödinger’s equation in quantum physics [Pfau et al. 2020; Hermann et al. 2020, 2022] and Navier-Stokes equations in fluid mechanics [Kochkov et al. 2021b; Brunton et al. 2020]. In these areas, simulators are employed to compute solutions of mathematical equations, and the results are used as data to train deep learning models. Once trained, these models can make predictions at a speed that is much faster than simulators. In addition to improved efficiency, deep learning models have been shown to exhibit better out-of-distribution (OOD) generalization, with scope extended to much wider practical settings, where training and unseen data usually follow different distributions.

In other areas such as biology, the underlying biophysical process is not completely understood and may not ultimately be described by mathematical equations. In these cases, experimentally generated data can be used to train deep learning models in order to model the underlying biophysical process. For example, in biology, AI systems, such as AlphaFold [Jumper et al. 2021], RoseTTAFold [Baek et al. 2021], and ESMFold [Lin et al. 2022a], trained on experimentally acquired 3D structures, enable the computational prediction of protein 3D structures at an accuracy comparable to experimental results. In addition to technical challenges, a key element in these areas is the availability of large amounts of experimentally generated data. For example, the success of AlphaFold, RoseTTAFold, and ESMFold highly relies on the large amount of protein 3D structure data generated using experiments and deposited into databases, such as the Protein Data Bank.

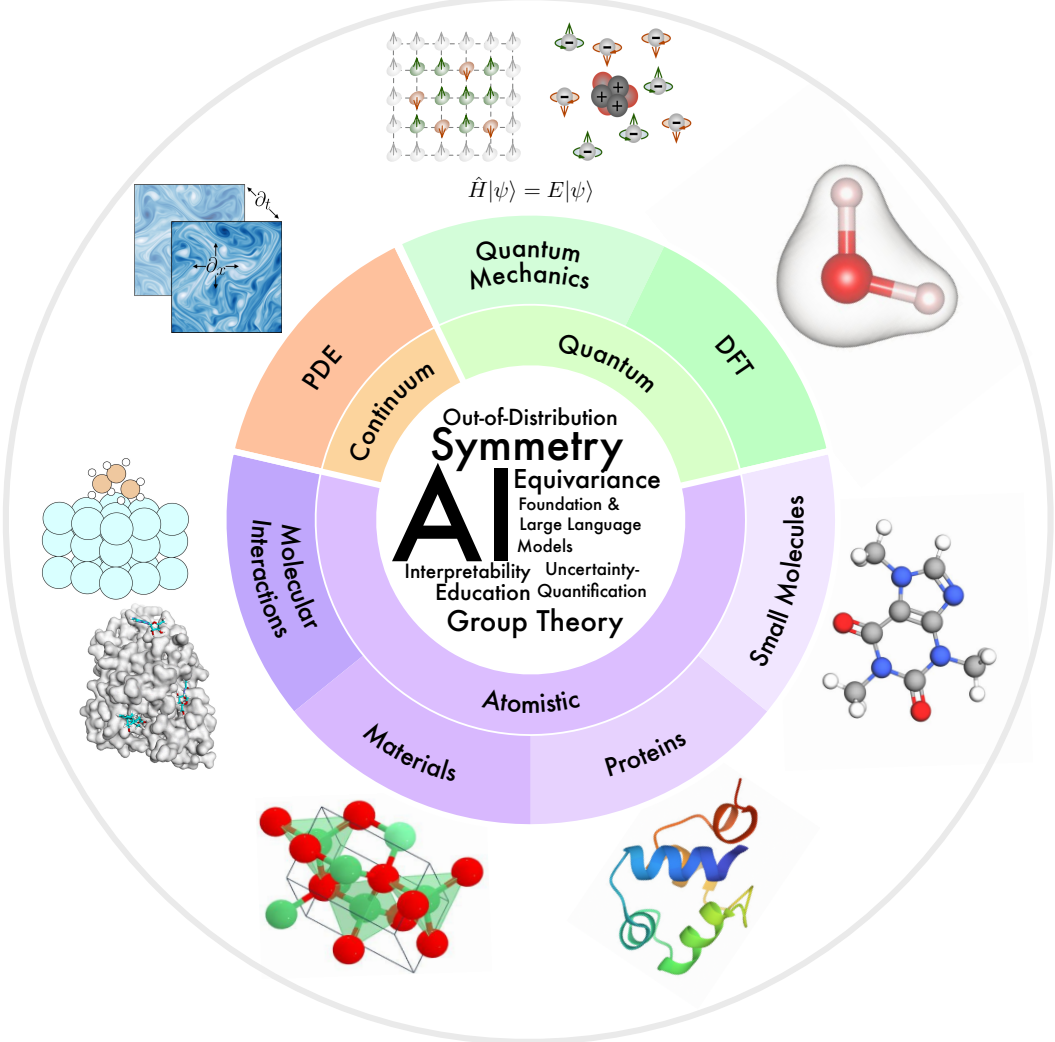


Fig. 1. An integrative overview of the selected research areas in AI for science. As described in Section 1.1, we focus on AI for *quantum mechanics*, *DFT*, *small molecules*, *proteins*, *materials*, *molecular interactions*, and *PDE*. We visually depict these diverse areas in the outermost circle. These areas are arranged by their respective spatial and temporal scales of physical world modeling, highlighting *quantum*, *atomistic*, and *continuum* systems. Notably, as summarized in Section 1.2, a set of common technical considerations and challenges, such as *symmetry*, *interpretability*, and *out-of-distribution generalization*, exist across these multiple AI for science research areas. We show these technical areas in the innermost circle.

1.1 Scientific Areas

In this work, we provide a technical and unified review of several research areas in AI for science that researchers have been working on during the past several years. We organize different areas of AI for science by the spatial and temporal scales at which the physical world is modeled. An overview of scientific areas we focus in this paper is given in Figure 1.

Quantum Mechanics studies physical phenomena at the smallest length scales using wavefunctions, which describe the complete dynamics of quantum systems. In quantum physics, wavefunctions are obtained by solving the Schrödinger equation, which incurs exponential complexity. In this work, we provide technical reviews on how to design advanced deep learning methods for learning neural wavefunctions efficiently.

Density Functional Theory (DFT) and *ab initio* quantum chemistry approaches are first-principles methods widely used in practice to calculate electronic structures and physical properties of molecules and materials. However, these methods are still computationally expensive, limiting their use in small systems ($\sim 1,000$ atoms). In this work, we present technical reviews on deep learning methods for accurately predicting quantum tensors, which in turn can be used to derive many other physical and chemical properties, including, electronic, mechanical, optical, magnetic, and catalytic properties of molecules and solids. We also touch on machine learning methods for density functional learning.

Small Molecules, also known as micromolecules, typically have tens to hundreds of atoms and play important regulatory and signaling roles in many chemical and biological processes. For example, 90% of approved drugs are small molecules, which can interact with target macromolecules (like proteins), altering the activity or function of the target. In recent years, significant progress has been made in using machine learning methods to accelerate scientific discoveries on small molecules at the atomistic level. In this work, we present in-depth technical reviews on small molecule representation learning, molecular generation, simulation, and dynamics.

Proteins are macromolecules that consist of one or more chains of amino acids. It is commonly believed that amino acid sequences determine protein structures, which in turn determines their functions. Proteins perform most of the biological functions, which include structural, catalytic, reproductive, metabolic, and transporting roles, etc. Recently, machine learning approaches have led to dramatic advances in protein structure prediction [Jumper et al. 2021; Baek et al. 2021; Lin et al. 2022a]. In this work, we provide technical reviews on how to learn representations from protein 3D structures, and how to generate and design novel proteins.

Materials Science studies the relationship of processing, structure, properties, and performance of materials. The intrinsic structure of materials from atomistic, to micro and continuum scale determine their quantum, electronic, catalytic, mechanical, optical, magnetic, and other properties through interplay with external stimuli/environment. Recently, machine learning methods have been developed to predict the properties of crystal materials and design novel crystal structures. In this work, we provide technical reviews on the property prediction and structure generation of crystal materials.

Molecular Interactions study how molecules interact with each other to carry out many of the physical and biological functions. Recent advances in machine learning have spurred the renaissance in modeling various molecular interactions, such as ligand-receptor and molecule-material interactions. In this work, we present in-depth and comprehensive reviews on such advances.

Continuum Mechanics models physical processes that evolve in time and space at the macroscopic level using partial differential equations (PDEs), including fluid flows, heat transfer, and electromagnetic waves, etc. However, solving PDEs using classic solvers suffers from several limitations, including low efficiency, difficulties in out-of-distribution generalization and multi-resolution analysis. In this work, we provide reviews on recent deep learning methods for surrogate modeling that addresses these limitations.

In each area, we provide a precise problem setup and discuss the key challenges of using AI to solve such problems. We then provide a survey of major approaches that have been developed. We also describe datasets and benchmarks that have been used to evaluate machine learning methods. Finally, we summarize the remaining challenges and propose several future directions in each research area. When applicable, we include the recommended prerequisite sections at the beginning of each subsection to indicate inter-section dependencies. The overall taxonomic structure is summarized as Figure 2. This paper presents a comprehensive taxonomy, anchored by the shared mathematical and physical principles of symmetry, equivariance, and group theory, delving into seven specific domains within the realm of AI for science, and discussing common technical challenges existing in multiple areas. This enables a comprehensive and structured exploration of AI for science.

1.2 Technical Areas of AI

We have observed that a set of common technical challenges exist in multiple areas of AI for science.

Symmetry: A common and recurring observation from many scientific problems is that objects or systems of interests usually contain geometric structures. In many cases, these geometric structures imply certain symmetries that the underlying physics obeys. For example, in molecular dynamics, molecules are represented as graphs in 3D space, and translating or rotating a molecule may not change its properties. Then the symmetry here is named translational or rotational invariance. Formally, a symmetry is defined as a transformation that, when applied on an object of interest, leaves certain properties of the object unchanged (invariant) or changed in a deterministic way (equivariant) [Bronstein et al. 2021]. Symmetries are very strong inductive biases, as P. Anderson (1972) stated that “*It is only slightly overstating the case to say that physics is the study of symmetry.*” [Anderson 1972]. Thus, a key challenge of AI for science is how to effectively integrate symmetries in AI models. We use symmetry as the main common thread to connect many of the topics in this paper. The required symmetries for each area are also summarized in Figure 3.

Interpretability: Science aims at understanding the governing rules of the physical worlds. Thus, the aims of AI for science are to (1) design models capable of modeling the physical world accurately, and (2) interpret models to verify or discover the governing physics [E et al. 2020]. Thus, interpretability is essential in AI for science.

Out-of-Distribution (OOD) Generalization and Causality: Traditional machine learning methods assume training and test data follow the same distribution. In reality, different distribution shifts may exist between training and test data, raising the need to identify causal factors capable of OOD generalization. OOD generalization is particularly relevant in scientific simulations as this avoids the need to generate training data for every different settings.

Foundation and Large Language Models: When labeled training data are not readily available, the capability to perform unsupervised or few-shot learning becomes important. Recently, foundation models [Bommasani et al. 2021] have demonstrated promising performance on natural language processing tasks. Typically, foundation models are large-scale models pre-trained under self-supervision or generalizable supervision, allowing a wide range of downstream tasks to be performed in few-shot or zero-shot manners. This paradigm is becoming increasingly popular due to the recent developments of large language models (LLM) such as GPT-4. We provide our perspectives on how such a paradigm could accelerate discoveries in AI for science.

Uncertainty Quantification (UQ) studies how to guarantee robust decision-making under data and model uncertainty, and is a critical part of AI for science. UQ has been studied in various disciplines of applied mathematics, computational and information sciences, including scientific

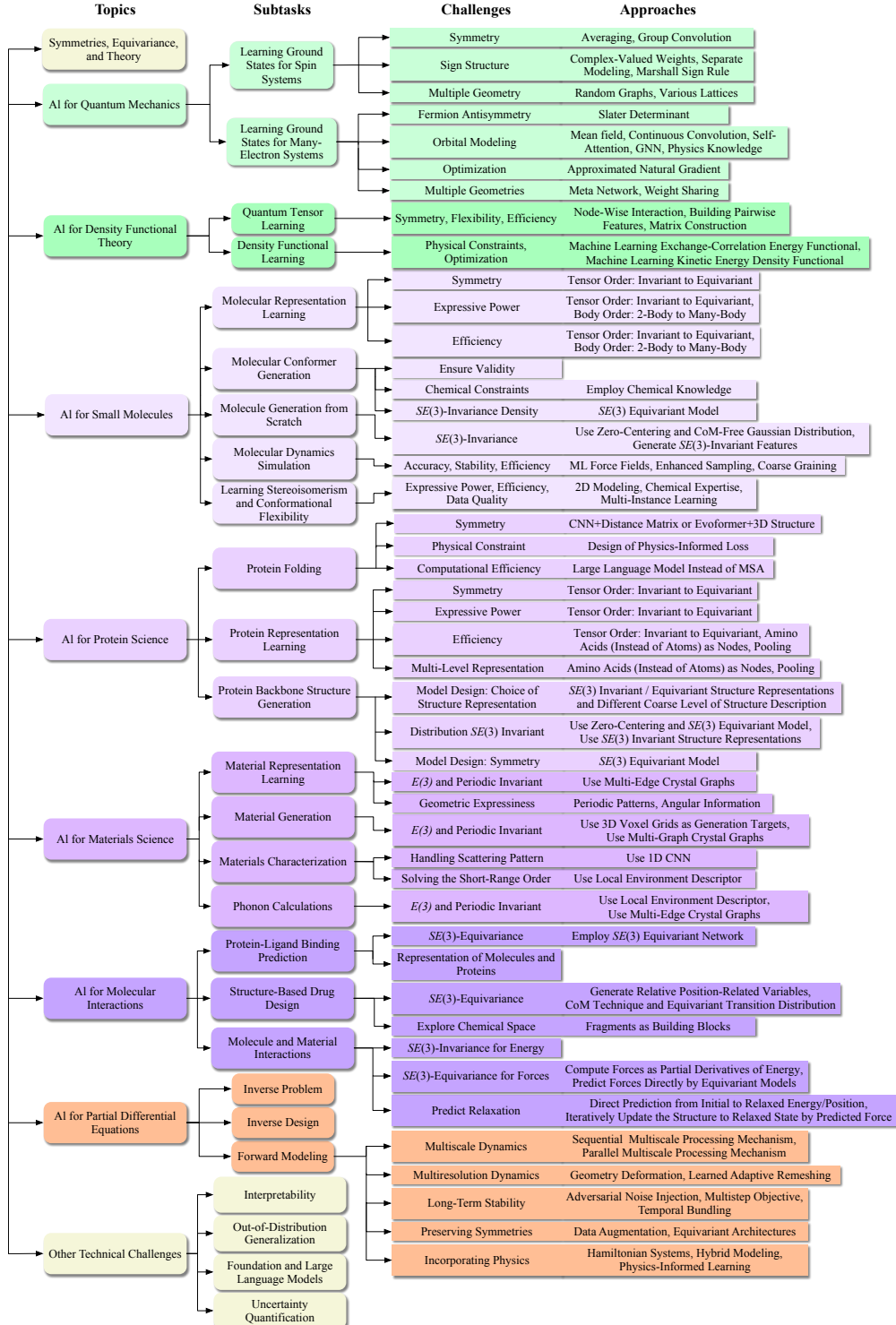


Fig. 2. The overall taxonomic structure of this paper. We outline the areas of AI for science included in this paper and summarize selected problems, central challenges, and the major approaches.

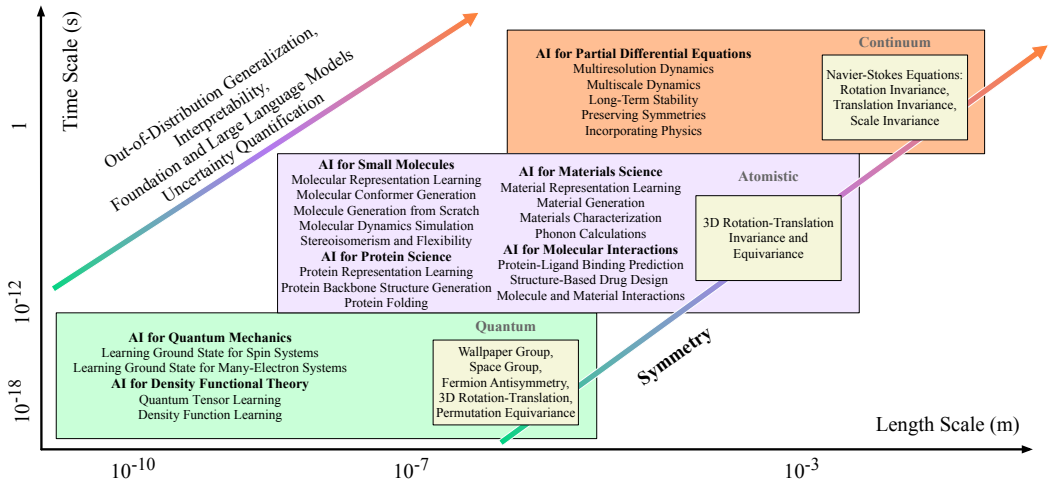


Fig. 3. Spatial and temporal scale of scientific areas. We explore the intersection of AI and various scientific disciplines within a continuum of spatial and temporal scales. This framework accommodates a diverse range of areas and problems, unified by their distinct symmetries and shared technical challenges. Symmetries, inherent to the structure of natural science and governed by mathematical and physical laws, manifest in numerous patterns across various scientific fields. This cross-disciplinary perspective provides a fresh lens through which we can address and investigate complex scientific problems with AI methods.

computation, statistic modeling, and more recently machine learning. We provide an up-to-date reviews of UQ in the context of scientific discoveries.

Education: AI for science is an emerging and rapidly developing area of research with many useful resources developed physically or online. To facilitate learning and education, we have compiled categorized lists of resources that we find to be useful. We also provide our perspectives on how the community can do better to facilitate the integration of AI with science and education.

1.3 Integrative Multi-Scale Analysis

In this survey, we conduct analysis at different levels, including quantum physics, density functional theory (DFT), molecular dynamics (MD), and continuum dynamics. There are notable differences in terms of the level of approximations and the scales they are dealing with. Specifically, quantum physics deals with the behavior and interactions of particles such as electrons, protons, and neutrons, as well as their quantum mechanical properties by solving the Schrödinger's equation for many-body interacting system. The spatial scale in quantum physics is typically on the order of the atomic and subatomic level, ranging from picometer (10^{-12} meters) to nanometer (10^{-9} meters) scale, depending on specific problems. DFT solves the Schrödinger's equation for electrons and ions using an alternative approach by mapping many-body interacting system to many-body non-interacting system, which therefore allows to provide insights into the electronic structure of realistic materials such as atoms, molecules, and solids ranging from angstroms (10^{-10} meters) to hundreds of angstroms. MD simulations operate at a larger scale, typically ranging from the nanometer (10^{-9} meter) to micrometer (10^{-6} meter) scale using empirical/semi-empirical force fields as well as the rising machine learning force fields. MD focuses on the motion and interactions of atoms and molecules over time under various thermodynamic ensembles, allowing for the investigation of dynamic behavior, structural changes, kinetic, and thermodynamic properties. In

comparison, quantum physics aims to solve many-body wavefunctions and Hamiltonian for many-body interacting system; DFT takes an alternative approach with practical applications for molecules and materials; MD simulations operate at a much larger spatial scale and longer time scale without explicitly dealing with spatial and spinor components of electronic wavefunctions. To address even larger scales and eliminate the discrete characteristics of particles, partial differential equations (PDE) are used to study the continuum system behaviors in scales ranging from micrometers (10^{-6} meter, such as the Kolmogorov microscale) in fluid dynamics to kilometers (10^3 meters) in climate dynamics. We compare the spatial and temporal scales of different systems in Figure 3. Accordingly, the focus areas in this work are clustered into quantum, atomistic, and continuum systems. The choice of the theoretical levels depends on the phenomena of interest and the computational complexity required for the study. Different analyses can benefit each other and lead to integrative analysis.

1.4 Online Resources

AI for science is an emerging and rapidly developing area of research. To enable continuous updates of this work, we have created an online portal (<https://air4.science/>), which will be maintained and updated regularly. The online portal contains our assets including a mindmap, which is designed to visualize the taxonomic structure of the various areas covered in our work. This mindmap serves as a comprehensive overview allowing users to navigate and will be updated regularly after the publication of this work to include new topics and significant advancements in the field. In addition, we include a feedback form (<https://air4.science/feedback>) on the portal. This form serves as a channel for individuals to contribute their thoughts, suggestions, and comments regarding this paper. We highly value input from the wider community to improve our work.

This paper is accompanied by a software library and benchmarks under the project repository “AIRS: AI Research for Science” (<https://github.com/divelab/AIRS/>), that we have developed as part of our scientific pursuits in these areas. A set of software libraries have been included and will be added continuously as our research progresses. We also maintain a curated list of literature and resources pertaining to each AI for science topics in the project repository. We welcome contributions from the wider community to both the library and literature via pull requests.

1.5 Scope and Feedback

AI research for science is an enormous and emerging field, and our focus in this work is on AI for quantum, atomistic, and continuum systems. Thus, our work is by no means comprehensive and only includes selected areas of AI for science related to physics, chemistry, biology, material science, molecular simulation and dynamics, and partial differential equations, etc. Given the evolving nature of this area, our work is by no means conclusive in any sense. We expect to continuously include more methods and benchmarks as the area develops. AI for science is highly interdisciplinary, and there is no doubt that we have missed relevant work in the literature, for which we must apologize. We welcome any feedback and comments from the community to improve our work. Readers are encouraged to submit their feedback to us via the above online portal.

1.6 Contributions and Authorship

This work was initiated and conceptualized by Shuiwang Ji, who also leads the distributed writing process and provides scientific and administrative support throughout the project. Each of the individual sections was written by a subset of authors, and authorship is given in each section. Given that all these sections are related, there have been extensive discussions across sections. Authorship is based on the amounts of direct contributions to each section, including texts, equations, figures, tables, discussions, and feedback, etc. Contributions are approximately quantified based on the

number of pages to which each author contributes in the final paper, slightly adjusted based on levels of difficulties and thus discussions required. Many authors have provided constructive discussions and feedback, which have also been considered. When multiple authors work on a part collaboratively, percentage of contributions from each author is estimated and used in the calculation. Authorship for the entire paper is determined based on the cumulative contributions made to all sections. All authors have made significant contributions to this work, and their orders should be interpreted only in an approximate sense.

1.7 Notations

We adopt standard mathematical notation in this work. Scalars are denoted by lowercase letters, such as a , while boldface lowercase letters, such as \mathbf{a} , are used to denote vectors. Matrices are denoted by uppercase letters, such as A , with their ij -th entry denoted as a_{ij} and their k -th column denoted as \mathbf{a}_k . Tuples or sets are denoted by calligraphic uppercase letters, such as \mathcal{A} . The rules hold for all notations except for those with special meanings, in which case we use their conventional forms. For example, the Hamiltonian matrix is denoted by H , the coefficient matrix in DFT by C , and energy scalars by E and V . We provide a summary of common notations shared by multiple sections followed by key notations for individual directions.

Notation of Particle Systems: We denote an n -body particles system, such as a molecule, material, and a protein, by a tuple of matrices $\mathcal{M} = (A, C)$, where A denotes the particle attributes and $C = [\mathbf{c}_1, \dots, \mathbf{c}_n] \in \mathbb{R}^{3 \times n}$ represents the Cartesian coordinates of particles in the system. Specifically, when only particle types are used as the attributes, we denote the system by $\mathcal{M} = (z, C)$, where $z \in \mathbb{Z}^n$ is a vector representing the types, such as atom charges. Additional attributes of a system can be included in the tuple, such as a material $\mathcal{M} = (z, C, L)$ with a lattice matrix L .

Notation of Transformations: We denote the rotation transformation by $R_\alpha : \mathbb{R}^{n \times n} \rightarrow \mathbb{R}^{n \times n}$ with an angle α , who can be represented by a rotation matrix $R \in \mathbb{R}^{3 \times 3}$. The corresponding order- ℓ Wigner-D matrix is denoted by $D^\ell(R)$. We represent the translation transformation by a vector $t \in \mathbb{R}^3$. Consequently, an $E(3)$ -transformation on C is denoted as $RC + t\mathbf{1}^T$.

Dirac Notation: Dirac notation, named after Paul Dirac, is commonly used in quantum physics to represent quantum states. In this notation, a quantum state is denoted by a ket vector, written as $|\psi\rangle$, a column vector in a complex vector space. The conjugate transpose of a ket vector is represented by a bra vector, written as $\langle\psi|$, which is a row vector. The inner product between a bra and a ket is denoted as $\langle\phi|\psi\rangle$, yielding a complex number. The outer product of a ket and a bra is represented as $|\psi\rangle\langle\phi|$, resulting in a complex matrix. Operators can be applied to quantum states by writing them to the left of the ket vector, such as $\hat{O}|\psi\rangle$, representing a matrix-vector multiplication.

Key Notations in Individual Sections: Other notations are defined individually for each area. We summarize the key notations in each direction in Table 1.

Table 1. Summary of key notations. Notations used in a single area are individually defined in the table and in each section.

Sections	Key notations
Sec. 2	Input signal $X \in \mathbb{R}^{s \times s}$, convolution kernel $W \in \mathbb{R}^{k \times k}$, convolution operator $*$. Spherical harmonics functions $Y^\ell(\cdot) : \mathbb{R}^3 \rightarrow \mathbb{R}^{2\ell+1}$, node feature $\mathbf{h}_i^{\ell_1} \in \mathbb{R}^{2\ell_1+1}$, message $\mathbf{m}_i^{\ell_3} \in \mathbb{R}^{2\ell_3+1}$, CG matrix $C_{\ell_1, \ell_2}^{\ell_3} \in \mathbb{R}^{(2\ell_3+1) \times (2\ell_1+1)(2\ell_2+1)}$, Widger-D matrix $D^\ell(R) \in \mathbb{R}^{(2\ell+1) \times (2\ell+1)}$.
Sec. 3	Wavefunction ψ or $ \psi\rangle$, a spin configuration $ \sigma^{(i)}\rangle$, number of spins N , number of electrons of a certain spin N^\uparrow, N^\downarrow . Electron coordinates $\mathbf{r} = [r_1, \dots, r_{N^\uparrow+N^\downarrow}]$. Set of possible molecules $\mathbb{M} = \{M = \{\mathbf{c}_i, z_i\}_{i=1}^{ M }, \mathbf{c}_i \in \mathbb{R}^3, z_i \in \mathbb{Z}\}$. Electron orbital network $\phi_\theta^\uparrow, \phi_\theta^\downarrow$, determinants: $\det [\dots]$, local energy E_{loc} , Hamiltonian matrix for spin systems $H \in \mathbb{C}^{2^N \times 2^N}$, Hamiltonian operator \hat{H} , potential energy V .
Sec. 4	Wavefunction ψ or $ \psi\rangle$, number of orbitals N_o , \mathbf{r}_i electron position, electronic wavefunction coefficients matrix $C_e \in \mathbb{R}^{N_o \times N_o}$ or $\mathbb{C}^{N_o \times N_o}$ (depending on the nature of physical systems), Hamiltonian matrix $H := H_{DFT} \in \mathbb{R}^{N_o \times N_o}$ or $\mathbb{C}^{N_o \times N_o}$ (depending on the nature of physical systems), overlap matrix $S \in \mathbb{R}^{N_o \times N_o}$, eigen energy diagonal matrix $\epsilon \in \mathbb{R}^{N_o \times N_o}$, electron density ρ , energy $E[\rho]$ which is a function of electron density, and external potential $V_{ext}(\mathbf{r})$.
Sec. 5	3D molecule $\mathcal{M} = (z, C)$, where z denotes atom types and C represents coordinates. Distance d_{ij} between two atoms i, j . Scalar feature $s \in \mathbb{R}^d$, vector feature $v \in \mathbb{R}^{d \times 3}$, order- ℓ feature \mathbf{h}_{icm}^ℓ , for node i , channel c , and representation index $-\ell \leq m \leq \ell$. 2D molecule (for conformer generation) $\mathcal{G} = (z, E)$, where z denotes atom types and $e_{ij} \in \mathbb{Z}$ denotes the edge type between node i and j . Generative model f_G , predictive model f_P , equilibrium ground-state geometry C_{eq} .
Sec. 6	Alpha-carbon C_α , coordinate matrix C , protein backbone structure $\mathcal{P}_{base} = (z, C^{C_\alpha})$ or $\mathcal{P}_{bb} = (z, C^{C_\alpha}, C^N, C^C)$, where $a_i \in \{k 1 \leq k \leq 20, k \in \mathbb{Z}\}$ denotes the type of the i -th amino acid and $C^{C_\alpha}, C^N, C^C \in \mathbb{R}^{3 \times n}$ are backbone atom coordinates.
Sec. 7	Material $\mathcal{M} = (z, C, L)$ with lattice matrix $L = [\mathbf{t}_1, \mathbf{t}_2, \mathbf{t}_3] \in \mathbb{R}^{3 \times 3}$, property prediction function $f : M \mapsto y$, material distribution p , periodic transformation $C' = C + LK$.
Sec. 8	Molecule $\mathcal{M} = (A, E, C)$, where A refers to the atomic properties, E denotes edge features, C denotes coordinates, and protein $\mathcal{P} = (B, S)$, where B refers to node types, S denotes coordinates, and binding pose prediction function $f_{pose} : (B, S, A, E) \mapsto [C_1, \dots, C_k]$, binding strength prediction function $f_{strength} : (A, E, C, B, S) \mapsto q$. Molecule-material pair $\mathcal{S} = (z, C)$ as an integrated system. Energy prediction function $f_E : \mathcal{S} \mapsto e$, force prediction function $f_F : \mathcal{S} \mapsto F$, relaxed energy prediction function $f_{RE} : \mathcal{S}_{init} \mapsto e_{rel}$, relaxed structure prediction function $f_{RS} : \mathcal{S}_{init} \mapsto C_{rel}$.
Sec. 9	Function $u : U \rightarrow \mathbb{R}^m$ of space and time to be solved, partial derivative with respect to space ∂_x and time ∂_t , differential operators \mathcal{B} and \mathcal{D} , spatial domain \mathbb{X} and its boundary $\partial\mathbb{X}$, temporal domain \mathbb{T} . Group action of group G on function f is denoted by $L_g f(x) := f(g^{-1}x)$.

2 SYMMETRIES, EQUIVARIANCE, AND THEORY

In many scientific problems, the objects of interest normally reside in 3D physical space. Any mathematical representation of these objects invariably relies on a reference coordinate frame, making representations coordinate-dependent. However, nature does not have a coordinate system, and so coordinate-independent representations are desired. Thus, one of the key challenges of AI for science is how to achieve invariance or equivariance. In this section, we provide a detailed review of the mathematical and physical foundations for achieving equivariance. To make the content friendly to readers, we organize this section by a progressive increase in complication, with the logic flow shown in Section 2.1. First, in Section 2.2, Section 2.3, and Section 2.4, we provide motivating examples for equivariance to discrete and continuous symmetry transformations, and describe how the tensor product is used in practice. After that, in Section 2.5, through concrete and intuitive examples, we try to elucidate the physical and mathematical foundations for the underlying theory, such as symmetry groups, irreducible representations, tensor products, spherical harmonics, etc. Then in Section 2.6 and Section 2.7, we further lay out the detailed and formal theory, which can be skipped for certain readers. We provide a more general form of equivariant networks in Section 2.8. Finally, we point out several open research directions that are worth exploring in the field in Section 2.9.

2.1 Overview

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Describing physical data necessitates making choices, such as establishing a reference frame. While these choices facilitate the numerical representation of physical phenomena within data, the resulting data now mirrors *both* the phenomenon under investigation *as well as such choices*. As choices for description, like the frame of reference, are essentially arbitrary, the represented phenomena should not be influenced by these selections. This concept is referred to as *symmetry*. Symmetries refer to aspects of physical phenomena that remain unchanged, or invariant, under transformations such as the change of reference frame. Understanding how to treat symmetries in data is therefore essential to artificial intelligence in science if we aspire to gain insight into the intrinsic, objective properties of the physical world, independent of our observational or representational biases.

If certain symmetries are present in the system, the predicted targets are naturally invariant or equivariant to the corresponding symmetry transformations. For instance, when predicting energies of 3D molecular structures, the predicted target remains unchanged even if the input 3D molecule is translated or rotated in 3D space. One possible strategy to achieve symmetry-aware learning is adopting data augmentation when training supervised learning models. Specifically, random symmetry transformations are applied to input data samples and labels to force the model to output *approximately* equivariant predictions. However, there are several drawbacks with data augmentation. First, to account for the additional degree of freedom from choosing a reference frame, more model capacity would be needed to represent patterns that would be relatively simple in a fixed reference frame. Second, many symmetry transformations, such as translation, can produce an infinite number of equivalent data samples, making it difficult for finite data augmentation operations to completely reflect the symmetries in data. Third, in some scenarios, we need to build a very deep model by stacking multiple layers to achieve good prediction performance. However, it would pose much more challenges to force the deep model to output approximately equivariant predictions by data augmentation if the model does not maintain equivariance at every layer. Last but not least, in some scientific problems such as molecular modeling, it is important to provide

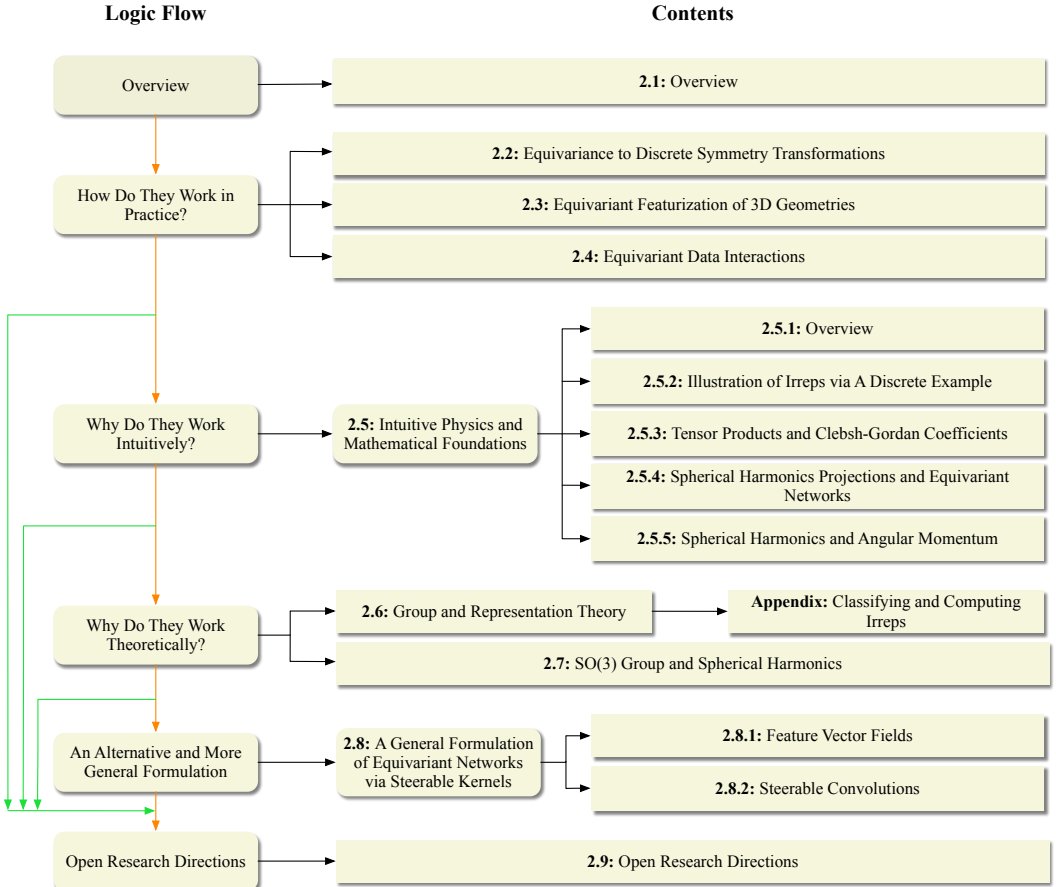


Fig. 4. The overall logic flow and the associated subsections for Section 2. Note the arrows in the logic flow show the dependencies among different subsections, and especially, the green skip connections indicate that some subsections can be skipped. The black arrows show the associations between the logic flow and the subsections, as well as the relationships between each subsection and its child sections. Note the purpose of this figure is for readers to quickly navigate to certain contents based on background and interest, e.g., certain readers may skip the subsections associated with “Why Do They Work Theoretically?”.

provably robust predictions under these transformations so that users can employ machine learning models in a reliable way.

Given the drawbacks of using data augmentation, an increasing number of studies focus on developing symmetry-adapted machine learning models that are designed to meet the underlying symmetry constraints. With symmetry-adapted architecture, no data augmentation is required for symmetry-aware learning, and models can focus solely on learning the target prediction task. Recently, such symmetry-adapted models have shown significant success in scientific problems for a variety of different systems, including molecules (see Section 5), proteins (see Section 6), and crystalline materials (see Section 7). In the following sections, we will elaborate on the symmetry transformations considered in the scientific problems discussed in this paper, and the equivariant operations in designing symmetry-adapted models for these symmetry transformations.

2.2 Equivariance to Discrete Symmetry Transformations

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In certain scientific problems, the prediction targets are internally equivariant to a finite set of discrete symmetry transformations. To be concrete and simple, we consider the case where the inputs are 2D scalar fields, and the symmetry transformations consist of rotating by the angles of 90°, 180° and 270° [Cohen and Welling 2016]. An example of these problems is simulating the dynamics of the fluid field (e.g., scalar vorticity or density) in a 2D square plane where we learn a mapping between the fluid field at the current time step to the fluid field at the next time step. The simulated fluid fields should rotate accordingly if the input 2D fluid field rotates by 90°, 180° or 270° in certain scenarios (see Section 9 for details). Formally, let $X \in \mathbb{R}^{s \times s}$ be the input signals defined on a $s \times s$ grid, and the function $f : \mathbb{R}^{s \times s} \rightarrow \mathbb{R}^{s \times s}$ maps X to the predicted field. We define the rotation by the angle of α as $R_\alpha : \mathbb{R}^{s \times s} \rightarrow \mathbb{R}^{s \times s}$. The set of all discrete symmetry transformations is $\{R_\alpha\}_{\alpha \in \mathcal{A}}$, where $\mathcal{A} = \{0^\circ, 90^\circ, 180^\circ, 270^\circ\}$. Specifically, R_{0° is the identity mapping. R_{90° rotates the input matrix by 90°, i.e., $A' = R_{90^\circ}(A)$ satisfies $A'_{i,j} = A_{j,n-i}$ for any $A \in \mathbb{R}^{n \times n}$ and $0 \leq i, j \leq n-1$ (zero-based index). R_{180° and R_{270° are compositions of two and three 90° rotations, respectively. In other words, $R_{180^\circ} = R_{90^\circ} \circ R_{90^\circ}$ and $R_{270^\circ} = R_{90^\circ} \circ R_{90^\circ} \circ R_{90^\circ}$. The equivariance to discrete symmetry transformations requires f to satisfy

$$f(R_\beta(X)) = R_\beta(f(X)), \quad \forall \beta \in \mathcal{A}. \quad (1)$$

To motivate the idea of achieving equivariance to discrete symmetry transformations in $\{R_\alpha\}_{\alpha \in \mathcal{A}}$, we first consider a minimal example of an equivariant *group convolutional neural networks* (G-CNNs) [Cohen and Welling 2016]. Our example consists of a so-called lifting convolution [Bekkers et al. 2018] which performs convolutions with kernels rotated by every angle in \mathcal{A} and then it applies a pooling operation over the newly introduced rotation axis. First, let us reconsider standard convolution. Given the input feature map $X \in \mathbb{R}^{s \times s}$ and a learnable convolution kernel $W \in \mathbb{R}^{k \times k}$, the standard convolution $X * W$ computes a $s \times s$ feature map, where the feature value at the i -th row, j -th column is computed as

$$(X * W)_{ij} = \sum_{p=0}^{k-1} \sum_{q=0}^{k-1} W_{pq} X_{i+p, j+q}, \quad 0 \leq i, j \leq s-k. \quad (2)$$

Here we omit paddings for simplicity (the actual output size in Equation (2) is $(s-k+1) \times (s-k+1)$).

Now consider the group equivariant lifting convolution, it consists of four standard convolutions with kernels rotated by angle α . This creates the stack $\{F_\alpha\}_{\alpha \in \mathcal{A}}$ of feature maps $F_\alpha = X * R_\alpha(W)$ in which the new α axis indexes the filter response for each rotation α . The output can thus be considered as a field of "rotation response vectors", which is a particular instance of a feature field with fibers that transform via the regular representation of the rotation group [Cesa et al. 2022b]. A discussion of feature fields is beyond the scope of this section, but will be picked up in Section 2.8. The main point here is that the output is not the standard scalar field which we would like when modeling e.g. scalar vorticity or density. As such, our simple network follows the lifting convolution with a max pooling over α -axis, i.e., we pool over the rotation responses. The simple architecture is then described as

$$\text{GCNN}(X; W) = \text{Pool}(\{F_\alpha\}_{\alpha \in \mathcal{A}}) = \text{Pool}(\{X * R_\alpha(W)\}_{\alpha \in \mathcal{A}}), \quad (3)$$

noting that $\text{Pool}(\cdot)$ pools over the rotation axis.

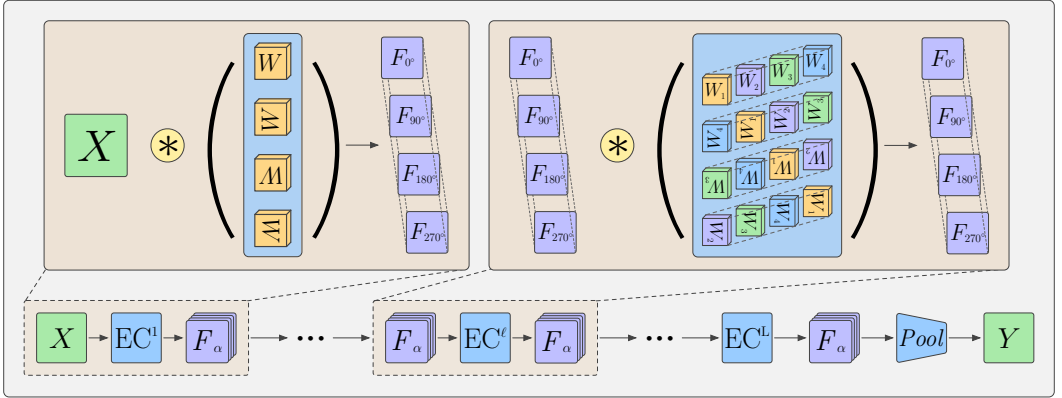


Fig. 5. Illustration of equivariant convolutional neural networks as in G-CNNs [Cohen and Welling 2016]. The network passes input through an equivariant convolution layer (EC), resulting in a set of four output feature maps. If the input X is rotated by 90° , 180° , or 270° , these feature maps will be rotated by β and their ordering permuted. Following this, L group convolution layers are applied, and a final pooling layer is added to account for the feature map permutations. This network is equivariant to rotations by 90° , 180° , or 270° .

The simultaneous use of four convolution operations with rotated kernels in combination with the pooling ensures that the overall G-CNN is equivariant, meaning

$$\text{GCNN}(R_\beta(X); W) = R_\beta(\text{GCNN}(X; W)), \quad \forall \beta \in \mathcal{A}. \quad (4)$$

First, as shown in Equation (3), the four convolution operations rotate the kernel W by 0° , 90° , 180° and 270° , separately, and produce four feature maps F_{0° , F_{90° , F_{180° , F_{270° by performing convolution operations on X with these four kernels. From the calculation process of convolution in Equation (2), we can show that if the input X is rotated by any $\beta \in \mathcal{A}$, the four output feature maps F_{0° , F_{90° , F_{180° , F_{270° will be rotated by the same angle β and change their permutation order, i.e.,

$$\{R_\beta(X) * R_\alpha(W)\}_{\alpha \in \mathcal{A}} = \{R_\beta(X * R_\beta^{-1}(R_\alpha(W)))\}_{\alpha \in \mathcal{A}} \quad (5)$$

$$= \{R_\beta(X * (R_{\alpha-\beta}(W)))\}_{\alpha \in \mathcal{A}} \quad (6)$$

$$= \{R_\beta(F_{(\alpha-\beta) \bmod 360^\circ})\}_{\alpha \in \mathcal{A}}. \quad (7)$$

Second, the pooling operation $\text{Pool}(\cdot)$ over the rotation axis is invariant to permutations within this axis and it preserves rotation equivariance over the spatial axes. We thus have

$$\text{Pool}(\{R_\beta(F_\alpha)\}_{\alpha \in \mathcal{A}}) = R_\beta(\text{Pool}(\{F_\alpha\}_{\alpha \in \mathcal{A}})), \quad \forall \beta \in \mathcal{A}. \quad (8)$$

When Equations (7) and (8) hold, equivariance property in Equation (4) will always be true.

The above simple G-CNN creates locally rotation invariant feature fields, and can be used to build deep equivariant networks with [Andrzejczyk et al. 2019]. However, it's intermediate features would not carry any directional information because of the rotation-axis pooling. Instead, full group equivariant convolutional networks (G-CNNs) [Cohen and Welling 2016] typically start with a lifting convolution, which, as explained above, adds an extra rotation axis to the feature maps (hence often named lifting convolution), followed by group convolution layers that *maintain the extra rotation axis* in the feature maps in order to be able to detect advanced patterns of features in terms of their relative positions and orientations, in which sense the kernels represent part-whole hierarchies [Bekkers 2020]. The typical architecture then starts with a lifting convolution, followed by multiple equivariant group convolution layers before ending with a pooling layer over the α -axis

(see Figure 5 for model illustrations). In each of these intermediate layers, four convolution kernels W_1, W_2, W_3, W_4 are used jointly to map the four input feature maps $F_{0^\circ}^{\text{in}}, F_{90^\circ}^{\text{in}}, F_{180^\circ}^{\text{in}}, F_{270^\circ}^{\text{in}}$ to the four output feature maps $F_{0^\circ}^{\text{out}}, F_{90^\circ}^{\text{out}}, F_{180^\circ}^{\text{out}}, F_{270^\circ}^{\text{out}}$ as

$$F_\alpha^{\text{out}} = \sum_{i=1}^4 F_{(\alpha+i*90^\circ) \bmod 360^\circ}^{\text{in}} * R_\alpha(W_i), \quad \alpha \in \mathcal{A}. \quad (9)$$

It can be shown that if the model uses the lifting convolution in the first layer, and full group convolutions as in Equation (9), the output feature maps at each layer are always equivariant to rotations. Additionally, due to the use of pooling over the rotation axis at the output end of the model, the prediction output of the model is ensured to have the equivariance property in Equation (1). It can further be shown that a linear operator is equivariant *if and only if* it is a group convolution [Bekkers 2020, Thm. 1]. It shows the importance of group convolutions as the essential building blocks for building equivariant G-CNNs; as such, in the work [Cohen et al. 2019, Thm. 3.1] the theorem is stated as (group) convolution *is all you need!*

2.3 Equivariant Featurization of 3D Geometries

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In other scientific problems, the symmetry transformations to be considered are not discrete but *continuous*. Particularly, for many science problems discussed in this paper, we focus on continuous $SE(3)$ transformations in 3D structures of chemical compounds, including translations and 3D rotations, where $SE(3)$ stands for the special Euclidean group in 3D space. In these problems, we aim to predict certain target properties from chemical compounds. A 3D point cloud is used to represent a chemical compound, where every basic unit of the chemical compound (e.g., every atom in the molecule) corresponds to a point in the 3D point cloud, and each point is associated with a 3D Cartesian coordinate. The target properties are usually constrained to be equivariant to $SE(3)$ transformations, *i.e.*, rotations and translations. Note that different from the discrete rotations discussed in Section 2.2, rotations in $SE(3)$ transformations are continuous, meaning that the 3D point cloud can rotate by any angle in 3D space. Formally, let $C = [c_1, \dots, c_n] \in \mathbb{R}^{3 \times n}$ be the coordinate matrix of 3D point cloud with n nodes where c_i is the coordinate of the i -th point, $f : \mathbb{R}^{3 \times n} \rightarrow \mathbb{R}^{2\ell+1}$ be a function mapping coordinate matrices to $(2\ell + 1)$ -dimensional property vector that is $SE(3)$ equivariant with order ℓ . The reason of involving an odd dimensionality of $2\ell + 1$ in f is related to irreducible representations and will be detailed in Section 2.5. Here, order- ℓ equivariance requires f to satisfy

$$f(RC + t\mathbf{1}^T) = D^\ell(R)f(C), \quad (10)$$

where $t \in \mathbb{R}^3$ is the translation vector and $\mathbf{1} \in \mathbb{R}^n$ is a vector whose elements are all equal to one, which broadcasts the vector t to all n input coordinates so that $t\mathbf{1}^T \in \mathbb{R}^{3 \times n}$. $R \in \mathbb{R}^{3 \times 3}$ is the rotation matrix satisfying $R^T R = I$ and $|R| = 1$. $D^\ell(R) \in \mathbb{R}^{(2\ell+1) \times (2\ell+1)}$ is the (real) Wigner-D matrix of R . Here we assume f to be translation-invariant since most physics properties of a system only depend on the relative positions of its components instead of their absolute positions. For example, the energy of a molecule can be completely determined from its interatomic distances. Wigner-D matrices are high-order rotation matrices for 3D rotation transformation in physics. When $\ell = 0$, $D^\ell(R) = [1]$, and f corresponds to the properties that are invariant to $SE(3)$ transformations, such as total energy, Hamiltonian eigenvalues, band gap, *etc.* When $\ell = 1$, $D^\ell(R) = R$, and f corresponds to the properties that will rotate accordingly in 3D space if C is rotated, such as force fields. When $\ell > 1$, $\ell \in \mathbb{N}_+$, $D^\ell(R) \in \mathbb{R}^{(2\ell+1) \times (2\ell+1)}$, f corresponds to properties to be rotated in space with a

higher dimension beyond 3D space if C is rotated, such as spherical harmonics functions with degree $\ell > 1$ and Hamiltonian matrix blocks.

To develop machine learning models for predicting such $SE(3)$ -equivariant properties, we need advanced methods to encode geometric information in C into $SE(3)$ -equivariant features. A commonly used $SE(3)$ -equivariant geometric feature encoding in physics and many existing machine learning methods is the spherical harmonics function. Generally, (real) spherical harmonics function $Y^\ell(\cdot) : \mathbb{R}^3 \rightarrow \mathbb{R}^{2\ell+1}$ maps an input 3D vector to a $(2\ell + 1)$ -dimensional vector representing the coefficients of order- ℓ spherical harmonics bases (see Section 2.5 for an introduction about the physical meaning of spherical harmonics bases). A nice property of the spherical harmonics function is that it is equivariant to order- ℓ rotations, or so-called order- ℓ $SO(3)$ transformations:

$$Y^\ell(R\mathbf{c}) = D^\ell(R)Y^\ell(\mathbf{c}), \quad (11)$$

where $D^\ell(R)$ is the same Wigner-D matrix as in Equation (10). Given the coordinates $\mathbf{c}_i, \mathbf{c}_j$ of two points i, j in a 3D point cloud, spherical harmonics function can be used to encode their relative position $\mathbf{c}_i - \mathbf{c}_j$ to an order- ℓ $SE(3)$ -equivariant feature vector.

2.4 Equivariant Data Interactions

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Recently, many $SE(3)$ -equivariant operations based on spherical harmonics function have been proposed and applied to machine learning models, where spherical harmonics are used to *featurize* 3D geometries into higher dimensions such that they can directly interact with high dimensional features that reside on the geometries (e.g., node features in a graph). In this section, we review methods of data interactions and operations that preserve equivariance.

2.4.1 Equivariant Data Interactions via Tensor Product.

There are many different ways to featurize local geometry via spherical harmonic related operations. One widely used operation is message passing [Gilmer et al. 2017] based on tensor product (TP) operations [Thomas et al. 2018]. For an n -node point cloud with coordinates $C = [\mathbf{c}_1, \dots, \mathbf{c}_n]$, we assume that each node i is associated with an order- ℓ_1 $SE(3)$ -equivariant node features $\mathbf{h}_i^{\ell_1} \in \mathbb{R}^{2\ell_1+1}$. The TP based message passing first computes a message $\mathbf{m}_i^{\ell_3} \in \mathbb{R}^{2\ell_3+1}$, then update $\mathbf{h}_i^{\ell_1}$ to new node feature $\mathbf{h}'_i^{\ell_1}$. This process can be formally described as

$$\begin{aligned} \mathbf{m}_i^{\ell_3} &= \sum_{j \in \mathcal{N}(i)} \mathbf{m}_{j \rightarrow i}^{\ell_3} = \sum_{j \in \mathcal{N}(i)} \text{TP}_{\ell_1, \ell_2}^{\ell_3} (\mathbf{c}_i - \mathbf{c}_j, \mathbf{h}_j^{\ell_1}), \\ \mathbf{h}'_i^{\ell_1} &= \mathbf{U}(\mathbf{h}_i^{\ell_1}, \mathbf{m}_i^{\ell_3}), \end{aligned} \quad (12)$$

where $\text{TP}_{\ell_1, \ell_2}^{\ell_3}(\cdot, \cdot)$ is the TP operation, $\mathcal{N}(i)$ is the neighboring node set of the node i , $\mathbf{U}(\cdot, \cdot)$ is the node feature updating function. $\mathcal{N}(i)$ is commonly defined as the set of nodes whose distances to i are smaller than a radius cutoff r , i.e., $\mathcal{N}(i) = \{j : \|\mathbf{c}_i - \mathbf{c}_j\|_2 \leq r\}$. The TP operation in Equation (12) uses order- ℓ_2 spherical harmonics function as the kernel to compute the message $\mathbf{m}_{j \rightarrow i}^{\ell_3}$ propagated from every node j in $\mathcal{N}(i)$ to the node i . The detailed calculation process can be described as

$$\text{TP}_{\ell_1, \ell_2}^{\ell_3} (\mathbf{c}_i - \mathbf{c}_j, \mathbf{h}_j^{\ell_1}) = C_{\ell_1, \ell_2}^{\ell_3} \text{vec} \left(F(d_{ij}) Y^{\ell_2}(\mathbf{r}_{ij}) \otimes \mathbf{h}_j^{\ell_1} \right). \quad (13)$$

Here, $F(d_{ij})$ is a multi-layer perceptron (MLP) model that takes the distance $d_{ij} = \|\mathbf{c}_i - \mathbf{c}_j\|_2$ as input, $\mathbf{r}_{ij} = \frac{\mathbf{c}_i - \mathbf{c}_j}{d_{ij}}$, \otimes is the vector outer product operation, i.e., $\mathbf{a} \otimes \mathbf{b} = \mathbf{a}\mathbf{b}^T$, $\text{vec}(\cdot)$ is the operation that flattens a matrix to a vector, and $C_{\ell_1, \ell_2}^{\ell_3}$ is Clebsch-Gordan (CG) matrix with $2\ell_3 + 1$ rows and

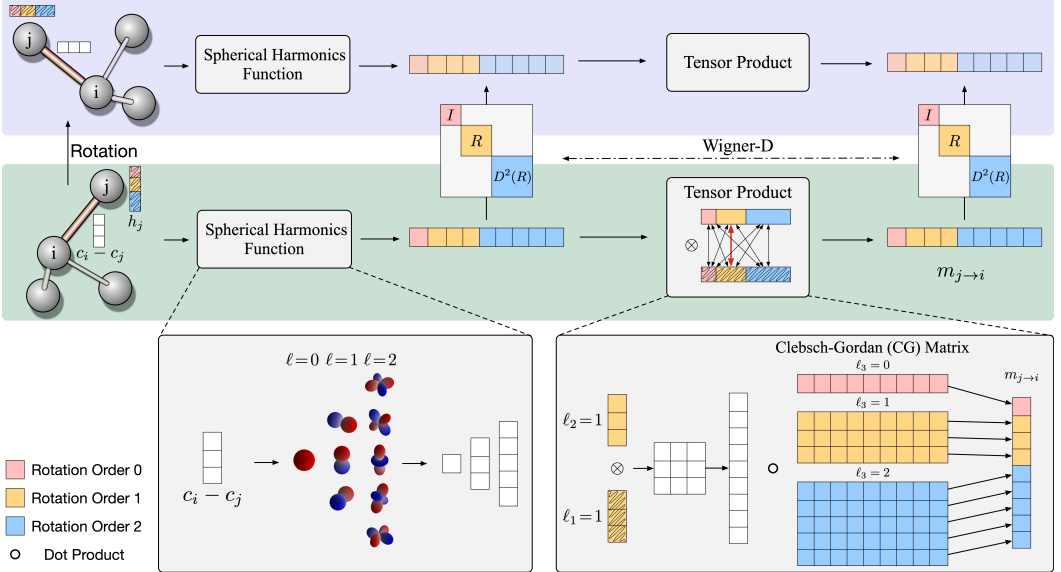


Fig. 6. Illustrations of tensor product operations [Thomas et al. 2018]. Here we show how to compute a message $\mathbf{m}_{j \rightarrow i}$ from node j to node i , assuming the rotation orders are up to 2. Given the coordinates \mathbf{c}_i and \mathbf{c}_j of node i and node j in a 3D point cloud, their relational position $\mathbf{c}_i - \mathbf{c}_j$ is first encoded into an $SE(3)$ equivariant feature vector using spherical harmonics functions. A tensor product is then performed between the computed feature vector and $SE(3)$ -equivariant node features \mathbf{h}_j of node j to compute the message $\mathbf{m}_{j \rightarrow i}$. The resulting message $\mathbf{m}_{j \rightarrow i}$ is $SE(3)$ -equivariant, rotating with the 3D point cloud via the corresponding Wigner-D matrix.

$(2\ell_1 + 1)(2\ell_2 + 1)$ columns. Particularly, CG matrix is widely used in physics to ensure that for $|\ell_1 - \ell_2| \leq \ell_3 \leq \ell_1 + \ell_2$, the $TP_{\ell_1, \ell_2}^{\ell_3}(\cdot, \cdot)$ operation is always $SE(3)$ -equivariant as

$$TP_{\ell_1, \ell_2}^{\ell_3} \left(R\mathbf{c}_i - R\mathbf{c}_j, D^{\ell_1}(R)\mathbf{h}_i^{\ell_1} \right) = D^{\ell_3}(R)TP_{\ell_1, \ell_2}^{\ell_3} \left(\mathbf{c}_i - \mathbf{c}_j, \mathbf{h}_i^{\ell_1} \right). \quad (14)$$

Hence, the message $\mathbf{m}_i^{\ell_3}$ is naturally $SE(3)$ -equivariant. We refer to e.g. Appendix A.5 of Brandstetter et al. [2022a] for derivation and Section 2.5.3 for an intuitive example. Also, for the node feature update function $U(\cdot, \cdot)$ in Equation (12), a linear operation or another TP operation can be used to maintain $SE(3)$ -equivariance of the new node feature $\mathbf{h}'_i^{\ell_1}$. Since all calculations of TP-based message passing are $SE(3)$ -equivariant, we can develop a powerful $SE(3)$ -equivariant model by stacking multiple such message passing layers. Note that in the discussed message passing operation here, both the input node feature and output message have a single rotation order. In practice, a complete node feature \mathbf{h}_i is composed of $SE(3)$ -equivariant features with multiple rotation orders. Multiple messages with different rotation orders are computed by TP operations and concatenated to the message $\mathbf{m}_{j \rightarrow i}$ from the node j to i and the aggregated message \mathbf{m}_i . Then, \mathbf{m}_i is used to update \mathbf{h}_i to new node feature \mathbf{h}'_i . We illustrate the tensor product operations of calculating $\mathbf{m}_{j \rightarrow i}$ with rotation orders up to 2 in Figure 6.

2.4.2 Approximately Equivariant Data Interactions via Spherical Channel Networks.

In addition to linear or TP operations, the node feature \mathbf{h}_i can also be updated on the spherical surface in a nonlinear way by spherical channel networks (SCNs) [Zitnick et al. 2022; Passaro and

Zitnick 2023] to achieve equivariance. An SCN considers all feature values in \mathbf{h}_i as coefficients of spherical harmonics bases, and \mathbf{h}_i represents a spherical function that maps a unit vector on the spherical surface to a real value. This spherical function can be described as a linear combination of spherical harmonics bases $f(\theta, \varphi) = \sum_{m,\ell} h_{i,m}^{\ell} Y_m^{\ell}(\theta, \varphi)$, where ℓ traverses the $SE(3)$ -equivariant feature vectors with different rotation orders in \mathbf{h}_i , and $-\ell \leq m \leq \ell$ traverses the elements in an order- ℓ $SE(3)$ -equivariant feature vector. Here, Y^{ℓ} is the same spherical harmonics function defined in Section 2.3, but its input vector is defined by the polar angle θ and the azimuthal angle φ in the spherical coordinate system. With $f(\theta, \varphi)$, an operation $G(\mathbf{h}_i)$ samples multiple (θ, φ) pairs on the spherical surface and produces a feature map from their corresponding function values $f(\theta, \varphi)$. This feature map can be used as a representation of spherical functions. In SCNs, a similar feature map is constructed from the message \mathbf{m}_i by G , and the node feature \mathbf{h}_i is updated to \mathbf{h}'_i by the point-wise convolution F_c on $G(\mathbf{m}_i)$, $G(\mathbf{h}_i)$ and the inverse operation of G as

$$\mathbf{h}'_i = \mathbf{h}_i + G^{-1}(F_c(G(\mathbf{m}_i), G(\mathbf{h}_i))). \quad (15)$$

Here, the inverse operation G^{-1} transfers feature map values to coefficients of spherical harmonics bases by performing a dot-product between feature values and spherical harmonics bases.

Following SCN, the equivariant spherical channel network (eSCN) [Passaro and Zitnick 2023] proposes a novel equivariant convolution that efficiently reduces the complexity of tensor products. For each edge \mathbf{r} , a specific rotation matrix R is applied to rotate the primary axis, thereby aligning \mathbf{y} axis with the direction of the edge shown as $R \cdot \mathbf{r} = (0, 1, 0)$. As a result, the spherical harmonic bases, denoted as $Y_m^{\ell}(R \cdot \mathbf{r})$, are equal to 1 when $m = 0$, and 0 otherwise. Thus, a significant computational cost reduction can be obtained since the calculation for $m \neq 0$ can be omitted in tensor product. Subsequently, an inverse of Wigner-D matrix is applied to the message to transform it back to original coordinate system, maintaining the equivariance. To further improve efficiency of tensor product, eSCN only considers non-zero entries in the large but sparse Clebsch-Gordan matrix by implementing an $SO(2)$ convolution comprised of two linear layers. Then, a point-wise non-linearity on the spherical surface is performed to obtain the message for each edge. Lastly, the eSCN adopts the same message aggregation as the SCN to update the node feature \mathbf{h}_i .

Note that in both SCN and eSCN, the aggregation operation is not strictly but *approximately* equivariant. Equivariance can only be maintained if the input node features are rotated by the angles that are exactly sampled in constructing the spherical grid. However, due to the continuous nature of the rotation, achieving this ideal condition is not always feasible.

2.5 Intuitive Physics and Mathematical Foundations

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In the above Section 2.2, Section 2.3, and Section 2.4, we provide applications of equivariance to discrete and continuous symmetry transformations in recent research, and describe how the tensor product is used in practice. In this section, we expect that, through some simple and intuitive examples, readers would understand the underlying theory in a reasonably short time. Specifically, in Section 2.5.1, we provide a sketch of group and representation theory, *i.e.*, the introduction of irreducible representations (irreps), and how equivariant neural network produce irreps through tensor product; we try to explain the intuitions of symmetry groups and irreps through a simple and discrete case, a square with four nodes, in Section 2.5.2; in Section 2.5.3, we provide an effortless example for readers to understand tensor products and Clebsch-Gordan coefficients; in Section 2.5.4, we introduce spherical harmonics projection, a concrete application of spherical harmonics; we further manifest the idea of spherical harmonics functions from the angular momentum perspective in Section 2.5.5.

2.5.1 Overview.

In this section, we give concrete examples to elucidate the fundamentals of group representation theory. Consider the vector space of polynomials Z , spanned by $(x^2, xy, xz, y^2, yz, z^2)$ from the direct product $(x, y, z) \times (x, y, z)$. When the original space (x, y, z) is transformed by a 3×3 rotation matrix, the vector space Z will be transformed by a 6×6 matrix. If we look at random $SO(3)$ rotations on this vector space, the 6×6 rotation matrices are dense; they do not look like they have independent vector spaces. However, if we perform a change of basis to $(x^2+y^2+z^2, xy, yz, 2z^2-x^2-y^2, xz, x^2-y^2)$, then the rotation matrices take on a striking pattern. Factually, the original space can be decomposed into two independent subspaces $L_0 = (x^2 + y^2 + z^2)$ which is invariant (the group representations for all elements take the form of $I = [1]$) and $L_2 = (xy, yz, 2z^2 - x^2 - y^2, xz, x^2 - y^2)$. This actually describes how to decompose a reducible representation into irreducible representations (irreps). To further elucidate this, we give an example in Section 2.5.2 for a discrete case, which might be easier for starters to understand.

This transformation is significant, as it means any vector space can be described as a concatenation of these fundamental vector spaces. In principle, it requires that, when conducting “representation” learning with machine learning, if the vector space we learn changes predictably under group action, *e.g.*, rotations, then our “learned” vector space must be comprised of irreps, no matter how complex it may be. In the equivariant neural network literature, the term tensor product is used to define a tensor product plus decomposition operation, *i.e.*, direct product two representations (reducible or irreducible) to produce (generally) a reducible representation and then decompose the reducible representation into irreps. A more detailed description of tensor product and such decomposition is provided in Section 2.5.3, where we show in a more general setting for the tensor product of two different 3D vectors.

Additionally, the abstract structure from polynomials can be directly extended to geometrical concepts. In fact, the vector space L_2 may look familiar to some readers as in fact, this is the vector space spanned by the angular frequency $\ell = 2$ real spherical harmonics (modulo normalization factors), which form a vector space of functions that transform as the irreps of $SO(3)$. Similarly, the L_0 vector space is proportional to the $\ell = 0$ spherical harmonic, which is a constant for all points on the sphere, $s \in \mathbb{S}^2$. We will introduce an easy-to-understand application to manifest spherical harmonics in Section 2.5.4, and also provide a detailed description in Section 2.7. Just briefly and intuitively, spherical harmonics form an orthogonal basis for functions on the sphere. This means that any function in 3D space with a unique origin can be separated into radial and angular degrees of freedom because these degrees of freedom are orthogonal under 3D rotation. In fact, spherical harmonics are the basis functions for performing a Fourier transform on the sphere, which must have integer frequencies due to periodic boundary conditions (analogous to Fourier transforms over periodic spatial domains). As a result, spherical harmonics have a wide range of uses, from lighting in computer graphics, signal processing of sound waves, and description of physical systems, *e.g.*, analyzing the cosmic microwave background and describing atomic orbitals.

2.5.2 Illustration of Irreducible Representations via A Discrete Example.

In Section 2.5.1, we provide a sketch of group and representation theory, *i.e.*, the introduction of irreps, and how equivariant neural network produce irreps through tensor product. In this section, we explain symmetry groups and irreducible representations through a simple example. We further elucidate the motivation of equivariant neural networks to incorporate these symmetries for effective learning. Note Section 2.5.1 gives a continuous form, which could be more generalizable. However, we believe it’s easier for readers to understand the concepts through discrete group transformations as follows.

Consider a square where each of the four nodes has a scalar feature $a_{1\sim 4}$. The symmetry group of the square, called C_{4v} , contains a 90° rotation, reflections along the vertical and horizontal axes, and reflections along the two diagonals. Under these symmetry transformations, the four scalar features transform into each other, forming a four-dimensional representation of the C_{4v} group. This representation is reducible and can be decomposed into three irreps: $a_1 + a_2 + a_3 + a_4$, $a_1 - a_2 + a_3 - a_4$, and $(a_1 - a_3, a_2 - a_4)$. The first two are one-dimensional irreps, and the third is a two-dimensional irrep. One can check that each irrep is closed under the C_{4v} transformations. The first irrep is invariant under all symmetry transformations. The second irrep changes sign under a 90° rotation and reflections along the vertical and horizontal axes. The third irrep transforms as a 2D vector.

To ensure equivariance, the learning outcome must also be classified into the irreps, which transforms accordingly under the group transformation. Then an equivariant neural network is a function that maps irreps to irreps, which is strongly constrained by the underlying symmetry group. Based on group theory, the C_{4v} group has five distinct irreps, four 1D irreps denoted as A_1 , A_2 , B_1 , B_2 , and one 2D irrep denoted as E . The A_1 irrep corresponds to the invariant irrep, such as $a_1 + a_2 + a_3 + a_4$ mentioned earlier. The A_2 irrep remains unchanged under rotation but changes sign under both reflections. The simplest A_2 is $(a_1 - a_2 + a_3 - a_4)(a_1 - a_3)(a_2 - a_4)$, which has a cubic order in a_i . The B_1 irrep changes sign under 90° rotation and both horizontal and vertical reflections, such as $a_1 - a_2 + a_3 - a_4$. The B_2 irrep changes sign under 90° rotation and diagonal reflections, such as $(a_1 - a_3)(a_2 - a_4)$. The C_{4v} group only has one 2D irrep, denoted as E , which transforms as a 2D vector, for instance, $(a_1 - a_3, a_2 - a_4)$.

The irreps impose strongly restricts the form of equivariant learning outcome f from the four scalar feature $a_{1\sim 4}$. For simplicity, let f be a linear function of the input feature a_i . A learning outcome invariant under symmetry transformations must be proportional to $a_1 + a_2 + a_3 + a_4$. On the other hand, if f is expected to be equivariant as a 2D vector, it must be proportional to $(a_1 - a_3, a_2 - a_4)$ up to a constant rotation.

Classifying quadratic and higher order learning outcomes into different irreps involving the product of irreps. In the case of C_{4v} , the product between any 1D irrep and the 2D irrep becomes a 2D irrep. For example, a 2D quadratic f must be $(a_1 + a_2 + a_3 + a_4)(a_1 - a_3, a_2 - a_4)$ or $(a_1 - a_2 + a_3 - a_4)(a_1 - a_3, a_4 - a_2)$. On the other hand, the product of two 2D irreps decomposes into three 1D irreps: $(a_1 - a_3)^2 + (a_2 - a_4)^2$, $(a_1 - a_3)^2 - (a_2 - a_4)^2$ and $(a_1 - a_3)(a_2 - a_4)$. The first one is invariant and is the A_1 irrep, same as $a_1 + a_2 + a_3 + a_4$. The second one, changing sign under the rotation and horizontal/vertical reflections, is the B_1 irrep, same as $a_1 - a_2 + a_3 - a_4$. The third one, on the other hand, changes sign under diagonal reflections, is the B_2 irrep. If the learning outcome is expected to transform as the A_2 irrep, which remains the same under 90° rotation but changes sign under reflections, it must at least be of the cubic order of the input features, and the simplest form is $(a_1 - a_2 + a_3 - a_4)(a_1 - a_3)(a_2 - a_4)$. Note that up to now, we describe for C_{4v} an ideal case where the product of two irreps may produce an irrep. However, more generally, in practical cases like equivariant neural networks, tensor product takes two irreps and produces a reducible representation, which is further decomposed to irreps as inputs to the next layer, as mentioned in Section 2.5.1. Essentially, this lays the foundation of achieving equivariance in modern equivariant neural networks.

This example illustrates how the group structure imposes significant constraints on the functions that map input data to the desired learning outcomes, based on their irreps. Equivariant neural networks aim to incorporate these constraints into the network architecture explicitly. By doing so, equivariant neural networks can leverage the inherent symmetries and transformations present in the data, leading to more effective and efficient learning.

2.5.3 Tensor Products and Clebsh-Gordan Coefficients.

Mathematically, the tensor product is defined to represent bilinear maps, which generalizes the scalar multiplication to vectors (tensors). Let us consider two 3D vectors $\mathbf{x}, \mathbf{y} \in \mathbb{R}^3$. Let $f: \mathbb{R}^3 \times \mathbb{R}^3 \rightarrow \mathbb{R}$ be a map taking two 3D vectors as inputs, being bilinear means when fixing one input, the restricted map $f(\cdot, \mathbf{y})$ or $f(\mathbf{x}, \cdot)$ is linear w.r.t. the other input. All such bilinear maps can be written as $f(\mathbf{x}, \mathbf{y}) = \sum_{ij} c_{ij} x_i y_j$, where x_i and y_j are elements in \mathbf{x} and \mathbf{y} , and c_{ij} are the coefficients defining different maps. The tensor product between \mathbf{x} and \mathbf{y} is defined as $\mathbf{x} \otimes \mathbf{y} = [x_1 y_1, x_1 y_2, x_1 y_3, x_2 y_1, x_2 y_2, x_2 y_3, x_3 y_1, x_3 y_2, x_3 y_3]^T \in \mathbb{R}^9$. If we define a coefficient vector $\mathbf{c} = [c_{11}, c_{12}, c_{13}, c_{21}, c_{22}, c_{23}, c_{31}, c_{32}, c_{33}]^T \in \mathbb{R}^9$, then any bilinear map can be expressed as $f(\mathbf{x}, \mathbf{y}) = \mathbf{c}^T (\mathbf{x} \otimes \mathbf{y})$. Consequently, f is uniquely represented by its coefficient vector \mathbf{c} . Thus, f lives in a 9-dimensional vector space whose basis can be defined through tensor product. Concretely, the basis can be defined as $\{\mathbf{e}_i \otimes \mathbf{e}_j\}_{i,j \in \{1,2,3\}}$ where \mathbf{e}_i and \mathbf{e}_j are the canonical basis vectors of the original 3D space, e.g., $\mathbf{e}_1 = [1, 0, 0]^T$. Since $\mathbf{e}_i \otimes \mathbf{e}_j$ are vectors with 1 at the $(3i + j)$ -th position and 0 elsewhere, they are orthogonal to each other.

An important property of tensor product is its equivariance. When \mathbf{x} and \mathbf{y} undergo a global rotation defined by a rotation matrix $R \in \mathbb{R}^{3 \times 3}$, each element in the tensor product $R\mathbf{x} \otimes R\mathbf{y}$ is in the form of $\sum_{ij} c_{ij} x_i y_j$, where c_{ij} is computed from elements in R . Thus, the tensor product $\mathbf{x} \otimes \mathbf{y}$ is also transformed by a matrix. Let that matrix be R^\otimes , we have $R^\otimes(\mathbf{x} \otimes \mathbf{y}) = R\mathbf{x} \otimes R\mathbf{y}$. R^\otimes then defines how the rotation transforms in the tensor product space. Note that R^\otimes is a 9×9 matrix and the dimension expands quickly with the dimensions of input spaces. We thus wish to identify smaller building blocks to efficiently describe how $\mathbf{x} \otimes \mathbf{y}$ transforms under rotations. Fortunately, this can be achieved for 3D rotations. For example, we know that when applying a global rotation, the dot product of two vectors is not changed. The dot product is a bilinear map defined as $f_{\text{dot}}(\mathbf{x}, \mathbf{y}) = x_1 y_1 + x_2 y_2 + x_3 y_3$. Expressed with the tensor product basis, the dot product can be defined by the coefficient vector $\mathbf{c}_{\text{dot}} = [1, 0, 0, 1, 0, 0, 0, 1]^T$. The rotation invariance of dot product gives $\mathbf{c}_{\text{dot}}^T (R\mathbf{x} \otimes R\mathbf{y}) = \mathbf{c}_{\text{dot}}^T R^\otimes(\mathbf{x} \otimes \mathbf{y}) = \mathbf{c}_{\text{dot}}^T (\mathbf{x} \otimes \mathbf{y})$. Since this holds for all pairs of \mathbf{x} and \mathbf{y} (e.g., $\mathbf{x} \otimes \mathbf{y}$ can be any basis vector $\mathbf{e}_i \otimes \mathbf{e}_j$), we have $\mathbf{c}_{\text{dot}}^T R^\otimes = \mathbf{c}_{\text{dot}}^T$. Hence, the space spanned by the dot product (i.e., $\lambda \mathbf{c}_{\text{dot}}$, where $\lambda \in \mathbb{R}$) defines a 1-dimensional stable subspace for R^\otimes .

Another stable subspace is the space spanned by the cross product. From the geometric interpretation, we know that the cross product is equivariant to rotation. The cross product can be expressed as a stack of 3 bilinear maps (vector output) as

$$f_{\text{cross}}(\mathbf{x}, \mathbf{y}) = \begin{bmatrix} x_2 y_3 - x_3 y_2 \\ x_3 y_1 - x_1 y_3 \\ x_1 y_2 - x_2 y_1 \end{bmatrix}, \quad (16)$$

which can be expressed as the coefficient matrix

$$C_{\text{cross}} = \begin{bmatrix} 0 & 0 & 0 & 0 & 0 & 1 & 0 & -1 & 0 \\ 0 & 0 & -1 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 1 & 0 & -1 & 0 & 0 & 0 & 0 & 0 \end{bmatrix}^T, \quad (17)$$

which is 9×3 for 3 output dimensions. The equivariance of cross product gives $f_{\text{cross}}(R\mathbf{x}, R\mathbf{y}) = R f_{\text{cross}}(\mathbf{x}, \mathbf{y})$, which writes in the tensor product basis as $C_{\text{cross}}^T (R\mathbf{x} \otimes R\mathbf{y}) = C_{\text{cross}}^T R^\otimes(\mathbf{x} \otimes \mathbf{y}) = R C_{\text{cross}}^T (\mathbf{x} \otimes \mathbf{y})$, which holds for all pairs of \mathbf{x} and \mathbf{y} . Thus we have

$$C_{\text{cross}}^T R^\otimes = R C_{\text{cross}}^T. \quad (18)$$

We can show the space spanned by the cross product defines a 3-dimensional stable subspace for R^\otimes . To show that, let $\mathbf{v} = C_{\text{cross}} \lambda^T$ be a vector in the tensor product basis, defined as a linear combination of the columns in C_{cross} , where $\lambda \in \mathbb{R}^3$ and $\mathbf{v} \in \mathbb{R}^9$. We have $\mathbf{v}^T R^\otimes = \lambda C_{\text{cross}}^T R^\otimes = \lambda R C_{\text{cross}}^T := \mathbf{u}^T$,

where $\mathbf{u} = C_{\text{cross}}^T (R^T \boldsymbol{\lambda}^T) \in \mathbb{R}^9$ is still a linear combination of the columns in C_{cross} . Hence, we have proven that the 3-dimensional space spanned by the columns in C_{cross} is stable to R^\otimes .

To have a complete view of this decomposition, we can wrap the coefficient vector \mathbf{c} for a bilinear map into a matrix as

$$\hat{C} = \begin{bmatrix} c_1 & c_2 & c_3 \\ c_4 & c_5 & c_6 \\ c_7 & c_8 & c_9 \end{bmatrix}.$$

Then the coefficient space spanned by the dot product can be written as $\lambda_1 \hat{C}_{\text{dot}} = \lambda_1 \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}, \forall \lambda_1 \in \mathbb{R}$. The coefficient space spanned by the cross product can be written as

$$\lambda_2 \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix} + \lambda_3 \begin{bmatrix} 0 & 0 & -1 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix} + \lambda_4 \begin{bmatrix} 0 & 1 & 0 \\ -1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, \forall \lambda_2, \lambda_3, \lambda_4 \in \mathbb{R}.$$

When projecting any coefficient \hat{C} onto the space spanned by the dot product, the trace of \hat{C} is extracted. One can verify that the space spanned by the cross product represents the space of all antisymmetric matrices, *i.e.*, $A^T = -A$. The remaining degrees of freedom in the 9-dimensional space of \hat{C} results in the space of all symmetric matrices with trace equal to 0, *i.e.*, $A^T = A, \sum_i A_{ii} = 0$, which is a 5-dimensional space. To summarize, we rewrite any \hat{C} as the summation

$$\hat{C} = \underbrace{\begin{bmatrix} \lambda_1 & 0 & 0 \\ 0 & \lambda_1 & 0 \\ 0 & 0 & \lambda_1 \end{bmatrix}}_{\text{Trace}} + \underbrace{\begin{bmatrix} 0 & \lambda_4 & -\lambda_3 \\ -\lambda_4 & 0 & \lambda_2 \\ \lambda_3 & -\lambda_2 & 0 \end{bmatrix}}_{\text{Antisymmetric}} + \underbrace{\begin{bmatrix} \lambda_5 & \lambda_6 & \lambda_7 \\ \lambda_6 & -\lambda_5 - \lambda_9 & \lambda_8 \\ \lambda_7 & \lambda_8 & \lambda_9 \end{bmatrix}}_{\text{Symmetric traceless}}. \quad (19)$$

To show the symmetric traceless part is indeed 5-dimensional, we can expand the basis and write it as

$$\lambda_5 \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 0 \end{bmatrix} + \lambda_6 \begin{bmatrix} 0 & 1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} + \lambda_7 \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix} + \lambda_8 \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & 0 \end{bmatrix} + \lambda_9 \begin{bmatrix} 0 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 1 \end{bmatrix}, \forall \lambda_{5-9} \in \mathbb{R}.$$

Translating to the tensor product basis, we can derive the function f_{5D} as

$$f_{5D}(\mathbf{x}, \mathbf{y}) = \begin{bmatrix} x_1 y_1 - x_2 y_2 \\ x_1 y_2 + x_2 y_1 \\ x_1 y_3 + x_3 y_1 \\ x_2 y_3 + x_3 y_2 \\ -x_2 y_2 + x_3 y_3 \end{bmatrix}. \quad (20)$$

Importantly, Equation (19) means the 9-dimensional coefficient space can be viewed a direct sum of a 1D, a 3D and a 5D vector spaces and each of them is stable to arbitrary global rotations. The decomposition can be conceptually written as

$$3 \otimes 3 = 1 \oplus 3 \oplus 5.$$

It is worth noting the in general such decomposition depends on the choice of the transformation. Here the transformation is the 3D rotation ($SO(3)$ group). The decomposition would be different if we choose another transformation. For example, for the trivial transformation (group $G = \{e\}$), the decomposition would result in 9 1-dimensional trivial subspaces.

One important property of these subspaces is that they cannot be further decomposed and remain stable to global rotations (*i.e.*, they are irreducible). The 1D subspace spanned by dot product transforms under rotation as scalar and is irreducible by definition. The 3D subspace spanned by cross product transforms as vector and we can prove that it is also irreducible. Concretely, by Equation (18), the 3-dimensional space spanned by C_{cross} is transformed by the same rotation matrix R under rotations. Since any 3D vector (under any basis) can be transformed to be colinear

with any other 3D vector with some 3D rotation, there is no smaller subspace in the cross product space that is stable under arbitrary rotations. For the 5D subspace, an intuitive proof for its irreducibility requires more advanced theories such as the angular momentum in physics, or the character theory in mathematics. Nevertheless, we can gain some intuition about the behaviour of f_{5D} by noticing that one of its component $x_1y_1 - x_2y_2$ changes sign under 90° degree rotation around the z axis, i.e., $(x_1, x_2) \leftarrow (-x_2, x_1)$ and $(y_1, y_2) \leftarrow (-y_2, y_1)$. More generally, f_{5D} corresponds to a representation with $\ell = 2$. Intuitively, an $\ell = 2$ object is something that returns to itself after a 180° rotation.

Generally, for any input dimensions, we can identify all such stable subspaces so that when the inputs undergo a global rotation, the subspaces in tensor product space will not mix with each other. By changing to the direct sum basis of these stable subspaces, one can efficiently express R^\otimes in a block diagonal form. The matrices for performing such a change of basis are the Clebsch-Gordan (CG) coefficients. In summary, tensor products define a basis for all bilinear maps between two vector spaces, which is particularly suitable for studying equivariance when a global transformation is applied, since equivariance essentially describes maps between transformations in an input-independent way.

2.5.4 Spherical Harmonics Projections and Equivariant Networks.

(Real) spherical harmonics $Y_m^\ell : \mathbb{S}^2 \rightarrow \mathbb{R}$ are special functions defined on the surface of a unit sphere \mathbb{S}^2 . They form a set of complete orthogonal bases for functions defined on \mathbb{S}^2 . Thus every function on \mathbb{S}^2 can be expanded as a linear combination of those spherical harmonics. This expansion is reminiscent of Fourier expansion of $v \in V$ based on a set of complete orthogonal bases $\{\mathbf{u}_1, \dots, \mathbf{u}_n\}$ of vector space V as

$$\mathbf{v} = \sum_{i=1}^n \langle \mathbf{u}_i, \mathbf{v} \rangle \mathbf{u}_i. \quad (21)$$

Similarly, a spherical function $f(\cdot) : \mathbb{S}^2 \rightarrow \mathbb{R}$ can be expanded by spherical harmonics such that

$$f(\Omega) = \sum_{\ell,m} a_{\ell,m} Y_m^\ell(\Omega), \quad (22)$$

where $a_{\ell,m} = \langle Y_m^\ell, f \rangle = \int Y_m^\ell(\Omega) f(\Omega) d\Omega$.

We use the Dirac delta function

$$\delta(x) = \begin{cases} \infty, & x = 0 \\ 0, & x \neq 0 \end{cases}, \quad (23)$$

as an example to illustrate the idea of spherical harmonics expansion. Let $\Omega, \Omega' \in \mathbb{S}^2$ and $f = \delta(\Omega - \Omega')$, we then obtain

$$a_{\ell,m} = \langle Y_m^\ell, \delta \rangle = \int Y_m^\ell(\Omega) \delta(\Omega - \Omega') d\Omega = Y_m^\ell(\Omega'). \quad (24)$$

As a result, the spherical harmonics expansion of the Dirac delta function is

$$f = \delta(\Omega - \Omega') = \sum_{\ell,m} a_{\ell,m} Y_m^\ell(\Omega) = \sum_{\ell,m} Y_m^\ell(\Omega') Y_m^\ell(\Omega). \quad (25)$$

The above delta function expansion is the basis of the spherical harmonics projection, which is widely used in local equivariant descriptors and convolution operations in equivariant neural networks. Specifically, to project a geometry vector to spherical harmonics, it contains two parts: a radial basis function to embed the length of the vector; the spherical harmonics expansion of delta function to embed the direction of the vector. Let a set of geometry vectors $\{\mathbf{r}_i \in \mathbb{R}^3\}_{i=1}^n$ and

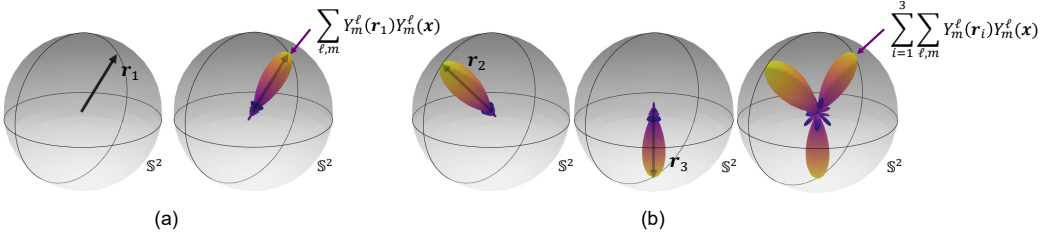


Fig. 7. Illustration of spherical harmonics projection for (a) a single unit vector and (b) multiple unit vectors. **(a)** A unit vector \mathbf{r}_1 and its reconstruction from the spherical harmonics projection are plotted. A unit vector is mathematically modeled as a Dirac delta function on the unit sphere \mathbb{S}^2 , with non-zero value only on the direction it points to. Spherical harmonics functions define a basis set for functions on \mathbb{S}^2 and the delta function is reconstructed as a linear combination of spherical harmonics where the linear coefficients gives the embedding of the function. The spherical harmonics projection contains infinite number of terms and only the terms with $\ell \leq 6$ are used in the plotted reconstruction. To visualize a function on the sphere clearly, the reconstruction is plotted as a 3D blob where the distance to the origin represents the function magnitude on the sphere along that direction. Additionally, the maximum amplitude in the reconstruction is scaled to one for visualization. **(b)** Multiple vectors can be embedded using spherical harmonics projection by summing up individual projections. Here, we show an example with two additional vectors \mathbf{r}_2 and \mathbf{r}_3 , where $\mathbf{r}_1, \mathbf{r}_2, \mathbf{r}_3$ are first individually embedded by spherical harmonics projection and their embeddings are then summed together to form an overall embedding, which is visualized as three blobs in a sphere. This example is adapted from lecture notes by Tess Smidt with permission.

spherical harmonics functional input $\mathbf{x} \in \mathbb{R}^3$, $\|\mathbf{x}\|_2 = 1$. The spherical harmonics projection is given as

$$\sum_{i=1}^n \phi(\|\mathbf{r}_i\|_2) \sum_{\ell,m} Y_m^\ell \left(\frac{\mathbf{r}_i}{\|\mathbf{r}_i\|_2} \right) Y_m^\ell(\mathbf{x}), \quad (26)$$

where $\phi(\cdot) : \mathbb{R} \rightarrow [0, \infty)$ is the radial basis function providing scaling of projection. Since ℓ is often defined within a radius instead of over the whole non-negative integers due to computational efficiency, the summation

$$\sum_{\ell,m} Y_m^\ell \left(\frac{\mathbf{r}_i}{\|\mathbf{r}_i\|_2} \right) Y_m^\ell(\mathbf{x})$$

approximates the Dirac delta function rather than exactly evaluates it. Assume $\ell \leq 6$ and $\|\mathbf{r}_i\|_2 = 1$, the spherical harmonics projection forms a blob around the vector \mathbf{r}_i , as illustrated in Figure 7 (a). When \mathbf{x} is closer to \mathbf{r}_i , the projection value is larger and the distance to the sphere center is longer. Multiple vectors can be embedded by summing up every single projection, resulting in multiple blobs in a sphere, as illustrated in Figure 7 (b).

2.5.5 Spherical Harmonics Functions and Angular Momentum.

The aforementioned spherical harmonics-based feature encoding and TP operation are actually tightly related to physical science, particularly quantum mechanics. In physics, spherical harmonics bases are commonly used in solving partial differential equations. Specifically, for single-electron hydrogenic atoms such as Hydrogen, the eigen wavefunctions of the electron are a set of analytic solutions of the Schrödinger equation, given by the product of the radial part $R_{nl}(r)$ and complex spherical harmonics. More details of spherical harmonics can be found in Section 2.7. The latter can be transformed into real spherical harmonics $Y_m^\ell(\theta, \varphi)$ which are often used in first-principles DFT, quantum chemistry, and recent deep learning models. The set of real spherical harmonics,

denoted by $Y_m^\ell(\theta, \varphi) : \mathbb{S}^2 \rightarrow \mathbb{R}$ where $\ell \in \mathbb{N}$ is the orbital angular momentum quantum number and $m \in \mathbb{Z}, -\ell \leq m \leq \ell$ is the magnetic quantum number, forms a complete orthogonal basis set that can be used to expand any spherical functions. Additionally, in physical systems, the TP operation is usually used in angular momentum coupling. Specifically, when we consider two electrons in the system with Coulomb forces, the angular momentum of the coupled wavefunctions can be deduced from the TP of the separate angular momentum. Besides the use of spherical harmonics for feature representations, they are also demonstrated for quantum tensor learning in Section 4, such as quantum Hamiltonian learning.

2.6 Group and Representation Theory

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Equivariant neural networks are formulated in the language of group and representation theory, the basics of which are briefly introduced in this section. After some elementary definitions in Section 2.6.1, we explain in Section 2.6.2 how groups can act on other objects and define invariant and equivariant functions w.r.t. such actions. In the context of deep learning, symmetry groups act on data and features and the network layers are constrained to be invariant or equivariant. The networks' feature spaces are usually vector spaces. Group actions on vector spaces are described by group representation theory, which is discussed in Section 2.6.3.

2.6.1 Symmetry Groups.

Groups are algebraic objects which formalize symmetry transformations like, e.g., translations, rotations or permutations. To motivate their formal definition, note first that we can always combine any two transformations into a single transformation. This composition of transformations is naturally obeying certain properties which characterize the algebraic structure of groups. Consider, for instance, the case of planar rotations. Each rotation can be identified with a rotation angle, and any two rotations by α and β are composed to a combined rotation by $\alpha + \beta$ (modulo 2π). Note that a rotation by any angle α can be undone by another rotation by the negated angle $-\alpha$. There is furthermore a trivial “identity” transformation, the rotation by $\alpha = 0$ degrees, which does not do anything. Finally, given rotations by three angles α, β and γ , the order of composition of the rotations is irrelevant, that is, $(\alpha + \beta) + \gamma = \alpha + (\beta + \gamma)$. Symmetry groups are defined exactly as sets of transformations whose composition satisfies these three properties.

DEFINITION 1 (GROUP). *Let G be a set and $\bullet : G \times G \rightarrow G$ be a binary operation that takes two elements from G and maps them to another element. If (G, \bullet) satisfy the following three axioms, they form a group that is:*

- (1) *Inverse: for any $g \in G$ there exists an inverse element $g^{-1} \in G$ satisfying $g \bullet g^{-1} = g^{-1} \bullet g = e$;*
- (2) *Identity: there exists an identity element $e \in G$ which satisfies $e \bullet g = g \bullet e = g$ for any $g \in G$;*
- (3) *Associativity: $(g \bullet h) \bullet k = g \bullet (h \bullet k)$ for any $g, h, k \in G$.*

For brevity, one often refers to the set G instead of (G, \bullet) as group and drops the composition in the notation, writing gh for $g \bullet h$. We will in the following make use of these abbreviations whenever the meaning is unambiguous.

The composition of planar rotations obeys actually yet another property: for any two angles α and β , the order of composition is irrelevant, since $\alpha + \beta = \beta + \alpha$. This commutativity of group elements is not included in the definition above since it does not apply to all symmetry groups. For instance, non-planar rotations in 3D do not commute with each other, rotations to not commute with translations or reflections, and permutations do in general not commute. Groups like planar rotations, whose elements do commute, are called *abelian*.

DEFINITION 2 (ABELIAN GROUP). Let G be a group. If all of its elements commute, that is, if $gh = hg$ for any $g, h \in G$, the group is called abelian.

An important class of groups are matrix groups, which are sets of square matrices that are composed via matrix multiplications and satisfy the three group axioms. Associativity holds hereby by the definition of matrix multiplications; the identity element is given by the identity matrix; and the set is required to be closed under matrix inversion. To give an example, consider the set of all invertible $n \times n$ matrices $GL(\mathbb{R}^n) := \{g \in \mathbb{R}^{n \times n} \mid \det(g) \neq 0\}$, which is called *general linear group* and is geometrically interpreted as the group of all possible basis changes of \mathbb{R}^n . As matrix multiplications are in general not commutative, this group is not abelian.

Groups may contain subsets which are themselves forming groups. They are therefore called subgroups.

DEFINITION 3 (SUBGROUP). Let G be a group and $H \subseteq G$ be a subset of transformations. If H is still forming a group, it is called a subgroup of G .

One can show that it is sufficient to check that H is closed under compositions; that is, $gh \in H$ for any $g, h \in H$, and under taking inverses, i.e., $g^{-1} \in H$ for any $g \in H$.

As an example, we consider the matrix subgroup $SO(n) := \{g \in \mathbb{R}^{n \times n} \mid \det(g) = 1\}$ of $GL(\mathbb{R}^n)$. It does not contain all $n \times n$ matrices with non-zero determinant, but only the subset of those with unit determinant. That it is indeed a subgroup of $GL(\mathbb{R}^n)$ is clear since it is closed under composition, $\det(gh) = \det(g)\det(h) = 1$ for $g, h \in SO(n)$, and under inversion, $\det(g^{-1}) = \det(g)^{-1} = 1$. The groups $SO(n)$ are called *special orthogonal groups* since they consist of rotation matrices which transform between orthogonal bases of \mathbb{R}^n . There are larger (non-special) orthogonal subgroups $O(n) := \{g \in \mathbb{R}^{n \times n} \mid \det(g) = \pm 1\}$ of $GL(\mathbb{R}^n)$ which contain not only rotations but also reflections.

2.6.2 Group Actions and Equivariant Maps.

The abstract definition of symmetry groups above captures their algebraic properties under composition, but does not yet allow to describe *transformations of other objects*. One and the same group can, indeed, act on various different objects, for instance, different feature spaces. Consider, for instance, the rotation group $SO(2)$ in two dimensions. It acts naturally on 2-dimensional vectors in \mathbb{R}^2 via matrix multiplication, but 2-dimensional rotations may also act on \mathbb{R}^3 by rotating around different axes, or may even transform images or point clouds by rotating them in space.

Besides having a symmetry group G , we therefore also need to consider a set or space X and need to specify how the group acts on it. This action should certainly satisfy that a consecutive transformations by two group elements g and h should equal a single transformation by the composed group element gh . It is furthermore desirable that the identity group element e leaves any object that it acts on invariant. These observations give rise to the following definition.

DEFINITION 4 (GROUP ACTION). Assume some group G and denote by X a set to be acted on. A (left) group action is then defined as a map

$$\triangleright : G \times X \rightarrow X, \quad (g, x) \mapsto g \triangleright x \tag{27}$$

which satisfies the following two conditions:

- (1) *Associativity*: for any $g, h \in G$ and $x \in X$, the combined action decomposes as $(gh) \triangleright x = g \triangleright (h \triangleright x)$; and
- (2) *Identity*: for any $x \in X$, the identity element $e \in G$ acts trivially, that is, $e \triangleright x = x$.

A set (or space) X that is equipped with a G -action is called G -set (or G -space).

In general, a function $f: X \rightarrow Y$ maps between sets X and Y . Invariant and equivariant functions map more specifically between G -sets and respect their group actions in the sense that they

commute with them. In the case of invariant functions, the output does not change at all when the input is transformed.

DEFINITION 5 (INVARIANT MAP). *Let $f : X \rightarrow Y$ be a function whose domain X is acted on by a G -action \triangleright_X . This function is called G -invariant if its output does not change under transformations of its input; that is, when*

$$f(g \triangleright_X x) = f(x) \quad \text{for any } g \in G, x \in X. \quad (28)$$

This definition is captured graphically by demanding that the following diagram commutes for any $g \in G$, which means that following the top arrow yields the same result as following the bottom path:

$$\begin{array}{ccc} X & & Y \\ g \triangleright_X \downarrow & \nearrow f & \\ X & \xrightarrow{f} & Y \end{array} \quad (29)$$

Many objects in deep learning should be group invariant. For instance, image classification should often be translation invariant, or the ionization energy of a molecule should be invariant under rotations and reflections of the molecule.

Equivariance generalizes this definition by allowing for the output to co-transform with the input: any G -transformation of the function's input leads to a corresponding G -transformation of the output.

DEFINITION 6 (EQUIVARIANT MAP). *Let $f : X \rightarrow Y$ be a function whose domain X and codomain Y are acted on by G -actions \triangleright_X and \triangleright_Y , respectively. This function is called G -equivariant if its output transforms according to transformations of its input; that is, when*

$$f(g \triangleright_X x) = g \triangleright_Y f(x) \quad \text{for any } g \in G, x \in X. \quad (30)$$

The corresponding commutative diagram is given by:

$$\begin{array}{ccc} X & \xrightarrow{f} & Y \\ g \triangleright_X \downarrow & & \downarrow g \triangleright_Y \\ X & \xrightarrow{f} & Y \end{array} \quad (31)$$

As an example of an equivariant map in deep learning, consider a neural network that predicts a magnetic moment of a molecule. Since the underlying laws of physics are rotation invariant, a rotation of the molecule should result in a corresponding rotation of the predicted magnetic moment, that is, the mapping is required to be rotation equivariant. The standard example of an equivariant network layer is the convolution layer: as is easily checked, translations of their input feature map result in corresponding translations of output feature maps. G -steerable convolutions generalize this behavior to more general symmetry groups [Weiler et al. 2018a; Thomas et al. 2018; Weiler and Cesa 2019a].

2.6.3 Group Representations.

Group representation theory describes specifically how symmetry groups act on *vector spaces*. A group representation $\rho_X(g)$ can be thought of as a set of matrices parameterized by group elements $g \in G$ that act on vector space X via matrix multiplication, $\rho_X(g) : X \rightarrow X$.¹ For example, for a vector space of a single 3D Cartesian vector, commonly referred to as (x, y, z) , the

¹More generally, $\rho_X(g)$ can be a linear operator acting on a vector space. If X is finite-dimensional one can always express such operators in terms of matrices relative to some choice of basis.

representation of 3D rotations $SO(3)$ takes the familiar form of 3×3 matrices, which themselves can be parameterized in many ways, e.g., axis-angle, Euler angles, or quaternions are all valid parameterizations of $g \in SO(3)$. Confusingly, group representation colloquially can refer to the matrix representation of the group G on a specific vector space $\rho_X(g)$, the vector space X that the group acts on, or the pair (ρ_X, X) .

The definition of a group puts specific constraints on these matrix representations: they must be invertible with $\rho_X(g^{-1}) = \rho_X(g)^{-1}$, any multiplication of two elements of the representation must also be a representation of the group, and a group representation will always contain the identity matrix \mathbb{I} , the representation of what is commonly referred to as the group element e in group theory literature.

DEFINITION 7 (GROUP REPRESENTATION). Consider a group G and a vector space X . A group representation of G on X is a pair (ρ_X, X) where

$$\rho_X : G \rightarrow GL(X) \quad (32)$$

is a group homomorphism from G to the general linear group $GL(X)$ of X , i.e., to the group of invertible linear maps from X to itself. That ρ_X is a homomorphism means that

$$\rho_X(gh) = \rho_X(g)\rho_X(h) \quad \forall g, h \in G, \quad (33)$$

which ensures that the group composition on the l.h.s. is compatible with the matrix multiplication on the r.h.s.

It is easy to show that $\rho_X(g^{-1}) = \rho_X(g)^{-1}$ and $\rho_X(e) = \mathbb{I}$ follow from this definition.

2.6.4 Irreducible Representations.

Group representations are not unique, and we have the following definition:

DEFINITION 8 (ISOMORPHIC REPRESENTATIONS). Let ρ_X and ρ_Y be representations of group G which act on vector spaces X and Y respectively. Then ρ_X and ρ_Y are said to be isomorphic if there exists a vector space isomorphism $Q : X \rightarrow Y$ such that for all $g \in G$

$$Q\rho_X(g) = \rho_Y(g)Q. \quad (34)$$

If Q is invertible such that $\rho_Y(g) = Q^{-1}\rho_X(g)Q$, then this can be thought of as a change of basis. If Q is unitary, then this is simply a “rotation” of the vector space basis.

One of the most powerful results from group representation theory is that there are reducible and irreducible representations (irreps). A reducible representation contains multiple independent irreps. The vector spaces spanned by different irreps do not mix under group action, i.e., they are independent.

DEFINITION 9 (REDUCIBLE AND IRREDUCIBLE REPRESENTATIONS). A representation ρ_X of group G is said to be reducible if it contains a nontrivial G -invariant subspace. In other words, there exists $V \subset X$ where $V \neq 0$ such that $\rho_X(g)V = V$ for all $g \in G$.

If no such subspace exists then the representation is said to be irreducible (commonly abbreviated as an irrep).

In most cases, when a representation $\rho_X(g)$ is reducible, then there exists similarity transform $Q\rho_Y(g) = \rho_X(g)Q$ such that $\rho_Y(g)$ is block diagonal.

In equivariant neural networks, the symmetry group considered usually acts in some well-defined way on our data. For example, the coordinates of atoms on a molecule would transform under rotation matrices. Hence, our input data is in the vector space of some representations. Since the representations can be broken up into a direct sum of irreducible ones for most groups, we can

specify the way our data transforms as a list of these irreps. In other words, the irreps are the natural data types in equivariant neural networks.

However, there can be multiple representations of the same group which are isomorphic (equivalent). Hence, we have to make a choice when specifying the irreps of our group. Further, we would like a way to label our irreps which is independent of our specific choice of matrices. For the finite groups, one can do so using characters. This is essentially the trace of the matrices in our irreps and is why character tables are used extensively (though there are usually other naming conventions for the irreps of point groups). More details about characters and finding irreps of finite groups can be found in the finite groups part of [Classifying and Computing Irreducible Representations](#) in Appendix.

In the case of infinite groups, using characters is infeasible since there are infinite group elements. Instead, there is well-understood theory on the irreps of semisimple Lie groups we can use. For the case of $SO(3)$ (and $SU(2)$), this essentially gives rise to the degree or angular momentum quantum number ℓ . In general, the irreps are labelled by what are called dominant integral weights. This is the result of a very important theorem called the theorem of highest weights. A brief introduction of the representation theory of semisimple Lie groups can be found in the semisimple Lie groups part of [Classifying and Computing Irreducible Representations](#) in Appendix.

2.7 $SO(3)$ Group and Spherical Harmonics

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The discrete group C_{4v} is one of the simplest non-abelian point groups and has a finite number of irreps. On the other hand, the 3D rotation group $SO(3)$, which is relevant to many scientific domains, is continuous. It has an infinite number of irreps labeled by a positive integer, $\ell = 0, 1, 2, 3, \dots$, which are known as angular momentum, with each irreps having a dimension of $2\ell + 1$.

The irreps of $SO(3)$ are expressed using spherical harmonics, denoted as $Y_m^\ell(\theta, \phi)$ or $Y_m^\ell(\hat{r})$, where $-\ell \leq m \leq \ell$. For a fixed angular momentum ℓ , the $2\ell + 1$ spherical harmonics form the corresponding irreps of the $SO(3)$ group. These spherical harmonics are obtained by solving the 3D Laplace equation in spherical coordinates. The Laplace equation can be written as:

$$\vec{\nabla}^2 f(\vec{r}) = 0. \quad (35)$$

In spherical coordinates, it takes the form:

$$\frac{1}{r^2} \frac{\partial}{\partial r} \left(r^2 \frac{\partial f}{\partial r} \right) - \frac{1}{r^2} \hat{\ell}^2 f = 0, \quad (36)$$

where $\hat{\ell}^2$ represents the angular part:

$$\hat{\ell}^2 f = -\frac{1}{\sin \theta} \frac{\partial}{\partial \theta} \left(\sin \theta \frac{\partial f}{\partial \theta} \right) - \frac{1}{\sin^2 \theta} \frac{\partial^2 f}{\partial \phi^2}. \quad (37)$$

Since the Laplace equation is invariant under rotations in $SO(3)$ as well as the radial variable r , the operator $\hat{\ell}^2$, which depends only on the angular variables, is also rotationally invariant. By employing the method of separation of variables, we can separate the solution $f(\vec{r})$ into the radial part $G(r)$ and the angular part $Y(\theta, \phi)$ such that $f(\vec{r}) = G(r)Y(\theta, \phi)$. Substituting this into the Laplace equation, we obtain two equations:

$$\frac{1}{r^2} \frac{\partial}{\partial r} \left(r^2 \frac{\partial G(r)}{\partial r} \right) - \frac{1}{r^2} \lambda G(r) = 0, \quad \hat{\ell}^2 Y(\theta, \phi) = \lambda Y(\theta, \phi). \quad (38)$$

Let us focus on the second equation, which solely depends on the angular variables. It represents the eigenvalue equation of $\hat{\ell}^2$. Due to boundary conditions, the eigenvalue can only be $\ell(\ell + 1)$,

where ℓ takes values $\ell = 0, 1, 2, 3, \dots$. It turns out that for a given ℓ , there exist $2\ell + 1$ linearly independent solutions, which are the spherical harmonics, denoted as $Y_m^\ell(\hat{r})$, where m is an integer from $-\ell$ to ℓ that labels the $2\ell + 1$ solutions.

The eigenvalue equation of $\hat{\ell}^2$ is rotationally invariant. Consequently, the solutions transform equivariantly under rotations. If $Y_m^\ell(\hat{r})$ is a solution with the eigenvalue $\ell(\ell+1)$, the rotated function $Y_m^\ell(R\hat{r})$ is also a solution with the same eigenvalue. Therefore the rotated function can be expressed as a linear combination of different m values, while keeping the same ℓ value. In other words, we have:

$$Y_m^\ell(R\hat{r}) = \sum_{m'=-\ell}^{\ell} D_{mm'}(R) Y_{m'}^\ell(\hat{r}), \quad (39)$$

where $D_{mm'}(R)$ is a matrix that depends on the rotation R . This matrix represents the transformation of the vector space spanned by the $2\ell+1$ solutions under 3D rotations. Therefore, the $2\ell+1$ solutions, which are characterized by the same ℓ but different m values, form a vector space that is closed under 3D rotations. This vector space corresponds to an irrep of the $SO(3)$ group.

The spherical harmonics are important to many scientific domains as they are the angular part of the solutions to arbitrary rotationally invariant partial differential equations. For instance, adding a potential term $V(r)$ to the Laplace equation only affects the radial equation but not the angular eigenequation. The spherical harmonics are also the solution to the Schrödinger equation of atoms, providing a quantum mechanical description of electrons' wavefunction. In this context, the values of the angular momentum quantum number ℓ correspond to the different types of atomic orbitals: $s(\ell = 0)$, $p(\ell = 1)$, $d(\ell = 2)$, and $f(\ell = 3)$ orbitals.

Conventionally, the spherical harmonics are complex functions taking the following form,

$$Y_m^\ell(\theta, \phi) = \sqrt{\frac{2\ell+1}{4\pi} \frac{(\ell-m)!}{(\ell+m)!}} P_\ell^m(\cos \theta) e^{im\phi} \quad (40)$$

where P_ℓ^m is a polynomial function called the associated Legendre function. It satisfies the relation $P_\ell^{-m}(\cos \theta) = (-1)^m (\ell-m)! / (\ell+m)! P_\ell^m(\cos \theta)$. The spherical harmonics form a complete orthogonal basis for functions defined on a sphere, and any spherical function can be expanded using this basis

$$f(\theta, \phi) = \sum_{\ell,m} a_{\ell,m} Y_m^\ell(\theta, \phi) \quad (41)$$

similar to the Fourier series. Following the orthonormal condition,

$$\int Y_m^{\ell *}(\theta, \phi) Y_{m'}^{\ell'}(\theta, \phi) d\cos \theta d\phi = \delta_{\ell,\ell'} \delta_{m,m'}, \quad (42)$$

The coefficient $a_{\ell,m}$ is $\int Y_m^{\ell *}(\theta, \phi) f(\theta, \phi) d\cos \theta d\phi$. The finer details of $f(\theta, \phi)$ are captured by higher-order spherical harmonics. The spherical harmonics of $\ell = 0, 1, 2$ are listed:

$$\begin{aligned} Y_0^0 &= \sqrt{\frac{1}{4\pi}} \\ Y_{-1}^1 &= \sqrt{\frac{4}{8\pi}} \sin \theta e^{-i\phi}, \quad Y_0^1 = \sqrt{\frac{4}{8\pi}} \cos \theta, \quad Y_1^1 = -\sqrt{\frac{4}{8\pi}} \sin \theta e^{i\phi} \\ Y_{-2}^2 &= \frac{1}{4} \sqrt{\frac{15}{2\pi}} \sin^2 \theta e^{-2i\phi}, \quad Y_{-1}^2 = \frac{1}{2} \sqrt{\frac{15}{2\pi}} \sin \theta \cos \theta e^{-i\phi}, \quad Y_0^2 = \frac{1}{4} \sqrt{\frac{5}{\pi}} (3 \cos^2 \theta - 1), \\ Y_1^2 &= -Y_{-1}^{2 *} \quad Y_2^2 = Y_{-2}^{2 *} \end{aligned} \quad (43)$$

The complex spherical harmonics are convenient to use as it only gains a phase under rotation around the z axis, *i.e.*, $Y_m^\ell(\theta, \phi + \gamma) = e^{im\gamma} Y_m^\ell(\theta, \phi)$. In Cartesian coordinates, it is sometimes more intuitive to consider the real spherical harmonics \mathcal{Y}_m^ℓ which is a linear combination of the complex ones. Notice that $Y_m^\ell(\theta, \phi) = (-1)^m Y_{-m}^{\ell*}(\theta, \phi)$ from the property of the associated Legendre function. The real spherical harmonics are constructed as

$$\mathcal{Y}_m^\ell \equiv \begin{cases} \frac{(-1)^m}{\sqrt{2}} (Y_m^\ell + Y_m^{\ell*}) & m > 0 \\ Y_0^\ell & m = 0 \\ \frac{(-1)^m}{i\sqrt{2}} (Y_{|m|}^\ell - Y_{|m|}^{\ell*}) & m < 0. \end{cases} \quad (44)$$

The $\ell = 0$ real spherical harmonics is the same as the complex one,

$$\mathcal{Y}_0^0 = \sqrt{\frac{1}{4\pi}}, \quad (45)$$

which is a uniform function on the sphere. This is also called the s orbital in atomic physics. The $\ell = 1$ real spherical harmonics are

$$\mathcal{Y}_1^1 = \sqrt{\frac{3}{4\pi}} \sin \theta \cos \phi = \sqrt{\frac{3}{4\pi}} \frac{x}{r}, \quad \mathcal{Y}_{-1}^1 = \sqrt{\frac{3}{4\pi}} \sin \theta \sin \phi = \sqrt{\frac{3}{4\pi}} \frac{y}{r}, \quad \mathcal{Y}_0^1 = \sqrt{\frac{3}{4\pi}} \cos \theta = \sqrt{\frac{3}{4\pi}} \frac{z}{r}$$

Since $(\mathcal{Y}_1^1, \mathcal{Y}_{-1}^1, \mathcal{Y}_0^1) \propto (x, y, z)$, it is clear that $\ell = 1$ real spherical harmonics transform as a 3D vector under rotations. This is one of the advantages of using real spherical harmonics instead of complex ones. In atomic physics, these are called p orbitals. For completeness, the $\ell = 2$ real spherical harmonics are provided below:

$$\mathcal{Y}_{-2}^2 = \sqrt{\frac{15}{4\pi}} \frac{xy}{r^2}, \quad \mathcal{Y}_{-1}^2 = \sqrt{\frac{15}{4\pi}} \frac{yz}{r^2}, \quad \mathcal{Y}_1^2 = \sqrt{\frac{15}{4\pi}} \frac{xz}{r^2}, \quad \mathcal{Y}_0^2 = \sqrt{\frac{5}{16\pi}} \frac{2z^2 - x^2 - y^2}{r^2}, \quad \mathcal{Y}_2^2 = \sqrt{\frac{5}{16\pi}} \frac{x^2 - y^2}{r^2}.$$

In atomic physics, these are also called d orbitals. In the rest of the paper, we mostly employ the real spherical harmonics and simply refer to them as Y_m^ℓ , instead of \mathcal{Y}_m^ℓ for the sake of convenience.

2.8 A General Formulation of Equivariant Networks via Steerable Kernels

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All of the equivariant convolution operations discussed above can be unified in a comprehensive representation theoretic language, the theory of *steerable CNNs* [Cohen and Welling 2017a; Weiler et al. 2018b; Weiler and Cesa 2019b; Cohen et al. 2019; Lang and Weiler 2020; Jenner and Weiler 2022; Cesa et al. 2022b; Weiler et al. 2021]. The feature spaces are in this formulation explained as spaces of *feature vector fields*, whose transformation laws are prescribed by some choice of group representation ρ . The central result is that *any* equivariant linear map between such feature maps is given by conventional convolutions, however, with symmetry constrained “*steerable kernels*”. An implementation of such steerable convolutions for any isometry groups in two and three dimensions is available in [Cesa et al. 2022a].

2.8.1 Feature Vector Fields.

Instead of focusing on a single symmetry group, for instance $E(3) = (\mathbb{R}^3, +) \rtimes O(3)$, steerable CNNs consider any group $\text{Aff}(G) = (\mathbb{R}^d, +) \rtimes G$ of affine transformations of d -dimensional Euclidean space \mathbb{R}^d .² They always contain translations in $(\mathbb{R}^d, +)$, which can be shown to necessitate convolution operations. $G \leq GL(\mathbb{R}^d)$ is any (sub)group of $d \times d$ matrices, including, *e.g.*, rotations, reflections,

²The operation \rtimes is a *semidirect product*, here combining the translation group with transformations in G (*e.g.*, rotations).

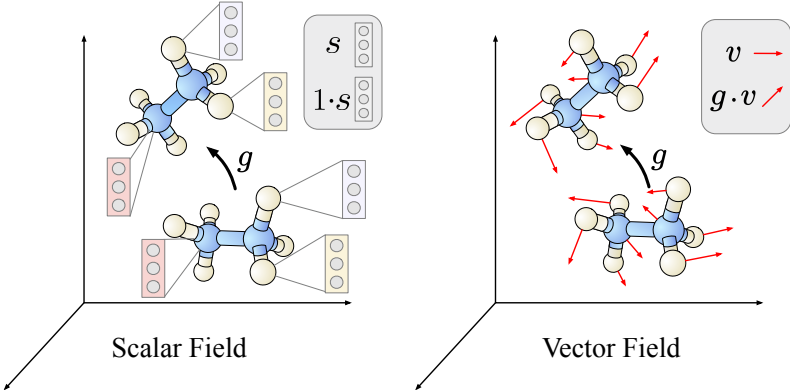


Fig. 8. Scalar and vector fields as simple examples of feature vector fields. Affine groups act on such fields by (1) moving features across space (black arrow), and (2) transforming the features themselves via some group representation ρ . For the trivial representation $\rho(g) = 1$, this explains scalar fields, while $\rho(g) = g$ describes vector fields. All of the feature spaces in the example above correspond to some choice of G -representation, e.g., Wigner-D matrices $\rho = D^\ell$ for tensor field networks and $G = SO(3)$. Steerable CNNs are built from layers which map in an equivariant way between feature fields, for instance from scalar to vector fields or vice versa. Linear equivariant maps are necessarily convolutions, however, with additionally symmetry constrained “steerable kernels”. This figure is adapted from [Weiler and Cesa \[2019b\]](#) with permission.

scaling or shearing. Affine group elements can always be written tg , where $t \in (\mathbb{R}^d, +)$ is a translation and $g \in G$ is a matrix group element.

As mentioned above, steerable CNNs operate on spaces of feature vector fields, which are functions

$$f : \mathbb{R}^d \rightarrow \mathbb{R}^c \quad (46)$$

that assign c -dimensional feature vectors $f(x) \in \mathbb{R}^c$ to any point of Euclidean space $x \in \mathbb{R}^d$. This definition is made in continuous space, however it can ultimately be discretized, e.g., on pixel grids or point clouds.

Recall that equivariant network layers are by definition commuting with group actions – feature fields are therefore not yet fully specified by Equation (46), but are additionally equipped with actions of $\text{Aff}(G)$, examples of which are visualized in Figure 8. The details of these actions are specified by a choice of *field type*. Before stating the general definition of such actions, let’s look at some simple examples:

- *Scalar fields* $s : \mathbb{R}^d \rightarrow \mathbb{R}$ consist of $c = 1$ dimensional features, i.e., scalars. Under pure translations $t \in (\mathbb{R}^d, +)$ they transform like $[t \triangleright s](x) := s(t^{-1}x) = s(x - t)$, i.e., the scalar values are shifted across space.³ This is the transformation behavior of the feature maps of conventional translation equivariant CNNs.
- More general affine group elements tg act according to $[tg \triangleright s](x) := s((tg)^{-1}x) = s(g^{-1}x - t)$ on scalar fields. This adds, for instance, spatial rotations or reflections $g \in O(d)$ of the scalar field, see Figure 8 (left).
- *Tangent vector fields* are functions $v : \mathbb{R}^d \rightarrow \mathbb{R}^d$. As visualized in Figure 8 (right), the transformations $g \in G$ do not only move the vectors to new spatial locations, but act on the

³We use \triangleright to denote group actions on fields. See Definition 4 for a general definition of group actions.

individual vectors themselves, for example by rotating them when $G = SO(d)$. Mathematically, this action is given by $[tg \triangleright v](x) := g \cdot v((tg)^{-1}x)$.

In general, the *field type* is specified by any G -representation $\rho : G \rightarrow GL(\mathbb{R}^c)$, which explains the action of G on individual feature vectors in \mathbb{R}^c . The corresponding action on the feature field as a whole becomes⁴

$$[tg \triangleright f](x) := \rho(g)f((tg)^{-1}x). \quad (47)$$

Note how scalar and tangent vector fields are recovered when choosing the trivial representation $\rho(g) = 1$ or the defining representation $\rho(g) = g$, respectively. Other examples are tensor product representations $\rho(g) = (g^{-\top})^{\otimes r} \otimes g^{\otimes s}$, which correspond to order (r, s) tensor fields, or irreducible representations, explaining e.g., the $2\ell+1$ -dimensional features of tensor field networks when $G = SO(3)$. The group convolutions from Section 2.2 correspond to the regular representation of the cyclic group $G = C_4$ (consisting of 90° rotations). It is given by permutation matrices that shift the field's four channels in a cyclic fashion:

$$\rho(0^\circ) = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}, \quad \rho(90^\circ) = \begin{pmatrix} 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \end{pmatrix}, \quad \rho(180^\circ) = \begin{pmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{pmatrix}, \quad \rho(270^\circ) = \begin{pmatrix} 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 \end{pmatrix}. \quad (48)$$

It can be shown that group convolutions are generally explained by regular G -representations.

2.8.2 Steerable Convolutions.

So far we only described the feature spaces and their group actions, but not the equivariant layers that map between them. Specifically for linear layers Weiler et al. [2018b] showed that the most general linear equivariant maps from input fields f_{in} of type ρ_{in} and output fields f_{out} of type ρ_{out} are given by *convolutions*

$$f_{\text{out}}(x) = [K * f_{\text{in}}](x) = \int_{\mathbb{R}^d} K(x - y) f_{\text{in}}(y) \, dy \quad (49)$$

with convolution kernels

$$K : \mathbb{R}^d \rightarrow \mathbb{R}^{c_{\text{out}} \times c_{\text{in}}}, \quad (50)$$

that are additionally required to be *G -steerable*, i.e., need to satisfy the symmetry constraint

$$K(gx) = \frac{1}{|\det g|} \rho_{\text{out}}(g) K(x) \rho_{\text{in}}(g)^{-1} \quad \forall g \in G, x \in \mathbb{R}^d. \quad (51)$$

Intuitively, the convolution operation ensures translational equivariance, while G -steerability adds equivariance under G -actions, thereby ensuring that the operation is mapping between the specified field types ρ_{in} and ρ_{out} . Note that a scalar convolution kernel would assign a single scalar to each point of \mathbb{R}^d , however, as we are mapping between fields of c_{in} and c_{out} -dimensional feature vectors, the kernels are $c_{\text{out}} \times c_{\text{in}}$ matrix valued.⁵

Performant implementations of convolution operations are readily available, such that the main difficulty in implementing equivariant convolutions is to parameterize the steerable kernels. To this end, observe that kernels form a vector space and that the kernel constraint is *linear* – steerable kernels live therefore in a vector subspace, and it is sufficient to solve for a basis in terms of which steerable kernels are expanded with learnable coefficients. Such bases were derived for $SO(3)$ irreps

⁴This action is known as *induced representation*. Specifically, ρ is a G -representation acting on feature vectors and induces an $\text{Aff}(G)$ -representation which acts on feature fields as a whole.

⁵This is also the case in non-equivariant convolutions. For example, discretized implementations on planar pixel grids in $d = 2$ dimensions represent kernels as arrays of shape $(s_1, s_2, c_{\text{out}}, c_{\text{in}})$, where the first two and the last two axes model the domain and codomain of the continuous kernel $K : \mathbb{R}^2 \rightarrow \mathbb{R}^{c_{\text{out}} \times c_{\text{in}}}$, respectively.

[Weiler et al. 2018b], general representations of any $G \leq O(2)$ [Weiler and Cesa 2019b], and, later, all representations of arbitrary compact groups G (*i.e.*, $G \leq O(d)$) [Lang and Weiler 2020; Cesa et al. 2022b]. They are implemented in the escnn library, which is available for PyTorch and jax [Cesa et al. 2022a].

To clarify the kernel constraint and to demonstrate how steerable CNNs relate to the equivariant models in the previous sections, we turn to explicit examples.

- The simplest example is when ρ_{in} and ρ_{out} are trivial representations, that is, when the kernel maps between scalar fields. Then $K : \mathbb{R}^d \rightarrow \mathbb{R}^{1 \times 1} = \mathbb{R}$ is a scalar kernel satisfying $K(gx) = \frac{1}{|\det g|} K(x)$. For orthogonal group $G \leq O(d)$, *i.e.*, rotations and reflections, the volume scaling factor drops out, and the constraint requires that the kernel is G -invariant (*e.g.*, rotation or reflection invariant).
- For $d = 2$, ρ_{in} being trivial and ρ_{out} being the regular representation of C_4 as defined in Equation (48), the kernel has the signature $K : \mathbb{R}^2 \rightarrow \mathbb{R}^{4 \times 1}$. The constraint becomes $K(gx) = \rho_{\text{out}}(g)K(x)$ which means that the G -rotated kernel on the left hand side should agree with the original kernel after shifting its four channels in a cyclic fashion. This is exactly the construction of kernels from Section 2.2, visualized in the left part of Figure 5.
- We adapt the last example, now requiring both $\rho_{\text{in}} = \rho_{\text{out}}$ to be given by the regular C_4 -representation. The kernel $K : \mathbb{R}^2 \rightarrow \mathbb{R}^{4 \times 4}$ should then satisfy $K(gx) = \rho_{\text{out}}(g)K(x)\rho_{\text{in}}(g)^{-1}$, which means that a spatial rotation equals a simultaneous shift of its rows and columns. The corresponding operation is a regular group convolution, whose kernel is shown in the right part of Figure 5.
- Let now ρ_{in} be trivial and $\rho_{\text{out}} = D^\ell$ be an irrep of $SO(3)$. The corresponding kernels $K : \mathbb{R}^3 \rightarrow \mathbb{R}^{(2\ell+1) \times 1}$ need to satisfy $K(gx) = D^\ell(g)K(x)$, which is solved by kernels whose angular parts are spherical harmonics and whose radial parts are freely learnable. This explains those TP operations in Equation (13) where the input features $\mathbf{h}_j^{\ell_1} := f_{\text{in}}(x_j)$ are of scalar order $\ell_1 = 0$ (trivial) and $\ell_2 = \ell_3 := \ell$.
- If $\rho_{\text{in}} = D^{\ell_1}$ and $\rho_{\text{out}} = D^{\ell_3}$ are both irreps of $SO(3)$ we get $K : \mathbb{R}^3 \rightarrow \mathbb{R}^{(2\ell_3+1) \times (2\ell_1+1)}$. The constraint $K(gx) = D^{\ell_3}(g)K(x)D^{\ell_1}(g)^{-1}$ is then equivalent to $\text{vec}K(gx) = (D^{\ell_1} \otimes D^{\ell_3})(g)\text{vec}K(x)$. Using a Clebsch-Gordan decomposition of the irrep tensor product it is easy to show that such steerable kernels correspond exactly to the general TP operation in Equation (13); see [Weiler et al. 2018b] or [Lang and Weiler 2020] for details.
- The $SO(3)$ -equivariant spherical channel networks (SCNs) from Section 2.4.2 operate on infinite-dimensional feature vectors that are functions on the 2-sphere \mathbb{S}^2 . From a representation theoretic viewpoint, these are just *quotient representations* as described in [Weiler and Cesa 2019b] and [Cesa et al. 2022b]. As an extension to standard steerable CNNs, the steerable kernels used in SCNs are themselves computed from the data via messages.

What is the advantage of the formulation in terms of steerable CNNs?

- (1) It explains equivariant convolutions in a general setting, independent from specific choices of spaces, symmetry groups or group representations. It clarifies thereby how the different approaches in the previous sections relate.
- (2) The previous approaches were introduced by proposing certain operations, which were subsequently shown to be equivariant w.r.t. specific group actions. Steerable CNNs are, conversely, fixing the group actions and subsequently deriving equivariant linear maps between them. While *e.g.*, the TP operations of tensor field networks turn out to be in one-to-one relation to steerable kernel solutions, the kernel constraint formulation allows to prove the *completeness* of these solutions. In many other cases it could be shown that the

authors were only using a subset of all admissible kernels, thus unnecessarily restricting the networks' expressive power [Weiler et al. 2021].

- (3) The approaches above describe only a single field type per model (or class of field types, like irreps). Steerable CNNs allow to build hybrid models whose feature spaces operate simultaneously on feature vectors of regular, irrep, quotient or any other field type.

The abstract representation theoretic formulation suggests natural generalizations to further spaces. Specifically, Cohen et al. [2019] extended steerable CNNs to *homogeneous spaces*, including e.g., spherical convolutions. Weiler et al. [2021] showed that coordinate independent convolutions on *Riemannian manifolds* are similarly requiring G -steerable kernels. This formulation is actually a *gauge field theory*, which proves in particular that the equivariant networks in this section are not only equivariant under global transformations but also under more general local gauge transformations.

Steerable kernels have an interesting connection to the scalar, vector or spherical tensor operators appearing in quantum mechanics. Both are formalized as so-called representation operators, which are described by the famous *Wigner–Eckart theorem* [Jeevanjee 2011b; Wigner 1931]. Lang and Weiler [2020] proved this connection and showed how it allows to solve the kernel constraint in general.

Jenner and Weiler [2022] extended steerable CNNs to the Schwartz distributional setting. This covers in particular *steerable partial differential operators* (PDOs), which explains how the PDOs that appear ubiquitously in the physical sciences respect symmetries.

2.9 Open Research Directions

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In addition to the aforementioned areas, in this section, we highlight several research directions that are among the most cutting-edge and exciting categories. As the field is growing rapidly, we expect to enrich each of the mentioned directions as well as include more topics in the future.

2.9.1 Symmetry Breaking.

Spontaneous symmetry breaking is crucial for explaining many natural phenomena such as magnetism, superconductivity, and even the Higgs mechanism [Beekman et al. 2019; Strocchi 2005], and has been related to neural network training [Ziyin and Ueda 2022]. In such cases, we have a highly symmetric input and desire to predict a lower symmetry output. It is desirable for equivariant networks to deal with this behavior, however, they are fundamentally limited.

Suppose our equivariant model is the function $f : X \rightarrow Y$. For an input x , suppose it is symmetric under a group G . Then for any $g \in G$, $f(gx) = f(x)$. This means the output must also be invariant under G , so it must have the same or higher symmetry. This means we can never predict a single lower symmetry output in an equivariant way. If we try to, the model will just average out all the degenerate outcomes, which might be useless.

There are two perspectives one can take in resolving the symmetry breaking problem for equivariant models. The first is that there is actually one particular degenerate solution we want to predict. In this case, we know from symmetry that we are missing information to perform the task. It turns out by using the gradient of the loss function, we can infer what type of additional input is required to break the symmetry [Smidt et al. 2021].

The second perspective is that all of the lower symmetry outputs are equally valid. In this case, we would like to represent all outputs simultaneously and/or randomly sample from them with equal probability. Treating this case properly is still an open problem.

2.9.2 Empirical Benefits and Expense of Equivariance versus Invariance.

Equivariance has been observed to give measurable benefit over invariance. Increasing the order of features (*i.e.*, the maximum spherical harmonic degree) in $SE(3)$ and $E(3)$ equivariant models has been demonstrated to improve performance [Batzner et al. 2022; Musaelian et al. 2023a; Owen et al. 2023; Yu et al. 2023c,b].

The computational expense of equivariance is dominated by the tensor product (including decomposition into irreps), which involves the contraction of two inputs with the three index Clebsch-Gordan tensor $C_{(l_1, m_1)(l_2, m_2)}^{(l_3, m_3)} X_{(l_1, m_1)} Y_{(l_2, m_2)} = Z_{(l_3, m_3)}$. In voxel models, this contraction can be precomputed for “traditional” convolutional filters, which reduces the computational cost. Otherwise, it must be computed explicitly, *e.g.*, “traditional” point wise convolutions and direct tensor product of features.

It is likely these expenses can be overcome through algorithmic workarounds (*e.g.*, eSCN-like operations [Passaro and Zitnick 2023] as mentioned in Section 2.4.2) and optimization of tensor product operation, whether that be via optimized kernels, domain-specific compilers, or more tailored hardware.

2.9.3 Universality of Equivariant Neural Architectures.

The previous sections discussed in detail how to tailor neural architectures such that they can only represent invariant or equivariant functions, no matter what weights are learned. Although the fundamental goal of this endeavor is to advantageously *restrict* the family of learnable functions to a subfamily known to contain the ground-truth solution, it is important to understand just how expressive a given architecture is within the family of equivariant functions. For instance, is the ground-truth solution still contained in the set of equivariant functions expressible by the architecture family? Clearly, this is an important sanity check.

Informally, an equivariant architecture family is said to be universal if, for any continuous equivariant function and error threshold ϵ , there exists a network in the family, typically that is “large” enough in some sense (*e.g.*, sufficiently many channels, layers, or orders), that approximates that function within error ϵ , according to some functional norm. Happily, prior work has established that many equivariant architectures are universal. In brief, [Yarotsky 2018] first proved that equivariant networks based on polynomial invariants and equivariants are universal, while Bogatskiy et al. [2022] showed that most architectures based on tensor products of irreducible representations of a Lie group are universal. Dym and Maron [2021] also demonstrated that $SE(3)$ -transformers and tensor field networks, two popular architectures operating on point cloud inputs, are universal. However, characterizing the expressivity of graph neural networks is an active research area, and they are in general not universal. Foundational work [Xu et al. 2018] connected the expressivity of message-passing architectures to the Weisfeiler-Lehman hierarchy of graph isomorphism tests, and Joshi et al. [2023] recently began extending this work to *geometric* graph networks (*i.e.*, graphs embedded in 3D space, which is often how point clouds are processed after connecting each point to its nearest neighbors). Such analyses of universality are not sufficient for predicting the relative performances of different equivariant architectures, but are a worthwhile criterion to evaluate when selecting an appropriate equivariant learning method for a given scientific task.

2.9.4 Frame Averaging as an Alternative for Equivariance.

As described in the previous section, most equivariant architectures are therefore expressive within the class of continuous equivariant functions. However, a key drawback of current tensor-based architectures, such as those discussed in Section 2.4 and Section 2.9.2, is their scalability. For example, a point cloud architecture following the template of tensor field networks [Thomas et al. 2018] naively takes time $O(L^6)$ for a single forward pass, where L is the maximum spherical

harmonic index [Passaro and Zitnick 2023]. Recently, frame averaging has emerged as a lightweight alternative to constrained architectures for enforcing equivariance in a learning pipeline.

Formally, a *moving frame* was first defined in 1937 by mathematician Élie Cartan as a smooth, equivariant map $\rho : \mathcal{M} \rightarrow G$, where \mathcal{M} is a manifold on which the Lie group G acts smoothly [Cartan 1937]. The equivariance property ensures that $\rho(gm) = g\rho(m) \forall m \in \mathcal{M}$. Although Cartan defined these objects for the purpose of studying invariants of submanifolds, they provide an intuitive method for enforcing equivariance in a learning pipeline.

First, suppose we are given a function $f : \mathcal{M} \rightarrow \mathcal{Y}$, where \mathcal{Y} is some target space, and a moving frame ρ . We can use ρ to make f invariant (known as the invariantization of f) as

$$f'(m) := f(\rho(m)^{-1}m) \quad \forall m \in \mathcal{M}.$$

It is easy to check that f' is invariant if

$$f'(hm) = f(\rho(hm)^{-1}hm) = f((h\rho(m))^{-1}hm) = f(\rho(m)^{-1}m) = f'(m).$$

Quite similarly, we can use ρ to make f equivariant as

$$f''(m) := \rho(m)f(\rho(m)^{-1}m).$$

One can again check that f'' is equivariant if

$$f''(hm) = \rho(hm)f(\rho(hm)^{-1}hm) = \rho(hm)f(\rho(m)^{-1}m) = h\rho(m)f(\rho(m)^{-1}m) = hf''(m).$$

Here, note that the input and output group actions are the same. To make f equivariant with respect to a different group action on \mathcal{Y} , we simply need another moving frame ρ' that is equivariant with respect to that group action, and can define $f''(m) := \rho'(m)f(\rho(m)^{-1}m)$ instead. Moreover, although moving frames were initially defined for Lie groups acting on manifolds, the straightforward reasoning above applies to any group acting on any space \mathcal{M} .

A straightforward method for equivariant machine learning is therefore to learn the function f using an *arbitrary* architecture, and make it invariant or equivariant using the moving frame constructions above. One must backpropagate through the moving frame, necessitating a degree of smoothness, but the end-to-end framework produces an equivariant function while (1) not requiring any specialization to the group G besides the fixed moving frame, and (2) allowing for an efficient, standard architecture f . Intuitively, the frames method turns the arbitrary function f into an equivariant function by only relying on its behavior at a fixed point on each orbit. Puny et al. [2021] generalize this framework to allow for *averaging* over an equivariant set of points on each orbit instead. Concretely, they define a frame \mathcal{F} more generally as a set-valued function, $\mathcal{F} : \mathcal{M} \rightarrow 2^G \setminus \emptyset$, which is equivariant: $\mathcal{F}(gm) = g\mathcal{F}(m)$, where the equality is between sets. It is then easy to check that the following “frame-averaged” function is equivariant if

$$\langle f \rangle_{\mathcal{F}}(x) := \frac{1}{|\mathcal{F}(x)|} \sum_{g \in \mathcal{F}(x)} f(g^{-1}x).$$

We note that, when the frame maps to $G \in 2^G$ for all elements of the input space \mathcal{M} , then frame-averaging reduces to the well-known Reynolds operator for group-averaging functions (which projects a given function to the closest equivariant function in an L_2 sense, see e.g., [Elesedy and Zaidi 2021]). Moreover, this formulation recovers the classical frame perspective when \mathcal{F} always maps to a set containing exactly one group element. Regardless of the particular choice of \mathcal{F} , it is worth noting that the resultant equivariant pipelines is capable of resulting any equivariant function, so long as the generic architecture is itself universal.

The trade-offs between frames and equivariant architectures remain an active area of research. For example, Pozdnyakov and Ceriotti [2023] motivate frame-averaging as superior to choosing a single frame for rotational equivariance, by observing that methods which canonicalize point clouds

to a single coordinate system are often not *smooth*, in the sense that, adding or removing one point, or changing its position slightly, may drastically change the choice of coordinate system. Instead, they propose computing a weighted average over the frames defined by all pairs of neighbors of one central point, where the weights are specifically chosen to ensure smoothness. However, this procedure is computationally intensive. Duval et al. [2023] address the computational challenge of averaging over a smaller set of coordinate systems defined by principal component analysis, opting to randomly sample a coordinate system at each forward pass during training, sacrificing guaranteed train-time equivariance for efficiency. They demonstrate promising performance-time tradeoffs on materials science tasks, including In light of the difficulty established by these two papers of finding a “good” coordinate frame, or set of frames, over which to average, one promising direction proposed by Kaba et al. [2022] is to *learn* the coordinate frame using a very lightweight equivariant architecture. Finally, several diverse and recent architectures can be interpreted as establishing local coordinate frames [Passaro and Zitnick 2023; Pozdnyakov and Ceriotti 2023], including the structure module of AlphaFold2 [Jumper et al. 2021], and applying the frame-based method for equivariance to local neighborhoods is a promising direction (as it encodes an inductive bias towards not just global, but also local, equivariance). Going forward, frames may provide an appealing alternative to equivariant architectures in applications for which computational efficiency is paramount.

2.9.5 Approximate Equivariance.

Sometimes physical problems do not adhere exactly to group symmetries, but nonetheless symmetries provide a helpful approximation (e.g., if the ground-truth function is still *close* to an equivariant function). Such so-called “approximate symmetries” can arise for a variety of reasons, including boundary effects, discretization error, or something more inherent to the problem, like a partial equivariance or symmetry-breaking property. For example, digit classification is invariant to small-angle rotations, but rotating a “6” yields a “9”, so the problem is not truly rotation-invariant. As a more scientific example, variations in the diffusion coefficient of plate may break the rotational isotropy of heat diffusion [Wang et al. 2022i]. In such problems, an inductive bias towards even approximate symmetry can still advantageously reduce the search space of neural nets.

Residual Pathway Priors [Finzi et al. 2021] first suggested relaxing exact equivariance constraints by parametrizing the learnable weight matrices as sums of equivariant and unconstrained matrices, where the loss function ensures that equivariance is favored. On tasks in vision, synthetic dynamical systems, and reinforcement learning, they demonstrate that their approach is superior in settings with approximate symmetry, yet does not significantly degrade in cases with exact or no symmetry.

More recently, Wang et al. [2022i] proposes a generalization of group CNNs (as well as steerable CNNs, the details of which we omit here but are analogous to the G-CNN case), as shown below:

$$\text{Ordinary group-convolution: } (f *_G \phi)(g) = \sum_{h \in G} f(h)\phi(g^{-1}h).$$

$$\text{Relaxed group-convolution: } (f \widetilde{*}_G \phi)(g) = \sum_{h \in G} f(h)\phi(g, h), \text{ where } \phi(g, h) := \sum_{l=1}^L w_l(h)\phi_l(g^{-1}h).$$

Above, f and ϕ are functions from G to $\mathbb{R}^{c_{in}}$ and $\mathbb{R}^{c_{in} \times c_{out}}$, respectively. Intuitively, such formulations allow the convolutional filter to be location-dependent. The choice of parameter L , the number of filter banks, influences the extent to which the learned function can stray from full symmetry. To encourage symmetry, the network is initialized to ordinary group-convolution (which is a special case of relaxed group convolution), and a term in the loss function discourages variation in each

w_l . They demonstrate superior performance on synthetic smoke plume and experimental jet flow datasets, relative to both perfectly equivariant and generic (not at all symmetric) architectures. Note that these tradeoffs are also justified theoretically in recent work [Petrache and Trivedi 2023].

Others works have presented alternative relaxations of group convolution. van der Ouderaa et al. [2022] instead relax $\phi(g^{-1}h)$ very generally to $\phi(g^{-1}h, h)$, which they parameterize using a few tricks (the group’s Lie algebra and Fourier features). Romero and Lohit [2022] instead relax group convolutions by learning a non-uniform measure over the group.

The previous pipelines were all motivated by, and tested on, data that only approximately adhered to a group symmetry. It is still an open question, however, whether these approximately equivariant networks will offer any long-term advantage over perfectly equivariant networks in tasks with a *genuine* group symmetry. Spherical channel networks for point cloud data (discussed in Section 2.4), for example, achieved state of the art results on the Open Catalyst dataset at the time of their release, despite not having perfect rotation equivariance. However, they have since been surpassed by fully equivariant networks [Passaro and Zitnick 2023]. Nonetheless, for real-world data with noise, approximate symmetry, or even slightly misspecified symmetry, these approaches interpolate advantageously between strictly symmetric and unconstrained architectures.

3 AI FOR QUANTUM MECHANICS

In this section, we provide technical reviews on how to design advanced deep learning methods for learning neural wavefunctions efficiently. In Section 3.1, we give an overview of the definition and how to solve quantum many-body problems in general. In Section 3.2, we introduce methods of learning ground states for quantum spin systems. In Section 3.3, we introduce methods of learning ground states for many-electron systems. An overview of the tasks and representative methods is shown in Figure 9.

3.1 Overview

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Quantum mechanics is the branch of physics that describes the laws governing atoms and subatomic particles [Feynman et al. 1965]. It is of fundamental importance in explaining the physical phenomena of quantum systems in the microscopic domain, ranging from a single particle to molecules and materials [Feynman et al. 2011; Griffiths and Schroeter 2018; Sakurai and Napolitano 2020]. A quantum state contains all the information about a quantum system and is represented as a wavefunction $|\psi\rangle$. Given a set of variables describing the system, such as the position and momentum of its particles, as inputs, the wavefunction $|\psi\rangle$ outputs a complex number that represents the probability amplitude for each possible outcome of a measurement of the system. The wavefunction $|\psi\rangle$ is a high dimensional function that requires an exponential amount of information to fully define. Obtaining the wavefunction of a quantum system is a challenging problem known as the quantum many-body problem. The wavefunction $|\psi\rangle$ is governed by the Schrödinger equation

$$\hat{H}|\psi\rangle = E|\psi\rangle, \quad (52)$$

where \hat{H} is the Hamiltonian operator that describes the motion and interaction of particles in the quantum system, and E is the total energy of that system. In the discrete case, the Hamiltonian operator \hat{H} can be represented as a Hamiltonian matrix H . In principle, all eigenvalues and eigenvectors of H can be obtained through eigenvalue decomposition. Then, the smallest eigenvalue is the ground-state energy of the system, and the corresponding eigenvector is known as the ground state, which is the lowest-energy stationary state. At zero temperature, the ground state fully determines all the properties of the quantum system. Therefore, we focus on how to obtain the ground state of a given quantum system.

The dimension of the Hamiltonian matrix grows exponentially with the size of the quantum systems, such as the number of particles in the system. For instance, the Hamiltonian matrix has a size of $2^N \times 2^N$ for a spin system with size N . Therefore, it is not feasible to obtain the ground state through direct eigendecomposition, even for relatively small systems. An alternative way to *approximately* obtain the ground state and its energy is the variational principle. Consider a parameterized function $|\psi(\theta)\rangle$ that represents a quantum state, where θ are learnable parameters. According to the variational principle, the energy of $|\psi(\theta)\rangle$ must be larger or equal to the ground state energy, which is the smallest eigenvalue of the Hamiltonian matrix. Consequently, to approximate the ground state by θ , one can optimize the variational parameters θ by minimizing the energy of the state. Formally, the expectation value of the energy can be written as

$$E(\theta) = \frac{\langle\psi(\theta)|\hat{H}|\psi(\theta)\rangle}{\langle\psi(\theta)|\psi(\theta)\rangle} = \frac{\int |\psi(s;\theta)|^2 \frac{\hat{H}\psi(s;\theta)}{\psi(s;\theta)} ds}{\int |\psi(s;\theta)|^2 ds} \geq E_0, \quad (53)$$

where E_0 is the ground state energy and E is the energy associate with the quantum state $|\psi(\theta)\rangle$. $\langle\psi(\theta)|$ is the conjugate transpose of $|\psi(\theta)\rangle$, and $\langle\psi(\theta)|\psi(\theta)\rangle$ denotes the dot product of these two

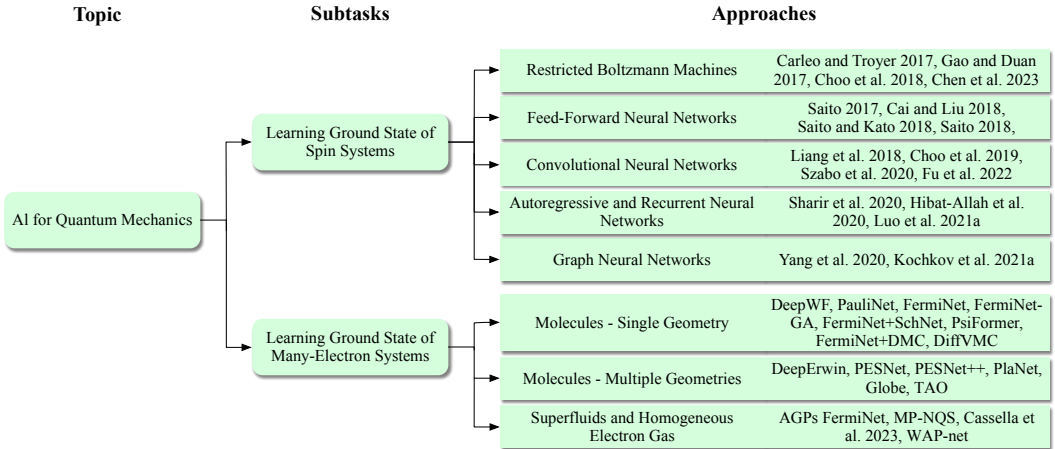


Fig. 9. An overview of the tasks and methods in AI for quantum mechanics. In this section, we focus on two subtasks including learning ground states of spin systems and learning ground states of many-electron systems. The methods for learning ground states of spin systems are grouped in terms of the category of the network they use to represent the quantum state. Specifically, Carleo and Troyer [2017], Gao and Duan [2017], Choo et al. [2018], and Chen et al. [2023] use restricted Boltzmann machines. Cai and Liu [2018], Saito and Kato [2018], Saito [2018], and Saito [2017] use feed-forward neural networks. Liang et al. [2018], Choo et al. [2019], Szabó and Castelnovo [2020], and Fu et al. [2022c] use convolutional neural networks. Sharir et al. [2020], Hibat-Allah et al. [2020], and Luo et al. [2023a] use autoregressive and recurrent neural networks. Yang et al. [2020], and Kochkov et al. [2021a] use graph neural networks. For learning ground states of many-electron systems, one important application is molecules. One category of methods, including DeepWF [Han et al. 2019] PauliNet [Hermann et al. 2020], FermiNet [Pfau et al. 2020], FermiNet-GA [Lin et al. 2023b], FermiNet+SchNet [Gerard et al. 2022], PsiFormer [von Glehn et al. 2023], FermiNet+DMC [Ren et al. 2023; Wilson et al. 2021], and DiffVMC [Zhang et al. 2023c], aim to optimize single geometry of a molecule at a time. Another category of methods, including DeepErwin [Scherbela et al. 2022], PESNet [Gao and Günemann 2021], PESNet++ & PlaNet [Gao and Günemann 2023b], Globe [Gao and Günemann 2023a], and TAO [Scherbela et al. 2023], aim to optimize multiple geometries of the same molecule or even among different molecules simultaneously. Beyond molecules, AGPs FermiNet [Lou et al. 2023] is developed for superfluids. MP-NQS [Pescia et al. 2023], Cassella et al. [2023], and WAP-net [Wilson et al. 2023] are developed for homogeneous electron gas.

vectors. The expectation value of the energy is the mean value of the quantity $\hat{H}\psi(\mathbf{s}; \boldsymbol{\theta})/\psi(\mathbf{s}; \boldsymbol{\theta})$, denoted as the local energy, with respect to a probability distribution $p(\mathbf{s}) = \frac{|\psi(\mathbf{s}; \boldsymbol{\theta})|^2}{\int |\psi(\mathbf{s}; \boldsymbol{\theta})|^2 d\mathbf{s}}$.

The mean value of E_{loc} cannot be obtained exactly due to the high dimensionality of the probability distribution. Instead, one can approximate it by sampling the probability distribution using the Monte Carlo method. In addition, the gradient $\partial E / \partial \boldsymbol{\theta}$ can also be obtained through sampling and is used to optimize the parameters $\boldsymbol{\theta}$ to decrease the energy. This method combining the variational principle and Monte Carlo sampling is called variational Monte Carlo (VMC), outlined in Figure 10.

To sample input configurations according to the probability distribution $p(\mathbf{s})$, Metropolis-Hastings (MH) algorithm is used to create a Markov Chain of input configurations that converges to the stationary distribution p . Specifically, with an input configuration \mathbf{s} on the Markov Chain, a new input configuration \mathbf{s}' is proposed according to the proposal distribution $g(\mathbf{s}'|\mathbf{s})$. And then, \mathbf{s}'

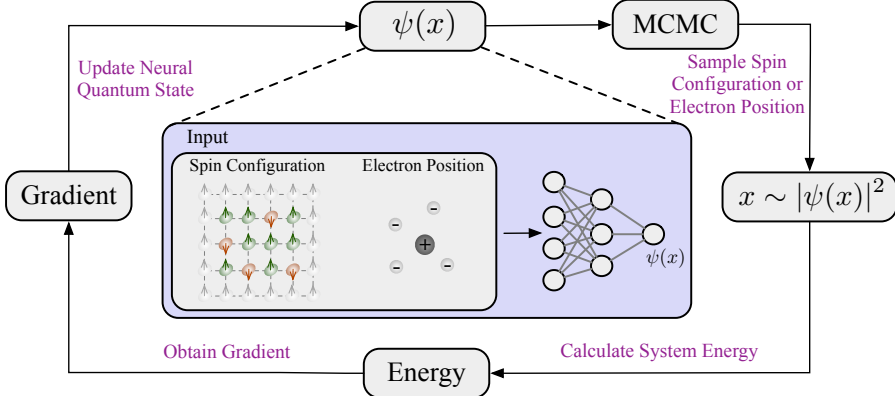


Fig. 10. Pipeline of variational Monte Carlo (VMC). The neural quantum state takes as input a spin configuration or electron positions and outputs the wavefunction value. In VMC, spin configurations or electron positions are sampled using Markov chain Monte Carlo (MCMC) according to the probability distribution determined by the wavefunction. And then, energy is calculated from these samples, and the neural quantum state is updated by the gradient of the energy.

is accepted or rejected according to the acceptance distribution $A(s', s)$. Formally,

$$A(s', s) = \min\left\{1, \frac{p(s')g(s|s')}{p(s)g(s'|s)}\right\}. \quad (54)$$

If s' is rejected, the next input configuration on the Markov Chain is still s . Once the Markov Chain converges to the stationary distribution, samples can be drawn from the Markov Chain, and they are ensured to satisfy the desired distribution.

After input configurations are sampled, we can approximate the system energy as the average of local energy, shown as below:

$$E \approx \frac{1}{N} \sum_{i=1}^N E_{loc}^{(i)}. \quad (55)$$

Then we can use energy E as the loss function and optimize the variational parameters θ to make the system energy as low as possible. Then, the optimized function $|\psi(\theta)\rangle$ with the lowest energy can be seen as a good approximation of the ground state. In this section, we review methods that use neural networks to represent quantum states for learning ground states of quantum spin systems and many-electron systems.

3.2 Learning Ground States for Quantum Spin Systems

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A quantum spin model is a many-body model that describes interacting spins on a lattice resulting from spins of electrons tightly bound to atoms. These spin interactions can result in various magnetic ground states of the system, such as ferromagnetism, anti-ferromagnetism, and even spin liquid, which is an exotic magnetic state that holds promise for topological quantum computing. Understanding the ground state of the quantum spin model provides valuable insight into magnetic materials that are integral to modern technology.

3.2.1 Problem Setup.

In a quantum spin system, each spin can be in two states, spin-up \uparrow , spin-down \downarrow , or their superposition. Any quantum state of N spins can be expressed as a superposition of 2^N spin configurations. All the combinations of spins constitute a computational basis. Specifically, a quantum state can be written as

$$|\psi\rangle = \sum_i \psi(\sigma^{(i)}) |\sigma^{(i)}\rangle, \quad (56)$$

where $|\sigma^{(i)}\rangle$ represents an array of spin configurations of N spins, e.g., $\uparrow\uparrow\downarrow\cdots\downarrow$, and $\psi(\sigma^{(i)})$ is the wavefunction value for the spin configuration $|\sigma^{(i)}\rangle$. The goal is to use neural networks to parameterize the wavefunction and obtain the ground state wavefunction using the variational Monte Carlo described in Section 3.1.

3.2.2 Technical Challenges.

Learning the ground states of quantum spin systems faces several key challenges, including incorporating symmetries of the wavefunction, learning ground state sign structures, and extending approaches to diverse lattice geometries.

Preserving Symmetries: In spin systems, the learned ground state should satisfy certain symmetric structures. Quantum spin systems exhibit rich and intriguing symmetries that are not present in traditional deep learning tasks, such as image object detection. Different from images, lattices are periodic grids with additional symmetries, such as rotations and reflections, which can be classified into 17 wallpaper groups, namely, 17 different plane symmetry groups that make various planar patterns invariant to the corresponding transformations. While most powerful neural networks can learn these symmetries automatically from data according to the universal approximation theorem, this is often hard to achieve due to the enormous solution space and the difficulty of optimization. Incorporating symmetries of the ground state into the neural network structure can guarantee the symmetries of the learned ground state and improve the data efficiency and facilitate finding the optimal solution.

Learning Sign Structures: In quantum mechanics, the sign structure of a wavefunction, in general, refers to the phase of the complex probability amplitude associated with a quantum state. It is challenging to learn the accurate sign structure of the ground state. Sometimes the ground state of quantum spin systems exhibits severe sign problems, where small changes in the spin configuration can cause a change in the sign of the wavefunction, making it difficult for neural quantum states to converge. This phenomenon is even more severe in a frustrated regime and makes it challenging for neural networks to capture complex sign structures of the ground state.

Multiple Geometries: Most existing methods only work for 1D chains or 2D square lattices. However, the lattice geometry of a magnetic material can be far richer than a simple square lattice and has significant effects on its ground state and thus its magnetic properties. The resulting magnetic frustration from this rich geometry provides a host for more exotic magnetic properties to emerge. Therefore, it is crucial to extend the neural network to handle various lattice geometries.

3.2.3 Existing Methods.

Neural quantum states (NQS) have emerged as a powerful variational ansatz for approximating the ground states of quantum many-body systems. NQS can be classified into five different categories based on the type of neural networks, as shown in Figure 9. Carleo and Troyer [2017] propose a pioneering work that uses restricted Boltzmann machine (RBM) to represent quantum states. Due to the success of using RBM as a variational ansatz [Gao and Duan 2017; Deng et al. 2017; Chen

Table 2. Summary of different works on how to address several challenges in solving quantum many-body problems for spin systems, including incorporating symmetries of wavefunctions, learning sign structures, and processing multiple geometries. To consider symmetric ground state structures, solutions include averaging output over transformed inputs according to symmetries or using group convolution. For learning sign structures, solutions include using complex-valued networks to implicitly consider phase, separately modeling amplitude and phase, or incorporating the known Marshall sign rule as the reference sign structure in some special cases. For application on multiple geometries, solutions include processing random graphs and various lattice geometries.

Challenges	Solutions	Methods
Symmetry	Averaging	[Nomura and Imada 2021] [Nomura 2021] [Ferrari et al. 2019]
	Group Convolution	[Choo et al. 2018] [Choo et al. 2019] [Chen et al. 2023] [Roth and MacDonald 2021]
Sign Structure	Complex-Valued	[Carleo and Troyer 2017] [Choo et al. 2019] [Sharir et al. 2020]
	Separate Modeling	[Cai and Liu 2018] [Szabó and Castelnovo 2020] [Kochkov et al. 2021a] [Fu et al. 2022c]
	Marshall Sign Rule	[Choo et al. 2019]
Multiple Geometries	Random Graphs	[Yang et al. 2020] [Kochkov et al. 2021a]
	Various Lattices	[Roth and MacDonald 2021] [Fu et al. 2022c]

et al. 2018a; Choo et al. 2018; Chen et al. 2023], researchers start exploring more expressive deep learning methods to represent quantum states, such as feed-forward neural networks [Cai and Liu 2018; Saito and Kato 2018; Saito 2018, 2017]. Later on, convolutional neural networks (CNNs) [Liang et al. 2018; Choo et al. 2019; Szabó and Castelnovo 2020] are applied to 2D square lattices and are found to represent highly entangled systems effectively. However, CNN cannot be naturally used on non-grid lattices or even random graphs, which necessitated the exploration of graph neural networks (GNNs) [Yang et al. 2020; Kochkov et al. 2021a] for dealing with arbitrary geometric lattices. Moreover, autoregressive and recurrent neural networks (RNNs) are applied to represent quantum states, enabling direct sampling of spin configurations [Sharir et al. 2020; Hibat-Allah et al. 2020; Luo et al. 2023a].

In addition to the different neural network types that affect the expressiveness of neural quantum states, various methods also focus on addressing some of the challenges mentioned above, as shown in Table 2. Incorporating the symmetries of ground states in neural networks can help reduce the hypothesis space. Effectively capturing sign structures of wavefunctions is crucial for neural quantum states to converge easily to optimal solutions. Moreover, the development of single neural quantum states that can function across multiple lattices could significantly enhance their practical usefulness and versatility.

Preserving Symmetries: To capture the symmetry of ground states, most work [Nomura and Imada 2021; Nomura 2021; Ferrari et al. 2019; Choo et al. 2018, 2019; Chen et al. 2023] use the symmetry-averaging technique, which involves transforming the input according to the symmetry group transformation and then taking the average of each output as the final predicted ground state value. Another approach to preserve symmetry is to use group equivariant convolution proposed in [Cohen and Welling 2016]. In GCNN [Roth and MacDonald 2021], authors propose a general framework to use group equivariant convolution to consider the full wallpaper groups and demonstrate the effectiveness of GCNN on square and triangular lattices. GCNN can be mapped to symmetry-averaging models by masking some filters between hidden layers.

Learning Sign Structures: In addition to capturing amplitudes of the ground state, sign structure is also crucial to be learned. Some works learn the amplitude and phases jointly by using a single neural network with complex-valued parameters [Carleo and Troyer 2017; Choo et al. 2019; Sharir et al. 2020]. Choo et al. [2019] uses Marshall sign rule as a reference sign structure and incorporate

it into the network design. The Marshall sign rule provides a simple sign structure that is known for bipartite graphs in some extremal limits, such as $J_1 = 0$ or $J_2 = 0$ for the $J_1 - J_2$ Heisenberg model. However, for ground states in more complex frustrated regimes, there's no such simple prior sign structure to use. [Cai and Liu \[2018\]](#) modifies the feed-forward neural network into two branches to separately predict amplitude and sign of ground states, which are then multiplied together. They use the cosine function as the activation function for predicting the sign, which is suitable for capturing the oscillating features of the input spins. [Kochkov et al. \[2021a\]](#) predicts log amplitude and phase of wave functions separately and shows that predicting phase directly enables effective generalization of the learned sign structure. [Szabó and Castelnovo \[2020\]](#) models amplitude and sign structure using two real-valued neural networks. Specifically, they compute the global phase by summing over predicted phasors for each local spin. Additionally, they adopt a two-stage optimization approach. First, they keep the amplitude of wave functions of all the spin configurations to be the same and only optimize the phase to minimize the system energy. This stage could provide a good initial sign structure since optimal sign structures depend weakly on amplitudes [[Szabó and Castelnovo 2020](#); [Marshall 1955](#)]. And then, the sign structure and amplitude are optimized simultaneously during the second stage.

Multiple Geometries: Most work mentioned above only use the square lattice as the test bed. A practical useful wavefunction ansatz should be applicable and work well across different lattice geometries. GNA [[Yang et al. 2020](#)] proposes universal wavefunction ansatz and conducts experiments on hard-core Boson systems over 2D Kagome lattices, triangular lattices, and randomly connected graphs. [Kochkov et al. \[2021a\]](#) designs another GNN-based ansatz that uses sublattice encoding to denote the node's location in a unit cell that respects the lattice symmetries. In addition to using GNN to achieve applicability on arbitrary lattices, LCN [[Fu et al. 2022c](#)] proposes lattice convolution that uses virtual vertices to augment original lattices to transform them into square lattices, so that a regular CNN can be applied.

3.2.4 Optimization Methods.

There are several ways to optimize the neural network quantum state. The straightforward approach is to calculate the system energy directly as the loss function and use gradient descent methods in deep learning, such as SGD and Adam, to update the network parameters [[Roth and MacDonald 2021](#); [Fu et al. 2022c](#)]. The energy gradient is given as

$$\Delta E_k = \langle E_{loc} O_k^* \rangle - \langle E_{loc} \rangle \langle O_k^* \rangle, \quad (57)$$

where $O_k = \frac{\partial \log \psi(\sigma; \theta)}{\partial \theta_k}$ is the variational derivative with respect to the k -th network parameter, and O_k^* is the complex conjugate of O_k . And $E_{loc} = \sum_j H_{ij} \frac{\psi(\sigma^{(j)}; \theta)}{\psi(\sigma^{(i)}; \theta)}$ is the local energy with respect to spin configuration $\sigma^{(i)}$. $\langle \cdot \rangle$ denotes the expectation value over all the sampled spin configurations. To sample spin configurations from the desired probability distribution $p(\sigma^{(i)}) = \frac{|\psi(\sigma^{(i)})|^2}{\sum_i |\psi(\sigma^{(i)})|^2}$ that is defined by the wavefunction ψ , we can use the Markov Chain Monte Carlo (MCMC) method described in Section 3.1. For the spin system, the proposed spin configuration in Markov Chain can be obtained by randomly flipping a spin in a lattice. So the proposal probability is symmetric, such that $g(\sigma' | \sigma) = g(\sigma | \sigma')$. Thus, the acceptance probability can be simplified as follows:

$$A(\sigma', \sigma) = \min \left\{ 1, \frac{p(\sigma')}{p(\sigma)} \right\}. \quad (58)$$

Another approach to optimize the neural network quantum state is to use stochastic reconfiguration (SR) [[Sorella et al. 2007](#)] that represents an imaginary-time evolution process in the variational space. When a quantum state undergoes imaginary time evolution, it eventually converges to the

ground state of the system. In stochastic reconfiguration, network parameters are updated as

$$\theta \leftarrow \theta - \eta S^{-1} \Delta E, \quad (59)$$

where η is the learning rate, ΔE is the energy gradient, and $S_{ij} = \langle O_i^* O_j \rangle - \langle O_i^* \rangle \langle O_j \rangle$. The only difference from the gradient descent is the presence of a covariance matrix S . Stochastic reconfiguration is generally more robust and less sensitive to the learning rate. However, the limitation of SR is that the size of matrix S equals the number of neural network parameters, making it computationally expensive to compute its inverse for neural networks with large parameters. An alternative optimization method proposed by Kochkov and Clark [2018] that can overcome the limitation of SR is imaginary time supervised wavefunction optimization (IT-SWO). IT-SWO interpolates between the energy gradient and stochastic reconfiguration methods [Kochkov et al. 2021a]. It optimizes the wavefunction ansatz to maximize the overlap between the current variational state $\psi(\sigma; \mathbf{w})$ and the imaginary-time evolved state $(I - \beta H)\psi(\sigma; \mathbf{r})$, where \mathbf{w} and \mathbf{r} represent the parameters of current state and the state at the end of last optimization iteration. At each optimization iteration, IT-SWO first updates the target state and keep it fixed during the current iteration, and then performs multiple inner steps with stochastic gradient descent to update the current state.

3.2.5 Datasets and Benchmarks.

In contrast to traditional machine learning tasks, models used to determine the ground state of quantum spin systems cannot be trained on a pre-existing dataset. Instead, the model is trained for a specific quantum spin system, which is defined by the lattice and Hamiltonian. During each step of the training process, data are dynamically sampled from the wavefunction (neural network) of a quantum system. This approach is known as concurrent machine learning as described by E et al. [2020]. Typically, a variety of lattice systems are considered, such as square, honeycomb, triangular, and kagome lattices. The most commonly used Hamiltonian is J_1 - J_2 quantum Heisenberg model, which is the prototypical model for studying the magnetic properties of quantum materials. In terms of evaluation metrics, the energy of the systems serves as a measure of how closely the approximated ground state aligns with the true ground state. A lower energy indicates a more accurate approximation.

3.2.6 Open Research Directions.

Neural network quantum states have shown promise in representing the ground state of quantum spin systems, but several challenges still need further exploration. First, we need more expressive and efficient wavefunction ansatz that can scale to arbitrarily large systems in order to be practically useful. As the size of the system increases, the intricacy of spin interactions also increases. Second, designing neural wavefunctions with provable sufficient expressiveness remains an open problem, especially for quantum systems exhibiting highly frustrated regimes and strong correlations. Third, fermionic lattice systems have a unique antisymmetry property of wavefunctions, which makes it difficult for neural quantum states to efficiently encode this symmetry. Finally, in variational Monte Carlo, Markov Chain Monte Carlo (MCMC) is commonly used to sample spin configurations from the probability distribution determined by wavefunctions and then calculate the system energy. However, performing exact sampling with MCMC is difficult, and the samples may still exhibit correlations, leading to inaccurate energy estimations. Usually, to decrease autocorrelation between samples, N annealing MCMC steps are added between two samples, where N represents the size of the system. However, this makes the sampling process computationally expensive for larger systems. A potential solution to this challenge is proposed in [Sharir et al. 2020]. They use an autoregressive model to represent quantum states, which bypass the MCMC sampling and can support more efficient and exact sampling.

3.3 Learning Ground States for Many-Electron Systems

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Compared to the quantum spin system, a more general scenario in quantum mechanics is when particles can move freely in space, such as electrons moving in molecules and solids. Studying many-electron systems is at the core of quantum chemistry, where properties of molecules are directly computed from first principles based on quantum physics. Specifically, accurately describing the ground states of molecules is of great interest because the ground state determines the most stable state of a molecule and is important to the understanding of its structural and chemical properties. Compared to quantum spin systems, the spin of the electrons does not occur in the Hamiltonian and, thus, can be fixed a priori [Foulkes et al. 2001]. As a result, the wavefunction only acts on the spatial coordinates in \mathbb{R}^3 of each of the N electrons. However, the search space for suitable wavefunctions grows exponentially with the number of electrons N . Moreover, the fermionic nature of electrons significantly increases the difficulty of the problem [Ceperley 1991]. For example, it has been shown that solving the sign problem, which can arise for fermions due to Pauli exclusion, is NP-hard [Troyer and Wiese 2005]. In this section, we mainly discuss learning ground states of molecules with quantum Monte Carlo (QMC) approaches. Additionally, similar methods have been applied to superfluid and homogeneous electron gas (HEG), which we will also review briefly for completeness.

3.3.1 Problem Setup.

Molecules are composed of electrons and atomic nuclei. Within the Born-Oppenheimer approximation [Born and Oppenheimer 1927], atomic nuclei are treated as fixed particles, hence quantum states are completely determined by electrons' spins and 3D coordinates. At ground states, spins of electrons can be determined by chemical rules, such as the Aufbau principle, the Hund's rule and the Pauli exclusion principle. Hence, we are able to define the wavefunction solely in terms of the electron coordinates. Formally, given N^\uparrow electrons with spin-up, N^\downarrow electrons with spin-down. The set of their 3D Cartesian coordinates is defined as $\mathbf{r} = [\mathbf{r}_1, \dots, \mathbf{r}_{N^\uparrow+N^\downarrow}] \in \mathbb{R}^{(N^\uparrow+N^\downarrow) \times 3}$, where the first N^\uparrow electrons have spin-up and the last N^\downarrow electrons have spin-down. A wavefunction $\psi : \mathbb{R}^{(N^\uparrow+N^\downarrow) \times 3} \rightarrow \mathbb{R}$ maps the set of coordinates to a scalar value. In the continuous case, the Hamiltonian operator $\hat{H} : (\mathbb{R}^{(N^\uparrow+N^\downarrow) \times 3} \rightarrow \mathbb{R}) \rightarrow (\mathbb{R}^{(N^\uparrow+N^\downarrow) \times 3} \rightarrow \mathbb{R})$ is a function that maps a wavefunction to another function, defining the energy of a molecule, and is defined as

$$[\hat{H}\psi](\mathbf{r}) = - \sum_i \frac{1}{2} \nabla_i^2 \psi(\mathbf{r}) + V(\mathbf{r}), \quad (60)$$

where the first term represents the kinetic energy and the second term represents the Coulomb potential, which is defined as

$$V(\mathbf{r}) = \sum_{i < j} \frac{1}{\|\mathbf{r}_i - \mathbf{r}_j\|_2} - \sum_{i,I} \frac{z_I}{\|\mathbf{r}_i - \mathbf{c}_I\|_2} + \sum_{I < J} \frac{1}{\|\mathbf{c}_I - \mathbf{c}_J\|_2}, \quad (61)$$

where \mathbf{c}_I denotes the coordinate of an atomic nucleus and z_I denotes its atomic charge. The terms define Coulomb potential between electron-electron pairs, electron-atom pairs, and atom-atom pairs, respectively. Note that Hamiltonian is Hermitian and therefore the wavefunction ψ is real-valued. Additionally, it is noteworthy that the Hamiltonian does not depend on electron spins. Thus, one can fix the spins a priori. Given the Hamiltonian, the local energy $E_{loc}(\mathbf{r}) = \frac{[\hat{H}\psi](\mathbf{r})}{\psi(\mathbf{r})}$ can

be expressed as

$$E_{loc}(\mathbf{r}) = -\frac{1}{2} \sum_i^{(N^\uparrow + N^\downarrow) \times 3} [\partial_i^2 \log |\psi(\mathbf{r})| + (\partial_i \log |\psi(\mathbf{r})|)^2] + V(\mathbf{r}), \quad (62)$$

where i goes through all $(N^\uparrow + N^\downarrow) \times 3$ spatial coordinates.

A fundamental constraint for a many-electron system is that its wavefunction must be antisymmetric upon permutation of two electrons with the same spin, a concept originating from Pauli exclusion. In quantum mechanics, exchanging two indistinguishable particles does not affect the probability density of particles. In our case, two electrons cannot be distinguished if they have the same spin. Hence, $\psi(\dots, \mathbf{r}_i, \dots, \mathbf{r}_j, \dots)^2 = \psi(\dots, \mathbf{r}_j, \dots, \mathbf{r}_i, \dots)^2$, for any (i, j) with same spins. Further, indistinguishable particles are classified into bosons, such as photons, and fermions, such as electrons, according to their exchange symmetry [Feynman et al. 1965], which refers to whether the wavefunction ψ remains unchanged or changes sign upon exchanging the positions of two particles. Electrons are fermions, so the wavefunction must be antisymmetric upon permutation of two electrons with the same spin, i.e., $\psi(\dots, \mathbf{r}_i, \dots, \mathbf{r}_j, \dots) = -\psi(\dots, \mathbf{r}_j, \dots, \mathbf{r}_i, \dots)$. This antisymmetry property leads to the Fermi-Dirac statistics in particle distributions, and thus fundamentally changes the behavior of fermions. Consequently, the task of finding ground states can be formulated as a constrained optimization problem. In the context of variational Monte Carlo, the wavefunction ψ is approximated by a parametrized class of functions ψ_θ . In this case, learning ground states is equivalent to the following optimization problem:

$$\psi_\theta : \mathbb{R}^{(N^\uparrow + N^\downarrow) \times 3} \rightarrow \mathbb{R} \quad (63)$$

$$\min_{\theta} \mathbb{E}_{p_\theta}[E_{loc}(\mathbf{r}; \theta)], \quad p_\theta \propto \psi_\theta^2 \quad (64)$$

$$\text{s.t. } \psi_\theta(\dots, \mathbf{r}_i, \dots, \mathbf{r}_j, \dots) = -\psi_\theta(\dots, \mathbf{r}_j, \dots, \mathbf{r}_i, \dots), \text{ where} \quad (65)$$

$$\text{pair } (i, j) \text{ satisfies } 1 \leq i, j \leq N^\uparrow \text{ or } N^\uparrow + 1 \leq i, j \leq N^\uparrow + N^\downarrow.$$

In the above optimization objective, the energy expectation is calculated as an average over samples obtained using Monte Carlo sampling. By the variational principle mentioned previously (Equation (53)), the energy expectation of any wavefunction is guaranteed to be larger than the ground state energy, and the lower bound is attained when ψ_θ converges to the ground state wavefunction.

The above formulation provides a framework for obtaining the ground-state energy for a single molecule. In this section, we additionally consider the setting for jointly optimizing for multiple geometries. For example, we are usually interested in studying the change in energy based on structural changes in a molecule. Joint optimization improves computational efficiency by eliminating the need to optimize again for every nucleus configuration. Formally, we define the potential energy surface (PES) as a function $E : \mathbb{M} \rightarrow \mathbb{R}$, where $\mathbb{M} = \{M = \{\mathbf{c}_i, z_i\}_{i=1}^{|\mathbb{M}|}, \mathbf{c}_i \in \mathbb{R}^3, z_i \in \mathbb{Z}\}$ is the set of possible molecules, maps from the molecular structure (coordinates and charges of nuclei) to the energy. Classically, to obtain a PES, one needs to repeat single structure calculation multiple times. The advent of neural network-based solution makes it possible to model the ab-initio solutions for PES with a single model. Concretely, in this setting, one is interested in finding a wavefunction $\psi_\theta : \mathbb{R}^{(N^\uparrow + N^\downarrow) \times 3} \times \mathbb{M} \rightarrow \mathbb{R}$, where the wavefunction is now also dependent on the molecular structure, in addition to the electron coordinates. Following Gao and Günemann [2023a], we call such ψ_θ the *generalized wavefunction*. Note that this formulation should not be confused with the Schrödinger equation without the Born-Oppenheimer approximation where the nuclei are treated as waves and are thus considered as part of the wavefunction. This is not the case here,

we still only model the electronic wavefunction but condition the wavefunction on the molecular structure. Finally, using the generalized wavefunction, the potential energy surface can be derived as $E(M) = \int \psi_\theta(\mathbf{r}, M) \hat{H}_M \psi_\theta(\mathbf{r}, M) d\mathbf{r}$ where \hat{H}_M refers to the Hamiltonian of molecule M .

Additionally, superfluids and homogeneous electron gas can also be modeled as fermions in continuous space. However, these problems do not involve nuclei and use a different potential energy in the Hamiltonian. Another major difference is that these problems are periodic in space. Nevertheless, the general approaches for the single-molecule setting can still be applied.

In our discussion, the wavefunction directly takes the continuous electron coordinates as input, as a result, the antisymmetric constraint must be encoded explicitly in ψ_θ . Methods with such continuous representation are called the first-quantization methods. Alternatively, in the second-quantization methods [Barrett et al. 2022; Choo et al. 2020; Herzog et al. 2022], wavefunctions are expressed in a discrete basis of antisymmetric functions. The comparison between the first and second quantization is discussed in Hermann et al. [2022]. First-quantization methods have gained more popularity recently because of their flexibility beyond the choice of an antisymmetric basis. Furthermore, to get a more complete description of many-electron systems, excited states [Entwistle et al. 2023; Feldt and Filippi 2020] can also be studied using similar approaches as the ground states. However, the details are out of the scope of our discussion in this section.

3.3.2 Technical Challenges.

There are several challenges related to finding many-electron ground states with QMC, including satisfying the fermion antisymmetry constraint, designing expressive neural networks for individual electrons (orbitals), achieving good optimization, and effectively learning generalized wavefunctions for multiple geometries to improve computational efficiency.

Fermion Antisymmetry: As introduced in Section 3.3.1. Fermion antisymmetry is a hard constraint imposed by quantum physics, and a neural wavefunction for electrons must adhere to it strictly. Failing to encode the antisymmetry constraint will void the variational guarantees and result in unphysically lower energies. Although deep neural networks can approximate arbitrarily complex functions, imposing such hard constraints poses a unique challenge.

Orbital modeling: Electrons interact with each other via the Coulomb potential and Pauli exclusion, which can result in highly non-linear landscapes in wavefunctions. Therefore, the networks must have a strong capacity to model the wavefunction of each electron (dubbed as orbitals) while accounting for the interactions with other electrons. Additionally, quantum physics gives us some prior knowledge of the system, which may be hard to model directly with neural networks. Thus, incorporating physics knowledge in orbital modeling is important to obtain solutions that respect physics.

Optimization: Although in principle we can get arbitrary approximation accuracy with VMC, it is challenging to achieve effective optimization of neural wavefunctions towards ground states. This is in part due to high accuracy requirement of the problem. The chemical accuracy is defined as 1 kcal/mol (1.594 mE_h or 0.043 eV) [Pfau et al. 2020], which is very small compared to the total energy. For example, the N_2 molecules, the error in energy estimation must be lower than 0.2% to be useful for chemical applications [Gerard et al. 2022]. Consequently, effective optimization methods are crucial to obtain accurate and stable optimization.

Multiple Geometries: There are some unique challenges related to the multiple geometries setting. Firstly, special considerations are necessary to make the learned wavefunctions adaptable to various molecular configurations, including different nucleus positions and variable numbers of nuclei and electrons (e.g., ionic systems) while respecting the fermion antisymmetry. Secondly, as the PES E is an observable metric, it is invariant to the Euclidean group $E(3)$, i.e., translations,

rotations, and reflections of the molecule, as well as the permutation group S_M . However, as an abstract concept, the electronic wavefunction does not exhibit such symmetries. Thus, the challenge is to design generalized wavefunctions that result in invariant energies. It can be shown that to obtain such a behavior one needs to design symmetry-breaking covariant wavefunctions. Thirdly, prior knowledge gives us additional constraints about limit behaviors. One such property is size-consistency, *i.e.*, the energy of a duplicated non-interacting system is twice the energy of the single system. Implementing such behaviors into wavefunctions remains a challenge to reduce the function search space. Lastly, while the wavefunction is directly linked to the energy, obtaining the energy from the wavefunction remains expensive as it requires numerical integration. Approximate inference methods promise to accelerate the process and enable high-resolution PES.

3.3.3 Existing Methods.

Recently, VMC-based neural networks have shown strong ability in modeling ground states of many-electron systems. Classical methods such as DFT or CCSD(T) either result in unreliable results in strongly correlated settings, *e.g.*, when bonds break, or scale unfavorably with the system size. VMC coupled with deep neural networks has shown to be able to outperform classical methods [Pfau et al. 2020; Gerard et al. 2022]. In particular, although DFT scales better than deep VMC methods ($O(N^3)$ v.s. $O(N^4)$), much higher accuracies can be achieved by deep VMC methods. On the other hand, deep VMC methods are faster than CCSD(T) and can achieve similar or higher accuracy. Moreover, although CCSD(T) can be applied to larger molecules, a smaller basis set must be picked. In the following, we briefly introduce how the challenges listed in Section 3.3.2 are resolved by existing methods. We first describe how to encode fermion antisymmetry in networks, in particular with Slater determinants. Next, we describe how networks are designed in existing methods. With these two components, we can already have a working neural wavefunction model. We then discuss how to effectively optimize the networks to reach ground states. Finally, we describe strategies to reuse and accelerate the computations via generalized wavefunctions. The challenges and existing methods are summarized in Table 3.

Fermion Antisymmetry: To design wavefunctions that satisfy the fermion antisymmetry (Equation (65)), a well-established method is the Slater determinant [Slater 1929]. The Slater determinant wavefunction is computed as the determinant of a matrix which is constructed by applying N molecular orbital functions to each of the N electrons so that each row of the matrix encodes one electron. The key motivation is that when two electrons are swapped, the two corresponding rows in the matrix are also swapped, so its determinant will change sign. Formally, let ϕ^\uparrow and $\phi^\downarrow : \mathbb{R}^3 \times \mathbb{R}^{(N^\uparrow + N^\downarrow) \times 3} \rightarrow \mathbb{R}^{N^\uparrow + N^\downarrow}$ be two single-orbital functions that map the 3D electron coordinates to a $(N^\uparrow + N^\downarrow)$ -dimensional feature vector, where ϕ^\uparrow is used to encode spin-up electrons and ϕ^\downarrow is used to encode spin-down electrons. ϕ^\uparrow and ϕ^\downarrow take an electron coordinate as well as all electron coordinates as input and produce a $(N^\uparrow + N^\downarrow)$ -dimensional vector. The objective of single orbital functions is to generate an embedding for each electron and the information from all electrons is used to provide context information. After encoding each electron with ϕ^\uparrow or ϕ^\downarrow , a set of $N^\uparrow + N^\downarrow$ feature vectors is obtained, each containing $N^\uparrow + N^\downarrow$ elements. The features are stacked into a matrix where each row represents one electron. The Slater determinant wavefunction ψ is

Table 3. Summary of challenges and existing methods for learning many-electron ground states. For electrons, a special challenge arises from fermion antisymmetry imposed by quantum physics. Most existing wavefunction models solve it through Slater determinants but have different network designs to model orbital functions. Moreover, to make learning accurate and practical, it is crucial to achieve effective optimization. Finally, the diversity and flexibility of molecules require methods to handle multiple geometries to increase computational efficiency.

Challenges	Fermion Antisymmetry	Orbital Modeling	Optimization	Multiple Geometries
Methods	Main method: Slater determinant Others: Explicit construction Pairwise construction AGP	PauliNet FermiNet FermiNet+SchNet PsiFormer Moon WAP-net MP-NQS	Framework: VMC DMC DiffVMC Optimizer: KFAC CG	DeepErwin PESNet PESNet++ PlaNet Globe TAO

then computed as the determinant of that matrix:

$$\psi(\mathbf{r}) = \det \begin{bmatrix} \phi^\uparrow(\mathbf{r}_1; \mathbf{r})^T \\ \vdots \\ \phi^\uparrow(\mathbf{r}_{N^\uparrow}; \mathbf{r})^T \\ \phi^\downarrow(\mathbf{r}_{N^\uparrow+1}; \mathbf{r})^T \\ \vdots \\ \phi^\downarrow(\mathbf{r}_{N^\uparrow+N^\downarrow}; \mathbf{r})^T \end{bmatrix}. \quad (66)$$

Note that the orbital function for spin-up and spin-down electrons are different so that the antisymmetry is present only exchanged.

For example, when $N^\uparrow = 2$ and $N^\downarrow = 1$ (Li atom), fermion antisymmetry is ensured by Slater determinants when \mathbf{r}_2 and \mathbf{r}_3 are exchanged, as demonstrated below:

$$\psi(\mathbf{r}_1, \mathbf{r}_2, \mathbf{r}_3) = \det \begin{bmatrix} \phi_1^\uparrow(\mathbf{r}_1) & \phi_2^\uparrow(\mathbf{r}_1) & \phi_3^\uparrow(\mathbf{r}_1) \\ \phi_1^\uparrow(\mathbf{r}_2) & \phi_2^\uparrow(\mathbf{r}_2) & \phi_3^\uparrow(\mathbf{r}_2) \\ \phi_1^\downarrow(\mathbf{r}_3) & \phi_2^\downarrow(\mathbf{r}_3) & \phi_3^\downarrow(\mathbf{r}_3) \end{bmatrix} = -\det \begin{bmatrix} \phi_1^\uparrow(\mathbf{r}_1) & \phi_2^\uparrow(\mathbf{r}_1) & \phi_3^\uparrow(\mathbf{r}_1) \\ \phi_1^\uparrow(\mathbf{r}_3) & \phi_2^\uparrow(\mathbf{r}_3) & \phi_3^\uparrow(\mathbf{r}_3) \\ \phi_1^\downarrow(\mathbf{r}_2) & \phi_2^\downarrow(\mathbf{r}_2) & \phi_3^\downarrow(\mathbf{r}_2) \end{bmatrix} = -\psi(\mathbf{r}_1, \mathbf{r}_3, \mathbf{r}_2). \quad (67)$$

To further increase expressiveness, multiple Slater determinants can be computed, each with a different set of orbital functions, and the final wavefunction is the linear combination of Slater determinants. When k Slater determinants are used, letting $w_p \in \mathbb{R}$ be the weights, the final wavefunction is computed as:

$$\psi(\mathbf{r}) = \sum_{p=1}^k w_p \det \begin{bmatrix} \phi^{\uparrow p}(\mathbf{r}_i; \mathbf{r})^T \\ \vdots \\ \phi^{\downarrow p}(\mathbf{r}_j; \mathbf{r})^T \\ \vdots \end{bmatrix}. \quad (68)$$

Besides Slater determinants, there are other ways to achieve antisymmetry. DeepWF [Han et al. 2019] and Pang et al. [2022] propose to enforce the antisymmetry to every electron pair. The wavefunction is defined in the form of $\prod_{i < j} (f(\mathbf{r}_i; \mathbf{r}) - f(\mathbf{r}_j; \mathbf{r}))$ where f outputs a scalar value. When a pair of $(\mathbf{r}_i, \mathbf{r}_j)$ is swapped, the sign of the product will be changed. This strategy

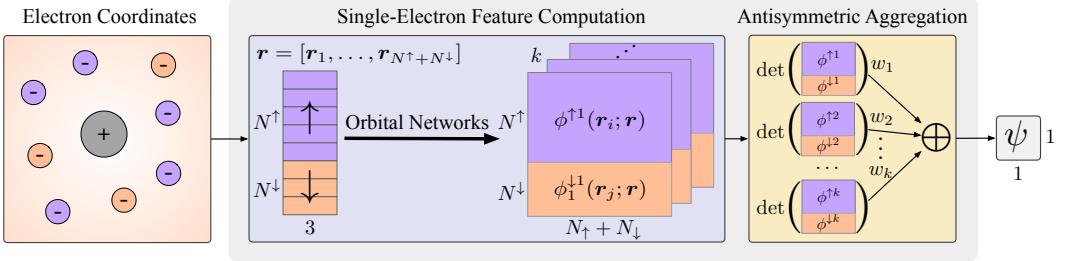


Fig. 11. Pipeline of many-electron wavefunction computation with Slater determinants, illustrated for molecules. The input to the network is a set of 3D electron coordinates with N^\uparrow electrons with spin-up and N^\downarrow electrons with spin-down. The spin structure (\uparrow or \downarrow) as well as the positions of atomic nuclei are fixed. A neural network is used to produce k ($N^\uparrow + N^\downarrow$)-dimensional features vectors for each electron, which are then concatenated into k ($N^\uparrow + N^\downarrow$) \times ($N^\uparrow + N^\downarrow$) matrices. Finally, the determinants of these matrices are computed and a linear combination of them gives the final wavefunction value.

is shown to be a special case of the Slater determinant (it can be written as the determinant of a Vandermonde matrix [Pang et al. 2022]) but with less computational cost. Lin et al. [2023b] generalizes the sum-of-product determinant computations by explicitly considering all possible permutations of electrons. The final wavefunction is the sum of results from all permutations, *i.e.*, $\sum_{\pi} \text{sign}(\pi)g(\pi(\mathbf{r}))$, where π iterates over all permutations $\text{sign}(\pi)$ gives the sign of each permutation, and g is a function maps the permuted \mathbf{r} to a scalar. This however leads to a factorial complexity. Finally, antisymmetric geminal power (AGP) wavefunctions have shown great success in modeling superfluids with neural-network wavefunctions [Lou et al. 2023]. There one uses pairwise orbital functions $\phi : \mathbb{R}^3 \times \mathbb{R}^3 \times \mathbb{R}^{(N^\uparrow \times N^\downarrow) \times 3} \rightarrow \mathbb{R}$, and constructs the wavefunction as $\psi(\mathbf{r}) = \det \Phi$, where $\Phi_{i,j} = \phi(r_i, r_j, \mathbf{r})$, $i \in \{1, \dots, N^\uparrow\}$, $j \in \{N^\uparrow + 1, \dots, N^\uparrow + N^\downarrow\}$.

Orbital Modeling: Although Slater determinants solves the exchange antisymmetry for many-electron systems, it does not provide any guarantee on the accuracy of the optimized wavefunction. To accurately model the ground-state wavefunction, the orbital functions ϕ^\uparrow and ϕ^\downarrow must stem from a flexible function family. Classically, orbital functions are modeled as single-particle orbitals from solutions for the single-particle Schrödinger's equation. FermiNet [Pfau et al. 2020] and PauliNet [Hermann et al. 2020] successfully use neural networks to model orbital functions while using the Slater determinant as antisymmetric aggregation, where all 3D electron coordinates are first encoded as a $(N^\uparrow + N^\downarrow) \times k$ -dimensional feature vector with permutation-equivariant neural networks ϕ_θ^\uparrow and ϕ_θ^\downarrow . The vectors are then concatenated into k ($N^\uparrow + N^\downarrow$) \times ($N^\uparrow + N^\downarrow$) matrices (*e.g.*, $k = 16$). Finally, the determinants of these matrices are computed and the final wavefunction value is the linear combination of determinants. As a result, the parameters in the network are the parameters in the orbital networks and the combination weights of determinants. Due to their prominent performances, follow-up works mostly follow the same pipeline, which is shown in Figure 11. To build a complete representation of the quantum state, one must consider all pairwise information between all particles, including electrons and nuclei. To this end, commonly used input features are relative vectors and distances between electron-electron and electron-nucleus pairs. Additionally, a symmetric part $e^{J(\mathbf{r}; \theta)}$ can be multiplied to the final wavefunction, which is called the *Jastrow factor*.

Orbital Modeling – Single-Electron Feature Computation: To capture the complex physics interactions, it is necessary that the orbital networks ϕ_θ^\uparrow and ϕ_θ^\downarrow consider all electron positions

collectively. Hence, when encoding one electron, all other electrons are also encoded to provide context information. As a result, ϕ_θ^\uparrow and ϕ_θ^\downarrow must be able to gather information from other electrons. PauliNet [Hermann et al. 2020] uses distance-based convolution to gather information from neighborhood electrons similar to SchNet [Schütt et al. 2018], where convolution weights are computed based on the relative distances: $\mathbf{h}'_i = \sum_j \mathbf{w}(\|\mathbf{r}_j - \mathbf{r}_i\|_2; \theta) \odot f(\mathbf{h}_j)$, where \odot denotes element-wise product and \mathbf{w} computes differently for different spins. A similar computation is also applied for nuclei. FermiNet [Pfau et al. 2020] uses mean field information about the electronic structure and pairwise distance features between each pair of electrons. Concretely, FermiNet maintains two computation branches that compute one-particle features and pairwise features, respectively. At each layer, both one-particle features \mathbf{h}_i and pairwise features \mathbf{h}_{ij} are averaged over electrons to get global representations for each spin, which are further concatenated to the one-particle features for the next layer $\mathbf{h}'_i = f([\mathbf{h}_i, \sum_{j, \sigma(j)=\uparrow} \mathbf{h}_j, \sum_{j, \sigma(j)=\downarrow} \mathbf{h}_j, \sum_{j, \sigma(j)=\uparrow} \mathbf{h}_{ij}, \sum_{j, \sigma(j)=\downarrow} \mathbf{h}_{ij}])$, where $\sigma(i)$ denotes the spin of the i -th electron and $[\cdot]$ denotes concatenation. FermiNet+SchNet [Gerard et al. 2022] replaces the simple interactions in FermiNet by integrating the convolutions from PauliNet. By abandoning the mean field of FermiNet, PsiFormer [von Glehn et al. 2023] achieves significantly lower energies by using an attention mechanism to capture all pairwise electron-electron interactions. In contrast, Moon [Gao and Günemann 2023a] replaces the global mean field of FermiNet by local mean fields on the nuclei by using continuous convolutions resulting in similar accuracy to PsiFormer. For superfluids and homogeneous electron gas, similar networks can be used but the input coordinates must be embedded with periodic functions to respect the periodicity of the problem [Pescia et al. 2022; Cassella et al. 2023]. MP-NQS [Pescia et al. 2023] models quantum states of HEG with message passing networks.

Fermion antisymmetry does not require ϕ_θ^\uparrow and ϕ_θ^\downarrow to be the same mappings. Hence, in the most general setting, they should be able to produce different computations. While traditionally choosing ϕ_θ^\uparrow and ϕ_θ^\downarrow to be different functions lead to better variational energies, for the ground state of a singlet-state system, *i.e.*, all electron spins are paired, we do have $\phi_\theta^\uparrow = \phi_\theta^\downarrow$. In the neural network setting, Gao and Günemann [2023b] has shown that implementing this constraint for singlet-state systems improves energies and accelerates optimization in neural network-based wavefunctions. However, in practice, most of the parameters of these two networks can be shared. Concretely, at each layer of the network, each electron is encoded as a feature $\mathbf{h}_i \in \mathbb{R}^d$ where d is the hidden dimension. The feature at the next layer is computed as $\mathbf{h}'_i = f(\mathbf{h}_i, \mathbf{h}^\sigma, \{\mathbf{h}_j\}_{j \neq i})$ where $\mathbf{h}^\sigma, \sigma \in \{\uparrow, \downarrow\}$ is a feature that which represents global information for different spins, which distinguishes spin-up and spin-down electrons. For example, \mathbf{h}^\uparrow and \mathbf{h}^\downarrow can be computed as the average of spin-up electron features and spin-down electron features, respectively. As a result, when two electrons with the same spins are swapped, \mathbf{h}^\uparrow and \mathbf{h}^\downarrow will not be influenced. Hence, their feature vectors will also be swapped. Otherwise, if two electrons with different spins are swapped, \mathbf{h}^\uparrow and \mathbf{h}^\downarrow will change and their feature vectors will become entirely different.

Orbital Modeling – Incorporating Physics: Recent studies show that the incorporation of physics knowledge can have an important impact on performance. Among those, wavefunction has decaying behavior at long distances, and envelope functions are used to ensure this fundamental behaviour. Essentially, the neural wavefunctions are multiplied with a mask function such that the wavefunction vanishes when electrons are far away from the nuclei. For example, FermiNet [Pfau et al. 2020] uses a simple exponential envelope. In the VMC setting, this also ensures to have a finite MCMC integration. Moreover, due to the singularities in the potential energy when two particles overlap, the wavefunctions must have discontinuous derivatives at such configurations, known as electron-electron cusp and electron-nuclear cusp. However, modeling such non-smooth behaviors is challenging for neural networks. A common way to handle electron-electron cusps is to include an

explicit divergent term in the wavefunction. For example, PauliNet [Hermann et al. 2020] handles electron-electron cusps by multiplying the wavefunction by $\exp\left(\sum_{i < j} -\frac{a_{ij}}{1 + \|\mathbf{r}_i - \mathbf{r}_j\|}\right)$, with a_{ij} being spin-dependent constants, which is part of the Jastrow factor. FermiNet [Pfau et al. 2020] proposes that the cusp conditions can be modeled by using distances as input features, e.g., $\|\mathbf{r}_i - \mathbf{r}_j\|$ since they are not differentiable when two particles overlap. For superfluids and homogeneous electron gas, periodic envelope functions can be used to improve convergence [Lou et al. 2023; Cassella et al. 2023].

Moreover, another commonly used strategy to incorporate physics is to make use of classical solutions. A commonly used strategy is to pretrain the orbital networks ϕ_θ to match the classical orbitals, such as the ones obtained with Hartree Fock methods. Although Gerard et al. [2022] shows that this is not always beneficial to pretrain with more accurate classical solutions. In contrast, PauliNet [Hermann et al. 2020] directly uses Hartree Fock solution as part of the orbital functions ϕ_θ instead of pretraining. For homogeneous electron gas, WAP-net [Wilson et al. 2023] multiplies the orbital function with Hartree-Fock plan wave orbitals, computed with transformed coordinates.

Optimization: Same to lattice systems, the neural wavefunction ψ_θ can be optimized via variational Monte Carlo (VMC). As defined in Equation (64), the objective to minimize the energy expectation, $\mathcal{L}(\mathbf{r}; \theta) = \mathbb{E}_{p_\theta}[E_{loc}(\mathbf{r}; \theta)]$, $p_\theta \propto \psi_\theta^2$. Since the expectation cannot be integrated analytically, we need to estimate the gradient from samples. One common way in machine learning to compute gradient through expectation is to use the stochastic gradient where the overall gradient is computed as the average gradient from each sample. However, the stochastic gradient cannot be used in our case because updating the parameters will also change the underlying probability distribution p_θ . To account for the distribution shift, a closed form gradient can be computed using the Hermitian property of the Hamiltonian [Ceperley et al. 1977], and is given by:

$$\nabla_\theta \mathcal{L}(\mathbf{r}; \theta) = 2\mathbb{E}_{p_\theta} \left[(E_{loc}(\mathbf{r}; \theta) - \mathbb{E}_{p_\theta}[E_{loc}(\mathbf{r}; \theta)]) \nabla_\theta \log |\psi_\theta(\mathbf{r}; \theta)| \right]. \quad (69)$$

Compared to the spin systems (Equation (57)), we omit the complex conjugate since ψ_θ is real-valued. From here we can use sample means to evaluate the expectations since we no longer need to differentiate through expectations and thus the gradient is unbiased. However, to estimate the expectations correctly, we need to generate samples following p_θ . To this end, MCMC with Metropolis-Hastings can be used to generate such samples. Given a batch of current samples \mathbf{r} (which are randomly initialized), we randomly perturb each of them with a Gaussian noise to get $\tilde{\mathbf{r}}_i = \mathbf{r}_i + \delta\mathbf{r}_i$, where $\delta\mathbf{r}_i \sim \mathcal{N}(0, \sigma)$. We then decide whether to accept the perturbed samples with Metropolis-Hastings rejection. Specifically, for each electron i , we compute the ratio $q = p_\theta(\tilde{\mathbf{r}}_i)/p_\theta(\mathbf{r}_i)$ and at the same time uniformly sample a random number a from $[0, 1]$. We then compare the value of q and a . If $q \geq a$, we keep the perturbed sample and let $\mathbf{r}_i = \tilde{\mathbf{r}}_i$. Otherwise, we reject the proposal and keep \mathbf{r}_i unchanged. We can prove that the samples generated with this procedure will converge to p_θ . σ controls how different the proposals are. Practically, we control σ so that the acceptance ratio is around 0.5.

Another important component in optimization is the choice of optimizer. Second-order optimizers such as natural gradients are found to be critical to achieve accurate optimization. Compared to first-order methods which update the parameters directly follow the reverse direction of gradients, which corresponds to the steepest direction in Euclidean space, natural gradient preconditions the gradient with the inverse of the Fisher information matrix so that the updates in the steepest direction in terms of the distribution. Concretely, the parameters are updated as $\theta \leftarrow \theta - \eta F^{-1} \nabla_\theta \mathcal{L}(\mathbf{r}; \theta)$. When dealing with unnormalized wavefunctions, the parameter update is equivalent to the stochastic reconfiguration with $F_{ij} \propto \mathbb{E}_{p_\theta} [(O_i - \mathbb{E}_{p_\theta}[O_i])(O_j - \mathbb{E}_{p_\theta}[O_j])]$ with $O_i = \frac{\partial \log |\psi_\theta(\mathbf{r})|}{\partial \theta_i}$. However, computing F^{-1} is infeasible for large models. Therefore approximations are needed. By

making certain assumptions to the Fisher matrix, KFAC [Martens and Grosse 2015] accelerates the computation by factorizing the Fisher matrix with Kronecker products. Alternatively, the conjugated gradient (CG) method can be used to approximate $F^{-1}\nabla_{\theta}\mathcal{L}(\mathbf{r}; \theta)$ [Gao and GÜNNEMANN 2021].

Besides VMC, there exist other methods to optimize neural wavefunctions. In diffusion Monte Carlo (DMC), each sample is additionally assigned with a weight such that the weighted average of sampled energies can be closer to the true ground state energy. To achieve this, the weights are computed based on imaginary time evolution. In DMC the sampling is done with Langevin dynamics, where the samples are generated following the quantum drift (or the score in machine learning), defined as $\nabla_{\mathbf{r}} \log \psi$. The process approximates the iterative application of the imaginary-time evolution operator $\psi \leftarrow e^{-\tau \hat{H}} \psi$, where τ is the evolution time. The score gives a 3D vector for each electron, which points towards the direction of higher probability density. Ren et al. [2023]; Wilson et al. [2021] first train a FermiNet with VMC and then use DMC to further approach the ground states. Moreover, DiffVMC [Zhang et al. 2023c] combines VMC and DMC by parameterizing the score directly. Instead of updating the weights for samples, DiffVMC updates the parametrized score function directly through a specially designed loss function based on score matching [Hyvärinen and Dayan 2005].

Generalized Wavefunctions for Multiple Geometries: Learning generalized Wavefunction that either covers the complete PES of a given molecule or across different compounds, faces a set of various challenges that do not apply to single structure calculations.

Multiple Geometries – Generalized Orbitals: In learning generalized wavefunctions, a key challenge is to adapt the molecular orbital function ϕ_i to the molecular structure. There have been various approaches on accomplishing this. The first work by Scherbela et al. [2022] (DeepErwin) tackles the problem by sharing most of the parameters across different structures and only retraining specific weights for each structure. Concurrently, Gao and GÜNNEMANN [2021] proposes a two-layered network approach to adapting the orbital functions called Potential Energy Surface Network (PESNet). In PESNet, the orbital functions are parametrized by another neural network that only acts on the nuclei, similar to supervised surrogate models. This avoids the need for retraining completely and has to be optimized only once to model a whole PES of a molecule. However, while the weight-sharing approach can be transferred to different sets of atoms, PESNet has no such capabilities.

When learning generalized orbitals for different compounds, *i.e.*, varying sets of nuclei, the problem of parameterizing molecular orbital functions $\{\phi_i\}_{i=1}^N$ is aggravated by the varying number of molecular orbitals N which corresponds to the number of electrons. While many weights can still be shared, one still needs to optimize the wavefunction separately for each molecule [Scherbela et al. 2022]. Two concurrent works tackle this problem and avoid the individual optimizations: Transferable Atomic Orbitals (TAOs) [Scherbela et al. 2023] and Graph-learned orbital embeddings (Globe) [Gao and GÜNNEMANN 2023a]. In TAO, the molecular orbital functions are constructed as linear combinations of atomic orbital functions $\phi_i = \sum_j a_j \phi_j$ similar to Hartree-Fock (HF) theory. In fact, TAO uses classical HF calculations to obtain the coefficients a_j . In contrast, Globe does not rely on classical HF calculations but builds on a two-layered network structure like PESNet and localizes orbitals as points in space. The parameters of the orbitals are then learned by spatial message passing similar to SchNet [Schütt et al. 2017b]. In their respective evaluations, the authors find TAO to perform better in transferring the wavefunction to new molecules as the HF calculation provides a strong inductive bias [Scherbela et al. 2023] while Globe shows strong capabilities in learning various compounds' ground states simultaneously [Gao and GÜNNEMANN 2023a].

Multiple Geometries – Symmetries: As the energy E is observable, it is invariant with respect to the Euclidean group $E(3)$. Concretely, let U_R be the unitary operator associated with the rotation matrix R , the rotated Schrödinger equation $U_R \hat{H} U_R^\dagger \psi = E\psi$ with $U_R \hat{H} U_R^\dagger = -\frac{1}{2} \sum_i \nabla_i^2 + \sum_{i < j} \frac{1}{\|r_i - r_j\|} - \sum_{i,I} \frac{z_I}{\|r_i - c_I R\|} + \sum_{I < J} \frac{1}{\|c_I - c_J\|}$ being the rotated Hamiltonian operator. From this formulation, one can see that rotating an eigenfunction ψ of \hat{H} solves the rotated Schrödinger equation, *i.e.*,

$$U_R \hat{H} U_R^\dagger U_R \psi = E U_R \psi, \quad (70)$$

$$U_R \hat{H} \psi = E U_R \psi, \quad (71)$$

$$\hat{H} \psi = E \psi. \quad (72)$$

Thus, to obtain invariant energies, one must have an equivariant wavefunction. A simple implementation would be invariant wavefunctions like PauliNet [Hermann et al. 2020] but as wavefunctions do not have to be invariant, this severely restricts the function class and typically results in higher energies. Instead, current approaches either rely on using equivariant coordinate frames that rotate with the nuclear structure [Gao and Günemann 2021; Gao et al. 2022; Gao and Günemann 2023a] or augmenting the training data by arbitrary rotations [Scherbela et al. 2023]. In addition to the Euclidean group, the PES is also invariant to the permutation group of the nuclei S_M . Integrating this symmetry is typically achieved by relying on summations over nuclei in the orbital functions rather than concatenations [Gao and Günemann 2021].

Multiple Geometries – Size-Consistency: Similar to symmetries which tell us the exact change of the wavefunction under certain actions, size consistency is prior information about the energy of the system. Specifically, size consistency refers to the change in energy depending on the size of the modeled system. In the limit case, where the system can be decomposed into two non-interacting subsystems, the energy of the whole system is simply the sum of the energies of the subsystems. Though, as it is only phrased in the limit case of non-interacting systems it cannot be phrased as a symmetry in a strict way. Nonetheless, restricting the functional form of neural wavefunctions by size consistency reduces the search space of potential function classes and results in better generalization [Gao and Günemann 2023a]. It can be shown that to implement size consistency, one needs to restrict orbital function ϕ to have decaying receptive fields such that particles do not interact with each other given sufficient distance [Gao and Günemann 2023a]. This is incompatible with the widely used FermiNet architecture which strongly relies on global averages. The Molecular orbital network (Moon) [Gao and Günemann 2023a] implements this by relying decaying spatial filters in a message-passing scheme similar to SchNet [Schütt et al. 2017b].

Multiple Geometries – Energy Surfaces: As the wavefunction is directly linked to the energy, one could for each structure solve the integral $\int \psi_\theta(\mathbf{r}, M) \hat{H}_M \psi_\theta(\mathbf{r}, M) d\mathbf{r}$ numerically to obtain the respective energy. However, due to the inherently expensive process of Monte Carlo integration, this proves costly if thousands to millions of states have to be evaluated. For single structure calculations, the final energy is often approximated as a running average over the energies observed during training, *e.g.*, the last 20% of observed training energies. However, this does not translate to the generalized setting where molecular structures are often sampled from some continuous distribution [Gao and Günemann 2021]. Thus, the structure will not be observed multiple times. Gao et al. [2022] tackles this issue by introducing Potential learning from ab-initio Networks (PlaNet). In PlaNet, one translates the idea of averaging training energies from the single-molecule setting to PES modeling by averaging the observed energy surfaces at each time step. In practice, this means that at each time step one fits a function to the observed energies. These functions are then temporally averaged in a first-order Taylor approximation by averaging the parameters of the function.

3.3.4 Datasets and Benchmarks.

Same as quantum spin systems (Section 3.2.5), the training data is sampled according to the distribution defined by the neural wavefunction. As a result, there is no need to generate a dataset beforehand. Instead, the data is defined in terms of the atom coordinates of a geometry. On the other hand, due to the variational nature of the optimization process, accuracies are evaluated by the average energy as well as the standard deviation estimated from samples, and a lower energy represents a more accurate result. Commonly tested systems are small or heavy atoms such as N or Fe, small or large molecules such as N₂ or CCl₄ [von Glehn et al. 2023], some special atomic configurations such as H₁₀, compound structures such as the Benzene dimer [Ren et al. 2023; von Glehn et al. 2023], as well as transition energies for molecular systems, defined as the difference between the ground state energies after and before a chemical process, such as the automerization of cyclobutadiene (C₄H₄) [Spencer et al. 2020] and the ionization of Fe [von Glehn et al. 2023].

3.3.5 Open Research Directions.

There are several remaining challenges for modeling many-electron systems with VMC. First, due to the fermion antisymmetry constraint, most existing methods use Slater determinants. However, optimizing through determinants could introduce extra difficulty. It is still to be seen whether we can effectively achieve fermion antisymmetry using methods other than Slater determinants. Second, currently most methods model wavefunctions explicitly. Modeling wavefunctions in real space is similar to modeling probability density in generative machine learning. Moving towards implicit modeling could be an interesting direction. Finally, one of the most pressing challenges lies in computational efficiency. As the computational complexity scales $O(N^4)$ with the number of electrons N , current calculations are limited to at most 80 electrons. it is important to further scale the methods. This could be achieved by more efficient sampling, better optimization, and enabling more effective weight sharing across systems. For example, the deep learning library Jax [Bradbury et al. 2018] has shown good accelerations for energy evaluation by improving implementations. Scaling is important to extend QMC methods to larger molecular systems or materials.

4 AI FOR DENSITY FUNCTIONAL THEORY

In this section, we first introduce the basic knowledge of density functional theory (DFT) in Section 4.1. Then in Section 4.2, we formulate the quantum tensor learning problem and describe a couple of state-of-the-art machine learning models. In Section 4.3, we review the recent progress of machine learning approaches for more accurate density functionals. We point out some promising future directions, *e.g.* quantum tensors as physics-based features for deep learning and machine learning density functionals. We summarize the structure of this section in Figure 12.

4.1 Overview

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Recommended Prerequisites: Section 3.1

In principle, modeling the quantum states of physical systems, such as molecules or materials, requires the explicit form of their wavefunctions $|\psi\rangle$ by solving the many-body Schrödinger equation. However, as the number of electrons in the system increases, the Hilbert space of the system grows exponentially, resulting in high computational and memory consumption. It becomes impractical to solve the Schrödinger equation for even small molecular systems with just tens of electrons. Therefore, for the time being, the methods in Section 3 cannot be directly used to obtain the many-body wavefunctions by solving the Schrödinger equation (Equation (52)) for large and complex systems.

4.1.1 Density Functional Theory.

In practice, first-principles density functional theory (DFT) [Hohenberg and Kohn 1964; Kohn and Sham 1965] and *ab initio* quantum chemistry methods [Szabo and Ostlund 2012] are widely used to solve the Schrödinger equation with different approximations and provide near-form wavefunctions in polynomial time complexity. In quantum chemistry methods such as Hartree-Fock theory, the total wavefunction of a system is represented by the Slater determinant of noninteracting electrons which satisfies the exact antisymmetry upon exchanging two fermionic electrons (*i.e.*, identical particles with spin 1/2). In the Hartree-Fock theory, the electron-electron Coulomb interaction and the exact exchange interaction are precisely taken into account, while the additional electron correlation energy beyond the exact exchange is not considered. Alternatively, the Hohenberg-Kohn DFT theorem [Hohenberg and Kohn 1964] shows that (HK-1) the ground-state electron density $\rho(\mathbf{r})$ (a three-dimensional quantity) uniquely determines the external potential (such as electron-nuclei interaction) and thus the Hamiltonian, and (HK-2) the ground-state total energy is a functional of electron density minimized by the ground-state density ρ_{gs} , *i.e.*, $E[\rho] \geq E[\rho_{\text{gs}}]$ for any trial/approximate density $\rho \neq \rho_{\text{gs}}$. In practice, the many-body interacting system may be mapped onto many one-body noninteracting systems within the DFT Kohn-Sham (KS) framework [Kohn and Sham 1965], where individual electrons are subject to a mean field that depends on the total electron density, as illustrated in Figure 13. The corresponding Kohn-Sham electronic total energy $E_{\text{KS}}[\rho]$ is given by

$$E_{\text{KS}}[\rho] = E_{\text{kin}}[\rho] + E_{\text{H}}[\rho] + E_{\text{ext}}[\rho] + E_{\text{xc}}[\rho], \quad (73)$$

where E_{kin} is the kinetic energy of noninteracting electrons, E_{H} is the Hartree term originating from electron-electron Coulomb interaction, E_{ext} denotes the external potential energy *e.g.* due to the interaction between electrons and nuclei, and E_{xc} denotes the exchange-correlation (XC) energy. Since the system is mapped onto a noninteracting one in the Kohn-Sham framework, the total electron density $\rho(\mathbf{r})$ can be obtained by summing over the contributions from individual

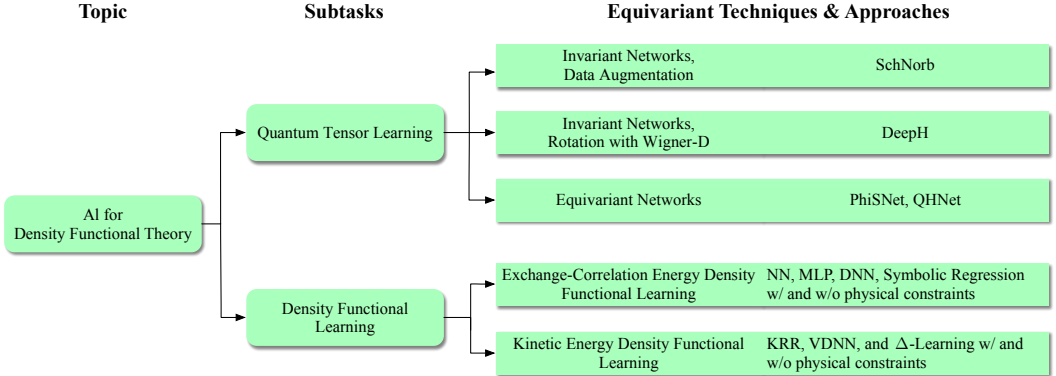


Fig. 12. An overview of tasks and methods in AI for density functional theory (DFT). In the quantum tensor learning subtask, invariant quantum tensor networks include SchNorb [Schütt et al. 2019] and DeepH [Li et al. 2022e]. SchNorb encourages the equivariance by training with data augmentation and DeepH guarantees the equivariance by rotation with Wigner D-matrix. Meanwhile, equivariant quantum tensor networks, including PhiSNet [Unke et al. 2021a] and QHNet [Yu et al. 2023c], intrinsically consider the equivariance of matrix by tensor product and tensor expansion. Another subtask is density functional learning, primarily focused on approximating exchange-correlation (XC) energy and kinetic energy. Several machine learning approaches have been employed for approximating exchange-correlation energy density functionals. These include neural network (NN) [Tozer et al. 1996; Dick and Fernandez-Serra 2019, 2020; Ryabov et al. 2020; Lei and Medford 2019; Gedeon et al. 2021], multiple layer perceptron (MLP) [Nagai et al. 2020], deep neural network (DNN) [Kirkpatrick et al. 2021; Pokharel et al. 2022], and symbolic regression [Ma et al. 2022], implemented both with and without physical constraints. In terms of approximating kinetic energy density functionals, approaches such as kernel ridge regression (KRR) [Snyder et al. 2012, 2015], voxel deep neural network (VDNN) [Ryczko et al. 2022b], and Δ-Learning [Ramakrishnan et al. 2015] have been used, again both with and without the inclusion of physical constraints. Alternatively, Ref. [Brockherde et al. 2017] aims to learn Hohenberg-Kohn map between external potential and electron density using KRR, thereby bypassing the Kohn-Sham equation.

non-interacting electrons in the orthonormal eigenstates ψ_i ,

$$\rho(\mathbf{r}) = \sum_i f_i \psi_i^*(\mathbf{r}) \psi_i(\mathbf{r}), \quad (74)$$

where f_i denotes the occupation number in state $\psi_i(\mathbf{r})$. Since the total number of electrons, N_e , is fixed for a given system, thus $\int \rho(\mathbf{r}) d\mathbf{r} = N_e$. For orthonormal eigenstates, $\langle \psi_i | \psi_j \rangle = \int \psi_i^*(\mathbf{r}) \psi_i(\mathbf{r}) d\mathbf{r} = \delta_{ij}$, hence $\sum_i f_i = N_e$. Correspondingly, the kinetic energy E_{kin} can be calculated by directly evaluating the expectation value of the kinetic energy operator $\hat{T} = -\frac{\hbar^2}{2m_e} \nabla^2$ for all occupied electronic states,

$$E_{\text{kin}}[\rho] = \sum_i f_i \langle \psi_i | \hat{T} | \psi_i \rangle = \sum_i f_i \int \psi_i^*(\mathbf{r}) \left[-\frac{\hbar^2}{2m_e} \nabla^2 \right] \psi_i(\mathbf{r}) d\mathbf{r}, \quad (75)$$

where \hbar stands for the reduced Planck's constant and m_e stands for the electron mass. Furthermore,

$$E_H[\rho] = \frac{e^2}{8\pi\epsilon_0} \iint \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} dr dr', \quad (76)$$

$$E_{\text{ext}}[\rho] = \int V_{\text{ext}}(\mathbf{r}) \rho(\mathbf{r}) d\mathbf{r}, \quad (77)$$

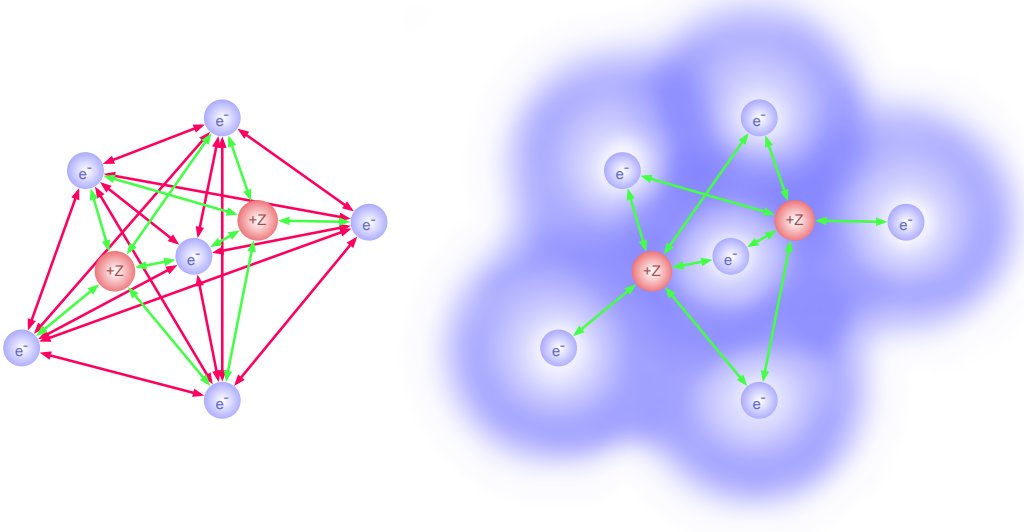


Fig. 13. An illustration contrasting the interacting many-body perspective (left) with the DFT perspective within the Kohn-Sham framework (right) for modeling electronic structure in an atomistic system (e.g., molecules). Red spheres represent nuclei and blue spheres represent electrons. Red and green edges represent interactions between electron-electron pairs and electron-nucleus pairs (e.g., via Coulomb potential), respectively. The blue shading on the right represents the electron density. **Left:** In the interacting many-body view, the wavefunction of the system is defined w.r.t. the coordinates of all particles. Hence the interactions between all electron-electron and electron-nucleus pairs are explicitly considered, leading to an exponential growth of the dimension for many-body wavefunctions with the increasing number of electrons. **Right:** In the DFT Kohn-Sham picture, the electron-electron interactions are replaced by the interaction between each electron and the average effect of all other electrons, modeled with an electron density. Since such interactions are equivalent for different electrons, modeling the electronic wavefunction of the system effectively reduces to modeling multiple single-electron wavefunctions. The lighter shading around each electron illustrates the exclusion of the electron itself in the electron density to avoid the self-interaction.

where e stands for the elementary charge, ϵ_0 stands for vacuum permittivity, and V_{ext} denotes the external potential. For the cases of molecules and materials, the external potential simply comes from the electron-nuclei Coulomb interaction,

$$V_{\text{ext}}(\mathbf{r}) = -\frac{e^2}{4\pi\epsilon_0} \sum_I \frac{Z_I}{|\mathbf{r} - \mathbf{R}_I|}, \quad (78)$$

where \mathbf{R}_I and Z_I denote the position and charge of the nuclei I , respectively. Similarly, the exchange-correlation (XC) energy $E_{\text{xc}}[\rho]$ can also be readily evaluated for a given electronic density $\rho(\mathbf{r})$.

4.1.2 The Kohn-Sham Equation.

According to the second Hohenberg-Kohn theorem, one can minimize Equation (73) by varying the electron density using the variational principle, thereby achieving the ground-state density and total energy. In practice, the method of Lagrange multipliers is applied under the constraint of total electrons N_e , where the Lagrange function is constructed as follows:

$$\mathcal{L}[|\psi_i\rangle] = E_{\text{KS}} - \sum_i \epsilon_i (\langle \psi_i | \psi_i \rangle - 1). \quad (79)$$

Thus at the stationary points of Lagrange function \mathcal{L} , we have

$$\frac{\delta \mathcal{L}}{\delta \langle \psi_i |} = H_{\text{KS}} |\psi_i\rangle - \epsilon_i |\psi_i\rangle = 0. \quad (80)$$

We then obtain the DFT Kohn-Sham equation,

$$H_{\text{KS}} |\psi_i\rangle = \epsilon_i |\psi_i\rangle, \quad (81)$$

where H_{KS} is the Kohn-Sham Hamiltonian, and ϵ_i and ψ_i are the corresponding Kohn-Sham eigen energies and eigen wavefunctions, respectively. More explicitly,

$$H_{\text{KS}} = -\frac{\hbar^2}{2m_e} \nabla^2 + V_{\text{H}}(\mathbf{r}) + V_{\text{XC}}(\mathbf{r}) + V_{\text{ext}}(\mathbf{r}), \quad (82)$$

where V_{H} is the Hartree potential as

$$V_{\text{H}}(\mathbf{r}) = \frac{e^2}{4\pi\epsilon_0} \int \frac{\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}'. \quad (83)$$

V_{XC} is the exchange-correlation potential as

$$V_{\text{XC}}(\mathbf{r}) \equiv \delta E_{\text{XC}}[\rho] / \delta \rho(\mathbf{r}) \quad (84)$$

and V_{ext} is the external potential defined above in Equation (77) for molecules and materials. Alternatively, the three potentials can be considered as an effective potential or self-consistent field (SCF) for individual electron, with $V_{\text{eff}}(\mathbf{r}) = V_{\text{SCF}}(\mathbf{r}) = V_{\text{H}}(\mathbf{r}) + V_{\text{XC}}(\mathbf{r}) + V_{\text{ext}}(\mathbf{r})$, thus

$$H_{\text{KS}} = -\frac{\hbar^2}{2m_e} \nabla^2 + V_{\text{SCF}}(\mathbf{r}). \quad (85)$$

More details of theoretical background, technical implementation, and applications of DFT can be found in many excellent books [Parr and Yang 1995; Engel and Dreizler 2011; Dreizler and Gross 2012; Fiolhais et al. 2003; Kaxiras and Joannopoulos 2019; Cohen and Louie 2016; Martin et al. 2016; Martin 2020; Koch and Holthausen 2001; Sholl and Steckel 2009; Yip 2005; Giustino 2014] and review papers [Payne et al. 1992; Kohn 1999; Kümmel and Kronik 2008; Jones 2015]. Nevertheless, with the exact exchange-correlation energy functional (though unknown by far), ground-state total energy as well as many other ground-state properties such as atomic forces and electric polarization can be derived exactly. Furthermore, although Kohn-Sham eigen wavefunctions $\psi_i(\mathbf{r})$ and eigen energies ϵ_i obtained from the DFT Kohn-Sham equation correspond to the non-interacting fictitious system, it turns out that electronic structure such as band structure, the density of states, etc. are fairly well described for many materials and compounds in practice, except strongly correlated materials, etc. It is therefore highly desirable to (1) develop machine learning approaches to predict full quantum mechanical Hamiltonian for arbitrary materials and molecules with arbitrary structures since it determines the underlying physical and chemical properties, and (2) develop more accurate density functionals beyond the state-of-the-art using machine learning methods under the known physical constraints.

4.1.3 Density Functional Theory In Practice.

Although DFT significantly simplifies the way of solving the many-body Schrödinger equation, the exact numerical solution of the Kohn-Sham equation in Equation (81) in principle needs infinite spatial grids to represent the exact wavefunctions. To avoid modeling the infinite spatial space, a set of predefined basis functions $\{\phi_j\}$ is introduced to approximate the single electronic wavefunctions. Commonly used basis sets include plane-wave basis, real space grids, wavelets, and localized atomic basis sets such as Slater-Type Orbitals (STO) [Slater 1930], Gaussian-Type Orbitals (GTO) [Boys and Egerton 1950], and Numerical Atomic Orbitals (NAO) [Koepernik and Eschrig 1999; Junquera

et al. 2001]. Here we focus on the localized atomic orbital basis functions $\{\phi_j\}$ with an analytical form which are often represented by the product of a radial function and spherical harmonics. Specifically, it is approximated using a linear combination of basis functions defined as

$$\psi_i(\mathbf{r}) = \sum_j c_{ij} \phi_j(\mathbf{r}), \quad (86)$$

where c_{ij} is the j -th coefficient of the electronic wavefunction ψ_i associated with basis function ϕ_j , forming wavefunction coefficient matrix C_e . C_e can be obtained by solving the DFT Kohn-Sham equation in the matrix form as a generalized eigenvalue problem,

$$HC_e = \epsilon SC_e, \quad (87)$$

where $H := H_{\text{DFT}} = H_{\text{KS}}$ (Equation (82)) is Hamiltonian with $h_{ij} \equiv \langle \phi_i | H | \phi_j \rangle = \int \phi_i^*(\mathbf{r}) H \phi_j(\mathbf{r}) d\mathbf{r}$, incorporating the interactions of different particles, $S \in \mathbb{R}^{N_o \times N_o}$ is the overlap matrix with $s_{ij} \equiv \langle \phi_i | \phi_j \rangle = \int \phi_i^*(\mathbf{r}) \phi_j(\mathbf{r}) d\mathbf{r}$, representing the integral of a pair of predefined orbital basis, and ϵ is a diagonal matrix where each diagonal element ϵ_{ii} represents the eigen energy for the corresponding eigen wavefunction ψ_i . Depending on the nature of the system, C_e and H may be $\in \mathbb{R}^{N_o \times N_o}$ or $\mathbb{C}^{N_o \times N_o}$, where N_o is the number of orbitals, and each atom may have multiple associated orbitals.

As discussed above, by mapping many-body interacting systems onto many one-body non-interacting systems using the Kohn-Sham approach [Kohn and Sham 1965], it is possible to compute electronic charge density using the single-particle electronic wavefunctions represented by wavefunction coefficients C_e and basis functions $\{\phi_j\}$. Consequently, the Hamiltonian matrix H_{KS} can be determined. To find C_e the solution of the Kohn-Sham equation (Equation (87)), the SCF algorithm [Payne et al. 1992; Cances and Le Bris 2000; Kudin et al. 2002] is commonly applied to improve the solutions C_e iteratively until the convergence is reached. When there are N_T steps in total, the time complexity of the DFT algorithm is $O(N_o^3 N_T)$, with each step being $O(N_o^3)$. However, running iterative SCF calculations for large systems is computationally expensive. To address this issue, deep learning models have been proposed to consider the interactions among the atoms and directly predict the Hamiltonian matrix. As shown in Figure 14, the final quantum tensors commonly obtained by self-consistent DFT calculations, such as Hamiltonian matrix, can be predicted by quantum tensor networks, which is discussed in detail in Section 4.2.

According to [Hohenberg and Kohn 1964] and [Kohn and Sham 1965], the ground-state total energy in Equation (73) is exact for many-body system if we have the exact exchange-correlation energy functional $E_{\text{XC}}[\rho]$. This can be more explicitly seen by re-writing the ground-state electronic total energy in Equation (73) using the Kohn-Sham eigen energies from the Kohn-Sham equation in Equations (81) or (87),

$$E_{\text{KS}} = \sum_i f_i \epsilon_i + E_{\text{XC}}[\rho] - \frac{e^2}{8\pi\epsilon_0} \iint \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r} d\mathbf{r}' - \int V_{\text{ext}}(\mathbf{r})\rho(\mathbf{r}) d\mathbf{r}. \quad (88)$$

The ground-state density $\rho(\mathbf{r})$ uniquely determines the Hamiltonian H_{KS} , thus determines Kohn-Sham eigen energies ϵ_i . Subsequently, the second, third, and last terms of the above Kohn-Sham total energy are completely decided. While eigen energies ϵ_i of the first term and the Hartree energy of the third term are easy to evaluate, the key challenge of the Kohn-Sham DFT lies in the unknown exchange-correlation energy functional $E_{\text{XC}}[\rho]$ and the corresponding exchange-correlation potential $V_{\text{XC}}(\mathbf{r}) \equiv \delta E_{\text{XC}}[\rho]/\delta\rho(\mathbf{r})$. Two main categories of XC energy functionals have been developed in the past, including the Local Density Approximation (LDA) [Ceperley and Alder 1980; Vosko et al. 1980; Perdew and Zunger 1981; Perdew and Wang 1992] where the XC energy depends on the local electron density $\rho(\mathbf{r})$ only, and Generalized Gradient Approximation (GGA) [Perdew et al. 1996, 1992; Becke 1988; Lee et al. 1988] where the XC energy depends on both

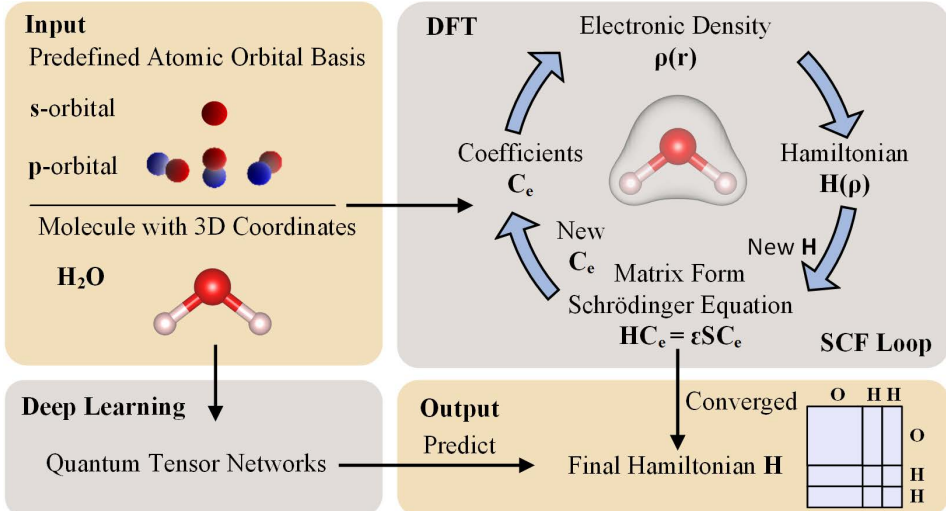


Fig. 14. The pipeline of the DFT calculations and deep learning methods to obtain the Hamiltonian matrix. The DFT calculation uses the predefined atomic orbital basis associated with the molecule and its coordinates to optimize Hamiltonian matrix $H(\rho)$ iteratively within the SCF loop until it reaches convergence towards the total energy minimum/minima. In contrast, deep learning method uses the proposed quantum tensor networks to directly predict the final Hamiltonian matrix, taking atomic types and coordinates as inputs. This eliminates the iterative optimization, thereby accelerating the DFT calculations.

the local electron density $\rho(\mathbf{r})$ and its gradient $\nabla\rho(\mathbf{r})$. In addition, hybrid XC functionals [Becke 1993; Heyd et al. 2003] have been proposed and widely used to (partially) include exact exchange, and meta-GGA functionals [Tao et al. 2003; Sun et al. 2015, 2016; Furness et al. 2020] have been proposed to include high-order gradients of electron density, kinetic energy density, etc. These methods gradually climb Jacob's ladder [Perdew and Schmidt 2001] with better accuracy at the price of higher computational cost. While all these approximations rely on the physical constraints and intuitions with great success in the fundamental materials and molecular research, the exact XC energy functional has not been achieved yet, presenting a unique opportunity for AI/ML approaches to tackle this challenge. We briefly summarize the recent progress in learning density functionals in Section 4.3 and discuss potential future directions in this area in Section 4.3.4.

4.2 Quantum Tensor Learning

Authors: Haiyang Yu, Zhao Xu, Limei Wang, Yaochen Xie, Xiaofeng Qian, Shuiwang Ji

In DFT calculations, quantum tensors, such as Hamiltonian matrix H and wavefunction coefficients C_e , can describe quantum states of physical systems and determine various critical physical properties, including total energy, charge density, electric polarization, etc. To accelerate the DFT calculations, various deep learning models [Schütt et al. 2019; Luya 2020; Unke et al. 2021a; Li et al. 2022e; Gong et al. 2023; Li et al. 2023e] have been proposed to directly predict the quantum tensors. The predicted quantum tensors are used to derive the physical properties at a reasonable level of accuracy, thereby accelerating the optimization process in the electronic structure calculations.

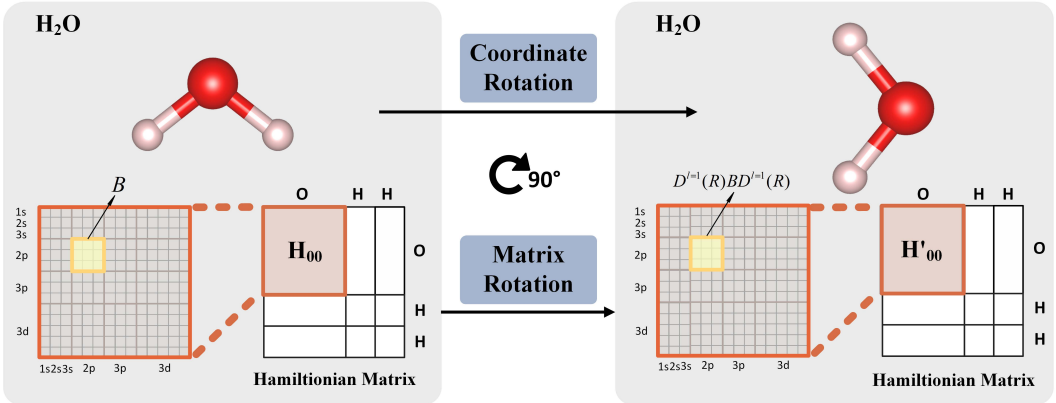


Fig. 15. The equivariance of Hamiltonian matrix \mathbf{H} . When the coordinates of the molecule are rotated using a rotation matrix R , the corresponding Hamiltonian matrix is rotated accordingly. Specifically, the Hamiltonian block B represents the orbital interaction between the oxygen $2p$ orbitals, and it is rotated to $D^{\ell=1}(R)BD^{\ell=1}(R)$ using Wigner D-matrix $D^{\ell}(R)$.

4.2.1 Problem Setup.

In this section, we focus on the task of predicting the Hamiltonian matrix, which is the key quantum tensors in accelerating the DFT algorithm. We denote the input 3D molecule as $\mathbf{M} = (\mathbf{z}, \mathbf{C})$ consisting of atom types $\mathbf{z} = (z_1, \dots, z_n) \in \mathbb{Z}^n$ and atom coordinates $\mathbf{C} = [c_1, \dots, c_n] \in \mathbb{R}^{3 \times n}$, where n is the number of atoms. We aim to develop deep learning models to predict target quantum tensors for input molecular geometries. Specifically, the Hamiltonian matrix $\mathbf{H} \in \mathbb{R}^{N_o \times N_o}$ is one of the prediction targets that can be used to derive various physical properties, where N_o represents the number of electronic orbitals.

4.2.2 Technical Challenges.

There are several challenges to tackle for quantum tensor learning. The first challenge is symmetry. Quantum tensor learning requires geometric deep learning models to guarantee the intrinsic permutation, translation, and rotation equivariance for quantum tensors. While geometric deep learning models usually maintain equivariant features, the predicted quantum tensors are generally composed of equivariant matrices. As shown in Figure 15, when the input molecule is rotated, the block B divided by orbitals in the Hamiltonian matrix is rotated to $D^{\ell_1}(R)BD^{\ell_2}(R)$, where $D^{\ell}(R)$ is the Wigner D-matrix for rotation R with rotation order ℓ . This raises the need to design equivariant architectures that build equivariant matrices from equivariant features. Another challenge is the flexibility of the model. Since the size of quantum tensors varies significantly with the chemical elements in the system, a flexible architecture is required to apply geometric deep learning models to diverse systems. Moreover, efficiency is also a challenge for equivariant networks. To maintain the equivariance, the operations in these models usually have a considerable computation cost compared to invariant networks.

4.2.3 Existing Methods.

Currently, several graph neural networks are proposed to learn the interactions among the atoms and predict the quantum tensor like SchNorb [Schütt et al. 2019], PhiSNet [Unke et al. 2021a], DeepH [Li et al. 2022e] and QHNet [Yu et al. 2023c]. They are composed of three parts, including establishing node-wise interaction, building pairwise features, and constructing the quantum matrix.

The node-wise interaction module encodes the atomic and geometric information between atoms, and builds the node-wise equivariant features using a message passing schema [Gilmer et al. 2017]. In addition, since the final Hamiltonian matrix is constructed with blocks representing the pairwise interaction of atoms, pairwise features are trained to learn such interactions. Finally, the matrix building module expands the pairwise features to matrices and then constructs the final quantum tensors corresponding to the atomic orbitals of the input atoms.

Node-Wise Interaction: The node-wise interaction module is used to construct representations of atoms by aggregating information from neighbors following the Message Passing Neural Networks (MPNNs) framework [Gilmer et al. 2017]. Specifically, the features \mathbf{h} of each node i in layer t are updated based on

$$\begin{aligned}\mathbf{m}_i^{t+1} &= \sum_{j \in \mathcal{N}(i)} M_t(\mathbf{h}_i^t, \mathbf{h}_j^t, \mathbf{h}_{ij}), \\ \mathbf{h}_i^{t+1} &= U_t(\mathbf{h}_i^t, \mathbf{m}_i^{t+1}).\end{aligned}\tag{89}$$

Here $\mathcal{N}(i)$ is the neighboring node set of node i , \mathbf{h}_{ij} is the edge feature between node i and node j , \mathbf{m}^{t+1} is the hidden variable at node i , and U_t and M_t denote the update and message functions at layer t . Note that the final prediction target, such as the Hamiltonian matrix, is an equivariant matrix, *i.e.*, if the input molecule is rotated by a rotation matrix R , each block B_{ij} of the Hamiltonian matrix \mathbf{H} should be transformed to $D^{\ell_i}(R)B_{ij}D^{\ell_j}(R)$ accordingly, as shown in Figure 15. Here $D^\ell(R) \in \mathbb{C}^{(2\ell+1) \times (2\ell+1)}$ is Wigner D-matrix of R . Therefore, it is crucial to ensure equivariance while constructing node features. One approach is to first construct invariant node features, followed by additional operations in the pairwise feature building and matrix construction steps to ensure or encourage equivariance. For example, SchNorb and DeepH construct invariant node features by aggregating the features and distances of neighboring nodes. Since the initial node features and distances are $SE(3)$ -invariant, the constructed node features are also invariant. Alternatively, another approach focuses on constructing equivariant node features directly. For example, PhiSNet and QHNet construct equivariant node features using spherical harmonics and tensor products, as introduced in Section 2.4, in each message passing layer, which ensures their equivariance to continuous symmetry transformations. Specifically, the features of each node are obtained by aggregating the tensor product between the features of each neighboring node and an equivariant filter. The filter depends on the spherical harmonics of the direction vector. With such operations, the methods can ensure equivariance at each layer. It is worth noting that the computational cost of a tensor product is significantly larger than a linear layer due to the need to multiply node features with CG matrix for each path. QHNet is much more efficient than PhiSNet by reducing the number of tensor products in the network.

Building Pairwise Features: Since the quantum matrix encodes interactions between atom pairs, it is critical to construct pairwise feature vectors for atom pairs. A typical approach is to process diagonal pairwise features f_{ii} and non-diagonal pairwise features f_{ij} separately, as they correspond to pairs of single or two atoms. However, it's also possible to process diagonal and non-diagonal pairs in the same way. Furthermore, pairwise features can be either invariant or equivariant to rotation, depending on how edge orientation is used. In DeepH, edge features are updated in its local coordinate message passing (LCMP) layer and then serve as pairwise features for atom pairs that are connected by these edges. Since self-loops are added in advance for diagonal atoms, LCMP can output both diagonal and non-diagonal features. Here, the obtained pairwise features are invariant to rotation ($l = 0$) because edge orientations have been converted to local coordinates before inputting to the LCMP layer. SchNorb computes invariant scalar for each edge and uses it to scale edge orientation in global coordinates. Hence, scaled edge features are equivariant and

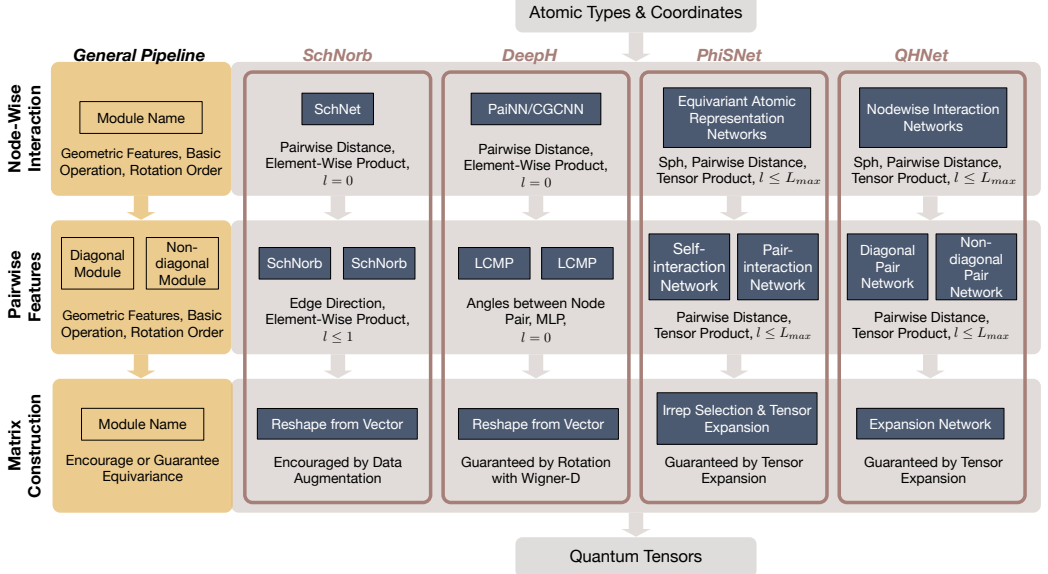


Fig. 16. An overview of quantum tensor networks and methods in AI for density functional theory. Quantum tensor networks take the atomic types and coordinates of a given molecule as input and output the predicted quantum tensor, such as the Hamiltonian matrix. Typically, quantum tensor networks consist of three sequential modules: the node-wise interaction module, pairwise feature building module, and matrix construction module. The node-wise interaction module updates the node features based on neighboring nodes within a cutoff distance. The pairwise feature module creates features to describe the relationship between atom pairs, with a diagonal module for pairs of a single atom and a non-diagonal module for pairs of two atoms. Pairwise features are then used to construct target matrices for node pairs, which are assembled to output the quantum tensor for the entire molecule. In node-wise interaction and pairwise feature modules, existing methods use different geometric features, basic operations, and rotation orders in their designs. Note that Sph denotes the spherical harmonics of edge direction. In the matrix construction module, the equivariance of the constructed matrix is either encouraged or guaranteed through various techniques. This figure provides module names and essential information about each module for existing methods, including SchNorb [Schütt et al. 2019], DeepH [Li et al. 2022e], PhiSNet [Unke et al. 2021a], and QHNet [Yu et al. 2023c].

of rotation order $l \leq 1$. Without self-loops, diagonal and non-diagonal pairwise features are then obtained from scaled edge features in different manners. Unlike the above two methods, QHNet and PhiSNet employ tensor products in their pairwise interaction modules, which leads to equivariant pairwise features of order $l \leq L_{max}$. Specifically, QHNet considers attentive scores of two atoms for non-diagonal pairs and uses tensor products for both diagonal and non-diagonal pairs. In contrast, PhiSNet uses the tensor product only for non-diagonal pairs and builds diagonal pairwise features like regular message passing. Corresponding module names, rotation orders, and key elements used for building pairwise features in the above methods are summarized in Figure 16.

Matrix Construction and Equivariance: After obtaining pairwise features, the final step is to build the quantum matrix, such as the Hamiltonian matrix. The molecular quantum matrix is composed of multiple pairwise blocks containing interactions between atoms. Here, the pairwise block is denoted by B_{ij} , where o_i and o_j are the numbers of orbitals of atom i and j , respectively. Depending on the nature of physical properties and systems, $B_{ij} \in \mathbb{R}^{o_i \times o_j}$ or $\mathbb{C}^{o_i \times o_j}$. Since pairwise features are vectors, we must convert them into block matrices. SchNorb and DeepH reshape the

flattened edge feature vector directly into a matrix, while QTNet and PhiSNet use tensor expansion to expand a single irrep vector into the matrix. Note that different atoms have varying numbers of orbitals. Thus, pairwise blocks B_{ij} are in different shapes, but outputting blocks with various shapes is challenging. To build blocks with various shapes, SchNorb, DeepH, and QTNet firstly construct immediate blocks with full orbitals and then extract exact blocks according to atom types. In contrast, PhiSNet maintains a record to guide which channel of irrep should be selected to build the block for each pair. Finally, blocks B_{ij} are assembled to construct the whole quantum matrix for the molecular system. Among the above-summarized methods, QTNet and PhiSNet adopt tensor expansion to ensure rotation equivariance of the quantum matrix. DeepH applies the inverse of the Wigner D-matrix to convert the predicted local quantum matrix back to global coordinates, thereby ensuring its rotation equivariance. SchNorb cannot guarantee the rotation equivariance, but it augments data via rotation to encourage the encoding of rotational symmetry. Figure 16 summarizes how the above methods construct the matrix and maintain matrix equivariance.

4.2.4 Datasets and Benchmarks.

For the quantum tensor learning task, MD17 [Schütt et al. 2019; Luya 2020] provides Hamiltonian matrices for the molecular geometries in the trajectories, with each trajectory corresponding to a single molecule. MD17 consists of 4 molecules: water, ethanol, Malondialdehyde, and Uracil. Furthermore, mixed MD17 [Yu et al. 2023c] combine four molecular trajectories in the MD17 dataset together and provide a quantum tensor dataset with multiple molecules in the training and testing sets.

However, the whole MD17 dataset contains only four distinct molecules. To enhance the generalization ability to the molecule space, Quantum Hamiltonian (QH9) dataset [Yu et al. 2023b] is a dataset of precise Hamiltonian matrices of molecular geometries. The open-source software PySCF (Python-based simulations of chemistry framework) [Sun et al. 2018] is used to compute the Hamiltonian matrix. QH9 consists of two kinds of datasets: static and dynamic.

- The QH-stable dataset consists of 130,831 stable molecular geometries, coming from a subset of the QM9 dataset [Ramakrishnan et al. 2014]. To explore the performance in both in-distribution and out-of-distribution (OOD) scenarios, two tasks are created on the dataset: (1) random split creates the QH-stable-iid; (2) split based on the number of constituting atoms would yield QH-stable-ood.
- The QH-dynamic dataset contains 2,399 molecular dynamics trajectories, where each trajectory has 60 geometries. Two split strategies on QH-dynamic yield two tasks: (1) QH-dynamic-mol splits training/validation/test set based on different molecules; (2) QH-dynamic-geo allows different geometries of the same molecule in training, validation, and test set.

The statistics of these datasets are shown in Table 4. Based on the curated QH9 dataset, Yu et al. [2023b] also demonstrates that the state-of-the-art method QHNet [Yu et al. 2023c] is capable of predicting Hamiltonian matrices for molecules of any kind and generalized well on unseen molecules. Specifically, QHNet trained on QH9 reached an MAE (Mean Absolute Error, between predicted Hamiltonian matrix and groundtruth) of $83.12 \times 10^{-6} E_h$ on the mixed MD17 dataset.

4.2.5 Open Research Directions.

Quantum tensors from DFT, such as Hamiltonian matrix H and density matrix D , are not only useful for computing the electronic structures, but also can serve as physics-based features to predict accurate molecular properties such as total energy and electronic polarization. The reason why PhiSNet, SchNorb, DeepH, QHNet/QTNet, etc. with small ℓ cutoff work so well for molecules and materials lies in the fact that most of these compounds are dominated by low-energy chemistry/physics where the eigen wavefunctions possess significant atomic-orbital like characteristics.

Table 4. Statistics of datasets for quantum tensor learning, including MD17 [Chmiela et al. 2017], mixed MD17 [Yu et al. 2023c], QH9 [Yu et al. 2023b].

Datasets	# geometries	# molecules	# training	# validation	# test
MD17-water	4,900	1	500	500	3900
MD17-ethanol	30,000	1	25,000	500	4,500
MD17-Malondialdehyde	26,978	1	25,000	500	1,478
MD17-Uracil	30,000	1	25,000	500	4,500
mixed MD17	91,878	4	75,500	2,000	14,378
QH-stable-iid	130,831	130,831	104,664	13,083	13,084
QH-stable-ood	130,831	130,831	104,001	17,495	9,335
QH-dynamic-geo	143,940	2,399	119,950	11,995	11,995
QH-dynamic-mol	143,940	2,399	115,140	14,340	14,460

For the same reason, one can directly construct atomic-like maximally localized Wannier functions (MLWFs) [Marzari and Vanderbilt 1997; Souza et al. 2001; Marzari et al. 2012] and quasi-atomic orbitals (QOs) [Qian et al. 2008, 2010] from molecular orbitals or eigen wavefunctions and obtain accurate Hamiltonians for the low-energy regime (e.g., from the lowest eigen energies to a few eVs above the Fermi level), which has enabled the discovery of novel materials and physics such as quantum spin Hall insulators [Qian et al. 2014; Marrazzo et al. 2018], Berry curvature memory effect [Wang and Qian 2019; Xiao et al. 2020], and nonlinear photocurrent [Wang and Qian 2020]. Accurate prediction of quantum tensors such as Hamiltonian will be highly crucial and valuable for accelerating the materials discovery in future, as recently demonstrated by DeepH [Li et al. 2022e; Gong et al. 2023; Li et al. 2023e]. Furthermore, existing work OrbNet [Qiao et al. 2020] employs the quantum tensors as node and edge features in the orbital graphs, resulting in a significant enhancement of molecular energy prediction performance. Due to the significant time and computational cost associated with DFT calculations, OrbNet uses GNF-xTB [Grimme 2013; Grimme and Bannwarth 2016], a fast semi-empirical method, to approximate the quantum tensors. The quality of these approximated quantum tensors directly influences the performance of deep learning models. Therefore, it is crucial to address the challenge of obtaining accurate quantum tensors within a reasonable time to build the physical-based input features [Bai et al. 2022]. Quantum tensor networks have the potential to address this challenge by accurately predicting quantum tensors to construct physics-based features for deep learning models, such as accurate deep learning force field for studying both electronic and structural phase transition in quantum materials [Li et al. 2021e].

4.3 Density Functional Learning

Authors: Alex Strasser, Xiaofeng Qian

Machine learning has been applied to model density functionals in order to predict the exchange-correlation energy functional, kinetic energy functional [Snyder et al. 2012] for orbital-free DFT, the universal functional, corrections to density functional, and more. The approaches vary widely, with most being numerical as well as symbolic approach [Ma et al. 2022]. Some made predictions starting from scratch, some start from the previously established functionals [Zheng et al. 2004], or from both [Ma et al. 2022], and other approaches incorporate exact physical constraints into the functional form [Hollingsworth et al. 2018; Pokharel et al. 2022; Nagai et al. 2022; Dick and

Fernandez-Serra 2021; Gedeon et al. 2021]. More details can be found in the recent perspectives and review articles [Burke 2012; Manzhos 2020; Kalita et al. 2021; Perdew 2021; Pederson et al. 2022; Fiedler et al. 2022; Kulik et al. 2022; Nagai and Akashi 2023].

4.3.1 Machine Learning Exchange-Correlation Energy Functionals.

The exchange-correlation (XC) energy (the last term in Equation (73)) is the most challenging part of the DFT-KS equation. Most widely used XC energy functionals are designed in analytical forms and fitted to various sets of known physical constraints, such as LDA and GGA mentioned above. Many of the calculations in applying ML to XC energy functionals implement a well-known quantum chemistry software called PySCF (The Python-based Simulations of Chemistry Framework) [Sun et al. 2018], e.g., Nagai et al. [2020, 2022]; Kirkpatrick et al. [2021]; Bystrom and Kozinsky [2022]. While great improvement has been demonstrated by incorporating more exact constraints [Tao et al. 2003; Sun et al. 2015, 2016; Furness et al. 2020], they are still approximations of the unknown exact XC energy functional. Machine learning can be used to enhance the accuracy of XC functionals, which are the primary source of error in typical KS-DFT calculations [Kim et al. 2013; Crisostomo et al. 2023], so we summarize some of the progress in this area.

The application of ML to XC functionals started with Tozer et al. [1996], which used a neural network to approximate the XC potential using the ZMP density inversion method [Zhao et al. 1994], resulting in geometries comparable to LDA and substantially more accurate eigenvalues. Another early study [Zheng et al. 2004] improves the widely-used B3LYP functional, using a neural network to optimize the three parameters used to determine the relative contributions of exact exchange functional, local spin density exchange functional, Becke88 exchange functional, as well as the LYP and VMN correlation energy functionals.

Recently, thanks to the rapid development of deep learning methods and their surprising capability to capture nonlinear patterns, data-driven approaches have been used to estimate the precise exchange-correlation energy functional [Bogojeski et al. 2020; Nagai et al. 2020; Dick and Fernandez-Serra 2021; Kirkpatrick et al. 2021; Bystrom and Kozinsky 2022; Trepte and Voss 2022; Sparrow et al. 2022; Bystrom and Kozinsky 2023; Dick and Fernandez-Serra 2019, 2020; Ryabov et al. 2020; Lei and Medford 2019], which demonstrates exceptional accuracy on main-group chemistry and represents a state-of-the-art achievement in the field. Unlike approximation techniques, data-driven approaches can learn a theoretically unbiased (exact) estimator of the XC energy functional from real data because they do not impose any approximations on the functional form. Specifically, to learn the exact XC energy functional from data, Nagai et al. [2020] builds a multiple layer perceptron (MLP) to learn the mapping from local density descriptor (human-curated feature) to local XC potential functional. However, this early attempt is an end-to-end paradigm, purely learning from data, and does not consider any physical constraints on the DFT system. It typically requires a large number of data points to reach desirable performance.

Since XC functionals that include exact constraints tend to have more predictive power and are more generalizable [Kaplan et al. 2023], identifying ways of incorporating these constraints into an ML-based XC functional is important. The two general approaches of imposing these exact constraints on an ML XC functional are 1) analytical, which guarantees adherence, and 2) data-driven, which primarily comes from training the ML model on data that obeys those constraints, such as data produced using the SCAN functional, and will not guarantee perfect adherence. Of the 17 known exact constraints for a semi-local XC functional, Pokharel et al. [2022] argues that six of these constraints would be conducive to an analytical application in ML models through post-processing steps, input restrictions, or choosing separate exchange and correlation models. These constraints include (for the exchange energy) negativity, spin-scaling, uniform density scaling, a

tight bound for two-electron densities, (for the correlation energy) non-positivity, and (for exchange and correlation together) the general Lieb-Oxford bound.

Several works combine data-driven XC energy fitting with exact physical constraints, including fractional electron constraint [Kirkpatrick et al. 2021], linear/nonlinear constraints [Sparrow et al. 2022], physical asymptotic constraints [Nagai et al. 2022], and others [Trepte and Voss 2022; Brown et al. 2021]. For example, one essential constraint of the DFT system is that electrons are treated as a continuous charge density rather than discrete particles. However, the continuous XC functionals cannot handle the derivative discontinuity of the XC energy at integer-electron numbers [Perdew et al. 1982; Perdew and Levy 1983]. To meet this physical constraint and address DFT's problematic delocalization error [Bryenton et al. 2023], Kirkpatrick et al. [2021] defines a fictitious system to enable fractional charge and spin, takes local features of electron density and trains a neural network to estimate the local energies, which are aggregated to obtain the XC energy. The resulting functional, DeepMind21 (DM21), demonstrates excellent performance on a bond-breaking dataset as well as across the QM9 [Ramakrishnan et al. 2014] and GMTKN55 [Goerigk et al. 2017] databases, superior to the three best hybrid functionals tested and reproducing multiple disassociation curves [Kirkpatrick et al. 2021]. Gedeon et al. [2021] proposes an approach to train a N_e neural network with a piece-wise linearity which reproduces the derivative discontinuity of the XC energy. Similarly, Sparrow et al. [2022] designs a novel set of bell-shaped spline functions as the basis to embed the linear and nonlinear constraints as well as incorporate the implicit smoothness constraint as a regularization term in the learning objective. One group examined the effects of imposing a spin-scaling constraint and the general Lieb-Oxford bound when attempting to reproduce the SCAN functional in a deep neural network, showing improvements from the constraints but limited generalizability when attempting to move from data without chemical bonding to chemically bound systems [Pokharel et al. 2022]. Furthermore, to satisfy physical asymptotic constraints, Nagai et al. [2022] breaks down the XC energy functional into different terms (e.g., spin-up exchange, spin-down exchange energy, correlation energy), analytically imposes asymptotic constraints on different neural modules and aggregates all the NNs' output. In total, they analytically imposed 10 constraints – 5 for the exchange part and 5 for the correlation part, and the physical constraints on the neural network enabled convergence in cases where the unconstrained NN did not converge. The application of the constraints was made easier by using the SCAN XC functional as a base, but they provide a way of imposing the same constraints for other base XC functionals. Bystrom and Kozinsky [2022, 2023] approximate the exchange energy functional using a nonparametric estimation that measures the similarity between the current data and existing data points using a Gaussian process model [Rasmussen and Williams 2006], while imposing the uniform scaling constraint. The resulting significant accuracy improvement when testing on the Minnesota Database 2015B [Yu et al. 2016a] and also shows promising generalizability to solid-state systems. Finally, Dick and Fernandez-Serra [2021] impose a local Lieb-Oxford bound, finding this constraint to aid generalizability along with the uniform scaling, spin-scaling, and non-negativity. Trained on the SCAN results of 21 molecules, the neural network was tested on the diet-GMTKN55 dataset [Gould 2018] and was competitive with SCAN and hybrid functionals.

Another development addresses the issue of only using the converged energies and densities to train a model by allowing information about each iteration of the self-consistent KS solution to backpropagate through a deep neural network – a Kohn-Sham regularizer [Li et al. 2021a]. With this extra data while training on only two exact energies and densities in a 1D H₂ dissociation curve, the model was able to achieve chemical accuracy for the entire dissociation curve. The authors later extend this model to include spin-density for spin-polarized systems and test on weakly correlated systems, the domain of standard DFT calculations [Kalita et al. 2022]. They find that incorporating spin-density while training on energies and densities on atoms substantially reduces the error and

improves convergence for equilibrium molecules, approaching chemical accuracy. As another way to go beyond converged energies in training, [Dick and Fernandez-Serra \[2021\]](#) assign an explicit function of iteration number in the loss function in order to penalize the slow convergence, leading to a smooth functional without convergence issues. The use of automatic differentiation enabled the extraction of more information contained in the electron density, thereby further expanding the training inputs. The same authors developed a metric for assessing XC functionals based on both energy and density errors since both are approximated in DFT calculations.

4.3.2 Machine Learning Kinetic Energy Functionals.

Rather than learning the XC functional, a different approach is to learn a density functional for the kinetic energy (KE), that is, KEDF. Compared to the KS approach, where the KE operator acts on the KS orbitals, data-driven machine learning KEDF instead allows one to neglect the KS orbitals entirely, resulting in the so-called orbital-free DFT (OF-DFT). While KS-DFT scales with $O(N_e^3)$, OF-DFT scales quasi-linearly. One difficulty with this approach is that the gradient descent method used to find the energy minimum requires an accurate gradient, but the gradient of the KE functional is noisy and not well-behaved. The functional derivative of KE arises from varying Equation (73) with respect to electron density, but retaining a general form of KE rather than the quantum mechanical KE operator as in Equation (75). In OF-DFT, the focus is approximating the KE functional, such as the very first DFT method using the Thomas-Fermi model. The KE functional derivative is used explicitly in the Euler-Lagrange Equation (90) in order to find the self-consistent density, given as

$$\frac{\delta T_s}{\delta \rho(\mathbf{r})} = \mu - V_{\text{eff}}(\mathbf{r}), \quad (90)$$

where T_s is the non-interacting KS kinetic energy, V_{eff} is the effective or KS potential, and μ is the chemical potential which ensures the constraint of total N_e electrons. A substantial amount of work has been done to address the noisy functional derivative problem, such as the development of nonlinear gradient denoising [[Snyder et al. 2015](#)]. One reason this is relevant is that the kinetic energy is in the same order as the total energy and is significantly larger than the XC energy, so an inaccurate approximation of the KE has a much larger impact than an inaccuracy in an XC approximation.

In the first pioneering work to apply ML for KE density functional approximation [[Snyder et al. 2012](#)], a kernel ridge regression was used to approximate a KE density functional, achieving chemical accuracy (mean absolute error below 1 kcal/mol) in a 1D analog to OF-DFT of noninteracting fermions in a 1D box. The same approach has been explored in more detail [[Li et al. 2016c](#)] and also applied to bond breaking for various 1D diatomic molecules, achieving chemical accuracy with 20 training data points, much better than usual OF-DFT errors [[Snyder et al. 2013](#)]. On ten atoms ranging from H to Ne and 19 molecules, [Seino et al. \[2018\]](#) uses density and its gradients up to the third order as explanatory variables in a neural network, demonstrating superiority over the majority of the other 27 semi-local KE density functionals in comparison. [Golub and Manzhos \[2019\]](#) show that using up to a fourth-order term in a gradient expansion of the kinetic energy density as an input into a neural network allows OF-DFT to reproduce the KS kinetic energy density very closely, for both solid state and molecular systems. Rather than using the density gradient directly, [Yao and Parkhill \[2016\]](#) show that the reduced density gradient, a dimensionless quantity, may be more informative as a neural network input, demonstrating strong predictive power of a convolutional neural network for seven alkanes with better performance than other GGAs at hydrocarbon bonding. One recent work [[Ryczko et al. 2022a](#)] uses slices of electron density in a voxel deep neural network (VDNN) to predict the kinetic energy of a graphene lattice within

chemical accuracy, and they also showed that one can use Monte Carlo-based optimization instead of gradient-based optimization for a 1D model system.

Just as in the XC case, KE functionals have exact constraints that must be applied to find physically motivated KE functionals, such as Pauli positivity, asymptotic limit, and coordinate scaling [Levy and Ou-Yang 1988; Holas and March 1995; Aldossari et al. 2023]. Similarly, there have been attempts to impose these exact constraints when developing ML KE functionals, the first of which applied the coordinate scaling condition in a 1D analog test [Hollingsworth et al. 2018]. Imposing the constraint for an ML functional by kernel ridge regression showed substantially improved accuracy for a 1D Hooke's atom case, but no improvement in the case of bond stretching in a 1D H₂ study. The Pauli positivity condition requires that the Pauli potential over all space is non-negative. It is satisfied if the KEDF only includes the Thomas-Fermi and von Weizsäcker terms, but it is not met in the fourth-order expansion used by Golub and Manzhos [2019]. Neither the scaling or asymptotic limit conditions are met by Yao and Parkhill [2016], although the authors point out that the physical constraints can be met in their approach via the training data fed to the convolutional neural network.

Finally, some studies aimed to learn a density functional other than XC or KE, such as the total energy functional, or learn corrections to a density functional rather than the functional itself [Bogojeski et al. 2020; Mezei and von Lilienfeld 2020]. One study developed a kernel ridge regression model to learn the difference (Δ -learning [Ramakrishnan et al. 2015]) in the energies from a low-level (e.g. DFT) calculation and a high-level calculation using coupled-cluster with single, double, and perturbative triple excitations (CCSD(T)) [Bogojeski et al. 2020]. This correction factor is a functional of the input DFT densities, and it allows for highly accurate predictions with the low computational cost of a standard KS-DFT calculation that scales with $\mathcal{O}(N_e^3)$ instead of $\mathcal{O}(N_e^7)$ for the CCSD(T) calculations. Another interesting approach uses machine learning to recommend the best density functional approximation for a given system, outperforming Δ -learning models as well as any of the other 48 tested approximations [Duan et al. 2023b]. Perhaps the most unique development is to learn the map from potential to density directly, called a Hohenberg-Kohn (HK) map, which can be done at a smaller computational cost and avoid the problem of the functional derivative [Brockherde et al. 2017].

4.3.3 Datasets and Benchmarks.

Compilations of highly accurate chemical data calculated at higher theory levels than DFT, such as CCSD(T), serve as an indispensable resource for the development of ML density functionals. These datasets can be used to train and test the accuracy of a density functional and compare with the performance of other functionals, especially useful for ML-based density functionals that require large amounts of highly accurate data for their training. Great efforts have been made to develop high quality datasets. For example, ACCDB is a collection of chemistry databases [Morgante and Peverati 2019], including five previously established databases (GMTKN, MGCD84 [Mardirossian and Head-Gordon 2017], Minnesota2015, DP284 [Hait and Head-Gordon 2018b,a], and W4-17), two new reaction energy databases automatically generated [Margraf et al. 2017] (W4-17-RE from W4-17 and MN-RE from Minnesota 2015B), and a new database for transition metals, which can be used as a benchmark for the development of density functionals. The GMTKN database consists of GMTKN55 [Goerigk et al. 2017] and MB08-165 [Korth and Grimme 2009]. The Minnesota database includes Database 2015 [Haoyu et al. 2015], Database 2015A [Yu et al. 2016b], and Database 2015B [Yu et al. 2016a]. Collectively, the ACCDB contains 10,049 structures and 8,656 unique reference data points (44,931 if the reaction energies are included), allowing for a substantial amount of high-quality data for training and testing. Another useful dataset is the SOL62 database [Trepte and Voss

2022; Zhang et al. 2018c], which consists of the cohesive energies of 62 solids (40 non-metals and 22 metals).

All these databases can be used as training and testing data for the development of machine-learned density functionals, as well as the evaluation of the functional. The accuracy of the prediction compared to those from higher theory levels around ten times more accurate enables a good benchmark for comparison, and several of these databases include the information of accuracy comparison for many other XC functionals. The datasets are split into various subsets to evaluate the functionals in different applications, such as atomization energies, barrier heights, bond energies, noncovalent interactions, and more.

In summary, many different machine learning approaches have been developed which provide more and more accurate density functionals (such as XC or KE functionals) and improve the predictions, even approaching chemical accuracy, with much lower computational cost compared to the higher level calculations. In Section 4.3.4 we discuss potential directions in this area for further exploration.

4.3.4 Open Research Directions.

ML-Based XC Functional: One exciting area of research is using symbolic regression to find the mathematical expression of density functionals that best fits the dataset, offering a mechanism of creating functionals from scratch or improving previously known functionals. Ma et al. [2022] is able to reconstruct a previously known functional from a starting point of a small library of mathematical instructions (e.g., multiplication, exponentiation, or building blocks of existing functionals) using an evolutionary search procedure. They used the same method to iterate the ω B97M-V functional through a regularized evolutionary algorithm in order to get a new functional – Google Accelerated Science 22 (GAS22), with improved error on the MGCD84 dataset. This fundamentally novel approach can be applied to other density functionals, such as KE functional for OF-DFT or universal functional, with a larger library of mathematical instructions for a wider search space which is more likely to find improved or novel functionals. The initial library was limited to only four (conveniently chosen) instructions in the first case, and for the second case the library included five arithmetic operations, six simple power operations, and the enhancement factors in the PBE, RPBE, B88, and B97 functionals. This library can be expanded to include more simple and complicated mathematical operations as well as the components of many previous functionals that were included in this study. Furthermore, the method can include density information, regularization, and exact constraints to restrict the functional form to be more physically motivated.

Another symbolic approach to ML for density functionals is through the use of automated feature engineering (AFE) and Q-learning, a reinforcement learning method. A recent investigation has used AFE to produce a feature generation tree where features are combined by mathematical operations into a physically meaningful equation, the exploration of which is guided by a deep Q-network (DQN) [Xiang et al. 2021]. The result demonstrated the improved classification and regression scores with less computation time for three material science databases compared to primary feature sets as well as another recent feature generation and selection technique, SISSO [Ouyang et al. 2018]. This novel method may be leveraged towards the discovery of an analytical XC functional by using AFE to produce equation-like features and applying the DQN to select the optimum features for further exploration. Thus, this AFE+DQN method offers another promising approach to the discovery of a symbolic XC functional, in addition to the evolutionary automated ML approach described above.

ML-Based KE Functional: With the substantial scaling advantage of OF-DFT to KS-DFT, there is significant interest in developing accurate kinetic energy density functionals (KEDFs), and machine

learning methods have demonstrated powerful towards that end. One area of future development in this area is to impose physical constraints on those KEDFs for ML models, whether that is through an analytical approach or a data-driven approach. It is still not entirely clear the full impact of imposing the exact constraints on the KEDFs on the accuracy of an ML model. In fact, because the KE functionals are much less well-explored compared to XC functionals, more work is needed to even establish these physical constraints by studying the behavior of the exact KEDF, *e.g.* addressing six open questions regarding the exact KEDF raised in a recent review article [Wrighton et al. 2023].

While there has been much work addressing the noisy functional derivative of the KEDF, there are more opportunities for avoiding gradient-based optimization entirely, such as using Monte Carlo-based optimization and erasing the need for evaluating a functional derivative of the KE, a method that still needs to be extended to the three-dimensional case [Ryczko et al. 2022b]. There is also room for exploring more methods that do not make use of the functional derivative [Brockherde et al. 2017]. Furthermore, exclusively learning on converged densities and energies limits the accuracy of ML models. While there have been steps forward to expand the kinds of training data used with a KS regularizer [Li et al. 2021a; Kalita et al. 2022] or automatic differentiation [Dick and Fernandez-Serra 2021], one further possibility is to find the converged external potentials to which the un converged densities correspond to. Some insightful suggestions have been made along these lines by [Ryczko et al. 2022b]. Another issue which is repeatedly raised is the non-uniform sampling and density distributions, which cause issues for ML models.

Overall, there has been substantial progress in ML for density functionals, among which the improvement that arises from the imposition of physical constraints is particularly interesting. One area for future improvement is that many of these functionals are fitted to or tested only on atomic or molecular systems, and their generalizability to solid-state systems needs to be further tested [Perdew 2021; Pokharel et al. 2022; Pederson et al. 2022]. Therefore, the use of datasets such as LC20 from [Sun et al. 2011] or SOL62 [Trepte and Voss 2022; Zhang et al. 2018c] for training and testing may advance ML density functionals for solid-state systems [Bystrom and Kozinsky 2023]. Similarly, most of the ML-based density functional predictors train their ML model with a small number of molecules, making their model hard to generalize in unseen molecules or solid-state systems that are different from the training data. For example, Nagai et al. [2020] incorporates three small molecules in the training set, which showed promising results across first- and second-row molecules, but it hasn't been tested on transition metal or extended systems yet. Since different kinds of molecules vary greatly in their properties, *e.g.*, organic and inorganic, small molecules and macromolecules, and open-shell and close-shell molecules, developing a ML-based density functional that generalizes well across different groups of molecules as well as solids is an important direction to explore in the future.

Going Beyond Atomistic Scale: Last but not least, the AI/ML development for quantum mechanics presented in Section 3 may allow efficient generation of more accurate datasets, which will in turn advance the development of the ML models towards the exact density functional. It is anticipated that quantum tensor learning and/or accurate ML density functionals will bring transformative impact to many other scientific fields such as organic and inorganic chemistry, condensed matter physics, materials design for electrical, mechanical, aerospace, nuclear, civil, and environmental engineering, as well as biological science and pharmaceutical research such as protein folding and drug design. In particular, it will enable the generation of more accurate datasets for the AI/ML development of molecules (Section 5), proteins (Section 6), materials (Section 7), and molecular docking (Section 8), *etc.*

5 AI FOR SMALL MOLECULES

In chemistry, a small molecule refers to a relatively low molecular weight organic compound. It typically comprises a small number of atoms, usually less than 100, and has a defined chemical structure. Small molecules are contrasted with macromolecules, such as proteins, nucleic acids, and polymers, which are much larger in size and often have complex structures. The use of AI approaches in small molecule learning allows for the development of more accurate and efficient methods for molecular predictive and generative tasks. In this section, we consider several key tasks in AI for molecular learning, including molecular representation learning, molecular conformer generation, molecule generation from scratch, molecular dynamics simulation, and representation learning of stereoisomerism and conformational flexibility, as summarized in Figure 17.

5.1 Overview

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Since machine learning approaches modeling 2D molecular graphs have been widely explored and achieved promising results [Gilmer et al. 2017; Wu et al. 2018; Yang et al. 2019; Hu et al. 2020; Wang et al. 2022h,e; Edwards et al. 2022; Veličković 2023], here, we focus on modeling 3D geometric molecules, a more challenging and practically meaningful perspective. The 3D geometry of a molecule plays a crucial role in many molecular predictive and generative tasks. To be specific, the 3D geometry of a molecule is a critical factor in determining many important properties, such as quantum properties [Ramakrishnan et al. 2014; Schütt et al. 2018; Liu et al. 2022f, 2021a, 2023a] and its binding affinity to a target protein, which is largely dependent on the complementary 3D shape of the molecule and the target protein [Anderson 2003]. Therefore, modeling 3D molecular geometries using predictive and generative AI approaches has immense potential. Particularly, it can significantly enhance the accuracy of predicting molecular properties and generate new molecules with desired properties.

In molecular representation learning, given 3D molecular geometries, our objective is to learn informative representations for various downstream tasks, such as molecule-level predictions and atom-level predictions. These representations are expected to capture accurate structural and chemical features of molecules. This task is fundamental since it is the basis for many advanced topics, such as drug discovery, materials design, and chemical reactions. The next two tasks we considered in this section are generative tasks. Specifically, molecular conformer generation is a conditional generation task where we aim to generate low-energy geometries or equilibrium ground-state geometries given a 2D molecular graph. This can provide an alternative to computationally expensive methods like density functional theory for obtaining 3D molecular geometries, thereby having significant potential in accelerating molecular simulation applications. Further, in certain applications, the desired 2D molecular graph is unknown, and we are interested in generating desired 3D molecules from scratch. Thus, in the task of molecule generation from scratch, our goal is to model the distribution over the 3D molecular geometry space with generative approaches. This can be used as the first step to generate novel molecules with desired properties for drug discovery, material science, and other applications. In addition to generic molecular representation learning, it is particularly important to learn to simulate molecular dynamics, which allows us to capture the time-dependent behavior of molecular systems, providing invaluable insights into their physical properties and structural transformations. AI approaches are facilitating the field of molecular dynamics simulations mainly through improving the accuracy of force fields, enhancing sampling methods, and enabling effective coarse-graining. Lastly, we consider the inherent complexity of

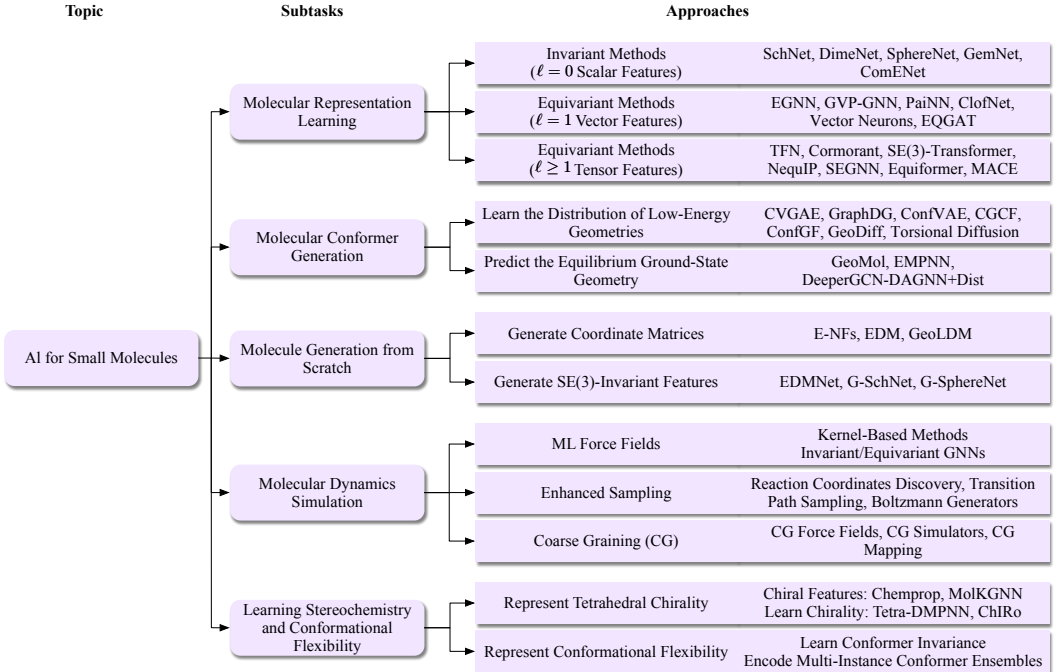


Fig. 17. An overview of the tasks and methods in AI for small molecules. In this section, we consider five tasks, including molecular representation learning, molecular conformer generation, molecule generation from scratch, molecular dynamics simulation, and learning stereochemistry and conformational flexibility. In molecular representation learning, the $\ell = 0$ case corresponds to invariant methods, including SchNet [Schütt et al. 2018], DimeNet [Gasteiger et al. 2020], SphereNet [Liu et al. 2022f], GemNet [Gasteiger et al. 2021], and ComENet [Wang et al. 2022g]. The $\ell = 1$ case corresponds to equivariant methods with order-1 vector features $v \in \mathbb{R}^{d \times 3}$, including EGNN [Satorras et al. 2021a], GVP-GNN [Jing et al. 2021], PaiNN [Schütt et al. 2021], ClofNet [Du et al. 2022], Vector Neurons [Deng et al. 2021], and EQGAT [Le et al. 2022]. The $\ell \geq 1$ case corresponds to equivariant methods with order- ℓ features $h^\ell \in \mathbb{R}^{d \times (2\ell+1)}$, including TFN [Thomas et al. 2018], Cormorant [Anderson et al. 2019], SE(3)-Transformer [Fuchs et al. 2020], NequIP [Batzner et al. 2022], SEGNN [Brandstetter et al. 2022a], Equiformer [Liao and Smidt 2023], and MACE [Batabia et al. 2022b]. In molecular conformer generation, one category of methods aims to learn the distribution of low-energy geometries, including CVGAE [Mansimov et al. 2019], GraphDG [Simm and Hernández-Lobato 2019], ConfVAE [Xu et al. 2021d], CGCF [Xu et al. 2021a], ConfGF [Shi et al. 2021], GeoDiff [Xu et al. 2022b], and Torsional Diffusion [Jing et al. 2022]. Another category of methods aims to predict only the equilibrium ground-state geometry, including GeoMol [Ganea et al. 2021], EMPNN [Xu et al. 2023c] and DeeperGCN-DAGNN+Dist [Xu et al. 2021b]. In molecule generation from scratch, one category of method aims to directly generate coordinate matrices of 3D molecules, including E-NFs [Satorras et al. 2021b], EDM [Hoogeboom et al. 2022], and GeoLDM [Xu et al. 2023a]. Another category of methods implicitly generates 3D atom positions from $SE(3)$ -invariant features, including EDMNet [Hoffmann and Noé 2019], G-SchNet [Gebauer et al. 2019], and G-SphereNet [Luo and Ji 2022]. In molecular dynamics simulation, research directions including ML force fields [Unke et al. 2021c], enhanced sampling [Sidky et al. 2020a], and coarse-graining approaches [Noid 2023] are briefly introduced. Learning stereochemistry has focused on encoding tetrahedral chirality by employing heuristic features (Chemprop [Yang et al. 2019], MolKGNN [Liu et al. 2022g]) or designing chiral message passing operations (Tetra-DMPNN [Pattanaik et al. 2020], ChiRo [Adams et al. 2021]). Representing conformational flexibility has involved learning conformer invariance [Adams et al. 2021] or explicitly encoding multi-instance conformer ensembles [Axelrod and Gomez-Bombarelli 2020; Chuang and Keiser 2020].

molecular structures for more effective representation learning. Specifically, we discuss the importance of molecular stereochemistry and conformational flexibility during molecular representation learning.

Since we are modeling molecules in 3D space, it is desired to take the underlying equivariance and invariance properties into consideration. Preserving the desired symmetry in 3D molecular learning tasks is crucial for obtaining accurate predictions and ensuring the physical constraints of the system. In addition, how to capture the 3D information accurately, such as distinguishing enantiomers of the same molecule, is another important consideration to achieve effective modeling.

5.2 Molecular Representation Learning

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Recommended Prerequisites: Sections 2.3, 2.4

In this section, we study the problem of molecular representation learning, which aims to learn informative representations of given input molecules. The learned representations can be used for various downstream tasks, such as molecule-level prediction and atom-level prediction. In addition, the representation learning models introduced in this section can be seen as backbones that enable more advanced applications, such as drug discovery and material design.

5.2.1 Problem Setup.

Molecular Graphs and Point Clouds: Molecules may be represented as 2D molecular graphs, which contain the graph topology (bonds between atoms) as well as node and edge features or as 3D molecular graphs, which additionally consider the 3D coordinates for each node. While the 2D representation suffices to describe the chemical identity of a molecule, the 3D configuration of the molecule (called *conformer*) is relevant for determining many experimentally relevant properties of the molecule, such as its energy or electric dipole moment. Thus, we focus on methods for working with 3D molecular graphs in the remainder of this section. Formally, we represent a 3D molecule as a point cloud with n atoms as $\mathcal{M} = (\mathbf{z}, \mathbf{C})$, where $\mathbf{z} = [z_1, \dots, z_n] \in \mathbb{Z}^n$ is the atom type vector and $\mathbf{C} = [c_1, \dots, c_n] \in \mathbb{R}^{3 \times n}$ is the atom coordinate matrix. To obtain a molecular graph from this point cloud, edges may then be added for example from the bonds (2D graph topology), from radial distance cut-offs or from the k nearest neighbors. Because edge construction differs between methods (further discussed below), we refer to a molecule as its point cloud $\mathcal{M} = (\mathbf{z}, \mathbf{C})$.

Task Formulations: We aim to learn latent representations of 3D molecules which can be used for downstream prediction tasks and applications. Two types of downstream prediction tasks are of interest: *molecule-level* predictions and *atom-level* predictions. For molecule-level property prediction tasks, we aim to learn a function $f(\mathcal{M})$ to predict a property y of any given molecule \mathcal{M} . Here, y can be a real number (regression problems such as the energy of a conformer), an integer (classification problems such as toxicity), or a tensor (such as the electric dipole vector, or the tensor of inertia). If the target property y is a scalar/tensorial quantity, it needs to be invariant/equivariant to changes in reference frame. For atom-level property prediction tasks, we aim to learn a function f to predict the property y_i of the i -th atom, such as per-atom forces for molecular simulation. Again, y may be a scalar or tensorial target property.

5.2.2 Technical Challenges.

Different from typical 2D graphs with topology only, the geometry of 3D structures poses unique challenges to 3D molecular modeling.

(1) The first challenge is that the learned representations correspond to physical geometric quantities and should follow the underlying symmetries for different applications [Bogatskiy et al. 2022]. To be specific, for tasks like energy prediction, the learned representations should be $SE(3)$ -invariant. This means that if the input molecule is rotated or translated, the learned representations should remain unchanged. For tasks like per-atom force prediction, the representations should be $SO(3)$ -equivariant. This is because if the input molecule is rotated, the prediction target (e.g., forces) should rotate accordingly.

(2) Another challenge is the theoretical expressive power of learned representations [Joshi et al. 2023], which instantiates itself as practical limitations of models at distinguishing different 3D geometries of molecules, such as enantiomers and different conformers of the same molecule. Learning expressive molecular representations is crucial for applications like drug design and molecular simulations [Pozdnyakov et al. 2020]. For example, the enantiomers of a chiral drug can interact very differently with other chiral molecules and proteins. Different conformers of the same molecule also have different potential energies and per-atom forces.

(3) Thirdly, efficiency is an important factor to consider when designing a model for molecular representation learning. High efficiency enables fast training and inference, reduces computational resources, and enhances scalability to large-scale, real-world datasets.

5.2.3 Overview of Existing Methods.

As indicated above, a 2D molecular graph contains the graph topology as well as the original node and edge features, base on which a 3D molecular graph further considers 3D coordinates for each node. Any geometric quantities, like distance, angle, and torsion angle, can be computed from the 3D coordinates. More generally, as introduced in Section 2, each node has an order- ℓ $SE(3)$ -equivariant node feature. From the perspective of tensor order, existing methods for 3D molecular representation learning can be categorized into invariant 3D graph neural networks (3D GNNs) with only $\ell = 0$ scalar-type features [Schütt et al. 2017a; Smith et al. 2017; Chmiela et al. 2017; Zhang et al. 2018a,b; Unke and Meuwly 2019; Schütt et al. 2018; Ying et al. 2021; Luo et al. 2023b; Gasteiger et al. 2020; Liu et al. 2022f; Gasteiger et al. 2021; Wang et al. 2022g], equivariant 3D GNNs with $\ell = 1$ vector-type features [Schütt et al. 2021; Jing et al. 2021; Satorras et al. 2021a; Du et al. 2022, 2023a; Thölke and Fabritiis 2022], and equivariant 3D GNNs with higher order $\ell \geq 1$ tensor features [Thomas et al. 2018; Fuchs et al. 2020; Liao and Smidt 2023; Batzner et al. 2022; Batatia et al. 2022a,b]. Specifically, invariant methods directly take invariant geometric features such as distances and angles as input, and thus, all internal features remain unchanged regardless of transformations like rotation and translation of the input molecule. In contrast, internal features in equivariant methods should transform accordingly the input molecule is rotated or translated.

In addition to tensor order, existing 3D GNN layers can be further categorized from the perspective of body order. Body order originates from the decomposition of potential energy surface (PES) as a linear combination of body-ordered functions. Traditional approaches [Brown et al. 2004; Braams and Bowman 2009] show that body order expansion as illustrated in Figure 18 leads to high accuracy and fast convergence in approximating the PES of molecular and material systems. As components of the total molecular energy, we can write the local energy of atoms in a body-ordered expansion:

$$E_i = f_1(\sigma_i) + \sum_{j_1}^N f_2(\sigma_i, \sigma_j) + \sum_{j_1, j_2} f_3(\sigma_i, \sigma_{j_1}, \sigma_{j_2}) + \dots + \sum_{j_1, \dots, j_v} f_v(\sigma_i, \sigma_{j_1}, \dots, \sigma_{j_v}), \quad (91)$$

where i is the index of the central atom, j_k is the index of one i 's neighbor node, N is the total number of 1-hop neighbors of atom i , and σ denotes the atom state, including atom attributes and

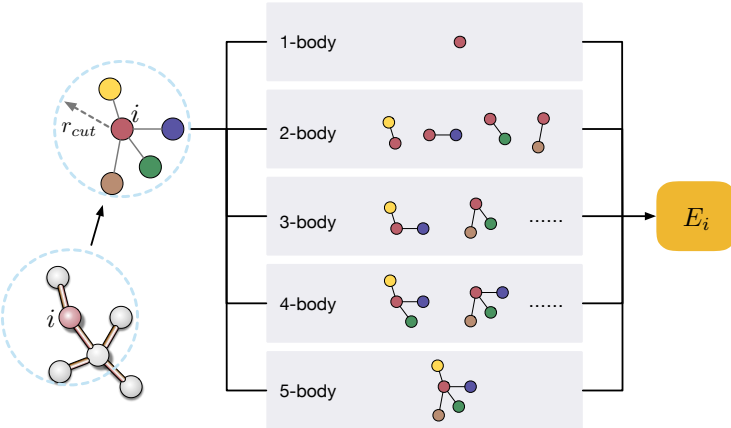


Fig. 18. An illustration of body order expansion in molecular energy prediction. First, neighboring nodes and edges of the central atom i are determined by a cutoff r_{cut} . Then, v body term considers all combinations of the central atom i and $v - 1$ of its 1-hop neighbors. Finally, the local energy of atom i is computed as a linear combination of all body-ordered functions. Compared to body order expansion [Brown et al. 2004; Braams and Bowman 2009], the standard message passing [Gilmer et al. 2017] only considers a body order of 2 as it solely involves the central atom and one neighboring atom in each message.

coordinates for the corresponding subscript atom. Generally, in body-order expansion, the v body term considers all combinations of the central atom and $v - 1$ of its 1-hop neighbors. Therefore, the standard message passing [Gilmer et al. 2017]

$$\begin{aligned} \mathbf{m}_i &= \sum_{j \in \mathcal{N}(i)} M(\mathbf{h}_i, \mathbf{h}_j, \mathbf{h}_{ij}), \\ \mathbf{h}'_i &= U(\mathbf{h}_i, \mathbf{m}_i) \end{aligned} \quad (92)$$

has a body order of 2 because each message involves the central atom and one neighbor. Here $\mathcal{N}(i)$ is the neighboring node set of node i , \mathbf{h}_{ij} is the edge feature between node i and node j , such as the edge length and edge type, and U and M are the update and message functions. Although standard message passing can further aggregate information from many nodes along edges through iterative layers, such aggregation is distinct from many-body interaction that is restricted within 1-hop of the central node. In this subsection, we discuss existing 3D molecular representation learning methods based on their tensor order as well as body order, as summarized in Figure 19.

5.2.4 Invariant Methods ($\ell = 0$ Scalar Features).

Invariant methods only maintain invariant node, edge, or graph features, which do not change if the input 3D molecule is rotated or translated. Invariant methods face a trade-off between improving their discriminative ability by considering many-body geometric features and maintaining their efficiency, as summarized in Table 5. Let n and k denote the number of nodes and the average degree in a molecule. Specifically, SchNet [Schütt et al. 2018] considers only pairwise distances as edge features \mathbf{h}_{ij} in the node-centered message passing schema shown in Equation (92), resulting in a complexity of $O(nk)$ and a body order of 2. DimeNet [Gasteiger et al. 2020] further considers

	Body Order				Example Operations	
	2	3	4	Many	Linear Layers	Others
Tensor Order	$\ell = 0$	SchNet	DimeNet GemNet-T	SphereNet ComENet		
	$\ell = 1$	EGNN	GVP-GNN PaiNN Vector Neurons EQGAT			
	$\ell \geq 1$	TFN Cormorant SE(3)-Transformer NequIP SEGNNet Equiformer			ACE Linear ACE MACE	

Fig. 19. An overview of existing methods for molecule representation learning. We categorize existing methods based on the tensor order of features and the body order of GNN layers, which are two key design choices for building maximally powerful 3D GNNs, as discussed in detail in Joshi et al. [2023] and Section 5.2.10. Invariant methods with $\ell = 0$ scalar features are described in detail in Section 5.2.4, equivariant methods with $\ell = 1$ vector features in Section 5.2.5, equivariant methods with $\ell \geq 1$ tensor features in Section 5.2.6, and higher body order methods in Section 5.2.7. In addition, different order features require specific operations to maintain $SE(3)$ equivariance. Here we list several example operations for methods with different tensor orders. Specifically, for the linear layers, each gray line between input and output features contains a learnable weight. The bias term can only be added to $\ell = 0$ scalar features, as it would break the equivariance of $\ell \geq 1$ features. Additionally, tensor product, introduced in 2.4 and illustrated in Figure 6, is another crucial operation for equivariant methods with $\ell \geq 1$ tensor features as it can maintain $SE(3)$ -equivariance of higher-order features. This figure is adapted from Joshi et al. [2023] with permission.

Table 5. Summary of existing invariant 3D graph neural networks ($\ell = 0$) for molecular representation learning, including SchNet [Schütt et al. 2018], DimeNet [Gasteiger et al. 2020], GemNet [Gasteiger et al. 2021], SphereNet [Liu et al. 2022f], and ComENet [Wang et al. 2022g]. Here n and k denote the number of nodes and the average degree in a molecule. The complexity depends on the calculation of the geometric features and the message passing schema.

Methods	Invariant Geometric Features	Body Order	Complexity	
SchNet		Pairwise distances d	2-body	$O(nk)$
DimeNet		$d + \text{Angles between edges } \theta$	3-body	$O(nk^2)$
GemNet		$d, \theta + \text{Angles between 4 nodes } \tau$	4-body	$O(nk^3)$
SphereNet		$d, \theta + \text{Angles between 4 nodes } \phi$	4-body	$O(nk^2)$
ComENet		d, θ, ϕ, τ	4-body	$O(nk)$

angles between each pair of edges with edge-centered message passing

$$\begin{aligned} \mathbf{m}_{ji} &= \sum_{k \in N(j) \setminus \{i\}} M(\mathbf{h}_{ji}, \mathbf{h}_{kj}, \mathbf{h}_{kji}), \\ \mathbf{h}'_{ji} &= U(\mathbf{h}_{ji}, \mathbf{m}_{ji}), \end{aligned} \quad (93)$$

and the complexity is $O(nk^2)$. Here $N(j) \setminus \{i\}$ is the set of neighboring nodes of node j except for node i , \mathbf{h}_{kji} is the feature of nodes k, j , and i , such as the angle θ_{kji} , and U and M are the update

and message functions. GemNet [Gasteiger et al. 2021] further considers two-hop dihedral angles, increasing body order to 4 and complexity to $O(nk^3)$. SphereNet [Liu et al. 2022f] computes local 4-body angles between two planes. To reduce the complexity, SphereNet does not incorporate all possible angles, instead reducing the number of angles by selecting reference nodes to construct reference planes while retaining $O(nk^2)$ complexity. ComENet [Wang et al. 2022g] defines complete geometric features that can distinguish all different 3D molecules that exist. Specifically, the distance and angles d, θ, ϕ are 2-body, 3-body, and 4-body geometric features and can be used to identify local structures. Here a local structure means a central node and its 1-hop neighborhood. This is because $d_{ij}, \theta_{ij}, \phi_{ij}$ can determine the relative position of node j in the local spherical coordinate system centered in i . In addition, The rotation angle τ further captures the remaining degree of freedom between local structures. Therefore, ComENet has the ability to generate a unique representation for each 3D molecule, able to distinguish all different 3D molecules in nature. Moreover, it follows the node-centered message passing schema in Equation (92), and the complexity is only $O(nk)$ by selecting reference nodes within 1-hop neighborhood.

In addition to the methods that convert equivariant 3D information to invariant features like distances and angles [Schütt et al. 2018; Gasteiger et al. 2020; Liu et al. 2022f; Gasteiger et al. 2021; Wang et al. 2022g], Du et al. [2022, 2023a] propose scalarization to obtain invariant features. Specifically, scalarization converts equivariant features into invariant features based on equivariant local frames. For example, given an equivariant frame $(\mathbf{e}_1, \mathbf{e}_2, \mathbf{e}_3)$, we can convert a 3D vector $\mathbf{r}_{ij} = \mathbf{c}_i - \mathbf{c}_j$ to $(\mathbf{r}_{ij} \cdot \mathbf{e}_1, \mathbf{r}_{ij} \cdot \mathbf{e}_2, \mathbf{r}_{ij} \cdot \mathbf{e}_3)$. Here $\mathbf{e}_1, \mathbf{e}_2, \mathbf{e}_3$ form an orthonormal basis. In addition to scalarization, Du et al. [2022, 2023a] also use tensorization to convert invariant features to equivariant features. Therefore, these methods can maintain both invariant and equivariant internal features and require both invariant operations and equivariant operations (see Section 5.2.5 and 5.2.6) to update the internal features.

5.2.5 Equivariant Methods ($\ell = 1$ Vector Features).

The first category of equivariant 3D GNNs [Satorras et al. 2021a; Du et al. 2022; Schütt et al. 2021; Deng et al. 2021; Jing et al. 2021; Thölke and Fabritiis 2022] uses order 1 vectors as intermediate features and propagates messages via a restricted set of operations that guarantee $E(3)$ or $SE(3)$ equivariance, as summarized in Table 6. Let us denote a scalar feature by $s \in \mathbb{R}^d$ and a vector by $v \in \mathbb{R}^{d \times 3}$. As summarized in Schütt et al. [2021] and Deng et al. [2021], operations on a vector v that can ensure equivariance include scaling of vectors $s \odot v$, summation of vectors $v_1 + v_2$, linear transformation of vectors Wv , scalar product $\|v\|^2$, $v_1 \cdot v_2$, and vector product $v_1 \times v_2$. Here \odot denotes element-wise multiplication. Note that $v_1 \cdot v_2 = \|v_1\| \|v_2\| \cos \theta$ and $v_1 \times v_2 = \|v_1\| \|v_2\| \sin \theta \vec{n}$, therefore, using scalar product and vector product can implicitly incorporate angular and directional information.

Existing methods use these operations to update both scalar and vector features by propagating scalar as well as vector messages. For example, EGNN [Satorras et al. 2021a] uses scaling of vectors and vector summation to ensure equivariance. To be specific, following the notation of Equation (92), an EGNN layer updates node representation \mathbf{h}_i and node coordinate \mathbf{c}_i as

$$\begin{aligned} \mathbf{m}_{ij} &= \phi_e(\mathbf{h}_i, \mathbf{h}_j, \|\mathbf{c}_i - \mathbf{c}_j\|^2, \mathbf{h}_{ij}), \\ \mathbf{c}'_i &= \mathbf{c}_i + C \sum_{j \neq i} (\mathbf{c}_i - \mathbf{c}_j) \phi_c(\mathbf{m}_{ij}), \\ \mathbf{h}'_i &= \phi_h \left(\mathbf{h}_i, \sum_{j \neq i} \mathbf{m}_{ij} \right), \end{aligned} \tag{94}$$

Table 6. Comparisons of equivariant methods using $\ell = 1$ vector features, including EGNN [Satorras et al. 2021a], ClofNet [Du et al. 2022], PaiNN [Schütt et al. 2021], GVP-GNN [Jing et al. 2021], Vector Neurons [Deng et al. 2021], and EQGAT [Le et al. 2022]. Here $s \in \mathbb{R}^d$ denotes a scalar feature, and $v \in \mathbb{R}^{d \times 3}$ denotes a vector feature. Existing methods use different operations to ensure equivariance.

Methods	Scaling $s \odot v$	Summation $v_1 + v_2$	Linear Transformation Wv	Scalar Product $\ v\ ^2, v_1 \cdot v_2$	Vector Product $v_1 \times v_2$
EGNN	✓	✓		✓	
ClofNet	✓	✓		✓	
PaiNN	✓	✓	✓	✓	
GVP-GNN	✓	✓	✓	✓	
Vector Neurons	✓	✓	✓	✓	
EQGAT	✓	✓	✓	✓	✓

where ϕ_e , ϕ_c , and ϕ_h denote learnable functions and C is a normalization factor. Different from EGNN which only considers a single vector for each edge, ClofNet [Du et al. 2022] employs complete frames that consist of three vectors for each edge. PaiNN [Schütt et al. 2021] further includes linear transformation and scalar product in the network. GVP-GNN [Jing et al. 2021] uses similar operations as PaiNN and was originally designed to learn representations for protein structures, but can also be adapted to molecules. Vector Neurons [Deng et al. 2021] is originally designed for point cloud data and can be applied to molecules. It also employs linear transformation to achieve the linear operator for order 1 vectors. In addition to linear operators, Vector Neurons incorporate carefully designed non-linear, pooling, and normalization layers that are tailored for order 1 vectors while ensuring the desired equivariance. Notably, it uses a learnable direction, which is equivariant, to split the domain into two half-spaces, and then non-linear layers such as ReLU can be defined to map such two spaces differently. In addition to the aforementioned operations, EQGAT [Le et al. 2022] uses cross product to update equivariant features during message passing. This enables interactions between type-1 vector features and allows for more comprehensive and expressive feature representations. Moreover, it uses attention mechanism to capture content and spatial information between nodes.

5.2.6 Equivariant Methods ($\ell \geq 1$ Tensor Features).

Another category of equivariant methods considers higher-order ($\ell \geq 1$) features that have been discussed in Section 2. Most existing methods under this category use tensor products (TP) of higher-order spherical tensors to build equivariant representations and follow the general architecture in Figure 20 to update features, and differing in body order and technical details. For example, TFN [Thomas et al. 2018] and NequIP [Batzner et al. 2022] follow the node-centered message passing scheme [Gilmer et al. 2017] to update node features based on messages from neighboring nodes. Since each message contains the information of the central atom and one neighbor, these methods naturally have a body order of 2. SE(3)-Transformer [Fuchs et al. 2020] and Equiformer [Liao and Smidt 2023] further enhance their model architectures with the attention mechanism. In addition, Cormorant [Anderson et al. 2019] and SEGNNs [Brandstetter et al. 2022a] introduce different designs of non-linearity on higher-order features.

Generally, higher order equivariant models build multiple feature vectors or "channels" for each rotation order $\ell \leq \ell_{max}$. Each channel of order ℓ has a length of $2\ell + 1$. As such, the features of node i can be indexed by \mathbf{h}_{icm}^ℓ , where ℓ is the rotation order, c is the channel index, and m is the

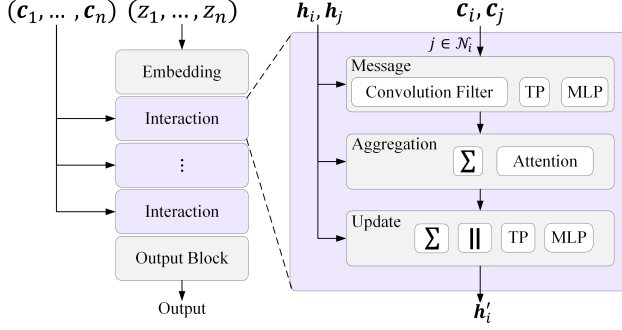


Fig. 20. A general architecture of higher order equivariant models. Each model consists of several interaction blocks which perform pairwise message passing between atoms. Here, Σ denotes summation, \parallel denotes feature concatenation, TP refers to the Tensor Product of feature vectors, and MLP refers to multilayer perceptrons. The specific operations used in each Message, Aggregation, and Update blocks differ between models, but existing methods such as TFN [Thomas et al. 2018], NequIP [Batzner et al. 2022], SE(3)-Transformer [Fuchs et al. 2020], Equiformer [Liao and Smidt 2023], Cormorant [Anderson et al. 2019], and SEGNNs [Brandstetter et al. 2022a] all fall under this framework.

representation index ($-\ell \leq m \leq \ell$). A general architecture of higher order equivariant models is given in Figure 20, and we describe each component below.

Nonlinear Functions: In order to preserve equivariance, the nonlinear functions used in these models are restricted to those which act as scalar transforms in the representation index m . The nonlinear functions used by various models are shown in Table 7. Notably, Cormorant uses the tensor product as the only nonlinear operation, and SE(3)-Transformer uses attention instead of nonlinear activations found in other models.

Table 7. The nonlinear functions used in higher order equivariant models. $\eta : \mathbb{R} \rightarrow \mathbb{R}$ is a nonlinear function such as SiLU or tanh, $\|\mathbf{h}_c^\ell\| = \sqrt{\sum_m |\mathbf{h}_{cm}^\ell|^2}$, and b_c^ℓ is a learnable bias.

Methods	Nonlinear Functions, $g(\mathbf{h}_c^\ell)$
TFN	$\eta (\ \mathbf{h}_c^\ell\ + b_c^\ell) \mathbf{h}_c^\ell$
NequIP	$\eta (\ \mathbf{h}_c^\ell\) \mathbf{h}_c^\ell$
SEGNN, Equiformer	$\eta (\mathbf{h}_c^0) \mathbf{h}_c^\ell$
Cormorant, SE(3)-Transformer	\mathbf{h}_c^ℓ

Linear Layers: The linear layers used in these models take the form as

$$W(\mathbf{h}^\ell) = \sum_{cr} W_{ccr}^\ell \mathbf{h}_{icm}^\ell. \quad (95)$$

The weights are constant across the m dimension, which is required to maintain equivariance. Optionally, biases can be added for $l = 0$ features.

Convolution Filters: These models generally build convolution filters as the product of a learnable radial function and spherical harmonics. The specific filters used by several models are shown in Table 8.

Table 8. The convolution filters used in higher order equivariant models. Here, d_{ij} is the distance between nodes.

Methods	Convolution Filter, $F(\cdot)_{cm}^\ell$
TFN, NequIP, Equiformer, $SE(3)$ -Transformer SEGNN Cormorant	$R_c^\ell(d_{ij})Y_m\left(\frac{c_i - c_j}{d_{ij}}\right)$
	$Y_m^\ell\left(\frac{c_i - c_j}{d_{ij}}\right)$
	$R_c^\ell(d_{ij}, \mathbf{h}_{icm}^\ell, \mathbf{h}_{jcm}^\ell)Y_m^\ell\left(\frac{c_i - c_j}{d_{ij}}\right)$

Message: Pairwise messages are then built using tensor products. All methods begin by taking the tensor product of the convolution filer and node features, however, some methods further augment these messages. The specific equations to compute messages in each model are shown in Table 9. In general, the tensor product of type ℓ_1 and ℓ_2 feature vectors produces outputs at all rotation orders $|\ell_1 - \ell_2| \leq \ell_3 \leq \ell_1 + \ell_2$. Section 2.4 describes the tensor product operations in more detail.

Table 9. The equations for message computing in higher order equivariant models. $\phi(\cdot) = g(W(\cdot))$. \parallel denotes concatenation of features. The ℓ , c , and m message indices are omitted for brevity.

Methods	Message \mathbf{m}_{ij}
TFN, NequIP, Cormorant, $SE(3)$ -Transformer SEGNN Equiformer	$TP(F(\cdot), \mathbf{h}_j)$
	$\phi_2(TP(F(\cdot), \phi_1(TP(F(\cdot), \mathbf{h}_i \parallel \mathbf{h}_j \parallel d_{ij}))))$
	$W(TP(F(\cdot)), \phi(TP(F(\cdot), W(\mathbf{h}_i) + W(\mathbf{h}_j)))))$

Aggregation: For each atom, messages are then aggregated over neighboring atoms. For all models, sum aggregation is used, however, $SE(3)$ -Transformer and Equiformer first weigh the incoming messages using attention. The aggregation functions used by each model are shown in Table 10. Note $SE(3)$ -Transformer uses dot-product attention. Equiformer uses more powerful MLP attention, however, in order to preserve equivariance, only $\ell = 0$ features are used to compute attention scores.

Table 10. The aggregation functions used in higher order equivariant models. α_{ij} are the attention scores.

Methods	Aggregated Message, \mathbf{m}_i
TFN, NequIP, Cormorant, SEGNN $SE(3)$ -Transformer, Equiformer	$\sum_{j \in N_i} \mathbf{m}_{ij}$ $\sum_{j \in N_i} \alpha_{ij} \mathbf{m}_{ij}$

Update: Finally, the aggregated message is used to update the features for each node. The specific update functions used in each model are shown in Table 11. In TFN and $SE(3)$ -Transformer, no residual connection is used between layers. However, later works have shown this connection is crucial in order to retain chemical information such as atom type.

5.2.7 Higher Body Order Methods.

Previous equivariant graph network layers consider two-body interactions using the spherical harmonics of the direction vectors from neighbors to the center node. Higher body order incorporates

Table 11. The update functions used in higher order equivariant models.

Methods	Updated Features, \mathbf{h}'_i
TFN, $SE(3)$ -Transformer	$\phi(\mathbf{m}_i)$
NequIP	$\mathbf{h}_i + \phi(\mathbf{m}_i) W(\mathbf{m}_i)$
Cormorant	$W(\mathbf{m}_i \ \mathbf{h}_i \ \text{TP}(\mathbf{h}_i, \mathbf{h}_i))$
SEGNN	$\mathbf{h}_i + W(\text{TP}(\phi(\text{TP}(\mathbf{h}_i \ \mathbf{m}_i, Y(\mathbf{c}_i))), Y(\mathbf{c}_i)))$

many-body interactions among neighbors. To better approximate the atomic potential, many-body decomposition [Drautz 2019] is introduced as

$$E_i(\mathbf{r}_{ij_1}, \dots, \mathbf{r}_{ij_n}) = \sum_j \sum_v c_v^{(1)} \phi_v(\mathbf{r}_{ij}) + \frac{1}{2} \sum_{j_1 j_2} \sum_{v_1 v_2} c_{v_1 v_2}^{(2)} \phi_{v_1}(\mathbf{r}_{ij_1}) \phi_{v_2}(\mathbf{r}_{ij_2}) \\ + \frac{1}{3!} \sum_{j_1 j_2 j_3} \sum_{v_1 v_2 v_3} c_{v_1 v_2 v_3}^{(3)} \phi_{v_1}(\mathbf{r}_{ij_1}) \phi_{v_2}(\mathbf{r}_{ij_2}) \phi_{v_3}(\mathbf{r}_{ij_3}) + \dots, \quad (96)$$

where ϕ_v is the single bond basis function, \mathbf{r}_{ij} denotes the direction vector from node i to j , and c is the coefficients. However, the complexity of many-body decomposition is $O(n^d)$, which indicates an exponential increase with respect to the number of neighbors n for high body order. Here, d corresponds to the body order. To decrease the complexity, atomic cluster expansion (ACE) [Drautz 2019; Dusson et al. 2022] uses the atomic basis $A_{iv} = \sum_j \phi_v(\mathbf{r}_{ij})$ to represent the atomic energy as

$$E_i(\mathbf{r}_{ij_1}, \dots, \mathbf{r}_{ij_n}) = \sum_v c_v^{(1)} A_{iv} + \sum_{v_1 v_2} c_{v_1 v_2}^{(2)} A_{iv_1} A_{iv_2} + \sum_{v_1 v_2 v_3} c_{v_1 v_2 v_3}^{(3)} A_{iv_1} A_{iv_2} A_{iv_3} + \dots. \quad (97)$$

Thus, a linear growth with the number of neighbors can be maintained. Following the ACE, many machine learning methods [Dusson et al. 2022; Drautz 2019; Batatia et al. 2022b,a; Musaelian et al. 2023a; Kondor 2018; Li et al. 2022h; Bigi et al. 2023; Kovacs et al. 2023; Musaelian et al. 2023b; Batatia et al. 2023] are developed to consider the higher body order interactions.

Linear ACE and MACE are developed based on the theory of “density trick”, Linear ACE sequentially builds particle basis, atomic basis, product basis, symmetrized basis, and finally uses the linear combination of symmetrized basis to construct high body-order features efficiently. The Linear ACE model [Kovács et al. 2021] consists of only one layer while MACE [Batatia et al. 2022b,a] leverages tensor product and further extends to multiple ACE layers to enlarge the receptive field so that semi-local information is also incorporated through message passing. In contrast to using ACE to obtain aggregated messages for nodes, Allegro [Musaelian et al. 2023a] focuses operations on edges. Specifically, Allegro performs tensor products between edges around the central node and increases the order of body interaction through a stack of layers. The many-body embeddings produced by Allegro are analogous to ACE’s symmetrized basis, although not strictly equivalent. In addition to the above methods, Wigner kernels [Bigi et al. 2023] develops body-ordered kernels calculated in a radial-element space with a cheaper cost that is linear to the maximum body order. N-body networks [Kondor 2018; Li et al. 2022h] is a hierarchical neural network that aims to learn atomic energies based on the decomposition of the many-body system. Among existing many-body methods, Linear ACE, MACE, and Allegro follow the general architecture summarized in Figure 20.

Convolution Filters: Convolution filters used in higher body order methods are summarized in Table 12. Similar to methods such as NequIP in Section 5.2.6, MACE builds convolution filters as

the product of a learnable radial function and spherical harmonics. Linear ACE and Allegro do not have convolution filters.

Table 12. The convolution filters used in higher body order methods.

Methods	Convolution Filter, $F(\cdot)$
Linear ACE, Allegro	-
MACE	$R_c^\ell(d_{ij}) Y_m^\ell\left(\frac{c_i - c_j}{d_{ij}}\right)$

Message: As shown in Table 13, linear ACE builds messages as the product of a radial basis and spherical harmonics, where the radial basis $R_{c,z_i z_j}^\ell$ is coupled with atomic types of node i and j . MACE builds messages as the tensor product of the convolution filter and a linear transformation of node features. The linear transformations used by MACE are the same as in Section 5.2.6. Allegro builds messages by passing invariant features from the previous layer \mathbf{x}_{ij}^{t-1} to an MLP, then multiplying the output with spherical harmonics.

Table 13. The equations for message computing in higher body order methods.

Methods	Message, \mathbf{m}_{ij}
Linear ACE	$R_{c,z_i z_j}^\ell(d_{ij}) Y_m^\ell\left(\frac{c_i - c_j}{d_{ij}}\right)$
MACE	$TP(F(\cdot), W(\mathbf{h}_j))$
Allegro	$MLP(\mathbf{x}_{ij}^{t-1}) Y_m^\ell\left(\frac{c_i - c_j}{d_{ij}}\right)$

Aggregation: The aggregation functions used in higher body order methods are summarized in Table 14. In linear ACE and MACE, the aggregation is implemented following atomic cluster expansion (ACE). According to ACE, message m_{ij} is the 2-body particle basis for atoms i and j . First, the two-body particle basis functions are summed over neighboring atoms to obtain the atomic basis for the central atom, i . Then, products of v atomic basis functions create a $v + 1$ -body product basis. In order to preserve rotational equivariance, the product basis is multiplied with the generalized Clebsch-Gordan coefficients to form the symmetrized basis. Finally, the aggregated many-body message for each atom is constructed by a linear combination of symmetrized basis features with different body orders. Here, ℓm denotes $(\ell_1 m_1, \dots, \ell_v m_v)$ and an additional index η_v is used to enumerate all paths of rotation orders (ℓ_1, \dots, ℓ_v) which result in the desired output rotation order. In Allegro, messages from neighboring edges are summed around the central atom i to obtain the aggregated message m_i for atom i .

Table 14. The aggregation functions used in higher body order methods.

Methods	Aggregation, \mathbf{m}_i
Linear ACE	$\sum_v \sum_{\eta_v} W_{\eta_v} \sum_{\ell m} C_{\eta_v, \ell m}^{\ell_o m_o} \prod_{\xi=1}^v \sum_{j \in N_i} \mathbf{m}_{ij}$
MACE	$\sum_v \sum_{\eta_v} W_{\eta_v} \sum_{\ell m} C_{\eta_v, \ell m}^{\ell_o m_o} \prod_{\xi=1}^v \sum_{j \in N_i} W_\xi \mathbf{m}_{ij}$
Allegro	$\sum_{j \in N_i} \mathbf{m}_{ij}$

Update: The update functions used in higher body order methods are summarized in Table 15. Linear ACE does not need to update node features because it only has a single layer. MACE updates the features for each node using the aggregated message and the node feature from the previous layer. In Allegro, the aggregated message m_i is used to update two features of edge e_{ij} . The first is an equivariant feature v_{ij} obtained by the tensor product of the aggregated message m_i and feature v_{ij} from the previous layer. Then, the invariant part of v_{ij} is extracted to update the invariant feature x_{ij} for edge e_{ij} .

Table 15. The update functions used in higher body order methods.

Methods	Update
Linear ACE	-
MACE	$\mathbf{h}_i^t = W(\mathbf{h}_i^{t-1}) + W(m_i)$
Allegro	$\mathbf{v}_{ij}^t = \text{TP}(\mathbf{m}_i, \mathbf{v}_{ij}^{t-1})$ $\mathbf{x}_{ij}^t = \phi(\mathbf{x}_{ij}^{t-1} \ \mathbf{v}_{ij}^{t, \ell m=00})$

Output: The output modules of many-body methods differ from other equivariant methods. Higher body order methods first compute local energies for each atom, then the total energy of the molecule is the sum of local energies over all atoms. In Linear ACE, the invariant part of the aggregated message is extracted as the local energy for each atom. In MACE, the local energy of an atom is the sum of a fixed term and a learnable term. The fixed term is determined by atomic type and corresponds to the isolated energy of that atom, which is precomputed using DFT. The learnable term is derived from the invariant part of node features obtained in each layer. In Allegro, the invariant edge feature from the final layer is used to obtain pairwise energies. Next, pairwise energies around the central atom are scaled with a scaling factor that depends on both atom types, $\theta_{z_i z_j}$. These are then summed to obtain the local energy for the central atom. The output functions used in higher body order methods are summarized in Table 16.

Table 16. The output functions used in higher body order methods.

Methods	Output
Linear ACE	$\sum_i \mathbf{m}_i^{\ell m=00}$
MACE	$\sum_i [E_{\text{iso},i} + \sum_{t=1}^{T-1} W\mathbf{h}_i^{t, \ell m=00} + \text{MLP}(\mathbf{h}_i^{T, \ell m=00})]$
Allegro	$\sum_i \sum_{j \in \mathcal{N}_i} \theta_{z_i z_j} \text{MLP}(\mathbf{x}_{ij}^T)$

5.2.8 Model Outputs.

Both invariant and equivariant methods should be able to deal with the symmetries for different tasks and applications. Invariant methods can produce $SE(3)$ -invariant features directly, and some equivariant features may also be achieved based on the final invariant features. For example, in order to predict per-atom forces that are $SO(3)$ -equivariant, invariant methods first predict the energy E and then use the gradient of the energy w.r.t. atom positions $f_i = -\frac{\partial E}{\partial \mathbf{c}_i}$ to compute the force of each atom, which can ensure energy conservation. Here \mathbf{c}_i is the coordinate of node i . Equivariant methods can also use the predicted energy to compute forces or predict forces

Table 17. Statistics of QM9 [Ramakrishnan et al. 2014], MD17 [Chmiela et al. 2017], rMD17 [Christensen and Von Lilienfeld 2020], MD17@CCSD(T) [Chmiela et al. 2018], ISO17 [Schütt et al. 2018], and Molecule3D [Xu et al. 2021b] datasets. We summarize the prediction tasks and the number of 3D molecule samples (# Samples), maximum number of atoms in one molecule (Maximum # atoms), and average number of atoms in one molecule (Average # atoms).

Datasets	Prediction Tasks	# Samples	Maximum # atoms	Average # atoms
QM9	Predict energetic, electronic, and thermodynamic properties	130,831	29	18.0
MD17	Predict energy and force	-	-	-
rMD17	Predict energy and force	-	-	-
MD17@CCSD(T)	Predict energy and force	-	-	-
ISO17	Predict energy and force	645,000	19	19
Molecule3D	Predict 3D geometry and energetic and electronic properties	3,899,647	137	29.1

directly. For other equivariant prediction targets like the Hamiltonian matrix discussed in Section 4, additional operations are necessary for invariant models to ensure equivariance, making equivariant methods more straightforward and suitable for such tasks.

5.2.9 Datasets and Benchmarks.

Molecular representation learning methods are evaluated on various tasks, such as energy prediction and per-atom force prediction. Table 17 summarizes commonly used datasets, including QM9 [Ramakrishnan et al. 2014], MD17 [Chmiela et al. 2017], rMD17 [Christensen and Von Lilienfeld 2020], MD17@CCSD(T) [Chmiela et al. 2018], ISO17 [Schütt et al. 2018], and Molecule3D [Xu et al. 2021b]. Typically, the mean absolute error and mean square error between the predicted and ground-truth values is used as evaluation metrics.

Specifically, QM9 dataset [Ramakrishnan et al. 2014] collects more than 130k small organic molecules with up to nine heavy atoms (CONF) from GDB-17 database [Ruddigkeit et al. 2012]. For each molecule, the dataset provides its 3D geometry for the stable state (minimal in energy), along with corresponding harmonic frequencies, dipole moments, polarizabilities, energies, enthalpies, and free energies of atomization. All properties were calculated at the B3LYP/6-31G(2df,p) level of quantum chemistry. Typically, a separate model is trained for each property.

MD17 dataset [Chmiela et al. 2017] includes molecular dynamic simulations of 8 small organic molecules, namely, aspirin, benzene, ethanol, malonaldehyde, naphthalene, salicylic acid, toluene, and uracil. For each molecule, the dataset provides hundreds of thousands of conformations and corresponding energies and forces. Revised MD17 (rMD17) dataset [Christensen and Von Lilienfeld 2020] is a recomputed version of MD17 to reduce numerical noise. For each molecule in the original MD17, 100,000 structures are taken, and the energies and forces are recalculated at the PBE/def2-SVP level of theory using very tight SCF convergence and very dense DFT integration grid. Therefore, the dataset is practically free from numerical noise. MD17@CCSD(T) [Chmiela et al. 2018] is calculated based on the more accurate and expensive CCSD or CCSD(T) method and contains fewer molecules. Typically, for MD17, rMD17, ad MD17@CCSD(T), a separate model is trained for each molecule, with the task of predicting the energy and force for each conformation. To ensure energy conservation, most methods compute the per-atom force from the predicted energy, as discussed in Section 5.2.8, and the commonly used loss function is a combination of the energy loss and force loss

$$\mathcal{L} = \lambda_E \mathcal{L}_E(\hat{E}, E) + \lambda_f \mathcal{L}_f(-\frac{\partial \hat{E}}{\partial C}, f). \quad (98)$$

Here, \hat{E} is the predicted energy, $-\frac{\partial \hat{E}}{\partial C}$ is the computed force, C is the atom coordinate matrix, E and f are the ground-truth energy and force, \mathcal{L}_E and \mathcal{L}_f are energy and force loss functions, such as

mean absolute error and mean square error, λ_E and λ_f are the weights for energy and force losses, which are often set to 1 and 1000, respectively.

ISO17 [Schütt et al. 2018] differs from MD17 in that it includes both chemical and conformational changes. The dataset contains molecular dynamics trajectories of 129 isomers with the same composition of $C_7O_2H_{10}$. Each trajectory consists of 5,000 conformations, resulting in a total of 645,000 samples. Unlike MD17 where a separate model is usually trained for each molecule, for ISO17, a typical setting is that a single model is trained across all 129 different molecules.

Molecule3D [Xu et al. 2021b] is a large-scale dataset with around 4 million molecules curated from PubChemQC [Nakata and Shimazaki 2017]. For each molecule, the dataset provides its precise ground-state 3D geometry derived from DFT at the B3LYP/6-31G* level, as well as molecular properties such as energies of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), the HOMO-LUMO gap, and total energy. Although Molecule3D is primarily designed for predicting 3D geometries from 2D molecular graphs (Section 5.3), in this subsection, we can directly take the ground-truth 3D geometries as input and test models' performance on property prediction tasks. It is worth noting that PCQM4Mv2 dataset [Hu et al. 2020, 2021a] is curated from PubChemQC as well, but only provides 3D geometries for the training data. Therefore, it is often used for tasks such as predicting properties from 2D molecules [Ying et al. 2021], pre-training [Zaidi et al. 2023], and 2D-3D joint-training [Luo et al. 2023b].

In addition to the datasets mentioned above, some of the molecular representation learning methods discussed in this section are also evaluated on larger molecules such as proteins, materials, DNA, and RNA. Example datasets include MD22 [Chmiela et al. 2023], Atom3D [Townshend et al. 2020], and OC20 [Chanussot* et al. 2021].

5.2.10 Open Research Directions.

Learning from Both 2D and 3D Information: Despite recent advances in molecular representation learning, several challenges require further exploration. One direction is the joint-training from both 2D and 3D information of molecules [Stärk et al. 2022a; Luo et al. 2023b]. While 3D information is crucial for accurately modeling the physical properties of molecules, it can be computationally expensive to calculate and hard to obtain experimentally. On the other hand, 2D information, such as the molecular graph, is computationally efficient to generate, but may not capture all the necessary information for accurate predictions. Therefore, exploring methods that can be jointly trained on both 2D and 3D information or transfer knowledge from 3D to 2D representations could lead to improved performance as well as efficiency in tasks such as property prediction and drug discovery.

Expressivity and Computational Efficiency: On the theoretical front, a challenge is developing provably expressive 3D GNNs that capture geometric interactions among atoms in a *complete* or universal manner [Pozdnyakov et al. 2020], as elaborated in Section 2.9.3. Towards this goal, Joshi et al. [2023] provide a theoretical upper bound on the expressive power of geometric GNNs in terms of discriminating non-isomorphic geometric graphs, and show that equivariant layers which propagate geometric information are more expressive than invariant ones, in general. They identify key design choices for building maximally powerful equivariant GNNs: (1) depth, (2) tensor order, and (3) scalarization body order. As highlighted in this section, body order controls how well a network manages to capture the local geometry in a neighbourhood of a node, while higher tensor order enables a network to have higher angular resolution when representing geometric information. Finally, network depth controls the receptive field of an architecture, and in many current equivariant architectures increased depth implicitly also leads to increased body order.

While increasing all three properties theoretically improves the expressive power of geometric GNNs, several practical challenges hinder provably expressive models. Computing higher-order tensors [Passaro and Zitnick 2023] and many-body interactions [Batatia et al. 2022b] scales the compute cost drastically, often limiting practically used networks to tensor order $l \leq 2$ and body order $v \leq 4$. Further, there is early evidence of geometric oversquashing with increasing depth [Alon and Yahav 2021]. Future research may focus on improving the efficiency of many-body and higher-order equivariant GNNs to scale to larger biomolecules as well as larger datasets.

Invariant Versus Equivariant Message Passing: Invariant GNNs are significantly more scalable than equivariant GNNs and can be as powerful when working with fully connected graphs [Joshi 2020] and pre-computing non-local features [Gasteiger et al. 2021; Wang et al. 2022g]. Similarly, some invariant GNNs build canonical reference frames to convert equivariant quantities into scalar features [Du et al. 2022; Duval et al. 2023], allowing non-linearities on all intermediate representations in the network. Investigating the trade-offs between invariant and equivariant message passing is another fruitful avenue of research on molecular representation learning.

5.3 Molecular Conformer Generation

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Recommended Prerequisites: Section 5.2

As discussed in Section 5.2, the role of 3D molecular geometries in molecular representation learning is integral as they significantly enhance the accuracy of property prediction compared to the use of 2D graphs solely. This enhancement of 3D information is attributed to the fact that the physical configuration of a molecule largely influences its numerous properties. For example, isomers with identical atomic compositions can have vastly different melting points due to variations in their molecular structures. In immunology, the shape of antibodies' binding site, specifically the complementarity determining regions (CDRs), precisely determines the antigen they can recognize and bind to, which is critical for immune response. Hence, spatial information of molecules is highly desirable when working on real-world applications such as molecular property prediction, molecular dynamics, and molecule-protein docking. However, the acquisition of accurate 3D geometries through Density Functional Theory (DFT) is significantly challenging due to its high computational cost, thus limiting the widespread application of 3D molecular geometries. Consequently, the employment of machine learning models for the reconstruction of 3D molecular geometries emerges as a promising alternative, offering the potential to mitigate computational cost and make 3D geometries more accessible.

5.3.1 Problem Setup.

Let the total number of atoms in the molecule be n . A 2D molecule is represented as $\mathcal{G} = (z, E)$, where $z = [z_1, \dots, z_n] \in \mathbb{Z}^n$ denotes atom type vector, and each $e_{ij} \in \mathbb{Z}$ in E denotes the edge type between nodes i and j . For a given 2D molecule \mathcal{G} , the corresponding 3D molecule further needs 3D geometries $C = [c_1, \dots, c_n] \in \mathbb{R}^{3 \times n}$ where c_i denotes the 3D coordinate of the i -th atom. One form of C is associated with a potential energy, sampled from the potential energy surface corresponding to the Boltzmann distribution, which dictates that states of lower potential energy are more probable in a given environment. Geometries that correspond to lower energy or high probability states are generally more stable and thus, are more likely to be corroborated by experimental observations. The geometry that minimizes the potential energy or maximizes the distribution, known as the equilibrium ground-state geometry, is the most stable and critical one. The problem of molecular geometry reconstruction can be bifurcated into two distinct tasks. The

first task, referred to as 3D geometry generation, involves training a generative model, denoted as f_G , with the aim of understanding the distribution $p(C|G)$ of low-energy geometries given the conditional 2D molecular graph G . On the other hand, the second task, known as 3D geometry prediction, seeks to train a predictive model f_P that is capable of directly estimating the equilibrium ground-state geometry C_{eq} based on its corresponding 2D graph G .

5.3.2 Technical Challenges.

The reconstruction of 3D molecular geometries from 2D molecular graphs poses three major challenges. The first challenge is to ensure that the obtained conformers are geometrically valid in 3D space. For instance, it is possible for symmetric graph nodes to have identical embeddings due to the permutation invariance inherent to GNNs, leading to invalid geometries. Therefore, it is essential to distinguish these symmetric atoms and enforce their reconstructed coordinates are distinct because atoms should not overlap in 3D space. Besides, existing works [Simm and Hernández-Lobato 2019; Xu et al. 2021b] consider Distance Geometry (DG) first and then reconstruct atom coordinates based on the distance matrix. In such cases, ensuring the 3D geometric validity of atom coordinates becomes particularly challenging due to the potential for the derived distance matrix to fail in constituting a valid Euclidean Distance Matrix (EDM). In addition to maintaining the 3D geometric validity of reconstructed conformers, the second challenge is to meet the chemical validity imposed on conformer fragments. For instance, aromatic rings or π bonds restrict all their atoms on a planar surface, while many macrocycles and small rings are non-planar [Wang et al. 2020b]. It is desirable to have reconstructed geometries obeying such quantum rules and being chemically valid. An additional challenge in reconstructing 3D molecular geometry arises from the inherent symmetry of the geometry density function. Given initial systems with zero centers of mass (CoM) [Xu et al. 2022b; Köhler et al. 2020], the generative geometry distribution of conformers is often modeled as an invariant distribution in order to draw asymptotically unbiased samples with respect to the ground truth distribution [Köhler et al. 2020]. Specifically, we must ensure the reconstructed conformer is subject to $SE(3)$ -invariant position distributions. Let rotation matrix $R \in SO(3) \subset \mathbb{R}^{3 \times 3}$ and translation vector $t \in \mathbb{R}^3$ and $1 \in \mathbb{R}^n$, the $SE(3)$ -invariant position distributions require $p(RC + t1^T|G) = p(C|G)$. In other words, rotated or translated conformers are regarded as identical because geometry reconstruction is independent of rotation and translation.

5.3.3 Existing Methods.

Generation-Based Methods: While there are a plethora of molecular generative models available, this section focuses exclusively on those that represent recent and significant contributions to the field. Many earlier generative models, such as CVGAE [Mansimov et al. 2019], GraphDG [Simm and Hernández-Lobato 2019], ConfVAE [Xu et al. 2021d] and CGCF [Xu et al. 2021a], are developed based on variational autoencoders (VAEs) or flow model as their fundamental theory. On the other hand, current state-of-the-art generative models mostly rely on score matching and probabilistic denoising diffusion models with $E(3)$ -equivariant/invariant modules. For instance, ConfGF [Shi et al. 2021] develops a 3D generative model that uses score matching and obtains scores by computing chain rule derivatives from positions to distances. To generate the position of each atom in a molecule, ConfGF applies an $E(3)$ -equivariant model to update atom positions during sampling. GeoDiff [Xu et al. 2022b] further extends ConfGF’s capabilities by incorporating a zero-centering $E(3)$ -equivariant diffusion probabilistic model. Although both methods can directly generate 3D coordinates of molecules, they do not consider chemical constraints, such as aromatic rings in which all atoms are at the same plane. As a result, they may produce chemically invalid conformers. In contrast, torsional diffusion [Jing et al. 2022] employs an $SE(3)$ -invariant diffusion model only to adjust conformers’ torsions while retaining all local structures like bond length and angle

Table 18. Summary of 3D outputs, model architecture, and distribution symmetry of several representative 3D molecular conformation generation methods. Among the various methods, CVGAE [Mansimov et al. 2019], GraphDG [Simm and Hernández-Lobato 2019], ConfVAE [Xu et al. 2021d] use conditional variational autoencoder to generate molecular conformers, where CVGAE directly generates 3D coordinates, and GraphDG and ConfVAE generate interatomic distances of molecular conformers. In the spirit of ConfVAE, CGCF [Xu et al. 2021a] generates interatomic distances by taking advantage of the flow generative model. Moreover, ConfGF [Shi et al. 2021] and GeoDiff [Xu et al. 2022b] employ zero-centering $E(3)$ -equivariant models to directly generate 3D coordinates, and achieve an $E(3)$ -invariant generative distribution. Conversely, Torsional Diffusion [Jing et al. 2022] applies an $SE(3)$ -invariant diffusion model to generate torsions exclusively, preserving local structures such as bond length and angle, which are generated by RDKit. Additionally, GeoMol [Ganea et al. 2021], DeeperGCN-DAGNN+Distance [Xu et al. 2021b], and EMPNN [Xu et al. 2023c] implement predictive strategies for the generation of molecular conformers.

Methods	3D Outputs	Architecture	Distribution Symmetry
CVGAE	Coordinates	VAE	-
GraphDG	Distances	VAE	$E(3)$ -Invariant
ConfVAE	Distances	VAE	$E(3)$ -Invariant
CGCF	Distances	Flow	$E(3)$ -Invariant
ConfGF	Coordinates	Score Matching	$E(3)$ -Invariant
GeoDiff	Coordinates	Diffusion	$E(3)$ -Invariant
Torsional diffusion	Torsions	Diffusion	$SE(3)$ -Invariant
GeoMol	Coordinates	Predictive Model	$SE(3)$ -Invariant
EMPNN	Coordinates	Predictive Model	-
DeeperGCN-DAGNN+Dist	Distances	Predictive Model	$E(3)$ -Invariant

generated by RDKit. By doing so, it takes advantage of the chemical knowledge introduced by RDKit. However, this approach heavily depends on RDKit-generated conformers and cannot refine local ring structures, such as macrocycles or small non-planar rings. Based on torsional diffusion, DiffDock [Corso et al. 2022] advances protein-ligand docking as a conformation generation process conditioned on proteins. We describe the details of DiffDock in Section 8.

Prediction-Based Methods: Alternatively, other existing works formulate the task as a predictive task, which focuses on predicting the equilibrium ground-state conformer. One such example is GeoMol [Ganea et al. 2021], which employs message passing neural networks (MPNNs) with geometric constraints to predict local structures to generate diverse conformers. To ensure geometrical validity, GeoMol applies a matching loss that effectively distinguishes symmetric atoms by searching for the best matching substructures to ground truths among all possible permutations of symmetric nodes. Another notable approach is DeeperGCN-DAGNN+Distance, as proposed in [Xu et al. 2021b]. This method aims to predict the full distance matrix and then directly use the distance matrix in downstream tasks because pairwise distances implicitly provide 3D information. On the contrary, EMPNN [Xu et al. 2023c] uses node indices to break node symmetries and explicitly outputs geometrically valid 3D coordinates of the ground-state conformer.

To provide a comprehensive overview of the various generative and predictive methods in the field, we summarize representative approaches in Table 18.

5.3.4 Datasets and Benchmarks.

Geometric Ensemble Of Molecules (GEOM) [Axelrod and Gómez-Bombarelli 2022] is a dataset of high-quality molecular conformers, where the 3D molecular structures are first initialized by

RDKit [Landrum 2010] and then optimized by ORCA [Neese 2012] and CREST [Grimme 2019] programs. It contains two subsets, GEOM-QM9 and GEOM-Drugs, which are the commonly used benchmark datasets for evaluating the performance of different molecular conformation generation methods. GEOM-QM9 dataset contains 133,258 small organic molecules from the original QM9 dataset [Ramakrishnan et al. 2014]. All molecules in this dataset have up to 9 heavy atoms and 29 total atoms including hydrogen atoms, with very smaller molecular mass and few rotatable bonds. The atomic types of heavy atoms are limited to carbon, nitrogen, oxygen, and fluorine. By contrast, GEOM-Drugs contains larger drug-like molecules, with a mean of 44.4 atoms (24.9 heavy atoms) and a maximum of 181 atoms (91 heavy atoms). Most importantly, these large molecules contain significant flexibility, e.g., with an average of 6.5 and up to a maximum of 53 rotatable bonds, which is challenging for learning the molecular conformation generation models.

5.3.5 Open Research Directions.

While significant progress has been made in ML-based molecular conformation generation models, there remain several promising directions for future research. Firstly, the current benchmark datasets are all simulated in the vacuum environment, while in reality, molecular conformations are different in surrounding solvent environments. Future research could focus on incorporating solvent effects into the generative models, allowing them to generate conformations that reflect the realistic behavior of molecules in given solvent environments. Second, learning the conformation generative models often requires large amounts of training data. However, in many cases, limited data is available for specific classes of molecules or compounds. Future research could also explore transfer learning and few-shot learning techniques to leverage knowledge learned from a broader set of molecules and apply it to generate conformations for less-studied or novel compounds. This could significantly reduce the data requirements and improve the generalization capabilities of the models. Thirdly, existing methods mainly focus on the generation of low-energy conformers due to their stability. However, exploring molecular conformers in high-energy transition states (TS) is equally significant, as they are pivotal to the progress of chemical reactions [Choi 2023; Duan et al. 2023a]. Hence, future research could also concentrate on generating the TS structures for reactants and products, facilitating an enhanced understanding of the kinetics and mechanisms of chemical reactions. Addressing these directions will significantly advance the field and contribute to the development of more effective, efficient, and practically relevant molecular conformation generation methods.

5.4 Molecule Generation from Scratch

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Recommended Prerequisites: Section 5.2

In Section 5.2, we study the problem of property predictions from given molecules. However, some other real-world problems, such as designing novel molecules for drugs, require us to model the reverse process, *i.e.*, to obtain target molecules with given properties. Exhaustively searching target molecules in chemical space is impossible because the number of candidate molecules can be very large, *e.g.*, there are around 10^{33} drug-like molecules [Polishchuk et al. 2013] in estimation. Recently, the significant progress in deep generative learning has motivated many researchers to generate novel molecules with advanced deep generative models, including variational auto-encoders (VAEs) [Kingma and Welling 2014], generative adversarial networks (GANs) [Goodfellow et al. 2014a], flow models [Rezende and Mohamed 2015] and diffusion models [Ho et al. 2020]. Some early studies [Jin et al. 2018; You et al. 2018a; Shi* et al. 2020] generate molecules in the form of 2D

molecular graphs. However, these methods do not generate 3D coordinates of atoms in molecules, so they cannot distinguish molecules with the same 2D graphs but different 3D geometries, such as spatial isomers. Actually, many molecular properties, such as quantum properties or biological activities, are determined by 3D geometries of molecules. Hence, in this section, we focus on the 3D molecule generation problem.

5.4.1 Problem Setup.

We represent a 3D molecule with n atoms as $\mathcal{M} = (z, C)$, where $z = [z_1, \dots, z_n] \in \mathbb{Z}^n$ is the atom type vector and $C = [c_1, \dots, c_n] \in \mathbb{R}^{3 \times n}$ is the atom coordinate matrix. Here, for the i -th atom, z_i is its atomic number and c_i is its 3D Cartesian coordinate. Our target is to learn a probability distribution p over the 3D molecule space with generative models and sample novel 3D molecules from p . Note that different from the molecular conformer generation or prediction problem discussed in Section 5.3, we do not generate 3D molecules from any conditional inputs like 2D molecular graphs, but instead generate them from scratch.

5.4.2 Technical Challenges.

The central challenge in generating 3D molecules from scratch lies in achieving $SE(3)$ -invariance when generating 3D atom positions. In other words, the generative model should assign the same probability to \mathcal{M} and \mathcal{M}' if \mathcal{M}' can be obtained by rotating or translating \mathcal{M} in 3D space. Generally, there are two strategies to generate 3D molecular structures. First, generative models may directly use the coordinate matrix as the generation targets or outputs. But the challenge is that the probabilistic modeling for coordinate matrices should be carefully designed to ensure invariance to $SE(3)$ transformations. Second, instead of directly generating coordinates, generative models may take some $SE(3)$ -invariant 3D features, such as distances or angles, as generation targets. This strategy removes the necessity of explicitly considering $SE(3)$ -invariance in generative models, but requires the generated 3D features to have valid values and complete 3D structure information so that 3D atom coordinates can be reconstructed from them.

5.4.3 Existing Methods.

Three representative methods of directly generating coordinate matrices of 3D molecules are E-NFs [Satorras et al. 2021b], EDM [Hoogeboom et al. 2022], and GeoLDM [Xu et al. 2023a]. They adopt multiple strategies to achieve $E(3)$ -invariance, where $E(3)$ is a superset of $SE(3)$, including translation, rotation, and reflection. Specifically, to remove the freedom of translation, any 3D molecules are always zero-centered by reducing the centroid, *i.e.*, the averaged 3D coordinates over all atoms, from each column of the coordinate matrix before being passed into generative models. In other words, the probability density captured by these approaches is non-zero only on coordinate matrices with zero centroids. In addition, the probability density of zero-centered coordinates is calculated by their corresponding latent variables, which are subject to CoM-free Gaussian distribution [Köhler et al. 2020]. Mathematically, CoM-free Gaussian distribution ensures that the probability density is invariant to rotation and reflection. Flow and diffusion models are used to map between zero-centered coordinate matrices and latent prior variables in E-NFs and EDM, respectively. GeoLDM further proposes to first encode the zero-centered atoms into a zero-centered latent space, where each atom is represented with latent invariant features and latent equivariant coordinates. Then instead of the original coordinate matrices, GeoLDM learns to map between the latent variable and prior Gaussian distribution via latent diffusion models [Rombach et al. 2022].

In contrast to E-NFs and EDM, some methods implicitly generate 3D atom positions from $SE(3)$ -invariant features. To represent complete structural information of a 3D molecules, one alternative to the coordinate matrix is Euclidean distance matrix that contains distances between every pairwise

Table 19. Summary of 3D outputs, model architecture, generation pipeline, distribution symmetry of several representative 3D molecule generation methods. Among these methods, E-NFs [Satorras et al. 2021b], EDM [Hoogeboom et al. 2022], and GeoLDM [Xu et al. 2023a] directly generate 3D coordinates of atoms in molecules. They achieve $E(3)$ -invariant by zero-centering coordinates and using CoM-free Gaussian distribution. On the other hand, EDMNet [Hoffmann and Noé 2019], G-SchNet [Gebauer et al. 2019], and G-SphereNet [Luo and Ji 2022] implicitly generate 3D positions of atoms by distances, angles, and torsion angles that are invariant to rotations and translations.

Methods	3D Outputs	Architecture	Pipeline	Distribution Symmetry
E-NFs	Coordinates	Flow	One-shot	$E(3)$ -Invariant
EDM	Coordinates	Diffusion	One-shot	$E(3)$ -Invariant
EDMNet	Distances	GAN	One-shot	$E(3)$ -Invariant
G-SchNet	Distances	Autoregressive model	Autoregressive	$E(3)$ -Invariant
G-SphereNet	Distances + Angles + Torsion angles	Flow	Autoregressive	$SE(3)$ -Invariant
GeoLDM	Coordinates	Latent Diffusion	One-shot	$E(3)$ -Invariant

atoms in the molecule. EDMNet [Hoffmann and Noé 2019] is the first work studying generating 3D molecular structures in the form of Euclidean distance matrices. In EDMNet, various of novel loss functions are used to train a GAN model to generate valid Euclidean distance matrices so that 3D Cartesian coordinates can be successfully reconstructed. Different from one-shot methods like EDMNet, other methods adopt an autoregressive procedure to generate 3D molecules through step-by-step placing atoms in 3D space. Two representatives of autoregressive methods are G-SchNet [Gebauer et al. 2019] and G-SphereNet [Luo and Ji 2022]. In both methods, a complete 3D molecule is generated by multiple steps, and only one new atom is generated and placed to the local region of a reference atom at each generation step. Specifically, G-SchNet places the new atom to one of the candidate grid positions of the reference atom through sampling from distance distributions predicted by an autoregressive generative model. On the other hand, G-SphereNet generates distances, line angles, and torsion angles by autoregressive flow models to determine the relative position of the new atom to the reference atom. Because of the use of torsion angles, G-SphereNet captures $SE(3)$ -invariant distributions. We summarize the key information of discussed 3D molecule generation methods in Table 19. Note that among these discussed methods, only EDM can take molecular properties as conditional input and perform property-oriented generation, while other methods can only use implicit strategies to generate molecules with desirable properties, such as optimizing latent representations. Some other methods may consider more complicated conditional inputs, such as protein pockets [Ragoza et al. 2022; Liu et al. 2022c], which is introduced in Section 8.

5.4.4 Datasets and Benchmarks.

Two benchmark datasets, QM9 [Ramakrishnan et al. 2014] and GEOM-Drugs [Axelrod and Gómez-Bombarelli 2022], are commonly used to evaluate the performance of different 3D molecule generation methods. QM9 dataset collects more than 130k small organic molecules from GDB-17 [Rudigkeit et al. 2012] database. All molecules in QM9 have up to 9 heavy atoms (29 atoms including hydrogen atoms), and the element type of any heavy atom is always one of carbon, nitrogen, oxygen, and fluorine. The 3D atom coordinates of molecules in QM9 are calculated at the B3LYP/6-31G (2df, p) level of quantum chemistry by Gaussian software [Frisch et al. 2009]. In addition to QM9, GEOM-Drugs is another dataset used to evaluate the performance of 3D molecule generation methods in generating larger and more complicated drug molecules. It collects 430k drug-like molecules with up to 181 atoms. The 3D atom coordinates of molecules in GEOM-Drugs are first

initialized by RDKit [Landrum 2010], then optimized by ORCA [Neese 2012] and CREST [Grimme 2019] software. In both datasets, the 3D coordinates of atoms in molecules are calculated by DFT.

5.4.5 Open Research Directions.

Despite that a lot of 3D molecule generation methods have been proposed in recent years, there exist several challenges hampering them to generate practically useful 3D molecules. First, most existing methods consider $E(3)$ symmetries, not $SE(3)$ symmetries, so they are also invariant to reflection. This invariance should be avoided in many biological and chemical applications where generative models are expected to discriminate 3D molecules with different chiralities. Additionally, it is crucial for the generated 3D molecules to meet chemical constraints in 3D positions of some local structures so that they are chemically valid and synthesizable. For instance, all atoms in a benzene ring are restricted to be in the same plane. However, it still remains challenging and under-explored to design generative models that satisfy all chemical constraints.

5.5 Molecular Dynamics Simulation

Authors: Xiang Fu, Tommi Jaakkola

Recommended Prerequisites: Section 5.2

Since its development in the 1950s, molecular dynamics (MD) simulation has evolved into a well-established and valuable technique for gaining atomistic insights into a wide range of physical and biological systems [Alder and Wainwright 1959; Rahman 1964; Frenkel and Smit 2001; Schlick 2010; Tuckerman 2010]. Through MD simulations, researchers can effectively characterize the potential energy surface (PES) that underlies the system and calculate macro-level observables based on the resulting MD trajectories. These observables play a crucial role in determining important material properties, such as the diffusivity of battery materials [Webb et al. 2015], and provide valuable insights into physical mechanisms, such as protein folding kinetics [Lane et al. 2011; Lindorff-Larsen et al. 2011]. However, the practical applicability of MD simulations is limited due to their high computational cost. This cost arises from two main factors: Firstly, in many applications that demand high accuracy, the energy and forces must be determined using quantum chemistry methods, which involve approximately solving the computationally expensive Schrödinger equation (Section 4). Secondly, when studying large and intricate systems like polymers and proteins, extensive simulations spanning nanoseconds to milliseconds are often necessary to investigate specific physical processes, while the time step size required for numerical stability is often at the femtosecond level. Conducting such simulations, even with less accurate classical force fields, incurs substantial computational expenses. In recent years, machine learning (ML) approaches have shown promise to accelerate MD simulations substantially. This section provides a brief overview of some forefronts of ML methods applied to MD simulations, encompassing ML force fields, ML augmented sampling methods, and ML-based coarse-graining methods. While we categorize this subsection under AI for small molecules, it is important to note that MD simulation is a versatile computational technique applicable to a wide range of molecules, including small organic molecules, biological macromolecules, and materials.

5.5.1 Problem Setup.

Simulating molecular dynamics involves integrating Newton's equation of motion: $\frac{d^2\mathbf{x}}{dt^2} = \mathbf{m}^{-1}\mathbf{F}(\mathbf{x})$. The forces necessary for this integration are obtained by differentiating a potential energy function: $\mathbf{F}(\mathbf{x}) = -\frac{\partial E(\mathbf{x})}{\partial \mathbf{x}}$. Here, \mathbf{x} represents the state configuration, \mathbf{m} represents the mass of the atoms, and F and E represent the force and potential energy function, respectively. To replicate desired

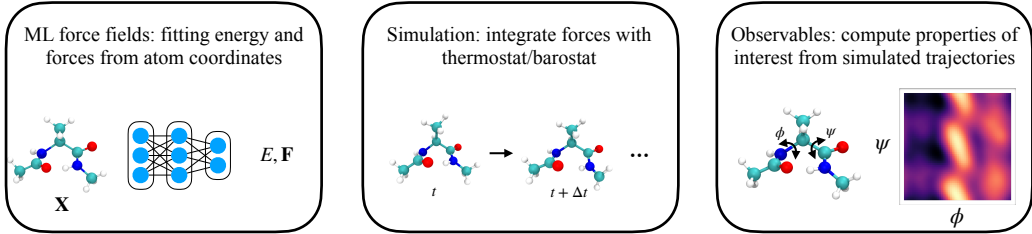


Fig. 21. Simulating molecular dynamics with machine learning. To replace expensive quantum mechanical calculations, an ML force field is learned to predict energy and forces from atomic coordinates. With a learned force field, we can simulate MD by pairing it with an appropriate thermostat/barostat. From the simulated trajectories, properties of interest can be computed.

thermodynamic conditions, such as constant temperature or pressure, an appropriate thermostat or barostat is selected to augment the equation of motion with additional variables. The choice of these conditions depends on the specific system and task at hand. Through the simulation, a time series of positions $\{\mathbf{x}_t \in \mathbb{R}^{N \times 3}\}_{t=0}^T$ (and velocities) is generated, where t denotes the temporal order index and T represents the total number of simulation steps, N is the number of atoms in the molecule. From the time series, observables $O(\mathbf{x}_t)$ such as radial distribution functions (RDFs), virial stress tensors, mean-squared displacement (MSD), and free energy surfaces with respect to key reaction coordinates, can be computed. These observables play a crucial role in studying the structural and dynamical properties of various physical and biological systems. Figure 21 summarizes the pipeline for using ML force fields (FFs) to simulate MD trajectories.

Obtaining the forces and energy for a given state requires classical or quantum mechanical calculations. While quantum mechanical calculation offers higher accuracy, it is computationally expensive. To accelerate MD simulation, one strategy is to fit a machine learning (ML) model that predicts $F(\mathbf{x})$ and $E(\mathbf{x})$ from the atomic coordinates. These models, known as machine learning force fields, are trained to approximate atom-wise forces and energies using a training dataset: $\{\mathbf{x}_i, \mathbf{F}_i, E_i\}_{i=1}^{N_{\text{data}}}$, where $\mathbf{x}_i \in \mathbb{R}^{N \times 3}$, $\mathbf{F}_i \in \mathbb{R}^{N \times 3}$, $E_i \in \mathbb{R}$, N_{data} is the number of data points. The learned force field can then be used to simulate molecular dynamics by replacing the computationally expensive quantum mechanical calculations for obtaining energy and forces.

In addition to ML force fields, it is important to note that the primary goal of MD simulation is to extract macroscopic observables that characterize system properties. Due to the chaotic nature of molecular dynamics, it is neither practical nor necessary to recover the trajectories given the initial states exactly. Therefore, many approaches focus on augmenting existing force fields to achieve more efficient sampling or coarse-graining that aims at reducing the system's complexity. The design of sampling and coarse-graining methods is often influenced by the specific system/observable of interest.

5.5.2 Technical Challenges.

First of all, the potential energy surfaces (PES) of molecular systems are often highly nonsmooth. Complex atomic interactions require expressive descriptors of the atomic environment. Ideally, the physical symmetry of energy ($E(3)$ -invariant) and forces ($E(3)$ -equivariant) should be respected. Expressive model architecture is a key technical problem in designing accurate ML force fields. The complex PES also poses technical challenges in effectively sampling diverse conformations, which motivates research in enhanced sampling methods. Secondly, while the simulation time step is usually at the femtosecond level, the observable of interest can often be at a much longer time

scale. Therefore, practically useful MD trajectories require simulations of millions to billions of steps to sample the dynamics. This practical need poses challenges to the efficiency, stability, and accuracy of the learned force fields. It is very hard to predict the performance of the learned force field in a simulation setting without actually running expensive simulations. Recent works have demonstrated that a lower force/energy prediction error does not imply a more stable and accurate simulation or observable calculation [Fu et al. 2023a]. The scale discrepancy between full-atom MD simulations and practical observables of interest also motivates research in coarse-graining approaches. In particular, the sampling of rare atomistic events is an important but difficult problem due to the combination of the two challenges stated above: the complex potential energy surface may have high energy barriers between different local minima, making rare events such as transitions between different metastable states hard to sample. Consequently, these transitions happen at a much longer time scale than the time scale a learned force field operates on. To summarize, the technical challenges of learning MD simulation root in the inherent complexity of the potential energy surface of atomistic systems and the computational complexity in calculating energy and forces for large spatiotemporal scales.

5.5.3 Existing Methods.

ML force fields [Behler and Parrinello 2007; Khorshidi and Peterson 2016; Smith et al. 2017; Artrith et al. 2017; Chmiela et al. 2017, 2018; Zhang et al. 2018a,b; Thomas et al. 2018; Jia et al. 2020; Gasteiger et al. 2020; Schoenholz and Cubuk 2020; Noé et al. 2020; Doerr et al. 2021; Kovács et al. 2021; Satorras et al. 2021b; Unke et al. 2021b; Park et al. 2021; Thölke and Fabritiis 2022; Gasteiger et al. 2021; Friederich et al. 2021; Liu et al. 2022f; Li et al. 2022d; Batzner et al. 2022; Takamoto et al. 2022b; Musaelian et al. 2023a] have attained incredible accuracy and data/compute efficiency that makes them promising for replacing quantum mechanical calculations in many applications. Different model architectures have been explored, including kernel-based methods, feed-forward neural networks, and message-passing neural networks. These models are designed to respect the physical symmetry principle, including the $E(3)$ -invariance of energy and $E(3)$ -equivariance of forces. Much of the molecular representation learning research has been motivated by MD applications. They have been covered with more details in Section 5.2.

In an effort to enhance the sampling process, machine learning (ML) methodologies have been employed to uncover crucial reaction coordinates [Sidky et al. 2020a; Mehdi et al. 2023] (also known as collective variables). These serve as prerequisites for implementing specific advanced sampling techniques, such as Meta Dynamics [Laio and Parrinello 2002; Barducci et al. 2008]. Identifying reaction coordinates can also play a significant role in elucidating the Molecular Dynamics (MD) process, notably in fitting a Markov state model for studying the transitions occurring between metastable states in protein molecules [Mardt et al. 2018]. Moreover, ML techniques are being harnessed for learning coarse-grained force fields [Husic et al. 2020; Wang et al. 2019b], coarse-grained and latent-space simulators [Fu et al. 2022b; Vlachas et al. 2021; Sidky et al. 2020b], and coarse-graining mapping [Wang and Gómez-Bombarelli 2019; Wang et al. 2022l; Köhler et al. 2023]. Simulation in the coarse-grained space is usually much more efficient but involves trade-offs over accuracy. Learning coarse-grained mapping encompasses the discovery of a coarse-graining scheme capable of preserving the essential information of the molecular state, as well as facilitating coarse-graining back mapping (predicting the distribution of fine-grained states corresponding to a coarse-grained state).

It's important to acknowledge that the research areas mentioned above possess extensive histories in their respective fields and continue to be vigorously developed. The materials and references explored here provide just a very preliminary overview. For a more in-depth understanding, we

direct interested readers towards more exhaustive surveys on these topics [Sidky et al. 2020a; Unke et al. 2021c; Noid 2023].

5.5.4 Datasets and Benchmarks.

Small molecules [Chmiela et al. 2017, 2023; Eastman et al. 2023] have been a popular testbed for ML force field development. The widely used MD17 dataset [Chmiela et al. 2017] contains MD data for eight small molecules generated from path-integral molecular dynamics simulations, with updated versions MD17@CCSD(T) [Chmiela et al. 2018] and rMD17 [Christensen and Von Lilienfeld 2020] that use higher levels of theory and are more accurate. Other systems of interest include bulk water [Zhang et al. 2018a], various crystalline solid materials (e.g., Li-ion electrolytes [Batzner et al. 2022]), and amorphous materials (e.g., polymer [Fu et al. 2022b]). With these datasets, force and energy prediction error over a test dataset is a common benchmarking strategy in existing work. Some papers [Stocker et al. 2022; Zhang et al. 2018a; Batzner et al. 2022] also study the stability of simulation and certain observables such as the distribution of interatomic distances, radial distribution function, diffusion coefficient, and so on. In particular, a recent benchmark study [Fu et al. 2023a] compared a series of existing ML force fields over a wide range of systems and tasks and found a misalignment between force/energy prediction performance and simulation performance, which shows the inefficacy of using force and energy prediction as the sole evaluation protocol.

For biomolecules, one focus of existing studies is to recover their free energy surface (FES) with respect to key reaction coordinates. Alanine dipeptide [Noé et al. 2020] is a standard benchmarking molecule due to its well-understood reaction coordinates and FES: there are two main conformational degrees of freedom: dihedral angle ϕ of C – N – C $_{\alpha}$ – C and dihedral angle ψ of N – C $_{\alpha}$ – C – N, with six FES minima over these two reaction coordinates. It has been studied in many papers focusing on sampling the Boltzmann distribution [Fu et al. 2023a], transition path sampling [Holdijk et al. 2022], and coarse-grained MD studies [Wang et al. 2019b; Vlachas et al. 2021; Greener and Jones 2021]. More complex biomolecules, such as the small protein Chignolin have also been studied in existing works [Husic et al. 2020; Wang et al. 2022]. For materials, past works have studied Li-ion battery electrolytes, such as LiPS [Batzner et al. 2022] and solid polymer electrolytes [Fu et al. 2022b], while looking at the radial distribution function and Li-ion diffusivity as the key observables. Finally, we note that MD simulation is a broad area with diverse applications and datasets available. We are only covering some of the most popular ones that were studied with ML methods.

5.5.5 Open Research Directions.

The precision and efficacy of current machine learning (ML) force fields present ample opportunities for further refinement. Predominantly, existing methodologies are built upon kernel-based or message-passing schemes across a graph formed with a predetermined radius cutoff. A promising avenue to enhance ML force fields' proficiency involves accurately and efficiently capturing long-range interactions. Given that the primary motivation behind ML force fields is to expedite Molecular Dynamics (MD) simulations, designing neural architectures that prioritize computational efficiency and parallelizability without compromising accuracy is critical. For instance, existing works have explored strictly local ML potentials [Musaelian et al. 2023a]. In practice, simulation instability is a recurrent issue when applying ML force fields. Active learning strategies [Vandermause et al. 2020; Ang et al. 2021] that focus on gathering new data from states where the learned model underperforms can help address these instability concerns and decrease the number of ground truth calculations required for training a dependable ML force field. As a crucial avenue for extending MD simulations to broader spatial and temporal scales, the implementation of coarse-graining

methods in both space (by converting atoms into coarse-grained beads) and time (by employing larger time steps, such as through learning time-integrated dynamics) are of great importance. Future research should strive to further comprehend and characterize the information retained and forfeited in a coarse-graining (CG) scheme. It's also imperative to explore ways to create more effective CG schemes that retain the maximum amount of information within a specified computational budget. Lastly, the endeavor to learn to sample rare events constitutes another vibrant area of research. In this domain, various methods such as ML-based collective variable discovery [Sidky et al. 2020a], transition path sampling [Holdijk et al. 2022], and deep generative models for modeling the Boltzmann distribution [Noé et al. 2019] represent promising directions to advance this research theme.

5.6 Learning Stereoisomerism and Conformational Flexibility

Authors: Keir Adams, Connor W. Coley

Recommended Prerequisites: Section 5.2

One potential advantage of 3D graph neural networks (GNNs) over their 2D counterparts is their capacity to natively model the structural differences between stereoisomers, molecules that share the same 2D molecular graph but have differing spatial arrangements of atoms in 3D. Stereoisomerism is commonly induced by tetrahedral chiral centers (e.g., carbon atoms with four non-equivalent bonded neighbors); double bonds with different E/Z (cis/trans) configurations; and chiral axes of atropisomers, allenes, or other helical molecules (Figure 22) [Eliel and Wilen 1994]. Notably, a molecular graph may have many different stereoisomers; a molecule with N tetrahedral chiral centers can have up to 2^N stereoisomers, without even considering E/Z isomerism or axial chirality. Two stereoisomers can be classified as either diastereomers, or enantiomers. Enantiomers are mirror-image chiral molecules that cannot be superimposed via thermodynamically-permissible conformational changes (e.g., rotations about chemical bonds). Diastereomers generally have distinct chemical properties altogether, while enantiomers exhibit identical physicochemical properties in many situations unless interacting with other chiral molecules (such as proteins), in which case they may exhibit wildly different properties [McConathy and Owens 2003; Chhabra et al. 2013]. Hence, the ability of graph neural networks to learn the subtle influence of stereoisomerism is crucial for practical applications across domains ranging from medicinal chemistry to chemical catalysis. Stereoisomerism has been overlooked as an aspect of molecular identity because the majority of benchmarks used for molecular property prediction do not require careful treatment given their high aleatoric uncertainty and underrepresentation of stereoisomers in the datasets.

Conformational isomerism is yet another form of stereochemistry which describes how a single molecule can adopt many different low-lying structures on the potential energy surface (PES), collectively called the conformer ensemble [Wolf 2007; Eliel and Wilen 1994]. Section 5.2 described representation learning on static and previously-known 3D molecular structures, such as DFT-optimized ground-state molecular geometries from the QM9 dataset [Ramakrishnan et al. 2014]. In reality, molecules are not static structures, but are instead constantly interconverting between different conformations through intramolecular motions such as chemical bond rotations and smaller vibrational perturbations. The energetic penalty for these motions is environment-dependent (e.g., solvent-dependent), and the rate of interconversion between conformers is highly temperature-dependent. For instance, at room temperature, cyclohexane undergoes a chair flip (10 kcal/mol energetic barrier) with a characteristic period of microseconds [Hendrickson 1961], whereas a bulky biaryl system like 1,1'-Binaphthyl interconverts between its (R)- and (S)- atropisomers (23 kcal/mol energetic barrier) on a time scale of hours [Meca et al. 2003]. Colloquially, two conformers

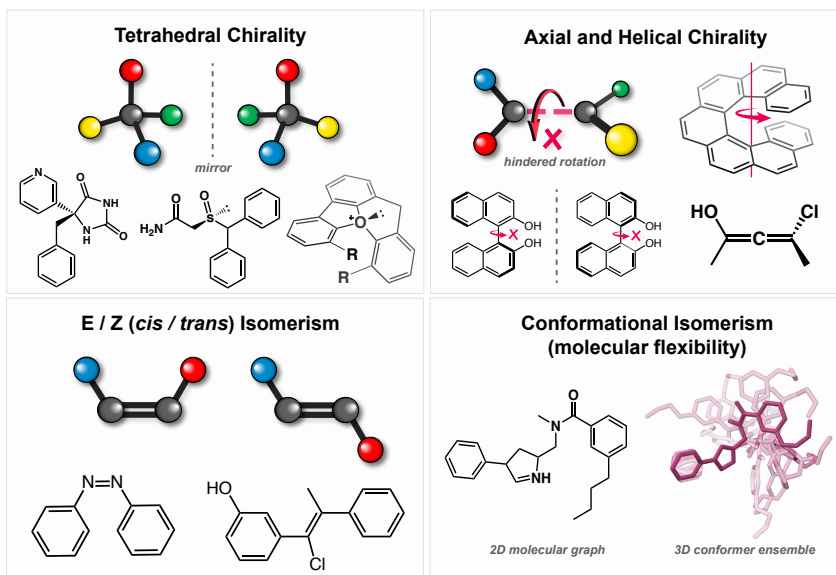


Fig. 22. Stereochemistry is an important yet often overlooked aspect of molecular identity that describes the differing orientations or arrangements of atoms in 3D space for molecules which share a common 2D graph topology. Stereoisomerism can be caused by a variety of local structural characteristics; shown here are common forms of stereochemistry that are particularly relevant for medicinal chemistry, chemical catalysis, and organic chemistry. *Tetrahedral chirality* (or point chirality) describes the differing orientations of four non-equivalent chemical groups around a stereogenic center. Tetrahedral chiral centers are inverted upon reflection, and hence induce enantiomerism. Tetrahedral chirality is often associated with carbon atoms, but can arise elsewhere, such as in sulfinyl or oxonium compounds. *Axial chirality* is caused by the (non-planar) orientations of four chemical substituents around a chiral axis. Axial chirality is found in allenes due to the propeller-like arrangement of adjoining double (π) bonds, and commonly in atropisomers, where bulky substituents restrict rotation about a single (σ) bond. Helical chirality is also a form of axial chirality, although its structural origin is different. Like tetrahedral chirality, axial chirality leads to enantiomerism. *E/Z isomerism* is caused by the differing *cis* or *trans* configurations of (planar) double bonds. Unlike tetrahedral or axial chirality, E/Z isomerism does not produce pairs of enantiomers. These forms of stereochemistry yield stereoisomers that typically cannot be interconverted on practical timescales without undergoing chemical reactions. Any given stereoisomer can also have a distribution of structurally-distinct but rapidly-interconverting conformational isomers, or *conformers*, owing to the molecule's 3D flexibility. Observable molecular properties are often related to thermodynamic averages over the entire conformer ensemble.

would be considered to belong to distinct “stereoisomers” if they do not thermally interconvert on a practical timescale (e.g., cannot be isolated at room temperature); two conformers that are mutually accessible would be described as corresponding to the same “stereoisomer”.

Many experimentally observable chemical properties depend on the full distribution of thermodynamically accessible conformers. On the other hand, some may depend on a particular (higher-energy) geometry that is not known *a priori*, such as the active binding pose of a ligand. The PES can also be substantially altered by intermolecular interactions (e.g., with solvent molecules), making it challenging to identify *a priori* which molecular structure(s) significantly contribute to observable properties without performing prohibitively expensive simulations. Although 3D GNNs have primarily been developed to encode individual 3D structures (Section 5.2), recent works have attempted to represent conformational flexibility by explicitly encoding conformer

ensembles [Axelrod and Gomez-Bombarelli 2020; Chuang and Keiser 2020]. This may be impactful for predicting distribution-dependent molecular properties such as Boltzmann-averaged ligand-protein binding affinities [Miller and Dill 1997; Gilson and Zhou 2007]; chemical reaction rates and selectivities [Hansen et al. 2016; Guan et al. 2018]; and entropic contributions to free energies [Mezei and Beveridge 1986; Chen et al. 2004].

5.6.1 Problem Setup.

For a given 2D molecular graph $\mathcal{G} = (\mathbf{z}, E)$, where \mathbf{z} is the vector of atom types (e.g., atomic numbers) and E denotes the graph adjacency matrix, we can formally describe its thermodynamically-accessible conformer ensemble as a set $C_{\mathcal{G}} = \{C_i\}_{i=1}^{|C_{\mathcal{G}}|}$ of structurally-distinct 3D molecular geometries $C_i \in \mathbb{R}^{3 \times n}$, each annotated with a (free) energy. Although the conformer ensemble is actually a continuous distribution, it is common to describe it with a discrete set of conformers by imposing a sub-Angstrom minimum root mean square distance (RMSD) threshold between any C_i and C_j . $C_{\mathcal{G}}$ can be divided into S disjoint subsets corresponding to the distribution of conformers available to each stereoisomer $C_{\mathcal{G}}^s = \{C_k^s\}_{k=1}^{|C_{\mathcal{G}}^s|}$ of the molecular graph so that $C_{\mathcal{G}} = C_{\mathcal{G}}^1 \cup C_{\mathcal{G}}^2 \cup \dots \cup C_{\mathcal{G}}^S$. The decision of which conformers belong to disjoint subsets can be somewhat subjective, but is typically based on their ability to interconvert on whatever timescale is relevant to the application at hand. Each conformer in a distribution can be assigned a statistical (Boltzmann) weight $p_{C_i^s} = \exp(\frac{-e_i}{k_B T}) / \sum_j \exp(\frac{-e_j}{k_B T})$ corresponding to its expected presence under experimental conditions, where e_i is the (free) energy of conformer C_i^s , k_B is the Boltzmann-constant, and T is the temperature. Some example stereochemical representation learning tasks include classifying a given conformer as one of many stereoisomers, contrasting the learned representations of conformers belonging to different stereoisomers, or training a supervised model to predict the properties of non-interconvertible stereoisomers from sampled conformers. This final supervised learning task aims to learn $\hat{f}(C_k^s \in C_{\mathcal{G}}^s; \theta) \approx f(C_k^s)$, where \hat{f} is a neural network with weights θ . Tasks related to learning on conformer ensembles include predicting Boltzmann-averaged properties $\langle y \rangle_{k_B} = \sum_i p_{C_i} f(C_i)$ from a small subset of the full conformer ensemble $\hat{f}(\{C_k^s\}_{k=1}^{K \ll |C_{\mathcal{G}}^s|}; \theta) \approx \langle y \rangle_{k_B}$, where $f(C_i)$ is a per-conformer property, or identifying a property-active conformer amongst a set of (non-active) decoy conformers.

5.6.2 Technical Challenges.

Learning molecular stereochemistry and conformational flexibility presents multifaceted modeling challenges. Because stereoisomers have the same molecular graph, 2D GNNs are inherently limited in their ability to distinguish stereoisomers with different chemical properties. Often, practitioners augment a molecular graph with simple atom (node) or bond (edge) features that store stereochemical information such as the handedness of chiral centers or the configuration of double bonds [Yang et al. 2019]. However, commonly used features (such as R/S chiral atom tags) are global properties that do not act in accordance with local graph convolutions [Pattanaik et al. 2020], have restricted representation learning power [Adams et al. 2021], and do not account for all forms of molecular chirality. 3D GNNs can also be limited in their ability to express certain stereochemistries according to their symmetry properties. For instance, many mathematically simple 3D GNNs with $E(3)$ -invariant features cannot distinguish the mirror-image structures of enantiomers. As a result, more complex networks with either 4-body interactions or equivariant features are often needed to robustly express chirality from 3D molecular structures [Liu et al. 2022f; Gasteiger et al. 2021; Thomas et al. 2018]. Further, adequately predicting properties of stereoisomers $f(C_{\mathcal{G}}^s)$ from a single 3D structure C_k^s requires the neural network to learn an invariance over 3D conformations to avoid confusing which conformers belong to which stereoisomer [Adams et al. 2021]. Meanwhile,

simultaneously encoding multiple conformers (at least) linearly scales the computational cost of training/inference while also making network optimization significantly more challenging [Axelrod and Gomez-Bombarelli 2020].

Modeling conformer ensembles and molecular flexibility raises additional challenges associated with the cost of obtaining high-quality conformer ensembles, especially at inference time. Namely, if a prediction model is trained with conformers obtained from expensive quantum chemical or molecular dynamics simulations, then the same simulations are likely required at inference time in order to avoid a domain shift reducing model accuracy. On the other hand, it may be difficult to accurately predict structure-sensitive properties of high-quality conformers when only encoding cheap conformers that are not faithful representations of the ground-truth conformers. For instance, it has been observed that encoding conformers optimized with molecular mechanics force fields to predict ground-state quantum properties of DFT-optimized molecules can lead to substantial loss in model accuracy compared to the case where ground-truth conformers are used directly [Stärk et al. 2022a; Pinheiro et al. 2022]. Similarly, it may be challenging to accurately predict properties of *unknown* property-active conformers when only modeling non-active conformers that are randomly sampled from the ensemble (e.g., predicting protein-ligand binding affinity without knowing the relevant ligand poses *a priori*). Although using a 2D GNN may side-step the challenges of obtaining quality conformers, 2D GNNs often cannot adequately learn functions that are highly sensitive to molecular geometry.

There are also challenges associated with collecting high-quality datasets for benchmarking and model development. When developing models to predict distribution-dependent properties obtained from simulated conformers, it is crucial that exhaustive conformer simulations are performed at a sufficiently high level of theory in order to avoid missing important (low-energy or property-active) conformers or assigning undue statistical weight to unrealistic geometries. These conformer searches should ideally be performed in a setting that reflects physical conditions, such as considering the influence of solvent molecules on the PES. Additionally, developing new models for stereochemical representation learning is often impeded by a lack of high-quality datasets that simultaneously 1) include properties of multiple stereoisomers for each molecular graph, 2) include properties that are sensitive to molecular stereochemistry, and 3) include properties with high signal-to-noise ratios.

5.6.3 Existing Methods.

Representing Molecular Stereochemistry: Existing works have chiefly focused on encoding tetrahedral chirality, and occasionally E/Z (or cis/trans) isomerism, through special tokens in molecular SMILES strings or through atom (node) and bond (edge) attributes in the 2D molecular graph [Yang et al. 2019]. SMILES strings can natively store the handedness of tetrahedral chiral centers via '@' and '@@' tokens that indicate whether the ordering of neighboring atoms (as provided in the string) is clockwise (CW) or counterclockwise (CCW). '\' and '/' tokens similarly store the configuration of double bonds. Hence, sequence encoders like transformers or recurrent neural networks can in principle express these forms of stereochemistry. Any 2D graph neural network that uses node and edge attributes may similarly include binary one-hot features that store the local configurations around tetrahedral centers or double bonds, based on a specified ordering of atoms. A related strategy creates graph convolution kernels that use the sign of the tetrahedral volume under a specific atom ordering [Liu et al. 2022g]. However, using atom or bond orderings that are sensitive to arbitrary bookkeeping breaks the permutation invariance of graph neural networks. Instead of relying on local atom orderings, the absolute R/S handedness of tetrahedral chiral centers and E/Z configuration of double bonds can instead be encoded via heuristic Cahn-Ingold-Prelog (CIP) rules that specify the global priority ranking of neighboring atoms [Cahn et al.

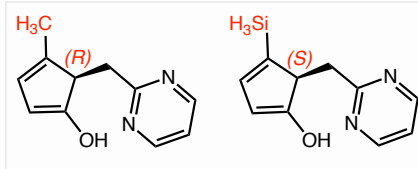


Fig. 23. Global (R/S) chiral atom tags do not conserve local 3D geometric features. Small changes to the molecular graph, such as swapping a carbon atom for a silicon atom, can flip the handedness of a tetrahedral chiral centers (according to CIP rules) without affecting the 3D molecular geometry or chemical reactivity.

1966]. However, small edits to a molecular graph (e.g., replacing a carbon atom with a silicon atom) can flip the parity of these global atom/bond tags without substantially changing the 3D molecular geometry (Figure 23), potentially making the learned stereochemical representations non-smooth.

Pattanaik et al. [2020] and Adams et al. [2021] introduce alternative methods of encoding tetrahedral chirality in graph neural networks without using heuristic rules or breaking permutation invariance. In each message passing layer of the GNN, Pattanaik et al. [2020] alters the sum-pooling operation (Equation (92)) to instead enumerate and encode all 12 permutations of the four neighboring atoms around each chiral center, thereby learning 2D atom representations that are sensitive to tetrahedral chirality. Adams et al. [2021] introduces ChIRo, which learns 3D representations of tetrahedral chirality by specially encoding the dihedral/torsion angles of each internal chemical bond. Although a conformer is required as input to the model, ChIRo is natively invariant to conformational changes caused by bond rotations, and using a cheap conformation generated by RDKit is shown to be sufficient. These approaches are empirically demonstrated to be superior to solely using local/global chiral atom tags when predicting chiral-dependent properties, at the expense of being computationally less efficient.

$SE(3)$ -invariant 3D GNNs, such as SphereNet [Liu et al. 2022f], can also learn representations that are sensitive to tetrahedral chirality. In this case, it is important that multiple conformers per molecule are used as training-time data augmentation in order to force the networks to learn an approximate conformer invariance [Adams et al. 2021]. However, in addition to being computationally demanding and requiring many conformers to train, 3D GNNs currently underperform in chiral property prediction tasks on small molecules compared to simply using 2D GNNs with chiral atom tags.

Representation Learning on Conformer Ensembles: The most common strategy for learning on conformer ensembles is to formulate the task as a multiple-instance learning (MIL) problem [Dietterich et al. 1997; Maron and Lozano-Pérez 1997; Ilse et al. 2018]. In this setting, multiple conformer instances of a given molecule are individually encoded into a set of fixed-length embeddings and then pooled or otherwise aggregated to obtain a single embedding of the entire conformer ensemble [Axelrod and Gomez-Bombarelli 2020; Chuang and Keiser 2020]. Formally, the ensemble embedding can be represented as

$$\begin{aligned} \mathbf{h}_k &= \hat{f}(C_k; \theta), \\ \mathbf{h}_{C_G} &= \sum_{k=1}^K a_k \mathbf{h}_k. \end{aligned} \tag{99}$$

Here, $\hat{f}(C_k; \theta)$ is any machine learning model that learns an embedding \mathbf{h}_k of a conformer C_k , such as a 3D GNN. For instance, Axelrod and Gomez-Bombarelli [2020] employs SchNet [Schütt et al. 2018], augmented with additional chemical features, as an underlying 3D GNN. Many other works employing MIL for molecular machine learning tasks have used non-neural models or

hand-crafted 3D feature vectors to encode \mathbf{h}_k [Zahrt et al. 2019; Zankov et al. 2021; Weinreich et al. 2021]. a_k can be set to a constant in order to weight each encoded conformer equally in the ensemble-level representation \mathbf{h}_{C_g} , as in sum- or mean-pooling. Alternatively, a_k can be learned attention coefficients that assign relative importance to each encoded conformer, which may be used to identify key conformer instances in the ensemble without needing to predict instance-level labels [Chuang and Keiser 2020]. Another approach uses max pooling to aggregate single-instance conformer representations [Liu et al. 2021a]. Because each conformer instance may be in a random reference frame, typically only $l = 0$ invariant representations are aggregated ($\mathbf{h}_k = \mathbf{h}_k^{l=0}$).

To avoid the cost of sampling or encoding multiple conformers at inference time, other works have attempted to implicitly model conformational flexibility by encoding a single “effective” structure obtained by averaging multiple conformers in structure-space [Weinreich et al. 2022], by learning conformer invariance via conformer-based data augmentation during training [Adams et al. 2021], or by considering multiple conformations during the collection of training labels [Suriana et al. 2023]. On the other hand, not explicitly encoding individual conformers from the ensemble may preclude the model from identifying key conformer instances, or otherwise reduce the model’s sensitivity to important 3D structures.

5.6.4 Datasets and Benchmarks.

Representing Molecular Stereochemistry: Enantiomers share common physio-chemical properties such as dipole moments or HOMO-LUMO gaps, meaning that popular benchmarks developed for 3D representation learning, like QM9 [Ramakrishnan et al. 2014] or PubChemQC [Nakata and Shimazaki 2017], cannot be used to evaluate the ability of models to learn chiral-sensitive representations even if the molecules in these datasets have well-defined chirality. On the other hand, biological measurements such as protein-ligand binding affinity or toxicity measurements can be influenced by chirality as well as other forms of stereochemistry. However, experimental datasets such as those contained in MoleculeNet [Wu et al. 2018] typically do not contain measurements for multiple stereoisomers of the same molecule, preventing straightforward evaluations of whether models learn the effects of stereochemistry. Further, the subtle effects of chirality may be obfuscated by experimental noise. As a result, existing works benchmark models on simulated datasets that have been specially curated to display acute sensitivity to molecular stereochemistry, particularly tetrahedral chirality.

Pattanaik et al. [2020] filters the D4 dopamine receptor protein-ligand docking screen performed by Lyu et al. [2019] to curate a dataset of 287,468 drug-like molecules with a Bemis-Murcko 1,3-dicyclohexylpropane scaffold that have at least one tetrahedral chiral center. Each molecule is also constrained to have a pair of enantiomers or diastereomers present in the dataset. They further subdivide this dataset into a subset containing enantiomer pairs with differences in docking scores above a threshold, and another subset containing enantiomer pairs with only one chiral center. Adams et al. [2021] also uses receptor-ligand docking to evaluate chirality-aware models. To control for stochasticity in docking simulations, they dock enantiomers from PubChem3D [Bolton et al. 2011] with low molecule weight and few rotatable bonds to a small docking box (PDB-ID: 4JBV), and only retain 34,560 pairs of enantiomers with differences in docking scores above a statistically significant threshold. In both works, models are evaluated based on their capability to correctly rank-order stereoisomer pairs by their predicted docking scores.

Beyond benchmarking on simulated datasets, Adams et al. [2021] and Mamede et al. [2020] curate datasets containing experimentally-measured optical activity (“L” versus “D” classifications) for one-chiral center enantiomers, sourced from the Reaxys database [Lawson et al. 2014]. “L” versus “D” classification is especially interesting as a benchmarking task because optical activity is difficult to simulate without expensive *ab initio* calculations, L/D labels have no correlation to

R/S chiral tags, and the optical rotation for one enantiomer can be directly inferred if its value is known for the other enantiomer. Moreover, predicting optical activity is practically useful for assigning the absolute configurations of chiral molecules. We envision that the collection and public dissemination of similar datasets containing chirality-sensitive properties of interest will be instrumental in furthering the field of chiral molecular representation learning.

Representation Learning on Conformer Ensembles: Few datasets have been developed to benchmark deep learning models on predicting the properties of conformer ensembles due to the resources required to obtain high-quality conformer ensembles and their associated properties for a large library of molecular compounds. [Axelrod and Gomez-Bombarelli \[2020\]](#) has benchmarked a handful of 2D, 3D, and 3D-ensemble models on their ability to classify bioactive hits from a library of 278,758 drug-like molecules from the GEOM-DRUGS dataset [[Axelrod and Gómez-Bombarelli 2022](#)], each annotated with experimental inhibition data against the SARS-CoV 3CL protease. Each molecule contains numerous conformers generated with the CREST program [[Pracht et al. 2020](#)], which are used to build models that encode ensembles each containing up to 200 conformers. On this task, models that explicitly encode multiple conformers in a multi-instance ensemble do not outperform baseline models that only encode a single conformation. The MIL models also require orders of magnitude more resources to train. It is important to note that this experimental dataset is very unbalanced, containing just 426 bioactive hits, which may complicate model optimization. [Chuang and Keiser \[2020\]](#) introduces a small synthetic dataset containing 1157 biaryl ligands with at most one rotatable bond, each containing an average of 13.8 conformers generated with OMEGA [[Hawkins et al. 2010](#)]. They evaluate the ability of a multi-instance attention model to identify whether an encoded ensemble contains a key conformer instance with a specific bidentate coordination geometry. On this toy task, however, simple random forest baselines using ECFP4 molecular fingerprints outperformed the MIL models.

Recent works have created new datasets containing large conformer ensembles that could potentially be used for conformer ensemble learning. In particular, [Grambow et al. \[2023\]](#) introduces the CREMP dataset, consisting conformer ensembles for 36,198 macrocyclic peptides generated with CREST. [Siebenmorgen et al. \[2023\]](#) introduces MISATO, a dataset containing 10-ns molecular dynamics traces for 16,972 protein-ligand complexes in explicit water solvent.

5.6.5 Open Research Directions.

Designing the next generation of geometric models to better represent molecular stereochemistry chiefly requires the development of new benchmarking datasets that contain multiple stereoisomers per molecule, labeled with properties that are sensitive to stereochemistry. In real-world applications, however, it may be impractical or infeasible to collect data on multiple stereoisomers due to experimental or computational budgets. Hence, another promising direction is to design models or training strategies that can learn stereochemistry-sensitive representations without needing to be exposed to multiple stereoisomers for each molecular graph during training time. Additionally, while current methods have focused solely on encoding tetrahedral chirality, future work could explore how to represent other forms of medicinally-relevant stereochemistry, such as atropisomerism.

Based on the limited number of existing studies, 3D conformer ensemble models are currently unable to outperform traditional 3D models that only encode a single conformer, despite the rich structural information contained in a conformer ensemble. Future work could investigate if this is due to the use of out-of-date 3D model architectures in existing studies, the inadequacy of the MIL framework to capture conformer flexibility, or the inherent difficulties of optimizing models that simultaneously encode structurally diverse ensembles, among other possible factors. The computational burden of both generating and encoding multiple conformers during training and inference also presents a practical barrier to the widespread adoption of MIL models for encoding

conformer ensembles. New strategies to efficiently account for conformational flexibility should be explored. Finally, because simulating high-quality conformer ensembles in physically realistic environments is often impractical for large virtual screens at inference time, future work could investigate the transferability of models that instead encode readily-available conformer ensembles obtained from inexpensive algorithms or generative models.

6 AI FOR PROTEIN SCIENCE

Proteins consist of a chain of amino acids at the primary level. They can fold into 3D structures to perform many biological functions. Recent breakthroughs in graph neural networks, diffusion models, and 3D geometric modeling enable the use of machine learning to boost the discovery of novel proteins. In this section, we focus on three AI for protein science topics, including protein folding (protein structure prediction) in Section 6.2, protein representation learning in Section 6.3, protein backbone structure generation in Section 6.4.

6.1 Overview

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We first give formal definitions of different levels of protein structures, and then introduce protein folding, protein representation learning, and protein backbone generation tasks. After that, we introduce geometric constraints that need to be considered for the latter two tasks. An overview of this section is shown in Figure 24.

We first describe the four-level structure of a protein. (1) Amino acid is the basic building block of protein. Proteins consist of chains of amino acids, with each amino acid containing nitrogen (N), alpha-carbon (C_α), carbon (C), and oxygen (O) atoms, as well as atoms in the side chain (known as R-group). The side chain determines the amino acid category. The amino acid chain is also known as the protein's primary structure; (2) based on the primary structure, secondary structures are locally folded structures that form based on interactions within the protein backbone; (3) tertiary structure is the three-dimensional structure of a single polypeptide chain (polypeptide chain refers to a string of amino acids connected together by peptide bonds); (4) quaternary structure describes the association between multiple polypeptide chains. They characterize the structure of a protein at different levels of complexity. In this paper, we mainly focus on primary and tertiary structures.

Notations of Protein Structures: Formally, a full-atom level protein structure can be represented as

$$\mathcal{P}_{\text{full}} = (\mathbf{z}, \mathbf{C}). \quad (100)$$

Here $\mathbf{z} = [z_1, \dots, z_n] \in \mathbb{Z}^n$ is the amino acid type vector, where each z_i denotes the type of the i -th amino acid and n denotes the number of amino acids in the protein. There are 20 commonly occurring amino acids that are used to build proteins in living organisms. The amino acid chain represents the primary structure of the protein and is folded into a 3D structure, where $\mathbf{C} = [C_1, \dots, C_n] \in \mathbb{R}^{n \times k \times 3}$ denotes the coordinate matrix of the protein. Note that distinct from other sections, we use \mathbf{C} to denote the coordinate matrix in this section in order to avoid notation conflict with carbon atom C . Each C_i includes the coordinates of all atoms in the amino acid i , including N , C_α , C , O , and side chain atoms. k is the maximum number of atoms in each amino acid. If we only consider the C_α atom in each amino acid, a protein structure can be represented as

$$\mathcal{P}_{\text{base}} = (\mathbf{z}, \mathbf{C}^{C_\alpha}), \quad (101)$$

where $\mathbf{C}^{C_\alpha} = [\mathbf{c}_1^{C_\alpha}, \dots, \mathbf{c}_n^{C_\alpha}] \in \mathbb{R}^{3 \times n}$ denotes the coordinate matrix of C_α atoms. Similarly, a protein backbone structure can be represented as

$$\mathcal{P}_{\text{bb}} = (\mathbf{z}, \mathbf{C}^{C_\alpha}, \mathbf{C}^N, \mathbf{C}^C), \quad (102)$$

where $\mathbf{C}^{C_\alpha} = [\mathbf{c}_1^{C_\alpha}, \dots, \mathbf{c}_n^{C_\alpha}] \in \mathbb{R}^{3 \times n}$, $\mathbf{C}^N = [\mathbf{c}_1^N, \dots, \mathbf{c}_n^N] \in \mathbb{R}^{3 \times n}$, and $\mathbf{C}^C = [\mathbf{c}_1^C, \dots, \mathbf{c}_n^C] \in \mathbb{R}^{3 \times n}$ denote coordinate matrices of C_α , N , C atoms. In the following parts, \mathcal{P} is used to denote a general representation of protein structures.

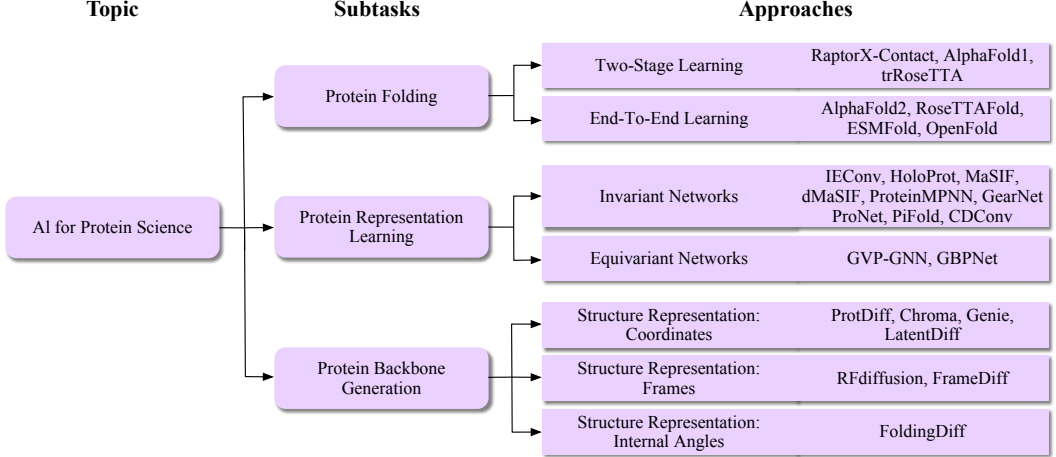


Fig. 24. An overview of the tasks and methods in AI for protein science. In this section, we focus on three subtasks, including protein folding, protein representation learning, and protein backbone generation. The methods for protein folding can be categorized into two classes, before and after AlphaFold2 [Jumper et al. 2021]: (1) two-stage learning: RaptorX-Contact [Wang et al. 2017a], AlphaFold1 [Senior et al. 2020], trRoseTTA [Du et al. 2021], (2) end-to-end learning: AlphaFold2 [Jumper et al. 2021], RoseTTAFold [Baek et al. 2021], ESMFold [Lin et al. 2023a], OpenFold [Ahdrizt et al. 2022]. The methods for protein representation learning are grouped into the invariant networks, including IEConv [Hermosilla et al. 2021], HoloProt [Somnath et al. 2021], MaSIF [Gainza et al. 2020, 2023], dMaSIF [Sverrisson et al. 2021], ProteinMPNN [Dauparas et al. 2022], GearNet [Zhang et al. 2023d], ProNet [Wang et al. 2023b], PiFold [Gao et al. 2023], and CDCConv [Fan et al. 2023], and equivariant networks, including GVP-GNN [Jing et al. 2021] and GBPNet [Aykent and Xia 2022]. For protein backbone generation, the methods are grouped in terms of structure representations they use. Specifically, ProtDiff [Wu et al. 2022d], Chroma [Ingraham et al. 2022], LatentDiff [Fu et al. 2023b], and Genie [Lin and AlQuraishi 2023] use 3D Euclidean coordinates as the structure representation for protein backbone structure, while RFdiffusion [Watson et al. 2022] and FrameDiff [Yim et al. 2023] use frame representations. Besides, FoldingDiff [Wu et al. 2022d] uses internal angles to represent protein backbone structures.

Protein Folding: The three-dimensional (3D) geometric structure of proteins plays a crucial role in determining their function. The specific arrangement and spatial organization of atoms within a protein molecule are essential for its interactions with other molecules, such as substrates, cofactors, ligands, and other proteins. Traditional X-ray crystallography is indeed considered an expensive and resource-intensive method for determining protein structures [Ilari and Savino 2008]. Machine learning methods were proposed automatically predict the protein structure based on the amino acid sequence. Protein folding, also known as protein structure prediction, aims to predict protein 3D structure (coordinates of all the atoms in both backbone and side chain, denoted C in Equation (100)) based on the amino acid sequence z .

Protein Representation Learning: Protein representation learning aims to learn informative representations for protein structures. The learned representations can be used for a wide range of

predication tasks, including enzyme reaction classification [Webb et al. 1992; Hermosilla et al. 2021; Hermosilla and Ropinski 2022; Zhang et al. 2023d; Fan et al. 2023], protein inverse folding [Ingraham et al. 2019; Jing et al. 2021; Hsu et al. 2022; Dauparas et al. 2022; Gao et al. 2023], and protein-ligand binding affinity prediction [Wang et al. 2004; Liu et al. 2015; Öztürk et al. 2018; Karimi et al. 2019; Somnath et al. 2021; Wang et al. 2023b], as shown in Figure 26. Protein representation learning can significantly speed up the processes of protein screening and new protein discovery.

Protein Backbone Structure Generation: As described above, a protein backbone consists of a chain of amino acid backbones, each of which contains the nitrogen (N), alpha-carbon (C_α), and carbon (C) atoms. These backbone atoms determine the secondary structure and overall shape of a protein, significantly affecting the protein functions. Hence, generating protein backbones is of great importance in *de novo* protein design. Specifically, the protein backbone generation task is to learn a generative model p_θ that can model the density distribution of real protein backbones $p_{\mathcal{P}_{bb}}$ and then we can sample a novel protein backbone $\hat{\mathcal{P}}_{bb}$ that satisfies $p_\theta(\hat{\mathcal{P}}_{bb}) \approx p_{\mathcal{P}_{bb}}(\hat{\mathcal{P}}_{bb})$. In practice, instead of jointly modeling the density of protein backbone atom positions and amino acid types, most studies formulate this problem as a conditional generation task, where atom positions are first sampled from the learned generative model, and then amino acid types are predicted from generated structures using a trained inverse folding model.

The goal for the protein backbone structure generation task is to create a model distribution that can easily generate samples to imitate the real data distribution. However, several challenges must be addressed to achieve this goal, including establishing a bijective mapping between data distribution and prior distributions such as Gaussians, ensuring distribution $E(3)/SE(3)$ -invariance, employing $E(3)/SE(3)$ -equivariant message passing, and efficient modeling of protein structures.

In this survey, for protein representation learning and protein backbone generation tasks, we mainly focus on structure-based instead of sequence-based predictive and generative methods for the following reasons. Firstly, protein representation learning is a complex task that requires the consideration of both protein structure and sequence. A significant proportion of protein functionalities are influenced by the structure and cannot be deduced directly from the sequence alone. And changes in the structure can result in different properties for the same protein sequence. Secondly, in protein generation, a key objective is to generate new protein structures that meet specific structure constraints, such as containing specific sub-structures, possessing particular secondary structures, and binding to particular molecules and antigens. These geometric constraints can be incorporated into protein structure generation methods as conditions but cannot be directly addressed from the protein sequence generation perspective. Particularly, protein generation using deep learning approaches is largely under-explored, and there is no much work published on this topic. Recent research trend shows diffusion models have great capacity and achieve the best performance. Thus, regarding deep learning approaches, we focus on diffusion models in this survey.

6.2 Protein Folding

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Different from small molecules that consist of a few atoms discussed in Section 5, proteins are macromolecules composed of a large number of atoms (mostly 1,000 to 10,000), posing greater challenges in estimating their native structure. In this section, we first formulate the protein folding problem, then identify the major challenges and discuss the existing methods and datasets. Finally, we point out a couple of potential directions for future work.

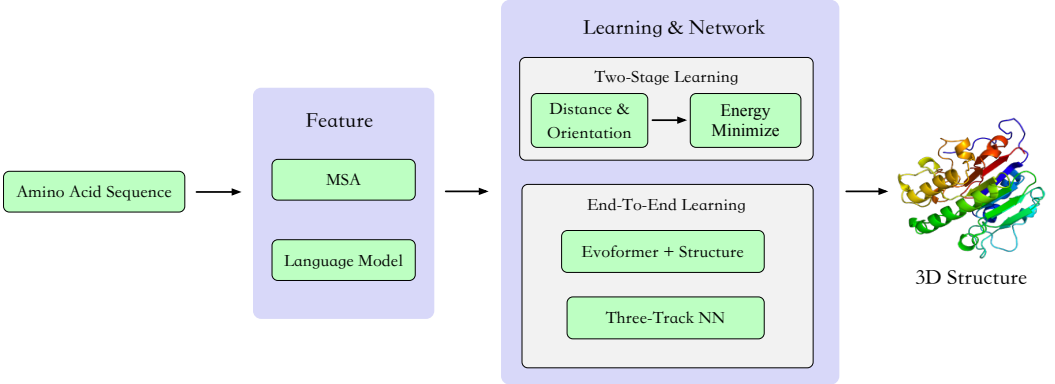


Fig. 25. Summarization of the protein folding algorithms. Existing methods, including RaptorX-Contact [Wang et al. 2017a], AlphaFold1 [Senior et al. 2020], trRoseTTA [Du et al. 2021], AlphaFold2 [Jumper et al. 2021], RoseTTAFold [Baek et al. 2021], AlphaFold-Multimer [Evans et al. 2021], ESMFold [Lin et al. 2023a], and OpenFold [Ahdritz et al. 2022], follow this pipeline and their respective modules are summarized in Table 20.

6.2.1 Problem Setup.

Protein folding, also known as protein structure prediction, aims to predict protein 3D structure (including coordinates of all the atoms in both backbone and side chain, denoted C in Equation (100)) based on the amino acid sequence z .

6.2.2 Technical Challenges.

Generator with $E(3)/SE(3)$ -Equivariance: The neural network is designed to maintain consistency when applied to protein structures undergoing $SE(3)$ transformations.

Physical Constraints: Protein folding is governed by fundamental physical principles that dictate the spatial arrangement of atoms within the protein structure. One of these principles is the concept of bond lengths, which refers to the average distances between atoms participating in chemical bonds. In protein molecules, the distances between bonded atoms are relatively fixed, meaning they have characteristic and well-defined values. These fixed bond lengths are determined by the types of atoms involved and the specific chemical bonds formed between them. Another principle is the distance between arbitrary paired atoms can not be too short to avoid a clash. It is necessary to incorporate these physical constraints into end-to-end models.

Computational Efficiency: Proteins can adopt an astronomical number of possible structures due to the flexibility of their backbone and side chains. Exploring this vast structural space to identify the most energetically favorable folded state is computationally demanding.

6.2.3 Existing Methods.

Existing work on protein folding can be classified into two categories, known as two-stage prediction and end-to-end prediction. Table 20 and Figure 25 summarize the major difference between existing approaches. Before the development of geometric deep learning, to circumvent the straightforward generation of 3D coordinates, most of the earlier methods leverage a two-stage learning process: the first stage is to predict the pairwise distance and orientation (e.g., torsion angles), while the second stage is to design a differentiable potential function as the optimization surrogate. The pairwise distance and orientation are invariant under $SE(3)$ transformation. Prominent approaches include RaptorX-Contact [Wang et al. 2017a], AlphaFold1 [Senior et al. 2020], trRosetta [Du et al.

2021], *etc.*, and are essentially non-end-to-end approaches. Then, the emergence of geometric deep learning, especially $E(3)/SE(3)$ neural networks, enables the buildup of an end-to-end system for protein structure prediction. Specifically, in 2020, AlphaFold2 [Jumper et al. 2021] has demonstrated remarkable accuracy in predicting the 3D structures of proteins in the 14-th CASP (Critical Assessment of Structure Prediction) competition (a biennial community-wide competition in the field of protein structure prediction). It concatenates Evoformer (a variant of the transformer) and the $SE(3)$ -equivariant structure module. The structure module aims to transform the representation into a 3D structure. It first generates the coordinates of the backbone sequentially: For each residue, instead of the global coordinate, it produces the relative position to the previous residue, which is parameterized by a rotation matrix (three learnable parameters) and transition vector (three learnable parameters). It was followed by a couple of works, including RoseTTAFold [Baek et al. 2021], AlphaFold-Multimer [Evans et al. 2021], ESMFold [Lin et al. 2023a], UniFold [Li et al. 2022c], OpenFold [Ahdritz et al. 2022], *etc.*

Specifically, drawing inspiration from AlphaFold2, RoseTTAFold [Lin et al. 2023a] introduced an innovative three-track neural network, enabling the joint modeling of 1D protein sequence, 2D distance map, and 3D coordinate information. By adopting this approach, impressive precision was achieved in predicting protein folding structures, on par with the performance of AlphaFold2. However, the original implementation of protein structure prediction is prohibitively time-consuming and resource-intensive. One major computational bottleneck lies in multiple sequence alignment (MSA). MSA is used for almost all the protein folding methods (before and after AlphaFold2) and plays a critical role in the final performance. The primary motivation for performing MSA is to identify and understand the functional and structural constraints on biological molecules. By aligning sequences from different species or within a single organism, researchers can identify conserved regions that are critical for maintaining the function of the molecule. MSA exhaustively searches over large-scale protein structure databases to identify similar amino acid sequences to reveal insight of biological evolutionary relationships and enhance the input feature. However, the MSA procedure is typically time-consuming and resource-demanding due to its brute-force essence. To alleviate this issue, ESMFold [Lin et al. 2023a] pretrains a large language model on amino acid sequences and uses it to replace MSA with a powerful neural representation, which is shown to accelerate the whole process significantly. The other neural architectures of ESMFold follow AlphaFold2. OpenFold [Ahdritz et al. 2022] develops a memory-efficient version of AlphaFold2, and curates OpenProteinSet, one of the largest public MSA databases (five million protein structures). In addition, OpenFold releases the code to benefit the whole community. AlphaFold-Multimer [Evans et al. 2021] enhances the prediction performance of AlphaFold in the context of multi-chain protein complex structure via incorporating more multi-chain proteins in training data.

6.2.4 Datasets and Benchmarks.

Protein Data Bank (PDB) [Berman et al. 2000] is the most well-known public protein structure database, where the protein structure is determined by X-ray crystallography [Ilari and Savino 2008]. The PDB collects, validates, and disseminates experimentally determined atomic coordinates and related information, such as experimental methods, resolution, and bibliographic references. The PDB houses over 180,000 protein structures, and the number of structures in the PDB is constantly growing as researchers determine and deposit new structures. For AlphaFold2, the dataset comes from two sources. Specifically, 75% of the training samples are from a self-distillation dataset from Uniclust30 [Mirdita et al. 2017]; 25% are from protein data bank (PDB) [Berman et al. 2000]. It removes some repetitive samples, and the final dataset contains around 475K protein structures.

The Critical Assessment of Protein Structure Prediction (CASP) is a biennial competition that aims to evaluate state-of-the-art methods in protein structure prediction. CASP provides a platform

Table 20. Summary of existing protein folding approaches, including RaptorX-Contact [Wang et al. 2017a], AlphaFold1 [Senior et al. 2020], trRoseTTA [Du et al. 2021], AlphaFold2 [Jumper et al. 2021], RoseTTAFold [Baek et al. 2021], AlphaFold-Multimer [Evans et al. 2021], UniFold [Li et al. 2022c], ESMFold [Lin et al. 2023a], and OpenFold [Ahdritz et al. 2022]. Two-stage learning typically consists of (1) prediction of pairwise distance and orientation and (2) energy minimization.

Methods	Feature	Learning	Network	Symmetry
RaptorX-Contact	MSA	Two-Stage	Residual CNN	$SE(3)$ -Invariant
AlphaFold1	MSA	Two-Stage	Residual CNN	$SE(3)$ -Invariant
trRoseTTA	MSA	Two-Stage	Residual CNN	$SE(3)$ -Invariant
AlphaFold2	MSA	End-To-End	Evoformer + Structure	$SE(3)$ -Equivariant
RoseTTAFold	MSA	End-To-End	Three-Track NN	$SE(3)$ -Equivariant
UniFold	MSA	End-To-End	Evoformer + Structure	$SE(3)$ -Equivariant
ESMFold	Language model	End-To-End	Evoformer + Structure	$SE(3)$ -Equivariant
OpenFold	MSA	End-To-End	Evoformer + Structure	$SE(3)$ -Equivariant
AlphaFold-Multimer	MSA	End-To-End	Evoformer + Structure	$SE(3)$ -Equivariant

for researchers and computational methods to assess their ability to predict the three-dimensional structure of proteins accurately. During CASP, participants are given a set of protein sequences for which the experimental structures have been determined but are kept confidential. The participants use their computational methods to predict the corresponding protein structures without any prior knowledge of the experimental structures. These predictions are then evaluated and compared to the experimental structures to assess the accuracy and quality of the predictions. AlphaFold1 dominates CASP13, which was held in 2018. After two years, in CASP14 held in 2020, AlphaFold2 achieved nearly 90 Global Distance Test (GDT) scores, roughly equivalent to X-ray crystallography’s accuracy. It became the first computational method to predict protein structures with near experimental accuracy and is called the gold standard of protein folding. Many follow-up works focus on reproducing AlphaFold2 and matching its performance, including RoseTTAFold [Baek et al. 2021], OpenFold [Ahdritz et al. 2022], ESMFold [Lin et al. 2023a], etc. For example, thanks to the use of the large language model instead of MSA, ESMFold achieves up to 60 \times speedup while maintaining accuracy [Lin et al. 2023a].

6.2.5 Open Research Directions.

A couple of challenges remain unsolved and hinder the practical use of protein folding. First, most current methods can accurately predict the structures of proteins with single chains and would degrade significantly for multi-chain proteins. The key reason is a multi-chain protein is more complex than a single-chain one with limited available data. Second, most of the current methods rely heavily on MSA and degrade significantly when dealing with proteins that are different from the training set and MSA database. Thus, enhancing the generalization ability to dissimilar proteins is a critical problem. Third, in the real world, proteins do not exist in isolation but often interact with other proteins, RNA, and DNA. Therefore, predicting protein structures in different contexts (e.g., RNA-protein complex, DNA-protein complex) is also an important challenge, where the available data is also limited. So, future research aims to predict the structure of a protein in these complicated scenarios.

6.3 Protein Representation Learning

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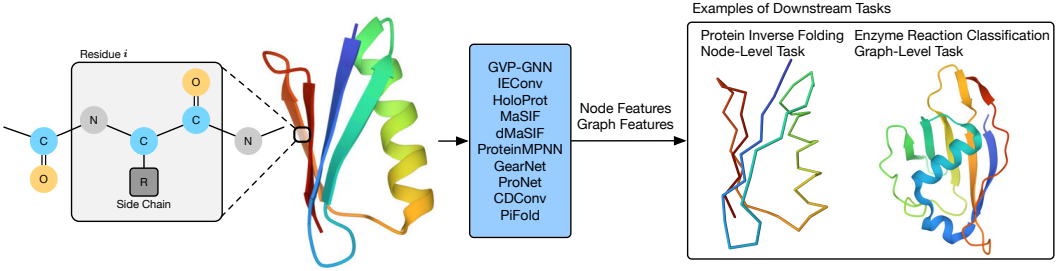


Fig. 26. Illustrations of protein structures, representation learning methods, and examples of downstream tasks. Proteins consist of one or more amino acid chains. When two or more amino acids bond to form a peptide, water molecules are removed, and the remaining part of each amino acid is called an amino-acid residue. The left part of the figure shows the detailed structure of a residue. In the middle, we list existing protein representation learning methods, including GVP-GNN [Jing et al. 2021], IEConv [Hermosilla et al. 2021], HoloProt [Somnath et al. 2021], MaSIF [Gainza et al. 2020, 2023], dMaSIF [Sverrisson et al. 2021], ProteinMPNN [Dauparas et al. 2022], GearNet [Zhang et al. 2023d], ProNet [Wang et al. 2023b], CDConv [Fan et al. 2023], and PiFold [Gao et al. 2023]. Different downstream tasks are shown in the right, including node-level tasks such as protein inverse folding [Ingraham et al. 2019] and graph-level tasks such as enzyme reaction prediction [Hermosilla et al. 2021].

Recommended Prerequisites: Sections 2.3, 2.4, 5.2

Different from small molecules discussed in Section 5, proteins are macromolecules with a large number of atoms, making protein representation learning more challenging. In this section, we highlight the challenges associated with protein representation learning. We also summarize existing methods designed specifically for protein structural learning.

6.3.1 Problem Setup.

The objective of protein representation learning is to learn a suitable representation that can encode important information about the given protein sequence and structure. Well-learned protein representations can be used to facilitate many downstream tasks, such as protein property prediction tasks that we focus on in this survey. Specifically, protein property prediction tasks can be classified into two categories, known as protein-level tasks and node-level tasks. For protein-level tasks like enzyme reaction classification, we aim to learn a function f to predict the property y of any given protein \mathcal{P} , and y can be a real number (regression problem) or categorical number (classification problem). For node-level tasks like inverse protein folding, we aim to learn a function f to predict the property y_i of the i -th amino acid.

6.3.2 Technical Challenges.

Complex protein structures pose several significant challenges that need to be addressed for protein representation learning as follows.

Computational Efficiency: A significant challenge in protein representation learning lies in the size of proteins. Proteins can contain hundreds or even thousands of amino acids, which makes them much larger than small molecules. As a result, computational efficiency is a critical bottleneck. Effective methods are expected to address this challenge by efficiently handling the large size of proteins without compromising the prediction accuracy.

Multi-Level Structures: Proteins are complex molecules made up of amino acids, and each amino acid comprises several atoms, as shown in Figure 26. Therefore, methods should capture the amino

Table 21. Summary of existing protein learning methods, including GearNet [Zhang et al. 2023d], CDConv [Fan et al. 2023], GVP-GNN [Jing et al. 2021], ProteinMPNN [Dauparas et al. 2022], PiFold [Gao et al. 2023], IEConv [Hermosilla et al. 2021], ProNet [Wang et al. 2023b], HoloProt [Somnath et al. 2021], MaSIF [Gainza et al. 2020, 2023], and dMaSIF [Sverrisson et al. 2021]. The complexity of a method is typically influenced by the number of nodes in the corresponding graph. Different methods can incorporate different levels of protein structures and have varying expressive power, which affects their ability to distinguish different protein structures.

Methods	Node (Complexity)	Level of Structures	Network	Symmetry
GearNet	Amino acid	C_α	$\ell = 0$	$E(3)$ -Invariant
CDConv	Amino acid + Pooling	C_α	$\ell = 0$	$E(3)$ -Invariant
GVP-GNN	Amino acid	Backbone	$\ell \leq 1$	$E(3)$ -Equivariant
ProteinMPNN	Amino acid	Backbone	$\ell = 0$	$E(3)$ -Invariant
PiFold	Amino acid	Backbone	$\ell = 0$	$SE(3)$ -Invariant
IEConv	Atom + Pooling	All-Atom	$\ell = 0$	$E(3)$ -Invariant
ProNet	Amino acid	All-Atom	$\ell = 0$	$SE(3)$ -Invariant
HoloProt	Amino acid + Surface	C_α + Surface	$\ell = 0$	$SE(3)$ -Invariant
MaSIF, dMaSIF	Surface	Surface	$\ell = 0$	$SE(3)$ -Invariant

acid level information and probably further the details at the atomic level to generate more accurate predictions. This requires a multi-level representation of proteins, which is challenging for the design of machine learning models.

Preserving Symmetries: Methods should follow the desired symmetry. Specifically, the output of the model should be $SE(3)$ -invariant, which means that the predicted results should not change with respect to the rotation and translation transformation of the protein structure. This is because the function and properties of a protein do not depend on its orientation or position in 3D space, but rather on its chemical composition and spatial arrangement of atoms.

Expressive Power: Accurately distinguishing different protein structures poses another significant challenge in protein representation learning. For protein structures that cannot be matched via $SE(3)$ transformation, effective methods should be able to distinguish them. This requires the methods to have great expressive power.

6.3.3 Existing Methods.

In Section 5.2, we discussed recent studies on representation learning for small molecules with 3D structures, where both invariant and equivariant methods were proposed to learn accurate representations. However, proteins are macromolecules with a large number of atoms and inherent multi-level structures, presenting significant challenges, as detailed in the previous section. Therefore, it is not practical to directly apply methods designed for small molecules to proteins. In this section, we summarize existing methods that are specifically designed to process protein structures, focusing on strategies for dealing with a large number of atoms and capturing inherent multi-level structures, as well as how to build more powerful and symmetry-aware representation learning methods, as summarized in Table 21, to tackle the above challenges.

Existing methods use different strategies to deal with the large number of atoms in proteins. For example, IEConv [Hermosilla et al. 2021] treats each atom as a node in a protein graph and employs several hierarchical pooling layers to reduce the number of nodes. In addition, the pooling operations enable multi-scale protein analysis and help the model learn different levels of protein representations. In contrast, methods including GearNet [Zhang et al. 2023d], ProNet [Wang et al.

Table 22. Some statistic information of CATH 4.2 curated by [Ingraham et al. \[2019\]](#), Fold dataset [[Hou et al. 2018; Hermosilla et al. 2021](#)], Enzyme Reaction dataset [[Hermosilla et al. 2021](#)], Enzyme Commission (EC) dataset [[Gligorijević et al. 2021](#)], and Gene Ontology dataset [[Gligorijević et al. 2021](#)]. We summarize the prediction tasks and the number of protein samples (# Samples), maximum number of amino acids in one protein (Maximum # amino acids), and average number of amino acids in one protein (Average # amino acids).

Datasets	Prediction Tasks	# Samples	Maximum # amino acids	Average # amino acids
CATH 4.2	Protein inverse folding, predict amino acid sequence	19,752	500	233
Fold	Protein fold classification	16,292	1419	168
Enzyme Reaction	Enzyme reaction classification	37,428	3,725	299
Gene Ontology	Gene Ontology (GO) term prediction	36,635	997	258
Enzyme Commission	Enzyme Commission (EC) number prediction	19,198	998	299

[2023b](#)], GVP-GNN [[Jing et al. 2021](#)], ProteinMPNN [[Dauparas et al. 2022](#)], PiFold [[Gao et al. 2023](#)], and CDCConv [[Fan et al. 2023](#)] treat each amino acid as a node in the graph and consider the information of atoms in each amino acid as special node and edge features. Since each amino acid contains many atoms, the graph size in these methods is significantly smaller than that of IEConv [[Hermosilla et al. 2021](#)], resulting in more efficient methods.

Furthermore, existing methods capture different levels of protein structure. For example, GearNet [[Zhang et al. 2023d](#)] considers only the C_α atom in each amino acid, and this structural encoder is trained by leveraging multiview contrastive learning and different self-prediction tasks. GVP-GNN [[Jing et al. 2021](#)] takes the unit vectors in the directions of $\mathbf{c}_i^N - \mathbf{c}_i^{C_\alpha}, \mathbf{c}_i^C - \mathbf{c}_i^{C_\alpha}, \mathbf{c}_j^{C_\alpha} - \mathbf{c}_i^{C_\alpha}$ as inputs, leading to a complete description of the protein backbone structure. IEConv [[Hermosilla et al. 2021](#)] treats each atom as a node and considers both the Euclidean distance between atoms and the shortest path with covalent or hydrogen bonds. Thus it can capture the full-atom structure of a protein. Similarly, ProNet [[Wang et al. 2023b](#)] can also capture the full-atom structure. The difference is that ProNet treats each amino acid, rather than each atom, as a node. It uses Euler angles between two backbone triangles as edge features and dihedral angles in the side chain as additional node features. This strategy effectively captures both the backbone and side-chain structures, leading to an expressive-powerful and efficient description of the protein all-atom structure. In addition to atomic coordinates, protein surfaces play a crucial role in understanding molecular interactions and protein functions. HoloProt [[Somnath et al. 2021](#)] goes beyond considering only C_α atoms and incorporates surface structures to capture coarser details of the protein. Similarly, MaSIF [[Gainza et al. 2020, 2023](#)] and dMaSIF [[Sverrisson et al. 2021](#)] specifically recognize the importance of protein surfaces in their respective approaches, highlighting their significant role in the analysis and understanding of protein interactions.

To build powerful and symmetry-aware representation learning methods, existing methods learn different order representations for each node. As shown in Table 21, most existing methods consider only scalar features, and the feature order is 0. For GVP-GNN [[Jing et al. 2021](#)] and GBPNet [[Aykent and Xia 2022](#)], the feature order is 1, as it considers directional vectors as node and edge features. The directional features are used to update the learned features for each node. However, currently, there are no higher-order methods designed for protein representation learning, mainly due to the large scale of protein structures. It would be interesting to explore the power of high-order (and many-body) methods discussed in Section 2 and Section 5.2 for protein representation learning.

6.3.4 Datasets and Benchmarks.

Representation learning methods for proteins with 3D structures are evaluated on various tasks, such as amino acid type prediction and protein function prediction. Table 22 summarizes commonly used datasets, including CATH 4.2 curated by [Ingraham et al. \[2019\]](#), Fold dataset [[Hou et al. 2018;](#)

Hermosilla et al. 2021], Enzyme Reaction dataset [Hermosilla et al. 2021], Enzyme Commission (EC) dataset [Gligorijević et al. 2021], and Gene Ontology dataset [Gligorijević et al. 2021].

CATH 4.2 dataset curated by Ingraham et al. [2019] is used for the task of inverse folding, also called computational protein design (CPD) or fixed backbone design, which aims to infer an amino acid sequence that can fold into a given structure. The dataset is collected based on the CATH hierarchical classification of protein structure [Orengo et al. 1997] and is split into 18,024 structures for training, 608 for validation, and 1,120 for testing. The evaluation metrics include perplexity and recovery. Perplexity measures the ability of the model to give a high likelihood to held-out sequences, and recovery evaluates predicted sequences versus the native sequences of templates. In addition to CATH 4.2, some other datasets are also used to test the performance of models on the inverse folding task, including CATH 4.3 [Hsu et al. 2022], TS 50 [Li et al. 2014; Jing et al. 2021], and TS 500 [Qi and Zhang 2020; Gao et al. 2023].

Fold dataset [Hou et al. 2018; Hermosilla et al. 2021] is a collection of 16,712 proteins with 3D structures curated from the SCOPe 1.75 database [Murzin et al. 1995], and each of the proteins is labeled with one of 1,195 fold classes. The fold classes indicate the secondary structure compositions, orientations, and connection orders of proteins. To evaluate the generalization ability of models, three test sets are used, namely Fold, Superfamily, and Family. Specifically, the Fold test set consists of proteins whose superfamily are unseen during training, the Superfamily test set consists of proteins whose family are unseen during training, and the Family test set consists of proteins whose family are present during training. Among these three test sets, Fold is the most challenging one as it differs most from the training data set. The dataset is divided into 12,312 proteins for training, 736 for validation, 718 for Fold, 1,254 for Superfamily, and 1,272 for Family. Accuracy is the evaluation metric for the fold classification task.

Enzyme Reaction dataset [Hermosilla et al. 2021] is a collection of enzymes, which are proteins that act as biological catalysts and can be classified with enzyme commission (EC) numbers [Webb et al. 1992] based on the reactions they catalyze. In total, this dataset contains 37,428 proteins with 3D structures from 384 classes, and the EC annotations are downloaded from the SIFTS database [Dana et al. 2019]. The dataset is divided into 29,215 proteins for training, 2,562 for validation, and 5,651 for testing, with each EC number represented in all three splits. Accuracy is used as the evaluation metric for this task.

Enzyme Commission (EC) dataset [Gligorijević et al. 2021] is also a collection of enzymes. However, unlike the enzyme reaction dataset that forms a protein-level classification task, this dataset forms 538 binary classification tasks based on the three-level and four-level 538 EC numbers [Webb et al. 1992]. Additionally, the enzymes collected in this dataset are different from those in the Enzyme Reaction dataset. In total, this dataset contains 19,198 proteins, with 15,550 for training, 1,729 for validation, and 1,919 for testing. This multi-label classification task is evaluated using two metrics, namely protein-centric maximum F-score (F_{\max}) and pair-centric area under precision-recall curve ($AUPR_{\text{pair}}$). For more detailed information on these metrics, please refer to relevant papers [Gligorijević et al. 2021; Wang et al. 2022a; Zhang et al. 2023d; Fan et al. 2023].

Gene Ontology dataset [Gligorijević et al. 2021] is used for the prediction of protein functions based on Gene Ontology (GO) terms [Ashburner et al. 2000] and forms multiple binary classification tasks. Specifically, GO classifies proteins into hierarchically related functional classes organized into three different ontologies, namely biological process (BP) with 1,943 classes, molecular function (MF) with 489 classes, and cellular component (CC) with 320 classes. The dataset is divided into 29,898 proteins for training, 3,322 for validation, and 3,415 for testing. The evaluation metrics are the same as those used for Enzyme Commission (EC) dataset [Gligorijević et al. 2021].

In addition to the datasets mentioned above, Atom3D [Townshend et al. 2020] is also commonly used to test the performance of protein representation learning methods. Specifically, Atom3D is

a unified collection of datasets concerning the 3D structures of biomolecules, including proteins, small molecules, and nucleic acids. It includes several datasets for protein-related tasks, such as Protein Interface Prediction (PIP), Residue Identity (RES), Mutation Stability Prediction (MSP), Ligand Binding Affinity (LBA), Ligand Efficacy Prediction (LEP), Protein Structure Ranking (PSR). The LBA task is described in detail in Section 8.

6.3.5 Open Research Directions.

Despite the recent advances in protein representation learning, several challenges remain unresolved, and certain directions remain underexplored. For example, while current methods can effectively capture the full-atom structure of proteins, it is still uncertain whether these methods can accurately capture the important local substructures, such as α -helix and β -sheet, in the secondary structure, and their spatial arrangement in the tertiary structure. Additionally, incorporating accessible surface area and protein domains into protein representation learning methods is important for a more comprehensive understanding of protein structure and function, potentially enhancing performance as well. For example, MaSIF [Gainza et al. 2020, 2023] focuses on solvent-excluded protein surfaces represented as meshes. On the other hand, dMaSIF [Sverrisson et al. 2021] employs oriented point clouds to model protein surfaces. Meanwhile, although both $\ell = 0$ and $\ell = 1$ equivariant methods have been proposed for protein representation learning, there is currently a lack of higher-order methods, which have been extensively investigated for small molecules.

In addition, protein dynamics is a significant and actively evolving field that focuses on studying the motions, conformational changes, and interactions of proteins over time, which are critical for understanding the function and behavior of proteins. One direction to explore protein dynamics is by leveraging temporal GNNs in conjunction with the protein representation learning methods we introduced. By incorporating temporal information and accounting for the dynamic nature of protein structures and interactions, temporal GNNs offer a promising avenue for unraveling the intricacies of protein dynamics. Another important aspect is protein circuit design, which holds great potential for advancing our understanding of protein function and behaviors.

6.4 Protein Backbone Structure Generation

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Recommended Prerequisites: Sections 5.3, 5.4

As mentioned above, we focus on diffusion models for protein backbone structure generation in this survey. Here, we first describe the aforementioned challenges for protein backbone structure generation in detail, then discuss how previous methods address these challenges from two perspectives, including protein structure representations and diffusion processes. The pipeline of protein generation with diffusion model is illustrated in Figure 27.

6.4.1 Problem Setup.

The protein backbone generation task is to learn a generative model p_θ that can model the density distribution of real protein backbones $p_{\mathcal{P}_{bb}}$ and then we can sample a novel protein backbone $\hat{\mathcal{P}}_{bb}$ that satisfies $p_\theta(\hat{\mathcal{P}}_{bb}) \approx p_{\mathcal{P}_{bb}}(\hat{\mathcal{P}}_{bb})$.

6.4.2 Technical Challenges.

Sophisticated Data Distribution: Generation tasks need the sampling of new data points from the data distribution. However, the distribution of real protein structures is unknown, sparse, and

intractable to sample from. Thus, establishing a bijective mapping between the data distribution and prior distributions, such as Gaussians, is crucial.

Distribution $E(3)/SE(3)$ -Invariance: Distribution $E(3)/SE(3)$ -invariance, which arises from the nature of real protein structure distribution, needs to be satisfied. Specifically, for a protein backbone structure \mathcal{P}_{bb} sampled from real data distribution $p_{\mathcal{P}_{bb}}$, if we apply 3D rotation transformations $R \in \mathbb{R}^{3 \times 3}$, $|R| = \pm 1$ for $E(3)$ and $R \in \mathbb{R}^{3 \times 3}$, $|R| = 1$ for $SE(3)$ and translation transformations $b \in \mathbb{R}^3$, the geometric structure of the given protein remains unchanged. Hence, we have $p_{\mathcal{P}_{bb}}(\mathcal{P}_{bb}) = p_{\mathcal{P}_{bb}}(R\mathcal{P}_{bb} + b)$ for real data distribution $p_{\mathcal{P}_{bb}}$. To better mimic the real data distribution and minimize bias in machine learning methods, distribution $E(3)/SE(3)$ -invariance needs to be encoded into the generation pipeline.

$E(3)/SE(3)$ -Equivariant Networks: In order to achieve distribution $E(3)/SE(3)$ -invariance, the neural networks should satisfy the equivariant property. As discussed above, two identical protein structures up to a group transformation should satisfy $p_{\mathcal{P}_{bb}}(\mathcal{P}_{bb}) = p_{\mathcal{P}_{bb}}(R\mathcal{P}_{bb} + b)$. For a protein in 3D space, first, we can let the coordinates have zero centroids by subtracting the mean values of the coordinates. In this way, the translational invariance for the distribution is naturally satisfied. Therefore, we only need to consider rotational invariance, that is, $p_{\mathcal{P}_{bb}}(\mathcal{P}_{bb}) = p_{\mathcal{P}_{bb}}(R\mathcal{P}_{bb})$ should be satisfied. Through the total probability rule, we have $p(\mathcal{P}_{bb}) = \int p(\mathcal{P}_{bb}|Z)p(Z)dZ$, where Z denotes the latent variables. Similarly, we also have $p(R\mathcal{P}_{bb}) = \int p(R\mathcal{P}_{bb}|RZ)p(RZ)dRZ$.

If we sample latent variables Z from zero-centered Gaussian distribution, then $p(RZ) = p(Z)$ can be easily satisfied. And in order to achieve $p(R\mathcal{P}_{bb}|RZ) = p(\mathcal{P}_{bb}|Z)$, we should make the networks mapping from Z to \mathcal{P}_{bb} to be equivariant to R . Thus, $p(R\mathcal{P}_{bb}) = p(\mathcal{P}_{bb})$ holds and distribution $E(3)/SE(3)$ -invariance is satisfied.

Computational Efficiency: Efficient modeling of protein structures is also crucial. In addition to the aforementioned challenges, it is worth noting that the exploration space of protein structures is incredibly vast, while known protein structures are limited. Therefore, elegant protein representations must be established not only to ease the modeling difficulty but also to minimize geometric bias in machine learning models.

6.4.3 Existing Methods.

Protein Structure Representations: To address the efficiency issue of protein structure modeling, recent approaches either generate protein backbone structures alone [Wu et al. 2022d; Trippe et al. 2022; Fu et al. 2023b; Lin and AlQuraishi 2023; Yim et al. 2023], or first generate protein backbone structures and then predict the full atom-level protein structures [Ingraham et al. 2022; Watson et al. 2022] due to high complexity and vast exploration space of complete protein structures. When modeling protein backbone structures, different 3D representations are used, resulting in different modeling costs and diffusion processes. Specifically, ProtDiff [Trippe et al. 2022], Chroma [Ingraham et al. 2022], LatentDiff [Fu et al. 2023b], and Genie [Lin and AlQuraishi 2023] represent protein backbone structures by alpha carbon positions \mathcal{P}_{bb} , without considering positions of N, C, O atoms. LatentDiff further encodes the protein backbone structure into the compact latent space to reduce modeling complexity. Both FoldingDiff [Wu et al. 2022d] and RFdiffusion [Watson et al. 2022] represent protein backbone structures based on positions of C_α, N, C atoms. However, FoldingDiff [Wu et al. 2022d] uses relative bond and torsion angles along amino acid chains, and RFdiffusion [Watson et al. 2022] uses the 3D positions of C_α and 3×3 rotation matrices representing the rigid-body orientation of each residue in a global reference frame. Recently, FrameDiff [Yim et al. 2023] further considers positions of O atoms and represents protein backbone structures using 3D positions of C_α , 3×3 rotation matrices, and rotation torsion angles of O . The

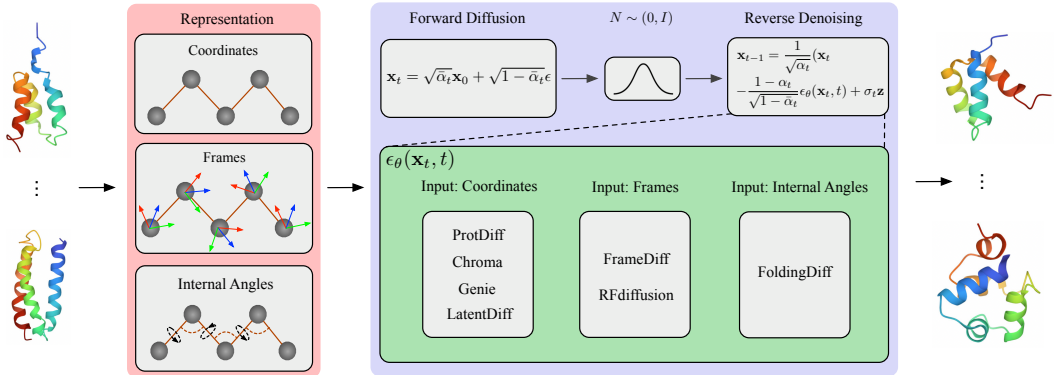


Fig. 27. Pipeline of protein generation with diffusion models. The process begins by diffusing protein backbone structures to Gaussian noise, which is then denoised using a learned denoising network to generate novel protein structures. Protein backbones can be represented in various ways, such as coordinates, frames, or internal angles. Different methods have been proposed for each representation. For coordinates representation, methods include ProtDiff [Trippe et al. 2022], Chroma [Ingraham et al. 2022], LatentDiff [Fu et al. 2023b] (coordinates are in the latent space), and Genie [Lin and AlQuraishi 2023]. For frames representation, methods include FrameDiff [Yim et al. 2023] and RFdiffusion [Watson et al. 2022]. For internal angles representation, the method is FoldingDiff [Wu et al. 2022d].

Table 23. Summary of protein 3D representations used by previous works as well as the corresponding levels of protein structural granularity and modeling space. N denotes the number of amino acids. f denotes the downsampling factor in LatentDiff. Among them, ProtDiff [Wu et al. 2022d], Chroma [Ingraham et al. 2022], and Genie [Lin and AlQuraishi 2023] only consider positions of alpha carbons, LatentDiff [Fu et al. 2023b] consider the node positions in the latent space, while FoldingDiff [Wu et al. 2022d] and RFdiffusion [Watson et al. 2022] further consider positions of carbon and nitrogen atoms in protein backbone structures. FrameDiff [Yim et al. 2023] considers all atoms in protein backbone structures.

Methods	Protein 3D Representations	Structural Granularity	Modeling Space
ProtDiff	Coordinates	C_α	$\mathbb{R}^{N \times 3}$
FoldingDiff	Bond and torsion angles	C_α, C, N	$[0, 2\pi]^{6N}$
Chroma	Coordinates	C_α	$\mathbb{R}^{N \times 3}$
RFdiffusion	Coordinates + Frame rotation angles	C_α, C, N	$\mathbb{R}^{N \times 3} SO(3)^N$
LatentDiff	Coordinates (latent space)	C_α	$\mathbb{R}^{\frac{N}{f} \times 3}$
Genie	Coordinates	C_α	$\mathbb{R}^{N \times 3}$
FrameDiff	Coordinates + Frame rotation angles	C_α, C, N, O	$\mathbb{R}^{N \times 3} SO(3)^N [0, 2\pi]^N$

representations, corresponding levels of structural granularity, and modeling spaces of previous works are summarized in Table 23.

Diffusion Models: Given different protein structure representations, corresponding diffusion processes need to be established to address challenges in protein backbone generation, including establishing the bijective mapping between data distribution and prior distribution to enable sampling, ensuring distribution $E(3)/SE(3)$ -invariance, and $E(3)/SE(3)$ -equivariant property of neural networks.

3D Euclidean coordinates of alpha carbons are used in ProtDiff [Trippe et al. 2022], LatentDiff [Fu et al. 2023b], and Genie [Lin and AlQuraishi 2023], with relatively simple diffusion process.

Specifically, ProtDiff, LatentDiff, and Genie use the zero-mean distribution to get rid of influences of translations in 3D space and achieve distribution translation invariance. Moreover, LatentDiff encodes protein structures into the latent space to reduce the modeling complexity. Beyond this, to address influences of rotation transformations, ProtDiff and LatentDiff use the $E(3)$ -equivariant network EGNN and achieve distribution $O(3)$ -invariance, while Genie achieves distribution $SO(3)$ -invariance which is sensitive to reflections by using $SE(3)$ -equivariant IPA layers. Corresponding forward and reverse diffusion processes for 3D coordinates similarly used in the image domain are established to enable the sampling of protein structures. One limitation of this structure representation is that only the positions of alpha carbons in the backbone structure are considered, and an additional generation step is needed to generate the positions of N , C , and O atoms.

As mentioned above, diffusion models need to transform original data into Gaussian noise and learn a denoising network to mimic and generate realistic data from Gaussian noise. Most methods transform Euclidean coordinates by adding isotropic Gaussian noise, which appears as an uncorrelated diffusion process and could break some common structure constraints of proteins. As a result, the models need to have extra designs to learn these correlations from data. To avoid this, Chroma [Ingraham et al. 2022] introduces a correlated diffusion process that transforms proteins into random collapsed polymers and uses designed covariance models to encode the chain and radius of gyration constraints. Additionally, Chroma designs a random graph neural network that can capture long-range information with sub-quadratic scaling, and the network predicts pairwise inter-residue geometries and then optimize 3D protein structures in a $SE(3)$ -equivariant manner.

Bond and torsion angles along the backbone structure are used by FoldingDiff [Wu et al. 2022d], with the assumption that the lengths of chemical bonds along the backbone chain follow practical constraints. Due to the $E(3)$ -invariant nature of bond and torsion angles, a simple sequence model is used to achieve distribution $E(3)$ -invariant. To enable the sampling process from Gaussian noise, FoldingDiff applies the forward and reverse diffusion processes similar to coordinates to bond angles and torsion angles, regardless of the fact that bond angles and torsion angles belong to compact Riemannian Manifolds instead of Euclidean space.

Frame representations consisting of 3D positions of alpha carbons and relative rotation angles of amino acid planes are used by RFdiffusion [Watson et al. 2022] and FrameDiff [Yim et al. 2023]. Specifically, to achieve distribution $SE(3)$ -invariance, FrameDiff uses the zero-mean distribution to get rid of influences of translations in 3D space and achieve distribution translation invariance for 3D positions of alpha carbons. Beyond this, to address influences of rotation transformations, FrameDiff achieves distribution $SO(3)$ -invariance which is sensitive to reflections by using $SE(3)$ -equivariant IPA layers. And RFdiffusion modifies RoseTTAFold, a powerful protein structure prediction method, as the denoising network and achieves $SE(3)$ -equivariance property. To enable the sampling process from Gaussian noise, different from previous works [Wu et al. 2022d; Lin and AlQuraishi 2023; Wu et al. 2022d] using diffusion processes established in Euclidean space, FrameDiff proposes solid $SO(3)$ diffusion forward and reverse processes for the rotation matrices of amino acid planes which belong to compact Riemannian Manifolds. Also, for the RFdiffusion, noise should be added to the rotation matrix, so Brownian motion on the manifold of $SO(3)$ is adopted. And the diffusion processes of alpha carbon positions in FrameDiff and RFdiffusion are similar to ProtDiff and Chroma. Moreover, RFdiffusion uses a self-conditioning mechanism that uses the denoising network output as the template input to the subsequent denoising step, which is similar to the recycling in AlphaFold2 [Jumper et al. 2021]. The diffusion space and achieved distribution symmetries of previous works are summarized in Table 24.

Table 24. Demonstration of diffusion processes for different protein representations and achieved distribution symmetries of previous works. Among them, ProtDiff [Wu et al. 2022d], FoldingDiff [Wu et al. 2022d], and LatentDiff [Fu et al. 2023b] achieve distribution $E(3)$ -invariance and treat chiral protein structures as the same, while Chroma [Ingraham et al. 2022], RFdiffusion [Watson et al. 2022], Genie [Lin and AlQuraishi 2023], and FrameDiff [Yim et al. 2023] achieve distribution $SE(3)$ -invariance and have better generation performances.

Methods	Network	Diffusion Space	Distribution Symmetry
ProtDiff	$E(3)$ -Equivariant	Euclidean	$E(3)$ -Invariance
FoldingDiff	$E(3)$ -Invariant	Angle	$E(3)$ -Invariance
Chroma	$E(3)$ -Equivariant	Euclidean	$SE(3)$ -Invariance
RFdiffusion	$SE(3)$ -Equivariant	Euclidean + $SO(3)$	$E(3)$ -Invariance
LatentDiff	$E(3)$ -Equivariant	Euclidean	$E(3)$ -Invariance
Genie	$SE(3)$ -Equivariant	Euclidean	$SE(3)$ -Invariance
FrameDiff	$SE(3)$ -Equivariant	Euclidean + $SO(3)$ + $SO(2)$	$SE(3)$ -Invariance

6.4.4 Datasets and Benchmarks.

For now, there are no standard benchmark datasets for the task of protein backbone structure generation. Early protein backbone generation methods are usually evaluated on selected protein structures from PDB [Berman et al. 2000] or other protein structure libraries. Specifically, ProtDiff uses 4269 single-chain protein structures with the number of amino acids in the range [40, 128], while FoldingDiff uses the CATH [Ingraham et al. 2019] dataset with 24316 structures for training, 3039 structures for validation, and 3040 structures for testing. Chroma [Ingraham et al. 2022] queries non-membrane X-ray protein structures with a resolution of 2.6 of Å or better, and an additional set of 1725 non-redundant antibody structures was added, resulting in 28819 structures in total. In RFdiffusion [Watson et al. 2022], RosettaFold (RF) is pre-trained on a mixture of several data sources, including monomer/homo-oligomer and hetero-oligomer structures in the PDB, AlphaFold2 data having pLDDT > 0.758, and negative protein-protein interaction examples generated by random pairing. Then, RFdiffusion is trained on monomer structures in the PDB used for RF training. LatentDiff curates about 100k training data from Protein Data Bank (PDB) [Berman et al. 2000] and Swiss-Prot data in AlphaFold Protein Structure Database (AlphaFold DB) [Jumper et al. 2021; Varadi et al. 2022]. Genie uses 8766 protein domains, with 3,942 domains having at most 128 residues from the Structural Classification of Proteins-extended (SCOPe) dataset, and FrameDiff uses 20312 protein backbones from PDB [Berman et al. 2000] for training. When evaluating model performance, the widely-used metrics include the scTM score (higher is better) and novelty of generated protein backbone structures compared with the training structures.

6.4.5 Open Research Directions.

Recent protein generative models mainly focus on the backbone level of protein structures but can not generate full atom-level protein structures in a one-step manner. Additionally, recent methods are mainly designed for random protein structure generation or conditional generation given protein substructures. Another potential direction beyond this will be generating protein structures satisfying desirable properties.

7 AI FOR MATERIALS SCIENCE

In this section, we discuss the applications of AI techniques in materials science. We first give an overview introduction about crystalline materials and elaborate formal definitions of physical symmetries of crystalline materials in Section 7.1. Next, we discuss two common and fundamental tasks, material representation learning problem and the material generation problem in Section 7.2 and 7.3, respectively. Furthermore, we include three advanced topics, including ordered crystalline materials characterization in Section 7.4, disordered crystalline materials characterization in Section 7.5, and phonon calculations in Section 7.6.

7.1 Overview

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In addition to small molecules and proteins, AI methods have been used for modeling crystalline materials, which are another family of large chemical compounds formed by periodic repetitions of atoms in 3D space. Crystalline materials are the foundation of many real-world industrial applications, such as semiconductor electronics, solar cells, and batteries [Butler et al. 2018]. Due to the dramatic demand of the industry, materials science has emerged to study a variety of fundamental research, such as predicting material properties (*e.g.*, formation energy) and designing novel materials with target properties. For a long time, the research progress in these problems was relatively slow due to heavy reliance on either expensive lab experiments or time-consuming materials simulations. Recently, inspired by the success of AI methods, especially machine learning models on molecules, many studies have tried to apply these models to crystalline materials related problems [Choudhary et al. 2023; Du et al. 2023b]. Nonetheless, different from molecules, the arrangement of atoms in crystalline materials has a complicated periodic arrangement of repeating unit cells and atoms. Hence, crystalline materials have very different physical symmetries from molecules, and developing effective AI models for them requires explicitly capturing these symmetries in models.

In this and the following sections, we describe the structure of a crystalline material by lattice vectors and one of its unit cells, *i.e.*, the smallest repeatable structures. Specifically, let the number of atoms in any unit cell be n , then a crystalline material \mathcal{M} is represented as $\mathcal{M} = (\mathbf{z}, \mathbf{C}, \mathbf{L})$. Here, $\mathbf{z} \in \mathbb{Z}^n$ is the atom type vector where the i -th element z_i of \mathbf{z} denotes the atom type (atomic number) of the i -th atom in the unit cell. $\mathbf{C} = [\mathbf{c}_1, \dots, \mathbf{c}_n] \in \mathbb{R}^{3 \times n}$ is the coordinate matrix where \mathbf{c}_i denotes the 3D coordinate of the i -th atom in the unit cell. $\mathbf{L} = [\boldsymbol{\ell}_1, \boldsymbol{\ell}_2, \boldsymbol{\ell}_3] \in \mathbb{R}^{3 \times 3}$ is the lattice matrix, and the three lattice vectors $\boldsymbol{\ell}_1, \boldsymbol{\ell}_2, \boldsymbol{\ell}_3$ describe the three periodicity vectors along which the atoms periodically repeat themselves. In physics, there exist several well-defined symmetry transformations for crystalline materials, including the following permutation, $E(3)$, and periodic transformations.

- **Permutation transformations** produce a new material by permuting the atom orders in $\mathcal{M} = (\mathbf{z}, \mathbf{C}, \mathbf{L})$, *i.e.*, exchanging the elements in \mathbf{z} and column vectors in \mathbf{C} with the same order.
- **$E(3)$ transformations**, or rigid transformations, change the 3D coordinates of $\mathcal{M} = (\mathbf{z}, \mathbf{C}, \mathbf{L})$ by translation, rotation, or reflection in 3D space. Specifically, \mathbf{C} and \mathbf{L} are replaced by $R\mathbf{C} + \mathbf{t}\mathbf{1}^T$ and $R\mathbf{L}$, respectively, where $\mathbf{t} \in \mathbb{R}^3$ is an arbitrary translation vector, $R \in \mathbb{R}^{3 \times 3}$ is an orthogonal matrix satisfying $R^T R = I$, and $\mathbf{1}$ is an n -dimensional vector whose elements are all 1s.
- **Periodic transformations** map the 3D coordinates of $\mathcal{M} = (\mathbf{z}, \mathbf{C}, \mathbf{L})$ to periodically equivalent coordinates, which can be formally described as replacing \mathbf{C} by a new coordinate matrix $\mathbf{C}' = \mathbf{C} + L\mathbf{K}$, where $\mathbf{K} \in \mathbb{Z}^{3 \times n}$ is an arbitrary integer matrix.

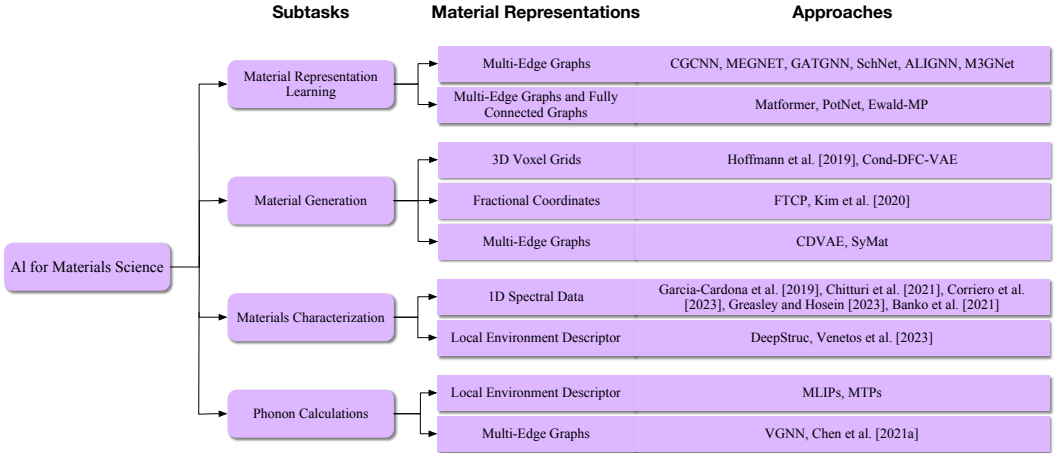


Fig. 28. An overview of the tasks and methods in AI for materials science. In this section, we consider four tasks, including material representation learning, material generation, materials characterization, and phonon calculations. In material representation learning, we discuss two categories of methods using different graph representations of materials. The first category of methods, including CGCNN [Xie and Grossman 2018], MEGNET [Chen et al. 2019b], GATGNN [Louis et al. 2020], SchNet [Schütt et al. 2018], ALIGNN [Choudhary and DeCost 2021], and M3GNet [Chen and Ong 2022], only use multi-edge graphs. The second category of methods, including Matformer [Yan et al. 2022], PotNet [Lin et al. 2023d], and Ewald-MP [Kosmala et al. 2023], use multi-edge graphs and fully connected graphs to capture periodic information. In material generation, we discuss three categories of methods using three different representations of 3D material structures as generation targets. The first category of methods, including Hoffmann and Noé [2019] and Cond-DFC-VAE [Court et al. 2020], generate 3D voxel grids. The second category of methods, including FTCP [Ren et al. 2022] and Kim et al. [2020], generate fractional coordinates. The third category of methods, including CDVAE [Xie et al. 2022a] and SyMat [Luo et al. 2023c], generate materials in the form of multi-edge graphs. In materials characterization, we discuss two categories of methods using different representations of materials. The first category of methods, including Garcia-Cardona et al. [2019], Chitturi et al. [2021], Corriero et al. [2023], Greasley and Hosein [2023] and Banko et al. [2021], uses 1D spectral data to represent materials and then to predict material structures. The second category of methods, including DeepStruc [Kjær et al. 2023] and Venetos et al. [2023], takes advantage of local environment descriptors of materials to model disordered materials. Note that both Section 7.4 and Section 7.5 describe materials characterization while Section 7.5 specifically addresses scenarios for disordered materials. For the sake of clarity, these two sections have been integrated into a single task, materials characterization, in this figure. In phonon calculations, we discuss two categories of methods using different representations of materials. The first category of methods, including MLIPs [Mortazavi et al. 2020] and MTPs [Zuo et al. 2020], applies local environment descriptors of materials to represent material structure. The second category of methods, including VGNN [Okabe et al. 2023] and Chen et al. [2021a], uses multi-edge graphs to capture periodic information.

In other words, for an arbitrary material $\mathcal{M} = (z, C, L)$, if \mathcal{M}' is obtained by applying one of the above transformations on \mathcal{M} , we should consider \mathcal{M} and \mathcal{M}' as different representations of the same material. Ideally, the learned property prediction function f and material distribution p should be symmetry-aware, i.e., satisfying $f(\mathcal{M}) = f(\mathcal{M}')$ and $p(\mathcal{M}) = p(\mathcal{M}')$. In the following subsections, we review and discuss existing AI methods for material representation learning and material generation, as well as emerging topics including ordered/disordered materials characterization and phonon calculation, and compare them mainly from the perspective of capturing symmetries. See an overview of our covered methods in Figure 28.

7.2 Material Representation Learning

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Recommended Prerequisites: Section 7.1

We first discuss material representation learning for property prediction in this section. The major challenge of developing material representation learning models lies in capturing symmetries in crystalline materials, particularly the invariance to periodic transformations. To overcome this challenge, existing studies have proposed numerous crystal graph representation construction methods and crystal graph neural network models. We elaborate on them in the next subsections.

7.2.1 Problem Setup.

Material representation learning requires learning a function f to predict the property y of any given material M , and y can be a real number (regression problem) or categorical number (classification problem).

7.2.2 Technical Challenges.

Crystalline material representation learning aims to predict physical and chemical properties of crystalline materials based on their lattice structures. As already illustrated in the above section, different from small molecules or proteins, crystalline materials consist of a smallest unit cell structure and corresponding periodic repeating patterns in 3D space. Thus, unique geometric symmetries and model designs need to be established for crystalline materials. Specifically, when rotation transformations are applied to C and L together, or when translation transformations are applied to C alone, the crystal structure remains unchanged, which is described as Unit Cell $E(3)$ invariant property by [Yan et al. \[2022\]](#). Beyond this, when periodic transformations are applied, the crystal structure remains the same and the corresponding graph representation should be the same, which is periodic invariant described in Matformer [\[Yan et al. 2022\]](#). Additionally, periodic patterns that indicate the orientations that a unit cell repeats in 3D space are also crucial for crystal structural modeling. For the crystal neural network design, due to the fact that crystal structures can have more than two hundred atoms in a unit cell, it remains challenging to consider higher body order interactions between atoms. However, higher-order interactions are arguably indispensable to achieve complete geometric representations for crystal structures. Additionally, powerful and efficient networks need to be designed for crystal structures as there could be more than two hundred atoms in the unit cell.

7.2.3 Existing Methods.

As shown in Figure 29, a typical procedure of material representation learning contains constructing crystal graph representations and employing crystal graph neural networks to predict properties. We discuss several representative crystal graph representation methods and crystal graph neural network models below.

Crystal Graph Representations: The periodic nature of crystals poses unique challenges for crystal representation learning as mentioned above. Specifically, to tackle challenges (1) and (2), CGCNN [\[Xie and Grossman 2018\]](#) proposes to represent the infinite crystal structure by multi-edge crystal graph built upon the unit cell structure, which has been verified to be periodic invariant by [Yan et al. \[2022\]](#). Concretely, multi-edge crystal graph maps a given atom and all its duplicates in 3D space as a single node, and models the interactions between node pairs by recording pairwise Euclidean distances as multiple edges. Due to the effectiveness and simplicity of the multi-edge crystal graph, it has been widely used by follow-up works, including MEGNET [\[Chen et al. 2019b\]](#),

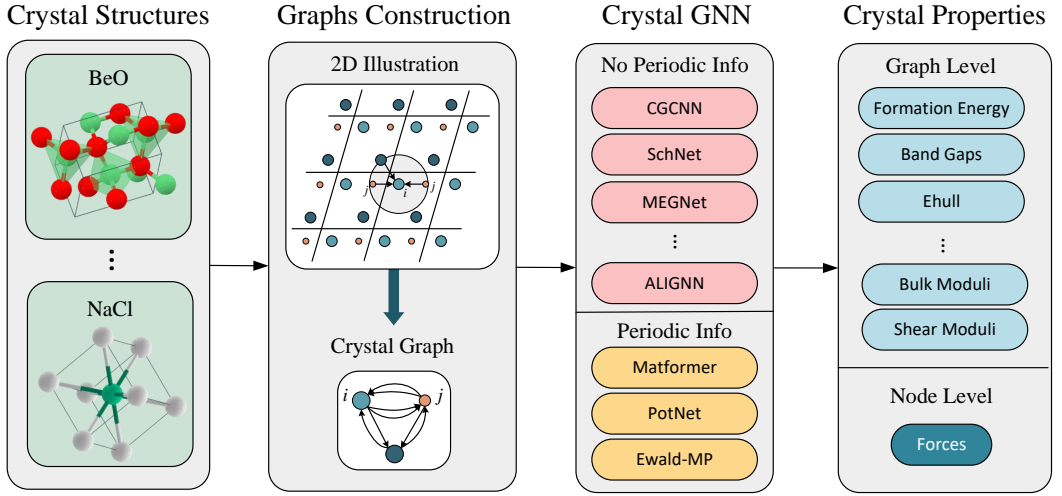


Fig. 29. Pipeline of material representation learning. A crystalline material is transformed into a crystal graph representation, subsequently serving as the input for a crystal graph message passing neural network. Then the models are trained to accurately predict the desired properties of the crystal. Notably, a message passing process can be distinguished based on its incorporation of periodic information, such as lattice lengths or infinite potential summations. Networks without periodic information include models such as CGCNN [Xie and Grossman 2018], SchNet [Schütt et al. 2018], MEGNet [Chen et al. 2019b], and ALIGNN [Choudhary and DeCost 2021], among others. Conversely, networks that incorporate periodic information include Matformer [Yan et al. 2022], PotNet [Lin et al. 2023d], and Ewald-MP [Kosmala et al. 2023]. Additionally, the predicted output can be classified into graph-level properties, including formation energies and band gaps, or node-level properties, which include forces.

Table 25. Summary of crystal graph representations used by previous works. It can be seen that all previous works satisfy periodic invariance and achieve $E(3)$ invariance. Among these methods, Matformer [Yan et al. 2022], PotNet [Lin et al. 2023d], and Ewald-MP [Kosmala et al. 2023] encode periodic patterns explicitly. CGCNN [Xie and Grossman 2018], MEGNET [Chen et al. 2019b], GATGNN [Louis et al. 2020], SchNet [Schütt et al. 2018], Matformer [Yan et al. 2022], and PotNet [Lin et al. 2023d] only use two body bond information, while ALIGNN [Choudhary and DeCost 2021] and M3GNet [Chen and Ong 2022] further use three body angle information.

Methods	Periodic Invariant	Symmetry	Periodic Pattern	Body Order	Complete
CGCNN	✓	$E(3)$ invariant	✗	2	✗
MEGNET	✓	$E(3)$ invariant	✗	2	✗
GATGNN	✓	$E(3)$ invariant	✗	2	✗
SchNet	✓	$E(3)$ invariant	✗	2	✗
Matformer	✓	$E(3)$ invariant	✓	2	✗
PotNet	✓	$E(3)$ invariant	✓	2	✗
Ewald-MP	✓	$E(3)$ invariant	✓	2	✗
ALIGNN	✓	$E(3)$ invariant	✗	3	✗
M3GNet	✓	$E(3)$ invariant	✗	3	✗

GATGNN [Louis et al. 2020], and ALIGNN [Choudhary and DeCost 2021]. Additionally, the use of

pairwise Euclidean distances enables multi-edge crystal graph invariant to $E(3)$ transformations including rotations, translations, and reflections in 3D space. Thus, multi-edge crystal graphs are periodic invariant and $E(3)$ invariant, only considering pairwise Euclidean distances with a body order of 2. Beyond this, to enable higher body order interactions, ALIGNN proposes to further include angles between bonds without breaking periodic invariance, and M3GNet includes three-body interactions in a similar way. However, these methods only consider local interactions of given atoms and cannot capture periodic patterns, which are the key differences between crystal structures and molecule structures. To explicitly capture the periodic patterns, Matformer [Yan et al. 2022] proposes to encode three periodic vectors by using six self-connecting edges, whose combination can fully capture the lengths of periodic vectors and angles between them. Additionally, recent two works take advantage of Ewald summations to capture periodic information. PotNet [Lin et al. 2023d] proposes to consider the infinite interactions between any two nodes in a crystal structure by using infinite summations of multiple types of potentials, and thus periodic patterns are captured by those summations. In a similar vein, Ewald-MP [Kosmala et al. 2023] tackles the problem of long-range interactions in periodic structures by applying the decomposition from Ewald summation. By decomposing the aggregation of message passing into a short-range signal, modeled in real space, and a long-range signal, modeled in Fourier space, periodic structure information is captured explicitly. A summary of crystal graph representations of previous works is shown in Table 25.

Crystal Graph Neural Networks: The effectiveness of modern graph neural networks in predicting material properties relies on the symmetry and high-order information they incorporate as well as the used message passing fashions. Existing networks such as CGCNN [Xie and Grossman 2018], MEGNET [Chen et al. 2019b], GATGNN [Louis et al. 2020], and SchNet [Schütt et al. 2018] employ radius crystal graphs and consider only two-body distances as edge features during message passing, similar to the methods used in molecular representation learning. Concretely, SchNet and MEGNET use common graph convolution networks, whereas CGCNN employs the sigmoid gate operation to the concatenation of node and edge features, and GATGNN employs the attention mechanism to weight node features during message passing. Moreover, as mentioned above, Matformer [Yan et al. 2022], PotNet [Lin et al. 2023d], and Ewald-MP [Kosmala et al. 2023] incorporate additional features during message passing to address the limitation of lacking periodic information of the previous methods. Specifically, Matformer explicitly encodes lattice structure information into self-connecting edges. In addition, PotNet considers infinite interatomic potential summations by summing up long-range and short-range interactions and encoding them into the edge features of fully-connected graphs. In contrast, Ewald-MP decouples the long-range interactions from the short-range message passing and combines the short and long-range node embeddings after each layer. To reduce complexity compared to fully-connected graphs, both message passings are built upon cutoff graphs, a distance cutoff in real space and a frequency cutoff in Fourier space. While most of the above methods are based on interatomic (two-body) information, incorporating three-body information can also enhance material representations, as demonstrated by ALIGNN [Choudhary and DeCost 2021] and M3GNet [Chen and Ong 2022]. These methods convert the crystal graph into a line graph, where the original edges become node features, and the angles between edges become edge features. They subsequently apply graph neural networks, similar to CGCNN, to learn material representations.

7.2.4 Datasets and Benchmarks.

There are three widely-used crystal property prediction benchmarks as shown in Table 26, including the Materials Project-2018.6.1 [Chen et al. 2019b], JARVIS [Choudhary and DeCost 2021], and MatBench [Dunn et al. 2020]. The Materials Project-2018.6.1 has four widely-used regression tasks

Table 26. Dataset statistics of the Materials Project-2018.6.1(MP) [Chen et al. 2019b], JARVIS [Choudhary and DeCost 2021], and MatBench [Dunn et al. 2020]. The number of crystals in the largest scale tasks of corresponding datasets, number of regression tasks, and number of classification tasks are summarized.

Datasets	Largest scale task	# regression tasks	# classification tasks
MP	69,239	4	2
JARVIS	55,722	29	10
MatBench	132,752	10	3

for the properties of formation energy, band gap, bulk moduli, and shear moduli. JARVIS includes 29 crystal property regression tasks and 10 crystal property classification tasks. The widely used regression tasks in JARVIS are formation energy, bandgap (OPT), bandgap (MBJ), Ehull, and total energy. MatBench consists of 10 regression tasks and 3 classification tasks. Most of the crystal properties are calculated by using Density Function Theory (DFT) based methods.

7.2.5 Open Research Directions.

First, current deep learning based crystal property prediction methods are mainly designed for regression and classification tasks. A possible future direction would be exploring the higher rotation order crystal properties including atomic forces, dielectric tensors, *etc.* Second, current works for crystal property prediction tasks are not geometrically complete, and geometric completeness for infinite crystal structures is a challenging yet important topic.

7.3 Material Generation

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Recommended Prerequisites: Section 7.1

In this section, we focus on the problem of generating crystalline materials. In this problem, a key challenge is achieving invariance to all symmetry transformations in probabilistic modeling frameworks of generative models. We elaborate the details of several existing crystalline material generation methods and their captured symmetries in the following subsections.

7.3.1 Problem Setup.

Material generation learns a probabilistic distribution p over the material space so that novel materials can be generated by sampling from p .

7.3.2 Technical Challenges.

For the crystalline material generation problem, we aim to learn a probability distribution p over the material space with generative models, and sample novel crystalline materials from p . The key challenge of this problem is incorporating all symmetries of crystalline materials into generative models. In other words, if two crystalline materials M and M' can be mutually transferred to each other by symmetry transformations, they should be assigned to the same probability by generative models. Particularly, the symmetry transformations of 3D material structures include not only $E(3)$ transformations that commonly exist in other chemical compounds, but also periodic transformations that are unique to crystalline materials. Hence, we cannot simply apply the existing 3D molecule generation methods (Section 5.4) to the crystalline material generation problem because they do not consider invariance to periodic transformations. It poses significant

challenges to incorporate the invariance to periodic transformations into existing $E(3)$ -invariant generative models in 3D molecule generation. Also, periodic transformations do not preserve the distances between every pair of two atoms so they are not Euclidean transformations. Hence, we cannot generate 3D material structures using 3D features like distances, angles, or torsion angles as generation targets.

7.3.3 Existing Methods.

Generally, two strategies can be used to ensure invariance to periodic transformations. First, we can implicitly determine the atom positions in materials by generating 3D features or representations (e.g., 3D voxel grids) that are internally invariant to periodic transformations. Two representative methods using this strategy are Cond-DFC-VAE [Court et al. 2020] and the method proposed in Hoffmann et al. [2019]. They both convert crystalline materials to 3D voxel grids, smooth 3D voxel grids to 3D density maps, and use 3D density maps as the generation targets. Specifically, a 3D voxel grid is obtained from the 3D crystalline material by extracting the information of all atoms in a 3D cube. For any grid point in the 3D voxel grid, its voxel value is non-zero if there is an atom in its corresponding position, and non-zero voxel values contain the information of atom types. Because 3D voxel grids are usually very sparse and not suitable to serve as the direct generation targets, they are smoothed to 3D density maps where most values are non-zero. Crystalline materials are generated by first generating 3D density maps with a VAE model [Kingma and Welling 2014], then segmented to 3D voxel grids by a U-Net model [Ronneberger et al. 2015]. 3D voxel grids or 3D density maps are invariant to periodic transformations, but not invariant to $E(3)$ transformations, so Cond-DFC-VAE and the method in Hoffmann et al. [2019] both fail to capture $E(3)$ symmetries.

In addition, the other strategy is to directly generate the lattice matrix L and coordinate matrix C , but tailored probabilistic modeling is needed for generative models so that for any integer matrix $K \in \mathbb{Z}^{3 \times n}$, $p(C) = p(C + LK)$ always holds. Two early methods, FTCP [Ren et al. 2022] and the method in Kim et al. [2020], propose to generate the lattice parameters and fractional coordinate matrices. Specifically, lattice parameters are ℓ_2 -norms of three lattice vectors and angles between every two lattice vectors, and for a crystalline material $M = (z, C, L)$, its fractional coordinate matrix is defined as $F = L^{-1}C$. It can be easily demonstrated that lattice parameters and fractional coordinate matrices are invariant to rotation and reflection transformations. However, fractional coordinate matrices assume an order among atoms so they are not permutation-invariant, and they are not invariant to translation and periodic transformations. Instead of directly generating fractional coordinate matrices, two recent methods, CDVAE [Xie et al. 2022a] and SyMat [Luo et al. 2023c], propose to generate crystalline materials in the form of 3D graphs. They use VAE models to generate the aforementioned lattice parameters, initialize atom coordinates randomly, and iteratively refine atom coordinates by score matching models [Song and Ermon 2019]. Particularly, in both methods, the atom coordinates refinement is done by $E(3)$ -equivariant graph neural network models on the multi-edge graph [Xie and Grossman 2018], a 3D graph representation of the crystalline material. Since multi-edge graphs do not assume the order of atoms and are invariant to periodic transformations, their probabilistic modeling of atom coordinates refinement ensures invariance to permutation and periodic transformations. Despite these similarities, CDVAE and SyMat apply score matching to different targets in the coordinate refinement process. CDVAE directly applies score matching to atom coordinates, which fails to achieve translation-invariant. Differently, SyMat applies score matching to pairwise distances between atoms so as to achieve invariance to all $E(3)$ transformations. We summarize the key information and the captured symmetries of all crystalline material generation methods discussed in this section in Table 27.

Table 27. Summary of 3D outputs, model architecture, and the captured symmetries in several representative crystalline material generation methods. Among these methods, Hoffmann and Noé [2019] and Cond-DFC-VAE [Court et al. 2020] generate 3D voxel grids, which are invariant to permutations and periodic transformations, but not invariant to rotations, reflections, and translations. FTCP [Ren et al. 2022] and Kim et al. [2020], generate fractional coordinates and only achieves invariance to rotations and reflections. CDVAE [Xie et al. 2022a] and SyMat [Luo et al. 2023c] generate materials in the form of multi-edge graphs. They both achieve invariance to permutations, rotations, reflections, and periodic transformations. However, CDVAE fails to achieve invariance to translations due to directly applying score matching to coordinates, while SyMat achieves it because it applies score matching to pairwise distances.

Methods	3D Outputs	Architecture	Permutation invariant	Rotation & Reflection invariant	Translation invariant	Periodic invariant
Hoffmann et al. [2019]	3D voxel grids	VAE & U-Net	✓	✗	✗	✓
Cond-DFC-VAE	3D voxel grids	VAE & U-Net	✓	✗	✗	✓
FTCP	Fractional coordinates	VAE	✗	✓	✗	✗
Kim et al. [2020]	Fractional coordinates	GAN	✗	✓	✗	✗
CDVAE	Multi-edge graphs	VAE & Score matching	✓	✓	✗	✓
SyMat	Multi-edge graphs	VAE & Score matching	✓	✓	✓	✓

Table 28. Some statistic information of Perov-5, Carbon-24, and MP-20 datasets [Xie et al. 2022a]. We summarize the number of 3D molecule samples (# Samples), maximum number of atoms in one molecule (Maximum # atoms), and average number of atoms in one molecule (Average # atoms).

Datasets	# Samples	Maximum # atoms	Average # atoms
Perov-5	18,928	5	5.0
Carbon-24	10,153	24	9.2
MP-20	45,231	20	10.4

7.3.4 Datasets and Benchmarks.

For a long time, there are no standard benchmark datasets for the material generation task. Early material generation methods are usually evaluated on manually selected materials from the Materials Project database [Jain et al. 2013] or other data libraries. Until recently, Xie et al. [2022a] curate three benchmark datasets Perov-5, Carbon-24, and MP-20 for the evaluation of different material generation methods. Perov-5 dataset collects 18,928 perovskite materials from an open material database for water splitting [Castelli et al. 2012b,a]. All materials in Perov-5 have 5 atoms in a unit cell. Carbon-24 dataset collects 10,153 materials whose 3D structures are optimized by AIRSS [Pickard and Needs 2006, 2011] at 10 GPa. All materials in Carbon-24 only contain carbon atoms and have up to 24 atoms in a unit cell. MP-20 dataset is composed of 45,231 materials whose energies above the hull and formation energies are smaller than 0.08 eV/atom and 2 eV/atom, respectively. All materials in MP-20 are obtained from Materials Project database and have up to 20 atoms in a unit cell. See Table 28 for some statistical information of these three datasets.

7.3.5 Open Research Directions.

Though several crystalline material generation methods have been proposed recently, some challenges remain unsolved and prevent them from practical use. First, recent methods including CDVAE and SyMat are based on score-matching models. They achieve better performance than earlier methods, but take much higher computational costs in refining atom coordinates for thousands of iterations by score-matching models. An efficient generative model that can generate good material samples with a reasonable time cost is desirable in real-world applications, but designing such a

model remains challenging. Second, it is important to ensure that the crystalline materials generated by models are practically synthesizable. However, to our knowledge, no standard evaluation metric has been used to measure the synthesizability of crystalline materials generated by generative models in the literature. Introducing such synthesizability metrics can be useful in filtering out materials that cannot be practically synthesized, and it is also interesting yet challenging to design novel material generation methods that can optimize the synthesizability metrics of their generated materials.

7.4 Materials Characterization

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Recommended Prerequisites: Section 7.1

In the last two sections, we described the ML methods for predicting materials' properties based on their crystal structures as well as for generating new crystal structures. However, perhaps a more fundamental challenge is the accurate and efficient experimental determination of crystal structures. Beyond long-range ordering, a spectrum of local disorders from short-range order in amorphous materials to correlated disorders in Prussian Blue analogs can manifest and influence materials' properties [Simonov and Goodwin 2020; Kholina et al. 2022]. Therefore, to determine the exact crystal structure of a material, typically, a combination of instruments and characterization techniques such as X-ray and neutron scattering and spectroscopies are used.

The measurements are usually done in laboratories or at large-scale facilities (e.g., synchrotron beamlines) and require time-consuming and careful modeling of the data. To give more perspective, modern X-ray detectors can generate as many as 1,000,000 images per day, and data post-processing, analysis, and interpretation of the experiments can take over a year [Doucet et al. 2020; Chen et al. 2021b; Wang et al. 2017b]. The ability of machine learning methods to process big data, and find patterns in complex data, and the computer vision algorithms for the autonomous detection of images can play a significant role in accelerating the existing workflows at beamline user facilities by providing immediate feedback during the experiments [Wang et al. 2017b; Doucet et al. 2020; Sullivan et al. 2019; Yanxon et al. 2023; Wang et al. 2017b; Banko et al. 2021; Özer et al. 2022; Venderley et al. 2022].

Here, we discuss some of the challenges and opportunities of integrating machine learning methods with characterization techniques. We specifically review the existing works on using scattering and spectroscopy techniques to predict the average and local crystal structures (and their inverse problem). We note that, due to the huge variation and complexity of the characterization methods, we only scratch the surface of possibilities.

7.4.1 Problem Setup.

As discussed in the overview, the potential applications of ML in materials characterization methods are huge due to the inherent diversity of materials characterization methods. Therefore, we focus our problem setup into two main categories:

- Crystal structure prediction: This category involves using the output of the experimental characterization techniques, such as the one-dimensional (1D) spectra of X-ray diffraction, to predict three-dimensional (3D) crystal structures, which can be described by atomic positions along with three lattice vectors or other crystal structure parameters such as crystal systems, Bravais lattice types, unit cell lengths, and cell angles.
- Its Inverse Problem: The inverse of the aforementioned procedure forms the second category, whereby predicting the output of the characterization methods using the crystal structure.

For example, using the crystal structure of materials, such as atomic positions and lattice vectors, to reconstruct one-dimensional (1D) and two-dimensional (2D) diffraction spectra.

7.4.2 Technical Challenges.

The process of reconstructing a 3D crystal structure or relevant parameters from a scattering pattern is not trivial. The scattering signal can be described by a complex wave function, where the amplitude represents the magnitude of the scattering and the phase represents the position and arrangement of the scattering centers. However, in practice, only the intensities (square of the amplitude) are measured or recorded. Without the phase information, a diffraction pattern may not map uniquely to the crystal structure and vice versa. This is known as the phase problem) [Sivia et al. 1991]. The problem is further complicated as scattering data is obtained on a per-material basis. The experimental data can vary depending on the material, quality of the sample, and what scientific questions experimentalists are asking. Thus, there could be many modalities of data per experiment. It is thus crucial to develop ML models that generalize to experimental as well as simulated data.

As earlier stated in this section, crystals are periodic and highly symmetric structures. For example, for the inverse problem of reconstructing scattering data from crystal structures, equivariance should be preserved (e.g., applying an element of a crystal's symmetry group to the input will lead to the same scattering output). Thus, these problems require the design of symmetry preserving and equivariant ML methods, which is challenging.

7.4.3 Existing Methods.

Spectral Data as Input: First, we describe existing work that provides spectral data as input to an ML algorithm. Most existing work uses this to either classify the crystal symmetry (crystal class, Bravais lattice, or space group) or to find the crystal lattice parameters in a regression problem. The structure of a crystalline material can be described by three lattice vectors and a unit cell. The unit cell can be specified according to six lattice parameters which are the lengths of the cell edges ($\|\ell_1\|, \|\ell_2\|, \|\ell_3\|$) and the angles between them (α, β, γ). In 3D space, there are seven crystal classes and 14 Bravais lattices.

Many previous studies have used CNNs for lattice type classification or lattice parameter regression. Garcia-Cardona et al. [2019] develop a CNN classifier to predict Perovskite crystal systems and lattice parameters $\{\|\ell_1\|, \|\ell_2\|, \|\ell_3\|, \alpha, \beta, \gamma\}$ from neutron scattering data. They then train a random forest regression model for each of the crystallographic symmetries studied to predict the lattice parameters. Both models yielded lower accuracy for lower symmetry crystal systems. They concluded that more sophisticated models were necessary to handle experimental data. Chitturi et al. [2021] assume that the crystal systems are already known and use 1D CNNs to predict lattice parameters for each crystal system. The 1D CNNs were able to predict unit cell lengths but unable to predict unit cell angles of monoclinic or triclinic systems (lower symmetry classes). Corriero et al. [2023] also use CNNs and random forests to predict the crystal system and space group.

While CNNs may be a popular choice, Greasley and Hosein [2023] demonstrate that more conventional supervised learning algorithms such as Support Vector Machine (SVM) and Complement Naive Bayes (CNB) classifiers performed equally as well to a neural network for multi-phase identification with experimental and simulated XRD spectra. Other groups have also employed variational autoencoder (VAE) architectures. The latent space of a VAE is a "compressed" representation of the training samples, so this can be an effective strategy for learning meaningful, continuous representations of materials properties from scattering data. For example, Banko et al. [2021] find that the latent space provides direct visual evidence of the clustering properties of the encoder model and distribution of the main reflection axes in the XRD patterns.

However, it is crucial to note that these methods all take 1D spectra as input. A CNN for example assumes translational invariance in the input data, a feature that may not be present in these spectra. Thus, the symmetries and assumptions of ML models should be considered before applying them to powder spectra. Additionally, it would be worthwhile to develop methods that utilize equivariance in a more meaningful way.

Spectral Data as Output: The inverse problem, such as using structures or other physical properties to predict diffraction patterns, has not been studied as extensively using ML techniques. [Cheng et al. \[2023b\]](#) develop an ML-based framework that can predict both one-dimensional and two-dimensional inelastic neutron scattering (INS) spectra from the structure (atomic coordinates and elemental species). This study extends work done in [Chen et al. \[2021a\]](#) using equivariant neural networks to predict the phonon density of states. They first employ an autoencoder to represent the 2D spectrum (a function of momentum and energy transfer) in latent space, as otherwise there would be 300 x 300 map in momentum/energy space to predict. They then use a Euclidean neural network for feature prediction in the latent space and reconstructed the 2D spectrum. Note this approach could also be applied to 1D spectra, perhaps without the autoencoder step. Equivariant neural networks thus represent a promising avenue for the inverse problem of reconstructing spectral data from crystal structures.

7.4.4 Datasets and Benchmarks.

Using synthetic data for training X-ray/neutron scattering ML models is inevitable due to the lack of available experimental data. However, experimental data are often quite different than simulated data. Simulated datasets with structural information include the Materials Project [\[Jain et al. 2013\]](#), the Inorganic Crystal Structure Database (ICSD) [\[D et al. 2019\]](#), and the Cambridge Structural Database (CSD) [\[Groom et al. 2016\]](#). One approach to developing more generalizable models is to perform data augmentation on simulated datasets to address possible experimental factors such as peak shift, broadening, texture, and noisy background. Another approach is to train on simulated data but test on experimental data (an avenue that, as expected, doesn't tend to produce satisfactory results). To our knowledge, there does not exist an experimental dataset for benchmarking performance of ML algorithms for X-ray/neutron scattering due to the wide variety of experimental setups and issues investigated. For developing robust ML methods, such benchmarking datasets should be created.

7.4.5 Open Research Directions.

In the future, it would be useful to build models that employ the principles of symmetry and Fourier transforms to effectively represent scattering data in either real space, reciprocal space, or both. A constant theme in scattering experiments is the acquisition of data in reciprocal space to inform something traditionally represented in real space. Neural networks that operate in the frequency domain have been developed. [Li et al. \[2021b\]](#) develop a neural network architecture defined in Fourier space and [Yi et al. \[2022\]](#) extend this to perform graph convolutions in the Fourier domain. This approach could potentially be synthesized with equivariant neural networks and applied to this problem. Due to the phase problem, an equivariant neural network that understands Fourier space and can exchange information with real space could be quite powerful.

Another future direction could be exploring different ways of representing materials through equivariant operations and how these relate to powder spectra. This could lend itself to a different equivariant ML approach. In general, current models perform worse with lower symmetry structures (classifying and predicting lattice parameters) as well as experimental data, so this should be addressed in future work.

7.5 Local Structure and Disordered Materials Characterization

Authors: Aria Mansouri Tehrani, Tuong Phung, Yuchao Lin, Tess Smidt

Recommended Prerequisites: Sections 7.1, 7.4

The crystallographic methods we have discussed in the last section are useful for creating structural models for the average atomic positions. However, they neglect that crystalline materials can possess disorders (random or correlated). Disorder has been shown to exist and significantly influence the property of crystalline materials [Simonov and Goodwin 2020; Kholina et al. 2022; Venetos et al. 2023; Cheetham et al. 2016]. Additionally, short-range order is even more vital in materials with limited or without long-range orders, such as nanostructures and amorphous materials. [Li et al. 2020c; Martin et al. 2002] One approach to probe the local structure uses atomic pair distribution function (PDF) analysis, which is the Fourier transform of total scattering from X-ray, neutron, or electron scattering of powder or single crystal samples [Young and Goodwin 2011; Kjær et al. 2023; Liu et al. 2019]. Alternatively, spectroscopies analysis such as nuclear magnetic resonance (NMR) can provide further insight into the local structures [Venetos et al. 2023].

7.5.1 Problem Setup.

In this task, models are taking advantage of the results of characterization techniques that probe the local structures of materials, e.g., the pair distribution function (PDF) and nuclear magnetic resonance (NMR), to predict or generate the structures of materials, including three-dimensional (3D) atomic positions within the laboratory coordinate system [Kjær et al. 2023], or chemical properties corresponding to the local atomic frame of reference, such as magnitude, anisotropy and orientation of NMR chemical shift tensor [Venetos et al. 2023], which describe how a nucleus is influenced by the electronic environment surrounding it.

7.5.2 Technical Challenges.

Solving the short-range order in materials is a notoriously challenging task experimentally and computationally. For example, incorporating even small disorders such as site-sharing or point defects in DFT calculations requires expensive supercell calculations of many different configurations. These calculations are almost impossible for amorphous materials, therefore hindering the possibility of creating large training data for ML. Consequently, ML models cannot rely on standard computational data. Unfortunately, experimental characterizations of short-range order are also not trivial. Some methods to gain insights into local structures are PDF analysis of powder diffraction, 3D-ΔPDF modeling of single crystal diffraction (using the Yell computer program), or merging scattering and spectroscopic techniques [Simonov et al. 2014; Venetos et al. 2023]. The total scattering techniques require large quantities of phase pure powder samples suitable for neutron or high-quality single crystal samples, high-intensity neutron or X-ray sources at large facilities, domain and beamline experts to perform the experiments, and arduous refinements of the diffuse scattering. On the ML side, it is therefore critical to construct representations that can effectively capture the local atomic structure, develop models that are data efficient, and can predict tensorial properties.

7.5.3 Existing Methods.

Ab initio solving of crystal structures from atomic pair distribution function is extremely challenging and so far has only been done for highly symmetric nanostructures [Juhás et al. 2006]. Recent work has developed a deep generative model called DeepStruc, that can solve a simple monometallic nanoparticle structure directly from a PDF using a conditional variational autoencoder [Kjær et al.

2023]. PDF, also known as $G(r)$, which represents the histogram of real-space interatomic distances, is defined as

$$G(r) = 2/\pi \int_{Q_{\min}}^{Q_{\max}} Q[S(Q) - 1] \sin(Qr) dQ,$$

where Q is the scattering vector, and $S(Q)$ is the total scattering structure function that depends on the measured X-ray scattering intensities and the atomic form factor. And the structures of monometallic nanoparticles are represented as graphs, $\mathcal{G} = (X, A)$, where $X \in \mathbb{R}^{N \times 3}$. Here, X is the node feature matrix, and the interatomic connections are described by the adjacency matrix $A \in \mathbb{R}^{N \times N}$. On this basis, they utilize conditional deep generative models to synthesize data conditioned on PDF by solving the unassigned distance geometry problem (uDFO) and in essence capturing the relationship between the atomic structures and PDF. Finally, they show that by using experimental data, their model can successfully predict the crystal structures of some simple monometallic nanoparticles [Juhás et al. 2006].

Beyond scattering techniques, nuclear magnetic resonance (NMR) is a powerful spectroscopy tool that is often used in conjunction with powder X-ray diffraction to elucidate the local environment of materials. The NMR chemical shift tensor encodes both the average electronic environment of an atom represented by chemical shift as well as anisotropies that contain additional structural data. These anisotropies are evident by the line shape in an NMR measurement and can be used to infer the local chemical bondings [Venetos et al. 2023]. Exploiting the advances in equivariant geometric deep learning methods to directly predict tensorial properties while preserving the input symmetries, researchers have recently developed a model to predict Si chemical shift tensors in silicates [Venetos et al. 2023]. In this work, the rotational equivariance is implemented using the MatTEN package, which utilizes the tensor field network and e3nn while, for comparison, symmetry-invariant models have also been constructed. The result shows that equivariant models outperform symmetry invariant ones by 53 %, highlighting the application of equivariant geometric deep learning models in predicting symmetry-dependent tensorial properties in characterization measurements.

7.5.4 Datasets and Benchmarks.

Since there is a lack of widely recognized benchmarks within this specialized field, datasets are often simulated and self-generated to develop ML models. An example is the PDF dataset [Kjær et al. 2023], where a total of 3,742 structures of monometallic nanoparticles were generated through the atomic simulation environment (ASE) alongside their corresponding PDF. Particularly, seven types of monometallic nanoparticle structures, such as simple cubic (sc), body-centered cubic (bcc), and face-centered cubic (fcc), among others, are included. These are constructed across a size range spanning from 5 to 200 atoms. Additionally, a subset of *ab initio* NMR chemical shift tensors of relaxed structures computed by Sun et al. [2020] is used in Venetos et al. [2023], which comprises 421 unique silicate structures, with 1,387 unique silicon sites and different numbers of bridging oxygen atoms. These examples underline the opportunity of creating more comprehensive datasets in the future, which would help facilitate this line of research.

7.5.5 Open Research Directions.

Since the local atomic environments in a material play a significant role in determining its overall properties, understanding the complex interplay between local environment geometries and material properties is crucial to designing next-generation materials with desired properties. We would like a local environment descriptor that is invariant to translation, rotation, and permutation of atoms of the same species, as these symmetry operations do not change physical properties.

Additionally, recent work has introduced a metric called local prediction rigidity (LPR) to assess to what extent the global quantities can be rigorously be assigned to the local, atom-centered contributions [Chong et al. 2023].

We can consider spherical harmonics (detailed in Section 2.7) as a natural starting point for coming up with such a descriptor. Spherical harmonics are a very nice set of basis functions for signals on the sphere and are well-suited for describing local atomic environments. This is because atoms don't like to be too close to each other, so they naturally spread across a sphere. Consequently, higher degrees of spherical harmonics are not needed in order to capture a local environment due to the angular spacing of atoms. They also transform as the irreducible representations of $SO(3)$, making them invariant under rotations. Considering this, spectra, which are quantities that can be computed from spherical harmonic coefficients, seem like a natural choice to characterize geometry.

Given a local environment, one can express it as a sum of radial Dirac delta functions as

$$\sum_{i=1}^N v_i \delta(\mathbf{r}_i).$$

From this function, a spherical harmonic signal \mathbf{x} is obtained by expanding this function into its spherical harmonic coefficients as

$$\mathbf{x} = a_{\ell,m} = \int \left(\sum_{i=1}^N v_i \delta(\mathbf{r}_i - \mathbf{x}) \right) Y_m^\ell(\mathbf{x}) d\mathbf{x} = \sum_{i=1}^N v_i Y_m^\ell(\mathbf{r}_i).$$

Taking this spherical harmonic signal x , one can calculate the spectra of order d by computing repeated symmetric tensor products ($x^{\otimes(d+1)}$) and extracting the scalar and pseudoscalar coefficients ($x^{\otimes(d+1)} \rightarrow (0, e) \oplus (0, o)$). The first, second, and third order spectra are more commonly known as the power spectrum, bispectrum, and trispectrum, respectively. The bispectrum in particular is effective in characterizing local environments, being more expressive than the power spectrum but less computationally expensive than the trispectrum.

Spectra also have other nice properties including smoothness (small perturbations to the original geometry lead to small perturbations in the resulting spectra), invertibility (the original geometry can be decoded up to a global rotation), and being fixed-length (for a given value of l). They can also be clustered (e.g., using k-means clustering), enabling the identification of geometric trends within a given class of materials or across different material classes.

Beyond further developing appropriate, expressive local representations, an essential avenue moving forward is to utilize the power of ML to accelerate the time-consuming characterization processes of disordered materials. For example, computer vision can be useful for the rapid detection of Bragg peak shapes, making corrections, and identifying artifacts, among others, while in some cases, the reliability of equivariant neural networks to preserve symmetry can be exploited to incorporate domain knowledge.

7.6 Phonon Calculations

Authors: Adriana Ladera, Tess Smidt

Recommended Prerequisites: Section 7.1

A phonon is the quantization of energy of vibrations in a lattice, analogous to photons being the quantization of the electromagnetic wave [Kittel 2004]. Calculations of these phonons are crucial to understanding the thermal and dynamical properties of materials. First principles phonon

calculations have become especially available due to advances in efficient density functional theory (DFT) codes and high-performance computing, but experimental and computational challenges in efficient phonon calculations still remain. In this section, we detail current obstacles (force prediction accuracy, limited resources, periodicity of crystallographic materials complicating training data and the learning of current models), existing methods (equivariant neural networks for direct prediction of phonon density of states, moment tensor potentials trained on *ab initio* molecular dynamics trajectories), and promising future directions for the integration of ML in phonon studies.

7.6.1 Problem Setup.

In phonon calculations, crystal structures, including atomic positions, atomic species, and lattice vectors of the unit cell, are given to calculate phonon properties, which comprise computations such as the phonon dispersion relation (PDR) (commonly known as the phonon band structure), and the phonon density of states (PDOS). PDR relates the phonon momentum, normally along a high symmetry path in the Brillouin zone, and the angular frequency of the phonon in each branch. It is significant for studying phonon-related properties, such as phonon-phonon interactions and electron-phonon coupling, with applications in thermal and electronic transport. PDOS summarizes phonon dispersions by integrating over the wave vector and summing over each branch. Formally, PDOS is defined as

$$g(\omega) = \frac{1}{N} \sum_{q_j} \delta(\omega - \omega_{q_j}), \quad (103)$$

where N is the number of unit cells in the crystal, q is the wave vector j is the band index, and ω_{q_j} is the phonon frequency [Togo and Tanaka \[2015\]](#). PDOS is important in understanding several material properties, such as superconductivity, electrical transport, and vibrational properties. When predicting PDOS, PDR, and other phononic calculations, graph neural network models often have no prior knowledge of interatomic forces and other characteristics are used, other than atomic positions, masses, and atomic species, whereas in ML interatomic potentials (MLIPs), the models are trained on *ab initio* molecular dynamics trajectories.

7.6.2 Technical Challenges.

For studying thermal properties and PDR, DFT simulations offer accurate approximations, but the computational cost quickly increases in the case of nanoporous and low-symmetry materials. Lower k-point grids and smaller supercells and plane-wave cutoff energy are often used to bypass computational constraints, but unsurprisingly resulting in the poor accuracy of the yielded PDR. Additionally, computational conditions can still produce nonphysical frequencies in phonon dispersion diagrams [\[Mortazavi et al. 2020\]](#). Obtaining PDOS via both experimental and computational methods poses a challenge, as *ab initio* calculations for complex materials demand high computational costs and inelastic scattering often requires limited resources [\[Chen et al. 2021a\]](#) such as high flux neutron sources or synchrotron X-rays [\[Hanus et al. 2021\]](#).

Several recent advances in machine learning for material science suggest a new paradigm for materials studies. However, the 3D nature of atomic systems and the periodicity and symmetry of crystallographic materials further complicate any potential learning of PDOS for a regular neural network, as this requires expensive data augmentation to learn different rotations and translations of the 3D coordinate systems. These problems therefore highlight the need for a more efficient strategy for obtaining PDOS. Turning towards ML methods, GNNs represent atoms and their bonds as graph nodes and edges, respectively, offering a natural representation of atomic systems. Symmetry-augmented GNNs [\[Geiger and Smidt 2022\]](#) additionally, hold an advantage due to the innate symmetry of crystal materials. Challenges in representation and neural network design choices, still persist, however, such as the difference between properties in real space and reciprocal

space, and the fixed length of output properties (*i.e.*, scalar outputs) [Delaire et al. 2009]. This is problematic for materials properties with varying degrees of dimensions, such as the number of phononic bands [Baroni et al. 2001].

7.6.3 Existing Methods.

In this section, we categorize approaches for efficient and accurate prediction and analysis of phononic properties of materials in terms of using local environment descriptors and geometric graph neural networks.

As a prime example of local environment descriptors, Mortazavi et al. [2020] utilize ML interatomic potentials (MLIPs) to train on computationally efficient *ab initio* molecular dynamics trajectories, providing an alternative and efficient method to DFT simulations. In MLIPs, the potential energy surface is described as a function of the local environment descriptors which are invariant to rotations, translations, and inversions of homonuclear atoms [Behler 2016]. Under the umbrella of MLIPs are moment tensor potentials (MTPs) [Zuo et al. 2020], which can approximate any interatomic interactions [Mortazavi et al. 2020], therefore able to evaluate phononic properties comparable to density functional perturbation theory methods without being computationally expensive. In addition, Ladygin et al. [2020] use MTPs to reproduce phonon properties with high accuracy compared to DFT-obtained data. Using active learning of MTPs as an advantage, training is conducted on molecular dynamics runs of Al, Mo, Ti, and U. The active learning approach automatically fits the MTP only on configurations in which there is significant extrapolation in data, which greatly reduces the number of DFT calculations required for training the MTP. The error between MTP and DFT results for PDOS of Al and U and phonon dispersion diagrams of Mo and Ti is far smaller than the error between DFT and experimental data. Similar errors are produced when comparing MTP and DFT for vibrational free energy and entropy results.

The geometric graph neural network learns material properties from 3D atomic positions, atomic species, and interatomic distances based on graph neural networks. For instance, Okabe et al. [2023] propose to augment GNNs with their Virtual Node Graph Neural Network (VGNN), which manages output properties with variable or even arbitrary dimensions. Okabe et al. [2023] present three versions of VGNN, each implemented with the symmetry-aware graph Euclidean convolutional neural networks [Geiger and Smidt 2022]. With the atomic positions and masses represented as a periodic graph, this input is passed through a series of convolution layers, which compute the tensor product of the input features, separated by nonlinear layers, which introduce the complexity to the model. Vector virtual nodes (VVN) is the simplest model with m -atom crystal structure input and $3 \times m$ Γ -phonon energies. The matrix virtual nodes (MVN) are slightly more accurate and computationally expensive for complex materials. Lastly, the momentum-dependent matrix virtual nodes (k-MVN) is the most complex and, given random k -points in the Brillouin zone, is able to predict the entire phonon band structure. This is done with virtual dynamical matrices 14 (matrices analogous to the phonon dynamical matrices) in the crystal graphs. Training data of the materials then enables the matrix elements to be learned from optimizing the neural network. Each of the listed models has no prior knowledge of interatomic forces and only takes the crystal structure as input. This methodology provides a computationally feasible strategy for obtaining full phonon band structures from the crystal structures of complex materials. Chen et al. [2021a] capture the main features of PDOS using a Euclidean neural network in 3 dimensions ($E(3)$ NN), as implemented by the $E(3)$ NN open-source repository [Geiger and Smidt 2022]. The $E(3)$ NN is equivariant to 3D rotations, translations, and inversion, and therefore is able to preserve all geometric inputs as well as their crystallographic symmetries. The dataset is a phonon database of 1,521 crystallographic semiconductor compounds, containing PDOS data-based density functional perturbation theory [Petretto et al. 2018]. $E(3)$ NN successfully reproduces key features in experimental data and predicts

a broad number of high phononic specific heat capacity materials directly from atomic structure without being computationally expensive.

7.6.4 *Datasets and Benchmarks.*

To the best of our knowledge, there does not exist a standard database for phonon calculations that are explicitly for ML methods. From the representative works, however, such datasets include a Γ -phonon database with approximately 146,000 materials from the Materials Project [Okabe et al. 2023] and a database of the PDOS data based on DPFT from 1,521 semiconductor compounds [Petretto et al. 2018]. However, the onset of ML applications in phonon calculations demands a standard database for material phonon data from which ML methods could be easily applied.

7.6.5 *Open Research Directions.*

While the main focus of lattice dynamics studies in the work of Ladygin et al. [2020] are single-component systems, there is promising work in multi-component systems as well. Specifically, MTP performs well in covalently and ionically bonded systems [Grabowski et al. 2019] as well as metallic systems [Novikov and Shapeev 2019]. Additionally, the Euclidean neural network workflow of Chen et al. [2021a] highlights a framework that could aid in high-throughput screening in the search for promising thermal materials candidates, while drawing connections between the structural symmetry of materials and their phononic properties.

8 AI FOR MOLECULAR INTERACTIONS

As described in Sections 5, 6, and 7, AI has revolutionized the field of molecular learning, protein science, and material science. While AI for individual molecules has been extensively studied, the physical and biological functions of molecules are often driven by their interactions with other molecules. In this section, we further introduce AI for molecular interactions, where we particularly consider the interactions between small molecules and proteins or materials.

8.1 Overview

Authors: Meng Liu, Shuiwang Ji

This research area focuses on using AI to gain insights into the molecular mechanisms that govern interactions between small molecules and other substances, which has great potential in advancing our understanding of molecular interactions and providing practical solutions for a wide range of challenges in life science and material science.

For both molecule-protein and molecule-material interactions, as illustrated in Figure 30, we categorize existing tasks into predictive tasks and generative tasks. Note that our categorization is based on the nature of the tasks, rather than the methods employed to perform them. To be specific, the predictive task related to the interaction of small molecules and proteins is binding (or docking) prediction. Binding refers to the process by which a small molecule, namely as a ligand, binds to a target protein based on their native shape complementary and chemical interactions [Fischer 1894], also known as the “lock-key” model. The target protein is usually associated with human disease. The drug molecule binds to the target protein to inhibit or activate it to treat human diseases. This task includes binding pose prediction and binding affinity prediction. On the other hand, the generative task for such interaction is to generate molecules that can bind to given target proteins, known as structure-based drug design [Anderson 2003]. Both protein-ligand binding prediction and structure-based drug design are fundamental and challenging problems in drug discovery. In terms of the interaction of small molecules and materials, to our knowledge, only predictive tasks have been investigated in existing works. Specifically, we are interested in predicting adsorption energy (S2E) and per-atom force (S2F) from a molecule-material pair structure. In addition, given the initial structure of a molecule-material pair, it is highly desired to predict the relaxed final structure (IS2RS) and the adsorption energy at its relaxed state (IS2RE). These tasks are critical for many problems in material science, such as electrocatalyst design for renewable energy storage [Zitnick et al. 2020]. The generative tasks for molecule-material interactions remain unexplored, and we discuss the potential opportunities in Section 8.4.5. An overview of the methods covered in this section is shown in Figure 31.

Ensuring the preservation of desirable symmetry in 3D space is a crucial aspect of molecular interaction tasks. This unique symmetry is distinct from the consideration of a single molecule, as it encompasses multiple instances involved in the interaction. In particular, it is essential to take into account the symmetry properties of each molecule within the context of the entire interaction system.

8.2 Protein-Ligand Binding Prediction

Authors: Hannes Stärk, Yuchao Lin, Shuiwang Ji, Regina Barzilay, Tommi Jaakkola

Recommended Prerequisites: Sections 5.2, 6.3

In this section, we study the protein-ligand binding problem. We aim to make inferences about how a small molecule, potentially a drug, interacts with a protein. For this purpose, we discuss

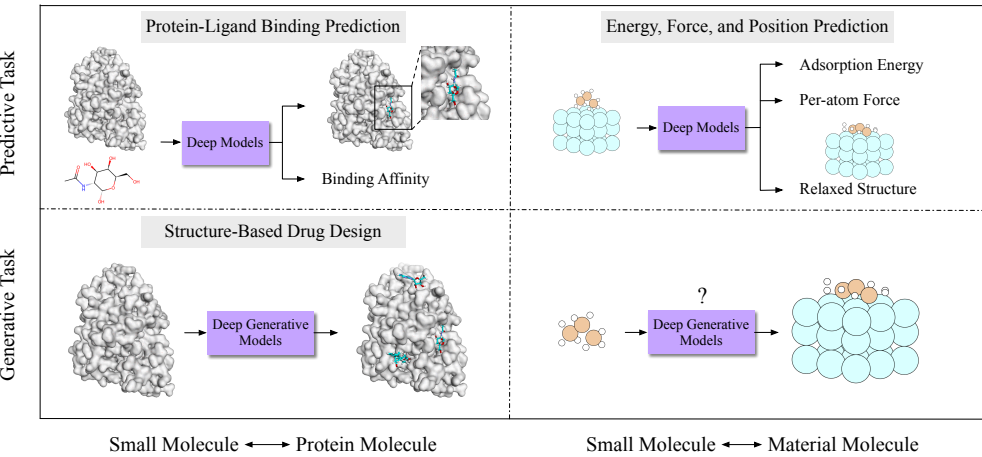


Fig. 30. An illustration of our covered tasks in molecular interactions. The bidirectional arrows represent interactions. For both molecule-protein interaction and molecule-material interaction, we categorize existing tasks into predictive tasks and generative tasks. In protein-ligand binding prediction, we aim to predict the binding pose of the ligand and the strength of the binding, namely binding affinity. In structure-based drug design, it is desired to generate 3D ligand molecules that can bind to the given target proteins. In the predictive tasks of molecule-material pairs, we are interested in predicting the adsorption energy and per-atom force vectors with given molecule-material pair structures. In addition, it is of interest to predict the relaxed final structure given the initial structure as input. The generative tasks for molecule-material interactions are unexplored and we discuss the possible directions in Section 8.4.5

molecular docking and binding affinity prediction, both of which are important for fields such as drug discovery or molecular biology.

8.2.1 Problem Setup.

In docking, we are given a protein structure (its amino acid identities and atom coordinates) and the molecular graph of a small molecule (ligand). The goal is to predict the atom positions with which the ligand most likely binds to the protein. This task can be divided into the scenario where the docking location (pocket) is approximately known and the blind docking scenario without any prior knowledge. Performing well at either means having a high fraction of approximately correct predictions. Meanwhile, with binding strength prediction, we refer to predicting a ranking or scalar binding affinity that indicates the strength with which a ligand binds to a protein; that is, roughly the fraction of times the ligand and protein can be observed in a bound vs. unbound state. The inputs for this could, e.g., be the bound protein-ligand structure or a protein structure and the ligand's molecular graph.

In both docking and binding strength prediction, it is important to consider the discussed nuances of the problem setup. Additionally, in both tasks, it is relevant whether one has access to the protein's structure when bound to the ligand (holo-structure), the unbound protein structure (apo-structure), the structure of the protein bound to another ligand, or only a computationally generated structure from, e.g., AlphaFold2. In docking evaluations, knowledge of the bound structure is often assumed, which is unrealistic for application purposes, and comparisons in the other scenarios are desirable.

To present docking and binding strength prediction in a formal context, the structure of a ligand can be expressed as $\mathcal{M} = (A, E, C)$, where $A = [a_1, \dots, a_n]$ refers to the atomic properties, for

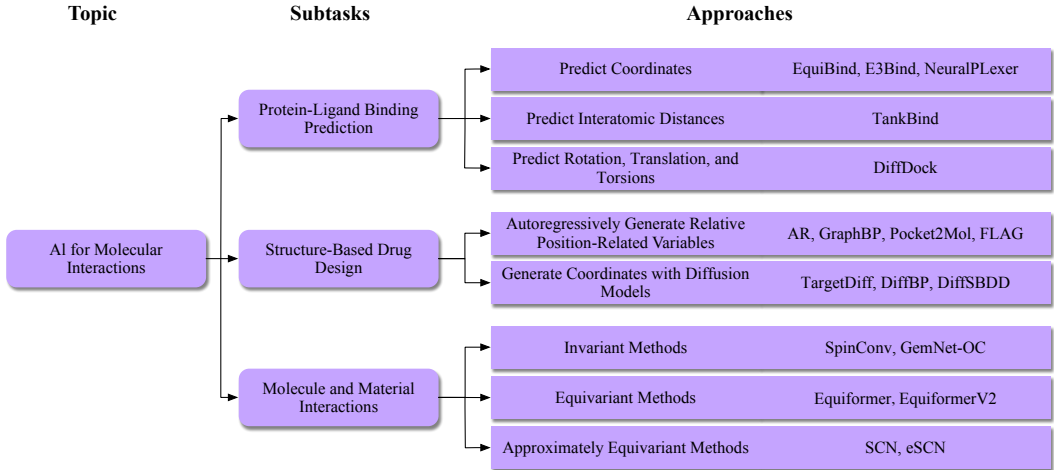


Fig. 31. An overview of the tasks and methods in AI for molecular interactions. This section considers three tasks, including protein-ligand binding prediction, structure-based drug design, and energy, force, and position prediction for molecule-material pairs. In protein-ligand binding prediction, one category of methods, including EquiBind [Stärk et al. 2022b], E3Bind [Zhang et al. 2023a], and NeuralPLexer [Qiao et al. 2023], aims to directly predict the 3D coordinates of ligands. In comparison, TankBind [Lu et al. 2022] predicts the interatomic distances between protein segments and ligands. Besides, DiffDock [Corso et al. 2022] generates the rotation, translation, and torsion angles of seed conformers given by RDKit. In structure-based drug design, one category of methods, including AR [Luo et al. 2021], GraphBP [Liu et al. 2022c], Pocket2Mol [Peng et al. 2022], and FLAG [Zhang et al. 2023b], aims to generate ligand atoms/fragments autoregressively by modeling their relative position-related variables. Another category of methods, including TargetDiff [Guan et al. 2023], DiffBP [Lin et al. 2022b], and DiffSBDD [Schneuing et al. 2022], considers generating 3D coordinates of all ligand atoms directly via diffusion models. In the prediction tasks for molecule-material pairs, the invariant methods are SpinConv [Shuaibi et al. 2021] and GemNet-OC [Gasteiger et al. 2022], and the equivariant methods include Equiformer [Liao and Smidt 2023] and EquiformerV2 [Liao et al. 2023]. Besides, SCN [Zitnick et al. 2022] and eSCN [Passaro and Zitnick 2023] are approximately equivariant methods.

instance, atomic types, of all n atoms contained in the molecule. The edge features, which could include bond types and bond lengths, are denoted by $E = [\mathbf{e}_1, \dots, \mathbf{e}_l]$ for all l chemical bonds in the molecule. Meanwhile, the 3D coordinate matrix is expressed as $C = [\mathbf{c}_1, \dots, \mathbf{c}_n] \in \mathbb{R}^{3 \times n}$. Similarly, the structure of a protein or a known pocket can be represented as $\mathcal{P} = (B, S)$. Here, $B = [\mathbf{b}_1, \dots, \mathbf{b}_m]$ represents the node features, including either amino acid types or atomic types, of all the amino acids or atoms within the designated protein or pocket, depending on the levels of granularity as detailed in Section 6. Additionally, the 3D coordinates corresponding to either the alpha-carbons of amino acids or the atoms within the structure are symbolized by $S = [\mathbf{s}_1, \dots, \mathbf{s}_m] \in \mathbb{R}^{3 \times m}$. The primary goal in docking is to sample from the distribution of docking poses based on the protein/pocket structure, which can be represented by $p_{\text{pose}}(C|B, S, A, E)$. Methods can either directly model the distribution $p_{\text{pose}}(C|B, S, A, E)$ and then sample, or predict k geometries of ligand conformers such that $f_{\text{pose}}(B, S, A, E) \mapsto [C_1, \dots, C_k]$. On the other hand, the objective of binding strength prediction is to estimate the binding affinity, to provide a ranking, or to make a binary binding vs. non-binding prediction. We unify these targets as q where $q \in \mathbb{R}$ or $q \in \mathbb{Z}^+$ or $q \in [0, 1]$, and the task is represented as $f_{\text{strength}}(A, E, C, B, S) \mapsto q$. Note that for binding strength prediction, the geometric information of ligands or the protein/pocket may not be given.

Table 29. Summary of deep learning methods for blind protein-ligand docking w.r.t. symmetry. EquiBind [Stärk et al. 2022b] and E3Bind [Zhang et al. 2023a] apply $E(3)$ -equivariant networks to predict ligand coordinates. TankBind [Lu et al. 2022] estimates interatomic distances between protein pocket candidates and the ligand together with an affinity for each candidate. DiffDock [Corso et al. 2022] employs an $SE(3)$ -equivariant diffusion model over rotations, translations, and torsions to sample candidates before ranking them. NeuralPLexer [Qiao et al. 2023] predicts a contact map and applies it to an $E(3)$ -equivariant diffusion model to generate ligand coordinates.

Methods	Outputs	Architecture	Network Symmetry
EquiBind	Coordinates	Regression-Based	$E(3)$ -Equivariant
E3Bind	Coordinates	Regression-Based	$E(3)$ -Equivariant
TankBind	Interatomic Distances	Regression-Based	$E(3)$ -Invariant
DiffDock	Rotation/Translation/Torsions	Generative	$SE(3)$ -Equivariant
NeuralPLexer	Coordinates	Generative	$E(3)$ -Equivariant

8.2.2 Technical Challenges.

In our description of molecular docking, we strive to predict the most likely binding pose of a ligand. This is only the global mode of the Boltzmann distribution that describes the probability of each possible ligand pose conditioned on the protein. Ideally, one would want a generative model that replicates this high-dimensional distribution with sparse support, which is especially challenging considering that there usually only is data for one of its modes. Producing the lowest energy (*i.e.*, highest probability) pose is already a difficult problem, given the large space of plausible ligand configurations.

A challenge for docking compared to protein structure prediction is that docking methods cannot rely on vast amounts of sequence data for evolutionary information to constrain the set of plausible structures which partially explains the success of protein structure prediction before geometric deep learning had large impacts on molecular docking. Data concerns also pose another technical challenge; the amount of easily accessible training data with reasonable quality is 20k samples. While there are more complexes in the PDB, it requires expert knowledge and is hard to clean this data from, *e.g.*, complexes that are only spurious interactions with very low affinities.

Similar data concerns impede progress in binding strength prediction with few (20k) data points for 3D structures and noisy measurements (also sequence-based data) preventing a successful affinity predictor for general protein-ligand combinations. However, an already currently useful strategy is training protein-specific predictors that only take ligands as input (under the condition that sufficient binding affinity data is available for the specific protein or can be gathered in an active learning setup). A potential direction towards general protein-ligand affinity predictions could be using geometric deep learning to aid or approximate statistical mechanics methods for calculating binding free energies. This, again, is hampered by the difficulty of modeling the Boltzmann distribution and partitions thereof.

Lastly, we discuss the symmetries involved in the two tasks. Docking is an $SE(3)$ -equivariant task; rotating or translating the input protein structure should result in a corresponding rotation and translation of the generated ligand poses. Technically, for a roto-translation $g \in SE(3)$ and its corresponding group action \triangleright , this task requires $p_{\text{pose}}(g \triangleright C|B, g \triangleright S, A, E) = p_{\text{pose}}(C|B, S, A, E)$ and $f_{\text{pose}}(B, g \triangleright S, A, E) = g \triangleright f_{\text{pose}}(B, S, A, E)$. In contrast, binding strength prediction is an $SE(3)$ -invariant task as rotation and translation to the system do not affect the prediction. Formally, for a roto-translation $g \in SE(3)$, it is desired that $f_{\text{strength}}(A, E, g \triangleright C, B, g \triangleright S) = f_{\text{strength}}(A, E, C, B, S)$.

8.2.3 Existing Methods.

We categorize the protein-ligand docking models into three distinct types: traditional search-based docking, regression-based docking, and generative docking. In order to highlight the recent progress of deep learning made in the field of blind protein-ligand docking, we provide an outline of blind protein-ligand docking methods, specifically with regard to their treatment of symmetry in Table 29.

Traditional Search-Based Docking: Traditional approaches employ an $E(3)$ -invariant scoring function that assigns likelihoods to ligand poses together with an optimization algorithm to find the scoring function's global minimum, *i.e.*, the most likely pose [Trott and Olson 2010; Koes et al. 2013; Halgren et al. 2004]. The most common scoring functions consist of physics-inspired terms of invariant quantities, such as interatomic distances, and use very few learned parameters. More recently, there have been deep learning parameterizations that employ, *e.g.*, 3D CNNs [McNutt et al. 2021]. Importantly, most of these methods are developed for docking to known pockets (with exceptions [Hassan et al. 2017]) and struggle with the larger search space in blind docking, leading to long inference times. Furthermore, their scoring functions are sensitive to deviations of the input protein from the bound structure [Corso et al. 2022; Karelina et al. 2023], which limits their ability for docking to computationally generated or unbound protein structures.

Regression-Based Docking: More recent deep learning methods significantly speed up blind docking by directly predicting ligand binding poses with $E(3)$ -equivariant/invariant GNN instead of parameterizing a scoring function for a search algorithm. Of the regression-based approaches, EquiBind [Stärk et al. 2022b] produces its prediction by finding key points in the protein that characterize the binding pocket and superimposing the ligand with them. Meanwhile, Tankbind [Lu et al. 2022] splits the protein into pocket candidates and predicts protein-ligand distances and an affinity score for each of them. E3Bind [Zhang et al. 2023a] uses the same pocket candidates but produces pair representations for them which are iteratively decoded by updating initial ligand coordinates. A disadvantage of these regression-based methods is that they are forced to predict a single pose even though multiple configurations are plausible and could have a significant likelihood under the Boltzmann distribution. This often leads to unphysical predictions with steric clashes and self-intersections [Corso et al. 2022].

Generative Docking: To resolve the mismatch between the docking task and regression-based solutions, the first proposed generative model is DiffDock [Corso et al. 2022]. Its diffusion model is parameterized by Tensor Field Networks [Thomas et al. 2018] and predicts updates to noisy ligand translations, rotations, and torsion angles before ranking generated samples with a confidence model. Meanwhile, NeuralPLexer [Qiao et al. 2023] predicts a contact map conditioned on which an $E(3)$ -equivariant diffusion model generates ligand coordinates and refolds the protein structure from, *e.g.*, an unbound structure to the bound structure. The mentioned generative models are also able to dock to unbound or computationally generated protein structures with a reasonable degree of accuracy.

8.2.4 Datasets and Benchmarks.

An important dataset of 3D structures of small molecules bound to proteins is PDDBind [Liu et al. 2017] which curates complexes from the Protein Data Bank (PDB) [Berman et al. 2003] if they have binding affinities available and meet additional quality criteria. It consists of 20k complexes with 4k unique proteins. A common dataset split [Stärk et al. 2022b] is based on time with complexes older than 2019 in the training data and newer ones as test data. Less stringent criteria than PDDBind for selecting complexes are applied by BindingMOAD [Wagle et al. 2023], which extracts 40k protein-ligand structures from PDB. APObind [Aggarwal et al. 2021] provides unbound protein structures

for each of its protein-ligand complexes. Helpful for approaches for peptides, Propedia [Martins et al. 2021] extracts protein-peptide complexes from PDB. While the amount of structure data is limited to these magnitudes, there is considerably more protein-ligand binding affinity data without structures in ChEMBL [Mendez et al. 2019] with 20 million activity measurements.

To evaluate docking predictions, it is common to estimate the fraction of correct predictions, which is defined as a generated ligand pose whose RMSDT to a ground truth structure is below a specified threshold. Additionally, the number of steric clashes in the generated structure is a relevant metric. To gauge the performance of binding strength prediction, the metrics depend on the task, such as accuracy for binary classification (binding vs. not binding), ranking correlation for correctly ranking a set of ligands' binding strengths, or MAE for a binding affinity prediction. For evaluation that resembles the real-world docking problem, it is desirable for the field to evaluate docking to unbound or computationally generated protein structures (apo-structures) instead of presuming the holo-structures as input.

8.2.5 Open Research Directions.

While the advances in molecular docking are impressive, the task is still far from solved with, e.g., 22% accuracy in DiffDock when docking to structures from ESMFold. Possible improvements could stem from better generative models for biophysical structures, more meaningful feature embeddings, or more expressive 3D architectures. Nevertheless, the structure prediction capabilities promise a path toward integrating them with downstream binding strength predictors. Unlocking such approaches or similar methods to jointly leverage the available structure data and the larger amounts of sequence-based binding affinity data is promising, and initial successes exist [Moon et al. 2023]. Additional help for accessing these larger amounts of data would be creative, problem-specific approaches for dealing with the noise in affinity measurements.

Furthermore, there is great value in extending molecular docking to additionally model the conformational change of the protein during binding. We think this is more meaningful and realistic as the binding usually changes the conformation of proteins in practice. Lastly, we wish to draw attention to the potential of generative models for statistical mechanics approaches for calculating or comparing protein-ligand interaction strengths/probabilities instead of relying on regressing on experimental affinity measurements.

8.3 Structure-Based Drug Design

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Recommended Prerequisites: Sections 5.2, 6.3

In this section, we consider structure-based drug design (SBDD), a generative task for protein-ligand interaction. In this task, we aim at generating 3D molecules, known as ligands, that can bind tightly to a specific protein (a.k.a. target protein), which can be formulated as a conditional generation problem.

8.3.1 Problem Setup.

Formally, following Section 8.2, we let $\mathcal{M} = (A, E, C)$ denote the ligand molecule and $\mathcal{P} = (B, S)$ denote a protein binding site. Overall, the goal of this task is to learn the conditional distribution $p(\mathcal{M}|\mathcal{P})$ from observed protein-ligand pairs.

8.3.2 Technical Challenges.

The unique symmetry challenge arises due to the fact that this generative task involves multiple molecules interacting with each other, rather than single molecules. This leads to a more complex

Table 30. Summary of existing methods for structure-based drug design in terms of adopted generative approaches, employed networks, level of modeled structures, and 3D output variables. Among these methods, AR [Luo et al. 2021], GraphBP [Liu et al. 2022c], and Pocket2Mol [Peng et al. 2022] generate atoms autoregressively by modeling relative position-related variables, which can be used to determine the position of the new atom. Further, instead of generating atoms, FLAG [Zhang et al. 2023b] considers generating fragments autoregressively. In comparison, TargetDiff [Guan et al. 2023], DiffBP [Lin et al. 2022b], and DiffSBDD [Schneuing et al. 2022] use diffusion models to directly generate 3D coordinates of all atoms in a one-shot schema.

Methods	Generative Approach	Network	Level of Structures	3D Outputs
AR	Autoregressive models	$\ell = 0$	Atom	Distribution of atom occurrence
GraphBP	Autoregressive flow	$\ell = 0$	Atom	Relative distances, angles, and torsions
Pocket2Mol	Autoregressive models	$\ell = 1$	Atom, bond	Relative coordinates
FLAG	Autoregressive models	$\ell = 0$	Fragment	Relative rotation angles
TargetDiff	Diffusion models	$\ell = 1$	Atom	Coordinates
DiffBP	Diffusion models	$\ell = 1$	Atom	Coordinates
DiffSBDD	Diffusion models	$\ell = 1$	Atom	Coordinates

symmetry challenge than modeling individual molecules. Particularly, the symmetries of individual molecules must be considered in the context of their relative positions and orientations with respect to each other. Specifically, if we rotate or translate the protein binding site, the generated molecules yielded by the generative models should be rotated or translated accordingly. Mathematically, the learned conditional distribution should satisfy $p(\mathcal{M}|\mathcal{P}) = p(g \triangleright \mathcal{M}|g \triangleright \mathcal{P})$, where $g \in SE(3)$ and \triangleright represents its corresponding group action. To achieve this, the molecule generated by the model should be equivariant to the $SE(3)$ transformation of the protein.

In addition, this task faces the challenge of an extremely vast search space. To be specific, the chemical space containing all possible molecules is estimated to exceed 10^{60} . In addition, the 3D molecules also have an additional conformation space. However, only a minuscule fraction of this space is relevant to drug discovery, as the molecules need to meet specific criteria to be considered “drug-like”. Thus, how to effectively and efficiently model and explore such space while considering the interactions with target proteins is a fundamental consideration in this task.

8.3.3 Existing Methods.

Early studies either generate molecular SMILES strings conditional on the 3D information of target proteins [Skalic et al. 2019; Xu et al. 2021c], or use estimated docking scores as the reward function to guide the molecule generative model [Li et al. 2021d; Fu et al. 2022a]. They do not explicitly model the crucial interactions between the ligand molecule and the target protein in 3D space. Ragoza et al. [2022] converts protein-ligand complex structures to atomic density grids and then uses generative approaches for 3D image data to tackle the task. A limitation is that the aforementioned equivariance property is not preserved, since 3D CNNs [Ji et al. 2013] is not an $SE(3)$ -equivariant operation for 3D grid data.

Recently, with the development of geometric deep learning and generative modeling, protein-ligand complexes are naturally modeled as 3D geometries and their intricate interactions and symmetry constraints can be effectively encoded. Generally, we can categorize these recent methods into two categories. The first type of method generates atoms autoregressively based on the current context, which includes the binding site and previously generated atoms. To preserve the aforementioned desired $SE(3)$ -equivariance property, at each autoregressive step, these methods consider modeling relative position-related variables of the new atom *w.r.t.* the current context, instead of generating its 3D coordinates directly. To be specific, AR [Luo et al. 2021] uses an

invariant 3D GNN (with feature order $\ell = 0$) to model the distributions of atom occurrence in 3D positions by taking the relative distances between the query position and the current context as input. Using such invariant distances *w.r.t.* the context and invariant 3D GNN together ensures the modeled distribution is equivariant to the rotation and translation of the context. In comparison, GraphBP [Liu et al. 2022c] and Pocket2Mol [Peng et al. 2022] model the relative position of the new atom *w.r.t.* the selected focal atom at each step. Specifically, GraphBP first constructs a local spherical coordinate system (SCS) at the focal atom, which is equivariant to the context’s rotation and translation. It then generates the invariant distance, angle, and torsion *w.r.t.* the reference SCS through an invariant 3D GNN. Pocket2Mol uses an equivariant neural network (with feature order $\ell = 1$) as the encoder and the obtained equivariant features of the focal atom can be used to generate the relative position of the new atom equivariantly. Pocket2Mol also explicitly generates bonds. Instead of using atoms as building blocks, FLAG [Zhang et al. 2023b] considers generating 3D molecules fragment-by-fragment. Such fragment vocabulary can be obtained from chemical priors and can help generate valid and realistic molecules. At each step, FLAG assembles the new fragment to the current context and then predicts the rotation angle of the new fragment *w.r.t.* the selected focal fragment. The $SE(3)$ -equivariance can be preserved by FLAG similarly to GraphBP. Among the above methods, AR, Pocket2Mol, and FLAG are trained via a mask-fill schema, in which atoms or fragments are randomly masked and the model is trained to recover them. In contrast, GraphBP is trained by maximizing the log-likelihood of the trajectory of atom placement steps, thanks to the exact likelihood computation of flow models.

Another line of methods, such as TargetDiff [Guan et al. 2023], DiffBP [Lin et al. 2022b], and DiffS-BDD [Schneuing et al. 2022], considers generating 3D coordinates of all atoms directly. Compared to the above autoregressive sampling methods, such a one-shot generation fashion does not require an order among atoms and can consider global interactions of the entire ligand molecule. Following the framework of EDM [Hoogeboom et al. 2022], these methods apply diffusion models [Ho et al. 2020] in continuous and discrete space to model atom coordinates and atom types, respectively. The denoising step is modeled by an equivariant GNN (with feature order $\ell = 1$) [Satorras et al. 2021b]. To circumvent the difficulty of maintaining translation equivariance in the diffusion process, they shift the Center of Mass (CoM) of the system to diffuse and denoise the coordinates in the linear subspace only. Moreover, with a loose notation, since the latent variables follow a rotationally invariant Gaussian distribution $p(\mathbf{r}_T)$ and the transition distribution $p(\mathbf{r}_{t-1}|\mathbf{r}_t, \mathcal{P})$ is equivariant, the aforementioned equivariance property can be achieved [Köhler et al. 2020].

In addition to deep generative models, reinforcement learning (RL) has also been used for structure-based drug design [Li et al. 2021d; Fu et al. 2022a], which formulates the drug molecule generation process as a Markov decision process (MDP). Unlike generative models that explicitly build the continuous data distribution, reinforcement learning selects the action from discrete space that would receive the maximal reward (*e.g.*, docking score in structure-based drug design). Specifically, DeepLigBuilder [Li et al. 2021d] builds a policy network to select the appropriate actions (whether/where to add atom, which atom to add) to grow the molecule in the target pocket; Reinforced genetic algorithm (RGA) [Fu et al. 2022a] leverages a policy network that selects the discrete action space (mutation, crossover position) of the genetic algorithm (GA) intelligently, which suppresses the random-walk behavior in genetic algorithm.

Worth to mention that earlier molecular optimization methods that are based on SMILES, SELF-IES, or molecular graphs can also achieve competitive performance in structure-based drug design if we incorporate docking score as the goal of optimization [Huang et al. 2021; Gao et al. 2022], including SMILES variational autoencoder (SMILES-VAE) [Gómez-Bombarelli et al. 2018], junction tree variational autoencoder (JTVAE) [Jin et al. 2018], graph convolutional policy network (GCPN) [You et al. 2018b], molecular graph-level genetic algorithm (Graph-GA) [Jensen 2019], graph

autoregressive flow (GraphAF) [Shi et al. 2019], Multi-constraint molecule sampling (MIMOSA) [Fu et al. 2021], etc.

8.3.4 Datasets and Benchmarks.

First, we briefly introduce several structure-based drug design datasets, including CrossDocked2020, PDBBind, DUD-E, and scPDB. (1) CrossDocked2020 [Francoeur et al. 2020] is a widely used benchmark dataset for evaluating the performance of various methods on structure-based drug design. CrossDocked2020 contains an extensive collection of 22,584,102 docked protein-ligand complexes. These complexes are generated through cross-docking, where ligands associated with a specific pocket are docked into each receptor assigned to that pocket by Pocketome [Kufareva et al. 2012], using the smina docking software [Koes et al. 2013]. There are a total of 2,922 pockets and 13,839 ligands covered in CrossDocked2020. Given the variability in the quality of these complexes, it is common in existing studies to include a filtering step. This step involves removing complexes with root-mean-squared deviation (RMSD) of the binding pose that exceeds a certain threshold. This aims to encourage the model to generate ligand molecules with higher binding affinity. (2) PDBBind is an extensive repository derived from the Protein Data Bank (PDB) [Berman et al. 2003], containing experimentally determined binding affinity data for protein-ligand complexes [Wang et al. 2004]. It comprises 19,445 protein-ligand pairs. (3) Directory of useful decoys, enhanced (DUD-E) provides a directory of useful decoys for protein-ligand docking [Mysinger et al. 2012]. It consists of 22,886 protein-ligand complexes and their affinities against 102 distinct protein targets. (4) scPDB is a refined version of the Protein Data Bank (PDB) specifically tailored for structure-based drug design, enabling the identification of optimal binding sites for protein-ligand docking [Meslamani et al. 2011]. It contains 16,034 protein-ligand pairs over 4,782 proteins and 6,326 ligands.

To assess the performance of different generative methods, several categories of metrics are commonly used. The first category involves measuring the quality of the generated molecules, including their chemical validity, novelty, and diversity. Additionally, comparing the distributions of specific variables, such as bond length [Ragoza et al. 2022; Liu et al. 2022c; Peng et al. 2022], bond angles [Ragoza et al. 2022], and the occurrence of different motifs [Peng et al. 2022], between the generated molecules and a reference set can provide further insights into the quality of the generated molecules. The second category aims to estimate the binding affinities between the generated molecules and the target proteins by using the Vina energy or deep learning-based scoring functions [Ragoza et al. 2022]. Comparing the binding affinities of the generated molecules to those of reference molecules helps assess their effectiveness in binding to the target. The last category includes measuring other important properties, such as drug-likeness QED (Quantitative Estimate of Drug-likeness) [Bickerton et al. 2012] and SA (synthesizability accessibility) [Ertl and Schuffenhauer 2009]. [Huang et al. 2021] incorporates an SBDD benchmark that compares five machine learning approaches under the same number of docking oracle calls (5K).

8.3.5 Open Research Directions.

Despite the progress of deep learning approaches in structure-based drug design, how to effectively and efficiently model the vast chemical space to generate valid and synthesizable molecules is still a predominant challenge. Incorporating essential chemical priors, such as motif fragments and scaffolds, could be a direction to tackle this challenge. For example, to enable fragment-based drug design, DiffLinker [Igashov et al. 2022] uses an $E(3)$ -equivariant 3D-conditional diffusion model similar to DiffSBDD to link disconnected molecular fragments (pharmacophores) into a single molecule, while it can take the surrounding protein pocket into consideration as conditional information. In addition, a recent work [Adams and Coley 2022] proposes a shape-based 3D

molecule generation approach, which could be another promising direction to narrow down the modeling space.

Considering that a molecule must satisfy many properties, such as solubility and permeability, to become drug-like [Bickerton et al. 2012], another remaining challenge is to simultaneously optimize multiple drug-like properties of generated drug candidates, while retaining its binding affinity for the target protein. To our knowledge, existing works do not explicitly optimize such properties during generative modeling.

8.4 Molecule and Material Interactions

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Recommended Prerequisites: Section 5.2

This subsection describes research problems related to interactions between molecules and materials, including predicting the adsorption energy and per-atom force of molecule-material pair structures (S2E and S2F) and predicting the relaxed final structure and the adsorption energy at its relaxed state (IS2RS and IS2RE). In addition, we discuss the unexplored generative tasks for molecule-material interactions in Section 8.4.5.

8.4.1 Problem Setup.

Let the total number of atoms in a molecule-material pair be n , and the paired structure \mathcal{S} is represented as $\mathcal{S} = (z, C)$. Here, $z \in \mathbb{Z}^n$ is the atom type vector indicating the atom type (atomic number) of all n atoms in the structure. $C = [c_1, \dots, c_n] \in \mathbb{R}^{3 \times n}$ is the coordinate matrix where c_i denotes the 3D coordinate of the i -th atom in the structure. The first problem that has garnered the attention of the research community is predicting the adsorption energy of molecule-material pair structures (S2E). This task is to learn a function f_E to predict the property $e \in \mathbb{R}$ for any given pair structure \mathcal{S} , where e is a real number. The second problem of interest is predicting per-atom force given a structure's atomic types and positions as input (S2F). Here, the goal is to learn a function f_F to predict force matrix $F \in \mathbb{R}^{3 \times n}$ for any given structure \mathcal{S} . Per-atom forces drive structure relaxation until the pair structure reaches its relaxed state with an energy minimum. The third problem aims to learn a function f_{RE} to predict the structure's adsorption energy $e_{rel} \in \mathbb{R}$ at its relaxed state, given its initial structure \mathcal{S}_{init} as input (IS2RE). Typically, the initial structure \mathcal{S}_{init} is heuristically determined. Similar to IS2RE, the last problem IS2RS aims to learn a function f_{RS} to predict the relaxed final structure given its initial structure as input. In this problem, the target $C_{rel} \in \mathbb{R}^{3 \times n}$ represents atomic positions at the relaxed state of the given structure. The Open Catalyst 2020 (OC20) dataset [Chanussot* et al. 2021] provides absorbate-catalyst pair structures and serves as a testbed for the problems related to molecule-material interactions described above.

8.4.2 Technical Challenges.

For different problems defined above, there exist distinct challenges. First, for the energy prediction problem (S2E), the model prediction has to be rotationally invariant because energy is a structure-level property that is invariant to the rotation of molecule-material pairs. In contrast to energy, the force prediction problem (S2F) aims to predict per-atom force vectors. Hence, force prediction has to be equivariant to the rotation of the structure. Similarly, the relaxed energy prediction problem (IS2RE) and the relaxed structure prediction problem (IS2RS) have the same challenge of maintaining invariance and equivariance, respectively. In addition to symmetry, IS2RE and IS2RS problems have another challenge that the initial structure only provides a rough hint about the

Table 31. Comparison of existing methods for energy, force, and position prediction of molecule-material pairs. Different methods focus on different tasks and use different pipelines to solve the IS2RE and IS2RS tasks. Note that the methods introduced in Section 5.2 can also be applied to the tasks presented in this section. However, for the sake of brevity, this table only includes several state-of-the-art methods, including ForceNet [Hu et al. 2021b], GNS+NoisyNode [Godwin et al. 2022], Equiformer [Liao and Smidt 2023], EquiformerV2 [Liao et al. 2023], SpinConv [Shuaibi et al. 2021], GemNet-OC [Gasteiger et al. 2022], SCN [Zitnick et al. 2022], and eSCN [Passaro and Zitnick 2023].

Methods	Task	Pipeline for IS2RE and IS2RS tasks	Network	Symmetry
ForceNet	S2F/IS2RS	Relax	-	-
GNS+NoisyNode	IS2RE/IS2RS	Direct	-	-
Equiformer	IS2RE	Direct	$\ell = 1$	$SE(3)/E(3)$ -Equivariant
EquiformerV2	S2EF/IS2RE/IS2RS	Relax	$\ell > 1$	$SE(3)/E(3)$ -Equivariant
SpinConv	S2EF/IS2RE/IS2RS	Direct/Relax	$\ell = 0$	Invariant
GemNet-OC	S2EF/IS2RE/IS2RS	Relax	$\ell = 0$	Invariant
SCN	S2EF/IS2RE/IS2RS	Direct/Relax	$\ell > 1$	Approximately equivariant
eSCN	S2EF/IS2RE/IS2RS	Relax	$\ell > 1$	Approximately equivariant

relaxed structure. Therefore, the model must consider structure relaxation, including molecule and material atoms, to obtain accurate predictions.

8.4.3 Existing Methods.

Table 31 provides a summary of existing methods, including their symmetry type, network order, tasks, and the pipeline used in the original paper. As discussed in Section 5.2, both invariant and equivariant methods can predict the $SE(3)$ -invariant energy e . Once the energy is predicted, the $SO(3)$ -equivariant force vectors F acting on atoms can be obtained using the formula $F_i = -\frac{\partial e}{\partial c_i}$. This formula is the only option to compute forces for invariant models like SpinConv [Shuaibi et al. 2021] and GemNet-OC [Gasteiger et al. 2022]. However, equivariant methods can also produce forces directly. As discussed in Section 5.2, one main challenge for invariant methods is efficiency. Existing methods suffer from high computational cost [Gasteiger et al. 2020, 2021] when incorporating more geometric information, like angles and torsion angles, into the network. On the other hand, one main challenge for equivariant methods is that, explicitly imposing physical constraints into the model architecture limits the capacity of the network [Schütt et al. 2021]. Consequently, recent studies try to design GNN models that are both efficient and expressive without explicit physical constraints. For example, ForceNet [Hu et al. 2021b] directly uses atom coordinates in a scalable manner, but implicitly imposes physical constraints by using data augmentation.

To address the challenge of IS2RE and IS2RS tasks, where only initial structures are given, methods that consider structure relaxation are needed. There are primarily two solutions. The first is to use a well-trained S2F model to iteratively update the structure from the initial one. Atom positions are updated step-by-step based on the predicted forces of the current structure until the predicted forces approach zero. While this method can accurately simulate structure relaxation, it requires numerous steps to achieve the final output. Recently, both SCN [Zitnick et al. 2022] and eSCN [Passaro and Zitnick 2023] use this indirect approach and note that they are approximately equivariant but with high order $\ell > 1$. Incorporating both eSCN and a recent direct method Equiformer [Liao and Smidt 2023], EquiformerV2 [Liao et al. 2023] achieves state-of-the-art performance among these indirect methods. It applies eSCN convolution layers to Equiformer network structure with several additional techniques, including separable \mathbb{S}^2 activation and layer normalization for vectors of $\ell = 0$ and those of $\ell > 0$. In contrast to indirect methods, the direct approach aims to model the relations between initial and relaxed structures, with the output typically being the difference

$C_{rel} - C_{init}$ [Godwin et al. 2022]. Hence, direct methods are faster in training and prediction but less accurate than indirect methods. Equiformer [Liao and Smidt 2023], an attention-incorporated equivariant architecture with $\ell = 1$, achieves state-of-the-art performance among direct methods. Currently, the NoisyNode technique proposed in [Godwin et al. 2022] is widely used in many other direct approaches. However, it's worth noting that both GNS+NoisyNode [Godwin et al. 2022] and ForceNet [Hu et al. 2021b] are based on the GNS framework, which is neither invariant nor equivariant. These methods take atom coordinates as input and use rotation data augmentation to improve performance.

8.4.4 Datasets and Benchmarks.

Open Catalyst 2020 (OC20) dataset [Chanussot* et al. 2021] is a valuable resource for testing machine learning models on the problems related to molecule-material interactions described in this subsection. This dataset provides adsorbate (molecule)-catalyst (material) pair structures and includes three tasks, namely Structure to Energy and Forces (S2EF), Initial Structure to Relaxed Energy (IS2RE), and Initial Structure to Relaxed Structure (IS2RS), which correspond to the tasks introduced in Section 8.4.1.

IS2RE and IS2RS tasks use the same dataset which contains the input initial structures and the output relaxed structures and energies. The dataset is originally split into training, validation, and test sets. The training set contains 460,328 structures. For both validation and testing sets, there are four subsets, namely in-domain (ID), out-of-domain adsorbate (OOD Ads), out-of-domain catalyst (OOD Cat), and out-of-domain adsorbate and catalyst (OOD Both). The ID set consists of structures from the same distribution as training. The OOD Ads set consists of structures with unseen adsorbates, the OOD Cat set consists of structures with unseen element compositions for catalysts, and the OOD Both set consists of structures with unseen catalysts and adsorbates. The four validation sets contain 24,943, 24,961, 24,963, and 24,987 structures, respectively. The testing sets contain 24,948, 24,930, 24,965, and 24,985 structures, respectively. Note that the labels for the test sets are not publicly available. Therefore, researchers need to submit their results to the Open Catalyst Project (OCP) leaderboard to evaluate their models on the test sets.

The dataset for S2EF contains more structures compared to the other two tasks. This is because the dataset for S2EF contains not only the initial and relaxed structures but also the intermediate structures in the relaxation trajectory, etc. The dataset comprises 133,934,018 structures for training, while for the ID, OOD Ads, OOD Cat, and OOD Both validation sets, it contains 999,866, 999,838, 999,809, and 999,944 structures, respectively. Similarly, the ID and OOD testing sets include 999,736, 999,859, 999,826, and 999,973 structures, respectively.

In addition to OC20 [Chanussot* et al. 2021], OC22 [Tran et al. 2023] is curated recently and focuses on oxide electrocatalysts. Similarly, it contains three tasks, namely Structure to Total Energy and Forces (S2EF-Total), Initial Structure to Total Relaxed Energy (IS2RE-Total), and Initial Structure to Relaxed Structure (IS2RS). One main difference between OC20 and OC22 is that OC22 employs total energy targets rather than adsorption energy targets. In general, this makes the models more general and enables the calculation of more properties, but it is more challenging.

8.4.5 Open Research Directions.

Though significant progress has been made over the last several years, there are still remaining challenges in modeling molecule-material interactions. Firstly, the relaxed energy/structure in the IS2RE/IS2RS challenge does not necessarily correspond to the global minimum adsorption energy for a molecule-material pair, and is sensitive to changes in the initial structure. Recently, [Lan* et al. 2022] used a brute force strategy to calculate relaxed energies across many initial configurations. In order to efficiently predict global minima, it will be necessary to use more advanced strategies for

sampling initial configurations or build models that do not depend on specified initial configurations. Secondly, although many materials in molecule-material interactions are periodic, to our knowledge, there do not exist any models which explicitly model such periodicity when modeling a molecule-material pair. Instead, models only consider in the material the unit cell closest to the small molecule. Finally, models are currently incapable of incorporating all physical properties of the system. Many models have seen improved performance by including molecular dynamics information, however, current works cannot model properties such as magnetic or charge effects which can significantly impact the relaxed energy/structure.

Generation of Molecule-Material Pairs: One promising direction for generative tasks in this field is to generate periodic materials conditioned on given molecules. This task involves training a generative model to produce new periodic material structures with specific properties, such as appropriate adsorption energies, when a specific molecule is present. To accomplish this, the generative model would need to be trained on a dataset of periodic material structures with different adsorbate molecules and their corresponding adsorption energies. Such a generative task has many potential applications, including designing new materials for electrocatalysts.

Although recent studies have explored unconditional material generation, as described in Section 7.3, developing a generative model that accurately captures the complex interactions between adsorbate molecules and periodic materials is challenging. It requires a deep understanding of the underlying physics and chemistry. Moreover, similar to the structure-based drug design task introduced in Section 8.3, the vast data space that needs to be modeled makes this task even more challenging. Additionally, this task presents a unique challenge in that the 3D geometry of the molecule is not static and will be influenced by the generated material molecule. Therefore, when generating periodic materials conditioned on given molecules, the generative model needs to account for the interplay between the molecule and the material, including the induced changes in the molecular geometry.

9 AI FOR PARTIAL DIFFERENTIAL EQUATIONS

In this section, we detail advances in the field of AI for solving Partial Differential Equations (PDEs). We overview the general formulation of PDE modeling and motivate machine learning methods in this context in Section 9.1. We discuss forward problems in Section 9.2 and inverse tasks in Section 9.3.

9.1 Overview

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A PDE mathematically describes the behavior of a system by an unknown multivariate function and its partial derivatives. PDEs are frequently applied in a variety of disciplines to model the space-time evolution of physical processes, such as airflow around an airfoil with the Navier-Stokes equations [Li et al. 2022b; Bonnet et al. 2022], global weather patterns with the shallow water equations [Gupta and Brandstetter 2022], or optical design with Maxwell's equations [Brandstetter et al. 2023]. Additional real-world applications of PDE modeling include weather forecasting [Pathak et al. 2022], carbon dioxide storage [Wen et al. 2022, 2023], seismic wave propagation [Yang et al. 2021a; Sun et al. 2022; Yang et al. 2023a; Sun et al. 2023], material sciences, and volcanic activities [Rahman et al. 2022a]. In many real-world applications, it may be intractable to obtain the functional form of the PDE solution, and therefore, the PDE must be solved numerically. Due to the widespread application of PDE modeling, over a century of work has been dedicated to developing numerical PDE solvers [Brandstetter et al. 2022c].

Classical solvers rely on discrete numerical approximations of derivatives [Quarteroni and Valli 2008; Bartels 2016], such as forward difference approximations of the form

$$\partial_x u(x, t) \approx \frac{u(x + \Delta x, t) - u(x, t)}{\Delta x}.$$

These approximations improve in accuracy as the step size Δx decreases to 0. This reduces the PDE to a set of (possibly non-linear) equations, which may be solved *explicitly* – where the solution at time t is determined using solutions at a previous time $t' < t$ – or *implicitly* – where the solutions are solved for all times at once. Explicit schemes are simpler to implement than implicit schemes, yet usually require smaller step sizes to achieve similar accuracy or even to converge for stiff problems which may exhibit sharp discontinuities [Courant et al. 1928].

While classical approaches such as the Finite Element Method (FEM) and the Finite Difference Method (FDM) Quarteroni and Valli [2008]; Bartels [2016] have been proven to be effective, they require high computational effort. Furthermore, they often need to be carefully tailored on a task-by-task basis to ensure numerical stability due to variability in the behavior of the PDE solutions dependent on the governing equations and setting. Large systems that are prevalent in industry applications of PDE modeling can require extensive computational resources and hundreds or even thousands of CPU hours [Lam et al. 2022].

To address these shortcomings, deep learning models have emerged as a general framework to produce solutions orders of magnitude faster than their numerical counterparts. This efficiency is primarily achieved via the ability of neural networks to take significantly larger time steps during inference, learn on more coarse spatial discretizations compared to classical solvers [Pfaff et al. 2021; Stachenfeld et al. 2021], and use explicit forward methods instead of implicit methods [Tang et al. 2020; Wu et al. 2022c].

Unlike time-consuming iterative methods for implicit schemes, neural solvers learn a direct mapping from past states to future states, with fewer restrictions on the resolution of the data [Kochkov et al. 2021b]. Additionally, neural solvers can easily be optimized and evaluated using parallelized

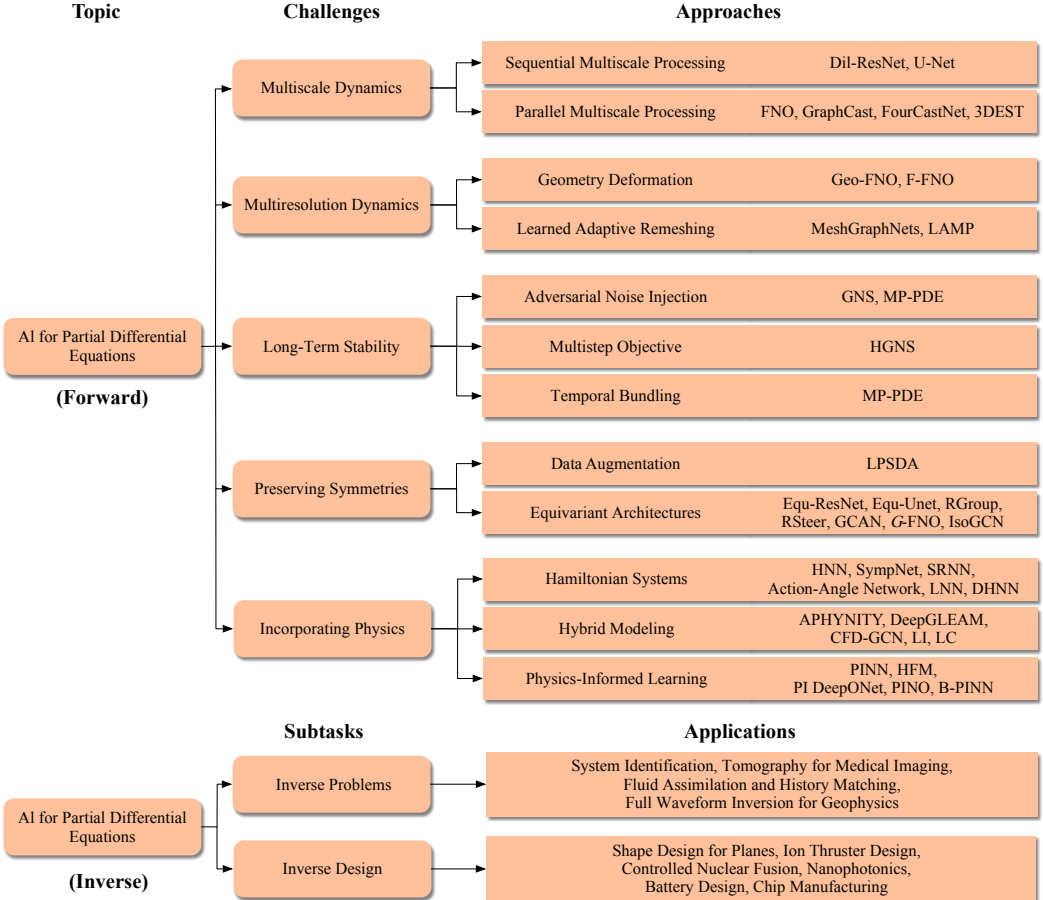


Fig. 32. Overview of AI for forward modeling and inverse modeling of partial differential equations (PDEs). In Section 9.2, we consider the forward modeling task, that is, mapping from the initial timesteps of the numerical solution of a PDE to later timesteps. We identify and detail four fundamental challenges that have defined the development of neural PDE solvers. Multiscale dynamics arise in systems where physics evolve on a continuum from local to global scales [Stachenfeld et al. 2021; Gupta and Brandstetter 2022; Li et al. 2021b; Liu et al. 2022b; Lam et al. 2022; Pathak et al. 2022; Bi et al. 2022; Rui Wang 2020; Li et al. 2021c], while multiresolution dynamics occur in systems with fast-evolving, isolated regions that require greater resources for stable simulation [Li et al. 2022b; Tran et al. 2021; Pfaff et al. 2021; Wu et al. 2022a]. Solvers that use explicit schemes, where the state of the system at the next time step is computed from the current time step, encounter error in their inputs introduced by previous predictions and thus must consider methods for maintaining rollout stability [Sanchez-Gonzalez et al. 2020; Brandstetter et al. 2022c; Wu et al. 2022c]. Equivariant architectures and training techniques enforce symmetries of the system, enabling improved generalization and sample complexity [Brandstetter et al. 2022b; Wang et al. 2021a, 2022i; Ruhe et al. 2023; Horie et al. 2021; Helwig et al. 2023]. Lastly, incorporating physics into architectures enables predictions to maintain physical consistency and reduces the difficulty of the learning task where models would otherwise be tasked with extracting physical laws from the training data [Greydanus et al. 2019; Jin et al. 2020; Chen et al. 2020b; Daigavane et al. 2022; Cranmer et al. 2020; Sosanya and Greydanus 2022; Yin et al. 2021; Wu et al. 2021; Belbute-Peres et al. 2020; Kochkov et al. 2021b; Tompson et al. 2017; Raissi et al. 2019, 2020; Wang et al. 2021b; Li et al. 2021g; Yang et al. 2021b]. In Section 9.3, we consider the inverse modeling task, including inverse problems and inverse design. Specifically, the task considered by inverse problems is to infer the unknown parameters of the system given observed dynamics, while the task for inverse design is to optimize the system based on a predefined objective. These two subtasks of inverse modeling have various applications across the science and engineering fields.

GPU operations, while the design of GPU-compatible classical solvers may offer limited benefit due to their iterative nature, and furthermore requires intimate familiarity with complex numerical methods. Most importantly, these models have the ability to adapt to the task at hand and can be trained to generalize across initial conditions [Li et al. 2021b; Gupta and Brandstetter 2022] and PDE parameters [Brandstetter et al. 2022c; Tran et al. 2021]. Further, unlike classical solvers, neural solvers can learn dynamics directly from observed data, an ability that is especially useful when the underlying equations are unknown [Lienen and Günemann 2022].

Motivated by their prominence in industrial applications of PDE modeling and their challenging nature, many works focus on designing neural solvers for the class of time-evolving PDEs. Formally, a time-evolving PDE is a system of equations relating the derivatives of an unknown function $u : U \rightarrow \mathbb{R}^m$ of space and time [Olver 2014], where $U = \mathbb{X} \times \mathbb{T}$ consists of the spatial domain \mathbb{X} and the temporal domain \mathbb{T} . Given U , we consider time-evolving PDEs given by a set of equations [Brunton and Kutz 2023; Evans 2022]

$$\begin{aligned} \partial_t u + \mathcal{D}(x, t, u, \partial_x u, \partial_{xx} u, \dots) &= \mathbf{0} & (x, t) \in U, \\ u(x, 0) &= u_0(x) & x \in \mathbb{X}, \\ \mathcal{B}u(x, t) &= \mathbf{0} & (x, t) \in \partial\mathbb{X} \times \mathbb{T}, \end{aligned} \quad (104)$$

where \mathcal{D} is an operator relating the partial derivatives of the solution u on the space-time domain U , \mathcal{B} is a differential operator relating derivatives on the boundary of the spatial domain $\partial\mathbb{X}$, and u_0 is the initial condition describing u at time $t = 0$. To solve this PDE, we must identify a function $u(x, t; \gamma)$ satisfying the constraints in Equation (104), either in analytical or numerical form, where $\gamma = (u_0, \mathcal{B}, \gamma_P)$ denotes the PDE configuration describing the initial condition u_0 , boundary condition \mathcal{B} , and PDE parameters γ_P . There are several deep learning frameworks that exist for approximating this solution. Approaches for the forward problem, discussed in Section 9.2, utilize a forecasting model as a learned solver to map past numerical solutions to future solutions. Alternatively, inverse problems and inverse design, detailed in Section 9.3, consider the reverse direction, where the task is instead to map from observed solution data to the initial conditions, boundary conditions, or PDE parameters of the underlying equations, or to optimize the design of a system based on some criterion.

9.2 Forward Modeling

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In this section, we overview the progress of machine learning models developed for forward PDE problems. We formalize the forward task for neural PDE solvers in Section 9.2.1 before outlining the primary challenges that have shaped their development in Section 9.2.2. In Sections 9.2.3 to 9.2.7, we discuss models and techniques that have emerged in response to these challenges, as well as datasets and benchmarks for these models in Section 9.2.8, before closing with a discussion of remaining challenges and future directions in Section 9.2.9.

9.2.1 Problem Setup.

In forward problems, models are tasked with predicting future states of the system given initial conditions or historical observations as inputs [Kovachki et al. 2021; Li et al. 2021g,c; Brandstetter et al. 2022c; Gupta and Brandstetter 2022; Li et al. 2021b; Sanchez-Gonzalez et al. 2020; Stachenfeld et al. 2021; Wen et al. 2023; Yang et al. 2023a]. Models are trained on a set of numerical solutions $\{u^{(j)}\}_{j=1}^n$, where $u^{(j)}(x, t) := u(x, t; \gamma^{(j)})$ and the PDE configurations $\gamma^{(j)}$ vary depending on the

setting. For example, the $\gamma^{(j)}$ may correspond to varying initial conditions [Li et al. 2021b; Rahman et al. 2022b; Gupta and Brandstetter 2022] or PDE parameters [Li et al. 2020b; Yang et al. 2021a; Brandstetter et al. 2022c; Tran et al. 2021].

Numerical PDE solutions discretize the solution domain U into a finite set of collocation points on which the solver will approximate the value of the solution function. For a PDE solution u , denote u_t as the t -th time step in a uniform discretization of the temporal domain \mathbb{T} with step size Δ_T , that is, $u_t(x) := u(x, t\Delta_T)$, where x is any point in the discretization of the spatial domain \mathbb{X} . Additionally, let the PDE solutions at consecutive time points be denoted as $u_{k:(k+K)} := \{u_k, u_{k+1}, \dots, u_{k+K}\}$. The dynamics forecasting task is then defined as:

$$\phi_\theta(u_{0:(k-1)}^{(j)}) = \widehat{u_{k:T}^{(j)}}, \quad (105)$$

where ϕ_θ is optimized based on a suitably chosen loss \mathcal{L} as

$$\phi_\theta = \arg \min_{\phi_\theta: \theta \in \Theta} \mathbb{E}_{u^{(j)}} \left[\mathcal{L} \left(\phi_\theta \left(u_{0:(k-1)}^{(j)} \right), u_{k:T}^{(j)} \right) \right]. \quad (106)$$

In many cases, the form of the PDE gives rise to a solution set closed to the action of a symmetry group G . That is, if u is a function that satisfies Equation (104), then for all group elements $g \in G$, $L_g u$ also satisfies Equation (104), where for a function f , $L_g f(x) := f(g^{-1}x)$ denotes f transformed by g . In such a case, it is desirable to constrain the model search space such that these symmetries are automatically respected, that is $\phi_\theta(L_g u_0) = L_g \phi_\theta(u_0)$. Such constraints have been shown to improve generalization and sample complexity for learned solvers via explicit encoding in equivariant architectures such as equivariant CNNs [Wang et al. 2021a; Helwig et al. 2023] and equivariant GCNs [Horie et al. 2021], and through data augmentation [Brandstetter et al. 2022b]. We discuss this further in Section 9.2.6.

9.2.2 Technical Challenges.

We next identify five key challenges encountered by neural solvers in the forward modeling setting, each of which have given rise to a variety of solutions that have shaped machine learning research in the field of PDE modeling.

Multiscale Dynamics (Section 9.2.3): As physics evolve on multiple spatial scales, capturing the interactions within and between dynamics on each scale is vital in producing high-quality numerical solutions to PDEs. However, it is challenging to do this effectively, particularly at global scales, without excessively trading off computational efficiency or sacrificing performance at local scales.

Multiresolution Dynamics (Section 9.2.4): Many systems possess isolated regions of fast-evolving dynamics that require higher resolution discretizations relative to others to maintain solver stability. Therefore, the ability to model irregular geometries balances a trade-off between simulation accuracy and computational effort by enabling resources to be allocated dynamically in space and time.

Long-Term Stability (Section 9.2.5): Evolving a system for many time steps can lead to an accumulation of error that causes the predicted rollout to diverge from the ground truth. As this error accumulation does not naturally occur until test time, it can be difficult to condition models to be stable during inference.

Preserving Symmetries (Section 9.2.6): Many PDEs have intrinsic symmetries which are often used to find reduced-order models and improve solution efficiency. For machine learning, symmetry may be used as an inductive bias to reduce the difficulty of the learning task and narrow the size of the model search space. Furthermore, Noether's theorem [Halder et al. 2018; Noether 1971]

establishes a connection between symmetries and conservation laws, which implies that models that uphold symmetries can produce physically consistent predictions.

Incorporating Physics (Section 9.2.7): Since machine learning models are fundamentally statistical, they are prone to make scientifically implausible predictions when trained solely on data without explicit constraints. Thus, leveraging known physical principles to guide deep learning models is crucial for learning the correct underlying dynamics instead of simply fitting the observed data, which may contain spurious, non-physical trends. This can be accomplished by imposing constraints on the loss function and the design of the architecture, or by appropriately augmenting traditional physics-based models with neural nets.

In the sections that follow, we discuss the above challenges in greater detail and advances made by previous works to address them, outlined in Figure 32.

9.2.3 Existing Methods: Multiscale Dynamics.

Dynamics evolve and interact at multiple scales in many physical systems. For example, turbulent flows exhibit a hierarchy of localized regions of turbulent motion of different sizes, known as *eddies*, in which energy from one scale is dispersed to eddies at the next smallest scale [Pope 2000]. While the behavior of particles is often most strongly associated with particles in the immediate neighborhood, architectures composed of layers that only consider local information such as ResNets [He et al. 2016] rely on stacks of many layers to propagate long-range signals, and therefore demonstrate inferior performance in evolving dynamics [Li et al. 2021b; Gupta and Brandstetter 2022; Ruhe et al. 2023]. Thus, a primary factor for faithful and efficient simulation via machine learning is the incorporation of multiscale processing mechanisms [Li et al. 2020b; Gupta and Brandstetter 2022; Rahman et al. 2022b; Wen et al. 2023] that balance the complexity tradeoff while maintaining sufficient local information flow.

Stachenfeld et al. [2021] implement this mechanism in their Dil-ResNet using blocks of convolution layers with sequentially increasing dilation rates to process information beginning from local up to global scales followed by sequentially decreasing dilation rates to process from global to local. Following a similar philosophy, Gupta and Brandstetter [2022] study several variants of the U-Net architecture [Ronneberger et al. 2015], which uses downsampling and upsampling in place of dilation to traverse between local and global scales. In both cases, the mechanism processes local and global information sequentially while managing complexity by increasing the receptive field with a fixed kernel size, which we visualize in Figure 33.

In contrast to sequential processing mechanisms, the Fourier Neural Operator (FNO) proposed by Li et al. [2021b] processes multiscale information in parallel [Gupta and Brandstetter 2022]. Neural operators are maps between function spaces [Kovachki et al. 2021] and de facto models for scientific computing and physics phenomena dealing with PDEs. Unlike the previously discussed Dil-ResNet and U-Net that represent operators parameterized in the physical domain, a subset of neural operators are designed to be theoretically discretization invariant [Kovachki et al. 2021]. Furthermore, the universal approximation theorem for operators theoretically guarantees the ability of neural operators to approximate any continuous operator between Banach spaces within arbitrary precision [Kovachki et al. 2021]. Among neural operator architectures, FNO deploys Fourier bases for internal integrations and is applicable to arbitrary function spaces with complex domains [Li et al. 2022b]. When dealing with regular grids and domains, the Fourier bases integration is often carried out using the Fast Fourier transform (FFT), making FNO among the most computationally efficient neural operator models. FNO has been successfully applied to many large-scale applications, including weather forecasting [Pathak et al. 2022] and climate mitigation acts [Wen et al. 2023]. In Fourier space, low frequency modes represent information on a global scale, with higher frequency modes containing local information, and thus, multiscale processing in the frequency domain occurs

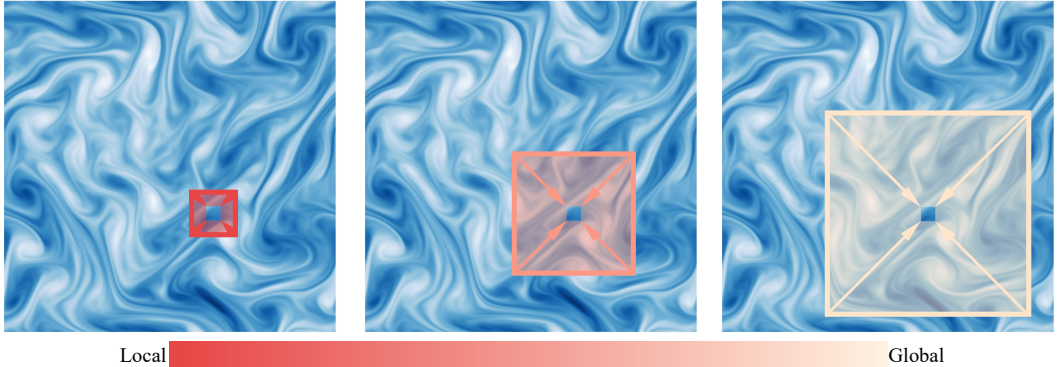


Fig. 33. Multiscale dynamics. Many systems exhibit dynamics consisting of interacting components with sizes ranging from local to global scales. A primary example is turbulent flows, which possess a hierarchy of eddies that decay down to the smallest scale, referred to as the Kolmogorov scale [Pope 2000]. Constructing machine learning models with multiscale processing mechanisms is therefore key for high-fidelity simulation. These mechanisms aggregate information on each scale to update the latent representation at each mesh point. Here, we visualize a mechanism that performs aggregation and updates on each scale sequentially, as considered by Stachenfeld et al. [2021] and Gupta and Brandstetter [2022], however, mechanisms such as those proposed by Li et al. [2021b] and Lam et al. [2022] act in parallel.

in parallel through the point-wise multiplication inherent to Fourier convolutions. To manage complexity, frequency modes above a fixed cutoff are set to zero, reducing the number of operations and parameters.

Guibas et al. [2022] extend FNOs into the vision transformer framework, which was later applied by Pathak et al. [2022] for the task of forecasting global weather. To efficiently leverage attention as a multiscale processing mechanism, Guibas et al. [2022] use the Fourier transform as an inexpensive token mixer. Bi et al. [2022] also build on vision transformers for weather forecasting, instead using patch embedding in their 3D Earth Specific Transformer (3DEST) to manage the quadratic complexity incurred by attention. Lam et al. [2022] propose GraphCast in the same setting, a GNN operating on a graph representing the state of the global weather, with the edge set containing seven different lengths of edges for efficiently passing messages long-range, ranging from a few edges spanning long distances to hundreds of thousands of localized short edges. As weather phenomena range from localized blizzards to heatwaves spanning multiple continents [Gupta and Brandstetter 2022], and the training data spans nearly a half-century [Hersbach et al. 2020], efficient multiscale processing is particularly important for the task considered by these works. In large part due to effective choice of this processing mechanism, each of these models outperform the numerical weather prediction model currently used for delivering real-world forecasts on various tasks while operating at a fraction of the cost [Bi et al. 2022; Lam et al. 2022; Pathak et al. 2022].

9.2.4 Existing Methods: Multiresolution Dynamics.

The ability to model non-uniform discretizations of the PDE domain is important for balancing the tradeoff between computational cost and solution accuracy. Classical numerical solvers rely on the assumption that the solution is sufficiently smooth between collocation points to maintain stability [Kochkov et al. 2021b]. However, phenomena such as shockwaves and solid objects impeding flows, as we visualize in Figure 34, introduce local regions of steep gradient that require expensive high-resolution discretizations to maintain this smoothness [Berger and Oliger 1984]. Uniform discretizations wastefully allocate the same high resolution required in these isolated

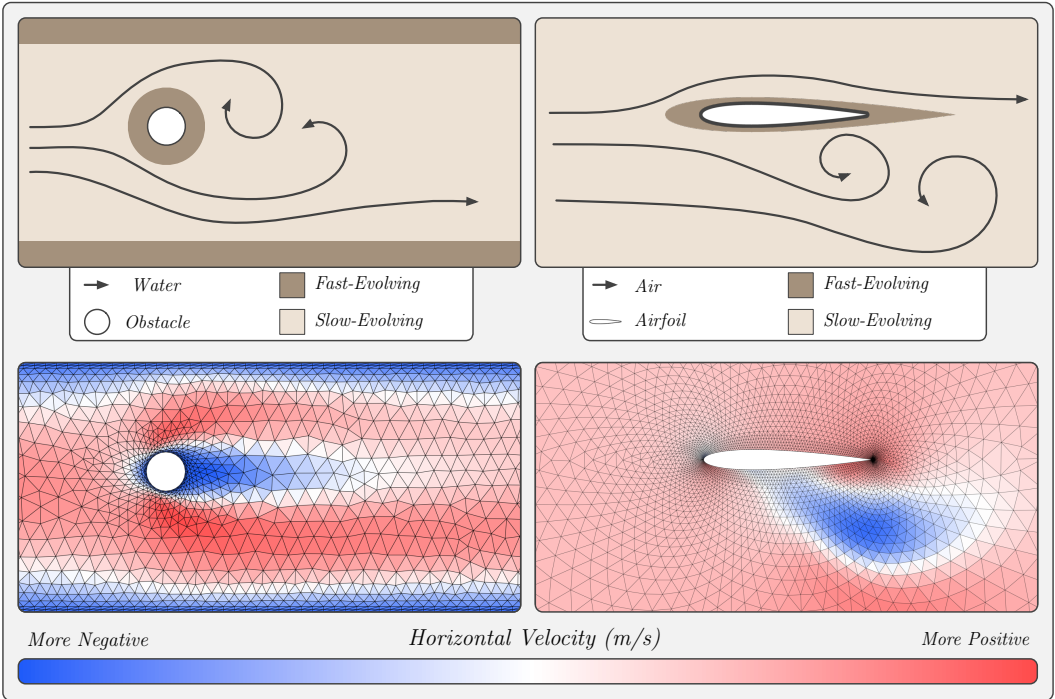


Fig. 34. Multiresolution dynamics, data from Pfaff et al. [2021]. Systems with localized regions of fast-evolving dynamics, such as fluid flow around a cylinder (left) or air flow around an airfoil (right), require high-resolution discretizations in these regions for dynamics to be stably resolved. Irregular discretizations of the domain can manage this cost by allocating high resolution in regions of high gradient and coarse resolution elsewhere. However, since irregularly discretized functions cannot be modeled by architectures such as CNNs, there has been a call for GNNs for simulating dynamics [Pfaff et al. 2021]. Furthermore, the location in space of these high gradient regions can shift as the system evolves, thus requiring the discretization to dynamically adapt. While traditional re-meshing algorithms can be expensive, Pfaff et al. [2021] and Wu et al. [2022a] propose learned alternatives for adaptive mesh refinement that reduce this cost.

regions even in areas where the dynamics are slower-evolving [Wu et al. 2022a]. To address this limitation, non-uniform meshes allocate fine resolution in high-gradient regions and coarse resolution elsewhere. Furthermore, the geometry of the mesh can adapt as high-gradient regions shift in space, as is commonly the case for time-evolving PDEs [Berger and Oliger 1984].

While machine learning methods have been shown to allow for coarser discretizations than numerical methods due to their ability to learn a direct mapping [Kochkov et al. 2021b; Stachenfeld et al. 2021], neural networks still benefit from a certain level of continuity in the solution space. In addition to the inefficiency for dynamics with isolated regions of high gradient, surrogate models built on the CNN architecture, such as those explored by Stachenfeld et al. [2021] and Gupta and Brandstetter [2022], cannot directly model non-rectangular domains, for example, fluid flow around a cylinder. These limitations have given rise to a number of approaches for modeling dynamics on non-uniform meshes and learned mesh adaptation.

Similar to classical CNNs, the use of the discrete Fourier transform (DFT) by the FNO [Li et al. 2021b] restricts the model to rectangular domains with regularly-spaced grid points. Li et al. [2022b] therefore propose the Geo-FNO, a framework later extended by Tran et al. [2021] with the F-FNO.

These architectures first deform the input mesh to a rectangular, uniform grid in latent space for Fourier convolutions with the FFT and finally apply the reverse deformation to produce the output on the irregular grid. However, this deformation-based approach comprises discretization invariance properties [Kovachki et al. 2021]. Furthermore, preserving the geometry of the domain is a primary factor enabling effective generalization and learning in convolutional architectures [Cohen et al. 2018].

GNNs present a geometry-preserving mechanism for learning functions on irregular meshes [Kipf and Welling 2017; Gilmer et al. 2017] that treat the input mesh as a graph on which the PDE solution is predicted at each node. Graph neural operators (GNO) [Li et al. 2020a], the neural operator generalization of GNNs, offer a theoretically discretization-invariant approach in this setting. Pfaff et al. [2021] take a similar approach with their MeshGraphNets framework, which additionally involves a second GNN that predicts the *sizing tensor* for each node. The predicted sizing tensor is then used by an adaptive mesh refinement algorithm to update mesh geometry as dynamics evolve. Note that updating the mesh geometry to allocate greater resources in higher gradient regions is distinct from the approach of Li et al. [2022b], which deforms the representation of the domain in latent space so that the collocation points in the mesh form a rectangular, uniform grid. However, this re-meshing algorithm is expensive and furthermore may not produce the optimal geometry in terms of the tradeoff between solution accuracy and cost of the mesh [Wu et al. 2022a]. Thus, Wu et al. [2022a] propose Learning controllable Adaptive simulation for Multi-resolution Physics (LAMP), a faster, data-driven approach to re-meshing using reinforcement learning. They jointly optimize the dynamics GNN and a mesh refinement policy, where the policy is selected by simultaneously minimizing the error of the dynamics GNN and the number of nodes in the mesh.

9.2.5 Existing Methods: Long-Term Stability.

Time-evolving PDEs are numerically solved by discretizing the temporal domain into time steps on which the solver produces solutions. This solution can be obtained using an explicit scheme, wherein the solution at a given time point is directly calculated using the preceding solution as $u_{t+1} = F(u_t)$ for some function F , or implicit schemes, which entail solving a system of (possibly non-linear) equations involving u_t and u_{t+1} [Olver 2014]. Although explicit schemes appear to require less computational effort than implicit, classical solvers that advance time using an explicit scheme can exhibit conditional stability, meaning that the discretization in time must be chosen sufficiently fine to prevent the solver from diverging [Courant et al. 1928; Olver 2014]. Such PDEs for which explicit methods require significantly finer time discretizations compared to the smoothness of the actual solution are termed *stiff*. As a result, implicit methods have been traditionally preferred for solving stiff PDEs. Nonetheless, many neural surrogates utilize explicit schemes for convenience, and have been shown to outperform classical solvers on computationally inexpensive coarse discretizations [Kochkov et al. 2021b; Stachenfeld et al. 2021]. However, explicit schemes inevitably introduce error to the inputs of the model, and thus, increasing the robustness of neural solvers to noisy inputs is key for enabling stable predictions over many time steps.

The task often considered in this setting is to predict the rollout up to time step T conditioned on the first k solutions, that is, the mapping $(u_0, u_1, \dots, u_{k-1}) \mapsto (u_k, u_{k+1}, \dots, u_T)$. In what follows, we take $k = 1$ for simplicity in notation such that the mapping to be learned is from the time 0 solution to the remaining $T - 1$ steps, but this is not necessary in general. For such a task, explicit schemes train ϕ_θ for a one-step prediction of u_{t+1} conditioned on the ground-truth solution at u_t , and at test time predict the full rollout by applying the trained network autoregressively T times [Brandstetter et al. 2022c; Sanchez-Gonzalez et al. 2020; Stachenfeld et al. 2021]. This one-step training strategy has been shown to be more effective than training recurrently to predict the full rollout u_1, u_2, \dots, u_T [Tran et al. 2021]. However, it is not representative of the task at test time,

since for $t > 1$, the input to the model will not be the ground truth u_t as in training, but rather $u_t + \varepsilon_t$, where ε_t is the error accumulated up to time t through autoregressive prediction of u_t .

Several works have addressed this using various techniques, including the Sobolev loss and physics properties such as dissipativity [Li et al. 2021c], the Lyapunov regularizer [Zheng et al. 2022] and adversarial noise injection [Sanchez-Gonzalez et al. 2020; Brandstetter et al. 2022c]. Noise injection approaches intentionally corrupt inputs during training with an approximation to ε_t . Sanchez-Gonzalez et al. [2020] apply this strategy in training their Graph Network-based Simulator (GNS) by assuming that ε_t follows a 0-mean Gaussian distribution with variance chosen as a hyperparameter. Although this approach is convenient since the noise distribution can be easily sampled, the normality assumption may not be valid, and furthermore, the variance hyperparameter must be carefully tuned. Brandstetter et al. [2022c] instead obtain the noise to train their Message Passing PDE solver (MP-PDE) directly from the model to reduce the distributional shift between the training noise and the test noise. They accomplish this by letting $\varepsilon_t = \phi_\theta(u_{t-1}) - u_t$ such that the input to the model is $u_t + \varepsilon_t = \phi_\theta(u_{t-1})$. Thus, the noise added to the inputs during training is directly sourced from the model, as it will be during testing.

Although this approach eliminates the distributional shift for rollouts of only 2 timesteps, the shift appears again for longer rollouts. Wu et al. [2022c] address this by optimizing their Hybrid Graph Network Simulator (HGNS) using a multi-step objective in recurrently predicting several steps ahead. The total loss is a weighted sum of the loss at each time step, with the one-step loss weighted heaviest so that the optimization initially targets short-term predictions before fine-tuning for longer-term predictions. This method is more stable than full-blown recurrent prediction since it only predicts several steps ahead, and not the full rollout.

While the previously discussed architectures predict only one step ahead, Brandstetter et al. [2022c] make the observation that since each forward propagation introduces some error, reducing the number calls to the model required to predict a rollout could reduce the total accumulated error. Instead of predicting only one time step ahead as $\phi_\theta(u_t) = \hat{u}_{t+1}$, they train their model to predict l steps ahead with one forward propagation as $\phi_\theta(u_t) = (\hat{u}_{t+1}, \hat{u}_{t+2}, \dots, \hat{u}_{t+l})$. For example, to predict 10 timesteps with $l = 2$, only 5 forward propagations are required instead of 10.

9.2.6 Existing Methods: Preserving Symmetries.

Dynamic systems are governed by the laws of physics, with symmetries of systems related to these laws through Noether's theorem [Noether 1971; Wang et al. 2021a]. The symmetry group of a PDE characterizes the transformations under which solutions remain solutions, e.g., for a PDE with rotation symmetry, rotating the solution function produces a function that is also a solution. Symmetries such as rotation invariance are understood intuitively as the lack of canonical reference frame that allows, for example, a 2-dimensional flow rotated by 90° to remain equally physically plausible. Other symmetries such as translation invariance arise in PDEs with infinite domains or periodic boundaries [Holmes et al. 2012]. Priors that enforce symmetries can improve generalization and sample complexity by reducing the size of the solution space [Raissi et al. 2019; Wang et al. 2021a; Brandstetter et al. 2022b].

As a method of instilling learned equivariance, Brandstetter et al. [2022b] propose to use data augmentation according to the symmetry group of the PDE. However, while data augmentation can be used to achieve an approximately equivariant architecture, the internal representations in general will not be equivariant [Worrall et al. 2017], which is a major factor contributing to effective learning in equivariant architectures. Furthermore, many systems exhibit dynamics with local symmetries that data augmentation cannot capture [Worrall et al. 2017].

Equivariant CNNs, which are composed of convolutional layers that automatically encode the desired symmetry [Cohen and Welling 2016, 2017b; Weiler et al. 2018a; Weiler and Cesa 2019a;

[Worrall and Welling 2019], address both of these shortcomings. Wang et al. [2021a] consider a variety of symmetries in constructing their Equ-ResNet and Equ-Unet for dynamics forecasting, including *exact* scale and rotation symmetries. However, dynamics often only exhibit approximate symmetries due to, for example, external forces [Wang et al. 2022i]. Wang et al. [2022i] thus relax equivariance constraints in constructing their RGroup and RSteer CNNs for approximately equivariant group and steerable convolutions, respectively [Cohen and Welling 2016, 2017b].

Instead of steerable or group convolution, Ruhe et al. [2023] take a different path to encode symmetries. They build on Clifford neural layers [Brandstetter et al. 2023] to propose Geometric Clifford Algebra Networks (GCANs) and, in achieving equivariance, leverage the fact that all isometries (translations, rotations, and reflections) can be decomposed into a sequence of reflections. In addition to demonstrating the effectiveness of their convolutional architecture as a neural solver compared to non-equivariant baselines, they show that their framework can be generalized to MLPs and GNNs.

While the previously discussed CNNs perform equivariant convolution in physical space, Helwig et al. [2023] extend group equivariant convolutions [Cohen and Welling 2016] to a frequency domain parameterization with the G-FNO architecture. In addition to the benefits brought in terms of multiscale processing as discussed in Section 9.2.3, this allows for superior generalization to discretizations with different resolution relative to physically parameterized alternatives [Li et al. 2021b].

9.2.7 Existing Methods: Incorporating Physics.

While deep neural networks are universal function approximators, practitioners often have deep insight into the behavior of physical systems. By carefully designing architectures to automatically respect physical laws that these systems obey, the learning task is simplified and the ability of the network to generalize over similar systems is improved. This is because these rules are difficult to learn from data directly, particularly in the small data regime. Further, encoding physical laws often increases the interpretability of network outputs, as they can be directly related to concepts practitioners are familiar with, which is in stark contrast to the usual treatment of neural networks as a black box modeling tools.

Hamiltonian Neural Networks (HNNs) [Greydanus et al. 2019] incorporate physics knowledge in the form of Hamiltonian mechanics for faithful modeling of Hamiltonian systems. In general, these systems are described by position $q(t)$ and canonical momenta $p(t)$, which evolve in time according to Hamilton's equations as

$$\dot{q} = \frac{\partial H}{\partial p}, \quad \dot{p} = -\frac{\partial H}{\partial q}, \quad (107)$$

where \dot{q} and \dot{p} are the derivatives of q and p with respect to time. Hamiltonian systems are everywhere – the motion of planets under the influence of gravity, particles impacted by electromagnetic forces, and blocks attached to springs all follow Hamiltonian mechanics. A key property of Hamiltonian systems is that as the system evolves over time, the Hamiltonian $H(q(t), p(t))$ is *conserved*, that is, it remains constant. Loosely, the Hamiltonian H captures the amount of energy in the system. HNNs [Greydanus et al. 2019] propose learning this Hamiltonian H directly from dynamics data. The time evolution of the Hamiltonian system is then computed by numerically integrating Equation (107) over time using an explicit Runge-Kutta method of order 4. The HNN model accurately learns the time evolution of simple Hamiltonian systems such as an oscillating pendulum without dissipating energy, making predictions that maintain consistency with Hamilton's equations. In contrast, a standard fully-connected neural network trained on the same data is unable to learn trajectories that conserve H , resulting in physically implausible predictions.

In theory, the current state $u(t) = (q(t), p(t))$ of a Hamiltonian system completely determines its state $u(t')$ at all future times $t' > t$. With this motivation, SympNets [Jin et al. 2020] directly learn the mapping from the current system state $u(t)$ to the future system state $u(t')$ using symplectic normalizing flows to avoid integrating over time. This makes SympNets more efficient over longer rollouts, as they avoid accumulation of numerical errors present during numerical integration. Symplectic Recurrent Neural Networks [Chen et al. 2020b] further improve upon Hamiltonian Neural Networks by using symplectic integrators such as the leapfrog method, which are a better fit for Hamiltonian systems than explicit Runge-Kutta schemes because they explicitly match the form of Equation (107). Thus, the symplectic integrator will conserve the learned Hamiltonian H when integrating over time up to numerical precision. Further, Chen et al. [2020b] propose training over longer rollouts produced by sampling the model recurrently instead of single-step predictions. This helps avoid the distributional shift problem inherent when recursively sampling from the model's predictions at each time step, as discussed in Section 9.2.5. Finally, to account for noise in the system observables, Chen et al. [2020b] update the initial state $u_0 = (q_0, p_0)$ via gradient descent. These modifications improve the accuracy of the HNN when modeling noisy, real-world systems.

Instead of general Hamiltonian systems, Action-Angle Networks [Daigavane et al. 2022] leverage properties of the special class of *integrable* Hamiltonian systems by learning a symplectic transformation of position q and momenta p to slow-varying action variables and fast-varying angle variables. For integrable systems, the dynamics in the action-angle space are effectively linear, which makes it both easier to learn and more efficient to numerically integrate compared to HNNs and Neural ODEs [Chen et al. 2019a].

As opposed to Hamiltonian systems, Lagrangian Neural Networks (LNN) [Cranmer et al. 2020] model Lagrangian systems where the form of the canonical momenta p is not necessarily known. Instead, Lagrangian mechanics provide the necessary insight to relate the time evolution of the position q as

$$\ddot{q} = \left(\frac{\partial^2 L}{\partial \dot{q}^2} \right)^{-1} \left(\frac{\partial L}{\partial q} - \dot{q} \frac{\partial^2 L}{\partial \dot{q} \partial q} \right), \quad (108)$$

where \dot{q} and \ddot{q} are the first and second derivatives of the position q with respect to time, and $L(q, \dot{q})$ is the Lagrangian of the system. Analogous to HNNs with the Hamiltonian H , LNNs model the Lagrangian L via a neural network. Then, by numerically integrating Equation (108), the time evolution of $q(t)$ can be obtained.

Finally, Sosanya and Greydanus [2022] augment HNNs to additionally predict a Rayleigh dissipation function D together with the Hamiltonian H . This allows the network to capture external forces such as friction which dissipate energy. Such forces cannot be captured in the original HNN framework because the HNN learns energy-conserving dynamics. The Dissipative HNN shows improved performance on predicting the time evolution of damped spring-block systems and the velocity fields of ocean surface currents.

While Hamiltonian mechanics can describe a large number of physical systems, in many real world scenarios, the system may not be sufficiently well understood or only partially observed. If the underlying PDEs describing the system are only partially known, physical knowledge can be leveraged in a hybrid setup. In this context, deep neural networks can be applied in conjunction with PDE-based methods to learn the residual between assumed governing equations and observed data. A representative example is the APHYNITY framework proposed by Yin et al. [2021], which operates on the premise that dynamics can be decomposed into physical (known) and augmented

(residual) components as

$$\partial_t u + (\mathcal{D} + \phi_\theta)(x, t, u, \partial_x u, \partial_{xx} u, \dots) = 0, \quad (109)$$

where ϕ_θ represents the data-driven component that complements the known operator \mathcal{D} . When learning the parameters of ϕ_θ , numerical integration is used to generate predictions at various steps based on $\mathcal{D} + \phi_\theta$ given an initial state. More importantly, it efficiently augments physical models with deep data-driven networks in such a way that the data-driven model only models what cannot be captured by the physical model. To achieve this, other than prediction loss, an additional L2 norm term $\|\phi_\theta\|_2$ is imposed on ϕ_θ . This avoids the situation that all or most of the dynamics could be captured by neural nets and the physics-based models contribute little to learning.

Similarly, DeepGLEAM [Wu et al. 2021] is a method used for predicting COVID-19 mortality by directly combining the mechanistic epidemic simulation model GLEAM with neural nets. GLEAM [Balcan et al. 2009] is a PDE-based model that characterizes complex epidemic dynamics based on meta-population age-structured compartmental models. DeepGLEAM employs a DCRNN [Li et al. 2017b] to learn the errors made by GLEAM, resulting in enhanced performance for one-week ahead COVID-19 death counts predictions.

Beyond applications to observed dynamics data, hybrid methods can be used to substitute the computationally intensive components of classical solvers or learn corrections for classical solvers applied on inexpensive but error-inducing coarse discretizations. Belbute-Peres et al. [2020] introduce a novel approach termed CFD-GCN that combines graph convolutional neural networks with a Computational Fluid Dynamics (CFD) simulator. This hybrid method aims to generate accurate predictions of high-resolution fluid flow. It runs a fast CFD simulator on a coarse triangular mesh to generate a lower-fidelity simulation, which is subsequently enhanced by upsampling it to finer meshes using the K-nearest neighbor interpolation technique. The fine-grained simulation is then processed by a graph convolutional neural network, which further refines the predictions for specific physical properties. Similarly, Kochkov et al. [2021b] utilize CNNs to perform learned interpolation and learned correction on coarse velocity components produced by classic numerical solvers, leading to significant speedup in simulating high-resolution fluid velocity fields. Moreover, Tompson et al. [2017] replace the numerical solver for solving Poisson's equations, which is the most computationally expensive step in the procedure of traditional Eulerian fluid simulation, with convolution networks. This approach results in significant speedup and demonstrates physically consistent predictions with strong generalization abilities.

As opposed to approximating the numerical PDE solution as in the previously discussed works, Physics-Informed Neural Networks (PINNs) aim to approximate the analytical solution by parameterizing the neural network ϕ_θ as the PDE solution [Raissi et al. 2019]. Using backpropagation, the spatial and temporal derivatives in Equation (104) can be exactly evaluated and used as a regularizing agent in an effort to ensure that the constraints prescribed by Equation (104) are approximately satisfied. Thus, for the operator \mathcal{T} defined as

$$\mathcal{T}\phi_\theta(x, t) = \partial_t \phi_\theta + \mathcal{D}(x, t, \phi_\theta, \partial_x \phi_\theta, \partial_{xx} \phi_\theta, \dots) \quad (x, t) \in U, \quad (110)$$

the network ϕ_θ can be optimized over the parameter space Θ as

$$\phi_\theta = \arg \min_{\phi_\theta: \theta \in \Theta} \lambda_{\mathcal{T}} \mathbb{E}_{x, t \in U} [\|\mathcal{T}\phi_\theta(x, t)\|] + \lambda_{\mathcal{B}} \mathbb{E}_{x, t \in \partial \mathbb{X} \times \mathbb{T}} [\|\mathcal{B}\phi_\theta(x, t)\|] + \mathbb{E}_{x \in \mathbb{X}} [\|\phi_\theta(x, 0) - u_0(x)\|]. \quad (111)$$

$\lambda_{\mathcal{T}}$ and $\lambda_{\mathcal{B}}$ are coefficients for balancing different loss terms, which require careful tuning. PINNs have found real-world applications in biomedical analyses of blood flow [Raissi et al. 2020] with the Hidden Fluid Mechanics framework, and have been coupled with neural operator architectures such as the Physics-Informed DeepONet [Wang et al. 2021b] and Physics-Informed Neural Operator [Li

Table 32. Selected PDE datasets for forward modeling. We highlight challenging datasets that have arisen from neural PDE solver benchmarks [Takamoto et al. 2022a; Gupta and Brandstetter 2022] and works introducing methodologies [Tran et al. 2021; Pfaff et al. 2021]. These datasets model a variety of fields across 1,2 and 3 spatial dimensions, and include challenging tasks such as fast moving and turbulent dynamics, large time step prediction, conditional prediction, and irregular geometries.

Dataset	Source	Fields Modeled	Spatial Dimensions	Task Details
Compressible Navier-Stokes	Takamoto et al. [2022a]	Density, Pressure, Velocity	1,2,3	Initial Mach number=1 and viscosity= 1×10^{-8} yield fast-moving and turbulent dynamics
Shallow Water	Gupta and Brandstetter [2022]	Density, Velocity, Vorticity	2	Global weather on a rectangular domain with 48 hour time step
Incompressible Navier-Stokes	Gupta and Brandstetter [2022]	Pressure, Velocity, Vorticity	2	Includes conditional task: advance the state of the system conditioned on variable timestep size and forcing term
TorusVis and TorusVisForce	Tran et al. [2021]	Vorticity	2	Variable viscosity coefficient and forcing term. Includes time-varying forcing term
CylinderFlow and AirFoil	Pfaff et al. [2021]	Momentum, Pressure, Density	2	Irregular mesh on Eulerian systems
DeformingPlate and FlagDynamic	Pfaff et al. [2021]	Position, von-Mises Stress	3	Irregular mesh on Lagrangian systems

et al. 2021g] to improve sample complexity and even allow for fully self-supervised training. Yang et al. [2021b] further propose B-PINNs, which extends the concept of PINNs into the Bayesian framework. Under this approach, a PINN is used as the prior for solving partial differential equations (PDEs), while the Hamiltonian Monte Carlo method is employed to draw samples from the resulting posterior distribution. Compared with PINNs, B-PINNs not only provide uncertainty quantification but also obtain more accurate predictions on noisy data due to their capability of avoiding overfitting.

9.2.8 Datasets and Benchmarks.

The rise of neural PDE solvers has elicited many datasets for forward PDE modeling, several of which we highlight here and summarize in Table 32. Takamoto et al. [2022a] put forth PDEBench, which contains numerical solution data for 8 different PDEs with varying spatial dimensions. Beyond forward problems, Takamoto et al. [2022a] also consider inverse problems, a task we discuss in Section 9.3. Perhaps the most challenging PDEBench dataset is the compressible Navier Stokes equations for modeling the density, pressure, and velocity fields of a compressible fluid, which Takamoto et al. [2022a] include in one, two and three spatial dimensions. Fluids with velocities approaching or exceeding the speed of sound must be considered compressible, that is, as having a non-constant density [Anderson 2017]. Thus, Takamoto et al. [2022a] have released versions of this data with the initial Mach number, quantifying the ratio between the velocity of the fluid to the speed of sound in the fluid [Anderson 2017], as high as 1. Further, the low viscosities considered by Takamoto et al. [2022a] produce highly turbulent dynamics which must be resolved at small scales for stable simulation [Kochkov et al. 2021b].

Gupta and Brandstetter [2022] consider a particularly difficult realization of the shallow water equations generated using the global atmospheric model developed by Klöwer et al. [2022]. This PDE is derived by depth-integrating the Navier-Stokes equations, and, despite the name, can model fluids beyond water, just like Navier-Stokes equations [Vreugdenhil 1994]. This dataset of over 5,000 trajectories models global pressure, wind velocity, and wind vorticity fields. Gupta and Brandstetter [2022] consider the task of advancing the system by 48 hour intervals, a coarse mapping that is especially challenging to learn. Gupta and Brandstetter [2022] also released data for modeling the incompressible Navier-Stokes equations generated by the Φ_{Flow} solver [Holl et al. 2020], and consider an interesting conditional task in which the learned solver makes predictions to future

timesteps conditioned on varying timestep sizes and forcing terms. In a similar vein, [Tran et al. \[2021\]](#) have also put forth data for modeling the vorticity field of a fluid with the incompressible Navier-Stokes equations while generalizing over the viscosity coefficients and forcing terms.

Lastly, for PDEs modeled on irregular geometries, [Pfaff et al. \[2021\]](#) have released several datasets which span both Lagrangian and Eulerian systems. The data for Eulerian systems are fluid flow around a cylinder and air flow around an airfoil, visualized in Figure 34, and differ from Lagrangian systems in that the mesh is fixed. By contrast, the Lagrangian datasets, where the mesh deforms with the system, are a flag blowing in the wind and a deformable metal plate.

We note that while these works have curated an impressive selection of challenging PDEs foundational to the study of numerical solvers, solvers performing well on these tasks may not immediately generalize well to real-world applications of PDE modeling. Dynamics encountered in industrial settings can involve an interplay with complex external forces and occur in large, irregularly-shaped domains. In such settings, the previously discussed challenges of multiscale processing (Section 9.2.3), multiresolution modeling (Section 9.2.4), and rollout stability (Section 9.2.5) increase both in difficulty and importance. Future benchmarks should design tasks to explicitly probe these areas in more realistic scenarios. We furthermore note that while the emphasis in all of these datasets has been on time-evolving PDEs, modeling steady-state problems is a relevant, yet under-explored area, with initial data in this area put forth by [Bonnet et al. \[2022\]](#) with the steady-state Navier-Stokes equations.

9.2.9 Open Research Directions.

We close this section on forward modeling with a discussion of challenges faced by neural solvers that have largely been unaddressed by current works.

A primary limitation of learned solvers is the requirement of an adequate number of training data generated by costly numerical solvers [[Raissi et al. 2019](#); [Brandstetter et al. 2022b](#)], which is particularly problematic at the industry scale. Thus, improving the ability to generalize and sample complexity of learned solvers is necessary to justify and reduce this cost for their adaptation to real-world settings. Toward this goal, a richer subfield in the learned solver literature targeting techniques for out-of-distribution (OOD) dynamics should develop. In contrast to many current works that train solvers to generalize over initial conditions or PDE parameters from the same distribution as the training set, the goal of this work should be to enable learned solvers to accurately infer dynamics beyond those observed during training. Models excelling in this OOD setting will permit the span of the PDE solutions in the training set to be a subspace of the span of those in the test set, thereby improving sample complexity. OOD settings have been examined in the literature, such as [Kochkov et al. \[2021b\]](#), who study the performance of their hybrid classical - neural solver under OOD domain size, external forcing, and PDE parameters. [Stachenfeld et al. \[2021\]](#) similarly study OOD domain size, as well as OOD initial conditions and rollout length. However, there is a gap in the literature around works developing principled approaches in the regime of OOD dynamics. Initial works in this area have treated the parameters of the differential equation as environments in the context of meta-learning and trained a model that can be transductively fine-tuned at test time to adapt to unseen, OOD environments [[Wang et al. 2022j](#); [Mouli et al. 2023](#); [Kirchmeyer et al. 2022](#)].

A second factor limiting applicability is that much of the literature focuses on problems with only one or two spatial dimensions despite the prevalence of three-dimensional problems in industrial applications of PDE modeling. While many architectures discussed in this section admit an immediate extension to three spatial dimensions, in practice, three-dimensional modeling presents obstacles in the form of limited memory that must be carefully handled [[Wu et al. 2022c](#); [Lam et al. 2022](#); [Bi et al. 2022](#)]. Beyond memory requirements, the optimization of a three-dimensional model

is naturally more challenging than its two-dimensional counterpart due to the increased size of the search space. Furthermore, in three dimensions, challenging dynamics not present in lower dimensions may be introduced, such as the prominent case of turbulent flows, where the transition to three dimensions induces chaos to a degree that is unseen in two-dimensional flows due to the energy cascade discussed in Section 9.2.3 [Lienen et al. 2023]. Thus, future works should look to design neural solvers that are both scalable to three spatial dimensions and that have sufficient inductive biases for the optimization to effectively navigate the search space while maintaining sufficient expressiveness to faithfully model more challenging dynamics.

Additionally, as neural networks struggle to model non-smooth functions, modeling systems with sudden changes, such as the trajectory of a ball bouncing off a wall, remains a challenge. Such problems represent an extreme version of *stiffness*, because the timescale of such drastic interactions is orders of magnitude smaller than the usual time step size for advancing the system. Chen et al. [2020b] propose a method to handle one-time interactions of such a kind by augmenting the update equation in their integrator with a *rebound* module. Kim et al. [2021] propose efficient methods for computing the gradients and appropriate normalization of Neural ODEs [Chen et al. 2019a] to model stiff systems. However, there is still a great need to identify more general solutions to model stiff systems.

We lastly find several of the settings generated from the intersections of the challenges discussed in Sections 9.2.3 to 9.2.6 to be relevant, yet understudied, problems. Dynamics that evolve on multiple scales (Section 9.2.3) and have localized regions of high gradient (Section 9.2.4) require models that can operate on irregularly-discretized functions with multiscale processing mechanisms. Janny et al. [2023] have taken initial steps in this setting with their mesh transformer, which uses graph attention as a multiscale processing mechanism and manages complexity by pooling nodes in the same neighborhood prior to computing attention scores. Additionally, multiscale processing mechanisms (Section 9.2.3) have been incorporated into equivariant neural solvers (Section 9.2.6) by Wang et al. [2021a] with their equivariant U-net and by Helwig et al. [2023] with their group convolutions in the frequency domain. Future work may explore adapting equivariant attention [Fuchs et al. 2020; Liao and Smidt 2023] as a multiscale processing mechanism in neural solvers. Finally, equivariant neural solvers (Section 9.2.6) that can operate on irregularly structured input (Section 9.2.4) are an interesting direction, with initial work in this area done by Horie et al. [2021] with their IsoGCN.

9.3 Inverse Problem and Inverse Design

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In Section 9.2, we have delved into the advances and challenges of neural PDE solvers for simulating the *forward* evolution of PDEs. The reverse direction is equally exciting, including (1) *inverse problems*, where the task is to infer the unknown parameters or state of the system given (partial) observations of the dynamics, and (2) the emerging direction of AI-assisted *inverse design*, where the task is to optimize the system (parameters or components such as initial or boundary conditions) based on a predefined objective. Both tasks are universal across science and engineering. In Figure 35, we conceptualize forward problems, inverse problems, and inverse design.

9.3.1 Problem Setup.

Let $u(x, t; \gamma)$ be a forward model that describes a physical process and is induced by the PDE configuration $\gamma = (u_0, \mathcal{B}, \gamma_P)$ describing the initial conditions u_0 , boundary conditions \mathcal{B} , and PDE parameters γ_P . Furthermore, let $\tilde{\gamma} \subset \gamma$ be the properties to be recovered (in inverse problems) or the design parameters to be optimized (in inverse design). Finally, let \mathcal{J} be an objective function

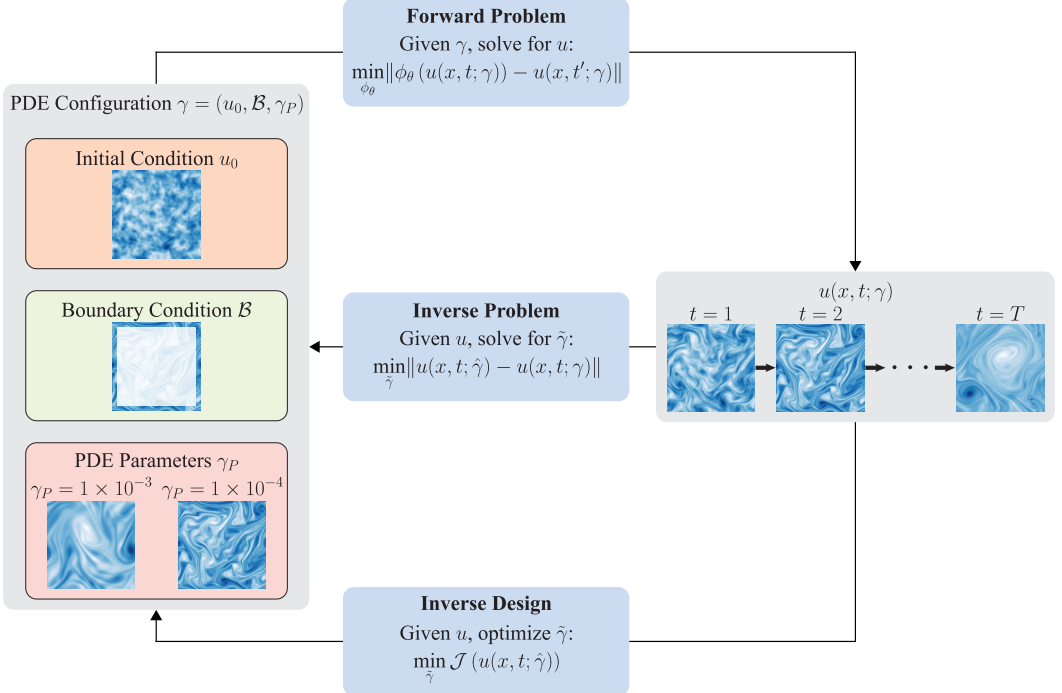


Fig. 35. Illustration and comparison of forward problems, inverse problems, and inverse design. The solution to a PDE $u(x, t; \gamma)$ sampled on grid points (x, t) discretizing space-time is induced by the PDE configuration $\gamma = (u_0, \mathcal{B}, \gamma_P)$ describing the initial conditions u_0 , boundary conditions \mathcal{B} , and PDE parameters γ_P . In forward problems, the task is to learn the mapping from the solution induced by a particular choice of γ at earlier time steps t to the solution at later time steps $t' > t$ using the forecasting model ϕ_θ . Conversely, inverse problems consider the task of identifying a subset of the PDE configuration $\hat{\gamma} \subset \gamma$, such as the initial conditions u_0 , that generated the observed rollout data. The data is assumed to have originated from a forward model $u(x, t; \gamma)$, and the estimated configuration $\hat{\gamma}$ is optimized by minimizing the discrepancy between $u(x, t; \hat{\gamma})$ and the observed data $u(x, t; \gamma)$, where $\hat{\gamma}$ denotes the union of the estimated components of the configuration $\tilde{\gamma}$ with the known components. Lastly, inverse design involves identifying $\tilde{\gamma} \subset \gamma$ such that the resulting rollout $u(x, t; \tilde{\gamma})$ optimizes some criterion \mathcal{J} , such as identifying the shape of an airplane wing that minimizes drag.

which evaluates the quality of recovery or design. The inverse problem and inverse design can then be formulated as an optimization problem [Lu et al. 2021b] as

$$\hat{\gamma} = \arg \min_{\tilde{\gamma}} \mathbb{E}_{x,t} [\mathcal{J}(u(x, t; \tilde{\gamma}))], \quad (112)$$

where $\hat{\gamma}$ represents the union of the components of the PDE configuration to be estimated $\tilde{\gamma}$ with the remaining components that are assumed to be known. For example, when modeling a dynamic system, u typically defines what the rollout would be when a certain initial condition and boundary condition is given and \mathcal{J} measures the difference between the simulated rollout induced by $\hat{\gamma}$ and the observed or targeted rollout. In the above formulation, u is fixed and can be modeled with a classical PDE solver. To accelerate and improve the optimization, u can also be a learned model and can be made to be differentiable. In this case, additional constraints on u may be required to ensure

physical consistency, and thus, the joint optimization of \hat{y} and u is constrained as:

$$\hat{y}, u = \arg \min_{\hat{y}, u : C(u, \hat{y}) \geq 0} \mathbb{E}_{x,t} [\mathcal{J}(u(x, t; \hat{y}))] \quad (113)$$

where C can be the constraints stemming from the PDE or other constraints from multi-objective optimization [Lu et al. 2021b].

Inverse Problems versus Inverse Design: Despite the similarity suggested by their name, inverse problems and inverse design have different meanings in the context of PDEs. An inverse problem refers to the setting where some or all of the initial conditions, boundary conditions, or coefficients of the PDE are unknown, where the objective is then to determine or recover these unknowns from the observed data. An inverse problem typically assumes that the observed data is physically plausible and represents the solution to the PDE. For instance, in fluid mechanics, the observed data might be the vorticity field, and only the initial condition is unknown. Then, the inverse problem would be to determine the initial condition u_0 that would produce such a vorticity field. Alternatively, inverse design refers more specifically to a design or optimization methodology in which a predefined objective is given, and the goal is to optimize the system configuration based on the objective. For instance, given a surrogate model u that can simulate forward fluid dynamics, the objective might be to design a surface that can guide the fluid flowing to the desired location. For inverse design, an exact solution may not necessarily exist, however, we may still want to optimize the proposed solution to satisfy the objective as much as possible. In some sense, inverse design can also be thought of as a specific type of inverse problem, where the goal is not just to determine unknown parameters or coefficients, but to design a system that behaves in a certain way.

Applications of Inverse Problems: Here we describe several examples of inverse problems that hold potential for AI to create new opportunities, which we outline in Figure 32.

- **Fluid dynamics grounding:** Learning a surrogate model of fluid dynamics typically requires the use of an expensive classical solver to obtain training data. An alternative approach is to consider an inverse problem, where the task is to infer the underlying dynamics solely based on a multi-view video of a 3D dynamical fluid scene [Guan et al. 2022].
- **System identification:** Traditionally, estimating the physical properties of an object requires conducting many physical experiments and the use of specifically designed algorithms. A promising inverse problem here is to infer the physical properties directly from visual observations [Li et al. 2023c].
- **Full waveform inversion for geophysics:** In geophysics, underground properties such as density or wave speed can be inferred from the measurement of seismic waves on the ground surface, a problem termed as *full waveform inversion* [Lin et al. 2023c]. These underground properties are important for applications such as energy exploration or earthquake early warning, which are otherwise difficult to measure due to the large scale of the problem.
- **Fluid assimilation and history matching:** Fluid assimilation aims to recover the entire fluid field from sparse observations in the spatio-temporal domain [Zhao et al. 2022]. Fluid assimilation can be applied to model underground flow. The geological model is adjusted such that the predictions match the historical observations, a task termed as *history matching* [Tang et al. 2021].
- **Tomography for medical imaging:** Tomography aims to recover internal structures of an object using only surface measurements. For example, in medical imaging, electrical impedance tomography (EIT) [Guo et al. 2023a] can infer the status of internal organs by measuring the voltage distribution on skin when an electrical current is injected, which avoids intrusive measurements or radiation exposure.

Applications of Inverse Design: Here we identify a few applications where AI-assisted inverse design can play a significant role and where vast opportunities lie.

- **Shape design for planes:** In aerodynamics, an important challenge is designing the shape of planes to minimize drag [Athanasopoulos et al. 2009]. This involves simulating the air fluid dynamics and its interaction with the boundary shape of the plane.
- **Ion thruster design:** In aerospace engineering, the design of efficient thrusters is highly important. For example, the Hall effect thruster (HET) is one of the most attractive electric propulsion (EP) technologies, since it has high specific impulse and high thrust density. One key question is how to design the shape and material arrangement of the thruster, given its complicated plasma dynamics [Hara 2019].
- **Controlled nuclear fusion:** Solving controlled nuclear fusion can pave the way for unlimited clean and cheap energy. In magnetic confinement with Tokamak, one of the two main approaches to controlled nuclear fusion, a key challenge is to optimize the external magnetic field and wall design in order to shape the plasma into configurations with good stability, confinement and energy exhaust [Ambrosino et al. 2009; Degrave et al. 2022].
- **Chip manufacturing:** Many processes in chip manufacturing involve inverse design. One important application is plasma deposition. Specifically, the problem is how to design the shape of the dielectric cell so that the deposition of the plasma onto a substrate is as smooth as possible [Hara et al. 2023].
- **Shape design for underwater robots:** In underwater robots, an important problem is to design the shape of the robots to achieve multiple objectives, including minimizing drag, improving energy efficiency, improving dirigibility, and improving certain acoustics properties [Saghafi and Lavimi 2020].
- **Addressing climate change:** Inverse design can play a significant role in many approaches to address climate change, including improving materials for buildings, optimizing carbon capture, solar geoengineering, and design of carbon credits and policy [Rolnick et al. 2022].
- **Nanophotonics:** Nanophotonics focuses on designing structures with a scale close to the wavelength of electromagnetic waves. Developing principled methods for designing micro-scale structures, nano-scale structures, or topological patterns to interact with light has important implications in applications such as laser generation, data storage, chip design, and solar cell design [Molesky et al. 2018].
- **Battery design:** Deep learning-enabled inverse design has vast potential in battery design. For example, it can be used for the inverse design of battery interphases, which is important for developing high-performance rechargeable batteries [Bhowmik et al. 2019]. Besides the battery itself, hyperparameter-searching techniques in machine learning can be used to accelerate the experimental exploration of high-cycle-life charging protocols of lithium-ion batteries [Attia et al. 2020], which is critical for electric cars.

9.3.2 Technical Challenges.

Common Challenges: As inverse problems and inverse design involve the forward modeling task to evaluate $u(x, t; \gamma)$ in Equations (112) and (113), challenges encountered in forward problems discussed in Section 9.2.2 are typically also present here. Another common challenge in both inverse problems and inverse design is adversarial modes. This can occur when the high-dimensional parameters are inferred or designed with a deep learning-based surrogate model. Parameters with noisy, adversarial modes can occur that are not physically plausible, but achieve an excellent loss [Zhao et al. 2022]. In the following, we illustrate several further unique challenges.

Challenges for Inverse Problems

- **Objective mismatch:** When the forward model and the inverse problem objective are jointly optimized, the forward model might sacrifice the physical constraints given by the underlying PDE in exchange for increased optimality in the inverse problem objective, resulting in physically inconsistent solutions.
- **Ill-posedness:** In many applications, a complete measurement is often not available, which makes the inverse problem ill-posed and the solution non-unique. For example, when modeling a fluid, it is not feasible to track the movement of every fluid element. Thus, sparse measurement must be used. Another example is in tomography, where the problem is fundamentally ill-posed, as we try to infer inner structure solely from measurements on the boundary.
- **Indirect observation:** In some scenarios, it is hard or expensive to conduct direct measurement of physical states of an object or solution field. Instead, we may only be able to afford to video the object moving and interacting with the environment. Inferring the unknown parameters only from visual observations then poses a significant challenge.
- **Incorporating Physics:** Just as it is challenging to incorporate physics principles into the forward problem, it is equally important to ensure that inverse models adhere to the desired physical laws. It is crucial to extract relevant physical knowledge from well-established theories and incorporate it into the design of inverse models, while still maintaining sample and training efficiency and accuracy without any compromises.

Challenges for Inverse Design

- **Complex design space:** A fundamental challenge in inverse design, especially for real-world applications, is that the design space is hierarchical, heterogeneous, and consisting of many components that may be combined in many different ways. Take rocket design as an example. On a high level, a rocket consists of an airframe, a propulsion system, and a payload, each of which may consist of hundreds of parts. Thus, it presents a significant challenge to *represent* the complex design space and to *optimize* with respect to the chosen representation.
- **Multiple (contradicting) objectives:** Real-world engineering design problems typically have multiple objectives that may contradict one another. For example, to design a phone battery, we simultaneously want the battery to have a long lifespan and be lightweight. These two objectives contradict each other, and thus, we must find a balanced trade-off.
- **Temporally changing importance for multiple objectives:** In different scenarios, the importance of objectives may differ from one another. For example, as a rocket launches and transitions from ground to space, it will encounter drastically different environments, resulting in varying importance for its objectives of air resistance, fuel efficiency, and structure durability.

9.3.3 Existing Methods.

Inverse Problem: Recently, neural radiance field (NeRF) [Guan et al. 2022] has been applied to fluid dynamics grounding and system identification. NeuroFluid infers the underlying fluid dynamics from sequential visual observations by jointly training a particle transition model and a particle-driven neural renderer. PAC-NeRF [Li et al. 2023c] designs a hybrid Eulerian-Lagrangian representation of the neural radiance field combined with a differentiable simulator for estimating both physical properties and geometries of dynamic objects from sequential visual observations.

Zhao et al. [2022] tackles the problem of fluid assimilation from sparse fluid rollout observations, as well as the full waveform inversion problem. The objective is to find an initial condition such that the simulated rollout is close to the observed rollout on sparse measurement locations. To enable an adaptive spatial resolution, a mesh-based data representation is used in conjunction

with a learned GNN model [Pfaff et al. 2021] as a forward model to forecast dynamics from the initial condition. To tackle the ill-posedness, Zhao et al. [2022] propose to learn a latent vector for the entire fluid field, then infer the quantities to be recovered at each mesh point from the concatenation of the latent vector and the mesh coordinate.

Another important class of methods to address inverse problems is Physics-Informed Neural Networks (PINNs) [Raissi et al. 2019]. PINNs are a class of methods addressing forward and inverse problems simultaneously. They parameterize the solution function as a neural network optimized (using backpropagation) with an objective that consists of both a data loss, which penalizes the discrepancy of the neural network solution with observed data, and a physics-informed loss, which penalizes the violation of the provided PDE. During training, unknown parameters of the PDE or the system can also be learned. Furthermore, Lu et al. [2021b] develop a new PINN method with hard constraints (hPINN) to solve PDE-constrained inverse design while avoiding commonly-seen optimization issues in PINNs [Krishnapriyan et al. 2021]. The hPINN method utilizes two different techniques to enforce hard constraints on PINNs. One is the penalty method that gradually increases coefficients of the loss terms of boundary conditions and PDEs throughout training. The second technique involves the augmented Lagrangian method, which employs carefully selected multipliers in each iteration to enforce the constraints effectively.

In situations where it becomes essential to identify the accurate governing equations for practical problem-solving, various attempts have been made to derive the precise mathematical formulation based on observed data. The conventional approach [Schaeffer 2017; Brunton et al. 2016; Kaiser et al. 2018] often involves selecting from a wide dictionary of potential candidate functions and finding the combination of a subset that minimizes the discrepancies between the model predictions and the observed data. Recently, many studies have also employed neural networks to augment the dictionary of candidate functions or to capture more intricate relationships between these functions. For instance, Rudy et al. [2017] utilize neural networks as supplementary candidate functions in addition to predefined basis functions to model more complex dynamics. Martius and Lampert [2016]; Sahoo et al. [2018] introduce EQL which utilizes neural nets to identify complex governing equations from observed data. Rather than relying on conventional activation functions, they employ predefined basis functions, including identity and trigonometric functions. Additionally, they integrate custom division units into the framework to capture division relationships within the potential governing equations. However, generalizability and over-reliance on high-quality measurement data remain critical concerns in this research area.

Inverse Design: Past methods to address inverse design are mostly based on domain-specific classical solvers, which are extremely computationally expensive. Recently, with the success of neural PDE solvers, AI-assisted inverse design has also emerged, but largely remains unexplored. One notable work is by Allen et al. [2022], which uses backpropagation through time (BPTT) over the entire differentiable physical simulation to design the boundary for particle-based simulations. However, this method is still computationally expensive, as it must compute the gradient three times for hundreds of steps of simulation in the input space. Wu et al. [2022a] introduce BPTT in the latent space for inverse design, which improves both the runtime and accuracy compared to inverse design in the input space. For Stokes flow, Du et al. [2020] develop a method to simulate and optimize Stokes systems governed by design specifications with different types of boundary conditions. Li et al. [2022a] further introduce an anisotropic constitutive model for topology optimization that can generate new topological features that differ drastically from the initial shapes and enable flexible modeling of both free-slip and no-slip boundary conditions.

The above initial works of AI-assisted inverse design are limited to relatively simple and idealized scenarios. Therefore, there is a massive gap between the tasks considered by these works and

those in real-world engineering in terms of the following aspects: (1) *Complexity of the physics*: The physics in real-world systems may be multiresolution or even multiscale, making efficient and accurate simulation difficult. (2) *Complexity of the design*: Real-world systems consist of many parts, requiring the system to be designed in a more hierarchical and structured way. (3) *Generality and diversity*: The tasks tested above are restricted to a specific domain, and are not diverse enough to test the methods' generality across multiple disciplines. These challenges provide great opportunities to develop novel neural representations and methods for proposing improved designs. A related work is by [Degrave et al. \[2022\]](#), which for the first time, employs deep reinforcement learning (RL) for shaping fusion plasma, and demonstrates that deep RL is able to control such complex systems. This work further demonstrates the feasibility of such a method on complex physical systems and serves as inspiration for the community to work on more challenging problems that have the potential to offer long-term beneficial impacts on humanity.

9.3.4 Datasets and Benchmarks.

For the NeRF-related inverse problem, datasets are multi-view images of a dynamic scene or object generated by simulation engines, such as MLS-MPM [[Hu et al. 2018, 2019a](#)] and DFSPH [[Bender and Koschier 2015](#)]. In the context of fluid assimilation, rollout data can be simulated using a classical solver such as finite element method solver [[Logg et al. 2012](#)] used by [Zhao et al. \[2022\]](#).

In the domain of inverse design, as introduced in Section 9.3.3, different works have tested their methods using their respective domain-specific datasets, such as the datasets considered by [Allen et al. \[2022\]](#) for designing shape for airfoil and surfaces for particle-based fluid flows, and the dataset employed by [\[Wu et al. 2022a\]](#) for designing boundaries to control smoke in fluid flow. However, there has not been a standard benchmark to evaluate different inverse design methods systematically. Furthermore, compared to real-world engineering tasks, the current datasets are significantly lacking in terms of the complexity of the physics and the difficulty of the design. This presents an excellent opportunity for the community to introduce more diverse and more complex benchmarks in terms of physics and the design task.

9.3.5 Open Research Directions.

For the inverse problem, there are several possible future directions to explore: (1) *Uncertainty quantification*: Many inverse problems are ill-posed, and this instability can lead to high uncertainty in the solution. Uncertainty quantification is therefore crucial in these cases, as it can help describe uncertainties associated with the solution. (2) *Improved training techniques*: Complex or ill-posed inverse problems present difficulty in training deep neural networks, motivating future research to develop novel training strategies and regularization techniques.

For inverse design, the challenges (Section 9.3.2) and limitations of current works (Section 9.3.3) also point toward exciting future directions. We identify several exciting opportunities. (1) *Developing novel representations*: The hierarchical, heterogeneous, and complex design space presents ample opportunity to design suitable representations that balance faithfulness and efficiency. (2) *Developing new optimization methods*: The design space is typically hybrid, consisting of discrete variables, such as the number for each part, and continuous variables, such as the shape for each part and how parts are composed. This complex space presents an exciting opportunity for the development of novel optimization methods. (3) *Developing more general methods across domains*: The diversity of real-world tasks also calls for more general methods to tackle multiple domains.

10 RELATED TECHNICAL AREAS OF AI

In addition to the challenges specific to individual science areas, there are several technical challenges that are shared across multiple domains in the field of AI for science. In particular, we identify the following four common technical challenges: out-of-distribution generalization, interpretability, foundation models powered by self-supervised learning, and uncertainty quantification. These challenges have long been recognized in the field of AI and machine learning, but they take on increased significance in the context of AI for science due to the unique characteristics of the data and tasks involved. In this section, we discuss the current limitations, existing approaches, and potential research opportunities related to these four challenges.

10.1 Interpretability

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Interpretability, despite its ubiquity in the machine learning field, lacks a unified mathematical definition. Its meaning can differ based on the context. It sometimes refers to a model's inherent ability to offer humanly understandable interpretations of its predictions, a characteristic commonly observed in models such as decision trees. On the other hand, interpretability can also refer to an in-depth understanding of intricate models. For example, an interpretation highlights how distinct input graph patterns, e.g., a substructure, can lead to a certain GNN behavior, such as maximizing a target prediction. Within the scope of this paper, we narrow our focus to instance-level interpretations which provide input-dependent explanations for each input graph. From this perspective, an interpretation sheds light on significant patterns or components of an input graph crucial for its prediction. Notably, different components of the input graphs may contribute to the model's predictions to varying extents. Thus, an effective interpretation method precisely identifies those components and patterns that significantly impact the predictions, enabling a comprehensive understanding of the underlying factors driving the predictions of models.

Geometric deep learning (GDL) models have demonstrated significant potential in solving various problems in quantum, molecular, material, and protein science. However, to assess the scientific plausibility of GDL model outcomes, it is essential to achieve interpretability of results. Unfortunately, most GDL models lack interpretability and are often treated as black boxes, which hampers their reliability and limits their applicability in scientific domains. Here we explore the importance of interpretability with the incorporation of explainable artificial intelligence (XAI) with models. XAI aims to track the contributions of specific components of the input instance to the final predictions and identify the parts that carry information indicative of the prediction label. By understanding how model outputs are determined, the trustworthiness of their predictions increases. Additionally, XAI can test if model predictions are faithful to physical laws, which in turn will help improve the quality of existing GDL models. Precise interpretation techniques of model weights and features provide domain experts with deeper insights into the underlying mechanisms learned by these models, allowing the acquired knowledge from the model to guide future research directions. Interpretability of models can be particularly valuable for design of new compounds through identification of important substructures in molecules, materials, and proteins for particular properties.

10.1.1 Existing XAI Methods.

While many XAI methods have been developed to study graph neural networks [Ying et al. 2019; Yuan et al. 2021; Gui et al. 2022b; Baldassarre and Azizpour 2019; Huang et al. 2022; Schnake et al. 2021; Xie et al. 2022b], they mainly focus on 2D graphs. According to Yuan et al. [2023], existing approaches can be mainly categorized into four classes, namely, gradients/feature-based methods,

perturbation-based methods, decomposition methods, and surrogate methods. Gradients/feature-based methods, which rely on either feature values or gradients to evaluate feature importance, have been particularly popular because of their simplicity and the intuition they provide about feature importance. Perturbation-based methods analyze the change in prediction when input features are perturbed to generate importance scores. Decomposition methods decompose prediction scores and back-propagate these scores layer by layer until the input space to compute importance scores. These approaches provide more insights into each layer of graph neural networks. Surrogate-based methods sample some similar data to a given input example and fit a simple and interpretable model like a decision tree. The explanations from the surrogate model are used to explain the original predictions. These techniques are valuable for interpreting the behavior of complex models. For a deeper understanding of graph XAI, we recommend referring to the recent surveys [Yuan et al. 2023].

Despite the progress made in XAI for 2D graph neural networks, XAI for GDL models or 3D graphs remains an underexplored field. Existing GDL methods [Wang et al. 2022d; Tubiana et al. 2022] aim to interpret their architecture through systematic analysis and visualization of the learned representations. These representations, categorized into distinct clusters, are aligned with specific physical or chemical properties. However, the prediction mechanisms of these models and the contribution of input graph components to predictions remain unknown. There are unique challenges and opportunities in this domain due to the higher dimensionality of the geometric data and the complexity of the models. Although gradient/feature-based and perturbation-based methods are useful, they are insufficient to provide a complete explanation for the importance of geometric features. On the other hand, decomposition methods and surrogate-based methods cannot be easily applied to GDL models. Recently, Miao et al. [2023] proposes a new perturbation-based method specifically designed for 3D points. This approach uses a learnable interpreter model to introduce random noise to each 3D point. The interpreter model is trained together with the GDL model used to predict labels. The amount of the learned random noise is then used to generate importance scores for each input point. However, this work only focuses on interpreting the GDL models with invariant predictions and doesn't consider the invariance and equivariance of the explanations [Crabbé and van der Schaar 2023]. Thus, there is a need for more XAI techniques specifically for equivariant GDL models.

10.1.2 Potential Application Scenarios.

The contributions of interpretability with XAI to research science can be broadly categorized into the following four perspectives, with several potential applicable scenarios for each perspective.

Improving Trustworthiness of GDL Models: XAI techniques aim to provide insight into model behaviour and predictions, such as identifying important features and substructures of inputs. Interpretability of model predictions allows researchers to better understand underlying model mechanisms and in turn promotes trustworthiness of models. In molecular property prediction, XAI could validate faithfulness to physical rules for outputs of GDL models, such as the role of the chemical structure and functional groups of molecules in determining molecular properties. Similarly, in protein fold classification, XAI techniques can help identify the most important amino acid residues or secondary structure elements for predicting a specific fold, thereby verifying whether GDL models capture secondary structural features and assisting scientists in using model outputs to make research decisions. In material property prediction, XAI could be used to validate if the model is focusing on the correct elements and structure in the material for prediction. For learning ground state of quantum spin systems, using XAI to probe how perturbations of spin configuration and electron positions change the energy of the system will assist in validating if the learned energy is physically consistent. Application of XAI to GDL models through identifying

substructures, feature importance, and effects of perturbations can be a valuable method for verifying if models are exhibiting scientifically consistent behaviour to promote trustworthiness of model predictions.

Enabling Further Scientific Knowledge Discovery: XAI may reveal patterns and insights from model predictions that can help researchers discover new hypotheses and research questions, potentially leading to discoveries of new scientific knowledge. For example, when performing molecule energy prediction, XAI can provide valuable insights into the importance of substructures and perturbations of features of different conformers of the same molecule and their corresponding energy levels, assisting future research on the generation of molecular conformers. Furthermore, for protein science it could identify key secondary structure, or amino acid residues responsible for a given predicted property. This could guide further investigation of the identified substructures of the proteins. In the scenario of studying complex systems such as quantum mechanisms and PDEs, XAI can be used to understand the behavior of the systems and identify the most important variables and factors that contribute to a system's behavior of interest. By using XAI to gain insights of how features and weights affect the model's internal representations and decision-making processes, scientists can gain insight into the underlying physical principles and test new hypotheses about unknown and under-explored systems.

Diagnosing and Improving Existing Models: XAI enables researchers to improve the existing models by examining and ensuring that GDL models satisfy physical rules. The presence of scientifically erroneous model explanations also helps expose potential biases or errors in the model, which in turn improves model quality. For example, in modeling global weather patterns using shallow water equations, it is important to ensure that the GDL models used to solve these equations satisfy physical laws such as the conservation of mass, momentum, and energy. In molecular ML researchers can also use XAI to probe feature importance and validate if it aligns with chemically expected features such as atomic number, bond angle, *etc.* XAI techniques can help researchers identify whether the predictions of GDL models adhere to these physical laws, and identify which physical constraints the model needs to better satisfy. This could be applicable in finding many-electron ground states, by checking if outputs satisfy fermion antisymmetry constraints. XAI of GDL models is important for validation of results by domain experts and is helpful for revealing limitations in predictions for improvement of existing models.

Facilitating Design of Drugs and Materials: XAI can identify the critical substructures or functional groups that contribute a desired property in drug discovery and material design. For example, in molecular interactions between small molecules and proteins, XAI techniques can identify critical parts of the protein and ligand that contribute to predicting binding affinity. By obtaining information about specific groups of amino acids that determine affinity and the location of binding sites, researchers can design drugs that are more selective and only interact with the desired protein, reducing the risk of off-target effects. In material science, identification of elements and packing orientations that lead to a molecular property prediction can help guide discovery of new materials with particular properties. For protein science, XAI may identify particular substructure of a protein linked to predicted properties that researchers could use to design *de novo* proteins with similar properties. Generative models for drugs and materials could also benefit from XAI to better understand proposed designs. For example, in generating a drug for a given protein, using XAI to highlight the importance of particular substructures in the protein and the generated drug for binding can be helpful for researchers to understand generated compounds and direct future design. XAI can guide the design of new compounds through identification of patterns and features that are important for a desirable or undesirable property.

10.2 Out-of-Distribution Generalization

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The out-of-distribution (OOD) [Gulrajani and Lopez-Paz 2020; Arjovsky et al. 2019] problem focuses on the common learning scenario where test distribution shifts from training distribution, which substantially degrades model performances in scientific discovery tasks, as shown in Figure 36. The mismatching of the distribution is commonly referred to as distribution shifts, including several concepts of covariate shift [Shimodaira 2000], concept shift [Widmer and Kubat 1996], and prior shift [Quiñonero-Candela et al. 2008]. This problem occurs in diverse application scenarios [Miller et al. 2020; Sanchez-Gonzalez et al. 2020; Myers et al. 2014; Gui et al. 2020] and is tied to various fields, such as transfer learning [Weiss et al. 2016; Torrey and Shavlik 2010; Zhuang et al. 2020], domain adaptation [Wang and Deng 2018], domain generalization [Wang et al. 2022c], causality [Pearl 2009; Peters et al. 2017], and invariant learning [Arjovsky et al. 2019; Ahuja et al. 2021].

Currently, OOD generalization methods and studies in AI for science can significantly improve overall task performances as well as generalization abilities across various domains. While numerous studies exist on general OOD generalization [Arjovsky et al. 2019; Peters et al. 2016; Tzeng et al. 2017; Lu et al. 2021c; Rosenfeld et al. 2020; Ahuja et al. 2021; Sun and Saenko 2016; Ganin et al. 2016] and non-Euclidean OOD generalization [Wu et al. 2022b; Chen et al. 2022b; Zhu et al. 2021; Bevilacqua et al. 2021; Li et al. 2023a; Gui et al. 2023], the realm of OOD methodologies for scientific applications remains largely uncharted. In this section, we aim to summarize the research on OOD approaches within the field of AI for science and emphasize the importance of further exploration.

10.2.1 Background and Settings.

The distribution shift problem is studied under various settings including transfer learning [Weiss et al. 2016; Torrey and Shavlik 2010; Zhuang et al. 2020], domain adaptation [Wang and Deng 2018], domain generalization [Wang et al. 2022c], causality [Pearl 2009; Peters et al. 2017], and invariant learning [Arjovsky et al. 2019; Ahuja et al. 2021].

In domain adaptation scenarios, we target transferring knowledge from one (source) domain to another (target) domain with distribution shifts between domains. Specifically, we can access both source domain samples with labels and target domain samples. According to the accessibility of labels in target domains, domain adaptation is typically categorized into semi-supervised and unsupervised settings. Unsupervised domain adaptation [Pan et al. 2010; Patel et al. 2015; Wilson and Cook 2020] is the most popular setting because it does not require any labeled samples in the target domains. The basic and most common idea is aligning the distributions between the source and target domains, mitigating the distribution shifts. This goal can be often done by discrepancy minimization [Long et al. 2015; Sun and Saenko 2016; Kang et al. 2019] and adversarial training [Ganin and Lempitsky 2015; Tsai et al. 2018; Ajakan et al. 2014; Ganin et al. 2016; Tzeng et al. 2015, 2017]. However, domain adaptation necessitates the pre-collected target domain samples, shrinking its application scope, e.g., privacy-sensitive applications.

Without the requirement of pre-collected target samples, domain generalization [Wang et al. 2022c; Li et al. 2017a; Muandet et al. 2013; Deshmukh et al. 2019] instead delves into the prediction for unseen domains, providing more practical solutions. Despite the prosperity of these areas, domain adaptation and generalization methods are still in need of robustly theoretical and intuitive analysis. Meanwhile, as the development of causality [Pearl 2009; Peters et al. 2017], one common sense is that generalization is logically implausible without interventions and inductive biases. Therefore, environment partitions [Ganin et al. 2016; Zhang et al. 2022] are generally used as the indicator to imply the interventions that distributions come from.

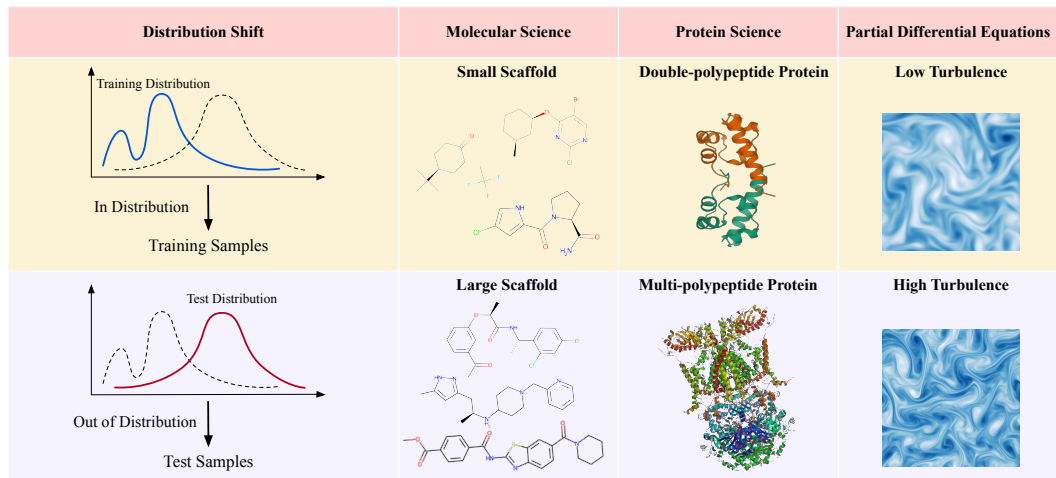


Fig. 36. Illustrations for OOD in the field of AI for science. The out-of-distribution (OOD) problem is universal among scientific tasks, where training and test samples are from different distributions. In molecular science, different molecule sizes and scaffolds are major sources of distribution shifts. In protein science, the complexity of 3D protein structures, along with the vast array of potential variations in composition and folding, renders the generalization to unseen distributions a formidable challenge. In PDEs, generalizing from higher viscosity to lower viscosity in time-evolving modeling is a difficult task since lower viscosity leads to more turbulent flows, giving rise to more chaotic dynamics and challenges in simulation.

Causality [Peters et al. 2016; Pearl 2009; Peters et al. 2017] and invariant learning [Arjovsky et al. 2019; Rosenfeld et al. 2020; Ahuja et al. 2021] can serve as the theoretical foundations for the out-of-distribution analysis, formulating various distribution shifts as graphical models or structural causal models (SCMs). Stemming from the independent causal mechanism assumption, the discovered causal correlations in SCMs are stable and ultimately endowed with physical laws. Therefore, learning causal mechanisms empowers deep models with generalization ability, leading to causality-based out-of-distribution analyses.

Peters et al. [2016] firstly introduces the concept of invariant predictions and proposes the learning strategy of optimal predictors invariant across all interventions. Motivated by the invariant learning principle, Arjovsky et al. [2019] formulates the interventions as environment partitions and proposes the invariant predictor learning strategy as an optimization process, namely, invariant risk minimization (IRM). IRM considers one of the most popular data generation assumptions, later known as the partially informative invariant feature (PIIF) assumption. Subsequently, numerous invariant learning works [Rosenfeld et al. 2020; Ahuja et al. 2021; Chen et al. 2022a; Lu et al. 2021c], endowed with causality, propose to solve distribution shifts formulated by various assumptions including fully informative invariant feature (FIIF) and anti-causal assumptions [Rosenfeld et al. 2020; Ahuja et al. 2021; Chen et al. 2022a], which makes these assumptions the popular basis of causally theoretical analyses for OOD problems.

10.2.2 OOD in AI for Quantum Mechanics.

In the domain of quantum mechanics, the OOD problem frequently emerges when determining the wavefunction of quantum systems [Yang et al. 2020; Kochkov et al. 2021a; Roth and MacDonald 2021; Fu et al. 2022c]. For example, as the sizes of quantum systems increase, the space needed to model the wavefunction grows exponentially, and the interactions between spins or particles

become more intricate. Additionally, different geometries of systems will also change the underlying physical interactions dramatically. It is challenging to apply a wavefunction ansatz to a larger lattice or a different molecule. Some works address OOD issues by better encoding intrinsic interaction modes that can be shared across system sizes and geometries. Botu and Ramprasad [2015] compare the fingerprints of new structures with those in the training dataset and mandate a fresh QM calculation if it is out of the predictable domain when one or more components of the structure's fingerprints lies outside the training range. QM-GNN [Guan et al. 2021] implements supplemental QM descriptors to facilitate the prediction of out-of-domain unseen examples. Caro et al. [2022] initiate a study of out-of-distribution generalization in Quantum Machine Learning (QML). They prove out-of-distribution generalization for the task of learning an unknown unitary, a fundamental primitive for a range of QML algorithms, with a broad class of training and test distributions, showing that one can learn the action of a unitary on entangled states having trained only product states.

10.2.3 OOD in AI for Density Functional Theory.

Within the realm of DFT, OOD situations commonly arise in the context of quantum tensor learning. For the task of quantum tensor prediction, current models are trained on systems with only tens of atoms due to computational complexity [Schütt et al. 2019; Unke et al. 2021a; Li et al. 2022e]. However, in practice, systems can contain hundreds even thousands of atoms. The size shift of quantum systems engenders prediction difficulties without a trivial solution. Several existing works put forward the problem of severe performance drop outside the defined applicability domain [Pereira et al. 2017; Li et al. 2016a], while few work offers feasible solution to address this issue. A general and realistic direction for future studies is to perform training using data of different sizes under the invariant risk minimization framework [Peters et al. 2016; Arjovsky et al. 2019] using size as the environment.

10.2.4 OOD in AI for Molecular Science.

The OOD challenge in molecular science arises from the vast and intricate chemical space that AI models must navigate, with many potential challenges stemming from data limitations, model architectures, and evaluation metrics [Gómez-Bombarelli et al. 2018; Chen et al. 2018b; Feinberg et al. 2018]. One primary challenge is the limited coverage of chemical space by training data, which can lead to biased predictions and model performance degradation on unseen molecules. This issue arises due to the immense size and complexity of the chemical space, with a virtually infinite number of potential compounds [Polishchuk et al. 2013]. For example, general GNNs are not capable of generalizing to large molecules when they are trained with small molecules. Motivated by pioneer causality-related invariant learning works [Arjovsky et al. 2019; Peters et al. 2016], Bevilacqua et al. [2021] propose to address the size shifts by introducing size-invariant graph representations. As many graph OOD learning methods [Wu et al. 2022b; Chen et al. 2022b; Gui et al. 2023] using subgraph-based graph modeling emerge recently, Yang et al. [2022] introduces a molecule-specific invariant learning method for drug discovery. In science fields, Sharifi-Noghabi et al. [2021] formulate the drug response prediction in cancers as an OOD problem and propose Velodrome under a semi-supervised setting. Besides OOD learning strategies, molecule OOD generation is also an emerging realistic direction. Current drug discovery experiments are expensive, and in-distribution generation will not provide innovative molecule structures. Therefore, OOD molecule generation is crucial for drug discovery. Recently, as the emergence of energy-based methods [Elflein 2023] and molecule generations [Liu et al. 2021d], Lee et al. [2022b] combine both energy-based generation and OOD detection to generate molecule out of the known molecular distribution. In addition, a score-based method molecular out-of-distribution diffusion (MOOD) [Lee et al. 2022a] is proposed

to generate novel and chemically meaningful molecule by utilizing gradients to guide the generation process to high property score regions.

10.2.5 OOD in AI for Protein Science.

OOD in AI for protein science is a critical research topic due to the immense diversity of protein structures and functions, as well as the continually evolving knowledge of protein sequence-structure-function relationships [Koehl and Levitt 2002; Petrey and Honig 2005]. The ability to generalize AI models for predicting protein structures, protein-protein interactions, or even protein-drug interactions beyond the training data distribution would accelerate progress in areas such as drug discovery, precision medicine [Ashley 2016], and protein engineering [Goldenzweig et al. 2016]. One primary challenge in this context is the limited availability of high-quality experimental data, since the lack of domains is critical for OOD generalization. Therefore, a potential solution is incorporating domain knowledge and physical principles into AI models [Jumper et al. 2021]. These approaches can help AI models learn more transferable and robust features that generalize better to novel protein sequences or complexes. ProGen [Madani et al. 2020] is an unsupervised protein sequence generation method by using language models which include non-trivial OOD performance evaluations. Gruver et al. [2021] find that ensemble models are more robust on OOD protein design than other methods. Kucera et al. [2022] propose an innovative protein sequence generation method with OOD generation evaluations. Finally, one possible direction is uncertainty estimation or OOD detection in protein science [Hamid and Friedberg 2018, 2019], which is underexplored.

10.2.6 OOD in AI for Material Science.

For material science, the OOD problem often arises due to the vast diversity of materials and their unique properties. Towards unseen OOD materials and compositions, the complexity of their structures, interactions, and properties present a significant challenge for AI-driven material discovery and optimization. Additionally, incorporating domain knowledge and physical principles into AI models can aid in learning more transferable and robust features that generalize better to novel materials and structures [Murdock et al. 2020]. Kailkhura et al. [2019] ensemble simple models and propose a transfer learning technique exploiting correlations among different material properties to reliably predict material properties from underrepresented and distributionally skewed data. Sutton et al. [2020] use subgroup discovery to determine domains of applicability of models within a materials class. Another aspect of material science is material design and discovery [Ghiringhelli et al. 2015; Xue et al. 2016a; Guo et al. 2019], which is influenced by the intricate nature of materials' structures and properties. Lastly, exploring OOD detection in material science [Musil et al. 2018], which remains largely unexplored, can be a promising future research direction.

10.2.7 OOD in AI for Chemical Interactions.

The OOD challenge is a critical issue in chemical interactions, particularly in the study of molecular interactions [Cai et al. 2022b,a], where models might struggle to generalize and provide accurate predictions when applied to new and unseen bindings. For instance, accurately predicting the docking efficacy of a drug candidate on a target protein that is significantly different from those in the training data is crucial for designing effective treatments. Recently, Zhang and Liu [2023] propose to consider protein-molecule interaction through subpocket-level similarities for drug generations, improving the model generalization ability. For drug-drug interactions (DDIs), Tang et al. [2023] devise a substructure interaction module, DSIL-DDI, to learn domain-invariant representations for DDI tasks, improving generalization ability and interpretability. To probe dark gene families, Cai et al. [2023] propose an innovative OOD meta-learning algorithm PortalCG to generalize from distinct gene families to dark gene family. Because of the challenge of the scarcity of receptor activity

data, Cai et al. [2022a] propose a self-supervised method DeepREAL to mitigate distribution shifts. To assess the OOD generalization ability of previous drug-target interaction works, Torrisi et al. [2022] provide a generalization ability evaluation by including systematic test sample separations.

10.2.8 OOD in AI for Partial Differential Equations.

In the field of neural PDE solvers, deriving training data from classical solvers can be prohibitively expensive. Therefore, a practically useful neural PDE solver should be able to generalize to different systems, including those with different initial conditions, boundary conditions, and PDE parameters. MAgNet [Boussif et al. 2022] enables zero-shot generalization to unseen meshes, solving PDEs at a different resolution from that seen during training. Brandstetter et al. [2022c] add noise during training to encourage stability and address the distribution shift problem. NCLaw [Ma et al. 2023] embeds a network architecture that strictly guarantees standard constitutive priors (including rotation equivariance and undeformed state equilibrium) inside a differentiable simulation and optimize based on the difference between the simulation and the motion observation. NCLaw can generalize to new geometries, initial/boundary conditions, temporal ranges, and even multi-physics systems after training on a single motion trajectory, achieving performance gains by orders-of-magnitude over previous neural network approaches on these typical OOD tasks. Other works [Kochkov et al. 2021b; Stachenfeld et al. 2021] study various OOD generalization abilities of learned models, including generalizing to conditions, rollout durations, and environment sizes outside the training distribution. Future works can incorporate prior physical knowledge into deep learning surrogate models to obey the underlying physical laws and capture invariant information, thereby improving generalization ability across different systems.

10.2.9 Datasets and Benchmarks.

To facilitate the development of OOD in scientific tasks, there have been prior benchmark works addressing the OOD problem in the scope of scientific tasks, providing schemes and evaluations for OOD learning on various real-world datasets. OGB [Hu et al. 2020] focuses on graph datasets, identifies and splits different distributions respecting multiple domains. Wilds [Koh et al. 2021; Sagawa et al. 2021] studies shifts on data collections from the wild covering multiple domains and data modalities. GOOD [Gui et al. 2022a] considers the completeness of distribution shifts and benchmarks diverse graph tasks with numerous datasets and methods. DrugOOD [Ji et al. 2022] and CardioTox [Han et al. 2021] focus on molecular graph OOD problems, and are curated based on a large-scale bioassay databases ChEMBL [Mendez et al. 2019], NCATS, and FDA [Siramshetty et al. 2020]. ImDrug [Li et al. 2022g] evaluates several drug discovery tasks for imbalanced learning. Further OOD studies for AI can benefit scientific tasks on the basis of these works.

10.2.10 Open Research Directions.

OOD scenarios are universal for AI in scientific fields, causing substantial deterioration in task performances; therefore, it is crucial to safeguard AI models in scientific domains from faltering in such situations to prevent adverse real-world consequences. We seek to underscore the significance of continued investigation and research into OOD strategies in the context of AI applications for scientific disciplines. For further studies, we point out that one promising direction is to identify and exploit causal factors [Peters et al. 2016] in the training data that can constrain the behavior of optimized models on unseen test data. The model can generalize to OOD if the nature of the target distribution shift is known a priori, for example, enabling generalization to OOD orientations with models built in $SE(3)$ equivariance.

10.3 Foundation and Large Language Models

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Supervised learning of deep models usually requires a large amount of labeled data. However, in the case of scientific discovery, obtaining labeled data can be especially challenging due to factors such as the need for expert domain knowledge, high computational or experimental costs, or physical limitations. For example, computing the energy of molecules using DFT methods can take hours to days per molecule, depending on its size. Additionally, experimentally obtaining positively labeled data for drug discovery is costly and time-consuming, making deep models less applicable for rapid drug discovery at the early stage of global pandemics such as COVID-19. This difficulty has led to an emerging research area focusing on self-supervised learning (SSL). SSL techniques enable deep models to leverage unlabeled data and learn realistic data priors, such as physical rules and symmetries, without relying on extensive labeled datasets. Based on SSL, foundation models push this idea of leveraging data with no task labels to an extreme by aiming to pretraining a single model over these data that is easy to adapt for all tasks [Bommasani et al. 2021]. It essentially allows knowledge to be transferred as pre-trained representations from a general, usually self-supervised, task to a wide range of specific tasks of interest with limited labeled data. Specifically, large language models (LLMs) are the most versatile and powerful foundation models so far thanks to the label-free and rich supervision contained in the text data. LLMs enable even more flexible knowledge capturing and transfer due to their strong knowledge acquisition and reasoning abilities in scientific domains, including Physics, Computer Science, Chemistry, Biology, Medical Science [Boiko et al. 2023; OpenAI 2023; Nori et al. 2023; Gupta et al. 2022], etc. One of the most exciting applications of LLMs in the sciences is generative modeling. While hallucination is a common problem for many LLM use-cases, it becomes a strength for discovering new drugs [Liu et al. 2021b], materials [Xie et al. 2023a], and research ideas [Wang et al. 2023b]. So far, SSL-powered foundation and large language models are among the most promising directions to address the challenges of label acquisition and enable AI applications to a broader range of scientific problems. In the following subsections, we discuss the current challenges, focuses, and progress of SSL techniques, single-modal foundation models, and LLMs in the domain of scientific discovery.

10.3.1 Self-Supervised Learning.

SSL aims to construct informative learning tasks by deriving labels from the data itself, based on the associations within it. According to Xie et al. [2023b], SSL methods can be broadly categorized into contrastive and predictive approaches, depending on whether paired data are required in the learning process. Specifically, contrastive approaches involve multiple data modalities or augmentations to obtain positive data pairs to be discriminated from randomly sampled negative pairs, whereas predictive approaches auto-generate easy-to-compute and informative labels from certain subsets of dimensions of the data as the learning targets. SSL has shown its effectiveness and necessity in various fields [Chen et al. 2020a; Devlin et al. 2019] in the paradigms of representation learning, pre-training, and auxiliary learning [Xie et al. 2023b].

SSL of Molecule and Protein Representations: In the context of AI for science, a majority of existing SSL work has focused on learning representation for molecules from their 2D graph formulations. In particular, general graph SSL work [You et al. 2020; Xie et al. 2022c] has considered molecule representation learning as an important use case. In contrast, other studies have developed SSL methods specifically for molecular graphs, which allows for the integration of domain knowledge such as functional groups (motifs) co-occurrence [Hu et al. 2019b; Rong et al. 2020; Li et al. 2021f], atom-bond associations [Rong et al. 2020], and reaction context [Wang et al. 2022e].

These approaches have proven to be effective in leveraging the topology of molecular graphs as indicated by the chemical bonds but may miss certain geometry information of higher significance for certain tasks such as quantum properties predictions. To further use the 3D geometry information of molecules, Liu et al. [2022e] and Stärk et al. [2022a] propose to construct SSL tasks for molecules based on trans-modal associations. Technically, these approaches learn to maximize the mutual information between representations of 2D and 3D modalities of a molecule so that the representations are informative for multiple downstream tasks. Moreover, the Noisy Nodes technique has been proposed as a predictive SSL method in both pre-training [Zaidi et al. 2022] and auxiliary learning [Godwin et al. 2022] paradigms for 3D molecules. Specifically, Noisy Nodes approaches provide self-supervision by corrupting the atom coordinates and training GNNs to estimate the injected noise, which is in line with the idea of denoising autoencoders [Vincent et al. 2008; Xie et al. 2020; Batson and Royer 2019]. The simple strategy is shown to be effective for 3D molecules and has been used in various following works [Luo et al. 2023b; Masters et al. 2022]. In addition to small molecules, there are also efforts on developing SSL approaches for proteins. Specifically, Yu et al. [2023a] use contrastive learning to train a model which can compare protein sequences against functional annotations, such as enzyme commission numbers, for functional understanding.

SSL of Neural PDE Solvers: SSL has also been used in training neural PDE solvers, where the cost of training data generated by expensive numerical methods is a primary limitation of supervised solvers. To reduce this cost, Raissi et al. [2019] propose the physics-informed neural network (PINN), which directly parameterizes the network as the PDE solution. The network is optimized with a self-supervised physics-informed loss derived using constraints on the solution specified by the PDE and was empirically validated by solving the Schrödinger equation in one spatial dimension. In a more challenging SSL setting, Raissi et al. [2020] infer the velocity and pressure field of a fluid flow using PINNs trained with constraints specified by the Navier-Stokes equations coupled with snapshots of the concentration of a scalar field such as dye advected by the flow. This application is particularly relevant in settings where pressure and velocity measurements are needed but only snapshots are accessible, such as biomedical analyses of blood flow to detect coronary stenoses [Raissi et al. 2020]. Additionally, neural solvers conceived in the supervised setting, such as DeepONet [Lu et al. 2019], have been extended to the SSL setting through the incorporation of a physics-informed loss [Wang et al. 2021b]. Unlike vanilla PINNs [Raissi et al. 2019], which are locked to one particular instance of the PDE, the physics-informed DeepONet proposed by Wang et al. [2021b] can generalize over a family of PDEs, e.g., over initial conditions, and even demonstrated successful performance in experiments on the OOD regime. Furthermore, Wang et al. [2021b] report the physics-informed DeepONet outperforms its supervised counterpart.

10.3.2 Single-Modal Foundation Models.

The success of SSL techniques has given rise to the development of foundation models in vision [Rombach et al. 2022; Kirillov et al. 2023; Li et al. 2022f; Wang et al. 2022f], language [Radford et al. 2018; Devlin et al. 2019], and medical [Moor et al. 2023] domains. Typically, foundation models are large-scale models pre-trained under self-supervision or generalizable supervision, allowing a wide range of downstream tasks to be performed in few-shot, zero-shot manners, with easy fine-tuning, or to be built upon learned embeddings. Similar to SSL techniques, they enable knowledge distillation and transfer from a large amount of unlabeled data to specific tasks with limited or even zero data. In this section, we focus on discussing foundation models that do not heavily rely on the natural language modality. Specifically, we explore the development of foundation models in the fields of protein and molecule analysis, where their versatility and potential impact are particularly evident.

Protein Discovery and Modeling: Foundation models have shown great potential in AI for Science to address various challenges related to protein discovery and analysis. AlphaFold [Jumper et al. 2021] and RoseTTAFold [Baek et al. 2021] are two foundation models that have made significant progress in predicting the geometry of protein folding. The trained models are then extended to perform more downstream tasks, including protein generation and protein-protein interaction (PPI). Specifically, RFdiffusion [Watson et al. 2022] fine-tunes RoseTTAFold to enable protein structure generation with a diffusion model. Instead of predicting structure from sequence, RFdiffusion performs unconditional generation from random noise, which can be further extended to conditional generation given certain functional motif or binding target. Similarly, Chroma [Ingraham et al. 2022] is developed as a foundation protein diffusion model to enable protein generations conditioned on desired properties, including substructures and symmetry, which facilitates multiple downstream applications such as therapeutic development. In addition, AlphaFold Multimer [Evans et al. 2021] and Humphreys et al. [2021] extend AlphaFold2 [Jumper et al. 2021] and RoseTTAFold [Baek et al. 2021], respectively, to perform prediction tasks of PPI, or protein complexes without further fine-tuning. In addition to modeling the protein structure and geometry, the language model has also shown to be effective in multiple tasks related to protein design [Madani et al. 2023; Melnyk et al. 2022; Zheng et al. 2023; Hie et al. 2023] even when only the sequential form of proteins is involved.

Molecule Analysis and Generation: For molecule-related tasks, while various self-supervised learning (SSL) techniques have been proposed, there is currently no dominant non-language-based foundation model in this field. However, two promising threads of research work have emerged, focusing on different modalities of molecules: the molecular graph, in terms of either the 2D structure or the 3D geometry, and the sequential representation in terms of SMILES [Weininger 1988; Weininger et al. 1989]. In the case of *2D molecular graphs*, researchers have extended the success of graph-based SSL studies [Wang et al. 2022k,e]. For example, Fifty et al. [2023] formulate molecules as graphs and pre-trains GNN models with a great amount of simulated data to predict the binding energies for interactions between molecules and protein targets in simulation. Compared to typical pre-training approaches, Fifty et al. [2023] demonstrate the potential of molecule foundation models in a wider range of downstream tasks, including few-shot docking and property predictions. Recent work also demonstrates the multi-tasking capability of foundation models built upon *3D molecular graphs* when encoded appropriately. Specifically, Flam-Shepherd and Aspuru-Guzik [2023] formulate 3D molecule-related tasks as the auto-regressive generation on the sequentialized 3D coordinates of atoms. This framework enables the use of language model architectures on multiple tasks, including molecule generation, material generation, and protein binding site prediction. On the other hand, existing work such as ChemGPT [Frey et al. 2022], ChemBERTa [Chithrananda et al. 2020; Ahmad et al. 2022], MolBert [Fabian et al. 2020], Schwaller et al. [2021], MegaMolBart [NVIDIA Corporation 2022], and Tysinger et al. [2023] focus on *string representations* of molecules and adapt pre-training techniques from language models to molecule representation learning from large collections of such strings. Language models are also shown to be capable of molecule generation by producing SMILES strings. However, since SMILES strings were not designed specifically for generative modeling, many generated SMILES strings are chemically invalid. New string representations [Grisoni 2023] for the generation purpose have been proposed, such as DeepSMILES [Krenn et al. 2020], which avoids ring and parenthesis closing issues, and SELFIES [Krenn et al. 2020], which proposes a formal grammar approach to ensure validity. SELFIES has been extended to incorporate groups [Cheng et al. 2023a] to better capture meaningful molecular motifs. These string-based studies are among the first attempts to explore the power of large language models and have shown great potential in various tasks. In spite of the

use of language models, these studies focus on the single modality of molecules and do not involve guidance and knowledge from natural language.

10.3.3 *Natural Language-Guided Scientific Discovery.*

Applying language models to the scientific domain is becoming increasingly popular due to its potential impact for accelerating scientific discovery [Hope et al. 2022]. A natural question is to ask why we want to integrate language into the scientific discovery process. Beyond the conspicuous and important task of extracting information from literature, there are a number of other compelling reasons. First, language enables scientists without computational expertise to leverage advances in AI. Second, language can enable high-level control over complex properties when designing novel artifacts (e.g., drug design going from low-level “logP” to high-level “antimalarial”). Third, language can serve as a “bridge” between modalities (e.g., cellular pathways and drugs) when data is scarce. Beyond these three reasons, language has been developed as *the* method by and for humans to abstractly reason about the world. In much the same way that science often relies on natural phenomenon (e.g., penicillin) for innovation, we can rely on linguistic phenomenon for abstraction and connection.

Traditionally, natural language processing (NLP) has been developed with a focus on core tasks including translation and sentiment analysis. In the scientific domain, NLP tasks have focused on extracting information from the literature, such as named entity recognition [Li et al. 2016d], entity linking [Lai et al. 2021a], relation extraction [Wei et al. 2016; Lai et al. 2021b], and event extraction [Zhang et al. 2021]. NLP models have advanced rapidly in recent years and hence resulted in strong foundational models which can be easily applied to most NLP tasks [Devlin et al. 2019; Raffel et al. 2020; Brown et al. 2020]. Further, these systems have, to some extent, commonsense [Bian et al. 2023] and reasoning [Wei et al. 2022; Yao et al. 2023a; Huang and Chang 2022] abilities which may further advance AI research for science. However, the variety and complexity of scientific text still pose challenges to these systems. Thus, considerable effort has gone into constructing domain-specific language model variants [Beltagy et al. 2019; Liu et al. 2021c; Michalopoulos et al. 2021; Gu et al. 2021; Meng et al. 2021; Yasunaga et al. 2022; Gupta et al. 2022; Luo et al. 2022; Taylor et al. 2022] to harness the valuable information contained therein. Building on these base models has powered a wide range of applications, ranging from large-scale information retrieval systems [Google 2004; Fricke 2018] to knowledge graph construction [Wang et al. 2020a; Zhang et al. 2021], and from analogical search engines for scientific creativity [Kang et al. 2022] to scientific paper generation [Wang et al. 2018, 2019a]. Although these applications are diverse, their common theme is the attempt to make sense of an overabundance of scientific information. Recently, work has further improved these scientific language models by introducing external knowledge from human-constructed databases into existing models [Lu et al. 2021a; Lai et al. 2023], applying distillation for data augmentation [Wang et al. 2023a], and augmenting models via retrieval [Naik et al. 2021; Zamani et al. 2022].

Current advances in LLMs for science generally focus on addressing two key challenges. First, science discovery problems usually involve complicated data modalities such as the geometric status of a particle system. It is hence crucial to develop effective approaches to encode and integrate scientific modalities with the language modality. Second, due to various task formulations and limited data and model availability, the adaptation of general-purposed LLMs to scientific domains is non-trivial in terms of the learning task formulations and paradigms. In this section, we discuss the frameworks and techniques of LLMs for science instantiated by existing works from the above two perspectives.

Multimodal Science with Language: To address the challenge of leveraging scientific data modalities, work has begun to investigate aligning natural language with modalities in the scientific

domain. While some cases explore different variations of the original modality, such as molecule string representations [Guo et al. 2021], significant interest has begun to grow in the integration of these naturally existing modalities and natural language for enabling control of the scientific discovery process. This is in large inspired by the success of models such as CLIP [Radford et al. 2021] (contrastive learning) and DALL-E [Ramesh et al. 2021] (joint sequence modeling) in the last two years. The high-level goal of integrating language with other modalities is to enable high-level function control (e.g., taste) rather than low-level property-specific control (e.g., solubility). The overall proposition is that similarly capable models in the scientific domain would vastly accelerate many aspects of the discovery process by enabling scientists to work with function rather than form in mind. Additionally, language is compositional by nature [Szabó 2020; Partee et al. 1984; Han et al. 2023], and therefore holds promise for composing these high-level properties [Liu et al. 2022d]. Such compositionality is shown in scientific tasks evaluated by Edwards et al. [2021, 2022]; Su et al. [2022]; Liu et al. [2022d].

Multimodal Science with Language — Determining Modalities: In order to determine the appropriate problem formulation and model design for a given application, it's first necessary to determine the relevant input and output modalities. For example, if one wants to extract reactions from the literature, a text-to-text model [Vaucher et al. 2020] should be sufficient. However, to develop a contextual understanding of the reactions we are extracting, we might additionally incorporate figures (vision) and molecular structures. In the case of drug molecule generation and editing with high-level instructions, incorporating language as an input would be appropriate [Edwards et al. 2022; Liu et al. 2023c; Fang et al. 2023]. In the case of retrieving relevant literature about a drug, we may choose a molecular graph as input which is used to retrieve from a corpus of papers. Generally, given sufficient data for training, adding modalities to language will likely be helpful simply by grounding the model's understanding into the real world. However, obtaining multimodal data can be challenging in practice. As a rule of thumb, one can ask themselves the following three questions for moving from single-modal solutions to multimodality: 1) is multimodality a core part of my task, such as molecule captioning? 2) Should I add language to Section 10.3.2 tasks? In other words, do I need the level of control and abstraction offered by natural language; or is there complementary information available as text? and 3) Will I meaningfully benefit from anything beyond language, or is all the information I need expressed as text?

Multimodal Science with Language — Integrating Modalities: Now that one has decided to pursue a multimodal approach to their application, it is crucial to develop a framework to integrate the modalities. There are two common approaches as shown in 37, namely, Bi-Encoder Models and Joint Representation Models. An analogy can be drawn here to the distinction between cross-encoders and bi-encoders in information retrieval [Reddy et al. 2023]; bi-encoders allow fast comparisons and can be less data-intensive to train, but cross-encoders allow more fine-grained interaction between modalities. We now discuss the two approaches in detail. *Bi-Encoder Models* consist of an encoder branch for text and a branch for the other modality such as molecules and proteins. They have the advantage of not requiring direct, early integration of the two modalities, allowing existing single-modal models to be integrated. Representative examples include Text2Mol [Edwards et al. 2021], which proposes a new task of retrieving molecules from natural language queries, and CLAMP [Seidl et al. 2023], which learns to compare molecules and textual descriptions of assays for drug activity prediction. BioTranslator [Xu et al. 2023b] takes this to the extreme by learning a latent representation between text, drugs, proteins, phenotypes, cellular pathways, and gene expressions. Generally, these bi-encoder models are effective for cross-modal retrieval [Edwards et al. 2021; Su et al. 2022; Liu et al. 2022d; Zhao et al. 2023], but they may also be integrated into molecule [Su et al. 2022; Liu et al. 2022d] and protein [Liu et al. 2023f] generation frameworks.

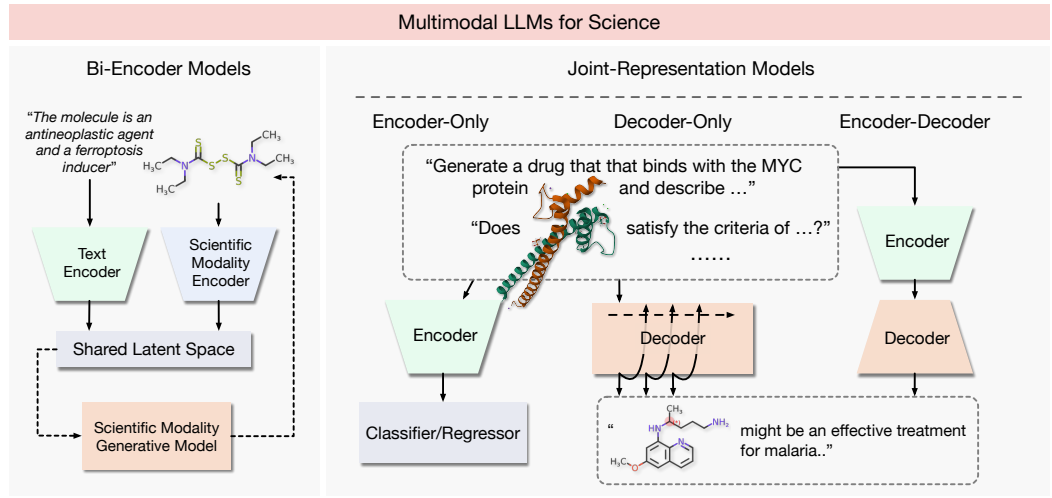


Fig. 37. High-level architectures for multimodal scientific NLP. Molecule-language multimodality is used as a source for examples. The dotted lines in the bi-encoder diagram indicate the possible extension to a generative framework. Example inputs and outputs are shown with “code-switched” modalities (*i.e.*, they are integrated into a single sequence). Encoders are used to generate an output representation which can be used for retrieval, classification, and regression, among other tasks. Decoder models are generally used for generative modeling applications. In some cases, non-LLM components can be used (such as for the decoder in an encoder-decoder model). We note that the extension of general-purpose LLMs via tools is another approach not shown in this figure.

Joint-Encoder Models, on the other hand, seeks to model interactions between multiple modalities inside the same network branch. These can be categorized by whether they incorporate a decoder or not. Encoder-only models can be used for prediction, regression, and (potentially slow) retrieval, but are unable to perform generative modeling. An example is KV-PLM [Zeng et al. 2022], which trains an encoder-only language model on literature data with molecule names replaced by SMILES strings. A second category is encoder-decoder [Edwards et al. 2022; Christofidellis et al. 2023] or decoder-only models [Liu et al. 2023e]. These can be used for cross-modal generative tasks, such as the “translation” between molecules and language proposed by Edwards et al. [2022], where molecules are generated to match a given textual description and vice versa. Interest has also arisen in using language to edit existing molecules for drug lead optimization [Liu et al. 2022d, 2023c]. Other work considers reaction sequences [Vaucher et al. 2020, 2021] or proteins [Gane et al. 2022].

Adapting LLMs to Science Domains: Existing LLMs have predominantly been studied and developed for general purposes. These LLMs can be leveraged and adapted to specific science domains or even particular tasks, capitalizing on their inherent knowledge and priors. When performing such an adaptation, it becomes crucial to meticulously design the formulation of the scientific task as a sequential generative task and incorporate essential domain-specific context and knowledge into the LLM, either during or prior to task inference. Moreover, given limited domain-specific data, one has to trade-off between the overwhelming irrelevant knowledge from general domain and the reasoning capability during the adaptation.

Adapting LLMs to Science Domains – Learning Task Formulations: LLMs are sequence-based models, so it is non-trivial to construct input and output sequences for different task formulations such as prediction, retrieval, and generation. As a general trend, however, most current language

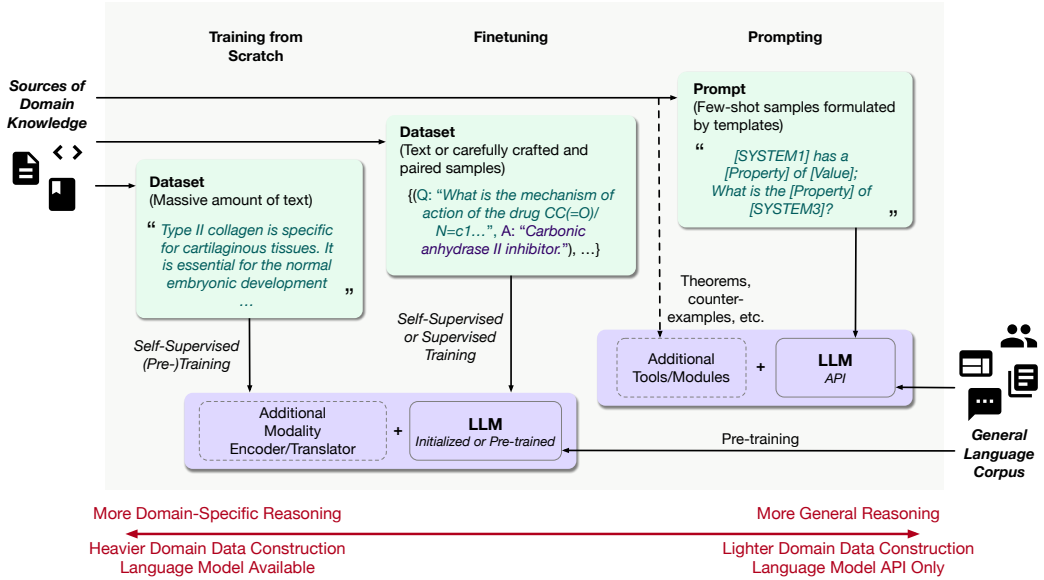


Fig. 38. Three paradigms of adapting LLMs to science domains. One can construct datasets consisting of massive amounts of text from science domains and train LLMs from scratch in a self-supervised manner. The trained model can be used directly or further fine-tuned for specific tasks. Alternatively, one can fine-tune a pre-trained general-purposed LLM with less amount of text data, in a self-supervised manner, or paired samples, in a supervised manner, from science domains. In cases of proprietary LLMs with API access, one can adapt the model by prompting with carefully designed templates, where domain knowledge are provided as few-shot samples in the prompt or as explicit knowledge with additional tools or modules. Dataset examples are from Galactica [Taylor et al. 2022] and ChEMBL [Liang et al. 2023], respectively.

models in scientific domains adapt pretraining procedures from core NLP such as BERT [Devlin et al. 2019] (e.g., KV-PLM [Zeng et al. 2022]) or GPT [Radford et al. 2018] (e.g., [Liu et al. 2023e]). As such, future work may find benefit in newer language model training objectives [Tay et al. 2023]. Some work, however, has attempted to use additional signals from known properties [Ahmad et al. 2022]. Additional work is often needed for designing multimodal learning formulations. Contrastive learning paradigms are widely applied in the case of bi-encoder models due to their multi-branch design. Learning tasks in joint-representation multimodal models are often designed to alleviate challenges with data scarcity. Strategies include multi-lingual [Edwards et al. 2022] and multi-task [Christofidellis et al. 2023] learning. In addition to training models for multimodal tasks, existing single-modal models can be integrated with multimodal extensions to avoid formulating a new learning objective. For example, bi-encoder models can be combined with flows [Su et al. 2022] or sequence generation models [Liu et al. 2022d, 2023c]. Further, classifier guidance can be used with an existing generative model [Ingraham et al. 2022].

Adapting LLMs to Science Domains - Learning Paradigms: Existing efforts have explored approaches to adapting LLMs for various specific applications. Taking into consideration varying levels of data availability and model accessibility, these adaptations can be achieved through several paradigms [Liu et al. 2023d; Wang et al. 2023c]. In the science domain, there are three typical adaptation paradigms, including training domain-specific LLMs from scratch, fine-tuning general-purposed LLMs, and few/zero-shot learning with prompting. Their learning frameworks, dataset construction, and model access are compared in Figure 38 and discussed in the following. LLM

architectures themselves usually contain certain prior about reasoning. Given an effective LLM architecture, it is natural to *train domain-specific LLMs from scratch* with highly customized data. Such an approach enables the highest flexibility with additional modules and blocks and helps learn better domain-specific knowledge given a fixed capacity of model. Once trained, those models can be further used in specific downstream tasks. However, to achieve a desired performance, a significant effort is to be made to construct the training dataset consisting of a large amount of text. For example, Galactica [Taylor et al. 2022] constructs a large scientific corpus with data collected from papers, code, knowledge base, etc. The LLM is then trained in a self-supervised manner [Devlin et al. 2019; Radford et al. 2018] on the collected domain-specific data, being able to tokenize math equations, SMILES, protein, and DNA strings. These approaches are capable of performing multiple downstream tasks such as drug discovery, repurposing, and interaction prediction. Taking advantage of the general reasoning capability of pre-trained language models, recent works also explore the *fine-tuning of pre-trained LLMs* with domain-specific datasets. BioMedLM [Bolton et al. 2022] and med-PALM [Singhal et al. 2023] are finetuned on biomedical domains from general LLMs GPT-2 [Radford et al. 2019] and PaLM [Chowdhery et al. 2022], respectively. They have shown promising performance on the medical question–answering tasks. Fine-tuning can also be performed with less amount of but paired data in the supervised fashion. For example, DrugChat [Liang et al. 2023] constructs an instruction-tuning dataset consisting of more than 143k manually crafted question-answer pairs and covering more than 10k drug compounds. The LLM is then trained together with a GNN module on the constructed dataset.

Due to the recent success and popularity achieved by the most advanced LLMs such as GPT-4 [OpenAI 2023], work has begun to adapt these general instruction-tuned models for the most challenging scientific discovery. As the advanced LLMs are mostly proprietary with API availability, their science domain adaptation is usually achieved by the paradigms of *few-shot or zero-shot learning*, also known as in-context learning, through prompting. In particular, domain knowledge can be provided as a context in the prompt in the form of theories, facts, or examples. This paradigm has demonstrated its effectiveness in subjects like Social Science [Zhong et al. 2023] and astronomy [Sotnikov and Chaikova 2023]. In the molecule domain, work has explored the chemical knowledge contained in these models in terms of language [Castro Nascimento and Pimentel 2023] and code generation [Hocky and White 2022; White et al. 2023]. Guo et al. [2023b] benchmark advanced LLMs on multiple tasks in the chemistry domain and demonstrate their competitive performance compared to task-specific machine learning approaches. Recent work, such as CancerGPT [Li et al. 2023d] and SynerGPT [Edwards et al. 2023], also explores the applications of language models for drug synergy prediction. SynerGPT proposes novel LLM training strategies for in-context learning to explore the higher-level “interactome” between drug molecules in a cell. They extend their model to inverse drug design and context optimization for standardized assays. The proposed training strategies may enable a new type of foundation model based on the drug interactome. Further, one particular promising route is augmenting LLMs with external tools such that even complex tasks become textual [Schick et al. 2023; Yao et al. 2023b], with scientific examples like using the Web APIs of the National Center for Biotechnology Information (NCBI) for answering genomics questions [Jin et al. 2023]. Existing LLMs are pretrained only from unstructured texts and fail to capture some domain knowledge. Recent solutions for domain knowledge-empowered LLMs include developing lightweight adapter framework to select and integrate structured domain knowledge to augment LLMs [Lai et al. 2023] and data augmentation for knowledge distillation from LLMs in the general domain to chemical domain [Wang et al. 2023a]. External domain tools can also be integrated into language model prompts to allow these “agents” to access external domain knowledge [Bran et al. 2023; Boiko et al. 2023; Liu et al. 2023c]. Specifically, ChatDrug [Liu et al. 2023c] enables LLM-powered drug editing in a few-shot manner by equipping LLMs with a

retrieval and domain feedback module. Work is also done in the few-shot setting for both regression and classification [Jablonka et al. 2023], as well as Bayesian optimization [Ramos et al. 2023].

Pushing this paradigm to the extreme, there are also emergent efforts on developing LLM-based agents for scientific discovery by connecting LLMs with tools for conducting experiments, such as in Chemistry [Boiko et al. 2023] and Machine Learning [Zhang et al. 2023e]. However, different science domains often rely on data in very different forms of modalities in practice, making it challenging for LLMs to be directly useful in many applications.

10.3.4 Open Research Directions.

There are still remaining challenges towards scientific discoveries with foundation and large language models. We identify and discuss the following three challenges and opportunities.

Data Acquisition for Foundation Models: Large-scale data acquisition presents a significant challenge when developing SSL and foundation models for scientific applications, mainly due to the specialized nature of scientific data compared to general internet data with no easy way around it. There are existing efforts such as collecting domain-specific text from the internet, image-caption pairs from PubMedCentral’s OpenAccess subset [Lin et al. 2023e], and scientific figures with captions from arXiv [Hsu et al. 2021]. However, most of these efforts focus on the image and text modalities and largely overlap with web data. More work is needed on curating more realistic scientific data with diverse modalities, such as sensory and tabular data, to support building more customized foundation models for science. Data scarcity is a key challenge for language-based scientific multimodality, such as models trained on molecule-text pairs. Existing work has attempted to alleviate this challenge by applying multilingual pretraining strategies [Edwards et al. 2022] or by using entity linking to extract large quantities of noisy molecule-sentence pairs from the literature [Zeng et al. 2022; Su et al. 2022]. However, improved extraction of less noisy and more complete data from the literature will greatly benefit these tasks. [Yang et al. 2023b] investigates the use of language models for extracting additional drug synergy training tuples from literature.

Addressing Algorithmic Challenges for SSL and Foundation Models: Aside from data, the main technical challenges for SSL and foundation models for science typically include incorporating diverse modalities in the architecture, designing customized pretraining techniques for these modalities, and addressing domain distribution shifts. Recent methods have mainly focused on combining text and image modalities [Liu et al. 2023b; Koh et al. 2023; Alayrac et al. 2022] and more well-studied scientific units like molecules and proteins, with limited recent SSL/foundation models works on other modalities like graph [Huang et al. 2023b], RNA expression [Rosen et al. 2023] and benchmark on even more rare modalities like bacterial genomics and particle physics [Tamkin et al. 2021]. Finally, the dynamic data change in the realistic scientific discovery process also forces the pretrained models to face domain distribution shifts, as exemplified in the Wilds benchmark [Koh et al. 2021]. More foundational work on designing robust SSL algorithms for diverse modalities is needed for applying AI for science in practice. For LLMs, since the knowledge from general domain is often overwhelming, developing better fusion models beyond perceivers can be a promising future direction.

Extending the Success to Broaden AIRS Topics: Self-supervised learning (SSL) and foundation models have demonstrated promising performance in domains such as small molecules, proteins, and continuum mechanics. However, their methodologies and applications in other areas have received less attention. For example, SSL has been relatively less explored in the context of quantum systems. Learning tasks in quantum systems often revolve around modeling wavefunctions, and the neural network architectures used tend to be specific to lattice structures, making knowledge transfer between systems or tasks challenging. Nonetheless, SSL holds significant potential in

these fields as unlabeled data distributions can contain valuable information about the underlying symmetry and physical rules. SSL can play a crucial role in learning these rules as a prior, thereby facilitating the discovery of fundamental principles across various systems. Furthermore, adapting foundation models presents another promising avenue for the discovery of emerging and less-explored domains with limited data. Particularly, as demonstrated by [Taylor et al. \[2022\]](#); [Xu et al. \[2023b\]](#), the text-based nature of LLMs enables them to capture and transfer knowledge among different systems more effectively and flexibly, bridging the gap between different domains and data modalities.

10.4 Uncertainty Quantification

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The capability of profiling and predicting properties of complex systems involved in previously discussed AI for Science tasks enables optimal and robust decision making for scientific discovery as well as automated generative capabilities. While developing deep forward prediction and generative models for inverse design under different conditions may have made significant advancements, reliable uncertainty quantification (UQ) in these physics constrained prediction and generative models, such as neural ODEs [Chen et al. 2019a] as well as DeepONet [Raissi et al. 2020], is critical to guarantee robust decision making under data and model uncertainty, however still requires investigation via collaboration of applied mathematics, computational science, and AI/ML researchers. Different UQ strategies have been developed in these research communities, from classical Bayesian model sensitivity analysis focusing on subsets of model parameters to the recent ensemble-based UQ in deep Bayesian learning. When integrating these UQ strategies into forward predictive and inverse generative models, scalability and efficiency are the utmost important factors to enable time-sensitive prediction and decision making in practice. Efficient and reliable approximate Bayesian computation and variational inference methods are to be developed to achieve desired performances of both predictive and computational criteria.

10.4.1 Uncertainty Quantification: Introduction and Background.

Uncertainty quantification (UQ) has been studied in various disciplines of applied mathematics, computational and information sciences, including scientific computation, statistic modeling, and more recently, machine learning. Traditional UQ aims at either quantitatively assessing prediction uncertainty or calibrating parameters of traditional physics-principled mechanistic models and data-driving machine learning models to address challenges of modeling complex systems due to enormous system complexity and data uncertainty [Kennedy and O'Hagan 2001; Psaros et al. 2023]. When modeling complex systems, the uncertainties of a computational model can be from multiple sources. First, dynamics of real-world complex systems are typically modulated by many potential internal and external factors. Abstract computational modeling often can not cover all these factors, due to either missing information or computation limitations. Some factors affecting the system outcomes may be unknown or ignored for model construction. Second, even if all of the influencing factors are included, due to lack of knowledge, especially for data-driven black-box machine learning models, the selected model itself can be mis-specified with potential inductive bias. Third, the systems dynamics itself to be modeled can be intrinsically stochastic and non-stationary. Fourth, significant data uncertainty has to be taken care of as the observed data themselves are inevitably noisy and even corrupted due to the inherent sensor noise or the random perturbations from uncontrollable environmental factors. Finally, due to the limited precision of the modern digital computer hardware, the numerical results from different models may still contain errors. All these above uncertain sources contribute to the uncertainty of the final system output or model prediction.

Aleatoric and Epistemic Uncertainty: Two types of uncertainties that have been identified and extensively investigated in computational modeling are the *aleatoric uncertainty* and *epistemic uncertainty* [Kendall and Gal 2017; Hüllermeier and Waegeman 2021]. Aleatoric uncertainty, also known as (a.k.a.) *stochastic uncertainty* or *data uncertainty*, refers to the uncertainty due to the intrinsic randomness of the physical process under investigation. For example, in a quantum spinning system, even if the quantum state of the system is known, the measurements with respect to the computational basis are typically random. In materials science experiments, since the noise of the

sensor measurements can hardly be removed completely, the experimental results under the same condition may differ with some degree. In molecular property prediction, the predicted molecular properties can have significant uncertainty if only the 2D structure information is provided due to the incomplete representation considering the actual 3D molecular geometry [Hirschfeld et al. 2020]. These uncertainties are irreducible even if more knowledge of the complex system or supplementary data become available. Epistemic uncertainty, a.k.a. *systematic uncertainty* or *model uncertainty*, represents the uncertainty due to the lack of knowledge of its physical process dynamics when modeling a complex system. Epistemic uncertainty can be reduced or removed as more and more knowledge or data becomes available, for which many Bayesian learning [Cohn et al. 1996; Lampinen and Vehtari 2001; Titterington 2004; Xue et al. 2016b; Qian and Dougherty 2016; Gal et al. 2017b; Goan and Fookes 2020; Boluki et al. 2020], UQ [Yoon et al. 2013; Lakshminarayanan et al. 2017; Huang et al. 2017; Sensoy et al. 2018; Ardywibowo et al. 2019; Amini et al. 2020; Wang et al. 2022b], experimental design methods [Kushner 1964; Mockus 2012; Mariet et al. 2020; Zhao et al. 2020; Lei et al. 2021; Griffiths 2023] have been developed as effective and efficient solution strategies.

Importance of Uncertainty Quantification: The uncertainty quantification problem is of great importance in various disciplines for complex system modeling and scientific discovery. Knowing the uncertainty associated with a certain prediction will help us develop more reliable models and making better decisions, especially for some safety-critical applications [McAllister 2017]. As some modern machine learning models such as deep neural networks have great approximation capacity and expressiveness, the aleatoric uncertainty needs to be taken great care to avoid over-fitting. Moreover, online machine learning strategies, such as *Bayesian active learning* [Cohn et al. 1996; Gal et al. 2017b; Zhao et al. 2021a,c,b] and *Bayesian optimization* [Kushner 1964; Mockus 2012], can be combined with the inverse uncertainty quantification to facilitate new material and compound discovery [Solomou et al. 2018; Talapatra et al. 2018, 2019; Lei et al. 2021].

10.4.2 Uncertainty Quantification in Computational Science.

Forward and Inverse Uncertainty Quantification: In computational science, the quantification of uncertainty is typically categorised into *forward uncertainty propagation* and *inverse uncertainty quantification*. The objective of *forward uncertainty propagation*, a.k.a. *sensitivity analysis* [Razavi et al. 2021; Rochman et al. 2014; Peherstorfer et al. 2018], is to measure how much the randomness of a certain input will result in the uncertainty of system output. By modeling input factors or model parameters as random variables with corresponding probability distributions, the randomness or uncertainty of the system output can be captured by forward uncertainty propagation. In many cases when the computational models are too complex such that the output random variable do not have the closed-form probability distribution, the forward uncertainty is often estimated by *Monte Carlo* (MC) sampling. Other forward UQ methods to alleviate the high computational cost of MC sampling include *Taylor approximation* [Fornasini 2008] and other *surrogate modeling* strategies [Box and Draper 2007], which have been extended to UQ in deep learning as discussed in the next section.

On the other hand, the *inverse uncertainty quantification*, a.k.a. *model calibration* [Malinverno and Briggs 2004; Nagel and Sudret 2016; Nagel 2019], aims at measuring how uncertain we are about the corresponding parameters of the system model or input factors that modulate the underlying physical process of the system, and then further reducing the relevant uncertainties.

One powerful method to solve the inverse UQ problem is Bayesian modeling. Compared to the frequentist approaches modeling parameters as deterministic variables and derive point estimates that best fit the selected model with the observed data, Bayesian approaches consider model

parameters as random variables and solve the Bayesian inverse problem to update the corresponding probabilities to derive predictive posterior belief of a certain outcome following the Bayes' theorem. As a simple illustration, assume that we want to quantify the uncertainty of the system input X , and let Y denote the corresponding system output, which can be noisy. Bayes' theorem states

$$p(X|Y) = \frac{p(Y|X)p(X)}{p(Y)}, \quad (114)$$

where $P(X)$ is the *prior distribution* representing our prior belief of X without any observation Y , $P(Y|X)$ is the *likelihood*, the probability distribution of system output being Y given X based on the adopted model assumptions. The denominator $P(Y)$ is often called the *evidence*, which is the marginal distribution $P(Y) = \int P(Y|X)p(X)dX$ over the randomness of Y . The *posterior distribution* $p(X|Y)$ captures our updated belief of X after observing the system output Y . The same idea can be applied to quantify the uncertainty of system parameters. We can quantify the inverse uncertainty by Bayesian inference based on Bayes's theorem to derive the probability distributions of the corresponding system input or model parameters.

Other Notions of Uncertainty: Although the Bayesian uncertainty has long been the primary notion of uncertainty in various applications for its simplicity and soundness in both applied mathematics and probability theory, there are also many other notions of uncertainty other than Bayesian uncertainty. Those includes other methods with probabilistic predictions [Nix and Weigend 1994], making *interval predictions* [Koenker 2005; Angelopoulos and Bates 2021], assigning each prediction with a *confidence score* [Jumper et al. 2021], as well as *distance-based uncertainty* [Sheridan et al. 2004; Liu and Wallqvist 2018; Hirschfeld et al. 2020]. More recently, a variant of Bayesian uncertainty quantification methods called uncertainty quantification of the 4th kind (UQ4K) [Bajgiran et al. 2022] has been proposed to alleviate “brittleness of Bayesian inference”, which is a phenomenon that Bayesian inference could be sensitive to the choice of prior [Owhadi et al. 2015]. In UQ4K, the authors have developed UQ in the game theory framework. Via a min-max game on the risk between the estimation of model parameters and the prior distribution, the authors promote a *hypothesis testing notion of uncertainty*, which gets rid of the choice of prior and does not suffer from the “Bayesian brittleness”. While those UQ methods are less explored compared to the Bayesian UQ approaches, they can be useful for certain applications with corresponding advantages over Bayesian UQ, for example, lower computational cost, better scalability, and solution properties with theoretical guarantee.

10.4.3 Uncertainty Quantification in Deep Learning.

In machine learning, most of the existing UQ methods are based on Bayesian statistics and probability theory. One specific example is *Bayesian linear regression* [Box and Tiao 2011], which is the corresponding Bayesian adaptation of *linear regression*. Bayesian linear regression similarly considers a linear parametrized model from the input \mathbf{x} to the output y with observation noise ϵ , but with model parameters \mathbf{w} treated as a random vector with Bayesian inference to update the corresponding posterior rather than solving for deterministic point estimates. With training data $\{\mathbf{x}, y\}_{n=1}^N$, the posterior update rule gives:

$$p(\mathbf{w}|\{\mathbf{x}, y\}_{n=1}^N) = \frac{p(\{y\}_{n=1}^N|\{\mathbf{x}\}_{n=1}^N, \mathbf{w})p(\mathbf{w})}{p(\{y\}_{n=1}^N|\{\mathbf{x}\}_{n=1}^N)}. \quad (115)$$

The uncertainty is well-preserved in the posterior distribution of \mathbf{w} , and can be quantified further for the posterior predictive distribution on \hat{y} given a new test point $\hat{\mathbf{x}}$. With \mathbf{x} projected into the kernel space, we can further extend to another popular Bayesian learning method: *Gaussian process regression* (GPR) [Rasmussen and Williams 2006], as the Bayesian adaptation of *kernel ridge*

regression. Although many machine learning models have been implicitly derived in the traditional frequentist fashion, most of them can be adapted into the Bayesian counterparts with the UQ capability.

There are also other non-Bayesian UQ methods in machine learning, including *conformal prediction* [Angelopoulos and Bates 2021] and *quantile regression* [Koenker 2005], with the objective to predict an interval of outcome predictions instead of either deriving point estimates or updating distributions.

UQ with Bayesian Neural Networks: *Bayesian neural networks* (BNNs) [Lampinen and Vehtari 2001; Titterington 2004; Goan and Fookes 2020] have been proposed as Bayesian counterparts of frequentist training of artificial neural networks (ANNs), by modelling the network parameters or activation as random variables. We here use a regression problem to illustrate the main idea, and assume that the model parameters θ are modeled as random variables.

Given a data point x and f_θ , which is a prespecified neural network architecture f with the model parameters θ , the *aleatoric uncertainty* is typically modeled as the y being the output of neural network $f_\theta(x)$ with a random noise ϵ added: $y = f_\theta(x) + \epsilon$, which implicitly specified a distribution $p(y|\theta, x, f)$ and further $p(\{y\}_{n=1}^N|\theta, \{x\}_{n=1}^N, f)$ under some independence assumption. The noise ϵ is typically modeled as a zero-mean Gaussian random variable with the noise being a hyperparameter, or data-dependent through a neural network, which is also termed as *mean variance estimation* (MVE) [Nix and Weigend 1994].

On the other hand, the *epistemic uncertainty* is the uncertainty of model parameters θ given the limited training data $\{x, y\}_{n=1}^N$. According to the Bayes' theorem, the posterior distribution of the model parameters θ can be derived as:

$$p(\theta|\{x, y\}_{n=1}^N, f) = \frac{p(\{y\}_{n=1}^N|\theta, \{x\}_{n=1}^N, f)p(\theta|f)}{\int p(\{y\}_{n=1}^N|\theta, \{x\}_{n=1}^N, f)p(\theta|f)d\theta}. \quad (116)$$

With the posterior distribution of model parameters $p(\theta|\{x, y\}_{n=1}^N, f)$, we can quantify the uncertainty of the model parameters θ . Given a new test data point \hat{x} , the model prediction is the marginal distribution of \hat{y} :

$$p(\hat{y}|\{x, y\}_{n=1}^N, \hat{x}, f) = \int p(\hat{y}|\theta, \hat{x}, f)p(\theta|\{x, y\}_{n=1}^N, f)d\theta. \quad (117)$$

The expectation $\mathbb{E}[\hat{y}|\{x, y\}_{n=1}^N, \hat{x}, f]$ can be used as the point prediction of \hat{y} . The total uncertainty of the forward prediction is the uncertainty of the marginal distribution $p(\hat{y}|\{x, y\}_{n=1}^N, \hat{x}, f)$. Different metrics can be used to quantify this uncertainty, such as the variance or the (differentiable) entropy. However, the denominator $\int p(\{y\}_{n=1}^N|\theta, \{x\}_{n=1}^N, f)p(\theta|f)d\theta$ is often intractable, which makes the computation of $p(\theta|\{x, y\}_{n=1}^N, f)$ difficult. Various inference methods based on either *Markov chain Monte Carlo* (MCMC) sampling with corresponding gradient-based variants [Welling and Teh 2011] or *variational inference* [Blei et al. 2017] have been developed to address the challenge. We refer the readers to Jospin et al. [2022] for a comprehensive review of different approximate inference methods.

Scalable Approximate Inference for Deep Neural Networks: Bayesian inference becomes more challenging with millions of model parameters in modern deep neural network (DNN) architectures. Many recent research efforts aim at scaling up the approximate inference algorithms, especially for efficient Bayesian inference with DNN architectures. Some commonly adopted approximation UQ methods include *Bayes-by-backprop* (BBB) [Blundell et al. 2015], *Monte Carlo dropout* (MC dropout) [Gal and Ghahramani 2016; Gal et al. 2017a], and *deep ensemble* (ensemble) [Lakshminarayanan et al. 2017; Huang et al. 2017]. BBB [Blundell et al. 2015] is an optimization trick which

can be coupled seamlessly with the backpropagation algorithm and the autodifferentiation training when the neural network parameters are modeled as random variables with reparameterizable variational distributions. MC dropout [Gal and Ghahramani 2016; Gal et al. 2017a] is a simple and efficient way to provide uncertainty estimation on any model trained with dropout [Srivastava et al. 2014], a regularization technique shutting down selected neurons randomly. By performing dropout at both training and testing time, MC dropout has been proven to be able to provide an approximation inference on random neural network weights. Deep ensemble [Lakshminarayanan et al. 2017; Huang et al. 2017] is another heuristic strategy to achieve effective predictive uncertainty by combining the models in different local optima from random initializations or multiple “snapshots” during training. Deep ensemble and its variants have been shown to have an interpretation in Bayesian perspective [Pearce et al. 2020]. Some other heuristic approaches to further scale up BNN inference have been developed by combining BNN with frequentist neural network training. For example, *Bayesian last layer* (BLL) [Brosse et al. 2020] has shown to be surprisingly effective for its simplicity by modelling only the last layer network parameters as random variables, which significantly reduces the number of uncertain parameters by fixing all the other parameters as point estimates except those in the final layer, under the premise that earlier layers are performing feature extraction and later layers are performing the final prediction task. These approximate inference methods can either be used alone or combined together to achieve better uncertainty estimation with improved scalability and computational efficiency.

Evidential Deep Learning: *Evidential deep learning* (EDL) [Sensoy et al. 2018; Amini et al. 2020] is a recent emerging UQ strategy for deep learning based on *Theory of Evidence*, a generalized Bayesian formulation. EDL explicitly considers the uncertainty due to the lack of evidence, which is the amount of support from data for certain prediction. The network output of a test data without the support of evidence is encouraged to be a predefined prior other than a confident prediction. To model the epistemic uncertainty, instead of considering the model parameters θ to be random, EDL alternatively introduces a random vector π , with the predictive distribution now becoming

$$p(\hat{y}|\theta, \hat{x}, f) = \int p(\hat{y}|\pi)p(\pi|\theta, \hat{x}, f)d\pi, \quad (118)$$

where $p(\hat{y}|\pi)$ is typically a categorical distribution for classification and Gaussian for regression, similar as most of the existing DNN models. The prior distribution of random vector π is typically a conjugate prior for $p(\hat{y}|\pi)$ and the neural network models the likelihood of $\pi : p(\hat{x}|\pi, \theta, f)$. The training procedure is similar as the usual neural network training except for an extra penalty term that increases as the posterior of π becomes far away from the prior distribution. Some of EDL models [Sensoy et al. 2018] can also be interpreted as a special case of BNN by performing amortized variational inference on the last layer activation. The EDL framework has been shown to be especially useful for out-of-distribution data detection in various classification [Sensoy et al. 2018; Stadler et al. 2021] and regression tasks [Amini et al. 2020]. We give a schematic illustration of EDL along with some other UQ methods for DNN including BNN and MVE in Figure 39.

Uncertainty Quantification for Graph Learning: When modeling dependency across different system components, graph neural networks (GNNs) have been developed with successes in diverse applications, including materials science, molecular biology, and quantum mechanics as detailed in previous sections. Existing UQ methods developed specifically for graph learning includes Graph DropConnect (GDC) [Hasanzadeh et al. 2020], Bayesian graph convolutional neural network (BGCNN) [Zhang et al. 2019b; Pal et al. 2020], variational inference for graph convolution networks (VGCN) [Hasanzadeh et al. 2019; Hajiramezanali et al. 2019; Elinas et al. 2020], graph posterior network (GPN) [Stadler et al. 2021], gaussian process with graph convolutional

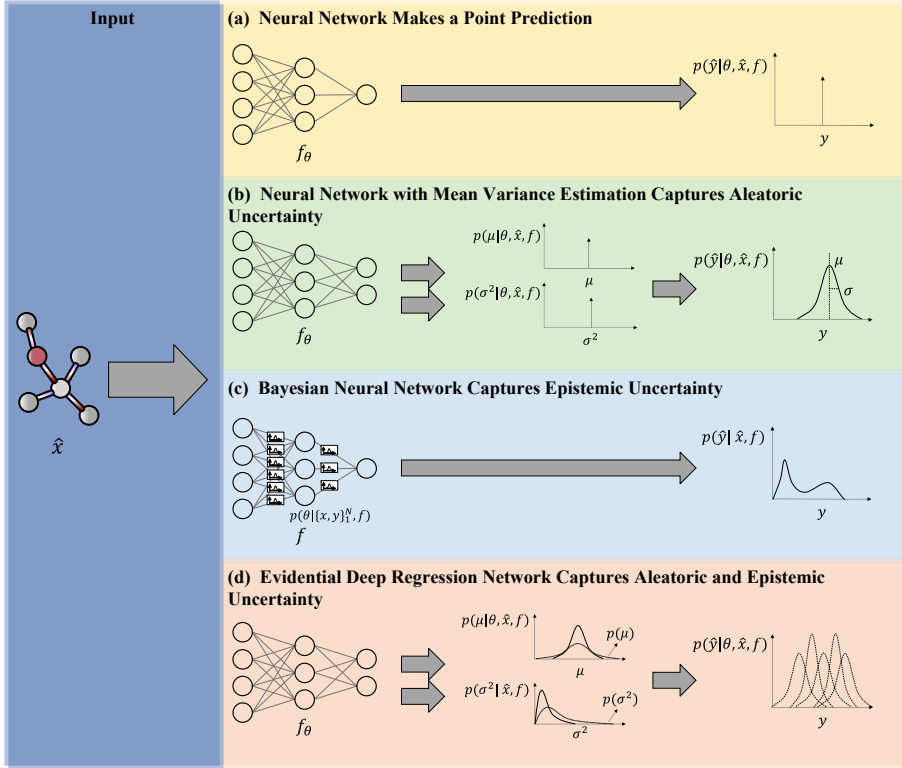


Fig. 39. Schematic illustration of different uncertainty quantification methods on a molecular energy prediction task, with \hat{y} denoting the predicted energy of a given molecule. Note that (c) Bayesian Neural Network can be further combined with Mean Variance Estimation to capture both aleatoric and epistemic uncertainty.

kernel (GPGC) [Fang et al. 2021], and conformalized GNN (CF-GNN) [Huang et al. 2023a]. Being a generalization of MC Dropout for GCNN with the dropout rates as learnable parameters, GDC [Hasanzadeh et al. 2020] provides UQ capability by modeling the neural network weights as Bernoulli random variables. BGCNN [Zhang et al. 2019b; Pal et al. 2020] further considers the topology uncertainty in graph structure by modeling the observed graph as a noisy observation of the true node relationship, and posterior inference on true node relationship is performed using the mixed membership stochastic block model (MMSBM). VGNC [Elinas et al. 2020] similarly models the structure uncertainty by performing variational inference on graph adjacency matrix and show this treatment can improve the model robustness to adversarial perturbation on graph structure. GPN [Stadler et al. 2021] is a UQ method for node classification task based on EDL, which is shown to be effective in OOD node detection. GPGC [Fang et al. 2021] is a graph Gaussian process model [Venkitaraman et al. 2020] with the kernel function defined over a deep GCNN, whose parameters are learned through the variational inducing point [Titsias 2009]. CF-GNN [Huang et al. 2023a] extends conformal prediction for the classification and regression tasks under an independence assumption to the graph-based models with theoretical analysis of validity conditions. The authors have shown satisfactory UQ performances with predicted intervals by CF-GNN covering the ground truth on various datasets.

Table 33. Existing research and benchmark studies on UQ for PDE surrogate solutions, DFT-related tasks, molecular property prediction, and compound-protein binding prediction.

Application	Method	Adopted UQ Approach(es)
Molecular Property	Ryu et al. [2019]	MVE, MC Dropout
	Zhang et al. [2019a]	BNN
	Scalia et al. [2020]	MC Dropout, Ensemble
	Hirschfeld et al. [2020]	MVE, Ensemble, GP, MC Dropout, RF [Ho 1995]
	Tran et al. [2020]	MVE, BNN, MC Dropout, Ensemble, GP
	Hie et al. [2020]	Ensemble, GP, BNN
	Soleimany et al. [2021]	EDL
	Yang and Li [2023]	MVE, Ensemble
	Greenman et al. [2023]	MVE, MC Dropout, Ensemble, GP, EDL
	Griffiths et al. [2022]	GP
Binding Affinity	Griffiths [2023]	GP
	Li et al. [2023b]	MVE, Ensemble, MC Dropout, BNN, Focal Loss [Lin et al. 2017; Mukhoti et al. 2020], SWAG [Maddox et al. 2019]
DFT	Wollschläger et al. [2023]	GP
	Hirschfeld et al. [2020]	GP
	Fowler et al. [2019]	MVE, Ensemble
PDE	Mahmoud et al. [2020]	GP
	Psaros et al. [2023]	GP, MCMC, Ensemble, MC Dropout, BNN, FP

Although all these previous efforts consider different uncertainty in graph learning, few of them actually quantify and analyze the estimated uncertainty. The appropriate treatment of uncertainty in graph learning is still an under-explored area despite recent success of GNNs on various graph learning tasks in material and protein property prediction.

10.4.4 Uncertainty Quantification in AI for Science.

Compared to other machine learning applications in computer vision and natural language processing, the problem of training under data scarcity is even more severe for scientific AI as the experiments typically being expensive and time-consuming to be conducted to collect meaningful data. Similar as any existing data-driven black-box model, the DNN-based scientific AI models can make erroneous but overconfident prediction for unseen input data [Hein et al. 2019], and are particularly vulnerable to adversarial attacks [Goodfellow et al. 2014b]. Therefore, there has been a growing interest in equipping those scientific AI models with the UQ capability. Many different benchmark studies and research efforts have been made to test the idea of using Gaussian process (GP) [Hie et al. 2020; Mahmoud et al. 2020; Hirschfeld et al. 2020; Tran et al. 2020; Griffiths et al. 2022; Griffiths 2023; Wollschläger et al. 2023; Li et al. 2023b; Psaros et al. 2023; Greenman et al. 2023], deep ensemble [Fowler et al. 2019; Scalia et al. 2020; Hirschfeld et al. 2020; Tran et al. 2020; Yang and Li 2023; Mariet et al. 2020; Hie et al. 2020; Greenman et al. 2023; Li et al. 2023b], MC dropout [Ryu et al. 2019; Zhang et al. 2019a; Scalia et al. 2020; Hirschfeld et al. 2020; Tran et al. 2020; Greenman et al. 2023; Li et al. 2023b], Bayesian neural network [Zhang et al. 2019a; Tran et al. 2020; Hie et al. 2020; Li et al. 2023b] and EDL [Soleimany et al. 2021; Greenman et al.

2023] to quantify the uncertainty of molecular property prediction [Ryu et al. 2019; Zhang et al. 2019a; Scalia et al. 2020; Hirschfeld et al. 2020; Tran et al. 2020; Hie et al. 2020; Soleimany et al. 2021; Griffiths et al. 2022; Yang and Li 2023; Greenman et al. 2023; Griffiths 2023; Li et al. 2023b; ?], compound-protein binding prediction [Hirschfeld et al. 2020], ground-state density prediction tasks [Fowler et al. 2019; Mahmoud et al. 2020], as well as PDE surrogate prediction tasks with physics-informed neural network (PINN) [Psaros et al. 2023], along with seismic inversion [Smith et al. 2020, 2022]. We summarize existing UQ research and benchmark studies applied to the above disciplines with the adopted UQ approaches in Table 33.

In particular, for chemical and protein molecular property prediction tasks, Ryu et al. [2019] have shown that MVE and MC Dropout based UQ methods can be used to assess the data quality. In Zhang et al. [2019a], Stein variational gradient descent (SVGD) [Liu and Wang 2016] inference algorithm has been implemented for BNN training to model the epistemic uncertainty and show that such an implementation can be used to mitigate the potential dataset bias and integrated into the active learning cycle to further improve the data efficiency. Soleimany et al. [2021] further test the idea of modeling the epistemic uncertainty via EDL, with the quantified uncertainty correlated with prediction error. Yang and Li [2023] also have tested MVE and Deep Ensemble based UQ on a molecular property prediction task, demonstrating the effectiveness in data noise identification and OOD data detection. Griffiths et al. [2022] and Griffiths [2023] use GP to model different types of uncertainties in molecular property prediction as well as molecular discovery tasks, which have been further extended to an open-source package to facilitate real-world scientific applications.

Among the DFT-related quantum mechanics computation tasks, Fowler et al. [2019] have estimated different types of uncertainty using MVE and deep ensemble in a task to predict ground state electron density and show that the quantified uncertainty is informative to help detect inaccurate predictions. Mahmoud et al. [2020] use a sparse Gaussian process to quantify the uncertainty in predicting the electronic density of states using the quasiparticle energy levels, and show that the predicted uncertainty can identify the problematic test structures. Tran et al. [2020] provide a benchmark study of UQ on a task to predict the adsorption energies of materials given atomic structures, with the best-performing model being a GP added at the end of a convolutional neural network. Wollschläger et al. [2023] propose six desiderata for UQ in molecular force field and further introduce localized neural kernel (LNK), a GNN-based deep kernel for GP-based uncertainty quantification, which is the first method to fulfill all of the six desiderata.

For deep models as surrogates for solving PDE problems, Psaros et al. [2023] have conducted a comprehensive study by applying different UQ methods on PINN and DeepONet with various evaluation metrics on forward PDE problems, mixed PDE with known and unknown noise, and operator learning problems. The authors also propose to combine generative adversarial network (GAN) and GP as a functional prior (FP) to harness historical data and reduce the computational cost. While the relative performance of different methods differs in different tasks, the quantified uncertainty is shown to be indicative of prediction error and informative for detecting OOD data.

There are also several benchmark studies aiming at comparing different UQ methods in terms of the informativeness to error, faithfulness of data fitting, and calibration in molecular property prediction tasks. Scalia et al. [2020] have benchmarked different UQ methods for chemical molecular property prediction tasks with the results in favor of ensemble-based UQ. In Hirschfeld et al. [2020], different UQ methods for small organic molecules property prediction have been tested showing that random forest (RF) [Ho 1995] and GP predictors on GNN-based features can provide the best UQ performance. Greenman et al. [2023] have provided another benchmark on UQ for protein property prediction with the results suggesting that no UQ method can perform consistently better than other competitive methods. Li et al. [2023b] also benchmark UQ methods for molecular property prediction with various training schemes, network architectures as well as post-hoc calibration

approaches, whose results suggest that different UQ methods may surpass others on different tasks with different experiment setups.

To summarize, by incorporating different UQ methods, existing scientific AI models for various tasks can get reasonable uncertainty without harming the predictive performance. Moreover, the quantified uncertainty can be useful for OOD data detection [Fowler et al. 2019; Mahmoud et al. 2020; Yang and Li 2023], data noise identification [Yang and Li 2023], and has the potential to be incorporated into active learning [Zhang et al. 2019a; Soleimany et al. 2021; Hie et al. 2020] and Bayesian experimental design [Hie et al. 2020] cycles for data-efficient model training and new molecular discovery.

10.4.5 *Open Research Directions.*

Evaluation of Uncertainty Quantification (UQ): One major difference between scientific AI modeling and other machine learning tasks is that often AI/ML models are considered as surrogates for more computationally expensive mechanistic models based on physics principles. Although existing research and benchmark studies have designed various UQ evaluation metrics by considering different aspects of uncertainty for AI/ML surrogates, comprehensive UQ evaluation is still challenging due to prohibitive computational cost to simulate all the underlying stochastic scenarios. It may be critical to construct new benchmark datasets based on corresponding high-fidelity computational methods and develop UQ evaluation metrics to help standardize UQ for scientific AI with guaranteed performance.

Development of UQ Methods with Domain Knowledge: As has been pointed out in many existing benchmark studies [Scalia et al. 2020; Hirschfeld et al. 2020; Psaros et al. 2023], even though some UQ methods may perform relatively well on a specific task, there is no existing UQ method that can consistently outperform other methods with different setups on different evaluation metrics. As there is not a general rule of thumb for applying UQ methods, many existing deep models for science with UQ capability are developed by empirically testing different existing UQ methods. There is a lack of UQ methods with the properties of physical or biological process explicitly considered. The development of new UQ methods for scientific AI models with the integrated domain knowledge is a research direction to be considered in the future.

Scalable UQ Approaches for Large Models: Most of the existing UQ methods for Deep Neural Networks are either too simplistic and restrictive to achieve satisfactory UQ performance, e.g., MC dropout, or computationally too expensive to be deployed in practice, e.g., deep ensemble and BNN. As the large models with billions of parameters start dominating more and more tasks in natural language processing and computer vision, there has been increasing interest in applying those large models on scientific discovery. Therefore, it is a promising research direction to develop new UQ methods, which is scalable for large models and can better trade-off between computational complexity and quality of the quantified uncertainty. Various types of approximate heuristics, stochastic gradient sampling variants, as well as variational inference techniques have been developed [Graves 2011; Welling and Teh 2011; Li et al. 2016b; Blundell et al. 2015; Shi et al. 2017; Gal and Ghahramani 2016; Gal et al. 2017a; Boluki et al. 2020; Dadaneh et al. 2020; Fan et al. 2020]. New approximation strategies, including recent subspace-based methods [Izmailov et al. 2019; Zhou et al. 2019; Dusenberry et al. 2020; Chen and Ghattas 2020; Boluki et al. 2023], may help further scale up UQ for large models.

11 LEARNING, EDUCATION, AND BEYOND

The advancement of AI holds immense promise for accelerating scientific discovery, driving innovations, and solving complex problems across various domains. However, to fully harness the potential of AI for scientific research, new challenges are faced on education, workforce development, and public engagements. In this section, we first collect existing resources for fundamentals of each AI and Science field. Next, we identify three main paradigm shifts, including discipline boundaries, communities, and educational resources. Finally, we point out recent progress and call for future actions to construct our new knowledge and community system to support the ever-growing AI for Science field.

11.1 Existing Resources for Fundamental AI and Science

Authors: Yuanqi Du, Yaochen Xie, Xiner Li, Shurui Gui, Tianfan Fu, Jimeng Sun, Xiaofeng Qian, Shuiwang Ji

In the ever-evolving landscape of AI and scientific fields, traditional learning materials such as books and courses have long served as the fundamental for knowledge acquisition. Conferences, particularly within AI and each scientific community, have been the traditional medium for fostering collaboration and sharing groundbreaking research. In this section, we lay out existing resources of different types covering fundamentals of each individual field of AI and Science in Table 34 and Table 36. Specifically, the most representative resource types are books, courses and libraries developed for computational purposes.

11.2 Paradigm Shifts in AI for Science

Authors: Yuanqi Du, Yaochen Xie, Xiner Li, Shurui Gui, Tianfan Fu, Jimeng Sun, Xiaofeng Qian, Shuiwang Ji

Despite the accumulating resources for individual AI and Science fields, the emerging field of AI for Science continues to grapple with substantial paradigm shifts. Educational resource types, community levels, and knowledge collection methods specific to AI for Science remain fragmented, necessitating a need for consolidation and further development. We identify three main paradigm shifts in this section.

Transcending Discipline Boundaries: The knowledge system in AI for Science should transcend disciplinary boundaries, promoting interdisciplinary collaborations to address multifaceted challenges and opportunities (Figure 40). Breaking traditional silos between scientific disciplines fosters a comprehensive understanding of complex problems and encourages innovative solutions. The integration of diverse perspectives from fields like physics, biology, chemistry, and artificial intelligence enhances knowledge breadth and facilitates the cross-pollination of ideas. This symbiotic relationship between AI and Science not only allows for shared problem-solving approaches but also enables principled advancements in AI research and development. By leveraging scientific principles, methodologies, and interdisciplinary techniques, breakthroughs in scientific discovery can be achieved to tackle pressing challenges in the AI for Science domain. Existing examples have already demonstrated the power of integrating AI and Science, such as interpreting neural networks with physical laws [Sorscher et al. 2022; Di Giovanni et al. 2023], designing generative models with dynamical system, control and optimal transport [Song et al. 2020; Xu et al. 2022a; Liu et al. 2022a; Berner et al. 2022], solving grand challenges like protein structure prediction [Jumper et al. 2021], and more. This collaborative approach will pave the way for breakthroughs in scientific discovery and enable us to tackle the most pressing challenges in the realm of AI for Science.

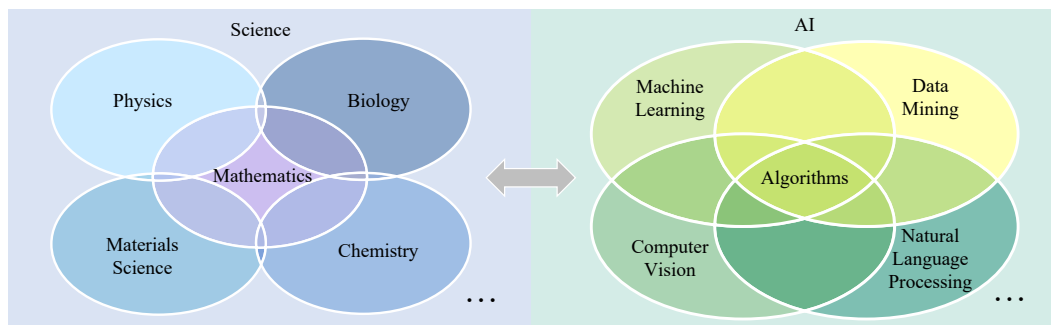


Fig. 40. Traditional scientific fields have recognized the power of interdisciplinary collaborations, leading to the emergence of new fields. Similarly, the intersection of AI and Science promises to forge new frontiers, as these two domains merge their strengths and synergize to tackle challenges in both AI and Science. Note that there are far more scientific fields and cross-domain overlappings that cannot be illustrated in this figure.

Fostering a Diverse and Agile Community: The AI for Science community celebrates diversity and flexibility, extending beyond the boundaries of AI and Science to include students, researchers, and practitioners with a shared interest in advancing the field. While traditional events like NeurIPS and ACS meetings have showcased the forefront of AI and Science research, there is a growing recognition of the need to expand beyond these established platforms. For example, there has been a significant increase in AI-related articles in chemical journals, with a 133% rise in ACS Omega from 2020 to 2021 [Imberti 2022]. Initiatives such as AI for Science workshops and symposiums have emerged, aiming to foster dialogue and collaboration between the AI and Science communities (Table 35). These events range from local gatherings to global conferences and facilitate collaborations between industry and research institutions (Figure 41). By embracing diversity, the community nurtures creativity, critical thinking, and unconventional problem-solving, harnessing a wealth of expertise and insights from individuals with diverse backgrounds. This collaborative environment propels the progress of AI for Science.

Enriching Educational Landscape: The educational resources in AI for Science are expanding beyond what we have seen before. As AI continues to revolutionize the scientific landscape, the demand for high-quality educational resources in AI for Science is growing rapidly. To meet this demand, numerous institutions and individuals are offering an array of educational resources, including summer schools, blogs, tutorials, paper reading groups, etc (Table 35). These resources cover a wide range of topics, from Fundamental concepts in AI to advanced techniques specific to scientific domains. Additionally, collaborations between academia and industry are enriching the educational landscape by providing real-world applications and case studies. However, despite the commendable efforts made to expand educational resources, challenges persist in establishing a systematic approach to learning AI for Science. The rapid pace of methodological advancements and the interdisciplinary nature of the field pose hurdles in curating a comprehensive curriculum. To address this, it becomes imperative to prioritize the development of structured educational programs that encompass the breadth and depth of AI for Science. Such programs should provide a well-rounded understanding of fundamental concepts, advanced methodologies, and their applications across scientific disciplines.

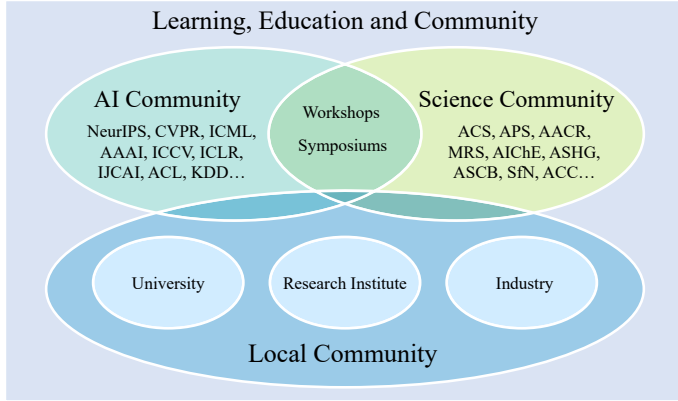


Fig. 41. Within the realm of AI and Science, there exists a vibrant global community encompassing researchers, experts, and enthusiasts from various backgrounds. This global network is complemented by local communities, including universities, research institutes, and industry partners. Through concerted efforts and collaboration, these communities form a powerful ecosystem, driving innovation, knowledge exchange, and transformative discoveries at both local and global scales.

11.3 Prospective and Proposed Actions

Authors: Yuanqi Du, Yaochen Xie, Xiner Li, Shurui Gui, Tianfan Fu, Jimeng Sun, Xiaofeng Qian, Shuiwang Ji

Significant progresses have been made to develop resources for AI for Science in recent years (Table 35 and Table 36). However, these resources often operate independently, lacking a cohesive and systematic road map. As the field undergoes paradigm shifts, it is crucial to develop a unified road map and resources to fill the missing gap. By recognizing the need for comprehensive educational materials, collaborative community platforms, and effective knowledge collection methods, we can better equip researchers, practitioners, and students with the tools and insights necessary to navigate the evolving landscape of AI for Science. Building on top of them, individuals are encouraged to contribute their expertise to subareas of AI for Science. It is through collective efforts that we can fully leverage the potential of AI for Science.

Table 34. Learning resources for fundamental AI or Science fields (open-source denotes that the code is publicly available and further development is permitted). Note that this table is by no means complete and only consists of a small set of available resources.

	Fundamental AI/Science	Type	Description
Symposiums/ Conferences	APS	Physics	American Physical Society
	ACS	Chemistry	American Chemical Society
	MRS	Materials	Materials Research Society
	AICHE	Chemistry	American Institute of Chemical Engineers
	TMS	Materials	The Minerals, Metals & Materials Society
	NeurIPS	AI	Neural Information Processing System
	ICLR	AI	Intl. Conf. on Learning Representations
	ICML	AI	Intl. Conf. on Machine Learning
	AAAI	AI	AAAI Conference on Artificial Intelligence
Courses	Computational Biology	Biology	-
	Quantum Physics	Physics	-
	Machine Learning	AI	-
	Deep Learning	AI	-
	Theoretical Chemistry	Chemistry	-
	Mechanical Engineering Analysis	Engineering	-
Software & Library	PySCF	Quantum Chemistry	Open-Source Quantum Chemistry Code
	PSI4	Quantum Chemistry	Open-Source Quantum Chemistry Code
	NWChem	Quantum Chemistry	Open-Source Quantum Chemistry Code
	CP2K	Quantum Chemistry	Open-Source Quantum Chemistry Code
	ORCA	Quantum Chemistry	Quantum Chemistry Code
	GAUSSIAN	Quantum Chemistry	Quantum Chemistry Code
	Q-Chem	Quantum Chemistry	Quantum Chemistry Code
	Quantum-ESPRESSO	First-Principles	Open-Source Electronic Structure Code
	ABINIT	First-Principles	Open-Source Electronic Structure Code
	GPAW	First-Principles	Open-Source Electronic Structure Code
	BerkeleyGW	First-Principles	Open-Source Electronic Structure Code
	WEST	First-Principles	Open-Source Electronic Structure Code
	Octopus	First-Principles	Open-Source Electronic Structure Code
	exciting	First-Principles	Open-Source Electronic Structure Code
	SIESTA	First-Principles	Open-Source Electronic Structure Code
	OpenMX	First-Principles	Open-Source Electronic Structure Code
	ABACUS	First-Principles	Open-Source Electronic Structure Code
	Wannier90	First-Principles	Open-Source Electronic Structure Code
	EPW	First-Principles	Open-Source Electronic Structure Code
	WIEN2k	First-Principles	Electronic Structure Code
	VASP	First-Principles	Electronic Structure Code
	FHI-aims	First-Principles	Electronic Structure Code
	CASTEP	First-Principles	Electronic Structure Code
	pymatgen	Materials	Open-Source Python Library for Materials Analysis
	ASE	Materials	Open-Source Python Library for Atomistic Simulations
	JARVIS-Tools	Materials	Software Package for Atomistic Data-Driven Materials Design
	PAOFLOW	Materials	Open-Source Code for Post-Processing First-Principles Calculations
	XtalOpt	Materials	Open-Source Crystal Structure Search Code
	CALYPSO	Materials	Crystal Structure Search Code
	USPEX	Materials	Crystal Structure Search Code
	Jmol	Atomistic	Open-Source Atomistic Visualization Software
	AtomEye	Atomistic	Open-Source Atomistic Visualization Software
	OVITO	Atomistic	Open-Source Atomistic Visualization Software
	Avogadro 2	Atomistic	Open-Source Atomistic Visualization Software
	VESTA	Atomistic	Atomistic Visualization Software
	PyMOL	Atomistic	Molecular Visualization Software
	RDKit	Cheminformatics	Open-Source Cheminformatics Software
	OpenBabel	Cheminformatics	Open-Source Cheminformatics Software
	AutoDock Vina	Cheminformatics	Open-Source Molecular Docking
	OpenMM	Molecular Dynamics	Open-Source Molecular Simulation Package
	GROMACS	Molecular Dynamics	Open-Source Molecular Simulation Package
	Amber	Molecular Dynamics	Molecular Simulation Package
	LAMMPS	Molecular Dynamics	Open-Source Molecular Simulation Package
	MDAnalysis	Molecular Dynamics	Open-Source Python Library for Molecular Dynamics Analysis
	Rosetta	Biology	Protein Structure Analysis
	Biotite	Biology	Open-Source Python Library for Computational Molecular Biology
	Biopython	Biology	Open-Source Python Library for Biological Computation
	ScanPy	Biology	Open-Source Python Library for Single-Cell Analysis
	PyClaw	PDE	Open-Source Finite Volume Numerical Solvers for PDE in Python

Table 35. Learning resources for AI for Science. Note that this table is by no means complete and only consists of resources commonly used by the authors.

	AI for Science	Type	Description
Workshops	AI4Science	General	AI for Science
	ML4PS	General	Machine Learning for Physical Sciences
	NSF AI4Science	General	AI-Enabled Scientific Revolution
	MLSB	Atomistic	Machine Learning for Structural Biology
	ML4Molecules	Atomistic	Machine Learning for Molecules
	AI4Mat	Atomistic	AI for Acc. Materials Design
	AIMS	Atomistic	Artificial Intelligence for Materials Science
Symposiums/Conferences	SimDL	Continuum	Deep Learning for Simulation
	AAAI Spring Symposium MoML	General Atomistic	Comp. Approaches to Scientific Discovery Molecular ML Conference
Research Institutes and Labs	IPAM	General	Institute for Pure & Applied Math. at UCLA
	CUAISci	General	Cornell University AI for Science Institute
	AI4Science	General	AI for Science Initiative at Caltech
	AI4ScienceLab	General	AI for Science Lab at UvA
	A3D3	General	Acc. AI Algo. for Data-Driven Discovery
	IAIFI	General	Institute for AI and Fundam. Interactions
	AI & Science	General	AI & Science Initiative at UChicago
	Molecule Maker Lab Institute	Atomistic	AI Institute for Molecule Discovery and Synthesis
	AI Institute in Dynamic Systems	Continuum	-
Tutorials & Blogs	AI4Science101 Blog Series	General	-
	AI4Science Tutorial Series	General	-
	Deep Learning and Quantum Many-Body Computation	Quantum	-
	Tutorial on Quantum Many-body problem	Quantum	-
	Neural Operator	Continuum	-
	Physics-Informed Neural Networks	Continuum	-
Reading Groups & Seminars	Scientific ML Webinar	General	Scientific Machine Learning Webinar Series
	AI4Science Seminar	General	AI for Science Seminar at Chalmers
	M2D2 Reading Group	Atomistic	Molecular Modeling & Drug Discovery
Courses	Data-driven Science and Engineering	General	-
	Group Equivariant Deep Learning	General	-
	Symmetry and its application to ML	General	-
	AI for Science Summer School	General	AI for Science Summer School at UChicago
	Crash Course on Neural Operators	Continuum	-
Software & Libraries	E3NN	General	Machine Learning and Symmetry Library
	DIG	General	Geometric Deep Learning Library
	NetKet	Quantum	Machine Learning for Quantum Physics
	DeepChem	Atomistic	Machine Learning for Molecules
	TDC	Atomistic	Machine Learning for Therapeutic Molecules
	DeePMD	Atomistic	Deep Learning Interatomic Potential and Force Field
	M ² Hub	Atomistic	Machine Learning for Materials Discovery
	Jax CFD	Continuum	Machine Learning for Computational Fluid Dynamics
	ΦFlow	Continuum	Open-Source Python PDE Solver Compatible with Popular Deep Learning Frameworks
Competitions & Benchmarks	Open Catalyst Project	Atomistic	Discover New Catalyst
	Open Graph Benchmark	Atomistic	Molecular Property Prediction
	PDEArena	Continuum	Operator Learning
	PDEBench	Continuum	Operator Learning
Review Papers	Machine Learning and Physical Sciences	General	-
	Quantum Chemistry in the Age of Machine Learning	Quantum	-
	Roadmap on Machine learning in electronic structure	Quantum	-
	Physics-Guided Deep Learning for Dynamical System	Continuum	-

Table 36. Recommended books for fundamental AI, Science and AI for Science fields. Note that this table is by no means complete and only consists of resources commonly used by the authors.

Title	Author	Domain	Info
Deep Learning	Ian Goodfellow, Yoshua Bengio, and Aaron Courville	AI	2016. MIT Press. [Goodfellow et al. 2016]
Pattern Recognition and Machine Learning	Christopher M. Bishop and Nasser M. Nasrabadi	AI	2006. Vol. 4. Springer. [Bishop and Nasrabadi 2006]
Machine Learning: A Probabilistic Perspective	Kevin P. Murphy	AI	2012. MIT Press. [Murphy 2012]
Advanced Engineering Mathematics	Erwin Kreyszig	Mathematics	2011. John Wiley & Sons. [Kreyszig 2011]
The Feynman Lectures on Physics: The New Millennium Edition	Richard Feynman, Robert Leighton, and Matthew Sands	Physics	2011. Basic Books. [Feynman et al. 2011]
Group Theory in a Nutshell for Physicists	Anthony Zee	Group Theory	2016. Vol.17. Princeton University Press. [Zee 2016]
Group Theory: Application to the Physics of Condensed Matter	Mildred S. Dresselhaus, Gene Dresselhaus, and Ado Jorio	Group Theory	2007. Springer Berlin, Heidelberg. [Dresselhaus et al. 2008]
Group Theory in Quantum Mechanics: An Introduction to Its Present Usage	Volker Heine	Group Theory	2007. Courier Corporation. [Heine 2007]
An Introduction to Tensors and Group Theory for Physicists	Nadir Jeevanjee	Group Theory	2011. Springer. [Jeevanjee 2011a]
Symmetry Principles in Solid State and Molecular Physics	Melvin Lax	Group Theory	2001. Courier Corporation. [Lax 2001]
Introduction to Quantum Mechanics	David J. Griffiths and Darrell F. Schroeter	Quantum Mechanics	2018. Cambridge University Press. [Griffiths and Schroeter 2018]
Modern Quantum Mechanics	J. J. Sakurai and J. Napolitano	Quantum Mechanics	2020. Cambridge University Press. [Sakurai and Napolitano 2020]
Quantum Theory of Angular Momentum	D. A. Varshalovich, A. N. Moskalev, and V. K. Khersonskii	Quantum Mechanics	1988. World Scientific. [Varshalovich et al. 1988]
Fundamentals of Condensed Matter Physics	Marvin L. Cohen and Steven G. Louie	Quantum Theory	2016. Cambridge University Press. [Cohen and Louie 2016]
Quantum Theory of Materials	Eftimios Kaxiras and John D. Joannopoulos	Quantum Theory	2019. Cambridge University Press. [Kaxiras and Joannopoulos 2019]
Electronic Structure: Basic Theory and Practical Methods	Richard M. Martin	Quantum Theory	2020. Cambridge University Press. [Martin 2020]
Modern Quantum Chemistry: Introduction to Advanced Electronic Structure Theory	Attila Szabo and Neil S. Ostlund	Quantum Chemistry	2012. Courier Corporation. [Szabo and Ostlund 2012]
Density-Functional Theory of Atoms and Molecules	Robert G. Parr and Weitao Yang	DFT	1995. Oxford University Press. [Parr and Yang 1995]
A Primer in Density Functional Theory	Carlos Fiolhais, Fernando Nogueira, and Miguel A. L. Marques	DFT	2003. Springer Berlin, Heidelberg. [Fiolhais et al. 2003]
Density Functional Theory: An Advanced Course	Eberhard Engel and Reiner M. Dreizler	DFT	2011. Springer Berlin, Heidelberg. [Engel and Dreizler 2011]
Density Functional Theory: An Approach to the Quantum Many-Body Problem	Reiner M. Dreizler and Eberhard K. U. Gross	DFT	2012. Springer Berlin, Heidelberg. [Dreizler and Gross 2012]
Interacting Electrons: Theory and Computational Approaches	Richard M. Martin, Lucia Reining, and David M. Ceperley	DFT	2016. Cambridge University Press. [Martin et al. 2016]
Density Functional Theory: A Practical Introduction	David S. Sholl and Janice A. Steckel	DFT	2009. John Wiley & Sons. [Sholl and Steckel 2009]
A Chemist's Guide to Density Functional Theory	Wolfram Koch and Max C. Holthausen	DFT	2001. John Wiley & Sons. [Koch and Holthausen 2001]
Materials Modelling using Density Functional Theory	Feliciano Giustino	Materials Modeling	2014. Oxford University Press. [Giustino 2014]
Handbook of Materials Modeling	Sidney Yip	Materials Modeling	2005. Springer Netherlands. [Yip 2005]
A Physical Introduction to Fluid Mechanics	Alexander J. Smits	Fluid Mechanics	2000. John Wiley & Sons Incorporated. [Smits 2000]
Lectures in Fluid Mechanics	Alexander J. Smits	Fluid Mechanics	2009. (MAE 553). [Smits 2009]
Turbulent Flows	Stephen B. Pope	Fluid Mechanics	2000. Cambridge University Press. [Pope 2000]
Turbulence, Coherent Structures, Dynamical Systems and Symmetry	Philip Holmes, John L. Lumley, Gahl Berkooz, and Clarence W Rowley	Fluid Mechanics	2012. Cambridge University Press. [Holmes et al. 2012]
Introduction to Partial Differential Equations	Peter J. Olver	PDE	2014. Vol.1. Springer. [Olver 2014]
Partial Differential Equations	Lawrence C. Evans	PDE	2022. Vol.19. American Mathematical Society. [Evans 2022]
Geometric Deep Learning: Grids, Groups, Graphs, Geodesics, and Gauges	Michael M. Bronstein, Joan Bruna, Taco Cohen, and Petar Veličković	AI & Geometry	arXiv preprint arXiv:2104.13478 (2021). [Bronstein et al. 2021]
Data-driven Science & Engineering: Machine learning, dynamical systems, and control	Steven L. Brunton and J. Nathan Kutz	AI & Engineering	2022. Cambridge University Press. [Brunton and Kutz 2022]
Deep Learning for Molecules & Materials	Andrew D. White	AI & Atomistic	LiveCoMS 3, 1499 (2021). [White 2021]

12 CONCLUSION

Advances in deep learning have revolutionized many artificial intelligence (AI) fields. Recently, deep learning has started to advance natural sciences by improving, accelerating, and enabling our understanding of natural phenomena, giving rise to a new area of research, known as AI for science. From our perspective and that of many others, AI for science opens a door for a new paradigm of scientific discovery and represents one of the most exciting areas of interdisciplinary research and innovation. Generally speaking, some scientific processes are described with equations that could be too complicated to be solvable, while others are understood from observable data acquired via (expensive) experiments. The mission of AI is to solve such scientific problems accurately and efficiently, along with many other parameters, such as symmetry in AI models, interpretability, out-of-distribution generalization and causality, uncertainty quantification, etc. In this paper, we provide a technical and unified review of several research areas in AI for science that researchers have been working on during the past several years. We organize different areas of AI for science by the spatial and temporal scales at which the physical world is modeled. In each area, we provide a precise problem setup and discuss the key challenges of using AI to solve such problems. We then provide a survey of major approaches that have been developed, along with datasets and benchmarks for evaluation. We further summarize the remaining challenges and point out several future directions for each area. Particularly, as AI for science is an emerging field of research, we have compiled categorized lists of resources in this paper to facilitate learning and education. We understand that given the evolving nature of this area, our work is by no means comprehensive or conclusive. Thus, we expect to continuously include more topics as the area develops and welcome any feedback and comments from the community.

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A CLASSIFYING AND COMPUTING IRREDUCIBLE REPRESENTATIONS

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Here we give a brief overview of the classification and computation of irreps for various types groups. A key tool for classifying and computing the irreps of various groups is Schur's lemma.

THEOREM 1 (SCHUR'S LEMMA). *Let ρ_X and ρ_Y be irreps of group G acting on vector spaces X and Y respectively. Let $Q : X \rightarrow Y$ be a linear map such that $Q\rho_X(g) = \rho_Y(g)Q$ for all $g \in G$. Then Q must be zero or an isomorphism. If $\rho_X = \rho_Y$ and $X = Y$ is finite-dimensional over an algebraically closed field, then Q must be a scalar multiple of the identity.*

Using Schur's lemma, it is possible to algorithmically decompose any reducible representation into irreps. Suppose we have some reducible representation ρ_X acting on space X . Consider the set of linear transformations $Q : X \rightarrow X$ such that $Q\rho_X(g) = \rho_X(g)Q$. Note that the condition $Q\rho_X(g) = \rho_X(g)Q$ can be rewritten as $Q\rho_X(g) - \rho_X(g)Q = 0$. Since $Q\rho_X(g) - \rho_X(g)Q$ is a linear operation on Q , this is just a nullspace problem. In particular, using standard linear algebra techniques we can find a basis Q_1, Q_2, \dots, Q_m spanning this nullspace.

Suppose $V \subset X$ is a subspace where $\rho_X|_V$ is an irrep and there are no other subspaces isomorphic to this one. Then by Schur's lemma, if we restrict all the Q_i to V , they must all either be a multiple of the identity or 0. In particular, for any linear combination $Q = \sum_{i=1}^r c_i Q_i$, this means that V must be an eigenspace of Q . The case where there are multiple copies of the same irrep is more complicated but a similar result holds. Hence, we can pick random coefficients r_i and compute $Q = \sum_{i=1}^r r_i Q_i$. The eigenspaces of this Q will with high probability give us a decomposition of X into irreps.

The method described above is extremely powerful since it gives us a way to decompose a representation into irreps without knowing what the irreps were in the first place.

Classifying irreps for Finite Groups: While the method described above is great for computing irreps, it does not tell us how to classify them. In particular, we would like a way to identify isomorphic irreps as the same. This motivates the concept of characters and character theory.

DEFINITION 10 (CHARACTER). *The character of a representation ρ_X is defined as*

$$\chi_{\rho_X}(g) = \text{Tr}[\rho_X(g)]. \quad (119)$$

One can check that isomorphic representations share the same character. It turns out the converse is true as well, representations with the same character must be isomorphic. Hence, characters give a better way of labelling representations. Further, since conjugation leaves a trace invariant, the character is the same across all elements of G in the same conjugacy class. So it is natural to list the character of a representation as a function of the conjugacy classes of the group. We refer the reader to a group theory textbook for more discussion on characters, such as *Group Theory: Application to the physics of Condensed Matter* [Dresselhaus et al. 2008].

In the case of finite groups, it turns out we can construct a representation which contains all the irreps. This representation is known as the regular representation.

DEFINITION 11 (REGULAR REPRESENTATION). *Let G be a group and let V be a vector space generated by the group (each element $g \in G$ is identified with a basis of V). The regular representation ρ is defined by*

$$\rho(g)h = gh. \quad (120)$$

We can construct the regular representation of any group using its multiplication table. Decomposing this representation into irreps would give us all possible irreps of the group.

Classifying irreps for the Semisimple Lie Groups: Classifying the irreps of infinite groups is in general a very hard problem. Using characters fail since we need characters for every group element and there would be infinite many of them. However, for the case of semisimple Lie algebras the classification is well known. This also covers the majority of groups used in scientific applications such as $SO(3)$. Here we give a brief overview of the mathematics involved using $SO(3)$ as an example. We begin by defining Lie groups and Lie algebras.

DEFINITION 12 (LIE GROUP). A Lie group is a group that is also a finite-dimensional smooth manifold. In particular, group multiplication and inversion are smooth maps.

A simple example of a Lie group is $SO(2)$, the group of 2D rotations. The corresponding manifold for $SO(2)$ is the circle where we can map the polar angle of each point to a counterclockwise rotation by that angle. Another example is $SO(3)$, the group of 3D rotations. The manifold corresponding to $SO(3)$ is more complicated and is the real projective space \mathbb{RP}^3 .

Because Lie groups are differentiable manifolds, one can instead study a local neighborhood rather than the entire manifold. One typically looks at the tangent space around the identity element in the group. The group multiplication induces a structure on this tangent space known as the Lie bracket. This tangent space along with the Lie bracket structure is known as a Lie algebra. Formally, a Lie algebra is defined as follows.

DEFINITION 13 (LIE ALGEBRA). A Lie algebra is a vector space \mathfrak{g} (over some field F) together with a binary operation called the Lie bracket $[\cdot, \cdot] : \mathfrak{g} \times \mathfrak{g} \rightarrow \mathfrak{g}$ which satisfies

(1) *Bilinearity*

$$[ax + by, z] = a[x, z] + b[y, z] \quad [z, ax + by] = a[z, x] + b[z, y]$$

for scalars $a, b \in F$ and $x, y, z \in \mathfrak{g}$

(2) *Alternativity*

$$[x, x] = 0$$

for $x \in \mathfrak{g}$

(3) *Jacobi identity*

$$[x, [y, z]] + [y, [z, x]] + [z, [x, y]] = 0$$

for all $x, y, z \in \mathfrak{g}$

(4) *Anticommutativity*

$$[x, y] = -[y, x]$$

for all $x, y \in \mathfrak{g}$.

For matrices, the Lie bracket is just the commutator $[x, y] = xy - yx$.

As an example, let us derive the Lie algebra $\mathfrak{so}(3)$ corresponding to $SO(3)$. Any rotation close to the identity can be written as a perturbation $I + \epsilon X$ where X is in the tangent space around the identity (Note: the physics convention adds an extra factor of i in front of X but the math convention does not). Our main condition is orthogonality, so we must have

$$(I + \epsilon X)^T (I + \epsilon X) = I + \epsilon(X^T + X) + \epsilon^2 X^T X = I.$$

Since ϵ^2 is small, the condition on X is $X^T + X = 0$ or antisymmetry. One set of bases for this vector space is

$$x = \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & -1 \\ 0 & 1 & 0 \end{pmatrix} \quad y = \begin{pmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{pmatrix} \quad z = \begin{pmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}. \quad (121)$$

One can check that the commutation relations are $[x, y] = z$, $[y, z] = x$, $[z, x] = y$. It is interesting to note that except for a factor of i due to the physics convention, the commutation relations of

$\mathfrak{so}(3)$ are exactly the commutation relations of the spin operators. This is not a coincidence, the group used in quantum mechanics to describe spin is $SU(2)$ and the corresponding Lie algebra $\mathfrak{su}(2)$ is the same as $\mathfrak{so}(3)$.

These commutation relations are what define the Lie algebra. A set of matrices such as the ones shown above which satisfy these relations are called representations of the Lie algebra.

DEFINITION 14 (LIE ALGEBRA REPRESENTATION). Consider a Lie algebra \mathfrak{g} and vector space X . A representation of \mathfrak{g} is a pair (ρ_X, X) where

$$\rho_X : \mathfrak{g} \rightarrow \mathfrak{gl}(V)$$

is an algebra homomorphism from \mathfrak{g} to the general linear algebra of X . In particular, that ρ_X is a homomorphism means that

$$\rho_X([A, B]_{\mathfrak{g}}) = [\rho_X(A), \rho_X(B)]_{\mathfrak{gl}(V)} = \rho_X(A)\rho_X(B) - \rho_X(B)\rho_X(A).$$

Note the similarity to group representations. The definitions of reducible and irreducible representations are analogous. It turns out for a class of Lie algebras known as semisimple Lie algebras, all representations can be reduced to a sum of irreducible ones. This is known as Weyl's theorem on complete reducibility. Hence, classifying irreps of the semisimple Lie algebras classifies all representations. Further, there is also Schur's lemma for irreducible representations of Lie algebras so we can use the algorithm described earlier to also decompose arbitrary Lie algebra representations into irreducible ones.

To classify the representations of the semisimple Lie algebras, one must take a close look at the vector space the representation acts on. Given any matrix and a vector space, one can always split the vector space into eigenspaces of the matrix. However, if there are repeated eigenvalues then this does not completely split the vector space. But if we have a set of commuting matrices, then we can possibly split the vector space more finely. For the semisimple Lie algebras, it turns out there is a particular subalgebra where all elements commute and lets us split the vector space of any representation as finely as possible. This subalgebra is known as a Cartan subalgebra.

DEFINITION 15 (CARTAN SUBALGEBRA). Suppose we have a Lie algebra \mathfrak{g} . A subalgebra \mathfrak{h} of \mathfrak{g} is a Cartan subalgebra if it is

(1) Nilpotent. That is the following sequences terminates in the zero subalgebra

$$\mathfrak{h} \geq [\mathfrak{h}, \mathfrak{h}] \geq [\mathfrak{h}, [\mathfrak{h}, \mathfrak{h}]] \geq [\mathfrak{h}, [\mathfrak{h}, [\mathfrak{h}, \mathfrak{h}]]] \geq \dots$$

(2) Self normalizing. That is for all $g \in \mathfrak{g}$ such that $[g, \mathfrak{h}] \subset \mathfrak{h}$, we must have $g \in \mathfrak{h}$.

For $SO(3)$, one choice of a Cartan subalgebra is that spanned by z or $\mathfrak{h} = Fz$. We can check that $[z, z] = 0$ so the subspace is nilpotent. Further, one can check that $[y, z] = x \notin \mathfrak{h}$ and $[x, z] = -y \notin \mathfrak{h}$. Note we could have chosen any 1 dimensional subspace such as x or y , however the choice of z matches up with the conventions used elsewhere and does not matter for purposes of classifying the representations.

With this subspace, we can then use the representations on this subspace to break up the vector space the representations act on.

DEFINITION 16 (WEIGHTS AND WEIGHT SPACES). Let \mathfrak{g} be a semisimple Lie algebra and let \mathfrak{h} be a Cartan subalgebra. Let (ρ_X, X) be a representation. Let λ be a linear functional $\lambda : \mathfrak{h} \rightarrow \mathbb{C}$. Then the weight space V_λ is the subspace

$$V_\lambda = \{v \in X : \forall h \in \mathfrak{h}, \rho(h)v = \lambda(h)v\}.$$

The linear functionals with nonzero weight space are called weights.

Essentially, the simultaneous eigenspaces are called weight spaces and the corresponding eigenvalues are called weights. Since λ is linear on \mathfrak{h} , we can fully specify λ with a list of its eigenvalues for a basis of \mathfrak{h} . In particular, if we have an orthonormal basis for \mathfrak{h} say b_1, b_2, \dots, b_n , then we can define an inner product on the dual space as $\langle \lambda_1, \lambda_2 \rangle = \sum_{i=1}^n \lambda_1(b_i)\lambda_2(b_i)$ so we view it as just a vector of eigenvalues. It turns out there is a unique (up to rescaling) way to define an inner product on the algebra which lets us construct this orthonormal basis. This uses something called an adjoint representation which is similar to the regular representation for finite groups.

DEFINITION 17 (ADJOINT REPRESENTATION, ROOTS, KILLING FORM). Let \mathfrak{g} be a Lie algebra. The adjoint representation is the representation $(\text{ad}_{\mathfrak{g}}, \mathfrak{g})$ such that

$$\text{ad}_{\mathfrak{g}}(A)B = [A, B]_{\mathfrak{g}}.$$

The nonzero weights of the adjoint representation are called roots and the weight spaces are called root spaces. We typically denote the set of roots as Φ .

The Killing form is an symmetric bilinear form on \mathfrak{g} defined by

$$K(A, B) = \text{Tr}[\text{ad}_{\mathfrak{g}}(A) \circ \text{ad}_{\mathfrak{g}}(B)]$$

and can be used to define an inner product.

One can check that the representation for $\mathfrak{so}(3)$ specified in (121) is in fact the adjoint representation. Using this, we can check that the eigenvalues of $\text{ad}_{\mathfrak{so}(3)}(z)$ are $1, 0, -1$ so $(1), (-1)$ are the roots (Note: we made a choice in our scaling of the eigenvalues. In principle we could have just as well picked $z/2$ and had $1/2, 0, -1/2$ but our choice is easier to work with). While in general, the weights can be any vector, there is an important class of weights called integral weights.

DEFINITION 18 (INTEGRAL WEIGHT). Let \mathfrak{g} be a Lie algebra and \mathfrak{h} be a Cartan subalgebra. A weight λ is called integral if for all roots $\alpha \in \Phi$,

$$2 \frac{\langle \lambda, \alpha \rangle}{\langle \alpha, \alpha \rangle}$$

is an integer, where $\langle \cdot, \cdot \rangle$ is the inner product on the dual space as described above.

In the case of $\mathfrak{so}(3)$, the roots are $(1), (-1)$. Hence $\langle \alpha, \alpha \rangle = 1$ so the integral weights are half integer valued.

While any representation can have multiple weights, it turns out all representations can be uniquely identified by a highest weight. To do this in general, we must pick a set of positive roots Φ^+ and use this to define a partial ordering on the weight space. We can then define a dominant weight as one which always has a nonnegative inner product with the positive roots. We refer the reader to a standard textbook on Lie algebras for more details on these concepts such as *Lie Groups, Lie Algebras, and Representations: An Elementary Introduction* [Hall and Hall 2013]. In the case of $\mathfrak{so}(3)$ however, this is easy. We can just pick the positive roots to be (1) . Since our vectors are 1-dimensional, we can order our weights just by numeric value. Further, the dominant weights are simply those with nonnegative numerical value.

We now present the main result known as the theorem of highest weight which classifies all irreducible representations of semisimple Lie algebras.

THEOREM 2 (THEOREM OF HIGHEST WEIGHT). Let \mathfrak{g} be a finite-dimensional semisimple Lie algebra. Then

- (1) Let (ρ_X, X) be an irreducible representation. Then (ρ_X, X) has a unique highest weight and the highest weight is dominant and integral
- (2) If two representations have the same highest weight, then they are isomorphic

- (3) *For every dominant integral weight λ , there is an irreducible representation with highest weight λ .*

Hence, the dominant integral weights classify all the irreducible representations. In the case of $\mathfrak{so}(3)$, this means we can list the representations by nonnegative half integers, corresponding to spin representations and why we can label them by a single number ℓ . This makes sense since $\mathfrak{so}(3)$ and $\mathfrak{su}(2)$ are equivalent. It turns out only the integer ones correspond to representations of $SO(3)$.

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