Universal Differential Equations for Scientific Machine Learning

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August 10, 2020

Abstract

In the context of science, the well-known adage “a picture is worth a thousand words” might well be “a model is worth a thousand datasets.” Scientific models, such as Newtonian physics or biological gene regulatory networks, are human-driven simplifications of complex phenomena that serve as surrogates for the countless experiments that validated the models. Recently, machine learning has been able to overcome the inaccuracies of approximate modeling by directly learning the entire set of nonlinear interactions from data. However, without any predetermined structure from the scientific basis behind the problem, machine learning approaches are flexible but data-expensive, requiring large databases of homogeneous labeled training data. A central challenge is reconciling data that is at odds with simplified models without requiring “big data”.

In this work we develop a new methodology, universal differential equations (UDEs), which augments scientific models with machine-learnable structures for scientifically-based learning. We show how UDEs can be utilized to discover previously unknown governing equations, accurately extrapolate beyond the original data, and accelerate model simulation, all in a time and data-efficient manner. This advance is coupled with open-source software that allows for training UDEs which incorporate physical constraints, delayed interactions, implicitly-defined events, and intrinsic stochasticity in the model. Our examples show how a diverse set of computationally-difficult modeling issues across scientific disciplines, from automatically discovering biological mechanisms to accelerating the training of physics-informed neural networks and large-eddy simulations,
can all be transformed into UDE training problems that are efficiently solved by a single software methodology.

Recent advances in machine learning have been dominated by deep learning which utilizes readily available “big data” to solve previously difficult problems such as image recognition [1, 2, 3] and natural language processing [4, 5, 6]. While some areas of science have begun to generate the large amounts of data required to train deep learning models, notably bioinformatics [7, 8, 9, 10, 11], in many areas the expense of scientific experiments has prohibited the effectiveness of these ground breaking techniques. In these domains, such as aerospace engineering, quantitative systems pharmacology, and macroeconomics, mechanistic models which synthesize the knowledge of the scientific literature are still predominantly deployed due to the inaccuracy of deep learning techniques with small training datasets. While these mechanistic models are constrained to be predictive by utilizing prior structural knowledge conglomerated throughout the scientific literature, the data-driven approach of machine learning can be more flexible and allows one to drop the simplifying assumptions required to derive theoretical models. The purpose of this work is to bridge the gap by merging the best of both methodologies while mitigating the deficiencies.

It has recently been shown to be advantageous to merge differential equations with machine learning. Physics-Informed Neural Networks (PINNs) utilize partial differential equations in the cost functions of neural networks to incorporate prior scientific knowledge [12]. While this has been shown to be a form of data-efficient machine learning for some scientific applications, the resulting model does not have the interpretability of mechanistic models. On the other end of the spectrum, machine learning practitioners have begun to make use of scientific structures as a modeling basis for machine learning. For example, neural ordinary differential equations are initial value problems of the form [13, 14, 15, 16]:

\[ u' = \mathcal{N}_\theta(u, t), \]

defined by a neural network \( \mathcal{N}_\theta \) where \( \theta \) are the weights. As an example, a neural network with two hidden layers can be written as

\[ \mathcal{N}_\theta(u, t) = W_3\sigma_2(W_2\sigma_1(W_1[u; t] + b_1) + b_2) + b_3, \]

where \( \theta = (W_1, W_2, W_3, b_1, b_2, b_3) \) where \( W_i \) are matrices and \( b_i \) are vectors of weights, and \( \sigma_1, \sigma_2 \) are the choices of activation functions. Because the embedded function is a universal approximator (UA), it follows that \( \mathcal{N}_\theta \) can learn to approximate any sufficiently regular differential equation. However, the resulting model is defined without direct incorporation of known mechanisms. The Universal Approximation Theorem (UAT) demonstrates that sufficiently large neural networks can approximate any nonlinear function with a finite set of parameters [17, 18, 19]. Our approach extends the previous data-driven neural ODE approaches to directly utilize mechanistic modeling simultaneously with UAs in order to allow for arbitrary data-driven model extensions. The objects of this semi-mechanistic approach, which we denote as Universal Differential
Equations (UDEs) for universal approximators in differential equations, are differential equation models where part of the differential equation contains an embedded UA, such as a neural network, Chebyshev expansion, or a random forest.

As a motivating example, the universal ordinary differential equation (UODE):

$$u' = f(u, t, U_\theta(u, t)),$$

(3)

denotes a known mechanistic model form $f$ with missing terms defined by some UA $U_\theta$. Similar generalizations to incorporate process noise, delayed interactions, and physics-based constraints are given by embedding UAs into stochastic, delay, and differential-algebraic equations respectively. In Section 1 we describe our methodology and software implementation for efficiently training UDEs of any of these forms in a way that covers stiffness, nonlinear algebraic constraints, stochasticity, delays, parallelism, and more. We then demonstrate that the following abilities within the UDE framework:

- In Section 2.1 we recover governing equations from much lesser data than prior methods and demonstrate the ability to accurately extrapolate from a short time series.
- In Section 2.2 we demonstrate the ability to utilize arbitrary conservation laws as prior knowledge in the discovery of dynamical systems.
- In Section 2.3 we discover the differential operator and nonlinear reaction term of a biological partial differential equation (PDE) from spatiotemporal data, demonstrating the interpretability of trained UDEs.
- In Section 3 we derive an adaptive method for automated solving of a 100-dimensional nonlinear Hamilton-Jacobi-Bellman PDE, the first adaptive method for this class of problems that the authors are aware of.
- In Section 4.1 we automate the discovery of fast, accurate, and physically-consistent surrogates to accelerate a large-eddy simulation commonly used in the voxels of a climate simulation.
- In Section 4.2 we approximate closure relations in viscoelastic fluids to accelerate the simulation of a system of 6 differential-algebraic equations by 2x, showing that this methodology is also applicable to small-scale problems.
- In Section 4.3 we demonstrate that discrete physics-informed neural networks fall into a subclass of universal ODEs and extend previous methods directly through this formalism.
1 Efficient Training of Universal Differential Equations via Differentiable Programming

Training a UDE amounts to minimizing a cost function $C(θ)$ defined on the current solution $u_θ(t)$, the current solution to the differential equation with respect to the choice of parameters $θ$. One choice is the Euclidean distance $C(θ) = \sum_i \|u_θ(t_i) - d_i\|$ at discrete data points $(t_i, d_i)$. When optimized with local derivative-based methods, such as stochastic gradient decent, ADAM [20], or L-BFGS [21], this requires the calculation of $\frac{dC}{dθ}$ which by the chain rule amounts to calculating $\frac{du}{dθ}$. Thus the problem of efficiently training a UDE reduces to calculating gradients of the differential equation solution with respect to parameters.

In certain special cases there exist efficient methods for calculating these gradients called adjoints [22, 23, 24, 25, 26]. The asymptotic computational cost of these methods does not grow multiplicatively with the number of state variables and parameters like numerical or forward sensitivity approaches, and thus it has been shown empirically that adjoint methods are more efficient on large parameter models [27, 28]. However, given the number of different families of UDE models we wish to train, we generalize to a differentiable programming framework with reverse-mode accumulation in order to allow for deriving on-the-fly approximations for the wide range of possible differential equation types.

Given a function $f(x) = y$, the pullback at $x$ is the function:

$$B^*_f(y) = y^T f'(x),$$

where $f'(x)$ is the Jacobian $J$. We note that $B^*_f(1) = (\nabla f)^T$ for a function $f$ producing a scalar output, meaning the pullback of a cost function computes the gradient. A general computer program can be written as the composition of discrete steps:

$$f = f^L \circ f^{L-1} \circ \ldots \circ f^1,$$

and thus the vector-Jacobian product can be decomposed:

$$v^T J = (...((v^T J_L) J_{L-1}) \ldots) J_1,$$

which allows for recursively decomposing a the pullback to a primitively known set of $B^*_f$:

$$B^*_f(A) = B^*_f\left(\ldots B^{z_{L-2}}_f \left( B^{z_{L-1}}_f( A) \right) \ldots \right),$$

where $x_i = (f^i \circ f^{i-1} \circ \ldots \circ f^1)(x)$. Implementations of code generation for the backwards pass of an arbitrary program in a dynamic programming language can vary. For example, building a list of function compositions (a tape) is provided by libraries such as Tracker.jl [29] and PyTorch [30], while other libraries perform direct generation of backward pass source code such as Zygote.jl [31], TAF [32], and Tapenade [33].
The open-source differential equation solvers of DifferentialEquations.jl [34] were developed in a manner such that all steps of the programs have a well-defined pullback when using a Julia-based backwards pass generation system. Our software allows for automatic differentiation to be utilized over differential equation solves without any modification to the user code. This enables the simulation software already written with DifferentialEquations.jl, including large software infrastructures such as the MIT-CalTech CLiMA climate modeling system [35] and the QuantumOptics.jl simulation framework [36], to be compatible with all of the techniques mentioned in the rest of the paper. Thus while we detail our results in isolation from these larger simulation frameworks, the UDE methodology can be readily used in full-scale simulation packages which are already built on top of the Julia SciML ecosystem.

The full set of adjoint options, which includes continuous adjoint methods and pure reverse-mode AD approaches, is described in Supplement S??. Methods via solving ODEs in reverse [16] are the common adjoint utilized in neural ODE software such as torchdiffeq and are O(1) in memory, but are known to be unstable under certain conditions such as on stiff equations [37]. Checkpointed interpolation adjoints [25] and continuous quadrature approaches are available which do not require stable reversibility of the ODEs while retaining a relatively low-memory implementation via checkpointing (in particular Section 2.2 and 4.1 are noted as a cases which are not stable under the reversed adjoint but stable under the checkpointing adjoint approach). These adjoint methods fall under the continuous optimize-then-discretize approach. Through the differentiable programming integration, discrete adjoint sensitivity analysis [38, 39] is implemented through both tape-based reverse-mode [40] and source-to-source translation [31], with computational trade-offs between the two approaches. The former can be faster on scalarized heterogeneous differential equations while the latter is more optimized for homogeneous vectorized functions calls like are demonstrated in neural networks and discretizations of partial differential equations. Full discretize-then-optimize is implemented using this package by utilizing the step-wise integrator interface in conjunction with these discrete adjoints of the steps. Continuous and discrete forward mode sensitivity analysis approaches are also provided and optimized for equations with smaller numbers of parameters.

Previous research has shown that the discrete adjoint approach is more stable than continuous adjoints in some cases [41, 37, 42, 43, 44, 45] while continuous adjoints have been demonstrated to be more stable in others [46, 43] and can reduce spurious oscillations [47, 48, 49]. This trade-off between discrete and continuous adjoint approaches has been demonstrated on some equations as a trade-off between stability and computational efficiency [50, 51, 52, 53, 54, 55, 56, 57, 58]. Care has to be taken as the stability of an adjoint approach can be dependent on the chosen discretization method [59, 60, 61, 62, 63], and our software contribution helps researchers switch between all of these optimization approaches in combination with hundreds of differential equation solver methods with a single line of code change.

As described in Supplement S??, these adjoints utilize reverse-mode auto-
matic differentiation for vector transposed Jacobian products within the adjoint definitions to reduce the computational complexity while supporting advanced features like constraint and conservation equations. In addition, the module DiffEqFlux.jl handles compatibility with the Flux.jl neural network library so that these vector Jacobian products are automatically replaced with efficient pullback implementations for embedded deep neural networks (also known as backpropogation) wherever neural networks are encountered in the right hand side of any differential equation definitions. This allows for common deep architectures, such as convolutional neural networks and recurrent neural networks, to be efficiently used as the basis for a UDE without any Jacobians being calculated in the full adjoint and without requiring any intervention from users.

Using this approach, the solvers are capable of building efficient gradient calculations for training ML-embedded UDEs of the classes:

- Universal Ordinary Differential Equations (UODEs)
- Universal Stochastic Differential Equations (USDEs), or universal differential equations with continuous process noise
- Universal Delay Differential Equations (UDDEs), or universal differential equations with delayed interactions
- Universal Differential-Algebraic Equations (UDAEs), or universal differential equations with constraint equations and conservation laws
- Universal Boundary Value Problems (UBVPs), or universal differential equations with final time point constraints
- Universal Partial Differential Equations (UPDEs)
- Universal Hybrid (Event-Driven) Differential Equations

as well as the combinations, such as stochastic delay differential equations, jump diffusions, and stochastic partial differential equations. A combination of over 300 solver methods cover the efficient training of stiff and non-stiff versions of each of these equations, with support for adaptivity, high-order, automatic stiffness detection, sparse differentiation with automatic sparsity detection and coloring [64], Newton-Krylov implicit handling, GPU compatibility, and multi-node parallelism via MPI compatibility. Thus together, semi-mechanistic UDEs of any form can embed machine learning models and be trained using this open-source library with the most effective differential equation solvers for that class of equations.

1.1 Features and Performance

We assessed the viability of alternative differential equation libraries for universal differential equation workflows by comparing the features and performance of the given libraries. Table 1 demonstrates that the Julia SciML ecosystem is
Table 1: Feature comparison of ML-augmented differential equation libraries. 
First first column corresponds to support for stiff ODEs, then DAEs, SDEs, DDEs, stabilized non-reversing adjoints, discretize-then-optimize methods, distributed computing, and multithreading. Sparse refers to automated sparsity handling in Jacobian calculations of implicit methods.

| Feature | Stiff | DAEs | SDEs | DDEs | Stabilized | DtO | GPU | Dist | MT | Sparse |
|---------|-------|------|------|------|------------|-----|-----|------|-----|--------|
| SciML   | ✓     | ✓    | ✓    | ✓    | ✓          | ✓   | ✓   | ✓    | ✓   | ✓      |
| torchdiffeq | 0  | 0    | 0    | 0    | 0          | ✓   | ✓   | 0    | 0   | 0      |
| torchsde | 0  | 0    | ✓    | 0    | 0          | ✓   | 0   | 0    | 0   | 0      |
| tfdiffeq | 0  | 0    | 0    | 0    | 0          | ✓   | 0   | 0    | 0   | 0      |

the only differential equation solver library with deep learning integration that supports stiff ODEs, DAEs, DDEs, stabilized adjoints, distributed and multithreaded computation. We note the importance of the stabilized adjoints in Section 4.1 as many PDE discretizations with upwinding exhibit unconditional instability when reversed, and thus this is a crucial feature when training embedded neural networks in many PDE applications. Table 2 demonstrates that the SciML ecosystem exhibits more than an order of magnitude performance when solving ODEs against torchdiffeq of up to systems of 1 million equations. Because the adjoint calculation itself is a differential equation, this also corresponds to increased training times on scientific models. To reinforce this result, Supplement S?? demonstrates a 100x performance difference over torchdiffeq when training the spiral neural ODE from [16, 41]. We note that the author of the tfdiffeq library has previous concluded “speed is almost the same as the PyTorch (torchdiffeq) codebase (±2%)”. Additionally, Supplement S?? demonstrates a 1,600x performance advantage for the SciML ecosystem over torchsde using the geometric Brownian motion example from the torchsde documentation [65]. Given the computational burden, the mix of stiffness, and non-reversibility of the examples which follow in this paper, these results demonstrate that the SciML ecosystem is the first deep learning integrated differential equation software ecosystem that can train all of the equations necessary for the results of this paper. Note that this does not infer that our solvers will demonstrate more than an order of magnitude performance difference on all equations, for example very non-stiff ODEs dominated by large dense matrix multiplications like in image classification neural ODEs, but it does demonstrate that on the equations generally derived from scientific models (ODEs derived from PDE discretizations, heterogeneous differential equation systems, and neural networks in sufficiently small systems) that an order of magnitude or more performance difference can exist.
Table 2: Relative time to solve for ML-augmented differential equation libraries (smaller is better). Standard non-stiff solver benchmarks from representative scientific systems were taken from [66] as described in Supplement S??, SciML stands for the optimal method choice out of the 300+ from the SciML, which for the first is DP5, for the second is VCABM, and for the rest is ROCK4.

| # of ODEs | SciML | SciML DP5 | torchdiffeq dopri5 | torchdiffeq adams |
|-----------|-------|-----------|--------------------|-------------------|
| 3         | 1.0x  | 1.0x      | 5,850x             | 7,600x            |
| 28        | 1.0x  | 1.0x      | 1.9x               | 1100x             |
| 768       | 1.0x  | 2.9x      | 2.9x               | 710x              |
| 3,072     | 1.0x  | 2.9x      | 2.9x               | 490x              |
| 12,288    | 1.0x  | 2.9x      | 420x               | 490x              |
| 49,152    | 1.0x  | 3.4x      | 31x                | 170x              |
| 196,608   | 1.0x  | 2.9x      | 41x                | 44x               |
| 786,432   | 1.0x  | 2.6x      | 38x                | 43x               |

2 Knowledge-Enhanced Model Reconstruction of Biological Models

Automatic reconstruction of models from observable data has been extensively studied. Many methods produce non-symbolic representations by learning functional representations [67, 68] or through dynamic mode decomposition (DMD, eDMD) [69, 70, 71, 72]. Symbolic reconstruction of equations has utilized symbolic regressions which require a prechosen basis [73, 74], or evolutionary methods to grow a basis [75, 76]. However, a common thread throughout much of the literature is that added domain knowledge constrains the problem to allow for more data-efficient reconstruction [77, 78]. Here we detail how a UA embedded workflow can augment existing symbolic regression frameworks to allow for reconstruction from partially known models in a more data-efficient manner.

2.1 Automated Identification of Nonlinear Interactions with Universal Ordinary Differential Equations

The SInDy algorithm [79, 80, 81] finds a sparse basis $\Xi$ over a given candidate library $\Theta$ minimizing the objective function $\|\dot{X} - \Theta \Xi\|_2 + \lambda \|\Xi\|_1$ using data for $\dot{X}$ generated by interpolating the trajectory data $X$. Here we describe a UDE approach to extend SInDy in a way that embeds prior structural knowledge.

As a motivating example, take the Lotka-Volterra system:

$$\dot{x} = \alpha x - \beta xy,$$
$$\dot{y} = \gamma xy - \delta y.$$  \hspace{1cm} (8)

Assume that a scientist has a short time series from this system but knows the birth rate of the prey $x$ and the death rate of the predator $y$. With only this information, a scientist can propose the knowledge-based UODE as:

$$\dot{x} = \alpha x + U_1(x,y),$$
$$\dot{y} = -\delta y + U_2(x,y),$$  \hspace{1cm} (9)
Figure 1: Automated Lotka-Volterra equation discovery with UODE-enhanced SInDy. (A) The error in the trained UODE against $x(t)$ and $y(t)$ in green and yellow respectively. (B) The measured values of the missing term $U_2(x, y)$ throughout the time series, with the neural network approximate in green and the true value $\gamma_{xy}$ in yellow. (C) The extrapolation of the knowledge-enhanced SInDy fit series. The green and yellow dots show the data that was used to fit the UODE, and the dots show the true solution of the Lotka-Volterra Equations 8 beyond the training data. The blue and purple lines show the extrapolated solution how the UODE-enhanced SInDy recovered equations.

which is a system of ordinary differential equations that incorporates the known structure but leaves room for learning unknown interactions between the the predator and prey populations. Learning the unknown interactions corresponds training the UA $U : \mathbb{R}^2 \rightarrow \mathbb{R}^2$ in this UODE.

While the SInDy method normally approximates derivatives using a spline over the data points or similar numerical techniques, here we have $U_\theta(x, y)$ as an estimator of the derivative for only the missing terms of the model and we can perform a sparse regression on samples from the trained $U_\theta(x, y)$ to reconstruct only the unknown interaction equations. As described in Supplement S??, we trained $U_\theta(x, y)$ as a neural network against the simulated data for $t \in [0, 3]$ and utilized a sparse regression techniques [79, 82, 83] on the neural network outputs to reconstruct the missing dynamical equations. Using a 10-dimensional polynomial basis extended with trigonometric functions, the sparse regression yields 0 for all terms except for the missing quadratic terms, directly learning the original equations in an interpretable form. Even though the original data
did not contain a full period of the cyclic solution, the resulting fit is then able to accurately extrapolate from the short time series data as shown in Figure 1. Supplement S?? further demonstrates the robustness of the discovery approach to noise in the data. Likewise, when attempting to learn full ODE with the original SInDy approach on the same trained data with the analytical derivative values, we were unable to recover the exact original equations from the sparse regression, indicating that the knowledge-enhanced approach increases the robustness of equation discovery.

We note that collaborators using the preprint of this manuscript have successfully demonstrated the ability to construct UODE models which improve the prediction of Li-ion battery performance [84] and for automated discovery of droplet physics directly from imaging data, effectively replicating the theoretical results of a one and a half year study with a UODE discovery process which trains in less than an hour [85].

2.2 Incorporating Prior Knowledge of Conservation Equations

The extra features of the SciML ecosystem can be utilized to encode more information into the model. For example, when attempting to discover a biological chemical reaction network or a chemical combustion network, one may only have prior knowledge of the conservation laws between the constituent substrates. As a demonstration, in the Robertson equation

\begin{align}
\frac{dy_1}{dt} &= -0.04y_1 + 10^4 y_2 y_3 \\
\frac{dy_2}{dt} &= 0.04y_1 - 10^4 y_2 y_3 - 3 \times 10^7 y_2^2 \\
1 &= y_1 + y_2 + y_3
\end{align}

one might only have prior knowledge of the conservation equation \(1 = y_1 + y_2 + y_3\). In this case, a universal DAE of the form:

\begin{align}
\frac{d[y_1, y_2]}{dt} &= U_\theta(y_1, y_2, y_3) \\
1 &= y_1 + y_2 + y_3
\end{align}

can be utilized to encode this prior knowledge. This can then be trained by utilizing a singular mass matrix in the form \(Mu' = f(u, p, t)\). Supplement S??’s derivation of the adjoint method describes a new initialization scheme for index-1 DAEs in mass matrix form which directly solves a linear system for new consistent algebraic variables in the adjoint pass without requiring the approximate nonlinear iterations of [86], thus further demonstrating the efficiency and accuracy of the SciML software’s methods for UDE workflows. Supplement S?? demonstrates the ability to learn this system utilizing the SciML tools through this universal DAE approach.
2.3 Reconstruction of Spatial Dynamics with Universal Partial Differential Equations

To demonstrate discovery of spatiotemporal equations directly from data, we consider data generated from the one-dimensional Fisher-KPP (Kolmogorov–Petrovsky–Piskunov) PDE [87]:

\[
\frac{\partial \rho}{\partial t} = r \rho (1 - \rho) + D \frac{\partial^2 \rho}{\partial x^2}, \tag{15}
\]

with \( x \in [0, 1] \), \( t \in [0, T] \), and periodic boundary condition \( \rho(0, t) = \rho(1, t) \). Here \( \rho \) represents population density of a species, \( r \) is the local growth rate and \( D \) is the diffusion coefficient. Such reaction-diffusion equations appear in diverse physical, chemical and biological problems [88]. To learn the generated data, we define the UPDE:

\[
\rho_t = \text{NN}_\theta(\rho) + \hat{D} \text{CNN}(\rho), \tag{16}
\]

where \( \text{NN}_\theta \) is a neural network representing the local growth term. The derivative operator is approximated as a convolutional neural network CNN, a learnable arbitrary representation of a stencil while treating the coefficient \( \hat{D} \) as an unknown. We encode in the loss function extra constraints to ensure the learned equation is physically realizable, i.e. the derivative stencil must be conservative (the coefficients sum to zero), as described in Supplement S??.

Figure 2 shows the result of training the UPDE against the simulated data, which recovers the canonical \([1, -2, 1]\) stencil of the one-dimensional Laplacian and the diffusion constant while simultaneously finding a neural representation of the unknown quadratic growth term. We note that the differentiable programming integration in conjunction with the Flux.jl deep learning framework allows for the adjoints to automatically utilize efficient backpropagation of the embedded convolutional neural networks and automatically utilizes the fast kernels provided by cudnn when trained using GPUs.

3 Computationally-Efficient Solving of High-Dimensional Partial Differential Equations

It is impractical to solve high dimensional PDEs with mesh-based techniques since the number of mesh points scales exponentially with the number of dimensions. Given this difficulty, mesh-free methods based on universal approximators such as neural networks have been constructed to allow for direct solving of high dimensional PDEs [89, 90]. Recently, methods based on transforming partial differential equations into alternative forms, such as backwards stochastic differential equations (BSDEs), which are then approximated by neural networks have been shown to be highly efficient on important equations such as the nonlinear Black-Scholes and Hamilton-Jacobi-Bellman (HJB) equations [91, 92, 93, 94].
Figure 2: Recovery of the UPDE for the Fisher-KPP equation. (A) Training data and (B) prediction of the UPDE for $\rho(x,t)$. (C) Curves for the weights of the CNN filter $[w_1, w_2, w_3]$ indicate the recovery of the $[1, -2, 1]$ stencil for the 1-dimensional Laplacian. (D) Comparison of the learned (blue) and the true growth term (orange) showcases the learned parabolic form of the missing nonlinear equation.
Here we will showcase how one of these methods, a deep BSDE method for semi-linear parabolic equations [92], can be reinterpreted as a universal stochastic differential equation (USDE) to generalize the method and allow for enhancements like adaptivity, higher order integration for increased efficiency, and handling of stiff driving equations through the SciML software.

Consider the class of semilinear parabolic PDEs with a finite time span $t \in [0, T]$ and $d$-dimensional space $x \in \mathbb{R}^d$ that have the form:

$$
\frac{\partial u}{\partial t}(t, x) + \frac{1}{2} \text{Tr} \left( \sigma \sigma^T (t, x) (\text{Hess}_x u) (t, x) \right) + \nabla u(t, x) \cdot \mu(t, x) + f \left( t, x, u(t, x), \sigma^T (t, x) \nabla u(t, x) \right) = 0,
$$

(17)

with a terminal condition $u(T, x) = g(x)$. Supplement S?? describes how this PDE can be solved by approximating by approximating the FBSDE:

$$
\begin{align*}
&dX_t = \mu(t, X_t) dt + \sigma(t, X_t) dW_t, \\
&dU_t = f \left( t, X_t, U_t, U^1_{\theta_1}(t, X_t) \right) dt + \left[ U^1_{\theta_1}(t, X_t) \right]^T dW_t,
\end{align*}
$$

(18)

where $U^1_{\theta_1}$ and $U^2_{\theta_2}$ are UAs and the loss function is given by the requiring that the terminating condition $g(X_T) = u(X_T, W_T)$ is satisfied.

3.1 Adaptive Solution of High-Dimensional Hamilton-Jacobi-Bellman Equations

A fixed time step Euler-Maryumana discretization of this USDE gives rise to the deep BSDE method [92]. However, this form as a USDE generalizes the approach in a way that makes all of the methodologies of our USDE training library readily available, such as higher order methods, adaptivity, and implicit methods for stiff SDEs. As a motivating example, consider the classical linear-quadratic Gaussian (LQG) control problem in 100 dimensions:

$$
dX_t = 2\sqrt{\lambda} c_t dt + \sqrt{2} dW_t,
$$

(19)

with $t \in [0, T]$, $X_0 = x$, and with a cost function $C(c_t) = E \left[ \int_0^T \|c_t\|^2 dt + g(X_T) \right]$ where $X_t$ is the state we wish to control, $\lambda$ is the strength of the control, and $c_t$ is the control process. Minimizing the control corresponds to solving the 100-dimensional HJB equation:

$$
\frac{\partial u}{\partial t} + \nabla^2 u - \lambda \| \nabla u \|^2 = 0
$$

(20)

We solve the PDE by training the USDE using an adaptive Euler-Maruyama method [95] as described in Supplement S??. Supplementary Figure ?? showcases that this methodology accurately solves the equations, effectively extending recent algorithmic advancements to adaptive forms simply be reinterpreting
the equation as a USDE. While classical methods would require an amount of memory that is exponential in the number of dimensions making classical adaptively approaches infeasible, this approach is the first the authors are aware of to generalize the high order, adaptive, highly stable software tooling to the high-dimensional PDE setting.

4 Accelerated Scientific Simulation with Automatically Constructed Closure Relations

4.1 Automated Discovery of Large-Eddy Model Parameterizations

As an example of directly accelerating existing scientific workflows, we focus on the Boussinesq equations [96]. The Boussinesq equations are a system of 3+1-dimensional partial differential equations acquired through simplifying assumptions on the incompressible Navier-Stokes equations, represented by the system:

\[ \nabla \cdot u = 0, \]
\[ \frac{\partial u}{\partial t} + (u \cdot \nabla) u = -\nabla p + \nu \nabla^2 u + b \hat{z}, \]
\[ \frac{\partial T}{\partial t} + u \cdot \nabla T = \kappa \nabla^2 T, \]

(21)

where \( u = (u, v, w) \) is the fluid velocity, \( p \) is the kinematic pressure, \( \nu \) is the kinematic viscosity, \( \kappa \) is the thermal diffusivity, \( T \) is the temperature, and \( b \) is the fluid buoyancy. We assume that density and temperature are related by a linear equation of state so that the buoyancy \( b \) is only a function \( b = \alpha g T \) where \( \alpha \) is the thermal expansion coefficient and \( g \) is the acceleration due to gravity.

This system is commonly used in climate modeling, especially as the voxels for modeling the ocean [97, 98, 99, 96] in a multi-scale model that approximates these equations by averaging out the horizontal dynamics \( T(z, t) = \int \int T(x, y, z, t) dx dy \) in individual boxes. The resulting approximation is a local advection-diffusion equation describing the evolution of the horizontally-averaged temperature \( \overline{T} \):

\[ \frac{\partial \overline{T}}{\partial t} + \frac{\partial \overline{wT}}{\partial z} = \kappa \frac{\partial^2 \overline{T}}{\partial z^2}. \]

(22)

This one-dimensional approximating system is not closed since \( \overline{wT} \) is unknown. Common practice closes the system by manually determining an approximating \( \overline{wT} \) from ad-hoc models, physical reasoning, and scaling laws. However, we can utilize a UDE-automated approach to learn such an approximation from data. Let

\[ \overline{wT} = U_\theta \left( P, \overline{T}, \frac{\partial \overline{T}}{\partial z} \right) \]

(23)
where $P$ are the physical parameters of the Boussinesq equation at different regimes of the ocean, such as the amount of surface heating or the strength of the surface winds [100]. We can accurately capture the non-locality of the convection in this term by making the UDE a high-dimensional neural network. Using data from horizontal average temperatures $T$ with known physical parameters $P$, we can directly reconstruct a nonlinear $P$-dependent parameterization by training a universal diffusion-advection partial differential equation. Supplementary Figure ?? demonstrates the accuracy of the approach using a deep UPDE with high order stabilized-explicit Runge-Kutta (ROCK) methods where the fitting is described in Supplement S??. To contrast the trained UPDE, we directly simulated the 3D Boussinesq equations under similar physical conditions and demonstrated that the neural parameterization results in around a 15,000x acceleration. This demonstrates that physical-dependent parameterizations for acceleration can be directly learned from data utilizing the previous knowledge of the averaging approximation and mixed with a data-driven discovery approach.

4.2 Data-Driven Nonlinear Closure Relations for Model Reduction in Non-Newtonian Viscoelastic Fluids

All continuum materials satisfy conservation equations for mass and momentum. The difference between an elastic solid and a viscous fluid comes down to the constitutive law relating the stresses and strains. In a one-dimensional system, an elastic solid satisfies $\sigma = G\gamma$, with stress $\sigma$, strain $\gamma$, and elastic modulus $G$, whereas a viscous fluid satisfies $\sigma = \eta \dot{\gamma}$, with viscosity $\eta$ and strain rate $\dot{\gamma}$. Non-Newtonian fluids have more complex constitutive laws, for instance when stress depends on the history of deformation,

$$\sigma(t) = \int_{-\infty}^{t} G(t-s)F(\dot{\gamma}(s)) \, ds, \quad (24)$$

alternatively expressed in the instantaneous form [101]:

$$\sigma(t) = \phi_1(t),$$
$$\frac{d\phi_1}{dt} = G(0)F(\dot{\gamma}) + \phi_2, \quad (25)$$
$$\frac{d\phi_2}{dt} = \frac{dG(0)}{dt}F(\dot{\gamma}) + \phi_3,$$
$$\vdots$$

where the history is stored in $\phi_i$. To become computationally feasible, the expansion is truncated, often in an ad-hoc manner, e.g. $\phi_n = \phi_{n+1} = \cdots = 0$, for some $n$. Only with a simple choice of $G(t)$ does an exact closure condition exist, e.g. the Oldroyd-B model. For a fully nonlinear approximation, we train a UODE according to the details in Supplement S?? to learn a closure relation:
\[ \sigma(t) = U_0(\dot{\gamma}, \phi_1, \ldots, \phi_N), \quad (26) \]
\[ \frac{d\phi_i}{dt} = U_i(\dot{\gamma}, \phi_1, \ldots, \phi_N), \quad \text{for } i = 1 \text{ to } N \quad (27) \]

from the numerical solution of the FENE-P equations, a fully non-linear constitutive law requiring a truncation condition [102]. Figure 3 compares the neural network approach to a linear, Oldroyd-B like, model for \( \sigma \) and showcases that the nonlinear approximation improves the accuracy by more than 50x. We note that the neural network approximation accelerates the solution by 2x over the original 6-state DAE, demonstrating that the universal differential equation approach to model acceleration is not just applicable to large-scale dynamical systems like PDEs but also can be effectively employed to accelerate small scale systems.

### 4.3 Efficient Discrete Physics-Informed Neural Networks as Universal ODEs

To further demonstrate the breadth of computational problems covered by the UODE framework, we note that the discrete physics-informed neural networks can be cast into the framework of UODEs. A physics-informed neural network is the representation of a PDE’s solution via a neural network, allowing machine learning training techniques to solve the equation [12]. These works note that the continuous PDE can be discretized in a single dimension to give rise to the discrete physics-informed neural network, simplified as:

\[
u^{n+c_i} = u^n - \Delta t \sum_{j=1}^{q} a_{ij} \mathcal{N}[u^{n+c_j}] \]  
\[
u^{n+1} = u^n - \Delta t \sum_{j=1}^{q} b_j \mathcal{N}[u^{n+c_j}] \]  

These results have demonstrated that the discrete form can enhance the computational efficiency of training physics-informed neural networks. However, we note that this directly corresponds to training the universal ODE \( u' = \mathcal{N}(u) \) using an explicit or implicit Runge-Kutta method in the SciML ecosystem. This directly gives rise to the further work on multistep discrete physics-informed neural networks [68, 78] by training the UODE via a multistep method, but also immediately gives the generalization to Runge-Kutta-Chebyshev, Rosenbrock, exponential integrator, and more formalizations which all are available via the SciML tools.

### 5 Discussion

While many attribute the success of deep learning to its blackbox nature, the key advances in deep learning applications have come by developing new architectures which directly model the structures that are attempting to be learned.
Figure 3: Convergence of neural closure relations for a non-Newtonian Fluid. (A) Error between the approximated $\sigma$ using the linear approximation Equation 7 and the neural network closure relation Equation 26 against the full FENE-P solution. The error is measured for the strain rates $\dot{\gamma} = 12 \cos \omega t$ for $\omega = 1, 1.2, \ldots, 2$ and tested with the strain rate $\dot{\gamma} = 12 \cos 1.5 t$. (B) Predictions of stress for testing strain rate for the linear approximation and UODE solution against the exact FENE-P stress.
For example, deep convolutional neural networks for image processing directly utilized the local spatial structure of images by modeling convolution stencil operations. Similarly, recurrent neural networks encode a forward time progression into a deep learning model and have excelled in natural language processing and time series prediction. Here we present a software that allows for combining existing scientific simulation libraries with neural networks to train and augment known models with data-driven components. Our results show that by building these hybrid mechanistic models with machine learning, we can arrive at similar efficiency advancements by utilizing all known prior knowledge of the underlying problem's structure. While we demonstrate the utility of UDEs in equation discovery, we have also demonstrated that these methods are capable of solving many other problems, and many methods of recent interest, such as discrete physics-informed neural networks, fall into the class of UDE methods and can thus be analyzed and efficiently computed as part of this formalization.

Our software implementation is the first deep learning integrated differential equation library to include the full spectrum of adjoint sensitivity analysis methods that is required to both efficiently and accurately handle the range of training problems that can arise from universal differential equations. We have demonstrated orders of magnitude performance advantages over previous machine learning enhanced adjoint sensitivity ODE software in a variety of scientific models and demonstrated generalizations to stiff equations, DAEs, SDEs, and more. While the results of this paper span many scientific disciplines and incorporate many different modeling approaches, together all of the examples shown in this manuscript can be implemented using the SciML software ecosystem in just hundreds of lines of code each, with none of the examples taking more than half an hour to train on a standard laptop. This both demonstrates the efficiency of the software and its methodologies, along with the potential to scale to much larger applications.

The code for reproducing the computational experiments can be found at:

https://github.com/ChrisRackauckas/universal_differential_equations

6 Acknowledgements

We thank Jesse Bettencourt, Mike Innes, and Lyndon White for being instrumental in the early development of the DiffEqFlux.jl library, Tim Besard and Valentin Churavy for the help with the GPU tooling, and David Widmann and Kanav Gupta for their fundamental work across DifferentialEquations.jl. Special thanks to Viral Shah and Steven Johnson who have been helpful in the refinement of these ideas. We thank Charlie Strauss, Steven Johnson, Nathan Urban, and Adam Gerlach for enlightening discussions and remarks on our manuscript and software. We thank Stuart Rogers for his careful read and corrections. We thank David Duvenaud for extended discussions on this work. We thank the author of the torchsde library, Xuechen Li, for optimizing the SDE benchmark code.
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Universal Differential Equations for Scientific Machine Learning: Supplemental Information

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August 10, 2020

1 DiffEqFlux.jl Pullback Construction

The DiffEqFlux.jl pullback construction is not based on just one method but instead has a dispatch-based mechanism for choosing between different adjoint implementations. At a high level, the library defines the pullback on the differential equation solve function, and thus using a differential equation inside of a larger program leads to this chunk as being a single differentiable primitive that is inserted into the back pass of Flux.jl when encountered by overloading the Zygote.jl \cite{1} and ChainRules.jl rule sets. For any ChainRules.jl-compliant reverse-mode AD package in the Julia language, when a differential equation solve is encountered in any Julia library during the backwards pass, the adjoint method is automatically swapped in to be used for the backpropogation of the solver. The choice of the adjoint is chosen by the type of the sensealg keyword argument which are fully described below.

1.1 Backpropagation-Accelerated DAE Adjoints for Index-1 DAEs with Constraint Equations

Before describing the modes, we first describe the adjoint of the differential equation with constraints. The constrained ordinary differential equation:

\[ u' = \tilde{f}(u, p, t), \]  
\[ 0 = c(u, p, t), \]

(1)  
(2)
can be rewritten in mass matrix form:

\[ Mu' = f(u, p, t). \]  

(3)

We wish to solve for some cost function \( G(u, p) \) evaluated throughout the differential equation, i.e.:

\[ G(u, p) = \int_{t_0}^{T} g(u, p, t) dt, \]  

(4)

To derive this adjoint, introduce the Lagrange multiplier \( \lambda \) to form:

\[ I(p) = G(p) - \int_{t_0}^{T} \lambda^*(M u' - f(u, p, t)) dt, \]  

(5)

Since \( u' = f(u, p, t) \), we have that:

\[ \frac{dG}{dp} = \frac{dI}{dp} = \int_{t_0}^{T} (g_p + g_u s) dt - \int_{t_0}^{T} \lambda^*(M s' - f_p) dt, \]  

(6)

for \( s_i \) being the sensitivity of the \( i \)th variable. After applying integration by parts to \( \lambda^* M s' \), we require that:

\[ M^* \dot{\lambda} = -\frac{df}{du}^* \lambda - \left( \frac{dg}{du} \right)^* s, \]  

(7)

\[ \lambda(T) = 0, \]  

(8)

to zero out a term and get:

\[ \frac{dG}{dp} = \lambda^*(t_0) M \frac{du}{dp}(t_0) + \int_{t_0}^{T} (g_p + \lambda^* f_p) dt. \]  

(9)

If \( G \) is discrete, then it can be represented via the Dirac delta:

\[ G(u, p) = \int_{t_0}^{T} \sum_{i=1}^{N} \|d_i - u(t_i, p)\|^2 \delta(t_i - t) dt, \]  

(10)

in which case \( g_u(t_i) = 2(d_i - u(t_i, p)) \),

(11)

at the data points \((t_i, d_i)\). Therefore, the derivative of an ODE solution with respect to a cost function is given by solving for \( \lambda^* \) using an ODE for \( \lambda^T \) in reverse time, and then using that to calculate \( \frac{dG}{dp} \). At each time point where discrete data is found, \( \lambda \) is then changed using a callback (discrete event handling) by the amount \( g_u \) to represent the Dirac delta portion of the integral. Lastly, we note that \( \frac{dG}{du_0} = -\lambda(0) \) in this formulation.
We have to take care of consistent initialization in the case of semi-explicit index-1 DAEs. We need to satisfy the system of equations

\[ M^* \Delta \lambda^d = h^*_u \lambda^a + g^*_u \]  
\[ 0 = h^*_u \lambda^a + g^*_u, \]  

where \( d \) and \( a \) denote differential and algebraic variables, and \( f \) and \( g \) denote differential and algebraic equations respectively. Combining the above two equations, we know that we need to increment the differential part of \( \lambda \) by

\[-h^*_u (h^*_u)^{-1} g^*_u + g^*_u \]  

at each callback. Additionally, the ODEs

\[ \mu' = -\lambda^* \frac{\partial f}{\partial p} \]  

with \( \mu(T) = 0 \) can be appended to the system of equations to perform the quadrature for \( \frac{\partial G}{\partial p} \).

1.2 Current Adjoint Calculation Methods

From this setup we have the following 8 possible modes for calculating the adjoint, with their pros and cons.

1. QuadratureAdjoint: a quadrature-based approach. This utilizes interpolation of the forward solution provided by DifferentialEquations.jl to calculate \( u(t) \) at arbitrary time points for doing the calculations with respect to \( \frac{df}{du} \) in the reverse ODE of \( \lambda \). From this a continuous interpolatable \( \lambda(t) \) is generated, and the integral formula for \( \frac{dG}{dp} \) is calculated using the QuadGK.jl implementation of Gauss-Kronrod quadrature. While this approach is memory heavy due to requiring the interpolation of the forward and reverse passes, it can be the fastest version for cases where the number of ODE/DAE states is small and the number of parameters is large since the QuadGK quadrature can converge faster than ODE/DAE-based versions of quadrature. This method requires an ODE or a DAE.

2. InterpolatingAdjoint: a checkpointed interpolation approach. This approach solves the \( \lambda(t) \) ODE in reverse using an interpolation of \( u(t) \), but appends the equations for \( \mu(t) \) and thus does not require saving the time-series trajectory of \( \lambda(t) \). For checkpointing, a scheme similar to that found in SUNDIALS [2] is used. Points \( (u_k, t_k) \) from the forward solution are chosen as the interval points. Whenever the backwards pass enters a new interval, the ODE is re-solved on \( t \in [t_{k-1}, t_k] \) with a continuous interpolation provided by DifferentialEquations.jl. For the reverse pass, the \( \text{tstops} \) argument is set for each \( t_k \), ensuring that no backwards integration step lies in two checkpointing intervals. This requires at most a total of
two forward solutions of the ODE and the memory required to hold the
interpolation of the solution between two consecutive checkpoints. Note
that making the checkpoints at the start and end of the integration in-
terval makes this equivalent to a non-checkpointed interpolated approach
which replaces the quadrature with an ODE/SDE/DAE solve for memory
efficiency. This method tends to be both stable and require a minimal
amount of memory, and is thus the default. This method requires an
ODE, SDE, or a DAE.

3. BacksolveAdjoint: a checkpointed backwards solution approach. Follow-
ing [3], after a forward solution, this approach solves the \( u(t) \) equation in
reverse along with the \( \lambda(t) \) and \( \mu(t) \) ODEs. Thus, since no interpo-
lations are required, it requires \( \mathcal{O}(1) \) memory. Unfortunately, many theoretical
results show that backwards solution of ODEs is not guaranteed to be sta-
ble, and testing this adjoint on the universal partial differential equations
like the diffusion-advection example of this paper showcases that it can be
divergent and is thus not universally applicable, especially in cases of stiff-
ness. Thus for stability we modify this approach by allowing checkpoints
\( (u_k, t_k) \) at which the reverse integration is reset, i.e. \( u(t_k) = u_k \), and the
backsolve is then continued. The \texttt{tstops} argument is set in the integrator
to require that each checkpoint is hit exactly for this resetting to occur.
By doing so, the resulting method utilizes \( \mathcal{O}(1) \) memory + the number
of checkpoints required for stability, making it take the least memory ap-
proach. However, the potential divergence does lead to small errors in the
gradient, and thus for highly stiff equations we have found that this is
only applicable to a certain degree of tolerance like \( 10^{-6} \) given reasonable
numbers of checkpoints. When applicable this can be the most efficient
method for large memory problems. This method requires an ODE, SDE,
or a DAE.

4. ForwardSensitivity: a forward sensitivity approach. From \( u' = f(u, p, t) \),
the chain rule gives \( \frac{du}{dp} = \frac{df}{du} \frac{du}{dp} + \frac{df}{dp} \) which can be appended to the
original equations to give \( \frac{du}{dp} \) as a time series, which can then be used
to compute \( \frac{dG}{dp} \). While the computational cost of the adjoint methods
scales like \( \mathcal{O}(N + P) \) for \( N \) differential equations and \( P \) parameters, this
approach scales like \( \mathcal{O}(NP) \) and is thus only applicable to models with
small numbers of parameters (thus excluding neural networks). However,
when the universal approximator has small numbers of parameters, this
can be the most efficient approach. This method requires an ODE or a
DAE.

5. ForwardDiffSensitivity: a forward-mode automatic differentiation approach,
using ForwardDiff.jl [4] to calculate the forward sensitivity equations,
i.e. an AD-generated implementation of forward-mode “discretize-then-
optimize”. Because it utilizes a forward-mode approach, the scaling matches
that of the forward sensitivity approach and it tends to have similar perfor-
mance characteristics. This method applies to any Julia-based differential equation solver.

6. TrackerAdjoint: a Tracker-driven taped-based reverse-mode discrete adjoint sensitivity, i.e. an AD-generated implementation of reverse-mode “discretize-then-optimize”. This is done by using the TrackedArray constructs of Tracker.jl \cite{5} to build a Wengert list (or tape) of the forward execution of the ODE solver which is then reversed. This method applies to any Julia-based differential equation solver.

7. ZygoteAdjoint: a Zygote-driven source-to-source reverse-mode discrete adjoint sensitivity, i.e. an AD-generated implementation of reverse-mode “discretize-then-optimize”. This utilizes the Zygote.jl \cite{1} system directly on the differential equation solvers to generate a source code for the reverse pass of the solver itself. Currently this is only directly applicable to a few differential equation solvers, but is under heavy development.

8. ReverseDiffAdjoint: A ReverseDiff.jl taped-based reverse-mode discrete adjoint sensitivity, i.e. an AD-generated implementation of reverse-mode “discretize-then-optimize”. In contrast to TrackerAdjoint, this methodology can be substantially faster due to its ability to precompile the tape but only supports calculations on the CPU.

For each of the non-AD approaches, there are the following choices for how the Jacobian-vector products $Jv$ (jvp) of the forward sensitivity equations and the vector-Jacobian products $v'J$ (vjp) of the adjoint sensitivity equations are computed:

1. Automatic differentiation for the jvp and vjp. In this approach, automatic differentiation is utilized for directly calculating the jvp's and vjps. ForwardDiff.jl with a single dual dimension is applied at $f(u + \lambda \epsilon)$ to calculate $\frac{df}{du} \lambda$ where $\epsilon$ is a dual dimensional signifier. For the vector-Jacobian products, a forward pass at $f(u)$ is utilized and the backwards pass is seeded at $\lambda$ to compute the $\lambda' \frac{df}{du}$ (and similarly for $\frac{df}{dp}$). Note that if $f$ is a neural network, this implies that this product is computed by starting the backpropogation of the neural network with $\lambda$ and the vjp is the resulting return. Three methods are allowed to be chosen for performing the internal vjp calculations:

(a) Zygote.jl source-to-source transformation based vjps. Note that only non-mutating differential equation function definitions are supported in this mode. This mode is the most efficient in the presence of neural networks.

(b) ReverseDiff.jl tape-based vjps. This allows for JIT-compilation of the tape for accelerated computation. This is the fastest vjp choice in the presence of heavy scalar operations like in chemical reaction networks, but is not compatible with GPU acceleration.
Tracker.jl with arrays of tracked real values is utilized on mutating functions.

The internal calculation of the vjp on a general UDE recurses down to primitives and embeds optimized backpropogations of the internal neural networks (and other universal approximators) for the calculation of this product when this option is used.

2. Numerical differentiation for the jvp and vjp. In this approach, finite differences is utilized for directly calculating the jvps and vjps. For a small but finite $\epsilon$, $(f(u + \lambda \epsilon) - f(u)) / \epsilon$ is used to approximate $\frac{df}{du} \lambda$. For vjps, a finite difference gradient of $\lambda' f(u)$ is used.

3. Automatic differentiation for Jacobian construction. In this approach, (sparse) forward-mode automatic differentiation is utilized by a combination of ForwardDiff.jl [4] with SparseDiffTools.jl for color-vector based sparse Jacobian construction. After forming the Jacobian, the jvp or vjp is calculated.

4. Numerical differentiation for Jacobian construction. In this approach, (sparse) numerical differentiation is utilized by a combination of DiffEqDiffTools.jl with SparseDiffTools.jl for color-vector based sparse Jacobian construction. After forming the Jacobian, the jvp or vjp is calculated.

In total this gives 48 different adjoint method approaches, each with different performance characteristics and limitations. A full performance analysis which measures the optimal adjoint approach for various UDEs has been omitted from this paper, since the combinatorial nature of the options requires a considerable amount of space to showcase the performance advantages and generality disadvantages between each of the approaches. A follow-up study focusing on accurate performance measurements of the adjoint choice combinations on families of UDEs is planned.

2 Benchmarks

2.1 ODE Solve Benchmarks

The three ODE benchmarks utilized the Lorenz equations (LRNZ) weather prediction model from [6] and the standard ODE IVP Testset [7, 8]:

$$\frac{dx}{dt} = \sigma(y-x)$$  \hspace{1cm} (16)

$$\frac{dy}{dt} = x(\rho - z) - y$$  \hspace{1cm} (17)

$$\frac{dz}{dt} = xy - \beta z$$  \hspace{1cm} (18)
The 28 ODE benchmarks utilized the Pleiades equation (PLEI) celestial mechanics simulation from [6] and the standard ODE IVP Testset [7, 8]:

\[ x''_i = \sum_{j \neq i} m_j (x_j - x_i) / r_{ij} \]  
\[ y''_i = \sum_{j \neq i} m_j (y_j - y_i) / r_{ij} \]

where

\[ r_{ij} = \left( (x_i - x_j)^2 + (y_i - y_j)^2 \right)^{3/2} \]

on \( t \in [0, 3] \) with initial conditions:

\[ u(0) = [3.0, 3.0, -1.0, -3.0, 2.0, -2.0, 2.0, 3.0, -3.0, 2.0, 0.0, -4.0, 4.0, 0.0, 0.0, 0.0, 0.0, 1.75, -1.5, 0.0, 0.0, -1.25, 1.0, 0.0] \]

written in the form \( u = [x_i, y_i, x'_i, y'_i] \).

The rest of the benchmarks were derived from a discretization a two-dimensional reaction diffusion equation, representing systems biology, combustion mechanics, spatial ecology, spatial epidemiology, and more generally physical PDE equations:

\[ A_t = D \Delta A + \alpha_A(x) - \beta_A A - r_1 AB + r_2 C \]  
\[ B_t = \alpha_B - \beta_B B - r_1 AB + r_2 C \]  
\[ C_t = \alpha_C - \beta_C C + r_1 AB - r_2 C \]

where \( \alpha_A(x) = 1 \) if \( x > 80 \) and 0 otherwise on the domain \( x \in [0, 100] \), \( y \in [0, 100] \), and \( t \in [0, 10] \) with zero-flux boundary conditions. The diffusion constant \( D \) was chosen as 100 and all other parameters were left at 1.0.

In this parameter regime the ODE was non-stiff as indicated by solves with implicit methods not yielding performance advantages. The diffusion operator was discretized using the second order finite difference stencil on an \( N \times N \) grid, where \( N \) was chosen to be 16, 32, 64, 128, 256, and 512. To ensure fairness, the torchdiffeq functions were compiled using torchscript. The code for reproducing the benchmark can be found at:

https://gist.github.com/ChrisRackauckas/cc6ac746e2df285c28e0584a2bfa320

2.2 Neural ODE Training Benchmark

The spiral neural ODE from [3] was used as the benchmark for the training of neural ODEs. The data was generated according from the form:

\[ u' = Au^3 \]

where \( A = [-0.1, 2.0; -2.0, -0.1] \) on \( t \in [0, 1.5] \) where data was taken at 30 evenly spaced points. Each of the software packages trained the neural ODE
for 500 iterations using ADAM with a learning rate of 0.05. The defaults using the SciML software resulted in a final loss of 4.895287e-02 in 7.4 seconds, the optimized version (choosing BacksolveAdjoint with compiled ReverseDiff vector-jacobian products) resulted in a final loss of 2.761669e-02 in 2.7 seconds, while torchdiffeq achieved a final loss of 0.0596 in 289 seconds. To ensure fairness, the torchdiffeq functions were compiled using torchscript. Code to reproduce the benchmark can be found at:

https://gist.github.com/ChrisRackauckas/4a4d526c15cc4170ce37da837bfc32c4

2.3 SDE Solve Benchmark

The torchsde benchmarks were created using the geometric Brownian motion example from the torchsde README. The SDE was a 4 independent geometric Brownian motions:

\[ dX_t = \mu X_t dt + \sigma X_t dW_t \] (28)

where \( \mu = 0.5 \) and \( \sigma = 1.0 \). Both software solved the SDE 100 times using the SRI method [9] with fixed time step chosen to give 20 evenly spaced steps for \( t \in [0, 1] \). The SciML ecosystem solvers solved the equation 100 times in 0.00115 seconds, while torchsde v0.1 took 1.86 seconds. We contacted the author who rewrote the Brownian motion portions into C++ and linked it to torchsde as v0.1.1 and this improved the timing to roughly 5 seconds, resulting in a final performance difference of approximately 1,600x. The code to reproduce the benchmarks and the torchsde author’s optimization notes can be found at:

https://gist.github.com/ChrisRackauckas/6a03e7b151c86b32d74af54d495c6

3 Sparse Identification of Missing Model Terms via Universal Differential Equations

The SINDy algorithm [10, 11, 12] enables data-driven discovery of governing equations from data. Notice that to use this method, derivative data \( \dot{X} \) is required. While in most publications on the subject this [10, 11, 12] information is assumed. However, for our studies we assume that only the time series information is available. Here we modify the algorithm to apply to only subsets of the equation in order to perform equation discovery specifically on the trained neural network, and in our modification the \( \dot{X} \) term is replaced with \( U_\theta(t) \), the output of the universal approximator, and thus is directly computable from any trained UDE.

After training the UDE, choose a set of state variables:

\[
X = \begin{bmatrix}
X^T(t_1) \\
X^T(t_2) \\
\vdots \\
X^T(t_m)
\end{bmatrix} = \begin{bmatrix}
x_1(t_1) & x_2(t_1) & \cdots & x_n(t_1) \\
x_1(t_2) & x_2(t_2) & \cdots & x_n(t_2) \\
\vdots & \vdots & \ddots & \vdots \\
x_1(t_m) & x_2(t_m) & \cdots & x_n(t_m)
\end{bmatrix}
\] (29)
and compute the action of the universal approximator on the chosen states:

\[
\dot{X} = \begin{bmatrix}
  x^T(t_1) \\
  x^T(t_2) \\
  \vdots \\
  x^T(t_m)
\end{bmatrix} = \begin{bmatrix}
  \dot{x}_1(t_1) & \dot{x}_2(t_1) & \cdots & \dot{x}_n(t_1) \\
  \dot{x}_1(t_2) & \dot{x}_2(t_2) & \cdots & \dot{x}_n(t_2) \\
  \vdots & \vdots & \ddots & \vdots \\
  \dot{x}_1(t_m) & \dot{x}_2(t_m) & \cdots & \dot{x}_n(t_m)
\end{bmatrix}
\]  

Then evaluate the observations in a basis \( \Theta(X) \). For example:

\[
\Theta(X) = [1 \ X \ X^{P_2} \ X^{P_3} \ \cdots \ \sin(X) \ \cos(X) \ \cdots]
\]

where \( X^{P_i} \) stands for all \( P_i \)th order polynomial terms such as

\[
X^{P_2} = \begin{bmatrix}
  x_1^2(t_1) & x_1(t_1)x_2(t_1) & \cdots & x_2^2(t_1) & \cdots & x_n^2(t_1) \\
  x_1^2(t_2) & x_1(t_2)x_2(t_2) & \cdots & x_2^2(t_2) & \cdots & x_n^2(t_2) \\
  \vdots & \vdots & \ddots & \vdots & \ddots & \vdots \\
  x_1^2(t_m) & x_1(t_m)x_2(t_m) & \cdots & x_2^2(t_m) & \cdots & x_n^2(t_m)
\end{bmatrix}
\]  

Using these matrices, find this sparse basis \( \Xi \) over a given candidate library \( \Theta \) by solving the sparse regression problem \( \hat{X} = \Theta \Xi \) with \( L_1 \) regularization, i.e. minimizing the objective function \( \| \hat{X} - \Theta \Xi \|_2 + \lambda \| \Xi \|_1 \). This method and other variants of SInDy applied to UDEs, along with specialized optimizers for the LASSO \( L_1 \) optimization problem, have been implemented by the authors and collaborators DataDrivenDiffEq.jl library for use with the DifferentialEquations.jl training framework.

### 3.1 Application to the Partial Reconstruction of the Lotka-Volterra Equations

On the Lotka-Volterra equations, we trained a UDE model against a trajectory of 31 points with a constant step size \( \Delta t = 0.1 \) starting from \( x_0 = 0.44249296, y_0 = 4.6280594 \) to recover the function \( U_\theta(x, y) \). The parameters are chosen to be \( \alpha = 1.3, \beta = 0.9, \gamma = 0.8, \delta = 1.8 \). The trajectory has been perturbed with additive noise drawn from a normal distribution scaled by \( 10^{-3} \).

The neural network consists of an input layer, one hidden layers with 32 neurons and a linear output layer. The input and hidden layer have hyperbolic tangent activation functions. We trained for 200 iterations with ADAM with a learning rate \( \gamma = 10^{-2} \). We then switched to BFGS with an initial stepnorm of \( \gamma = 10^{-2} \) setting the maximum iterations to 10000. Typically the training converged after 400-600 iterations in total. The loss was chosen was the L2 loss \( \mathcal{L} = \sum_i (u_\theta(t_i) - d_i)^2 \). The training converged to a final loss between \( 10^{-2} \) and \( 10^{-5} \) for the knowledge-enhanced neural network.

From the trained neural network, data was sampled over the original trajectory and fitted using the SR3 method with varying threshold between \( 10^{-7} \) and \( 10^3 \) logarithmically spaced to ensure a sparse solution. A pareto optimal
Figure 1: Loss trajectories and errors for 74 runs of the partially reconstruction of the Lotka Volterra equations under noisy data.

solution was selected via the L1 norm of the coefficients and the L2 norm of the corresponding error between the differential data and estimate. The knowledge-enhanced neural network returned zero for all terms except non-zeros on the quadratic terms with nearly the correct coefficients that were then fixed using an additional sparse regression over the quadratic terms. The resulting parameters extracted are $\beta \approx 0.9239$ and $\gamma \approx 0.8145$. Performing a sparse identification on the ideal, full solution and numerical derivatives computed via an interpolating spline resulted in a failure.

To ensure reproducible results, the training procedure has been evaluated 74 times given the initial trajectory. The training failed once (1.35%) to exactly identify the original equation in symbolic terms. The final loss has a mean of 0.0004 with a standard error of 0.0012, an upper and lower bound of 0.0103 and $3.3573 \times 10^{-5}$ respectively. The training converged between 241 and 615 iterations. The sum of the L2 error of the recovered model on the training data is centered around 0.1472 with a standard deviation of 0.1688. The results are shown in Figure 1.

3.2 Discovery of Robertson’s Equations with Prior Conservation Laws

On Robertson’s equations, we trained a UDAE model against a trajectory of 10 points on the timespan $t \in [0.0,1.0]$ starting from $y_1 = 1.0$, $y_2 = 0.0$, and
$y_3 = 0.0$. The parameters of the generating equation were $k_1 = 0.04$, $k_2 = 3e7$, and $k_3 = 1e4$. The universal approximator was a neural network with one hidden layers of size 64. The equation was trained using the BFGS optimizer to a loss of $9e-6$.

4 Model-based Learning for the Fisher-KPP Equations

To generate training data for the 1D Fisher-KPP equation, we take the growth rate and the diffusion coefficient to be $r = 1$ and $D = 0.01$ respectively. The equation is numerically solved in the domain $x \in [0,1]$ and $t \in [0,T]$ using a 2nd order central difference scheme for the spatial derivatives and the time-integration is done using the Tsitouras 5/4 Runge-Kutta method. We implement periodic boundary condition $\rho(x = 0,t) = \rho(x = 1,t)$ and initial condition $\rho(x,t = 0) = \rho_0(x)$ is taken to be a localized function given by

$$
\rho_0(x) = \frac{1}{2} \left( \tanh \left( \frac{x - (0.5 - \Delta/2)}{\Delta/10} \right) - \tanh \left( \frac{x - (0.5 + \Delta/2)}{\Delta/10} \right) \right),
$$

with $\Delta = 0.2$ which represents the width of the region where $\rho \simeq 1$. The data are saved at evenly spaced points with $\Delta x = 0.04$ and $\Delta t = 0.5$.

In the UPDE, the growth neural network $\text{NN}_\theta(\rho)$ has 4 densely connected layers with 10, 20, 20 and 10 neurons each and tanh activation functions. The diffusion operator is represented by a CNN that operates on an input vector of arbitrary size. It has 1 hidden layer with a $3 \times 1$ filter $[w_1, w_2, w_3]$ without any bias. To implement periodic boundary conditions for the UPDE at each time step, the vector of values at different spatial locations $[\rho_1, \rho_2, \ldots, \rho_{N_x}]$ is padded with $\rho_{N_x}$ to the left and $\rho_1$ to the right. This also ensures that the output of the CNN is of the same size as the input. The weights of both the neural networks and the diffusion coefficient are simultaneously trained to minimize the loss function

$$
\mathcal{L} = \sum_i (\rho(x_i, t_i) - \rho_{\text{data}}(x_i, t_i))^2 + \lambda |w_1 + w_2 + w_3|,
$$

where $\lambda$ is taken to be $10^2$ (note that one could also structurally enforce $w_3 = -(w_1 + w_2)$). The second term in the loss function enforces that the differential operator that is learned is conservative—that is, the weights sum to zero. The training is done using the ADAM optimizer with learning rate $10^{-3}$. 

11
5 Adaptive Solving for the 100 Dimensional Hamilton-Jacobi-Bellman Equation

5.1 Forward-Backwards SDE Formulation

Consider the class of semilinear parabolic PDEs, in finite time \( t \in [0,T] \) and \( d \)-dimensional space \( x \in \mathbb{R}^d \), that have the form:

\[
\frac{\partial u}{\partial t}(t,x) + \frac{1}{2} \text{tr} \left( \sigma \sigma^T(t,x) (\text{Hess}_x u)(t,x) \right) + \nabla u(t,x) \cdot \mu(t,x) + f(t,x,u(t,x),\sigma^T(t,x) \nabla u(t,x)) = 0,
\]

with a terminal condition \( u(T,x) = g(x) \). In this equation, \( \text{tr} \) is the trace of a matrix, \( \sigma^T \) is the transpose of \( \sigma \), \( \nabla u \) is the gradient of \( u \), and \( \text{Hess}_x u \) is the Hessian of \( u \) with respect to \( x \). Furthermore, \( \mu \) is a vector-valued function, \( \sigma \) is a \( d \times d \) matrix-valued function and \( f \) is a nonlinear function. We assume that \( \mu, \sigma, \) and \( f \) are known. We wish to find the solution at initial time, \( t = 0 \), at some starting point, \( x = \zeta \).

Let \( W_t \) be a Brownian motion and take \( X_t \) to be the solution to the stochastic differential equation

\[
dX_t = \mu(t,X_t)dt + \sigma(t,X_t)dW_t, \tag{36}
\]

with a terminal condition \( u(T,x) = g(x) \). With initial condition \( X(0) = \zeta \), has shown that the solution to ?? satisfies the following forward-backward SDE (FBSDE) \[13\]:

\[
u(t,X_t) - u(0,\zeta) =
- \int_0^t f(s,X_t,u(s,X_t),\sigma^T(s,X_t) \nabla u(s,X_t)) ds
+ \int_0^t [\nabla u(s,X_t)]^T \sigma(s,X_t) dW_s, \tag{37}
\]

with terminating condition \( g(X_T) = u(X_T,W_T) \). Notice that we can combine 36 and 37 into a system of \( d + 1 \) SDEs:

\[
dX_t = \mu(t,X_t)dt + \sigma(t,X_t)dW_t,

dU_t = f(t,X_t,U_t,\sigma^T(t,X_t) \nabla u(t,X_t)) dt + [\sigma^T(t,X_t) \nabla u(t,X_t)]^T dW_t, \tag{38}
\]

where \( U_t = u(t,X_t) \). Since \( X_0, \mu, \sigma, \) and \( f \) are known from the choice of model, the remaining unknown portions are the functional \( \sigma^T(t,X_t) \nabla u(t,X_t) \) and initial condition \( U(0) = u(0,\zeta) \), the latter being the point estimate solution to the PDE.
To solve this problem, we approximate both unknown quantities by universal approximators:

\[
\sigma^T(t, X_t) \nabla u(t, X_t) \approx U_1^3(t, X_t),
\]
\[
u(0, X_0) \approx U_2^2(X_0),
\]

Therefore we can rewrite 38 as a stochastic UDE of the form:

\[
\begin{align*}
dX_t &= \mu(t, X_t) dt + \sigma(t, X_t) dW_t, \\
dU_t &= f(t, X_t, U_t, U_1^3(t, X_t)) dt + \left[ U_1^3(t, X_t) \right]^T dW_t,
\end{align*}
\]

with initial condition \((X_0, U_0) = (X_0, U_2^2(X_0))\).

To be a solution of the PDE, the approximation must satisfy the terminating condition, and thus we define our loss to be the expected difference between the approximating solution and the required terminating condition:

\[
l(\theta_1, \theta_2 | X_T, U_T) = \mathbb{E} \left[ \| g(X_T) - U_T \| \right].
\]

Finding the parameters \((\theta_1, \theta_2)\) which minimize this loss function thus give rise to a BSDE which solves the PDE, and thus \(U_2^2(X_0)\) is the solution to the PDE once trained.

### 5.2 The LQG Control Problem

This PDE can be rewritten into the canonical form by setting:

\[
\begin{align*}
\mu &= 0, \\
\sigma &= \tilde{\sigma} I, \\
f(t, X_t) &= -\alpha \left\| \sigma^T(s, X_s) \nabla u(s, X_s) \right\|^2,
\end{align*}
\]

where \(\tilde{\sigma} = \sqrt{2}\), \(T = 1\) and \(X_0 = (0, ..., 0) \in \mathbb{R}^{100}\). The universal stochastic differential equation was then supplemented with a neural network as the approximator. The initial condition neural network was had 1 hidden layer of size 110, and the \(\sigma^T(t, X_t) \nabla u(t, X_t)\) neural network had two layers both of size 110.

For the example we chose \(\lambda = 1\) This was trained with the LambdaEM method of DifferentialEquations.jl [14] with relative and absolute tolerances set at \(1e^{-4}\) using 500 training iterations and using a loss of 100 trajectories per epoch.

On this problem, for an arbitrary \(g\), one can show with Itô’s formula that:

\[
u(t, x) = -\frac{1}{\lambda} \ln \left( \mathbb{E} \left[ \exp \left( -\lambda g(x + \sqrt{2} W_{T-t}) \right) \right] \right),
\]

which was used to calculate the error from the true solution.
Figure 2: Adaptive solution of the 100-dimensional Hamilton-Jacobi-Bellman equation. This demonstrates that as the universal approximators $U_{\theta_1}$ and $U_{\theta_2}$ converge to satisfy the terminating condition, $U_{\theta_2}$ network convergences to the solution of Equation ??.
6 Reduction of the Boussinesq Equations

As a test for the diffusion-advection equation parameterization approach, data was generated from the diffusion-advection equations using the missing function $\overline{wT} = \cos(\sin(T^4)) + \sin(\cos(T^2))$ with $N$ spatial points discretized by a finite difference method with $t \in [0, 1.5]$ with Neumann zero-flux boundary conditions. A neural network with two hidden layers of size 8 and tanh activation functions was trained against 30 data points sampled from the true PDE. The UPDE was fit by using the ADAM optimizer with learning rate $10^{-2}$ for 200 iterations and then ADAM with a learning rate of $10^{-3}$ for 1000 iterations. The resulting fit is shown in 3 which resulted in a final loss of approximately 0.007. We note that the stabilized adjoints were required for this equation, i.e. the backsolve adjoint method was unstable and results in divergence and thus cannot be used on this type of equation. The trained neural network had a forward pass that took around 0.9 seconds.

For the benchmark against the full Boussinesq equations, we utilized Oceananigans.jl [15]. It was set to utilize adaptive time stepping to maximize the time step according to the CFL condition number (capped at CFL $\leq 0.3$) and matched the boundary conditions, along with setting periodic boundary conditions in the horizontal dimension. The Boussinesq simulation used $128 \times 128 \times 128$ spatial points, a larger number than the parameterization, in order to accurately resolve the mean statistics of the 3-dimensional dynamics as is commonly required in practice [16]. The resulting simulation took 13,737 seconds on the same computer used for the neural diffusion-advection approach, demonstrating the approximate 15,000x acceleration.

Figure 3: Reduction of the Boussinesq equations. On the left is the comparison between the training data (blue) and the trained UPDE (orange) over space at the 10th fitting time point, and on the right is the same comparison shown over time at spatial midpoint.
7 Automated Derivation of Closure Relations for Viscoelastic Fluids

The full FENE-P model is:

\[ \sigma + g \left( \frac{\lambda}{f(\sigma)}\sigma \right) = \frac{\eta}{f(\sigma)} \dot{\gamma}, \]

\[ f(\sigma) = \frac{L^2 + \frac{\lambda(L^2 - 3)}{L^2} \text{Tr}(\sigma)}{L^2 - 3}, \]

where

\[ g(A) = \frac{DA}{Dt} - (\nabla u^T)A - A(\nabla u^T), \]

is the upper convected derivative, and \( L, \eta, \lambda \) are parameters [17]. For a one dimensional strain rate, \( \dot{\gamma} = \dot{\gamma}_{12} = \dot{\gamma}_{21} \neq 0, \dot{\gamma}_{ij} = 0 \) else, the one dimensional stress required is \( \sigma = \sigma_{12} \). However, \( \sigma_{11} \) and \( \sigma_{22} \) are both non-zero and store memory of the deformation (normal stresses). The Oldroyd-B model is the approximation:

\[ G(t) = 2\eta \delta(t) + G_0 e^{-t/\tau}, \]

with the exact closure relation:

\[ \sigma(t) = \eta \dot{\gamma}(t) + \phi, \]

\[ \frac{d\phi}{dt} = G_0 \dot{\gamma} - \phi/\tau. \]

As an arbitrary nonlinear extension, train a UDE model using a single additional memory field against simulated FENE-P data with parameters \( \lambda = 2, L = 2, \eta = 4 \). The UDE model is of the form,

\[ \sigma = U_0(\phi, \dot{\gamma}) \]

\[ \frac{d\phi}{dt} = U_1(\phi, \dot{\gamma}) \]

where \( U_0, U_1 \) are neural networks each with a single hidden layer containing 4 neurons. The hidden layer has a tanh activation function. The loss was taken as \( L = \sum (\sigma(t_i) - \sigma_{\text{FENE-P}}(t_i))^2 \) for 100 evenly spaced time points in \( t_i \in [0, 2\pi] \), and the system was trained using an ADAM iterator with learning rate 0.015. The fluid is assumed to be at rest before \( t = 0 \), making the initial stress also zero.

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