

# Integrating Physics-Based Modeling With Machine Learning: A Survey

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There is a growing consensus that solutions to complex science and engineering problems require novel methodologies that are able to integrate traditional physics-based modeling approaches with state-of-the-art machine learning (ML) techniques. This paper provides a structured overview of such techniques. Application areas for which these approaches have been applied are summarized, then classes of methodologies used to construct physics-guided ML models and hybrid physics-ML frameworks are described. We then provide a taxonomy of these existing techniques, which uncovers knowledge gaps and potential crossovers of methods between disciplines that can serve as ideas for future research.

Additional Key Words and Phrases: physics-guided, neural networks, deep learning, physics-informed, theory-guided, hybrid, knowledge integration

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## 1 INTRODUCTION

Machine learning (ML) models, which have already found tremendous success in commercial applications, are beginning to play an important role in advancing scientific discovery in domains traditionally dominated by mechanistic (e.g. first principle) models [22, 112, 117, 126, 127, 138, 214, 261]. The use of ML models is particularly promising in scientific problems involving processes that are not completely understood, or where it is computationally infeasible to run mechanistic models at desired resolutions in space and time. However, the application of even the state-of-the-art black box ML models has often met with limited success in scientific domains due to their large data requirements, inability to produce physically consistent results, and their lack of generalizability to out-of-sample scenarios [126]. Given that neither an ML-only nor a scientific knowledge-only approach can be considered sufficient for complex scientific and engineering applications, the research community is beginning to explore the continuum between mechanistic and ML models, where both scientific knowledge and data are integrated in a synergistic manner [2, 14, 126, 204].

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This paradigm is fundamentally different from mainstream practices in the ML community for making use of domain-specific knowledge, albeit in subservient roles, e.g., feature engineering or post-processing. In contrast to these practices that can only work with simpler forms of heuristics and constraints, this survey is focused on those approaches that explore a deeper coupling of ML methods with scientific knowledge.

Even though the idea of integrating scientific principles and ML models has picked up momentum just in the last few years [126], there is already a vast amount of work on this topic. This work is being pursued in diverse disciplines (e.g., earth systems [214], climate science [75, 135, 186], turbulence modeling [28, 179, 273], material discovery [40, 202, 226], quantum chemistry [222, 228], biological sciences [285], and hydrology [275]), and it is being performed in the context of diverse objectives specific to these applications. Early results in isolated and relatively simple scenarios have been promising, and the expectations are rising for this paradigm to accelerate scientific discovery and help address some of the biggest challenges that are facing humanity in terms of climate [75], health [259], and food security [118].

The goal of this survey is to bring these exciting developments to the ML community, to make them aware of the progress that has been made, and the gaps and opportunities that exist for advancing research in this promising direction. We hope that this survey will also be valuable for scientists who are interested in exploring the use of ML to enhance modeling in their respective disciplines. Please note that work on this topic has been referred to by other names, such as "physics-guided ML," "physics-informed ML," or "physics-aware AI," although it covers many scientific disciplines. In this survey, we also use the terms "physics-guided" or "physics," which should be more generally or interpreted as science or scientific knowledge.

The focus of this survey is on approaches that integrate mechanistic modeling with ML, using primarily ideas from physics and other scientific disciplines. This distinguishes our survey from other works that focus on more general knowledge integration into machine learning [1, 256] and other works covering physics integration into ML in specific domains (e.g., cyber-physical systems [204], chemistry [184]). This survey creates a novel taxonomy specific to science-based knowledge and covers a wide array of both methodologies and categories.

We organize the paper as follows. Section 2 describes different objectives that are being used through combinations of scientific knowledge and machine learning. Section 3 discusses novel ML methods and architectures that researchers are developing to achieve these objectives. Section 4 discusses opportunities for future research, and Section 5 contains concluding remarks. Table 1 categorizes the work surveyed in this paper by objectives and approaches.

## 2 OBJECTIVES OF PHYSICS-ML INTEGRATION

This section provides a brief overview of a diverse set of objectives where deep couplings of ML and scientific modeling paradigms are being pursued in the context of various applications. Though we enumerate these distinct categories in the following subsections, there are numerous cross-cutting themes. One is the use of ML as a surrogate model where the goal is to accurately reproduce the behavior of a mechanistic model at substantially reduced computational cost. Reduction in computational cost is a known characteristic of ML in comparison to the often high cost of numerical simulations, and ML-based surrogate models are relevant for many objectives. Also, though uncertainty quantification is its own subsection, it is relevant for most of the other objectives.

### 2.1 Improving predictions beyond that of state-of-the-art physical models

First-principles models are used extensively in a wide range of engineering and environmental applications. Even though these models are based on known physical laws, in most cases, they are

necessarily approximations of reality due to incomplete knowledge of certain processes, which introduces bias. In addition, they often contain a large number of parameters whose values must be estimated with the help of limited observed data, degrading their performance further, especially due to heterogeneity in the underlying processes in both space and time. The limitations of physics-based models cut across discipline boundaries and are well known in the scientific community (e.g., see Gupta et al. [103] in the context of hydrology).

ML models have been shown to outperform physics-based models in many disciplines (e.g., materials science [130, 220, 263], applied physics [15, 116], aquatic sciences [120], atmospheric science [185], biomedical science [245], computational biology [3]). A major reason for this success is that ML models (e.g., neural networks), given enough data, can find structure and patterns in problems where complexity prohibits the explicit programming of a system's exact physical nature. Given this ability to automatically extract complex relationships from data, ML models appear promising for scientific problems with physical processes that are not fully understood by researchers, but for which data of adequate quality and quantity is available. However, the black-box application of ML has met with limited success in scientific domains due to a number of reasons [120, 126]: (i) while state-of-the-art ML models are capable of capturing complex spatiotemporal relationships, they require far too much labeled data for training, which is rarely available in real application settings, (ii) ML models often produce scientifically inconsistent results; and (iii) ML models can only capture relationships in the available training data, and thus cannot generalize to out-of-sample scenarios (i.e., those not represented in the training data).

The key objective here is to combine elements of physics-based modeling with state-of-the-art ML models to leverage their complementary strengths. Such integrated physics-ML models are expected to better capture the dynamics of scientific systems and advance the understanding of underlying physical processes. Early attempts for combining ML with physics-based modeling in several applications (e.g., modeling the lake phosphorus concentration [109] and lake temperature dynamics [120, 128, 213]) have already demonstrated its potential for providing better prediction accuracy with a much smaller number of samples as well as generalizability in out-of-sample scenarios.

## 2.2 Downscaling

Complex first principle models are capable of capturing physical reality more precisely than simpler models, as they often involve more diverse components that account for a greater number of processes at finer spatial or temporal resolution. However, given the computation cost and modeling complexity, many models are run at a resolution that is coarser than what is required to precisely capture underlying physical processes. For example, cloud-resolving models (CRM) need to run at sub-kilometer horizontal resolution to be able to effectively represent boundary-layer eddies and low clouds, which are crucial for the modeling of Earth's energy balance and the cloud-radiation feedback [212]. However, it is not feasible to run global climate models at such fine resolutions even with the most powerful computers expected to be available in the near future.

Downscaling techniques have been widely used as a solution to capture physical variables that need to be modeled at a finer resolution. In general, the downscaling techniques can be divided into two categories: statistical downscaling and dynamical downscaling. Statistical downscaling refers to the use of empirical models to predict finer-resolution variables from coarser-resolution variables. Such a mapping across different resolutions can involve complex non-linear relationships that cannot be precisely modeled by simple empirical models. Recently, artificial neural networks have shown a lot of promise for this problem, given their ability to model non-linear relationships [232, 254]. In contrast, dynamical downscaling makes use of high-resolution regional simulations to dynamically simulate relevant physical processes at regional or local scales of interest. Due to

the substantial time cost of running such complex models, there is an increasing interest in using ML models as surrogate models (a model approximating simulation-driven input–output data) to predict target variables at a higher resolution [90, 237].

Although the state-of-the-art ML methods can be used in both statistical and dynamical downscaling, it remains a challenge to ensure that the learned ML component is consistent with established physical laws and can improve the overall simulation performance.

### 2.3 Parameterization

Complex physics-based models (e.g., for simulating phenomena in climate, weather, turbulence modeling, astrophysics) often use an approach known as *parameterization* to account for missing physics. In parameterization (note that this term has a different meaning when used in mathematics and geometry), specific complex dynamical processes are replaced by simplified physical approximations whose associated parameter values are estimated from data, often through a procedure referred to as parameter calibration. The failure to correctly identify parameter values can make the model less robust, and errors that result from imperfect parameterization can also feed into other components of the entire physics-based model and deteriorate the modeling of important physical processes. Hence, there is great interest in using ML models to learn new parameterizations directly from observations and/or high-resolution model simulation. Already, ML-based parameterizations have shown success in geology [43, 95] and atmospheric science [32, 33, 90, 186]. A major benefit of ML-based parameterizations is the reduction of computation time compared to traditional physics-based simulations. In chemistry, Hansen et al. [108] find that parameterizations of atomic energy using ML take seconds compared to multiple days for more standard quantum-calculation calculations, and Behler et al. [20] find that neural networks can vastly improve the efficiency of finding potential energy surfaces of molecules.

Most of the existing work uses standard black box ML models for parameterization, but there is an increasing interest in integrating physics in the ML models [23], as it has the potential to make them more robust and generalizable to unseen scenarios as well as reduce the number of training samples needed for training.

Note that both parameterization and downscaling described in Section 2.2 are used to replace components of larger models. To avoid confusion, we distinctly use downscaling to relate to resolutions (e.g., resolving certain physical processes to higher resolutions) and use parameterization for creating simplified parameterized approximations of complex dynamical processes.

### 2.4 Reduced-Order Models

Reduced-Order Models (ROMs) are computationally inexpensive representations of more complex models. Usually, constructing ROMs involves dimensionality reduction that attempts to capture the most important dynamical characteristics of often large, high-fidelity simulations and models of physical systems (e.g., in fluid dynamics [145]). This can also be viewed as a controlled loss of accuracy. A common way to do this is to project the governing equations of a system onto a linear subspace of the original state space using a method such as principal components analysis or dynamic mode decomposition [201]. However, limiting the dynamics to a lower-dimensional subspace inherently limits the accuracy of any ROM.

ML is beginning to assist in constructing ROMs for increased accuracy and reduced computational cost in several ways. One approach is to build an ML-based surrogate model for full-order models [46], where the ML model can be considered a ROM. Other ways include building an ML-based surrogate model of an already built ROM by another dimensionality reduction method [273] or building an ML model to mimic the dimensionality reduction mapping from a full-order model to a reduced-order model [179]. ML and ROMs can also be combined by using the ML model to learn

the residual between a ROM and observational data [257]. ML models have the potential to greatly augment the capabilities of ROMs because of their typically quick forward execution speed and ability to leverage data to model high dimensional phenomena.

One area of recent focus of ML-based ROMs is in approximating the dominant modes of the Koopman (or composition) operator, as a method of dimensionality reduction. The Koopman operator is an infinite-dimension linear operator that encodes the temporal evolution of the system state through nonlinear dynamics [36]. This allows linear analysis methods to be applied to nonlinear systems and enables the inference of properties of dynamical systems that are too complex to express using traditional analysis techniques. Applications span many disciplines, including fluid dynamics [233], oceanography [93], molecular dynamics [267], and many others. Though dynamic mode decomposition [178] is the most common technique for approximating the Koopman operator, many recent approaches have been made to approximate Koopman operator embeddings with deep learning models that outperform existing methods [150, 163, 171, 180, 188, 190, 244, 262, 286]. Adding physics-based knowledge to the learning of the Koopman operator has the potential to augment generalizability and interpretability, which current ML methods in this area tend to lack [190].

ROMs often lack robustness with respect to parameter changes within the systems they are representing [5], or are not cost-effective enough when trying to predict complex dynamical systems (e.g., multiscale in space and time). Thus, incorporating principles from physics-based models could potentially reduce the search space to enable more robust training of ROMs, and also allow the model to be trained with less data in many scenarios.

## 2.5 Inverse Modeling

The forward modeling of a physical system uses the physical parameters of the system (e.g., mass, temperature, charge, physical dimensions or structure) to predict the next state of the system or its effects (outputs). In contrast, inverse modeling uses the (possibly noisy) output of a system to infer the intrinsic physical parameters. Inverse problems often stand out as important in physics-based modeling communities because they can potentially shed light on valuable information that cannot be observed directly. One example is the use of x-ray images from a CT scan to create a 3D image reflecting the structure of part of a person's body. This can be viewed as a computer vision problem, where, given many training datasets of x-ray scans of the body at different capture angles, a model can be trained to inversely reconstruct textures and 3D shapes of different organs or other areas of interest. Allowing for better reconstruction while reducing scan time could potentially increase patient satisfaction and reduce overall medical costs.

Often, the solution of an inverse problem can be computationally expensive due to the potentially millions of forward model evaluations needed for estimator evaluation or characterization of posterior distributions of physical parameters [84]. ML-based surrogate models (in addition to other methods such as reduced-order models) are becoming a realistic choice since they can model high-dimensional phenomena with lots of data and execute much faster than most physical simulations.

Inverse problems are traditionally solved using regularized regression techniques. Data-driven methods have seen success in inverse problems in remote sensing of surface properties [59], photonics [197], and medical imaging [161], among many others. Recently, novel algorithms using deep learning and neural networks have been applied to inverse problems. While still in their infancy, these techniques exhibit strong performance for applications such as computerized tomography [47, 174], seismic processing [253], or various sparse data problems.

There is also increasing interest in the inverse design of materials using ML, where desired target properties of materials are used as input to the model to identify atomic structures that exhibit such

properties [137, 151, 202, 226]. Physics-based constraints and stopping conditions based on material properties can be used to guide the optimization process [151]. These constraints and similar physics-guided techniques have the potential to alleviate noted challenges in inverse modeling, particularly in scenarios with a small sample size and a paucity of ground-truth labels [127]. The integration of prior physical knowledge is common in current approaches to the inverse problem, and its integration into ML-based inverse models has the potential to improve data efficiency and increase its ability to solve ill-posed inverse problems.

## 2.6 Forward Solving Partial Differential Equations

In many physical systems, governing equations are known, but direct numerical solutions of partial differential equations (PDEs) using common methods, such as the Finite Elements Method or the Finite Difference Method [80], are prohibitively expensive. In such cases, traditional methods are not ideal or sometimes even possible. A common technique is to use an ML model as a surrogate for the solution to reduce computation time [65, 140]. In particular, NN solvers can reduce the high computational demands of traditional numerical methods into a single forward-pass of a NN. Notably, solutions obtained via NNs are also naturally differentiable and have a closed analytic form that can be transferred to any subsequent calculations, a feature not found in more traditional solving methods [140]. Especially with the recent advancement of computational power, neural networks models have shown success in approximating solutions across different kinds of physics-based PDEs [9, 131, 215], including the difficult quantum many-body problem [42] and many-electron Schrodinger equation [107]. As a step further, deep neural networks models have shown success in approximating solutions across high dimensional physics-based PDEs previously considered unsuitable for approximation by ML [106, 235]. However, slow convergence in training, limited applicability to many complex systems, and reduced accuracy due to unawareness of physical laws can prove problematic.

## 2.7 Discovering Governing Equations

When the governing equations of a dynamical system are known explicitly, they allow for more robust forecasting, control, and the opportunity for analysis of system stability and bifurcations through increased interpretability [217]. Furthermore, if the learned mathematical model accurately describes the processes governing the observed data, it therefore can generalize to data outside of the training domain. However, in many disciplines (e.g., neuroscience, cell biology, finance, epidemiology) dynamical systems have no formal analytic descriptions. Often in these cases, data is abundant, but the underlying governing equations remain elusive. In this section, we discuss equation discovery systems that do not assume the structure of the desired equation (as in Section 2.6), but rather explore a space a large space of possibly nonlinear mathematical terms.

Advances in ML for the discovery of these governing equations has become an active research area with rich potential to integrate principles from applied mathematics and physics with modern ML methods. Early works on the data-driven discovery of physical laws relied on heuristics and expert guidance and were focused on rediscovering known, non-differential, laws in different scientific disciplines from artificial data [92, 143, 144, 149]. This was later expanded to include real-world data and differential equations in ecological applications [72]. Recently, general and robust data-driven discovery of potentially unknown governing equations has been pioneered by [30, 227], where they apply symbolic regression to differences between computed derivatives and analytic derivatives to determine underlying dynamical systems. More recently, works have used sparse regression built on a dictionary of functions and partial derivatives to construct governing equations [37, 200, 216]. Lagergren et al. [141] expand on this by using ANNs to construct the dictionary of functions. These sparse identification techniques are based on the principle of Occam's

Razor, where the goal is that only a few equation terms be used to describe any given nonlinear system.

## 2.8 Data Generation

Data generation approaches are useful for creating virtual simulations of scientific data under specific conditions. For example, these techniques can be used to generate potential chemical compounds with desired characteristics (e.g., serving as catalysts or having a specific crystal structure). Traditional physics-based approaches for generating data often rely on running physical simulations or conducting physical experiments, which tend to be very time consuming. Also, these approaches are restricted by what can be produced by physics-based models. Hence, there is an increasing interest in generative ML approaches that learn data distributions in unsupervised settings and thus have the potential to generate novel data beyond what could be produced by traditional approaches.

Generative ML models have found tremendous success in areas such as speech recognition and generation [187], image generation [63], and natural language processing [100]. These models have been at the forefront of unsupervised learning in recent years, mostly due to their efficiency in understanding unlabeled data. The idea behind generative models is to capture the inner probabilistic distribution in order to generate similar data. With the recent advances in deep learning, new generative models, such as the generative adversarial network (GAN) and variational autoencoder (VAE), have been developed. These models have shown much better performance in learning nonlinear relationships to extract representative latent embeddings from observation data. Hence the data generated from the latent embeddings are more similar to true data distribution. In particular, the adversarial component of GAN consists of a two-part framework of a generative network and discriminative network, where the generative network's objective is to generate fake data to fool the discriminative network, while the discriminative network attempts to determine true data from fake data.

In the scientific domain, GANs can generate data like the data generated by the physics-based models. Using GANs often incurs certain benefits, including reduced computation time and a better reproduction of complex phenomenon, given the ability of GANs to represent nonlinear relationships. For example, Farimani et al. [78] have shown that Conditional Generative Adversarial Networks (cGAN) can be trained to simulate heat conduction and fluid flow purely based on observations without using knowledge of the underlying governing equations. Such approaches that use generative models have been shown to significantly accelerate the process of generating new data samples.

However, a well-known issue of GANs is that they incur dramatically high sample complexity. Therefore, a growing area of research is to engineer GANs that can leverage prior knowledge of physics in terms of physical laws and invariance properties. For example, GAN-based models for simulating turbulent flows can be further improved by incorporating physical constraints, e.g., conservation laws [288] and energy spectrum [269], in the loss function. Cang et al. [40] also imposed a physics-based morphology constraint on a VAE-based generative model used for simulating artificial material samples. The physics-based constraint forces the generated artificial samples to have the same morphology distribution as the authentic ones and thus greatly reduces the large material design space.

## 2.9 Uncertainty Quantification

Uncertainty quantification (UQ) is of great importance in many areas of computational science (e.g., climate modeling [64], fluid flow [55], systems engineering [195], among many others). At its core, UQ requires an accurate characterization of the entire distribution  $p(y|x)$ , where  $y$  is the

response and  $x$  is the covariate of interest, rather than just making a point prediction  $y = f(x)$ . This makes it possible to characterize all quantiles and skews in the distribution, which allows for analysis such as examining how close predictions are to being unacceptable, or sensitivity analysis of input features.

Applying UQ tasks to physics-based models using traditional methods such as Monte Carlo (MC) is usually infeasible due to the thousands or millions of forward model evaluations needed to obtain convergent statistics. In the physics-based modeling community, a common technique is to perform model reduction (described in Section 2.4) or create an ML surrogate model, in order to increase model evaluation speed since ML models often execute much faster [87, 170, 250]. With a similar goal, the ML community has often employed Gaussian Processes as the main technique for quantifying uncertainty in simulating physical processes [26, 211], but neither Gaussian Processes nor reduced models scale well to higher dimensions or larger datasets (Gaussian Processes scale as  $O(N^3)$  with  $N$  data points).

Consequently, there is an effort to fit deep learning models, which have exhibited countless successes across disciplines, as a surrogate for numerical simulations in order to achieve faster model evaluations for UQ that have greater scalability than Gaussian Processes [250]. However, since artificial neural networks do not have UQ naturally built into them, variations have been developed. One such modification uses a probabilistic drop-out strategy in which neurons are periodically "turned off" as a type of Bayesian approximation to estimate uncertainty [86]. There are also Bayesian variants of neural networks that consist of distributions of weights and biases [164, 296, 300], but these suffer from high computation times and high dependence on reliable priors. Another method uses an ensemble of neural networks to create a distribution from which uncertainty is quantified [142].

The integration of prior physics knowledge into ML for UQ has the potential to allow for a better characterization of uncertainty. For example, ML surrogate models run the risk of producing physically inconsistent predictions, and incorporating elements of physics could help with this issue. Also, note that the reduced data needs of ML due to constraints for adherence to known physical laws could alleviate some of the high computational cost of Bayesian neural networks for UQ.

### 3 PHYSICS-ML METHODS

Given the diversity of forms in which scientific knowledge is represented in different disciplines and applications, a diverse set of methodologies are needed to integrate physical principles into ML models. This section creates a taxonomy of five classes of methodologies to merge principles of physics-based modeling with ML; (i) physics-guided loss function, (ii) physics-guided initialization, (iii) physics-guided design of architecture, (iv) residual modeling, and (v) hybrid physics-ML models.

#### 3.1 Physics-Guided Loss Function

Scientific problems often exhibit a high degree of complexity due to relationships between many physical variables varying across space and time at different scales. Standard ML models can fail to capture such relationships directly from data, especially when provided with limited observation data. This is one reason for their failure to generalize to scenarios not encountered in training data. Researchers are beginning to incorporate physical knowledge into loss functions to help ML models capture generalizable dynamic patterns consistent with established physical laws.

One of the most common techniques to make ML models consistent with physical laws is to incorporate physical constraints into the loss function of ML models as follows [126],

$$\text{Loss} = \text{Loss}_{\text{TRN}}(Y_{\text{true}}, Y_{\text{pred}}) + \lambda R(W) + \gamma \text{Loss}_{\text{PHY}}(Y_{\text{pred}}) \quad (1)$$



where the training loss  $\text{Loss}_{\text{TRN}}$  measures a supervised error (e.g., RMSE or cross-entropy) between true labels  $Y_{\text{true}}$  and predicted labels  $Y_{\text{pred}}$ , and  $\lambda$  is a hyper-parameter to control the weight of model complexity loss  $R(W)$ . These first two terms are the standard loss of ML models. The addition of physics-based loss  $\text{Loss}_{\text{PHY}}$  aims to ensure consistency with physical laws and it is weighted by a hyper-parameter  $\gamma$ .

Steering ML predictions towards physically consistent outputs has numerous benefits. First, the computation of physics-based loss  $\text{Loss}_{\text{PHY}}$  requires no observation data and thus optimizing  $\text{Loss}_{\text{PHY}}$  allows including unlabeled data in training. Second, the regularization by physical constraints can reduce the possible search space of parameters. This provides possibility to learn with less labeled data while also ensuring the consistency with physical laws during optimization. Third, ML models which follow desired physical properties are more likely to be generalizable to out-of-sample scenarios, and thus become acceptable for use by domain scientists and stakeholders in physical science applications. Loss function terms corresponding to physical constraints are applicable across many different types of ML frameworks and objectives. In the following paragraphs, we demonstrate the use of physics-based loss functions for different objectives described in Section 2.

*Improving predictions.* In the context of complex natural systems, incorporation of physics-based loss has shown great success in improving prediction ability of ML models. In the context of lake temperature modeling, Karpatne et al. [128] includes an additional physics-based penalty that ensures that predictions of denser water predictions are at lower depths than predictions of less dense water, a known monotonic relationship. Jia et al. [119] and Read et al. [213] further extended this work to capture even more complex and general physical relationships that happen on a temporal scale. Specifically, they use a physics-based penalty for energy conservation in the loss function to ensure the lake thermal energy gain across time is consistent with the net thermodynamic fluxes in and out of the lake. A diagram of this model is shown in Figure 1. Note that the recurrent structure contains additional nodes (shown in blue) to represent physical variable (lake energy, etc) that are computed using purely physics-based equations. These are needed to incorporate energy conservation in the loss function. Similar structure can be used to model other physical laws such as mass conservation.

*Solving and Discovering PDEs and Governing Equations.* Another strand of work that involves loss function alterations is solving PDEs for dynamical systems modeling, in which adherence to the governing equations is enforced in the loss function. In Raissi et al. [208], this concept is developed and shown to create data-efficient spatiotemporal function approximators to both solve and find parameters of basic PDEs like Burgers Equation or Schrodinger Equation. Going beyond a simple feed-forward network, Zhu et al. [301] propose an encoder-decoder for predicting transient PDEs with governing PDE constraints. Geneva et al. [89] extended this approach to deep auto-regressive dense encoder-decoders with a Bayesian framework using stochastic weight averaging to quantify uncertainty.

Qualitative mathematical properties of dynamical systems modeling have also shown promise in informing loss functions for improved prediction beyond that of the physics model. Erichson et al. [74] penalize autoencoders based on physically meaningful stability measures in dynamical systems for improved prediction of fluid flow and sea surface temperature. They showed an improved mapping of past states to future states in time in both modeling scenarios in addition to improved generalizability to new data.

Physics-based loss function terms have also been used in the discovery of governing equations. Loiseau et al. [155] uses constrained least squares [96] to incorporate energy-preserving nonlinearities or to enforce symmetries in the identified equations to the equation learning process described

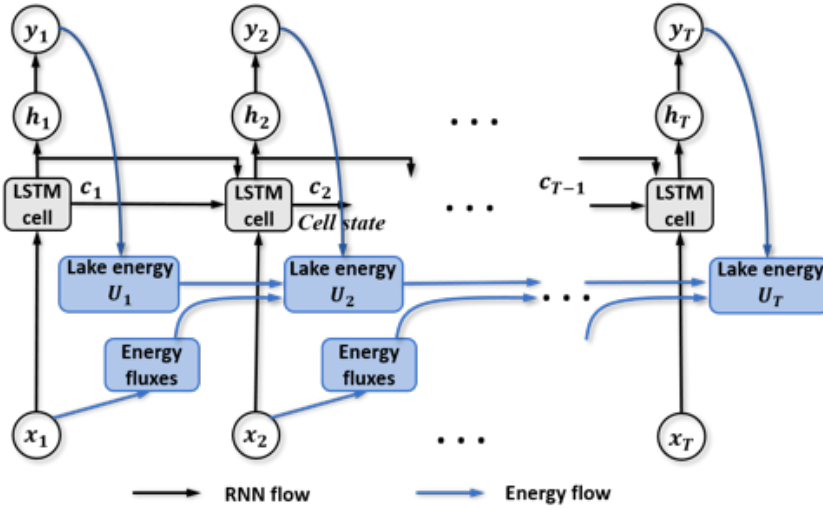


Fig. 1. The flow of the Physics-Guided Recurrent Neural Network model proposed in Jia et al. [120]. They include the standard RNN flow (black arrows) and the energy flow (blue arrows) in the recurrent process. Here  $U_T$  represents the thermal energy of the lake at time  $T$ , and both the energy output and temperature output  $y_T$  are used in calculating the loss function value. Figure adapted from Jia et al. [120].

in Section 2.7. Though these loss functions are mostly seen in common variants of NNs, they can also be seen in architectures such as echo state networks. In Doan et al. [66], they find integrating the physics-based loss from the governing equations in a Lorenz system, a commonly studied system in dynamical systems, strongly improves the echo state network's time-accurate prediction of the system and also reduces convergence time.

*Inverse modeling.* For applications in vortex induced vibrations, Raissi et al. [210] pose the inverse modeling problem of predicting the lift and drag forces of a system given sparse data about its velocity field. Kahana et al. [122] uses a loss function term pertaining to the physical consistency of the time evolution of waves for the inverse problem of identifying the location of an underwater obstacle from acoustic measurements. In both cases, the addition of physics-based loss terms made results more accurate and more robust to out-of-sample scenarios.

*Parameterization.* While ML has been used for parameterization, adding physics-based loss terms can further benefit this process by ensuring physically consistent outputs. Zhang et al. [292] parameterizes atomic energy for molecular dynamics using a NN with a loss function that, in addition to loss due to accuracy, takes into account atomic force, atomic energy, and terms relating kinetic and potential energy. Furthermore, in the context of climate modeling, Beucler et al. show that enforcing energy conservation laws improves prediction when emulating cloud processes [23, 24].

*Uncertainty quantification.* In Yang et al. [281] and Yang et al. [282], physics-based loss is implemented in a deep probabilistic generative model for uncertainty quantification based on adherence to the structure imposed by PDEs. To accomplish this, they construct probabilistic representations for the system states, and use an adversarial inference procedure to train them on data using a physics-based loss function that enforces adherence to the governing laws. This is expanded in Zhu et al. [301], where a physics-informed encoder-decoder network is defined in conjunction with a

conditional flow-based generative model for similar purposes. A similar loss function modification is performed in other works [89, 129, 278], but for the purpose of solving high dimensional stochastic PDEs with uncertainty propagation. In these cases, physics-guided constraints provide effective regularization for training deep generative models to serve as surrogates of physical systems where the cost of acquiring data is high and the data sets are small [282].

Another direction for encoding physics knowledge into ML UQ applications is to create a physics-guided Bayesian NN. This is explored in Yang et al. [277], where they use a Bayesian NN, which naturally encodes uncertainty, as a surrogate for a PDE solution. Additionally, they add a PDE constraint for the governing laws of the system to serve as a prior for the Bayesian net, allowing for more accurate predictions in situations with significant noise due to the physics-based regularization.

*Generative models.* In recent years, GANs have been used to efficiently generate solutions to PDEs and there is interest in using physics knowledge to improve them. Work by Yang et al. [279] shows GANs with loss functions based on PDEs can be used to solve stochastic elliptic PDEs in up to 30 dimensions. In a similar vein, Wu et al. [268] showed that physics-based loss functions in GANs can lower the amount of data and training time needed to converge on solutions of turbulence PDEs, while Shan et al. [231] saw similar results in the generation of microstructures satisfying certain physical properties in computational materials science.

### 3.2 Physics-Guided Initialization

Since many ML models require an initial choice of model parameters before training, researchers have explored different ways to physically inform a model starting state. For example, in NNs, weights are often initialized according to a random distribution prior to training. Poor initialization can cause models to anchor in local minima, which is especially true for deep neural networks. However, if physical or other contextual knowledge can be used to help inform the initialization of the weights, model training can be accelerated or improved [120]. One way to inform the initialization to assist in model training and escaping local minima is to use an ML technique known as *transfer learning*. In transfer learning, a model can be *pre-trained* on a related task prior to being fine-tuned with limited training data to fit the desired task. The pre-trained model serves as an informed initial state that ideally is closer to the desired parameters for the desired task than random initialization. One way to harness physics-based modeling knowledge is to use the physics-based model's simulated data to pre-train the ML model, which also alleviates data paucity issues. This is similar to the common application of pre-training in computer vision, where CNNs are often pre-trained with very large image datasets before being fine-tuned on images from the task at hand [243].

Jia *et al.* use this strategy in the context of modeling lake temperature dynamics [119, 120]. They pre-train their Physics-Guided Recurrent Neural Network (PGRNN) models for lake temperature modeling on simulated data generated from a physics-based model and fine tune the NN with little observed data. They showed that pre-training, even using data from a physical model with an incorrect set of parameters, can still significantly reduce the training data needed for a quality model. In addition, Read et al. [213] demonstrated that such models are able to generalize better to unseen scenarios than pure physics-based models.

Another application can be seen in computer vision in robotics, where images from robotics simulations have been shown to be sufficient without any real-world data for the task of object localization [248], while reducing data requirements by a factor of 50 for object grasping [31]. Then, in autonomous vehicle training, Shah et al. [230] showed that pre-training the driving algorithm in a simulator built on a video game physics engine can drastically lessen data needs. More generally,

we see that simulation pre-training of applications allows for significantly less expensive data collection than is possible with physical robots.

Another significant application of physics-guided initialization is in computational biophysics, where high computational cost of molecular dynamics simulations limits the number of mutant molecules that can be investigated from a given baseline molecule. Sultan et al. [239] showed that the molecular dynamics of a mutant protein can be estimated by training a variational autoencoder on the simulation of the protein from which the target protein mutated, allowing for surrogate models to offset the high cost of simulations. This method promises to be able to quickly sample related molecular systems to enable probing of the variation in large complex molecular structures, an otherwise very computationally expensive task. In a similar vein, in the context of predicting protein structures, Hurtado et al. [114] shows that a deep NN can be pre-trained on a large database of already-known protein structures to enhance performance.

Physics-guided model initialization has also been employed in chemical process modeling [158, 159, 276]. Yan et al. [276] uses Gaussian process regression for process modeling that has been transferred and adapted from a similar task. To adapt the transferred model, they use scale-bias correcting functions optimized through maximum likelihood estimation of parameters. Furthermore, Gaussian process models come equipped with uncertainty quantification which is also informed through initialization. A similar transfer and adapt approach is seen in Lu et al. [158], but for an ensemble of NNs transferred from related tasks. In both of the previous studies, the similarity metrics used to find similar systems are defined by considering various common process characteristics and behaviors.

### 3.3 Physics-Guided Design of Architecture

Although the physics-based loss and initialization in the previous sections helps constrain the search space of ML models during training, the ML architecture is often still a black box. In particular, they do not use any architectural properties to implicitly encode physical consistency or other desired physical properties. A recent research direction has been to construct new ML architectures that can make use of the specific characteristics of the problem being solved. . Though this section is focused largely on neural network architecture, we also include subsections on multi-task learning and structures of Gaussian processes. The modular and flexible nature of NNs in particular make them prime candidates for architecture modification. For example, domain knowledge can be used to specify node connections that capture physics-based dependencies among variables. Furthermore, incorporating physics-based guidance into architecture has the added bonus of also making the previously black box algorithm more interpretable, a desirable but typically missing feature of ML models used in physical modeling. In the following paragraphs, we discuss several contexts in which physics-guided ML architectures have been used.

*Intermediate Physical Variables.* One way to embed known physical principles into NN design is to ascribe physical meaning for certain neurons in the NN. It is also possible to declare physically relevant variables explicitly. In lake temperature modeling, Daw et al. [58] incorporate a physical intermediate variable as part of a monotonicity-preserving structure in the LSTM architecture as shown in Figure 2. This model produces physically consistent predictions in addition to appending a dropout layer to quantify uncertainty. In Muralidhar et al. [182], a similar approach is taken to insert physics-constrained variables as the intermediate variables in the convolutional neural network (CNN) architecture and achieve significant improvement over state-of-the-art physics-based models on the problem of predicting drag force on particle suspensions in moving fluids.

An additional benefit of adding physically relevant intermediate variables in an ML architecture is that they can help extract physically meaningful hidden representation that can be interpreted

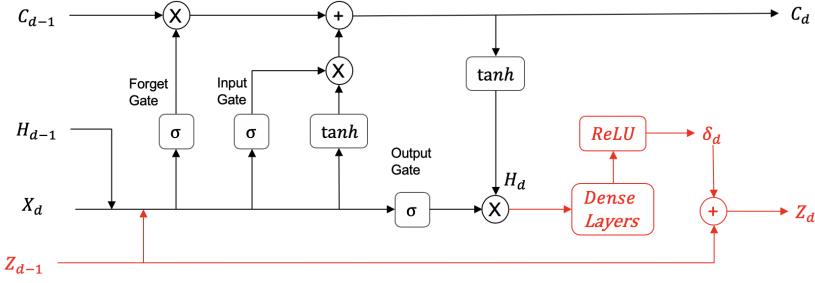


Fig. 2. Monotonicity-preserving LSTM Architecture. Components in red represent novel physics-informed innovations in LSTM.  $Z_d$  represents the density values at a specific depth  $d$ ,  $\delta_d$  is the increase of density from layer  $d - 1$  to  $d$ . The value of  $\delta$  is guaranteed to be non-negative as it is generated from an ReLU layer. Such an architecture ensures the monotonic increase of density values as depth increases. The following link contains the code for this paper: <https://github.com/arkadaw9/PNG>. Figure adapted from [58]

by domain scientists. This is particularly valuable, as standard deep learning models are limited in their interpretability since they can only extract abstract hidden variables using highly complex connected structure. This is further exacerbated given the randomness involved in the optimization process.

Another related approach is to fix one or more weights within the NN to physically meaningful values or parameters and make them non-modifiable during training. A recent approach is seen in geophysics where researchers use NNs for the waveform inversion modeling to find subsurface parameters from seismic wave data. In Sun et al. [240], they assign most of the parameters within a network to mimic seismic wave propagation during forward propagation of the NN, with weights corresponding to values in known governing equations. They show this leads to more robust training, in addition to creating a more interpretable NN with meaningful intermediate variables.

*Encoding invariances and symmetries.* In physics, there is a deep connection between symmetries and invariant quantities of a system and its dynamics. For example, Noether's Law, a common paradigm in physics, demonstrates a mapping between conserved quantities of a system and the system's symmetries (e.g. translational symmetry can be shown to correspond to the conservation of momentum within a system). Therefore, if an ML model is created that is translation-invariant, the conservation of momentum becomes more likely and the model's prediction becomes more robust and generalizable.

State-of-the-art deep learning architectures already encode certain types of invariance; for example, RNNs encode time invariance and CNNs can implicitly encode spatial translation, rotation, scale variance. In the same way, scientific modeling tasks may require other invariances based on physical laws. In turbulence modeling and fluid dynamics, Ling et al [152] defines a *tensor basis neural network* to embed the fundamental principle of rotational invariance into a NN for improved prediction accuracy. This solves a key problem in ML models for turbulence modeling because without rotational invariance, the model evaluated on identical flows with axes defined in other directions could yield different predictions. This work alters the NN architecture by adding a higher-order multiplicative layer that ensures the prediction lies on a rotationally invariant tensor basis. In a molecular dynamics application, Anderson et al. [6] show that a rotationally covariant NN architecture can learn the behavior and properties of complex many-body physical systems.

In a general setting, Wang et al. [260] show how spatiotemporal models can be made more generalizable by incorporating symmetries into deep NNs. More specifically, they demonstrated the encoding of translational symmetries, rotational symmetries, scale invariances, and uniform motion into NNs using customized convolutional layers in CNNs that enforce desired invariance properties. They also provided theoretical guarantees of invariance properties across the different designs and showed additional to significant increases in generalization performance.

Incorporating symmetries, by informing the structure of the solution space, also has the potential to reduce the search space of an ML algorithm. This is important in the application of discovering governing equations, where the space of mathematical terms and operators is exponentially large. Though in its infancy, physics-informed architectures for discovering governing equations are beginning to be investigated by researchers. In Section 2.7, symbolic regression is mentioned as an approach that has shown success. However, given the massive search space of mathematical operators, analytic functions, constants, and state variables, the problem can quickly become NP-hard. Udrescu et al. [251] designs a recursive multidimensional version of symbolic regression that uses a NN in conjunction with techniques from physics to narrow the search space. Their idea is to use NNs to discover hidden signs of "simplicity", such as symmetry or separability in the training data, which enables breaking the massive search space into smaller ones with fewer variables to be determined.

In the context of molecular dynamics applications, a number of researchers [21, 292] have used a NN per individual atom to calculate each atom's contribution to the total energy. Then, to ensure the energy invariance with respect to the possibility of interchanging two atoms, the structure of each NN and the values of each network's weight parameters are constrained to be identical for atoms of the same element. More recently, novel deep learning architectures have been proposed for fundamental invariances in chemistry. Schutt et al. [228] proposes continuous-filter convolutional (cfconv) layers for CNNs to allow for modeling objects with arbitrary positions such as atoms in molecules, in contrast to objects described by Cartesian-gridded data such as images. Furthermore, their architecture uses atom-wise layers that incorporate inter-atomic distances that enabled the model to respect quantum-chemical constraints such as rotationally invariant energy predictions as well as energy-conserving force predictions. Furthermore, Zhang et al. [293] propose an end-to-end modeling framework that preserves all natural symmetries of a molecular system using an embedding procedure that maps the input to symmetry-preserving components. This is further expanded for 3D point clouds in the "Tensor field networks" proposed by Thomas et al. [246], where they use continuous convolution layers on point cloud data, and also constrain the network filters to be the product of a learnable radial function and a spherical harmonic that make them rotation- and translation-equivariant. As we can see, because molecular dynamics often ascribes importance to different important geometric properties of molecules (e.g. rotation), network architectures dealing with invariances can be effective for improving performance and robustness of ML models.

Architecture modifications incorporating symmetry are also seen extensively in dynamical systems research involving differential equations. In a pioneering work by Ruthotto et al [218], three variations of CNNs are proposed for solving PDEs. Each variation uses mathematical theories to guide the design of the CNN based on fundamental properties of the PDEs. Multiple types of modifications are made, including adding symmetry layers to guarantee the stability expressed by the PDEs and layers that convert inputs to kinematic eigenvalues that satisfy certain physical properties. They define a parabolic CNN inspired by anisotropic filtering, a hyperbolic CNN based on Hamiltonian systems, and a second order hyperbolic CNN. Hyperbolic CNNs were found to preserve the energy in the system as intended, which set them apart from parabolic CNNs that smooth the output data, reducing the energy.

A recent direction also relating to conserved or invariant quantities is the incorporation of the Hamiltonian operator into NNs [54, 98, 249, 299]. The Hamiltonian operator in physics is the primary tool for modeling the time evolution of systems with conserved quantities, but until recently the formalism had not been integrated with NNs. Greydanus et al. [98] design a NN architecture that naturally learns and respects energy conservation and other invariance laws in simple mass-spring or pendulum systems. They accomplish this through predicting the Hamiltonian of the system and re-integrating instead of predicting the state of physical systems themselves. This is taken a step further in Toth et al. [249], where they show that not only can NNs learn the Hamiltonian, but also the abstract phase space (assumed to be known in Greydanus et al. [98]) to more effectively model expressive densities in similar physical systems and also extend more generally to other problems in physics. Recently, the Hamiltonian-parameterized NNs above have also been expanded into NN architectures that perform additional differential equation-based integration steps based on the derivatives approximated by the Hamiltonian network [51].

*Encoding other domain-specific physical knowledge.* Various other domain-specific physical information can be encoded into architecture that doesn't exactly correspond to known invariances but provides meaningful structure to the optimization process depending on the task at hand. This can take place in many ways, including using domain-informed convolutions for CNNs, additional domain-informed discriminators in GANs, or structures informed by the physical characteristics of the problem. For example, Sadoughi et al. [221] prepend a CNN with a Fast Fourier Transform layer and a physics-guided convolution layer based on known physical information pertaining to fault detection of rolling element bearings. A similar approach is used in Sturmfels et al. [238], but the added beginning layer instead serves to segment different areas of the brain for domain guidance in neuroimaging tasks. In the context of generative models, Xie et al. [274] introduce tempoGAN, which augments a general adversarial network with an additional discriminator network along with additional loss function terms that preserve temporal coherence in the generation of physics-based simulations of fluid flow. This type of approach, though found mostly in NN models, has been extended to non-NN models in Baseman et al. [17], where they introduce a physics-guided Markov Random Field that encodes spatial and physical properties of computer memory devices into the corresponding probabilistic dependencies.

A number of advances in encoding physical knowledge have been made in the context of quantum chemistry and molecular dynamics. In computational quantum chemistry, Pfau et al. [196] develop the paradigm of spin-based neural-network quantum states by defining a custom NN architecture to map electron states to the atom's wave function, which characterizes its atomic state. Key architecture components are intermediate layers in the NN to take the mean of the activation functions from each electron stream, and a final layer which applies a quantum spin-dependent linear transformation, weights the desired outputs as the solution wave function, and enforces any boundary conditions. In molecular kinetics, reduced order modeling frameworks are used for approximating the Koopman operator [171, 262]. Maradt et al. [171] describe a dual NN architecture that transforms the molecular configurations found at given time delays (e.g.  $t$  and  $t + \tau$ ) to Markov states along simulation trajectories to perform nonlinear dimensionality reduction. They show this outperforms state-of-the-art existing methods and also remains interpretable due to the mapping to realistic physical states. A similar approach is seen in [262], where they take a similar time-delay approach to improve over state-of-the-art but instead use autoencoders, which allows for the addition of a decoding step that can predict configurations in the original coordinate space from latent space points. This encoder-decoder architecture is further expanded on by Otto et al. [188], where they go beyond linear reconstruction using Koopman modes in the decoder step to an

architecture that does nonlinear reconstruction in order to learn ever lower dimensional Koopman subspaces.

Fan et al. [77] define new architectures to solve the inverse problem of electrical impedance tomography, where the goal is to determine the electrical conductivity distribution of an unknown medium from electrical measurements along its boundary. They define new NN layers based on a linear approximation of both the forward and inverse maps relying on the nonstandard form of the wavelet decomposition [25].

Architecture modifications are also seen in dynamical systems research encoding principles from differential equations. Chen et al. [48] develops a continuous depth NN based on the Residual Network [110] for solving ordinary differential equations. They change the traditionally discretized neuron layer depths into continuous equivalents such that hidden states can be parameterized by differential equations in continuous time. This allows for increased computational efficiency due to the simplification of the backpropagation step of training, and also creates a more scalable normalizing flow, an architectural component for solving PDEs. This is done by parameterizing the *derivative* of the hidden states of the NN as opposed to the states themselves. Then, in a similar application, Chang et al. [44] uses principles from the stability properties of differential equations in dynamical systems modeling to guide the design of the gating mechanism and activation functions in an RNN.

Currently, human experts have manually developed the majority of domain knowledge-encoded employed architectures, which can be a time-intensive and error-prone process. Because of this, there is increasing interest in automated neural architecture search methods [13, 73, 115]. A young but promising direction in ML architecture design is to embed prior physical knowledge into neural architecture searches. Ba et al. [12] adds physically meaningful input nodes and physical operations between nodes to the neural architecture search space to enable the search algorithm to discover more ideal physics-guided ML architectures.

*Auxiliary Task in Multi-Task Learning.* Domain knowledge can be incorporated into ML architecture as auxiliary tasks in a multi-task learning framework. Multi-task learning allows for multiple learning tasks to be solved at the same time, ideally while exploiting commonalities and differences across tasks. This can result in improved learning efficiency and predictions for one or more of the tasks. Therefore, another way to implement physics-based learning constraints is to use an auxiliary task in a multi-task learning framework. Here, an example of an auxiliary task in a multi-task framework might be related to ensuring physically consistent solutions in addition to accurate predictions. The promise of such an approach was demonstrated for a computer vision task by integrating auxiliary information (e.g. pose estimation) for facial landmark detection [297]. In this paradigm, a task-constrained loss function can be formulated to allow errors of related tasks to be back-propagated jointly to improve model generalization. Early work in a computational chemistry application showed that a NN could be trained to predict energy by constructing a loss function that had penalties for both inaccuracy *and* inaccurate energy derivatives with respect to time as determined by the surrounding energy force field [199]. In particle physics, De Oliveira et al. [62] uses an additional task for the discriminator network in a generative adversarial network (GAN) to satisfy certain properties of particle interaction for the production of jet images of particle energy.

*Physics-guided Gaussian process regression.* Gaussian process regression (GPR) [265] is a nonparametric, Bayesian approach to regression that is increasingly being used in ML applications. GPR has several benefits, including working well on small amounts of data and enabling uncertainty measurements on predictions. In GPR, first a Gaussian process prior must be assumed in the form of a mean function and a matrix-valued kernel or covariance function. One way to incorporate physical knowledge in GPR is to encode differential equations into the kernel [242]. This is a key



feature in Latent Force Models, which attempt to use equations in the physical model of the system to inform the learning from data [4, 160]. Alvarez et al. [4] draws inspiration from similar applications in bioinformatics [88, 146], which showed an increase in predictive ability in computational biology, motion capture, and geostatistics datasets. More recently, Glielmo et al. [94] propose a vectorial GPR that encodes physical knowledge in the matrix-valued kernel function. They show rotation and reflection symmetry of the interatomic force between atoms can be encoded in the Gaussian process with specific invariance-preserving covariant kernels. Furthermore, Raissi et al. [206] show that the covariance function can explicitly encode the underlying physical laws expressed by differential equations in order to solve PDEs and learn with smaller datasets.

### 3.4 Residual modeling

The oldest and most common approach for directly addressing the imperfection of physics-based models in the scientific community is residual modeling, where an ML model (usually linear regression) learns to predict the errors, or residuals, made by a physics-based model [82, 247]. A visualization is shown in Figure 3. The key concept is to learn biases of the physical model (relative to observations) and use it to make corrections to the physical model's predictions. However, one key limitation in residual modeling is their inability to enforce physics-based constraints (like in Section 3.1) because such approaches model the errors instead of the physical quantities in physics-based problems.

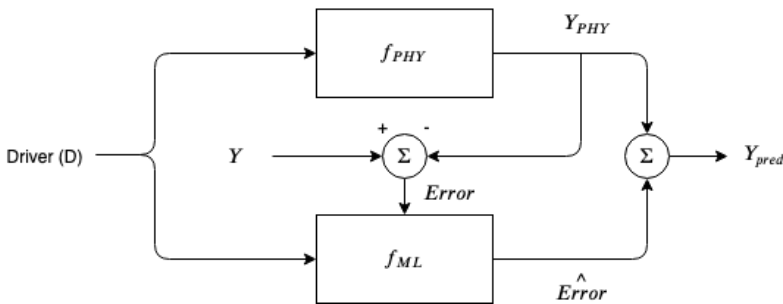


Fig. 3. Machine Learning model  $f_{ML}$  trained to model the error made by Physics model  $f_{PHY}$ . Figure adapted from [82]

Recently, a key area in which residual modeling has been applied is in reduced order models (ROMs) of dynamical systems (described in Section 2.4). After reducing model complexity to create a ROM, an ML model can be used to model the residual due to the truncation. ROM methods were created in response to the problem of many detailed simulations being too expensive to be used in various engineering tasks including design optimization and real-time decision support. In San et al. [223, 224], a simple NN used to model the error due to the model reduction is shown to sharply reduce high error regions when applied to known differential equations. Also, in Wan et al. [257], an RNN is used to model the residual between a ROM for prediction of extreme weather events and the available data projected to a reduced-order space.

As another example, in Kani et al. [125] a physics-driven "deep residual recurrent neural network (DR-RNN)" is proposed to find the residual minimiser of numerically discretized PDEs. Their architecture involves a stacked RNN embedded with the dynamical structure of the PDEs such that each layer of the NN solves a layer of the residual equations. They showed that DR-RNN sharply reduces both computational cost and time discretization error of the reduced order modeling framework.

### 3.5 Hybrid Physics-ML Models

Residual modeling is just one way to combine physics-based models with ML where both are operating simultaneously. We will call a generalized version of this *Hybrid Physics-ML Models*.

One straightforward method to combine physics-based and ML models is to feed the output of a physics-based model as input to an ML model. Karpatne et al [128] showed that using the output of a physics-based model as one feature in an ML model along with inputs used to drive the physics-based model for lake temperature modeling can improve predictions. Visualization of this method is shown in Figure 4. As we discuss below, there are multiple other ways of constructing a hybrid model, including replacing part of a larger physical model, or weighting predictions from different modeling paradigms depending on context.

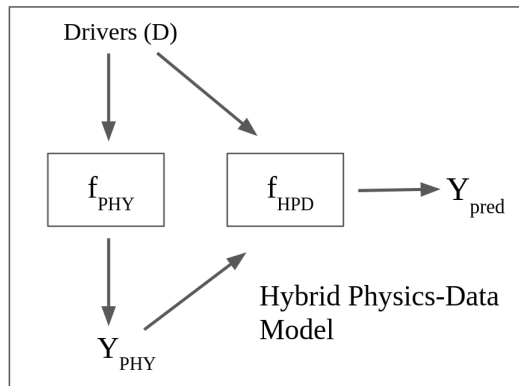


Fig. 4. Diagram of a hybrid physics-ML model. (Figure adapted from Karpatne et al. [128]).  $f_{PHY}$  represents the physics-based model which converts the input drivers  $D$  to simulated outputs  $Y_{PHY}$ ,  $f_{HPD}$  represents the hybrid physics-ML model that combines  $D$  and  $Y_{PHY}$  as input to make final predictions.

In one variant of hybrid physics-data models, ML models are used to replace one or more components of a physics-based model or to predict an intermediate quantity that is poorly modeled using physics. For example, to improve predictions of the discrepancy of Reynolds-Averaged Navier–Stokes (RANS) solvers in fluid dynamics, Parish et al. [192] propose a NN to estimate variables in the turbulence closure model to account for missing physics. They show this correction to traditional turbulence models results in convincing improvements of predictions. In Hamilton et al. [105], a subset of the mechanistic model’s equations are replaced with data-driven nonparametric methods to improve prediction beyond the baseline process model. In Zhang et al. [294], a physics-based architecture for power system state estimation embeds a deep learning model in place of traditional predicting and optimization techniques. To do this, they substitute NN layers into an unrolled version of an existing solution framework which drastically reduced the overall computational cost due to the fast forward evaluation property of NNs, but kept information of the underlying physical models of power grids and of physical constraints.

ML models for parameterization (see Section 2.3) can also be viewed as a type of hybrid modeling. Most of these efforts use black box ML models (citations covered in first column of Table 1), but some of these parameterization models can use more sophisticated physics-guided versions of ML. For example, in geodynamics modeling, Goldstein et al. [95] showed that an ML model for oscillatory flow ripples can be substituted into an established model for geologic material being moved by fluid flow. They showed that the hybrid model was able to capture dynamics previously absent from the model, which led to the discovery of two new observed pattern modes in the geology. In

chemical physics, Manzhos et al. [169] first showed that NNs could be used to represent component functions within high dimensional model representations (HDMR) of energy potentials. They show this improves the quality of the final representation and reduces the cost of the fits of component functions for HDMR traditionally done with explicit equations. Furthermore, the generality of the hybrid framework allows the HDMR to be constructed in a black box, molecule-independent way.

In another class of hybrid frameworks, the overall prediction is a combination of predictions from a physical model and an ML model, where the weights depend on prediction circumstance. For example, long-range interactions (e.g. gravity) can often be more easily modeled by classical physics equations than more stochastic short-range interactions (quantum mechanics) that are better modeled using data-driven alternatives. Hybrid frameworks like this have been used to adaptively combine ML predictions for short-range processes and physics model predictions for long-range processes for applications in chemical reactivity [284] and seismic activity prediction [191]. Estimator quality at a given time and location can also be used to determine whether a prediction comes from the physical model or the ML model, which was shown in Chen et al. [50] for air pollution estimation and in Vlachas et al. [255] for dynamical system forecasting more generally. In the context of solving PDEs, Malek et al [166] showcases a hybrid NN and traditional optimization technique to find the closed analytical form of the solution of a PDE. In this hybrid solver, there exist two terms, a term described by the NN and a term described by traditional optimization techniques.

Moreover, in inverse modeling, there is a growing use of hybrid models that first use physics-based models to perform the direct inversion, then use deep learning models to further refine the inverse model's predictions. Multiple works have shown an effective application for this in computed tomography (CT) reconstructions [38, 121]. Another common technique in inverse modeling of images (e.g. medical imaging, particle physics imaging), is the use of CNNs as *deep image priors* [252]. To simultaneously exploit data and prior knowledge, Senouf et al. [229] embed a CNN that serves as the image prior for a physics-based forward model for MRIs.

Table 1. Table of literature classified by objective and method

	Basic ML	Physics-Guided Loss Function	Physics-Guided Initialization	Physics-Guided Architecture	Residual Model	Hybrid Model
Improve prediction beyond physical model	[287] [35] [102] [104] [271]	[247] [4] [199] [139][128] [160] [183] [66] [74] [119] [153] [182] [213] [295] [109] [113] [165] [79] [285]	[158] [276] [162] [248] [31] [230] [239] [114] [119] [213] [276]	[152] [56] [17] [238] [6] [11] [183] [182] [193] [196] [221] [113] [260] [154] [293] [232] [292] [289] [291] [246] [228]	[247] [275] [258] [223] [257] [271] [153]	[95] [99] [222] [105] [128] [236] [50] [156] [191] [284] [294] [255] [280] [97] [165]
Parameterization	[198] [157] [203] [135] [169] [95] [43] [90] [186] [212] [29] [33] [189]	[292] [23] [24] [285]		[21] [23]		[294]
Reduced Order Models	[244] [179] [101] [273] [225] [286] [173]	[188] [10] [148]		[125] [262] [171] [74] [188] [190]	[125] [223] [224] [257] [101] [190]	[57]
Downscaling	[237] [254] [81] [232]					
Uncertainty quantification	[275] [142] [250] [283]	[270] [281] [266] [282] [301] [89]	[276] [162]	[58] [277] [266]		[67]
Inverse modeling	[59] [47] [161] [197]	[210] [122]		[27] [77] [240] [132]	[111]	[192] [105] [121] [241] [39] [38] [68] [229]
Discover Governing Equations	[30] [227] [37] [167] [168] [216] [123] [200][205] [52] [141] [290]	[207] [155]		[251]		
Solve PDEs	[147] [65] [140] [9] [215] [42] [264] [106] [235] [107] [131]	[209] [234] [279] [281] [60] [177] [208] [231] [301] [71] [89] [129] [268] [194] [176] [16]		[48] [218] [44] [60] [172] [235] [19] [51] [131] [76] [278] [194] [176] [206]		[166]
Data Generation	[187] [78]	[62] [40] [28] [133] [268] [288] [298]		[49] [274] [231]		

#### 4 DISCUSSION

Table 1 provides a systematic organization and taxonomy of the objectives and methods of existing physics-based ML applications. This table provides a convenient organization for papers discussed in this survey and other papers that could not be discussed because of space limitations. Importantly, analysis of works within our taxonomy uncovers knowledge gaps and potential crossovers of methods between disciplines that can serve as ideas for future research.

Indeed, there are a myriad of opportunities for taking ideas across applications, objectives, and methods, as well as bringing them back to the traditional ML discipline. For example, the physics-guided NN approach developed for aquatic sciences [109, 120] can be used in any application where an imperfect mechanistic model is available. Raissi et al. [210] take physics-guided loss function methods developed for solving PDEs and extend them to inverse modeling problems in fluid dynamics. Furthermore, Daw et al. [58] develop a monotonicity-preserving architecture for

lake temperature based on previous work done using loss function terms and a hybrid physics-ML architecture [128].

Note that some techniques for incorporating physical principles into ML models can be applied to a wide range of objectives. For example, the physics-guided loss function spans nearly all the rows of Table 1. Methods also have the potential for overlap, where, for example, physics-guided loss functions, physics-guided architecture, and physics-guided initialization could all be applied to an ML model used in a hybrid framework.

From Table 1, it is easy to see that several boxes in the Table 1 are rather sparse or entirely empty, many of which represent opportunities for future work. For example, ML models for parameterization are increasingly being used in domains such as climate science and weather forecasting [135], all of which can benefit from the integration of physical principles. Furthermore, principles from super-resolution frameworks, originally developed in the context of computer vision applications, are beginning to be applied to downscaling to create higher resolution climate predictions [254]. However, most of these do not incorporate physics (e.g., through additional loss function terms or an informed design of architecture). The fact that nearly all of the other methods (columns) except for hybrid modeling are applicable to this task shows that there is tremendous scope for new exploration, where research pursued in these columns in the context of other objectives can be applied for this objective. We also see a lot of opportunities for new research on physics-guided initialization, where, for example, pre-training an ML algorithm for inverse modeling or parameterization on simulation data related to the desired task could improve performance or lessen data needs.

We note that many boxes in Table 1 may not make sense for the pairing of an objective with a method. In particular, residual modeling (which is one of the oldest approaches for integrating physical models with statistical/machine learning models) cannot be naturally used for many of the objectives. For example, ML methods for solving PDEs often serve the purpose to reduce computation compared to an existing solving method rather than improve over a physical model, as in the case of residual modeling. Similarly, for discovering governing equations, there often isn't a known physical model to compare to for creating a residual model or for use in a hybrid approach. Data generation also does not make sense for residual modeling since the purpose is to simulate a data distribution rather than improve on a physical model.

Not all promising research directions are covered by Table 1 and the previous discussion. For instance, one promising direction is to forecast future events using ML and continuously updating model states by incorporating ideas from data assimilation [124]. An instance of this is pursued by Dua et al. [69] to build an ML model that predicts the parameters of a physical model using past time series data as an input. Another instance of this approach is seen in epidemiology, where Magri et al. [165] used a NN for data assimilation to predict a parameter vector that captures the time evolution of a COVID-19 epidemiological model. Such approaches can benefit from the ideas in Section 3 (e.g., physics-based loss, intermediate physics variables, etc.).

## 5 CONCLUDING REMARKS

Given the current deluge of sensor data and advances in ML methods, we envision that the merging of principles from ML and physics will play an invaluable role in the future of scientific modeling to address the pressing environmental and physical modeling problems facing society. Although both physics-based modeling and ML are rapidly advancing fields, knowledge integration for mutual gain requires clear lines of communication between disciplines. Currently, most of the diverse set of techniques for bringing ML and physical knowledge together have been developed in isolated applications by researchers working in distinct disciplines. Our hope is that this survey will accelerate the cross-pollination of ideas among these diverse research communities.

The contributions and structure provided in this survey also serve to benefit the ML community, where, for example, techniques for adding physical constraints to loss functions can be used to enforce fairness in predictive models, or realism for data generated by GANs. Further, novel architecture designs can enable new ways to incorporate prior domain information (beyond what is usually done using Bayesian frameworks) and can lead to better interpretability.

This survey focuses primarily on improving the modeling of engineering and environmental systems that are traditionally solved using physics-based modeling. However, the general ideas of physics-informed ML have wider applicability, and such research is already being pursued in many other contexts. For example, there are several interesting works in system control which often involves reinforcement learning techniques (e.g., combining model predictive control with Gaussian processes in robotics [7], informed priors for neuroscience modeling [91], physics-based reward functions in computational chemistry [53], and fluidic feedback control from a cylinder [134]). Other examples include identifying features of interest in the output of computational simulations of physics-based models (e.g., high-impact weather predictions [175], segmentation of climate models [181], and tracking phenomena from climate model data [85, 219]). There is also recent work on encoding domain knowledge for geometric deep learning [34, 41] that is finding increasing use in computational chemistry [61, 70, 83], physics [18, 45], geostatistics [8, 272], and neuroscience [136]. We expect that there will be a lot of potential for cross-over of ideas amongst these different efforts that will greatly expedite research in this nascent field.

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