Abstract

The classical development of neural networks has primarily focused on learning mappings between finite dimensional Euclidean spaces or finite sets. We propose a generalization of neural networks tailored to learn operators mapping between infinite dimensional function spaces. We formulate the approximation of operators by composition of a class of linear integral operators and nonlinear activation functions, so that the composed operator can approximate complex nonlinear operators. Furthermore, we introduce four classes of operator parameterizations: graph-based operators, low-rank operators, multipole graph-based operators, and Fourier operators and describe efficient algorithms for computing with each one. The proposed neural operators are resolution-invariant: they share the same network parameters between different discretizations of the underlying function spaces and can be used for zero-shot super-resolutions. Numerically, the proposed models show superior performance compared to existing machine learning based methodologies on Burgers’ equation, Darcy flow, and the Navier-Stokes equation, while being several order of magnitude faster compared to conventional PDE solvers.

Keywords: Deep Learning, Operator Inference, Partial Differential Equations, Navier-Stokes Equation.

1. Introduction

Learning mappings between infinite-dimensional function spaces is a challenging problem with numerous applications across various disciplines. Examples arise in numerous differential equation models in science and engineering, in robotics and in computer vision. In particular, any map where either the input or the output space, or both, can be infinite-dimensional; and in particular where the inputs and/or outputs are themselves functions. The possibility of learning such mappings opens up a new class of problems in the design of neural networks with widespread applicability. New ideas are required to build upon traditional neural networks which are mappings between finite-dimensional Euclidean spaces and/or sets of finite cardinality.

A naive approach to this problem is simply to discretize the (input or output) function spaces and apply standard ideas from neural networks. Instead we formulate a new class of deep neural network
based models, called neural operators, which map between spaces of functions on bounded domains \( D \subset \mathbb{R}^d, D' \subset \mathbb{R}^{d'} \). Such models, once trained, have the property of being discretization invariant: sharing the same network parameters between different discretizations of the underlying functional data. In contrast, the naive approach leads to neural network architectures which depend heavily on this discretization: new architectures with new parameters are needed to achieve the same error for differently discretized data. We demonstrate, numerically, that the same neural operator can achieve a constant error for any discretization of the data while standard feed-forward and convolutional neural networks cannot.

In this paper we experiment with the proposed model within the context of partial differential equations (PDEs). We study various solution operators or flow maps arising from the PDE model; in particular, we investigate mappings between functions spaces where the input data can be, for example, the initial condition, boundary condition, or coefficient function, and the output data is the respective solution. We perform numerical experiments with operators arising from the one-dimensional Burgers’ Equation (Evans, 2010), the two-dimensional steady state of Darcy Flow (Bear and Corapcioglu, 2012) and the two-dimensional Navier-Stokes Equation (Lemarié-Rieusset, 2018).

1.1 Background and Context

**PDEs.** “Differential equations [...] represent the most powerful tool humanity has ever created for making sense of the material world.” Strogatz (2009). Over the past few decades, significant progress has been made on formulating (Gurtin, 1982) and solving (Johnson, 2012) the governing PDEs in many scientific fields from micro-scale problems (e.g., quantum and molecular dynamics) to macro-scale applications (e.g., civil and marine engineering). Despite the success in the application of PDEs to solve real-world problems, two significant challenges remain:

- identifying the governing model for complex systems;
- efficiently solving large-scale non-linear systems of equations.

Identifying/formulating the underlying PDEs appropriate for modeling a specific problem usually requires extensive prior knowledge in the corresponding field which is then combined with universal conservation laws to design a predictive model. For example, modeling the deformation and fracture of solid structures requires detailed knowledge of the relationship between stress and strain in the constituent material. For complicated systems such as living cells, acquiring such knowledge is often elusive and formulating the governing PDE for these systems remains prohibitive, or the models proposed are too simplistic to be informative. The possibility of acquiring such knowledge from data can revolutionize these fields. Second, solving complicated non-linear PDE systems (such as those arising in turbulence and plasticity) is computationally demanding and can often make realistic simulations intractable. Again the possibility of using instances of data from such computations to design fast approximate solvers holds great potential for accelerating scientific and discovery and engineering practice.

**Learning PDE solution operators.** Neural networks have the potential to address these challenges when designed in a way that allows for the emulation of mappings between function spaces (Lu et al., 2019; Bhattacharya et al., 2020; Nelsen and Stuart, 2020; Li et al., 2020a b c; Patel et al., 2021, Opschoor et al. 2020; Schwab and Zech 2019; O’Leary-Roseberry et al. 2020). In PDE applications, the governing equations are by definition local, whilst the solution operator exhibits
non-local properties. Such non-local effects can be described by integral operators explicitly in the spatial domain, or by means of spectral domain multiplication; convolution is an archetypal example. For integral equations, the graph approximations of Nyström type (Belongie et al., 2002) provide a consistent way of connecting different grid or data structures arising in computational methods and understanding their continuum limits (Von Luxburg et al., 2008, Trillos and Slepčev, 2018, Trillos et al., 2020). For spectral domain calculations, there are well-developed tools that exist for approximating the continuum (Boyd, 2001, Trefethen, 2000). For these reasons, neural networks that build in non-locality via integral operators or spectral domain calculations are natural. This is governing framework for our work aimed at designing mesh invariant neural network approximations for the solution operators of PDEs.

1.2 Our Contribution

Neural Operators. We introduce the concept of neural operators by generalizing standard feed-forward neural networks to learn mappings between infinite-dimensional spaces of functions defined on bounded domains of $\mathbb{R}^d$. The non-local component of the architecture is instantiated through either a parameterized integral operator or through multiplication in the spectral domain. The methodology leads to the following contributions.

- We propose neural operators a concept which generalizes neural networks that map between finite-dimensional Euclidean spaces to neural networks that map between infinite-dimensional function spaces.

- By construction, our architectures share the same parameters irrespective of the discretization used on the input and output spaces done for the purposes of computation. Consequently, neural operators are capable of zero-shot super-resolution as demonstrated in Figure 1.
We propose four methods for practically implementing the neural operator framework: graph-based operators, low-rank operators, multipole graph-based operators, and Fourier operators. Specifically, we develop a Nyström extension to connect the integral operator formulation of the neural operator to families of graph neural networks (GNNs) on arbitrary grids. Furthermore, we study the spectral domain formulation of the neural operator which leads to efficient algorithms in settings where fast transform methods are applicable. We include an exhaustive numerical study of the four formulations.

Numerically, we show that the proposed methodology consistently outperforms all existing deep learning methods even on the resolutions for which the standard neural networks were designed. For the two-dimensional Navier-Stokes equation, when learning the entire flow map, the method achieves < 1% error with viscosity $10^{-3}$ and 8% error with Reynolds number 200.

The Fourier neural operator has an inference time that is three orders of magnitude faster than the pseudo-spectral method used to generate the data for the Navier-Stokes problem (Chandler and Kerswell, 2013) – 0.005s compared to the 2.2s on a 256 × 256 uniform spatial grid. Despite its tremendous speed advantage, the method does not suffer from accuracy degradation when used in downstream applications such as solving Bayesian inverse problems.

In this work, we propose the neural operator models to learn mesh-free, infinite-dimensional operators with neural networks. Compared to previous methods that we will discuss in the related work section 1.3 the neural operator remedies the mesh-dependent nature of standard finite-dimensional approximation methods such as convolutional neural networks by producing a single set of network parameters that may be used with different discretizations. It also has the ability to transfer solutions between meshes. Furthermore, the neural operator needs to be trained only once, and obtaining a solution for a new instance of the parameter requires only a forward pass of the network, alleviating the major computational issues incurred in physics-informed neural network methods Raissi et al. (2019). Lastly, the neural operator requires no knowledge of the underlying PDE, only data.

### 1.3 Related Works

We outline the major neural network-based approaches for the solution of PDEs. To make the discussion concrete, we will consider the family of PDEs in the form

$$
\begin{align*}
(L_a u)(x) &= f(x), & x \in D, \\
u(x) &= 0, & x \in \partial D,
\end{align*}
$$

for some $a \in \mathcal{A}$, $f \in \mathcal{U}^*$ and $D \subset \mathbb{R}^d$ a bounded domain. We assume that the solution $u : D \to \mathbb{R}$ lives in the Banach space $\mathcal{U}$ and $L : \mathcal{A} \to \mathcal{L}(\mathcal{U}; \mathcal{U}^*)$ is a mapping from the parameter Banach space $\mathcal{A}$ to the space of (possibly unbounded) linear operators mapping $\mathcal{U}$ to its dual $\mathcal{U}^*$. A natural operator which arises from this PDE is $\mathcal{G}^\dagger : \mathcal{A} \to \mathcal{U}$ defined to map the parameter to the solution $a \mapsto u$. A simple example that we study further in Section 5.2 is when $L_a$ is the weak form of the second-order elliptic operator $-\nabla \cdot (a \nabla)$ subject to homogeneous Dirichlet boundary conditions. In this setting, $\mathcal{A} = L^\infty(D; \mathbb{R}_+)$, $\mathcal{U} = H^1_0(D; \mathbb{R})$, and $\mathcal{U}^* = H^{-1}(D; \mathbb{R})$. When needed, we will
assume that the domain $D$ is discretized into $K \in \mathbb{N}$ points and that we observe $N \in \mathbb{N}$ pairs of coefficient functions and (approximate) solution functions $\{(a_j, u_j)\}_{j=1}^N$ that are used to train the model (see Section 2.1).

**Finite-dimensional operators.** An immediate approach to approximate $G^\dagger$ is to parameterize it as a deep convolutional neural network (CNN) between the finite-dimensional Euclidean spaces on which the data is discretized i.e. $G : \mathbb{R}^K \times \Theta \to \mathbb{R}^K$ (Guo et al., 2016; Zhu and Zabaras, 2018; Adler and Oktem, 2017; Bhatnagar et al., 2019). Khoo et al. (2017) concerns a similar setting, but with output space $\mathbb{R}$. Such approaches are, by definition, not mesh independent and need modifications to the architecture for different resolution and discretization of $D$ in order to achieve consistent error (if at all possible). We demonstrate this issue numerically in Section 6. Furthermore, these approaches are limited to the discretization size and geometry of the training data and hence it is not possible to query solutions at new points in the domain. In contrast for our method, we show in Section 6, both invariance of the error to grid resolution, and the ability to transfer the solution between meshes. The work Ummenhofer et al. (2020) proposed a continuous convolution network for fluid problems, where off-grid points are sampled and linearly interpolated. However the continuous convolution method is still constrained by the underlying grid which prevents generalization to higher resolutions. Similarly, to get finer resolution solution, Jiang et al. (2020) proposed learning super-resolution with a U-Net structure for fluid mechanics problems. However fine-resolution data is needed for training, while neural operators are capable of zero-shot super-resolution with no new data.

**DeepONet** Recently, a novel operator regression architecture, named DeepONet, was proposed by Lu et al. (2019, 2021) that designs a generic neural network based on the approximation theorem from Chen and Chen (1995). The architecture consists of two neural networks: a branch net applied on the input functions and a trunk net applied on the querying locations. Lanthaler et al. (2021) developed an error estimate on the DeepONet. The standard DeepONet structure is a linear approximation of the target operator, where the trunk net and branch net learn the coefficients and basis. On the other hand, the neural operator is a non-linear approximation, which makes it constructively more expressive. We include an detailed discussion of DeepONet in Section 3.2 and as well as a numerical comparison to DeepONet in Section 6.2.

**Physics Informed Neural Networks (PINNs).** A different approach is to directly parameterize the solution $u$ as a neural network $u : \overline{D} \times \Theta \to \mathbb{R}$ (E and Yu, 2018; Raissi et al., 2019; Bar and Sochen. 2019. Smith et al., 2020; Pan and Duraisamy, 2020). This approach is designed to model one specific instance of the PDE, not the solution operator. It is mesh-independent, but for any given new parameter coefficient function $a \in \mathcal{A}$, one would need to train a new neural network $u_a$ which is computationally costly and time consuming. Such an approach closely resembles classical methods such as finite elements, replacing the linear span of a finite set of local basis functions with the space of neural networks.

**ML-based Hybrid Solvers** Similarly, another line of work proposes to enhance existing numerical solvers with neural networks by building hybrid models (Pathak et al. 2020; Um et al., 2020a; Greenfeld et al., 2019). These approaches suffer from the same computational issue as classical methods: one needs to solve an optimization problem for every new parameter similarly to the PINNs setting. Furthermore, the approaches are limited to a setting in which the underlying PDE is known. Purely data-driven learning of a map between spaces of functions is not possible.
**Reduced basis methods.** Our methodology most closely resembles the classical reduced basis method (RBM) (DeVore, 2014) or the method of Cohen and DeVore (2015). The method introduced here, along with the contemporaneous work introduced in the papers (Bhattacharya et al., 2020; Nelsen and Stuart, 2020; Opschoor et al., 2020; Schwab and Zech, 2019; O’Leary-Roseberry et al., 2020; Lu et al., 2019), is, to the best of our knowledge, amongst the first practical supervised learning methods designed to learn maps between infinite-dimensional spaces. It remedies the mesh-dependent nature of the approach in the papers (Guo et al., 2016; Zhu and Zabaras, 2018; Adler and Oktem, 2017; Bhatnagar et al., 2019) by producing a single set of network parameters that can be used with different discretizations. Furthermore, it has the ability to transfer solutions between meshes. Moreover, it need only be trained once on the equation set \( \{a_j, u_j\}_{j=1}^N \). Then, obtaining a solution for a new \( a \sim \mu \) only requires a forward pass of the network, alleviating the major computational issues incurred in (E and Yu, 2018; Raissi et al., 2019; Herrmann et al., 2020; Bar and Sochen, 2019) where a different network would need to be trained for each input parameter. Lastly, our method requires no knowledge of the underlying PDE: it is purely data-driven and therefore non-intrusive. Indeed the true map can be treated as a black-box, perhaps to be learned from experimental data or from the output of a costly computer simulation, not necessarily from a PDE.

**Continuous neural networks.** Using continuity as a tool to design and interpret neural networks is gaining currency in the machine learning community, and the formulation of ResNet as a continuous time process over the depth parameter is a powerful example of this (Haber and Ruthotto, 2017; E, 2017). The concept of defining neural networks in infinite-dimensional spaces is a central problem that long been studied (Williams, 1996; Neal, 1996; Roux and Bengio, 2007; Globerson and Livni, 2016; Guss, 2016). The general idea is to take the infinite-width limit which yields a non-parametric method and has connections to Gaussian Process Regression (Neal, 1996; Matthews et al., 2018; Garriga-Alonso et al., 2018), leading to the introduction of deep Gaussian processes (Damianou and Lawrence, 2013; Dunlop et al., 2018). Thus far, such methods have not yielded efficient numerical algorithms that can parallel the success of convolutional or recurrent neural networks for the problem of approximating mappings between finite dimensional spaces. Despite the superficial similarity with our proposed work, this body of work differs substantially from what we are proposing: in our work we are motivated by the continuous dependence of the data, or the functions it samples, in a spatial variable; in the work outlined in this paragraph continuity is used to approximate the network architecture when the depth or width approaches infinity.

**Nyström approximation, GNNs, and graph neural operators (GNOs).** The graph neural operators (Section 4.1) has an underlying Nyström approximation formulation (Nyström, 1930) which links different grids to a single set of network parameters. This perspective relates our continuum approach to Graph Neural Networks (GNNs). GNNs are a recently developed class of neural networks that apply to graph-structured data; they have been used in a variety of applications. Graph networks incorporate an array of techniques from neural network design such as graph convolution, edge convolution, attention, and graph pooling (Kipf and Welling, 2016; Hamilton et al., 2017; Gilmer et al., 2017; Veličković et al., 2017; Murphy et al., 2018). GNNs have also been applied to the modeling of physical phenomena such as molecules (Chen et al., 2019) and rigid body systems (Battaglia et al., 2018) since these problems exhibit a natural graph interpretation: the particles are the nodes and the interactions are the edges. The work (Alet et al., 2019) performs an initial study that employs graph networks on the problem of learning solutions to Poisson’s equation, among
other physical applications. They propose an encoder-decoder setting, constructing graphs in the latent space, and utilizing message passing between the encoder and decoder. However, their model uses a nearest neighbor structure that is unable to capture non-local dependencies as the mesh size is increased. In contrast, we directly construct a graph in which the nodes are located on the spatial domain of the output function. Through message passing, we are then able to directly learn the kernel of the network which approximates the PDE solution. When querying a new location, we simply add a new node to our spatial graph and connect it to the existing nodes, avoiding interpolation error by leveraging the power of the Nyström extension for integral operators.

Low-rank kernel decomposition and low-rank neural operators (LNOs). Low-rank decomposition is a popular method used in kernel methods and Gaussian process (Kulis et al., 2006; Bach, 2013. Lan et al., 2017. Gardner et al., 2018). We present the low-rank neural operator in Section 4.2 where we structure the kernel network as a product of two factor networks inspired by Fredholm theory. The low-rank method, while simple, is very efficient and easy to train especially when the target operator is close to linear. Khoo and Ying (2019) similarly propose to use neural networks with low-rank structure to approximate the inverse of differential operators. The framework of two factor networks is also similar to the trunk and branch network used in DeepONet (Lu et al., 2019). But in our work, the factor networks are defined on the physical domain and non-local information is accumulated through integration, making our low-rank operator resolution invariant. On the other hand, the number of parameters of the networks in DeepONet(s) depend on the resolution of the data; see Section 3.2 for further discussion.

Multipole, multi-resolution methods, and multipole graph neural operators (MGNOs). To efficiently capture long-range interaction, multi-scale methods such as the classical fast multipole methods (FMM) have been developed (Greengard and Rokhlin, 1997). Based on the assumption that long-range interactions decay quickly, FMM decomposes the kernel matrix into different ranges and hierarchically imposes low-rank structures to the long-range components (hierarchical matrices) (Börm et al., 2003). This decomposition can be viewed as a specific form of the multi-resolution matrix factorization of the kernel (Kondor et al., 2014; Börm et al., 2003). For example, the works of Fan et al. (2019c, b); He and Xu (2019) propose a similar multipole expansion for solving parametric PDEs on structured grids. However, the classical FMM requires nested grids as well as the explicit form of the PDEs. In Section 4.3, we propose the multipole graph neural operator (MGNO) by generalizing this idea to arbitrary graphs in the data-driven setting, so that the corresponding graph neural networks can learn discretization-invariant solution operators which are fast and can work on complex geometries.

Fourier transform, spectral methods, and Fourier neural operators (FNOs). The Fourier transform is frequently used in spectral methods for solving differential equations since differentiation is equivalent to multiplication in the Fourier domain. Fourier transforms have also played an important role in the development of deep learning. In theory, they appear in the proof of the universal approximation theorem (Hornik et al., 1989) and, empirically, they have been used to speed up convolutional neural networks (Mathieu et al., 2013). Neural network architectures involving the Fourier transform or the use of sinusoidal activation functions have also been proposed and studied (Bengio et al. 2007; Mingo et al. 2004; Sitzmann et al., 2020). Recently, some spectral methods for PDEs have been extended to neural networks (Fan et al., 2019a, c; Kashinath et al. 2020). In Section
4.4. we build on these works by proposing the Fourier neural operator architecture defined directly in Fourier space with quasi-linear time complexity and state-of-the-art approximation capabilities.

1.4 Paper Outline

In Section 2, we define the general operator learning problem, which is not limited to PDEs. In Section 3, we define the general framework in terms of kernel integral operators and relate our proposed approach to existing methods in the literature. In Section 4, we define four different ways of efficiently computing with neural operators: graph-based operators (GNO), low-rank operators (LNO), multipole graph-based operators (MGNO), and Fourier operators (FNO). In Section 5 we define four partial differential equations which serve as a testbed of various problems which we study numerically. In Section 6, we show the numerical results for our four approximation methods on the four test problems, and we discuss and compare the properties of each method. In Section 7 we conclude the work, discuss potential limitations and outline directions for future work.

2. Learning Operators

2.1 Problem Setting

Our goal is to learn a mapping between two infinite dimensional spaces by using a finite collection of observations of input-output pairs from this mapping. We make this problem concrete in the following setting. Let $\mathcal{A}$ and $\mathcal{U}$ be Banach spaces of functions defined on bounded domains $D \subset \mathbb{R}^d$, $D' \subset \mathbb{R}^{d'}$ respectively and $\mathcal{G}^\dagger : \mathcal{A} \to \mathcal{U}$ be a (typically) non-linear map. Suppose we have observations $\{a_j, u_j\}_{j=1}^N$ where $a_j \sim \mu$ are i.i.d. samples drawn from some probability measure $\mu$ supported on $\mathcal{A}$ and $u_j = \mathcal{G}^\dagger(a_j)$ is possibly corrupted with noise. We aim to build an approximation of $\mathcal{G}^\dagger$ by constructing a parametric map

$$\mathcal{G}_\theta : \mathcal{A} \to \mathcal{U}, \quad \theta \in \mathbb{R}^p$$

(2)

with parameters from the finite-dimensional space $\mathbb{R}^p$ and then choosing $\theta^\dagger \in \mathbb{R}^p$ so that $\mathcal{G}_{\theta^\dagger} \approx \mathcal{G}^\dagger$.

We will be interested in controlling the error of the approximation on average with respect to $\mu$. In particular, assuming $\mathcal{G}^\dagger$ is $\mu$-measurable, we will aim to control the $L^2_\mu(\mathcal{A}; \mathcal{U})$ Bochner norm of the approximation

$$\|\mathcal{G}^\dagger - \mathcal{G}_\theta\|^2_{L^2_\mu(\mathcal{A}; \mathcal{U})} = \mathbb{E}_{a \sim \mu} \|\mathcal{G}^\dagger(a) - \mathcal{G}_\theta(a)\|^2_{\mathcal{U}} = \int_\mathcal{A} \|\mathcal{G}^\dagger(a) - \mathcal{G}_\theta(a)\|^2_{\mathcal{U}} d\mu(a).$$

(3)

This is a natural framework for learning in infinite-dimensions as one could seek to solve the associated empirical-risk minimization problem

$$\min_{\theta \in \mathbb{R}^p} \mathbb{E}_{a \sim \mu} \|\mathcal{G}^\dagger(a) - \mathcal{G}_\theta(a)\|^2_{L^2_\mu} \approx \min_{\theta \in \mathbb{R}^p} \frac{1}{N} \sum_{j=1}^N \|u_j - \mathcal{G}_\theta(a_j)\|^2_{L^2_\mu}$$

(4)

which directly parallels the classical finite-dimensional setting (Vapnik, 1998).

2.2 Discretization

Since our data $a_j$ and $u_j$ are, in general, functions, to work with them numerically, we assume access only to their point-wise evaluations. To illustrate this, we will continue with the example
of the preceding paragraph. For simplicity, assume $D = D'$ and suppose that the input and output function are real-valued. Let $D_j = \{x_j^{(1)}, \ldots, x_j^{(n_j)}\} \subset D$ be a $n_j$-point discretization of the domain $D$ and assume we have observations $a_j|_{D_j}, u_j|_{D_j} \in \mathbb{R}^{n_j}$, for a finite collection of input-output pairs indexed by $j$. In the next section, we propose a kernel inspired graph neural network architecture which, while trained on the discretized data, can produce the solution $u(x)$ for any $x \in D$ given an input $a \sim \mu$. That is to say that our approach is independent of the discretization $D_j$. We refer to this as being a function space architecture, a mesh-invariant architecture or a discretization-invariant architecture; this claim is verified numerically by showing invariance of the error as $n_j \to \infty$. Such a property is highly desirable as it allows a transfer of solutions between different grid geometries and discretization sizes with a single architecture which has a fixed number of parameters.

We note that, while the application of our methodology is based on having point-wise evaluations of the function, it is not limited by it. One may, for example, represent a function numerically as a finite set of truncated basis coefficients. Invariance of the representation would then be with respect to the size of this set. Our methodology can, in principle, be modified to accommodate this scenario through a suitably chosen architecture. We do not pursue this direction in the current work.

3. Proposed Architecture

3.1 Neural Operators

In this section, we outline the neural operator framework. We assume that the input functions $a \in \mathcal{A}$ are $\mathbb{R}^{d_a}$-valued and defined on the bounded domain $D \subset \mathbb{R}^d$ while the output functions $u \in \mathcal{U}$ are $\mathbb{R}^{d_u}$-valued and defined on the bounded domain $D' \subset \mathbb{R}^{d'}$. The proposed architecture $G_\theta : \mathcal{A} \to \mathcal{U}$ has the following overall structure:

1. **Lifting**: Using a pointwise function $\mathbb{R}^{d_a} \to \mathbb{R}^{d_{v_0}}$, map the input $\{a : D \to \mathbb{R}^{d_a}\} \mapsto \{v_0 : D \to \mathbb{R}^{d_{v_0}}\}$ to its first hidden representation. Usually, we choose $d_{v_0} > d_a$ and hence this is a lifting operation performed by a fully local operator.

2. **Iterative kernel integration**: For $t = 0, \ldots, T - 1$, map each hidden representation to the next $\{v_t : D_t \to \mathbb{R}^{d_{v_t}}\} \mapsto \{v_{t+1} : D_{t+1} \to \mathbb{R}^{d_{v_{t+1}}}\}$ via the action of the sum of a local linear operator, a non-local integral kernel operator, and a bias function, composing the sum with a fixed, pointwise nonlinearity. Here we set $D_0 = D$ and $D_T = D'$ and impose that $D_t \subset \mathbb{R}^{d_t}$ is a bounded domain.

3. **Projection**: Using a pointwise function $\mathbb{R}^{d_{v_T}} \to \mathbb{R}^{d_u}$, map the last hidden representation $\{v_T : D' \to \mathbb{R}^{d_{v_T}}\} \mapsto \{u : D' \to \mathbb{R}^{d_u}\}$ to the output function. Analogously to the first step, we usually pick $d_{v_T} > d_u$ and hence this is a projection step performed by a fully local operator.

The outlined structure mimics that of a finite dimensional neural network where hidden representations are successively mapped to produce the final output. In particular, we have

$$G_\theta := Q \circ \sigma_T(W_{T-1} + \mathcal{K}_{T-1} + b_{T-1}) \circ \cdots \circ \sigma_1(W_0 + \mathcal{K}_0 + b_0) \circ \mathcal{P}$$

(5)

where $\mathcal{P} : \mathbb{R}^{d_a} \to \mathbb{R}^{d_{v_0}}$, $Q : \mathbb{R}^{d_{v_T}} \to \mathbb{R}^{d_u}$ are the local lifting and projection mappings respectively, $W_t \in \mathbb{R}^{d_{v_t} \times d_{v_{t+1}}}$ are local linear operators (matrices), $\mathcal{K}_t : \{v_t : D_t \to \mathbb{R}^{d_{v_t}}\} \to \{v_{t+1} : D_{t+1} \to \mathbb{R}^{d_{v_{t+1}}}\}$ are non-local integral kernel operators, and $b_t \in \mathbb{R}^{d_{v_t}}$ are bias functions.
\( \mathbb{R}^{d_{t+1}} \) are integral kernel operators, \( b_t : D_{t+1} \to \mathbb{R}^{d_{t+1}} \) are bias functions, and \( \sigma_t \) are fixed activation functions acting locally as maps \( \mathbb{R}^{d_{t+1}} \to \mathbb{R}^{d_{t+1}} \) in each layer. The output dimensions \( d_v, \ldots, d_v \) as well as the input dimensions \( d_1, \ldots, d_{t-1} \) and domains of definition \( D_1, \ldots, D_{t-1} \) are hyperparameters of the architecture. By local maps, we mean that the action is pointwise, in particular, for the lifting and projection maps, we have \((P(a))(x) = P(a(x)) \) for any \( x \in D \) and \((Q(v_T))(x) = Q(v_T(x)) \) for any \( x \in D' \) and similarly, for the activation, \( (\sigma(v_{t+1}))(x) = \sigma(v_{t+1}(x)) \) for any \( x \in D_{t+1} \). The maps, \( P, Q, \) and \( \sigma_t \) can thus be thought of as defining Nemitskiy operators (Dudley and Norvaisa, 2011, Chapters 6, 7) when each of their components are assumed to be Borel measurable. This interpretation allows us to define the general neural network architecture when pointwise evaluation is not well-defined in the spaces \( \mathcal{A} \) or \( \mathcal{U} \) e.g. when they are Lebesgue, Sobolev, or Besov spaces.

The crucial difference between the proposed architecture (5) and a standard feed-forward neural network is that all operations are directly defined in function space and therefore do not depend on any discretization of the data. Intuitively, the lifting step locally maps the data to a space where the non-local part of \( G^1 \) is easier to capture. This is then learned by successively approximating using integral kernel operators composed with a local nonlinearity. Each integral kernel operator is the function space analog of the weight matrix in a standard feed-forward network since they are infinite-dimensional linear operators mapping one function space to another. We turn the biases, which are normally vectors, to functions and, using intuition from the ResNet architecture [CITE], we further add a local linear operator acting on the output of the previous layer before applying the nonlinearity. The final projection step simply gets us back to the space of our output function. We concatenate in \( \theta \in \mathbb{R}^p \) the parameters of \( P, Q, \{b_t\} \) which are usually themselves shallow neural networks, the parameters of the kernels representing \( \{K_t\} \) which are again usually shallow neural networks, and the matrices \( \{W_t\} \). We note, however, that our framework is general and other parameterizations such as polynomials may also be employed.

**Integral Kernel Operators** We define three version of the integral kernel operator \( K_t \) used in (5). For the first, let \( K^{(t)} \in C(D_{t+1} \times D_t; \mathbb{R}^{d_{t+1} \times d_v}) \) and let \( \nu_t \) be a Borel measure on \( D_t \). Then we define \( K_t \) by

\[
(K_t(v_t))(x) = \int_{D_t} K^{(t)}(x, y, v_t(y)) \, d\nu_t(y) \quad \forall x \in D_{t+1}.
\]

Normally, we take \( \nu_t \) to simply be the Lebesgue measure on \( \mathbb{R}^{d_t} \) but, as discussed in Section 4, other choices can be used to speed up computation or aid the learning process by building in \textit{a priori} information.

For the second, let \( K^{(t)} \in C(D_{t+1} \times D_t \times \mathbb{R}^d \times \mathbb{R}^d; \mathbb{R}^{d_{t+1} \times d_v}) \). Then we define \( K_t \) by

\[
(K_t(v_t))(x) = \int_{D_t} K^{(t)}(x, y, a(P^{D}_{t+1}(x)), a(P^{D}_{t}(y))) \, d\nu_t(y) \quad \forall x \in D_{t+1}.
\]

where \( \Pi^D_t : D_t \to D \) are fixed mappings. We have found numerically that, for certain PDE problems, the form (7) outperforms (6) due to the strong dependence of the solution \( u \) on the parameters \( a \). Indeed, if we think of (5) as a discrete time dynamical system, then the input \( a \in \mathcal{A} \) only enters through the initial condition hence its influence diminishes with more layers. By directly building in \( a \)-dependence into the kernel, we ensure that it influences the entire architecture.
Lastly, let \( \kappa^{(t)} \in C(D_{t+1} \times D_t \times \mathbb{R}^{d_{et}} \times \mathbb{R}^{d_{et}}; \mathbb{R}^{d_{t+1} \times d_{et}}) \). Then we define \( K_t \) by

\[
(K_t(v_t))(x) = \int_{D_t} \kappa^{(t)}(x, y, v_t(\Pi_t(x)), v_t(y)) v_t(y) \, dv_t(y) \quad \forall x \in D_{t+1}. \tag{8}
\]

where \( \Pi_t : D_{t+1} \to D_t \) are fixed mappings. Note that, in contrast to (6) and (7), the integral operator (8) is nonlinear since the kernel can depend on the input function \( v_t \). With this definition and a particular choice of kernel \( \kappa_t \) and measure \( \nu_t \), we show in Section 3.3 that neural operators are a continuous input/output space generalization of the popular transformer architecture (Vaswani et al. 2017).

**Single Hidden Layer Construction** Having defined possible choices for the integral kernel operator, we are now in a position to explicitly write down a full layer of the architecture defined by (5).

For simplicity, we choose the integral kernel operator given by (6), but note that the other definitions (7), (8) work analogously. We then have that a single hidden layer update is given by

\[
v_{t+1}(x) = \sigma_{t+1} \left( W_t v_t(\Pi_t(x)) + \int_{D_t} \kappa^{(t)}(x, y, v_t(\Pi_t(x)), v_t(y)) v_t(y) \, dv_t(y) + b_t(x) \right) \quad \forall x \in D_{t+1} \tag{9}
\]

where \( \Pi_t : D_{t+1} \to D_t \) are fixed mappings. We remark that, since we often consider functions on the same domain, we usually take \( \Pi_t \) to be the identity.

We will now give an example of a full single hidden layer architecture i.e. when \( T = 2 \). We choose \( D_1 = D \), take \( \sigma_2 \) as the identity, and denote \( \sigma_1 \) by \( \sigma \), assuming it is any activation function. Furthermore, for simplicity, we set \( W_1 = 0 \), \( b_1 = 0 \), and assume that \( \nu_0 = \nu_1 \) is the Lebesgue measure on \( \mathbb{R}^d \). We then have that (5) becomes

\[
(G_\theta(a))(x) = Q \left( \int_D \kappa^{(1)}(x, y) \sigma \left( W_0 P(a(y)) + \int_D \kappa^{(0)}(y, z) P(a(z)) \, dz + b_0(y) \right) \, dy \right) \tag{10}
\]

for any \( x \in D' \). In this example, \( P \in C(\mathbb{R}^{d_a}; \mathbb{R}^{d_{v0}}) \), \( \kappa^{(0)} \in C(D \times D; \mathbb{R}^{d_{v1} \times d_{v0}}) \), \( b_0 \in C(D; \mathbb{R}^{d_{v1}}) \), \( W_0 \in \mathbb{R}^{d_{v1} \times d_{v0}} \), \( \kappa^{(0)} \in C(D' \times D; \mathbb{R}^{d_{v2} \times d_{v1}}) \), and \( Q \in C(\mathbb{R}^{d_{v2}}; \mathbb{R}^{d_u}) \). One can then parametrize the continuous functions \( P, Q, \kappa^{(0)}, \kappa^{(1)}, b_0 \) by standard feed-forward neural networks (or by any other means) and the matrix \( W_0 \) simply by its entries. The parameter vector \( \theta \in \mathbb{R}^p \) then becomes the concatenation of the parameters of \( P, Q, \kappa^{(0)}, \kappa^{(1)}, b_0 \) along with the entries of \( W_0 \). One can then optimize these parameters by minimizing with respect to \( \theta \) using standard gradient based minimization techniques. To implement this minimization, the functions entering the loss need to be discretized; but the learned parameters may then be used with other discretizations. In Section 4, we discuss various choices for parametrizing the kernels and picking the integration measure and how these choices affect the computational complexity of the architecture.

**Preprocessing** It is often beneficial to manually include features into the input functions \( a \) to help facilitate the learning process. For example, instead of considering the \( \mathbb{R}^{d_a} \)-valued vector field \( a \) as input, we use the \( \mathbb{R}^{d+a} \)-valued vector field \( (x, a(x)) \). By including the identity element, information about the geometry of the spatial domain \( D \) is directly incorporated into the architecture. This allows the neural networks direct access to information that is already known in the problem and therefore eases learning. We use this idea in all of our numerical experiments in Section 6. Similarly, when dealing with a smoothing operators, it may be beneficial to include a smoothed version of the inputs \( a \) using, for example, Gaussian convolution. Derivative information may also be of interest and thus, as input, one may consider, for example, the \( \mathbb{R}^{d+2d_a+d_{da}} \)-valued vector field \( (x, a(x), a_x(x), \nabla_x a_x(x)) \). Many other possibilities may be considered on a problem-specific basis.
3.2 DeepONets are Neural Operators

We will now draw a parallel between the recently proposed DeepONet architecture in Lu et al. (2019) and our neural operator framework. In fact, we will show that a particular variant of functions from the DeepONets class is a special case of a single hidden layer neural operator construction. To that end, we work with (10) where we choose \( W_0 = 0 \) and denote \( b_0 \) by \( b \). For simplicity, we will consider only real-valued functions i.e. \( d_n = d_u = 1 \) and set \( d_{v_0} = d_{v_1} = d_{v_2} = n \in \mathbb{N} \). Define \( \mathcal{P} : \mathbb{R} \to \mathbb{R}^n \) by \( \mathcal{P}(x) = (x, \ldots, x) \) and \( \mathcal{Q} : \mathbb{R}^n \to \mathbb{R} \) by \( \mathcal{Q}(x) = x_1 + \cdots + x_n \). Furthermore let \( \kappa^{(1)} : D' \times D \to \mathbb{R}^{n \times n} \) be given as \( \kappa^{(1)}(x, y) = \text{diag}(\kappa_1^{(1)}(x, y), \ldots, \kappa_n^{(1)}(x, y)) \) for some \( \kappa_1^{(1)}, \ldots, \kappa_n^{(1)} : D' \times D \to \mathbb{R} \). Similarly let \( \kappa^{(0)} : D \times D \to \mathbb{R}^{n \times n} \) be given as \( \kappa^{(0)}(x, y) = \text{diag}(\kappa_1^{(0)}(x, y), \ldots, \kappa_n^{(0)}(x, y)) \) for some \( \kappa_1^{(0)}, \ldots, \kappa_n^{(0)} : D \times D \to \mathbb{R} \). Then (10) becomes

\[
(G_0(x))(x) = \sum_{j=1}^{n} \int_{D} \kappa_1^{(1)}(x, y)\sigma \left( \int_{D} \kappa_1^{(0)}(y, z)dz + b_j(y) \right)dy
\]

where \( b_j(y) = (b_1(y), \ldots, b_n(y)) \) for some \( b_1, \ldots, b_n : D \to \mathbb{R} \). Choose each \( \kappa_1^{(1)}(x, y) = w_j(y)\varphi_j(x) \) for some \( w_1, \ldots, w_n : D \to \mathbb{R} \) and \( \varphi_1, \ldots, \varphi_n : D' \to \mathbb{R} \) then we obtain

\[
(G_0(a))(x) = \sum_{j=1}^{n} G_j(a)\varphi_j(x)
\]

where \( G_1, \ldots, G_n : A \to \mathbb{R} \) are functionals defined as

\[
G_j(a) = \int_{D} w_j(y)\sigma \left( \int_{D} \kappa_1^{(0)}(y, z)dz + b_j(y) \right)dy.
\]

The set of maps \( \varphi_1, \ldots, \varphi_n \) form the trunk net while the functionals \( G_1, \ldots, G_n \) form the branch net of a DeepONet. The only difference between DeepONet(s) and (11) is the parametrization used for the functionals \( G_1, \ldots, G_n \). Following Chen and Chen (1995), DeepONet(s) define the functional \( G_j \) as maps between finite dimensional spaces. Indeed, they are viewed as \( G_j : \mathbb{R}^q \to \mathbb{R} \) and defined to map pointwise evaluations \( (a(x_1), \ldots, a(x_q)) \) of \( a \) for some set of points \( x_1, \ldots, x_q \in D \). We note that, in practice, this set of evaluation points is not known \( a \) priori and could potentially be very large. Indeed we show that (12) can approximate the functionals defined by DeepONet(s) arbitrary well therefore making DeepONet(s) a special case of neural operators. Furthermore (12) is consistent in function space as the number of parameters used to define each \( G_j \) is independent of any discretization that may be used for \( a \), while this is not true in the DeepONet case as the number of parameters grow as we refine the discretization of \( a \). We demonstrate numerically in Section 6 that the error incurred by DeepONet(s) grows with the discretization of \( a \) while it is remains constant for neural operators.

We point out that parametrizations of the form (11) fall within the class of linear approximation methods since the nonlinear space \( \mathcal{G}^+(A) \) is approximated by the linear space span\{\( \varphi_1, \ldots, \varphi_n \)\} DeVore (1998). The quality of the best possible linear approximation to a nonlinear space is given by the Kolmogorov \( n \)-width where \( n \) is the dimension of the linear space used in the approximation (Pinkus, 1985). The rate of decay of the \( n \)-width as a function of \( n \) quantifies how well the linear space approximates the nonlinear one. It is well know that for some problems such as the flow maps of advection-dominated PDEs, the \( n \)-widths decay very slowly; hence a very large \( n \) is needed for a
good approximation for such problems (Cohen and DeVore, 2015). This can be limiting in practice as more parameters are needed in order to describe more basis functions $\varphi_j$ and therefore more data is needed to fit these parameters.

On the other hand, we point out that parametrizations of the form (5), and the particular case (10), constitute (in general) a form of nonlinear approximation. The benefits of nonlinear approximation are well understood in the setting of function approximation, see e.g. (DeVore 1998), however the theory for the operator setting is still in its infancy (Bonito et al., 2020; Cohen et al., 2020). We observe numerically in Section 6 that nonlinear parametrizations such as (10) outperform linear one such as DeepONets or the low-rank method introduced in Section 4.2

3.3 Transformers are Neural Operators

We will now show that our neural operator framework can be viewed as a continuum generalization to the popular transformer architecture (Vaswani et al., 2017) which has been extremely successful in natural language processing tasks (Devlin et al., 2018; Brown et al., 2020) and, more recently, is becoming a popular choice in computer vision tasks (Dosovitskiy et al., 2020). The parallel stems from the fact that we can view sequences of arbitrary length as arbitrary discretizations of functions. Indeed, in the context of natural language processing, we may think of a sentence as a “word”-valued function on, for example, the domain $[0, 1]$. Assuming our function is linked to a sentence with a fixed semantic meaning, adding or removing words from the sentence simply corresponds to refining or coarsening the discretization of $[0, 1]$. We will now make this intuition precise.

We will show that by making a particular choice of the nonlinear integral kernel operator (8) and discretizing the integral by a Monte-Carlo approximation, a neural operator layer reduces to a pre-normalized, single-headed attention, transformer block as originally proposed in (Vaswani et al., 2017). For simplicity, we assume $d_{v_t} = n \in \mathbb{N}$ and that $D_t = D$ for any $t = 0, \ldots, T$, the bias term is zero, and $W = I$ is the identity. Further, to simplify notation, we will drop the layer index $t$ from (9) and, employing (8), obtain

$$u(x) = \sigma \left( v(x) + \int_D \kappa_v(x, y, v(x), v(y)) v(y) \, dy \right) \quad \forall x \in D$$

(13)

a single layer of the neural operator where $v : D \to \mathbb{R}^n$ is the input function to the layer and we denote by $u : D \to \mathbb{R}^n$ the output function. We use the notation $\kappa_v$ to indicate that the kernel depends on the entirety of the function $v$ and not simply its pointwise values. While this is not explicitly done in (8), it is a straightforward generalization. We now pick a specific form for kernel, in particular, we assume $\kappa_v : \mathbb{R}^n \times \mathbb{R}^n \to \mathbb{R}^{n \times n}$ does not explicitly depend on the spatial variables $(x, y)$ but only on the input pair $(v(x), v(y))$. Furthermore, we let

$$\kappa_v(v(x), v(y)) = g_v(v(x), v(y)) R$$

where $R \in \mathbb{R}^{n \times n}$ is matrix of free parameters i.e. its entries are concatenated in $\theta$ so they are learned, and $g_v : \mathbb{R}^n \times \mathbb{R}^n \to \mathbb{R}$ is defined as

$$g_v(v(x), v(y)) = \left( \int_D \exp \left( \frac{\langle Av(s), Bv(y) \rangle}{\sqrt{m}} \right) \, ds \right)^{-1} \exp \left( \frac{\langle Av(x), Bv(y) \rangle}{\sqrt{m}} \right) .$$
Here \( A, B \in \mathbb{R}^{m \times n} \) are again matrices of free parameters, \( m \in \mathbb{N} \) is a hyperparameter, and \( \langle \cdot, \cdot \rangle \) is the Euclidean inner-product on \( \mathbb{R}^m \). Putting this together, we find that (13) becomes

\[
\begin{align*}
  u(x) &= \sigma \left( v(x) + \int_D \exp \left( \frac{\langle Av(x), Bv(y) \rangle}{\sqrt{m}} \right) Rv(y) \, dy \right) \quad \forall x \in D. 
\end{align*}
\] (14)

Equation (14) can be thought of as the continuum limit of a transformer block. To see this, we will discretize to obtain the usual transformer block.

To that end, let \( \{ x_1, \ldots, x_k \} \subset D \) be a uniformly-sampled, \( k \)-point discretization of \( D \) and denote \( v_j = v(x_j) \in \mathbb{R}^n \) and \( u_j = u(x_j) \in \mathbb{R}^n \) for \( j = 1, \ldots, k \). Approximating the inner-integral in (14) by Monte-Carlo, we have

\[
\int_D \exp \left( \frac{\langle Av(s), Bv(y) \rangle}{\sqrt{m}} \right) \, ds \approx \frac{|D|}{k} \sum_{l=1}^k \exp \left( \frac{\langle Av_l, Bv(y) \rangle}{\sqrt{m}} \right).
\]

Plugging this into (14) and using the same approximation for the outer integral yields

\[
\begin{align*}
  u_j &= \sigma \left( v_j + \sum_{q=1}^k \frac{ \exp \left( \frac{\langle Av_j, Bv_q \rangle}{\sqrt{m}} \right) }{ \sum_{l=1}^k \exp \left( \frac{\langle Av_l, Bv_q \rangle}{\sqrt{m}} \right) } \right) Rv_q, \\
\quad j = 1, \ldots, k. 
\end{align*}
\] (15)

Equation (15) can be viewed as a Nyström approximation of (14). Define the vectors \( z_q \in \mathbb{R}^k \) by

\[
z_q = \frac{1}{\sqrt{m}} \langle Av_1, Bv_q \rangle, \ldots, \langle Av_k, Bv_q \rangle, \quad q = 1, \ldots, k.
\]

Define \( S : \mathbb{R}^k \rightarrow \Delta_k \), where \( \Delta_k \) denotes the \( k \)-dimensional probability simplex, as the softmax function

\[
S(w) = \left( \frac{\exp(w_1)}{\sum_{j=1}^k \exp(w_j)}, \ldots, \frac{\exp(w_k)}{\sum_{j=1}^k \exp(w_j)} \right), \quad \forall w \in \mathbb{R}^k.
\]

Then we may re-write (15) as

\[
\begin{align*}
  u_j &= \sigma \left( v_j + \sum_{q=1}^k S_j(z_q) Rv_q \right), \\
\quad j = 1, \ldots, k.
\end{align*}
\]

Furthermore, if we re-parametrize \( R = R^\text{out} R^\text{val} \) where \( R^\text{out} \in \mathbb{R}^{n \times m} \) and \( R^\text{val} \in \mathbb{R}^{m \times n} \) are matrices of free parameters, we obtain

\[
\begin{align*}
  u_j &= \sigma \left( v_j + R^\text{out} \sum_{q=1}^k S_j(z_q) R^\text{val} v_q \right), \\
\quad j = 1, \ldots, k.
\end{align*}
\]

which is precisely the single-headed attention, transformer block with no layer normalization applied inside the activation function. In the language of transformers, the matrices \( A, B, \) and \( R^\text{val} \) correspond to the queries, keys, and values functions respectively. We note that tricks such as layer
normalization (Ba et al., 2016) can be adapted in a straightforward manner to the continuum setting and incorporated into (14). Furthermore multi-headed self-attention can be realized by simply allowing \( \kappa \) to be a sum of over multiple functions with form \( g_{\kappa} \) all of which have separate trainable parameters. Including such generalizations yields the continuum limit of the transformer as implemented in practice. We do not pursue this here as our goal is simply to draw a parallel between the two methods.

While we have not rigorously experimented with using transformer architectures for the problems outlined in Section 5 we have found, in initial tests, that they perform worse, are slower, and are more memory expensive than neural operators using (6)-(8). Their high computational complexity is evident from (14) as we must evaluate a nested integral of \( v \) for each \( x \in D \). Recently more efficient transformer architectures have been proposed e.g. (Choromanski et al., 2020) and some have been applied to computer vision tasks. We leave as interesting future work experimenting and comparing these architectures to the neural operator both on problems in scientific computing and more traditional machine learning tasks.

4. Parameterization and Computation

In this section, we discuss various ways of parameterizing the infinite dimensional architecture (5). The goal is to find an intrinsic infinite dimensional parameterization that achieves small error (say \( \epsilon \)), and then rely on numerical approximation to ensure that this parameterization delivers an error of the same magnitude (say \( 2\epsilon \)), for all data discretizations fine enough. In this way the number of parameters used to achieve \( O(\epsilon) \) error is independent of the data discretization. In many applications we have in mind the data discretization is something we can control, for example when generating input/output pairs from solution of partial differential equations via numerical simulation. The proposed approach allows us to train a neural operator approximation using data from different discretizations, and to predict with discretizations different from those used in the data, all by relating everything to the underlying infinite dimensional problem.

We also discuss the computational complexity of the proposed parameterizations and suggest algorithms which yield efficient numerical methods for approximation. Subsections 4.1-4.4 delineate each of the proposed methods.

To simplify notation, we will only consider a single layer of (5) i.e. (9) and choose the input and output domains to be the same. Furthermore, we will drop the layer index \( t \) and write the single layer update as

\[
\forall x \in D \quad u(x) = \sigma \left( Wv(x) + \int_D \kappa(x, y)v(y) \, d\nu(y) + b(x) \right)
\]

where \( D \subset \mathbb{R}^d \) is a bounded domain, \( v : D \to \mathbb{R}^n \) is the input function and \( u : D \to \mathbb{R}^m \) is the output function. When the domain domains \( D \) of \( v \) and \( u \) are different, we will usually extend them to be on a larger domain. We will consider \( \sigma \) to be fixed, and, for the time being, take \( d\nu(y) = dy \) to be the Lebesgue measure on \( \mathbb{R}^d \). Equation (16) then leaves three objects which can be parameterized: \( W \), \( \kappa \), and \( b \). Since \( W \) is linear and acts only locally on \( v \), we will always parametrize it by the values of its associated \( m \times n \) matrix; hence \( W \in \mathbb{R}^{m \times n} \) yielding \( mn \) parameters.

The rest of this section will be dedicated to choosing the kernel function \( \kappa : D \times D \to \mathbb{R}^{m \times n} \) and the computation of the associated integral kernel operator. For clarity of exposition, we consider
only the simplest proposed version of this operator (6) but note that similar ideas may also be applied to (7) and (8). Furthermore, we will drop $\sigma$, $W$, and $b$ from (16) and simply consider the linear update
\[ u(x) = \int_D \kappa(x, y) v(y) \, d\nu(y) \quad \forall x \in D. \] (17)

To demonstrate the computational challenges associated with (17), let $\{x_1, \ldots, x_J\} \subset D$ be a uniformly-sampled $J$-point discretization of $D$. Recall that we assumed $d\nu(y) = dy$ and, for simplicity, suppose that $\nu(D) = 1$, then the Monte Carlo approximation of (17) is
\[ u(x_j) = \frac{1}{J} \sum_{l=1}^{J} \kappa(x_j, x_l) v(x_l), \quad j = 1, \ldots, J. \]

Therefore to compute $u$ on the entire grid requires $O(J^2)$ matrix-vector multiplications. Each of these matrix-vector multiplications requires $O(mn)$ operations; for the rest of the discussion, we treat $mn = O(1)$ as constant and consider only the cost with respect to $J$ the discretization parameter since $m$ and $n$ are fixed by the architecture choice whereas $J$ varies depending on required discretization accuracy and hence may be arbitrarily large. This cost is not specific to the Monte Carlo approximation but is generic for quadrature rules which use the entirety of the data. Therefore, when $J$ is large, computing (17) becomes intractable and new ideas are needed in order to alleviate this. Subsections 4.1-4.4 propose different approaches to the solution to this problem, inspired by classical methods in numerical analysis. We finally remark that, in contrast, computations with $W$, $b$, and $\sigma$ only require $O(J)$ operations which justifies our focus on computation with the kernel integral operator.

**Kernel Matrix.** It will often times be useful to consider the kernel matrix associated to $\kappa$ for the discrete points $\{x_1, \ldots, x_J\} \subset D$. We define the kernel matrix $K \in \mathbb{R}^{mJ \times nJ}$ to be the $J \times J$ block matrix with each block given by the value of the kernel i.e.
\[ K_{jl} = \kappa(x_j, x_l) \in \mathbb{R}^{m \times n}, \quad j, l = 1, \ldots, J \]
where we use $(j, l)$ to index an individual block rather than a matrix element. Various numerical algorithms for the efficient computation of (17) can be derived based on assumptions made about the structure of this matrix, for example, bounds on its rank or sparsity.

### 4.1 Graph Neural Operator (GNO)
We first outline the Graph Neural Operator (GNO) which approximates (17) by combining a Nyström approximation with domain truncation and is implemented with the standard framework of graph neural networks. This construction was originally proposed in Li et al. (2020c).

**Nyström approximation.** A simple yet effective method to alleviate the cost of computing (17) is employing a Nyström approximation. This amounts to sampling uniformly at random the points over which we compute the output function $u$. In particular, let $x_{k_1}, \ldots, x_{k_{J'}} \subset \{x_1, \ldots, x_J\}$ be $J' \ll J$ randomly selected points and, assuming $\nu(D) = 1$, approximate (17) by
\[ u(x_{k_j}) \approx \frac{1}{J'} \sum_{l=1}^{J'} \kappa(x_{k_j}, x_{k_l}) v(x_{k_l}), \quad j = 1, \ldots, J'. \]
We can view this as a low-rank approximation to the kernel matrix $K$, in particular,

$$K \approx K_{J,J'}K_{J',J'}K_{J,J} \quad (18)$$

where $K_{J,J'}$ is a $J' \times J'$ block matrix and $K_{J,J'}, K_{J'}$ are interpolation matrices, for example, linearly extending the function to the whole domain from the random nodal points. The complexity of this computation is $O(J'^2)$ hence it remains quadratic but only in the number of subsampled points $J'$ which we assume is much less than the number of points $J$ in the original discretization.

**Truncation.** Another simple method to alleviate the cost of computing (17) is to truncate the integral to a sub-domain of $D$ which depends on the point of evaluation $x \in D$. Let $s : D \to \mathcal{B}(D)$ be a mapping of the points of $D$ to the Lebesgue measurable subsets of $D$ denoted $\mathcal{B}(D)$. Define $d\nu(x,y) = 1_{s(x)}dy$ then (17) becomes

$$u(x) = \int_{s(x)} \kappa(x,y)v(y) \, dy \quad \forall x \in D. \quad (19)$$

If the size of each set $s(x)$ is smaller than $D$ then the cost of computing (19) is $O(c_s J^2)$ where $c_s < 1$ is a constant depending on $s$. While the cost remains quadratic in $J$, the constant $c_s$ can have a significant effect in practical computations, as we demonstrate in Section 6. For simplicity and ease of implementation, we only consider $s(x) = B(x,r) \cap D$ where $B(x,r) = \{y \in \mathbb{R}^d : \|y - x\|_d < r\}$ for some fixed $r > 0$. With this choice of $s$ and assuming that $D = [0,1]^d$, we can explicitly calculate that $c_s \approx r^d$.

Furthermore notice that we do not lose any expressive power when we make this approximation so long as we combine it with composition. To see this, consider the example of the previous paragraph where, if we let $r = \sqrt{2}$, then (19) reverts to (17). Pick $r < 1$ and let $L \in \mathbb{N}$ with $L \geq 2$ be the smallest integer such that $2^{L-1}r \geq 1$. Suppose that $u(x)$ is computed by composing the right hand side of (19) $L$ times with a different kernel every time. The domain of influence of $u(x)$ is then $B(x,2^{L-1}r) \cap D = D$ hence it is easy to see that there exist $L$ kernels such that computing this composition is equivalent to computing (17) for any given kernel with appropriate regularity. Furthermore the cost of this computation is $O(Lr^dJ^2)$ and therefore the truncation is beneficial if $r^d(\log_2 1/r + 1) < 1$ which holds for any $r < 1/2$ when $d = 1$ and any $r < 1$ when $d \geq 2$. Therefore we have shown that we can always reduce the cost of computing (17) by truncation and composition. From the perspective of the kernel matrix, truncation enforces a sparse, block diagonally-dominant structure at each layer. We further explore the hierarchical nature of this computation using the multipole method in subsection 4.3.

Besides being a useful computational tool, truncation can also be interpreted as explicitly building local structure into the kernel $\kappa$. For problems where such structure exists, explicitly enforcing it makes learning more efficient, usually requiring less data to achieve the same generalization error. Many physical systems such as interacting particles in an electric potential exhibit strong local behavior that quickly decays, making truncation a natural approximation technique.

**Graph Neural Networks.** We utilize the standard architecture of message passing graph networks employing edge features as introduced in Gilmer et al. (2017) to efficiently implement (17) on arbitrary discretizations of the domain $D$. To do so, we treat a discretization $\{x_1, \ldots, x_J\} \subset D$ as the nodes of a weighted, directed graph and assign edges to each node using the function $s : D \to \mathcal{B}(D)$ which, recall from the section on truncation, assigns to each point a domain of
integration. In particular, for $j = 1, \ldots, J$, we assign the node $x_j$ the value $v(x_j)$ and emanate from it edges to the nodes $s(x_j) \cap \{x_1, \ldots, x_J\} = \mathcal{N}(x_j)$ which we call the neighborhood of $x_j$. If $s(x) = D$ then the graph is fully-connected. Generally, the sparsity structure of the graph determines the sparsity of the kernel matrix $K$, indeed, the adjacency matrix of the graph and the block kernel matrix have the same zero entries. The weights of each edge are assigned as the arguments of the kernel. In particular, for the case of (17), the weight of the edge between nodes $x_j$ and $x_k$ is simply the concatenation $(x_j, x_k) \in \mathbb{R}^{2d}$. More complicated weighting functions are considered for the implementation of the integral kernel operators (7) or (8).

With the above definition the message passing algorithm of Gilmer et al. (2017), with averaging aggregation, updates the value $v(x_j)$ of the node $x_j$ to the value $u(x_j)$ as

$$u(x_j) = \frac{1}{|N(x_j)|} \sum_{y \in N(x_j)} \kappa(x_j, y)v(y), \quad j = 1, \ldots, J$$

which corresponds to the Monte-Carlo approximation of the integral (19). More sophisticated quadrature rules and adaptive meshes can also be implemented using the general framework of message passing on graphs, see, for example, Pfaff et al. (2020). We further utilize this framework in subsection 4.3.

**Convolutional Neural Networks.** Lastly, we compare and contrast the GNO framework to standard convolutional neural networks (CNNs). In computer vision, the success of CNNs has largely been attributed to their ability to capture local features such as edges that can be used to distinguish different objects in a natural image. This property is obtained by enforcing the convolution kernel to have local support, an idea similar to our truncation approximation. Furthermore, by directly using a translation invariant kernel, a CNN architecture becomes translation equivariant; this is a desirable feature for many vision models e.g. ones that perform segmentation. We will show that similar ideas can be applied to the neural operator framework to obtain an architecture with built-in local properties and translational symmetries that, unlike CNNs, remain consistent in function space.

To that end, let $\kappa(x, y) = \kappa(x - y)$ and suppose that $\kappa : \mathbb{R}^d \rightarrow \mathbb{R}^{m \times n}$ is supported on $B(0, r)$. Let $r^* > 0$ be the smallest radius such that $D \subseteq B(x^*, r^*)$ where $x^* \in \mathbb{R}^d$ denotes the center of mass of $D$ and suppose $r \ll r^*$. Then (17) becomes the convolution

$$u(x) = (\kappa * v)(x) = \int_{B(x, r) \cap D} \kappa(x - y)v(y) \, dy \quad \forall x \in D. \quad (20)$$

Notice that (20) is precisely (19) when $s(x) = B(x, r) \cap D$ and $\kappa(x, y) = \kappa(x - y)$. When the kernel is parameterized by e.g. a standard neural network and the radius $r$ is chosen independently of the data discretization, (20) becomes a layer of a convolution neural network that is consistent in function space. Indeed the parameters of (20) do not depend on any discretization of $v$. The choice $\kappa(x, y) = \kappa(x - y)$ enforces translational equivariance in the output while picking $r$ small enforces locality in the kernel; hence we obtain the distinguishing features of a CNN model.

We will now show that, by picking a parameterization that is *inconsistent* is functions space and applying a Monte Carlo approximation to the integral, (20) becomes a standard CNN. This is most easily demonstrated when $D = [0, 1]$ and the discretization $\{x_1, \ldots, x_J\}$ is equispaced i.e. $|x_{j+1} - x_j| = h$ for any $j = 1, \ldots, J - 1$. Let $k \in \mathbb{N}$ be an odd filter size and let $z_1, \ldots, z_k \in \mathbb{R}$ be the points $z_j = (j - 1 - (k - 1)/2)h$ for $j = 1, \ldots, k$. It is easy to see that $\{z_1, \ldots, z_k\} \subset B(0, (k - 1)h/2)$ which we choose as the support of $\kappa$. Furthermore, we parameterize $\kappa$ directly
by its pointwise values which are \( m \times n \) matrices at the locations \( z_1, \ldots, z_k \) thus yielding \( kmn \) parameters. Then (20) becomes

\[
\begin{align*}
    u(x)_p & \approx \frac{1}{k} \sum_{l=1}^{k} \sum_{q=1}^{n} \kappa(z_l)_p q \nu(x_j - z_l)_q, & j = 1, \ldots, J, \ p = 1, \ldots, m
\end{align*}
\]

where we define \( \nu(x) = 0 \) if \( x \not\in \{x_1, \ldots, x_J\} \). Up to the constant factor \( 1/k \) which can be re-absorbed into the parameterization of \( \kappa \), this is precisely the update of a stride 1 CNN with \( n \) input channels, \( m \) output channels, and zero-padding so that the input and output signals have the same length. This example can easily be generalized to higher dimensions and different CNN structures, we made the current choices for simplicity of exposition. Notice that if we double the amount of discretization points for \( \nu \) i.e. \( J \rightarrow 2J \) and \( h \rightarrow h/2 \), the support of \( \kappa \) becomes \( \bar{B}(0, (k-1)h/4) \) hence the model changes due to the discretization of the data. Indeed, if we take the limit to the continuum \( J \rightarrow \infty \), we find \( \bar{B}(0, (k-1)h/2) \rightarrow \{0\} \) hence the model becomes completely local. To fix this, we may try to increase the filter size \( k \) (or equivalently add more layers) simultaneously with \( J \), but then the number of parameters in the model goes to infinity as \( J \rightarrow \infty \) since, as we previously noted, there are \( kmn \) parameters in this layer. Therefore standard CNNs are not consistent models in function space. We demonstrate their inability to generalize to different resolutions in Section 6.

### 4.2 Low-rank Neural Operator (LNO)

By directly imposing that the kernel \( \kappa \) is of a tensor product form, we obtain a layer with \( \mathcal{O}(J) \) computational complexity that is similar to the DeepONet construction of Lu et al. (2019) discussed in Section 3.2, but parameterized to be consistent in function space. We term this construction the Low-rank Neural Operator (LNO) due to its equivalence to directly parameterizing a finite-rank operator. We start by assuming \( \kappa : D \times D \rightarrow \mathbb{R} \) is scalar valued and later generalize to the vector valued setting. We express the kernel as

\[
\kappa(x, y) = \sum_{j=1}^{r} \varphi^{(j)}(x) \psi^{(j)}(y) \quad \forall x, y \in D
\]

for some functions \( \varphi^{(1)}, \psi^{(1)}, \ldots, \varphi^{(r)}, \psi^{(r)} : D \rightarrow \mathbb{R} \) that are normally given as the components of two neural networks \( \varphi, \psi : D \rightarrow \mathbb{R}^r \) or a single neural network \( \Xi : D \rightarrow \mathbb{R}^{2r} \) which couples all functions through its parameters. With this definition, and supposing that \( n = m = 1 \), we have that (17) becomes

\[
\begin{align*}
    u(x) &= \int_{D} \sum_{j=1}^{r} \varphi^{(j)}(x) \psi^{(j)}(y) v(y) \ dy \\
         &= \sum_{j=1}^{r} \int_{D} \psi^{(j)}(y) v(y) \ dy \varphi^{(j)}(x) \\
         &= \sum_{j=1}^{r} (\psi^{(j)}, v) \varphi^{(j)}(x)
\end{align*}
\]
where $\langle \cdot, \cdot \rangle$ denotes the $L^2(D; \mathbb{R})$ inner product. Notice that the inner products can be evaluated independently of the evaluation point $x \in D$ hence the computational complexity of this method is $O(rJ)$ which is linear in the discretization.

We may also interpret this choice of kernel as directly parameterizing a rank $r \in \mathbb{N}$ operator on $L^2(D; \mathbb{R})$. Indeed, we have

$$u = \sum_{j=1}^{r} (\varphi^{(j)} \otimes \psi^{(j)}) v$$

which corresponds precisely to applying the SVD of a rank $r$ operator to the function $v$. Equation (21) makes natural the vector valued generalization. Assume $m, n \geq 1$ and $\varphi^{(j)} : D \to \mathbb{R}^m$ and $\psi^{(j)} : D \to \mathbb{R}^n$ for $j = 1, \ldots, r$ then, (21) defines an operator mapping $L^2(D; \mathbb{R}^m) \to L^2(D; \mathbb{R}^n)$ that can be evaluated as

$$u(x) = \sum_{j=1}^{r} \langle \psi^{(j)}, v \rangle_{L^2(D; \mathbb{R}^n)} \varphi^{(j)}(x) \quad \forall x \in D.$$ 

We again note the linear computational complexity of this parameterization. Finally, we observe that this method can be interpreted as directly imposing a rank $r$ structure on the kernel matrix. Indeed,

$$K = K_{Jr}K_{rJ}$$

where $K_{Jr}, K_{rJ}$ are $J \times r$ and $r \times J$ block matrices respectively. While this method enjoys a linear computational complexity, similar to the DeepONets of Lu et al. (2019), it also constitutes a linear approximation method which may not be able to effectively capture the solution manifold; see Section 3.2 for further discussion.

### 4.3 Multipole Graph Neural Operator (MGNO)

A natural extension to directly working with kernels in a tensor product form as in Section 4.2 is to instead consider kernels that can be well approximated by such a form. This assumption gives rise to the fast multipole method (FMM) which employs a multi-scale decomposition of the kernel in order to achieve linear complexity in computing (17); for a detailed discussion see e.g. (E, 2011, Section 3.2). FMM can be viewed as a systematic approach to combine the sparse and low-rank approximations to the kernel matrix. Indeed, the kernel matrix is decomposed into different ranges and a hierarchy of low-rank structures is imposed on the long-range components. We employ this idea to construct hierarchical, multi-scale graphs, without being constrained to particular forms of the kernel. We will elucidate the workings of the FMM through matrix factorization. This approach was first outlined in Li et al. (2020b) and is referred as the Multipole Graph Neural Operator (MGNO).

The key to the fast multipole method’s linear complexity lies in the subdivision of the kernel matrix according to the range of interaction, as shown in Figure 2

$$K = K_1 + K_2 + \ldots + K_L,$$  

where $K_{\ell}$ with $\ell = 1$ corresponds to the shortest-range interaction, and $\ell = L$ corresponds to the longest-range interaction; more generally index $\ell$ is ordered by the range of interaction. While the uniform grids depicted in Figure 2 produce an orthogonal decomposition of the kernel, the decomposition may be generalized to arbitrary discretizations by allowing slight overlap of the ranges.
The kernel matrix $K$ is decomposed with respect to its interaction ranges. $K_1$ corresponds to short-range interaction; it is sparse but full-rank. $K_3$ corresponds to long-range interaction; it is dense but low-rank.

![Hierarchical matrix decomposition](image)

**Multi-scale discretization.** Words seem to repeat what is written around (22) please simplify We construct $L \in \mathbb{N}$ levels of discretization, where the finest grid corresponds to the shortest-range interaction $K_1$, and the coarsest discretization corresponds to the longest-range interaction $K_L$. In general, the underlying discretization can be arbitrary and we produce a hierarchy of $L$ discretizations with a decreasing number of nodes $J_1 \geq \ldots \geq J_L$ and increasing kernel integration radius $r_1 \leq \ldots \leq r_L$. Therefore, the shortest-range interaction $K_1$ has a fine resolution but is truncated locally, while the longest-range interaction $K_L$ has a coarse resolution, but covers the entire domain. This is shown pictorially in Figure 2. The number of nodes $J_1 \geq \ldots \geq J_L$, and the integration radii $r_1 \leq \ldots \leq r_L$ are hyperparameter choices and can be picked so that the total computational complexity is linear in $J$.

A special case of this construction is when the grid is uniform. Then our formulation reduces to the standard fast multipole algorithm and the kernels $K_l$ form an orthogonal decomposition of the full kernel matrix $K$. Assuming the underlying discretization $\{x_1, \ldots, x_J\} \subset D$ is a uniform grid with resolution $s$ such that $s^d = J$, the $L$ multi-level discretizations will be grids with resolution $s_l = s/2^{l-1}$, and consequentially $J_l = s_l^d = (s/2^{l-1})^d$. In this case $r_l$ can be chosen as $1/s$ for $l = 1, \ldots, L$. To ensure orthogonality of the discretizations, the fast multipole algorithm sets the integration domains to be $B(x, r_l) \setminus B(x, r_{l-1})$ for each level $l = 2, \ldots, L$, so that the discretization on level $l$ does not overlap with the one on level $l - 1$. Details of this algorithm can be found in e.g. Greengard and Rokhlin (1997).

**Recursive low-rank decomposition.** The coarse discretization representation can be understood as recursively applying an give canonical citation for inducing points inducing points approximation: starting from a discretization with $J_1 = J$ nodes, we impose inducing points of size $J_2, J_3, \ldots, J_L$ which all admit a low-rank kernel matrix decomposition of the form (18). The original $J \times J$ kernel matrix $K_l$ is represented by a much smaller $J_l \times J_l$ kernel matrix, denoted by $K_{l,l}$. As shown in Figure 2, $K_1$ is full-rank but very sparse while $K_L$ is dense but low-rank. This is repeating something stated in previous paragraph. Such structure can be achieved by applying equation (18) recursively to equation (22), leading to the multi-resolution matrix factorization (Kondor et al., 2014):

$$K \approx K_{1,1} + K_{1,2}K_{2,2}K_{2,1} + K_{1,2}K_{2,3}K_{3,3}K_{3,2}K_{2,1} + \cdots$$

(23)

where $K_{1,1} = K_1$ represents the shortest range, $K_{1,2}K_{2,2}K_{2,1} \approx K_2$, represents the second shortest range, etc. The center matrix $K_{l,l}$ is a $J_l \times J_l$ kernel matrix corresponding to the $l$-level of the
discretization described above. The matrices $K_{l+1,l}, K_{l,l+1}$ are $J_l \times J_l$ and $J_l \times J_{l+1}$ wide and long respectively block transition matrices. Denote $v_l \in \mathbb{R}^{J_l \times n}$ for the representation of the input $v$ at each level of the discretization for $l = 1, \ldots, L$, and $u_l \in \mathbb{R}^{J_l \times n}$ for the output (assuming the inputs and outputs has the same dimension). We define the matrices $K_{l+1,l}, K_{l,l+1}$ as moving the representation $v_l$ between different levels of the discretization via an integral kernel that we learn. Combining with the truncation idea introduced in subsection 4.1, we define the transition matrices as discretizations of the following integral kernel operators:

$$K_{l,l} : v_l \mapsto u_l = \int_{B(x,r_{l,l})} \kappa_{l,l}(x,y) v_l(y) \, dy$$  \hspace{1cm} (24)

$$K_{l+1,l} : v_l \mapsto u_{l+1} = \int_{B(x,r_{l+1,l})} \kappa_{l+1,l}(x,y) v_l(y) \, dy$$  \hspace{1cm} (25)

$$K_{l,l+1} : v_{l+1} \mapsto u_l = \int_{B(x,r_{l,l+1})} \kappa_{l,l+1}(x,y) v_{l+1}(y) \, dy$$  \hspace{1cm} (26)

where each kernel $\kappa_{l,l} : D \times D \to \mathbb{R}^{n \times n}$ is parameterized as a neural network and learned.

**V-cycle Algorithm**  We present a V-cycle algorithm, see Figure 3 for efficiently computing (23). It consists of two steps: the **downward pass** and the **upward pass**. Denote the representation in downward pass and upward pass by $\tilde{v}$ and $\hat{v}$ respectively. In the downward step, the algorithm starts from the fine discretization representation $\tilde{v}_1$ and updates it by applying a downward transition $\tilde{v}_{l+1} = K_{l+1,l} \tilde{v}_l$. In the upward step, the algorithm starts from the coarse presentation $\hat{v}_L$ and updates it by applying an upward transition and the center kernel matrix $\hat{v}_l = K_{l+1,l} \hat{v}_{l+1} - K_{l,l} \hat{v}_l$. Notice that applying one level downward and upward exactly computes $K_{1,1}, K_{1,2} K_{2,2} K_{2,1}$, and a full $L$-level V-cycle leads to the multi-resolution decomposition (23).

Employing (24)-(26), we use $L$ neural networks $\kappa_{1,1}, \ldots, \kappa_{L,L}$ to approximate the kernel operators associated to $K_{l,l}$, and $2(L - 1)$ neural networks $\kappa_{1,2}, \kappa_{2,1}, \ldots$ to approximate the transitions $K_{l+1,l}, K_{l,l+1}$. Following the iterative architecture (5), we introduce the linear operator $W \in \mathbb{R}^{n \times n}$ (denoting it by $W_l$ for each corresponding resolution) to help regularize the iteration, as well as the nonlinear activation function $\sigma$ to increase the expensiveness. Since $W$ acts pointwise (requiring
remains the same for input and output), we employ it only along with the kernel $K_{l,l}$ and not the transitions. At each layer $t = 0, \ldots, T - 1$, we perform a full V-cycle as:

- **Downward Pass**
  
  For $l = 1, \ldots, L$:
  \[
  \tilde{v}_{l+1}^{(t+1)} = \sigma(\tilde{v}_{l+1}^{(t)} + K_{l+1,l} \tilde{v}_{l}^{(t+1)})
  \]  
  \[(27)\]

- **Upward Pass**
  
  For $l = L, \ldots, 1$:
  \[
  \tilde{v}_{l}^{(t+1)} = \sigma((W_{l} + K_{l,l})\tilde{v}_{l}^{(t+1)} + K_{l,l-1} \tilde{v}_{l-1}^{(t+1)}).
  \]  
  \[(28)\]

Notice that one full pass of the V-cycle algorithm defines a mapping $v \mapsto u$.

**Multi-level graphs.** We emphasize that we view the discretization $\{x_1, \ldots, x_J\} \subset D$ as a graph in order to facilitate an efficient implementation through the message passing graph neural network architecture. Since the V-cycle algorithm works at different levels of the discretization, we build multi-level graphs to represent the coarser and finer discretizations. We present and utilize two constructions of multi-level graphs, the orthogonal multipole graph and the generalized random graph.

The orthogonal multipole graph is the standard grid construction used in the fast multiple method which is adapted to a uniform grid, see e.g. (Greengard and Rokhlin, 1997). In this construction, the decomposition in (22) is orthogonal in that the finest graph only captures the closest range interaction, the second finest graph captures the second closest interaction minus the part already captured in the previous graph and so on, recursively. In particular, the ranges of interaction for each kernel do not overlap. While this construction is usually efficient, it is limited to uniform grids which may be a bottleneck for certain applications. Our second construction is the generalized random graph as shown in Figure 2 where the ranges of the kernels are allowed to overlap. The generalized random graph is very flexible as it can be applied on any domain geometry and discretization. Further it can also be combined with random sampling methods to work on problems where $J$ is very large or combined with active learning method to adaptively choose the regions where a finer discretization is needed.

**Linear complexity.** Each term in the decomposition (22) is represented by the kernel matrix $K_{l,l}$ for $l = 1, \ldots, L$, and $K_{l+1,l}, K_{l,l+1}$ for $l = 1, \ldots, L - 1$ corresponding to the appropriate sub-discretization. Therefore the complexity of the multipole method is \[\sum_{l=1}^{L} O(J_{l}^2 r_{l}^d) + \sum_{l=1}^{L-1} O(J_{l} J_{l+1}^d) = \sum_{l=1}^{L} O(J_{l}^2 r_{l}^d).\] By designing the sub-discretization so that $O(J_{l}^2 r_{l}^d) \leq O(J)$, we can obtain complexity linear in $J$. For example, when $d = 2$, pick $r_{l} = 1/\sqrt{J_{l}}$ and $J_{l} = O(2^{-l} J)$ such that $r_{L}$ is large enough so that there exists a ball of radius $r_{L}$ containing $D$. Then clearly $\sum_{l=1}^{L} O(J_{l}^2 r_{l}^d) = O(J)$. By combining with a Nyström approximation, we can obtain $O(J')$ complexity for some $J' \ll J$.

**4.4 Fourier Neural Operator (FNO)**

Instead of working with a kernel directly on the domain $D$, we may consider its representation in Fourier space and directly parameterize it there. This allows us to utilize Fast Fourier Transform (FFT) methods in order to compute the action of the kernel integral operator (17) with almost linear complexity. The method we outline was first described in Li et al. (2020a) and is termed the Fourier Neural Operator (FNO). For simplicity, we will assume that $D = \mathbb{T}^d$ is the unit torus and all
(a) The full architecture of neural operator: start from input \( a \). 1. Lift to a higher dimension channel space by a neural network \( P \). 2. Apply \( T \) (typically \( T = 4 \)) layers of integral operators and activation functions. 3. Project back to the target dimension by a neural network \( Q \). Output \( u \).

(b) Fourier layers: Start from input \( v \). On top: apply the Fourier transform \( F \); a linear transform \( R \) on the lower Fourier modes which also filters out the higher modes; then apply the inverse Fourier transform \( F^{-1} \). On the bottom: apply a local linear transform \( W \).

Figure 4: top: The architecture of the neural operators; bottom: Fourier layer.

functions are complex-valued. Let \( \mathcal{F} : L^1(D; \mathbb{C}^n) \to L^1(D; \mathbb{C}^n) \) Fourier transform maps \( L^2 \) into itself but maps \( L^1 \) into \( L^\infty \); maybe formulate on \( L^2 \)? denote the Fourier transform of a function \( v : D \to \mathbb{C}^n \) and \( \mathcal{F}^{-1} \) its inverse

\[
\begin{align*}
(Fv)_j(k) &= \int_D v_j(x)e^{-2\pi i \langle x, k \rangle} dx \\
(F^{-1}v)_j(x) &= \int_D v_j(k)e^{2\pi i \langle x, k \rangle} dk 
\end{align*}
\]

for \( j = 1, \ldots, n \) where \( i = \sqrt{-1} \) is the imaginary unit and \( \langle \cdot, \cdot \rangle \) denotes the Euclidean inner product on \( \mathbb{R}^d \). By letting \( \kappa(x, y) = \kappa(x - y) \) for some \( \kappa : D \to \mathbb{C}^{m \times n} \) in (17) and applying the convolution theorem, we find that

\[
u(x) = \mathcal{F}^{-1}(\mathcal{F}(\kappa) \cdot \mathcal{F}(v))(x) \quad \forall x \in D.
\]

We therefore propose to directly parameterize \( \kappa \) by its Fourier coefficients. We write

\[
u(x) = \mathcal{F}^{-1}(R_\phi \cdot \mathcal{F}(v))(x) \quad \forall x \in D
\]

where \( R_\phi \) is the Fourier transform of a periodic function \( \kappa : D \to \mathbb{C}^{n \times n} \) parameterized by some \( \phi \in \mathbb{R}^p \).

For frequency mode \( k \in D \), we have \( (\mathcal{F}v)(k) \in \mathbb{C}^n \) and \( R_\phi(k) \in \mathbb{C}^{m \times n} \). Notice that since we assume \( \kappa \) is periodic, it admits a Fourier series expansion, so we may work with the discrete modes \( k \in \mathbb{Z}^d \). We pick a finite-dimensional parameterization by truncating the Fourier series at a maximal number of modes \( k_{\text{max}} = |Z_{k_{\text{max}}} | = | \{ k \in \mathbb{Z}^d : |k_j| \leq k_{\text{max},j}, \text{ for } j = 1, \ldots, d \} | 
\)

We thus parameterize \( R_\phi \) directly as complex-valued \( (k_{\text{max}} \times m \times n) \)-tensor comprising a collection of truncated Fourier modes and therefore drop \( \phi \) from our notation. In the case where we have
real-valued $v$ and we want $u$ to also be real-valued, we impose that $\kappa$ is real-valued by enforcing conjugate symmetry in the parameterization i.e.

$$R(-k)_{j,l} = R^*(k)_{j,l}, \quad \forall k \in Z_{k_{\text{max}}}, \quad j = 1, \ldots, m, \quad l = 1, \ldots, n.$$ 

We note that the set $Z_{k_{\text{max}}}$ is not the canonical choice for the low frequency modes of $v_t$. Indeed, the low frequency modes are usually defined by placing an upper-bound on the $\ell_1$-norm of $k \in \mathbb{Z}^d$. We choose $Z_{k_{\text{max}}}$ as above since it allows for an efficient implementation. Figure 4 gives a pictorial representation of an entire Neural Operator architecture employing Fourier layer.

The discrete case and the FFT. Assuming the domain $D$ is discretized with $J \in \mathbb{N}$ points, we can treat $v \in \mathbb{C}^{J \times n}$ and $\mathcal{F}(v) \in \mathbb{C}^{J \times n}$. Since we convolve $v$ with a function which only has $k_{\text{max}}$ Fourier modes, we may simply truncate the higher modes to obtain $\mathcal{F}(v) \in \mathbb{C}^{k_{\text{max}} \times n}$. Multiplication by the weight tensor $R \in \mathbb{C}^{k_{\text{max}} \times m \times n}$ is then

$$(R \cdot (\mathcal{F}v_t))_{k,l} = \sum_{j=1}^{n} R_{k,l,j}(\mathcal{F}v)_{k,j}, \quad k = 1, \ldots, k_{\text{max}}, \quad l = 1, \ldots, m. \quad (30)$$

When the discretization is uniform with resolution $s_1 \times \cdots \times s_d = J$, $\mathcal{F}$ can be replaced by the Fast Fourier Transform. For $v \in \mathbb{C}^{J \times n}$, $k = (k_1, \ldots, k_d) \in \mathbb{Z}_{s_1} \times \cdots \times \mathbb{Z}_{s_d}$, and $x = (x_1, \ldots, x_d) \in D$, the FFT $\hat{\mathcal{F}}$ and its inverse $\hat{\mathcal{F}}^{-1}$ are defined as

$$\hat{\mathcal{F}} (v)_l(k) = \sum_{x_1=0}^{s_1-1} \cdots \sum_{x_d=0}^{s_d-1} v_l(x_1, \ldots, x_d) e^{-2i\pi \sum_{j=1}^{d} \frac{x_j k_j}{s_j}},$$

$$\hat{\mathcal{F}}^{-1} (v)(x) = \sum_{k_1=0}^{s_1-1} \cdots \sum_{k_d=0}^{s_d-1} v_l(k_1, \ldots, k_d) e^{2i\pi \sum_{j=1}^{d} \frac{x_j k_j}{s_j}}$$

for $l = 1, \ldots, n$. In this case, the set of truncated modes becomes

$$Z_{k_{\text{max}}} = \{(k_1, \ldots, k_d) \in \mathbb{Z}_{s_1} \times \cdots \times \mathbb{Z}_{s_d} \mid k_j \leq k_{\text{max},j} \text{ or } s_j - k_j \leq k_{\text{max},j}, \text{ for } j = 1, \ldots, d\}.$$

When implemented, $R$ is treated as a $(s_1 \times \cdots \times s_d \times m \times n)$-tensor and the above definition of $Z_{k_{\text{max}}}$ corresponds to the “corners” of $R$, which allows for a straight-forward parallel implementation of (30) via matrix-vector multiplication. In practice, we have found the choice $k_{\text{max},j} = 12$, which yields $k_{\text{max}} = 12^d$ parameters per channel, to be sufficient for all the tasks that we consider.

Choices for $R$. In general, $R$ can be defined to depend on $(\mathcal{F}a)$, the Fourier transform of the input $a \in \mathcal{A}$ to parallel our construction (7). Indeed, we can define $R_\phi : \mathbb{Z}^d \times \mathbb{C}^{d_a} \rightarrow \mathbb{C}^{m \times n}$ as a parametric function that maps $(k, (\mathcal{F}a)(k))$ to the values of the appropriate Fourier modes. We have experimented with the following parameterizations of $R_\phi$:

- **Direct.** Define the parameters $\phi_k \in \mathbb{C}^{m \times n}$ for each wave number $k$:

  $$R_\phi(k, (\mathcal{F}a)(k)) := \phi_k.$$

- **Linear.** Define the parameters $\phi_{k_1} \in \mathbb{C}^{m \times n \times d_a}, \phi_{k_2} \in \mathbb{C}^{m \times n}$ for each wave number $k$:

  $$R_\phi(k, (\mathcal{F}a)(k)) := \phi_{k_1}(\mathcal{F}a)(k) + \phi_{k_2}.$$
• *Feed-forward neural network.* Let $\Phi_{\phi} : \mathbb{Z}^d \times \mathbb{C}^{d_a} \rightarrow \mathbb{C}^{m \times n}$ be a neural network with parameters $\phi$:

$$R_{\phi}(k, (\mathcal{F}a)(k)) := \Phi_{\phi}(k, (\mathcal{F}a)(k)).$$

We find that the linear parameterization has a similar performance to the direct parameterization above, however, it is not as efficient both in terms of computational complexity and the number of parameters required. On the other hand, we find that the neural network parameterization has a worse performance. This is likely due to the discrete structure of the space $\mathbb{Z}^d$. Our experiments in this work focus on the direct parameterization presented above.

**Invariance to discretization.** The Fourier layers are discretization-invariant because they can learn from and evaluate functions which are discretized in an arbitrary way. Since parameters are learned directly in Fourier space, resolving the functions in physical space simply amounts to projecting on the basis $e^{2\pi i \langle x, k \rangle}$ which are well-defined everywhere on $\mathbb{C}^d$.

**Quasi-linear complexity.** The weight tensor $R$ contains $k_{\text{max}} < J$ modes, so the inner multiplication has complexity $O(k_{\text{max}})$. Therefore, the majority of the computational cost lies in computing the Fourier transform $\mathcal{F}(v)$ and its inverse. General Fourier transforms have complexity $O(J^2)$, however, since we truncate the series the complexity is in fact $O(J k_{\text{max}})$, while the FFT has complexity $O(J \log J)$. Generally, we have found using FFTs to be very efficient, however, a uniform discretization is required.

4.5 Summary

We summarize the main computational approaches presented in this section and their complexity:

• **GNO:** Subsample $J'$ points from the $J$-point discretization and compute the truncated integral

$$u(x) = \int_{B(x, r)} \kappa(x, y)v(y) \, dy$$

at a $O(J J')$ complexity.

• **LNO:** Decompose the kernel function tensor product form and compute

$$u(x) = \sum_{j=1}^{r} \langle \psi^{(j)}, v \rangle \varphi^{(j)}(x)$$

at a $O(J)$ complexity.

• **MGNO:** Compute a multi-scale decomposition of the kernel

$$K = K_{1,1} + K_{1,2}K_{2,2}K_{2,1} + K_{1,2}K_{2,3}K_{3,3}K_{3,2}K_{2,1} + \cdots$$

$$u(x) = (K v)(x)$$

at a $O(J)$ complexity.

• **FNO:** Parameterize the kernel in the Fourier domain and compute the using the FFT

$$u(x) = \mathcal{F}^{-1}(R_{\phi} \cdot \mathcal{F}(v))(x)$$

at a $O(J \log J)$ complexity.
5. Test Problems

A central application of neural operators is learning solution operators defined by parametric partial differential equations. In this section, we define four test problems for which we numerically study the approximation properties of neural operators. To that end, let \((A, U, F)\) be a triplet of Banach spaces. The first two problem classes considered are derived from the following general form

\[
L_a u = f
\]

for some \(f \in F\) and appropriate boundary conditions. For every \(a \in A\), \(L_a : U \to F\) is a, possibly nonlinear, partial differential operator, and \(u \in U\) corresponds to the solution of the PDE (35). The second class will be evolution equations with initial condition \(a \in A\) and solution at every time \(t > 0\) \(u(t) \in U\). We seek to learn the map from \(a\) to \(u := u(\tau)\) for some fixed time \(\tau > 0\); we will also study maps on paths (time-dependent solutions).

Our goal will be to learn the mappings \(G^\dagger : a \mapsto u\) or \(G^\dagger : f \mapsto u\); we will study both cases, depending on the test problem considered. We will define a probability measure \(\mu\) on \(A\) or \(F\) which will serve to define a model for likely input data. Furthermore, measure \(\mu\) will define a topology on the space of mappings in which \(G^\dagger\) lives, using the Bochner norm (3).

We will assume that each of the spaces \((A, U, F)\) are spaces of functions defined on a bounded domain \(D \subset \mathbb{R}^d\). All reported errors will be Monte-Carlo estimates of the relative error

\[
\mathbb{E}_{a \sim \mu} \left\| G^\dagger(a) - G_\theta(a) \right\|_{L^2(D)} \left/ \|G^\dagger(a)\|_{L^2(D)} \right.
\]

or equivalently replacing \(a\) with \(f\) in the above display and with the assumption that \(U \subset L^2(D)\). The domain \(D\) will be discretized, usually uniformly, with \(J \in \mathbb{N}\) points.

5.1 Poisson Equation

First we consider the one-dimensional Poisson equation with a zero boundary condition. In particular, (35) takes the form

\[
- \frac{d^2}{dx^2} u(x) = f(x), \quad x \in (0,1) \tag{36}
\]

\[
u(0) = u(1) = 0
\]

for some source function \(f \in H^{-1}((0,1);\mathbb{R})\). In particular, for \(D(L) := H^1_0((0,1);\mathbb{R}) \cap H^2((0,1);\mathbb{R})\), we have \(L : D(L) \to L^2((0,1);\mathbb{R})\) and note that \(L\) has no dependence on the parameter \(a \in A\).

We will consider the weak form of (36) and therefore the solution operator \(G^\dagger : H^{-1}((0,1);\mathbb{R}) \to H^1_0((0,1);\mathbb{R})\) defined as

\(G^\dagger : f \mapsto u\).

We define the probability measure \(\mu = N(0,C)\) where

\[
C = \left( L + I \right)^{-2},
\]

27
defined through the spectral theory of self-adjoint operators. Since $\mu$ charges a subset of $L^2((0, 1); \mathbb{R})$, we will learn $G^\dagger : L^2((0, 1); \mathbb{R}) \to H^1_0((0, 1); \mathbb{R})$ in the topology induced by (3).

In this setting, $G^\dagger$ has a closed-form solution given as

$$G^\dagger(f) = \int_0^1 G(\cdot, y) f(y) \, dy$$

where

$$G(x, y) = \frac{1}{2} (x + y - |y - x|) - xy, \quad \forall (x, y) \in [0, 1]^2$$

is the Green’s function. Note that while $G^\dagger$ is a linear operator, the Green’s function $G$ is non-linear as a function of its arguments. We will consider only a single layer of (5) with $\sigma_1 = \text{Id}$, $\mathcal{P} = \text{Id}$, $Q = \text{Id}$, $W_0 = 0$, $b_0 = 0$, and

$$K_0(f) = \int_0^1 \kappa_\theta(\cdot, y) f(y) \, dy$$

where $\kappa_\theta : \mathbb{R}^2 \to \mathbb{R}$ will be parameterized as a standard neural network with parameters $\theta$.

The purpose of the current example is two-fold. First we will test the efficacy of the neural operator framework in a simple setting where an exact solution is analytically available. Second we will show that by building in the right inductive bias, in particular, paralleling the form of the Green’s function solution, we obtain a model that generalizes outside the distribution $\mu$. That is, once trained, the model will generalize to any $f \in L^2((0, 1); \mathbb{R})$ that may be outside the support of $\mu$. For example, as defined, the random variable $f \sim \mu$ is a continuous function, however, if $\kappa_\theta$ approximates the Green’s function well then the model $G^\dagger$ will approximate well the solution to (36) even for discontinuous inputs.

Solutions to (36) are obtained by numerical integration using the Green’s function on a uniform grid with 85 collocation points. We use $N = 1000$ training examples.

5.2 Darcy Flow

We consider the steady state of Darcy Flow in two dimensions which is the second order elliptic equation

$$-\nabla \cdot (a(x) \nabla u(x)) = f(x), \quad x \in D$$

$$u(x) = 0, \quad x \in \partial D$$

(37)

where $D = (0, 1)^2$ is the unit square. In this setting $\mathcal{A} = L^\infty(D; \mathbb{R}_+)$, $\mathcal{U} = H^1_0(D; \mathbb{R})$, and $\mathcal{F} = H^{-1}(D; \mathbb{R})$. We fix $f \equiv 1$ and consider the weak form of (37) and therefore the solution operator $G^\dagger : L^\infty(D; \mathbb{R}_+) \to H^1_0(D; \mathbb{R})$ defined as

$$G^\dagger : a \mapsto u.$$  

(38)

Note that while (37) is a linear PDE, the solution operator $G^\dagger$ is nonlinear. We define the probability measure $\mu = T_\sharp N(0, C)$ where

$$C = (-\Delta + 9I)^{-2}$$

with $D(-\Delta)$ defined to impose zero Neumann boundary on the Laplacian. We view $T$ as a Nemyskii operator defined through the map $T : \mathbb{R} \to \mathbb{R}_+$ defined as

$$T(x) = \begin{cases} 12, & x \geq 0 \\ 3, & x < 0 \end{cases}.$$
The random variable $a \sim \mu$ is a piecewise-constant function with random interfaces given by the underlying Gaussian random field. Such constructions are prototypical models for many physical systems such as permeability in sub-surface flows and (in a vector generalization) material microstructures in elasticity.

Solutions to (37) are obtained using a second-order finite difference scheme on a uniform grid of size $421 \times 421$. All other resolutions are downsampled from this data set. We use $N = 1000$ training examples.

5.3 Burgers’ Equation

We consider the one-dimensional viscous Burgers’ equation

$$\frac{\partial}{\partial t} u(x, t) + \frac{1}{2} \frac{\partial}{\partial x} (u(x, t))^2 = \nu \frac{\partial^2}{\partial x^2} u(x, t), \quad x \in (0, 2\pi), t \in (0, \infty)$$

$$u(x, 0) = u_0(x), \quad x \in (0, 2\pi)$$

(39)

with periodic boundary conditions and a fixed viscosity $\nu = 0.1$. Let $\Psi : L^2_{\text{per}}((0, 2\pi); \mathbb{R}) \times \mathbb{R}_+ \to H^s_{\text{per}}((0, 2\pi); \mathbb{R})$, for any $s > 0$, be the flow map associated to (39), in particular,

$$\Psi(u_0, t) = u(\cdot, t), \quad t > 0.$$ 

We consider the solution operator defined by evaluating $\Psi$ at a fixed time. In particular, we let $G^\dagger : L^2_{\text{per}}((0, 2\pi); \mathbb{R}) \to H^s_{\text{per}}((0, 2\pi); \mathbb{R})$ be defined as

$$G^\dagger : u_0 \mapsto \Psi(u_0, 1).$$ 

(40)

We define the probability measure $\mu = N(0, C)$ where

$$C = 625 \left( -\frac{d^2}{dx^2} + 25 I \right)^{-2}$$

with domain of the Laplacian defined to impose periodic boundary conditions. We model the initial conditions $u_0 \sim \mu$ to (39) as $\mu$ charges a subset of $L^2_{\text{per}}((0, 2\pi); \mathbb{R})$.

Solutions to (39) are obtained using a pseudo-spectral split step method where the heat equation part is solved exactly in Fourier space then the non-linear part is advanced using a forward Euler method with a very small time step. We use a uniform spatial grid with $2^{13} = 8192$ collocation points and subsample all other resolutions from this data set. We use $N = 1000$ training examples.

5.4 Navier-Stokes Equation

We consider the two-dimensional Navier-Stokes equation for a viscous, incompressible fluid

$$\partial_t u(x, t) + u(x, t) \cdot \nabla u(x, t) + \nabla p(x, t) = \nu \Delta u(x, t) + f(x), \quad x \in \mathbb{T}^2, t \in (0, \infty)$$

$$\nabla \cdot u(x, t) = 0, \quad x \in \mathbb{T}^2, t \in [0, \infty)$$

$$u(x, 0) = u_0(x), \quad x \in \mathbb{T}^2$$

(41)

where $\mathbb{T}^2$ is the unit torus i.e. $[0, 1]^2$ equipped with periodic boundary conditions, and $\nu \in \mathbb{R}_+$ is a fixed viscosity. Here $u : \mathbb{T}^2 \times \mathbb{R}_+ \to \mathbb{R}^2$ is the velocity field. $p : \mathbb{T}^2 \times \mathbb{R}_+ \to \mathbb{R}^2$ is the pressure field. $f : \mathbb{T}^2 \to \mathbb{R}$ is a fixed forcing function. $\nu$ is viscosity.
Equivalently, we study its vorticity form to automatically impose the divergence-free condition
\[
\partial_t w(x,t) + u(x,t) \cdot \nabla w(x,t) = \nu \Delta w(x,t) + \nabla \times f(x), \quad x \in \mathbb{T}^2, t \in (0, \infty)
\]
\[
w(x,0) = w_0(x), \quad x \in \mathbb{T}^2
\]
(42)
where \( w = \nabla \times u : \mathbb{T}^2 \times \mathbb{R}_+ \to \mathbb{R} \) is the vorticity field. We define the forcing term as
\[
\nabla \times f(x_1, x_2) = 0.1(\sin(2\pi(x_1 + x_2)) + \cos(2\pi(x_1 + x_2))), \quad \forall (x_1, x_2) \in \mathbb{T}^2.
\]
The corresponding Reynolds number is estimated as \( Re = \frac{\sqrt{0.1}}{\nu(2\pi)^{3/2}} \) (Chandler and Kerswell 2013).

Let \( \Psi : L^2(\mathbb{T}^2; \mathbb{R}) \times \mathbb{R}_+ \to H^s(\mathbb{T}^2; \mathbb{R}), \) for any \( s > 0, \) be the flow map associated to (42), in particular,
\[
\Psi(w_0, t) = w(\cdot, t), \quad t > 0.
\]
Notice that this is well-defined for any \( w_0 \in L^2(\mathbb{T}; \mathbb{R}). \) We can see this through the stream-function formulation where the stream-function satisfies
\[
-\Delta \varphi_0(x) = w_0(x), \quad \forall x \in \mathbb{T}^2
\]
and the initial velocity is defined as
\[
u(x, 0) = \nabla \perp \varphi_0(x), \quad \forall x \in \mathbb{T}^2
\]
hence
\[
\nabla \cdot u(\cdot, 0) = \nabla \cdot \nabla \perp \varphi_0 = 0
\]
is divergence free.

We will define two notions of the solution operator. In the first, we will proceed as in the previous examples, in particular, \( G^\dagger : L^2(\mathbb{T}^2; \mathbb{R}) \to H^s(\mathbb{T}^2; \mathbb{R}) \) is defined as
\[
G^\dagger : w_0 \mapsto \Psi(w_0, T)
\]
(43)
for some fixed \( T > 0. \) In the second, we will map an initial part of the trajectory to a later part of the trajectory. In particular, we define \( G^\dagger : L^2(\mathbb{T}^2; \mathbb{R}) \times C((0, 10]; H^s(\mathbb{T}^2; \mathbb{R})) \to C((10, T]; H^s(\mathbb{T}^2; \mathbb{R})) \) by
\[
G^\dagger : (w_0, \Psi(w_0, t)|_{t \in (0, 10)}) \mapsto \Psi(w_0, t)|_{t \in (10, T]}
\]
(44)
for some fixed \( T > 10. \) We define the probability measure \( \mu = N(0, C) \) where
\[
C = T^{3/2}(-\Delta + 49I)^{-2.5}
\]
with periodic boundary conditions on the Laplacian. We model the initial vorticity \( w_0 \sim \mu \) to (42) as \( \mu \) charges a subset of \( L^2(\mathbb{T}^2; \mathbb{R}). \)

Solutions to (42) are obtained using a pseudo-spectral split step method where the viscous terms are advanced using a Crank–Nicolson update and the nonlinear and forcing terms are advanced using Heun’s method. Dealiasing is used with the 2/3 rule. For further details on this approach see (Chandler and Kerswell, 2013). Data is obtained on a uniform \( 256 \times 256 \) grid and all other resolutions are subsampled from this data set. We experiment with different viscosities \( \nu, \) final times \( T, \) and amounts of training data \( N. \)
5.4.1 BAYESIAN INVERSE PROBLEM

As an application of operator learning, we consider the inverse problem of recovering the initial vorticity in the Navier-Stokes equation (42) from partial, noisy observations at a much later time. Consider the first solution operator defined in subsection 5.4, in particular, \( G^\dagger : L^2(T^2; \mathbb{R}) \to H^s(T^2; \mathbb{R}) \) defined as

\[
G^\dagger : w_0 \mapsto \Psi(w_0, 50)
\]

where \( \Psi \) is the flow map associated to (42). We then consider the inverse problem

\[
y = O(G^\dagger(w_0)) + \eta
\]

of recovering \( w_0 \in L^2(T^2; \mathbb{R}) \) where \( O : H^s(T^2; \mathbb{R}) \to \mathbb{R}^{49} \) is the evaluation operator on a uniform \( 7 \times 7 \) interior grid, and \( \eta \sim \mathcal{N}(0, \Gamma) \) is observational noise with covariance \( \Gamma = (1/\gamma^2)I \) and \( \gamma = 0.1 \). We view (45) as the Bayesian inverse problem of recovering the posterior measure \( \pi^y \) from the prior measure \( \mu \). In particular, \( \pi^y \) has density with respect to \( \mu \), given by the Randon-Nikodym derivative

\[
\frac{d\pi^y}{d\mu}(w_0) \propto \exp(\|y - O(G^\dagger(w_0))\|_\Gamma^2)
\]

where \( \| \cdot \|_\Gamma = \| \Gamma^{-1/2} \cdot \| \) and \( \| \cdot \| \) is the Euclidean norm in \( \mathbb{R}^{49} \). For further details on the Bayesian approach, see e.g. (Cotter et al., 2009; Stuart, 2010).

We solve (45) by computing the posterior mean \( E_{w_0 \sim \pi^y}[w_0] \) using a Markov Chain Monte Carlo (MCMC) algorithm in order to draw samples from the posterior \( \pi^y \). We use the pre-conditioned Crank–Nicolson (pCN) method of Cotter et al. (2013) for this task. We employ pCN with both \( G^\dagger \) evaluated with the pseudo-spectral method described in section 5.4 and \( G_\theta \), the neural operator approximating \( G^\dagger \). After a 5,000 sample burn-in period, we generate 25,000 samples from the posterior using both approaches and use them to compute the posterior mean.

5.4.2 SPECTRA

Because of the constant in timeforcing term the energy reaches a non-zero equilibrium in time which is statistically reproducible for different initial conditions. To compare the complexity of the solution to the Navier-Stokes problem outlined in subsection 5.4 we show, in Figure 5, the Fourier spectrum of the solution data at time \( t = 50 \) for three different choices of the viscosity \( \nu \). We find that the rate of decay of the spectrum is \(-5/3\), matching what is expected in the turbulent regime (Kraichnan, 1967). Furthermore, we find that the energy does not decay in time due to the constant forcing term.

6. Numerical results

In this section, we compare the proposed neural operator with other supervised learning approaches, using the four test problems outlined in Section 5. In Subsection 6.1 we study the Poisson equation, and learning a Greens function; Subsection 6.2 consider the coefficient to solution map for steady Darcy flow, and the initial condition to solution at positive time map for Burgers equation. In Subsection 5.4 we study the Navier-Stokes equation.

We compare with a variety of architectures found by discretizing the data and applying finite-dimensional approaches, as well as with other operator-based approximation methods. We do not
The spectral decay of the Navier-stokes equation data. The y-axis is represents the value of each mode; the x-axis is the wavenumber $|k| = k_1 + k_2$. From left to right, the solutions have viscosity $\nu = 10^{-3}, 10^{-4}, 10^{-5}$ respectively.

Figure 5: Spectral Decay of Navier-Stokes.

compare against traditional solvers (FEM/FDM/Spectral), although our methods, once trained, enable evaluation of the input to output map orders of magnitude more quickly than by use of such traditional solvers on complex problems. We demonstrate the benefits of this speed-up in a prototypical application, Bayesian inversion, in Subsubsection 6.3.4.

All the computations are carried on a single Nvidia V100 GPU with 16GB memory. The code is available at https://github.com/zongyi-li/graph-pde and https://github.com/zongyi-li/fourier_neural_operator.

Setup of the four methods: We construct the neural operator by stacking four integral operator layers as specified in (5) with the ReLU activation. No batch normalization is needed. Unless otherwise specified, we use $N = 1000$ training instances and 200 testing instances. We use Adam optimizer to train for 500 epochs with an initial learning rate of 0.001 that is halved every 100 epochs. We set the channel dimensions $d_{v0} = \cdots = d_{v3} = 64$ for all one-dimensional problems and $d_{v0} = \cdots = d_{v3} = 32$ for all two-dimensional problems. The kernel networks $\kappa^{(0)}, \ldots, \kappa^{(3)}$ are standard feed-forward neural networks with three layers and widths of 256 units. We use the following abbreviations to denote the methods introduced in Section 4.

- **GNO:** The method introduced in subsection 4.1 truncating the integral to a ball with radius $r = 0.25$ and using the Nyström approximation with $J' = 300$ sub-sampled nodes.
- **LNO:** The low-rank method introduced in subsection 4.2 with rank $r = 4$.
- **MGNO:** The multipole method introduced in subsection 4.3. On the Darcy flow problem, we use the random construction with three graph levels, each sampling $J_1 = 400, J_2 = 100, J_3 = 25$ nodes respectively. On the Burgers’ equation problem, we use the orthogonal construction without sampling.
- **FNO:** The Fourier method introduced in subsection 4.4. We set $k_{\text{max},j} = 16$ for all one-dimensional problems and $k_{\text{max},j} = 12$ for all two-dimensional problems.

Remark on the resolution. Traditional PDE solvers such as FEM and FDM approximate a single function and therefore their error to the continuum decreases as the resolution is increased. The figures we show here exhibit something different: the error is independent of resolution, once enough resolution is used, but is not zero. This reflects the fact that there is a residual approximation error,
in the infinite dimensional limit, from the use of a finite-parametrized neural operator. Invariance of
the error with respect to (sufficiently fine) resolution is a desirable property that demonstrates that
an intrinsic approximation of the operator has been learned, independent of any specific discretiza-
tion. See Figure 7. Furthermore, resolution-invariant operators can do zero-shot super-resolution,
as shown in Subsubsection 6.3.1.

6.1 Poisson Equation

Recall the Poisson equation (36) introduced in subsection 5.1. We use a zero hidden layer neural
operator construction without lifting the input dimension. In particular, we simply learn a kernel
\( \kappa_\theta : \mathbb{R}^2 \rightarrow \mathbb{R} \) parameterized as a standard feed-forward neural network with parameters \( \theta \). Using
only \( N = 1000 \) training examples, we obtain a relative test error of \( 10^{-7} \). The neural operator gives
an almost perfect approximation to the true solution operator in the topology of (3).

To examine the quality of the approximation in the much stronger uniform topology, we check
whether the kernel \( \kappa_\theta \) approximates the Green’s function for this problem. To see why this is
enough, let \( K \subset L^2([0,1];\mathbb{R}) \) be a bounded set i.e.
\[
\|f\|_{L^2([0,1];\mathbb{R})} \leq M, \quad \forall f \in K
\]
and suppose that
\[
\sup_{(x,y) \in [0,1]^2} |\kappa_\theta(x,y) - G(x,y)| < \frac{\epsilon}{M},
\]
for some \( \epsilon > 0 \). Then it is easy to see that
\[
\sup_{f \in K} \|G^\dagger(f) - G_\theta(f)\|_{L^2([0,1];\mathbb{R})} < \epsilon,
\]
in particular, we obtain an approximation in the topology of uniform convergence over bounded
sets, while having trained only in the topology of the Bochner norm (3). Figure 6 shows the results
from which we can see that \( \kappa_\theta \) does indeed approximate the Green’s function well. This result
implies that by constructing a suitable architecture, we can generalize to the entire space and data
that is well outside the support of the training set.

6.2 Darcy and Burgers Equations

In the following section, we compare our four methods with different benchmarks on the Darcy
flow problem introduced in Subsection 5.2 and the Burgers’ equation problem introduced in Sub-
section 5.3. The solution operators of interest are defined by (38) and (40). We use the following
abbreviations for the methods against which we benchmark.

- **NN** is a standard point-wise feedforward neural network. It is mesh-free, but performs badly
due to lack of neighbor information.

- **FCN** is the state of the art neural network method based on Fully Convolution Network Zhu
and Zabaras (2018). It has a dominating performance for small grids \( s = 61 \). But fully
convolution networks are mesh-dependent and therefore their error grows when moving to a
larger grid.
left: learned kernel function; right: the analytic Green’s function.

This is a proof of concept of the graph kernel network on 1 dimensional Poisson equation and the comparison of learned and truth kernel.

Figure 6: Kernel for one-dimensional Green’s function, with the Nystrom approximation method

- **PCA+NN** is an instantiation of the methodology proposed in Bhattacharya et al. (2020): using PCA as an autoencoder on both the input and output data and interpolating the latent spaces with a neural network. The method provably obtains mesh-independent error and can learn purely from data, however the solution can only be evaluated on the same mesh as the training data.

- **RBM** is the classical Reduced Basis Method (using a PCA basis), which is widely used in applications and provably obtains mesh-independent error DeVore (2014). It has the good performance but the solutions can only be evaluated on the same mesh as the training data and one needs knowledge of the PDE to employ it.

- **DeepONet** is the Deep Operator network Lu et al. (2019) that has a nice approximation guarantee. We use the unstacked version with width 200.

6.2.1 Darcy Flow

The results of the experiments on Darcy flow are shown in Figure 7 and Table 1. The Fourier neural operator (FNO) obtains the lowest relative error compared to any of the benchmarks. Further, the error is invariant with respect to the resolution $s$, while the error of convolution neural network based methods (FCN) grows with the resolution. Compared to other neural operator methods such as GNO and MGNO that use Nyström sampling in physical space, the Fourier neural operator is both more accurate and more computationally efficient.

6.2.2 Burgers’ Equation

The results of the experiments on Burgers’ equation are shown in Figure 7 and Table 2. As for the Darcy problem, the Fourier neural operator obtains nearly one order of magnitude lower relative error compared to any benchmarks. The Fourier neural operator’s standard deviation is 0.0010 and its mean training error is 0.0012. If one replace the ReLU activation by GeLU, the test error will
(a) benchmarks on Burgers equation; (b) benchmarks on Darcy Flow for different resolutions; Train and test on the same resolution. For acronyms, see Section 6; details in Tables 2-1.

Figure 7: Benchmark on Burger’s equation and Darcy Flow

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<th>Networks</th>
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<th>$s = 141$</th>
<th>$s = 211$</th>
<th>$s = 421$</th>
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<td>0.1716</td>
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<td>0.0428</td>
<td>0.0420</td>
</tr>
<tr>
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<td><strong>0.0108</strong></td>
<td><strong>0.0109</strong></td>
<td><strong>0.0109</strong></td>
<td><strong>0.0098</strong></td>
</tr>
</tbody>
</table>

Table 1: Benchmarks on 2-d Darcy Flow on different resolutions $s$

Further reduce from **0.0018** to **0.0007**. We again observe the invariance of the error with respect to the resolution.

6.2.3 Zero-shot super-resolution.

The neural operator is mesh-invariant, so it can be trained on a lower resolution and evaluated at a higher resolution, without seeing any higher resolution data (zero-shot super-resolution). Figure 8 shows an example of the Darcy Equation where we train the GNO model on $16 \times 16$ resolution data in the setting above and transfer to $256 \times 256$ resolution, demonstrating super-resolution in space.
Table 2: Benchmarks on 1-d Burgers’ equation on different resolutions $s$

Graph kernel network for the solution of (5.2). It can be trained on a small resolution and will generalize to a large one. The Error is point-wise absolute squared error.

Figure 8: Darcy, trained on $16 \times 16$, tested on $241 \times 241$

6.3 Navier-Stokes Equation

In the following section, we compare our four methods with different benchmarks on the Navier-Stokes equation introduced in subsection 5.4. The operator of interest is given by (44). We use the following abbreviations for the methods against which we benchmark.

- **ResNet**: 18 layers of 2-d convolution with residual connections He et al. (2016).
- **U-Net**: A popular choice for image-to-image regression tasks consisting of four blocks with 2-d convolutions and deconvolutions Ronneberger et al. (2015).
- **TF-Net**: A network designed for learning turbulent flows based on a combination of spatial and temporal convolutions Wang et al. (2020).
- **FNO-2d**: 2-d Fourier neural operator with an auto-regressive structure in time. We use the Fourier neural operator to model the local evolution from the previous 10 time steps to the next one time step, and iteratively apply the model to get the long-term trajectory. We set and $k_{\text{max},j} = 12, d_v = 32$. 

<table>
<thead>
<tr>
<th>Networks</th>
<th>$s = 256$</th>
<th>$s = 512$</th>
<th>$s = 1024$</th>
<th>$s = 2048$</th>
<th>$s = 4096$</th>
<th>$s = 8192$</th>
</tr>
</thead>
<tbody>
<tr>
<td>NN</td>
<td>0.4714</td>
<td>0.4561</td>
<td>0.4803</td>
<td>0.4645</td>
<td>0.4779</td>
<td>0.4452</td>
</tr>
<tr>
<td>GCN</td>
<td>0.3999</td>
<td>0.4138</td>
<td>0.4176</td>
<td>0.4157</td>
<td>0.4191</td>
<td>0.4198</td>
</tr>
<tr>
<td>FCN</td>
<td>0.0958</td>
<td>0.1407</td>
<td>0.1877</td>
<td>0.2313</td>
<td>0.2855</td>
<td>0.3238</td>
</tr>
<tr>
<td>PCANN</td>
<td>0.0398</td>
<td>0.0395</td>
<td>0.0391</td>
<td>0.0383</td>
<td>0.0392</td>
<td>0.0393</td>
</tr>
<tr>
<td>DeepONet</td>
<td>0.0569</td>
<td>0.0617</td>
<td>0.0685</td>
<td>0.0702</td>
<td>0.0833</td>
<td>0.0857</td>
</tr>
<tr>
<td>GNO</td>
<td>0.0555</td>
<td>0.0594</td>
<td>0.0651</td>
<td>0.0663</td>
<td>0.0666</td>
<td>0.0699</td>
</tr>
<tr>
<td>LNO</td>
<td>0.0212</td>
<td>0.0221</td>
<td>0.0217</td>
<td>0.0219</td>
<td>0.0200</td>
<td>0.0189</td>
</tr>
<tr>
<td>MGNO</td>
<td>0.0243</td>
<td>0.0355</td>
<td>0.0374</td>
<td>0.0360</td>
<td>0.0364</td>
<td>0.0364</td>
</tr>
<tr>
<td>FNO</td>
<td><strong>0.0018</strong></td>
<td><strong>0.0018</strong></td>
<td><strong>0.0018</strong></td>
<td><strong>0.0019</strong></td>
<td><strong>0.0020</strong></td>
<td><strong>0.0019</strong></td>
</tr>
</tbody>
</table>
The learning curves on Navier-Stokes $\nu = 10^{-3}$ with different benchmarks. Train and test on the same resolution. For acronyms, see Section 6; details in Tables 3.

Figure 9: Benchmark on the Navier-Stokes

- **FNO-3d**: 3-d Fourier neural operator that directly convolves in space-time. We use the Fourier neural operator to model the global evolution from the initial 10 time steps directly to the long-term trajectory. We set $k_{max,j} = 12, d_v = 32$.

Table 3: Benchmarks on Navier Stokes (fixing resolution $64 \times 64$ for both training and testing)

<table>
<thead>
<tr>
<th>Config</th>
<th>Parameters</th>
<th>Time per epoch</th>
<th>$\nu = 10^{-3}$</th>
<th>$\nu = 10^{-4}$</th>
<th>$\nu = 10^{-4}$</th>
<th>$\nu = 10^{-5}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>$T = 50$</td>
<td>$N = 1000$</td>
<td>$T = 30$</td>
<td>$N = 1000$</td>
</tr>
<tr>
<td>FNO-3D</td>
<td>6,558,537</td>
<td>38.99s</td>
<td>0.0086</td>
<td>0.1918</td>
<td><strong>0.0820</strong></td>
<td>0.1893</td>
</tr>
<tr>
<td>FNO-2D</td>
<td>414,517</td>
<td>127.80s</td>
<td>0.0128</td>
<td><strong>0.1559</strong></td>
<td>0.0834</td>
<td><strong>0.1556</strong></td>
</tr>
<tr>
<td>U-Net</td>
<td>24,950,491</td>
<td>48.67s</td>
<td>0.0245</td>
<td>0.2051</td>
<td>0.1190</td>
<td>0.1982</td>
</tr>
<tr>
<td>TF-Net</td>
<td>7,451,724</td>
<td>47.21s</td>
<td>0.0225</td>
<td>0.2253</td>
<td>0.1168</td>
<td>0.2268</td>
</tr>
<tr>
<td>ResNet</td>
<td>266,641</td>
<td>78.47s</td>
<td>0.0701</td>
<td>0.2871</td>
<td>0.2311</td>
<td>0.2753</td>
</tr>
</tbody>
</table>

As shown in Table 3, the FNO-3D has the best performance when there is sufficient data ($\nu = 10^{-3}, N = 1000$ and $\nu = 10^{-4}, N = 10000$). For the configurations where the amount of data is insufficient ($\nu = 10^{-4}, N = 1000$ and $\nu = 10^{-5}, N = 1000$), all methods have $> 15\%$ error with FNO-2D achieving the lowest. Note that we only present results for spatial resolution $64 \times 64$ since all benchmarks we compare against are designed for this resolution. Increasing it degrades their performance while FNO achieves the same errors.
**2D and 3D Convolutions.** FNO-2D, U-Net, TF-Net, and ResNet all do 2D-convolution in the spatial domain and recurrently propagate in the time domain (2D+RNN). The operator maps the solution at previous time steps to the next time step (2D functions to 2D functions). On the other hand, FNO-3D performs convolution in space-time. It maps the initial time interval directly to the full trajectory (3D functions to 3D functions). The 2D+RNN structure can propagate the solution to any arbitrary time \( T \) in increments of a fixed interval length \( \Delta t \), while the Conv3D structure is fixed to the interval \([0, T]\) but can transfer the solution to an arbitrary time-discretization. We find the 3-d method to be more expressive and easier to train compared to its RNN-structured counterpart.

<table>
<thead>
<tr>
<th>Networks</th>
<th>( s = 64 )</th>
<th>( s = 128 )</th>
<th>( s = 256 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>FNO-3D</td>
<td>0.0098</td>
<td>0.0101</td>
<td>0.0106</td>
</tr>
<tr>
<td>FNO-2D</td>
<td>0.0129</td>
<td>0.0128</td>
<td>0.0126</td>
</tr>
<tr>
<td>U-Net</td>
<td>0.0253</td>
<td>0.0289</td>
<td>0.0344</td>
</tr>
<tr>
<td>TF-Net</td>
<td>0.0277</td>
<td>0.0278</td>
<td>0.0301</td>
</tr>
</tbody>
</table>

2D Navier-Stokes Equation with the parameter \( \nu = 10^{-3} \), \( N = 200 \), \( T = 20 \).

Table 4: Resolution study on Navier-stokes equation

### 6.3.1 Zero-shot super-resolution.

The neural operator is mesh-invariant, so it can be trained on a lower resolution and evaluated at a higher resolution, without seeing any higher resolution data (zero-shot super-resolution). Figure 1 shows an example where we train the FNO-3D model on \( 64 \times 64 \times 20 \) resolution data in the setting above with \((\nu = 10^{-4}, N = 10000)\) and transfer to \( 256 \times 256 \times 80 \) resolution, demonstrating super-resolution in space-time. The Fourier neural operator is the only model among the benchmarks (FNO-2D, U-Net, TF-Net, and ResNet) that can do zero-shot super-resolution; the method works well not only on the spatial but also on the temporal domain.

### 6.3.2 Spectral analysis

Figure 10 shows that all the methods are able to capture the spectral decay of the Navier-Stokes equation. Notice that, while the Fourier method truncates the higher frequency modes during the convolution, FNO can still recover the higher frequency components in the final prediction. Due to the way we parameterize \( R_\phi \), the function output by (29) has at most \( k_{\text{max},j} \) Fourier modes per channel. This, however, does not mean that the Fourier neural operator can only approximate functions up to \( k_{\text{max},j} \) modes. Indeed, the activation functions which occurs between integral operators and the final decoder network \( Q \) recover the high frequency modes. As an example, consider a solution to the Navier-Stokes equation with viscosity \( \nu = 10^{-3} \). Truncating this function at 20 Fourier modes yields an error around 2% as shown in Figure 11, while the Fourier neural operator learns the parametric dependence and produces approximations to an error of \( \leq 1\% \) with only \( k_{\text{max},j} = 12 \) parameterized modes.

### 6.3.3 Non-periodic boundary condition.

Traditional Fourier methods work only with periodic boundary conditions. However, the Fourier neural operator does not have this limitation. This is due to the linear transform \( W \) (the bias term)
The spectral decay of the predictions of different models on the Navier-Stokes equation. The y-axis is the spectrum; the x-axis is the wavenumber. Left is the spectrum of one trajectory; right is the average of 40 trajectories.

Figure 10: The spectral decay of the predictions of different methods

The error of truncation in one single Fourier layer without applying the linear transform $R$. The y-axis is the normalized truncation error; the x-axis is the truncation mode $k_{max}$.

Figure 11: Spectral Decay in term of $k_{max}$

which keeps the track of non-periodic boundary. As an example, the Darcy Flow and the time domain of Navier-Stokes have non-periodic boundary conditions, and the Fourier neural operator still learns the solution operator with excellent accuracy.

6.3.4 BAYESIAN INVERSE PROBLEM

As discussed in Section 5.4.1, we use the pCN method of Cotter et al. (2013) to draw samples from the posterior distribution of initial vorticities in the Navier-Stokes equation given sparse, noisy observations at time $T = 50$. We compare the Fourier neural operator acting as a surrogate model with the traditional solvers used to generate our train-test data (both run on GPU). We generate 25,000 samples from the posterior (with a 5,000 sample burn-in period), requiring 30,000 evaluations of the forward operator.

As shown in Figure 12, FNO and the traditional solver recover almost the same posterior mean which, when pushed forward, recovers well the later-time solution of the Navier-Stokes equation.
In sharp contrast, FNO takes 0.005s to evaluate a single instance while the traditional solver, after being optimized to use the largest possible internal time-step which does not lead to blow-up, takes 2.2s. This amounts to 2.5 minutes for the MCMC using FNO and over 18 hours for the traditional solver. Even if we account for data generation and training time (offline steps) which take 12 hours, using FNO is still faster. Once trained, FNO can be used to quickly perform multiple MCMC runs for different initial conditions and observations, while the traditional solver will take 18 hours for every instance. Furthermore, since FNO is differentiable, it can easily be applied to PDE-constrained optimization problems in which adjoint calculations are used as part of the solution procedure.

The top left panel shows the true initial vorticity while bottom left panel shows the true observed vorticity at $T = 50$ with black dots indicating the locations of the observation points placed on a $7 \times 7$ grid. The top middle panel shows the posterior mean of the initial vorticity given the noisy observations estimated with MCMC using the traditional solver, while the top right panel shows the same thing but using FNO as a surrogate model. The bottom middle and right panels show the vorticity at $T = 50$ when the respective approximate posterior means are used as initial conditions.

Figure 12: Results of the Bayesian inverse problem for the Navier-Stokes equation.

6.4 Discussion and Comparison of the Four methods

In this section we will compare the four methods in term of expressiveness, complexity, refinability, and ingenuity.

6.4.1 Ingenuity

First we will discuss ingenuity, in other words, the design of the frameworks. The first method, GNO, relies on the Nyström approximation of the kernel, or the Monte Carlo approximation of the integration. It is the most simple and straightforward method. The second method, LNO, relies on the low-rank decomposition of the kernel operator. It is efficient when the kernel has a near low-rank structure. The third method, MGNO, is the combination of the first two. It has a hierarchical,
multi-resolution decomposition of the kernel. The last one, FNO, is different from the first three; it restricts the integral kernel to induce a convolution.

GNO and MGNO are implemented using graph neural networks, which helps to define sampling and integration. The graph network library also allows sparse and distributed message passing. The LNO and FNO don’t have sampling. They are faster without using the graph library.

<table>
<thead>
<tr>
<th>scheme</th>
<th>graph-based</th>
<th>kernel network</th>
</tr>
</thead>
<tbody>
<tr>
<td>GNO</td>
<td>Nyström approximation</td>
<td>Yes</td>
</tr>
<tr>
<td>LNO</td>
<td>Low-rank approximation</td>
<td>No</td>
</tr>
<tr>
<td>MGNO</td>
<td>Multi-level graphs on GNO</td>
<td>Yes</td>
</tr>
<tr>
<td>FNO</td>
<td>Convolution theorem; Fourier features</td>
<td>No</td>
</tr>
</tbody>
</table>

Table 5: Ingenuity

6.4.2 Expressiveness

We measure the expressiveness by the training and testing error of the method. The full $O(J^2)$ integration always has the best results, but it is usually too expensive. As shown in the experiments 6.2.1 and 6.2.2. GNO usually has good accuracy, but its performance suffers from sampling. LNO works the best on the 1d problem (Burgers equation). It has difficulty on the 2d problem because it doesn’t employ sampling to speed-up evaluation. MGNO has the multi-level structure, which gives it the benefit of the first two. Finally, FNO has overall the best performance. It is also the only method that can capture the challenging Navier-Stokes equation.

6.4.3 Complexity

The complexity of the four methods are list in Table 6. GNO and MGNO have sampling. Their complexity depends on the number of nodes sampled $J'$. When using the full nodes. They are still quadratic. LNO has the lowest complexity $O(J)$. FNO, when using the fast Fourier transform, has complexity $O(J \log J)$.

In practice. FNO is faster then the other three methods because it doesn’t have the kernel network $\kappa$. MGNO is relatively slower because of its multi-level graph structure.

<table>
<thead>
<tr>
<th>Complexity</th>
<th>Time per epochs in training</th>
</tr>
</thead>
<tbody>
<tr>
<td>GNO</td>
<td>$O(J'^2 r^2)$</td>
</tr>
<tr>
<td>LNO</td>
<td>$O(J)$</td>
</tr>
<tr>
<td>MGNO</td>
<td>$\sum_i O(J_i^2 r_i^2) \sim O(J)$</td>
</tr>
<tr>
<td>FNO</td>
<td>$(J \log J)$</td>
</tr>
</tbody>
</table>

The theoretical time complexity and the empirical time complexity. Roundup to second (on a single Nvidia V100 GPU).

Table 6: Complexity
6.4.4 Refinability

Refineability measures the number of parameters used in the framework. Table 7 lists the accuracy of the relative error on Darcy Flow with respect to different number of parameters. Because GNO, LNO, and MGNO have the kernel networks, the slope of their error rates are flat: they can work with a very small number of parameters. On the other hand, FNO does not have the sub-network. It needs at a larger magnitude of parameters to obtain an acceptable error rate.

<table>
<thead>
<tr>
<th>Number of parameters</th>
<th>$10^3$</th>
<th>$10^4$</th>
<th>$10^5$</th>
<th>$10^6$</th>
</tr>
</thead>
<tbody>
<tr>
<td>GNO</td>
<td>0.075</td>
<td>0.065</td>
<td>0.060</td>
<td>0.035</td>
</tr>
<tr>
<td>LNO</td>
<td>0.080</td>
<td>0.070</td>
<td>0.060</td>
<td>0.040</td>
</tr>
<tr>
<td>MGNO</td>
<td>0.070</td>
<td>0.050</td>
<td>0.040</td>
<td>0.030</td>
</tr>
<tr>
<td>FNO</td>
<td>0.200</td>
<td>0.035</td>
<td>0.020</td>
<td>0.015</td>
</tr>
</tbody>
</table>

The relative error on Darcy Flow with respect to different number of parameters. The errors above are approximated value roundup to 0.05. They are the lowest test error achieved by the model, given the model’s number of parameters $|\theta|$ is bounded by $10^3, 10^4, 10^5, 10^6$ respectively.

Table 7: Refinability

7. Conclusions

We have introduced the concept of Neural Operator, the goal being to construct a neural network architecture adapted to the problem of mapping elements of one function space into elements of another function space. The network is comprised of four steps which, in turn, (i) extract features from the input functions, (ii) iterate a recurrent neural network on feature space, defined through composition of a sigmoid function and a nonlocal operator, and (iii) a final mapping from feature space into the output function.

We have studied four nonlocal operators in step (iii), one based on graph kernel networks, one based on the low-rank decomposition, one based on the multi-level graph structure, and the last one based on convolution in Fourier space. The designed network architectures are constructed to be mesh-free and our numerical experiments demonstrate that they have the desired property of being able to train and generalize on different meshes. This is because the networks learn the mapping between infinite-dimensional function spaces, which can then be shared with approximations at different levels of discretization. A further advantage of the integral operator approach is that data may be incorporated on unstructured grids, using the Nyström approximation; these methods, however, are quadratic in the number of discretization points; we describe variants on this methodology, using low rank and multiscale ideas, to reduce this complexity. On the other hand the Fourier approach leads directly to fast methods, linear-log linear in the number of discretization points, provided structured grids are used. We demonstrate that our method can achieve competitive performance with other mesh-free approaches developed in the numerical analysis community, and that it beats state-of-the-art neural network approaches on large grids, which are mesh-dependent. The methods developed in the numerical analysis community are less flexible than the approach we introduce here, relying heavily on the structure of an underlying PDE mapping input to output; our method is entirely data-driven.
7.1 Future directions

We foresee three main directions in which this work will develop: firstly as a method to speed-up scientific computing tasks which involve repeated evaluation of a mapping between spaces of functions, following the example of the Bayesian inverse problem 6.3.4, or when the underlying model is unknown as in computer vision or robotics; and secondly the development of more advanced methodologies beyond the four approximation schemes presented in Section 4 that are more efficient or better in specific situations; thirdly, the development of an underpinning theory which captures the expressive power, and approximation error properties, of the proposed neural network.

7.1.1 New Applications

The proposed neural operator is a blackbox surrogate model for function-to-function mappings. It naturally fits into solving PDEs for physics and engineering problems. In the paper we mainly studied three partial differential equations: Darcy Flow, Burgers’ equation, and Navier-Stokes equation, which cover a board range of scenarios. Due to its blackbox structure, the neural operator is easily applied on other problems. We foresee applications on more challenging turbulent flows such as climate models, sharper coefficients contrast raising in geological models, and general physics simulation for games and visual effects. The operator setting leads to an efficient and accurate representation, and the resolution-invariant properties make it possible to training on a smaller resolution dataset, and be evaluated on arbitrarily large resolution.

The operator learning setting is not restricted to scientific computing. For example, in computer vision, images can naturally be viewed as real-valued functions on 2-d domains and videos simply add a temporal structure. Our approach is therefore a natural choice for problems in computer vision where invariance to discretization is crucial. We leave this as an interesting future direction.

7.1.2 New Methodologies

There is much room for improvement upon the current methodologies given their excellent performance. The full $O(J^2)$ integration method still outperforms the Fourier method by about 40%. It is worth while to develop more advanced integration techniques or approximation schemes that follows the neural operator framework. For example, one can use adaptive graph or probability estimation in the Nyström approximation. It is also possible to use other basis than the Fourier basis such as the PCA basis and Chebyshev basis.

Another direction for new methodologies is to combine the neural operator in other settings. The current problem is set as a supervised learning problem. Instead, one can combine the neural operator with solvers Pathak et al. (2020); Um et al. (2020b), augmenting and correcting the solvers to get faster and more accuracy approximation. One can also combine neural operator with Physics-informed neural networks (PINNs) Raissi et al. (2019), using neural operators to generate a context grid that help the PINN.
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References


Steven Strogatz. Loves me, loves me not (do the math). 2009.


## Appendix A.

<table>
<thead>
<tr>
<th>Notation</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Operator learning</strong></td>
<td></td>
</tr>
<tr>
<td>$D \subset \mathbb{R}^d$</td>
<td>The spatial domain for the PDE</td>
</tr>
<tr>
<td>$x \in D$</td>
<td>Points in the the spatial domain</td>
</tr>
<tr>
<td>$a \in \mathcal{A} = (D; \mathbb{R}^{d_a})$</td>
<td>The input coefficient functions</td>
</tr>
<tr>
<td>$u \in \mathcal{U} = (D; \mathbb{R}^{d_u})$</td>
<td>The target solution functions</td>
</tr>
<tr>
<td>$D_j$</td>
<td>The discretization of $(a_j, u_j)$</td>
</tr>
<tr>
<td>$\mathcal{G}^T : \mathcal{A} \to \mathcal{U}$</td>
<td>The operator mapping the coefficients to the solutions</td>
</tr>
<tr>
<td>$\mu$</td>
<td>A probability measure where $a_j$ sampled from.</td>
</tr>
<tr>
<td><strong>Neural operator</strong></td>
<td></td>
</tr>
<tr>
<td>$v(x) \in \mathbb{R}^{d_v}$</td>
<td>The neural network representation of $u(x)$</td>
</tr>
<tr>
<td>$d_a$</td>
<td>Dimension of the input $a(x)$.</td>
</tr>
<tr>
<td>$d_u$</td>
<td>Dimension of the output $u(x)$.</td>
</tr>
<tr>
<td>$d_v$</td>
<td>The dimension of the representation $v(x)$.</td>
</tr>
<tr>
<td>$t = 0, \ldots, T$</td>
<td>The time steps (layers)</td>
</tr>
<tr>
<td>$\mathcal{P}, \mathcal{Q}$</td>
<td>The pointwise linear transformation $\mathcal{P} : a(x) \mapsto v_0(x)$ and $\mathcal{Q} : v_T(x) \mapsto u(x)$.</td>
</tr>
<tr>
<td>$\mathcal{K}$</td>
<td>The integral operator in the iterative update $v_t \mapsto v_{t+1}$.</td>
</tr>
<tr>
<td>$\kappa : \mathbb{R}^{2(d+1)} \to \mathbb{R}^{d_v \times d_v}$</td>
<td>The kernel maps $(x, y, a(x), a(y))$ to a $d_v \times d_v$ matrix</td>
</tr>
<tr>
<td>$K \in \mathbb{R}^{n \times n \times d_v \times d_v}$</td>
<td>The kernel matrix with $K_{xy} = \kappa(x, y)$.</td>
</tr>
<tr>
<td>$W \in \mathbb{R}^{d_v \times d_v}$</td>
<td>The pointwise linear transformation used as the bias term in the iterative update.</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>The activation function.</td>
</tr>
</tbody>
</table>

In the paper, we will use lowercase letters such as $v, u$ to represent vectors and functions; uppercase letters such as $W, K$ to represent matrices or discretized transformations; and calligraphic letters such as $\mathcal{G}, \mathcal{F}$ to represent operators.

Table 8: Table of notations: operator learning and neural operators