The Mathematics of Artificial Intelligence

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Abstract

We currently witness the spectacular success of artificial intelligence in both science and public life. However, the development of a rigorous mathematical foundation is still at an early stage. In this survey article, which is based on an invited lecture at the International Congress of Mathematicians 2022, we will in particular focus on the current “workhorse” of artificial intelligence, namely deep neural networks. We will present the main theoretical directions along with several exemplary results and discuss key open problems.

1 Introduction

Artificial intelligence is currently leading to one breakthrough after the other, both in public with, for instance, autonomous driving and speech recognition, and in the sciences in areas such as medical diagnostics or molecular dynamics. In addition, research on artificial intelligence and, in particular, on its theoretical foundations, is progressing at an unprecedented rate. One can envision that according methodologies will in the future drastically change the way we live in numerous respects.

1.1 The Rise of Artificial Intelligence

Artificial intelligence is however not a new phenomenon. In fact, already in 1943, McCulloch and Pitts started to develop algorithmic approaches to learning by mimicking the functionality of the human brain, through artificial neurons which are connected and arranged in several layers to form artificial neural networks. Already at that time, they had a vision for the implementation of artificial intelligence. However, the community did not fully recognize the potential of neural networks. Therefore, this first wave of artificial intelligence was not successful and vanished. Around 1980, machine learning became popular again, and several highlights can be reported from that period.

The real breakthrough and with it a new wave of artificial intelligence came around 2010 with the extensive application of deep neural networks. Today, this model might be considered the “workhorse” of artificial intelligence, and in this article we will focus predominantly on this approach. The structure of deep neural networks is precisely the structure McCulloch and Pitts introduced, namely numerous consecutive layers of artificial neurons. Today two main obstacles from previous years have also been eliminated; due to the drastic improvement of computing power the training of neural networks with hundreds of layers in the sense of deep neural networks is feasible, and we are living in the age of data, hence vast amounts of training data are easily available.

1.2 Impact on Mathematics

The rise of artificial intelligence also had a significant impact on various fields of mathematics. Maybe the first area which embraced these novel methods was the area of inverse problems, in particular, imaging science where such approaches have been used to solve highly ill-posed problems such as denoising, inpainting, superresolution, or (limited-angle) computed tomography, to name a few. One might note that due to the lack of a precise mathematical model of what an image is, this area is particularly suitable for learning methods. Thus, after a few years, a change of paradigm could be observed, and novel solvers are typically at least to some extent based on methods from artificial intelligence. We will discuss further details in Subsection 4.1.
The area of partial differential equations was much slower to embrace these new techniques, the reason being that it was not per se evident what the advantage of methods from artificial intelligence for this field would be. Indeed, there seems to be no need to utilize learning-type methods, since a partial differential equation is a rigorous mathematical model. But, lately, the observation that deep neural networks are able to beat the curse of dimensionality in high dimensional settings led to a change of paradigm in this area as well. Research at the intersection of numerical analysis of partial differential equations and artificial intelligence therefore accelerated since about 2017. We will delve further into this topic in Subsection 4.2.

1.3 Problems of Artificial Intelligence

However, as promising as all these developments seem to be, a word of caution is required. Besides the fact that the practical limitations of methods such as deep neural networks have not been explored at all and at present neural networks are still considered a “jack of all trades”, it is even more worrisome that a comprehensive theoretical foundation is completely lacking. This was very prominently stated during the major conference in artificial intelligence and machine learning, which is NIPS (today called NeurIPS) in 2017, when Ali Rahimi from Google received the Test of Time Award and during his plenary talk stated that “Machine learning has become a form of alchemy”. This raised a heated discussion to which extent a theoretical foundation does exist and is necessary at all. From a mathematical viewpoint, it is crystal clear that a fundamental mathematical understanding of artificial intelligence is inevitably necessary, and one has to admit that its development is currently in a preliminary state at best.

This lack of mathematical foundations, for instance, in the case of deep neural networks, results in a time-consuming search for a suitable network architecture, a highly delicate trial-and-error-based (training) process, and missing error bounds for the performance of the trained neural network. One needs to stress that, in addition, such approaches also sometimes unexpectedly fail dramatically when a small perturbation of the input data causes a drastic change of the output leading to radically different—and often wrong—decisions. Such adversarial examples are a well-known problem, which becomes severe in sensitive applications such as when a minor alterations of traffic signs, e.g, the placement of stickers, causes autonomous vehicles to suddenly reach an entirely wrong decision. It is evident that such robustness problems can only be tackled by a profound mathematical approach.

1.4 A Need for Mathematics

These considerations show that there is a tremendous need for mathematics in the area of artificial intelligence. And, in fact, one can currently witness that numerous mathematicians move to this field, bringing in their own expertise. Indeed, as we will discuss in Subsection 2.4, basically all areas of mathematics are required to tackle the various difficult, but exciting challenges in the area of artificial intelligence.

One can identify two different research directions at the intersection of mathematics and artificial intelligence:

- **Mathematical Foundations for Artificial Intelligence.** This direction aims for deriving a deep mathematical understanding. Based on this it strives to overcome current obstacles such as the lack of robustness or places the entire training process on solid theoretical feet.

- **Artificial Intelligence for Mathematical Problems.** This direction focuses on mathematical problem settings such as inverse problems and partial differential equations with the goal to employ methodologies from artificial intelligence to develop superior solvers.

1.5 Outline

Both research directions will be discussed in this survey paper, showcasing some novel results and pointing out key future challenges for mathematics. We start with an introduction into the mathematical setting, stating the main definitions and notations (see Section 2). Next, in Section 3 we delve into the first main direction, namely mathematical foundations for artificial intelligence, and discuss the research threads of expressivity, optimization,
generalization, and explainability. Section 4 is then devoted to the second main direction, which is artificial intelligence for mathematical problems, and we highlight some exemplary results. Finally, Section 5 states the seven main mathematical problems and concludes this article.

2 The Mathematical Setting of Artificial Intelligence

We now get into more details on the precise definition of a deep neural network, which is after all a purely mathematical object. We will also touch upon the typical application setting and training process, as well as on the current key mathematical directions.

2.1 Definition of Deep Neural Networks

The core building blocks are, as said, artificial neurons. For their definition, let us recall the structure and functionality of a neuron in the human brain. The basic elements of such a neuron are dendrites, through which signals are transmitted to its soma while being scaled/amplified due to the structural properties of the respective dendrites. In the soma of the neuron, those incoming signals are accumulated, and a decision is reached whether to fire to other neurons or not, and also with which strength.

This forms the basis for a mathematical definition of an artificial neuron.

Definition 2.1. An artificial neuron with weights \( w_1, \ldots, w_n \in \mathbb{R}, \) bias \( b \in \mathbb{R}, \) and activation function \( \rho : \mathbb{R} \to \mathbb{R} \) is defined as the function \( f : \mathbb{R}^n \to \mathbb{R} \) given by

\[
f(x_1, \ldots, x_n) = \rho \left( \sum_{i=1}^{n} x_i w_i - b \right) = \rho (\langle x, w \rangle - b),
\]

where \( w = (w_1, \ldots, w_n) \) and \( x = (x_1, \ldots, x_n) \).

By now, there exists a zoo of activation functions with the most well-known ones being as follows:

1. Heaviside function \( \rho(x) = \begin{cases} 1, & x > 0, \\ 0, & x \leq 0. \end{cases} \)
2. Sigmoid function \( \rho(x) = \frac{1}{1 + e^{-x}}. \)
3. Rectifiable Linear Unit (ReLU) \( \rho(x) = \max\{0, x\}. \)

We remark that of these examples, the by far most extensively used activation function is the ReLU due to its simple piecewise linear structure, which is advantageous in the training process and still allows superior performance.

Similar to the structure of a human brain, these artificial neurons are now being concatenated and arranged in layers, leading to an (artificial feed-forward) neural network. Due to the particular structure of artificial neurons, such a neural network consists of compositions of affine linear maps and activation functions. Traditionally, a deep neural network is then defined as the resulting function. From a mathematical standpoint, this bears the difficulty that different arrangements lead to the same function. Therefore, sometimes a distinction is made between the architecture of a neural network and the corresponding realization function (see, e.g., [6]). For this article, we will however avoid such technical delicacies and present the most standard definition.

Definition 2.2. Let \( d \in \mathbb{N} \) be the dimension of the input layer, \( L \) the number of layers, \( N_0 := d, N_\ell, \ell = 1, \ldots, L, \) the dimensions of the hidden and last layer, \( \rho : \mathbb{R} \to \mathbb{R} \) a (non-linear) activation function, and, for \( \ell = 1, \ldots, L, \) let \( T_\ell \) be the affine-linear functions

\[
T_\ell : \mathbb{R}^{N_{\ell-1}} \to \mathbb{R}^{N_\ell}, \quad T_\ell x = W^{(\ell)} x + b^{(\ell)},
\]
with $W^{(ℓ)} \in \mathbb{R}^{N_l \times N_{l-1}}$ being the weight matrices and $b^{(ℓ)} \in \mathbb{R}^{N_l}$ the bias vectors of the $ℓ$th layer. Then $\Phi : \mathbb{R}^d \to \mathbb{R}^{N_L}$, given by

$$\Phi(x) = T_L \rho(T_{L-1} \rho(\ldots \rho(T_1(x))))$$

is called (deep) neural network of depth $L$.

Let us already mention at this point that the weights and biases are the free parameters which will be learned during the training process. An illustration of the multilayered structure of a deep neural network can be found in Figure 1.

![Deep neural network](image)

**Figure 1:** Deep neural network $\Phi : \mathbb{R}^d \to \mathbb{R}$ with depth 5.

## 2.2 Application of a Deep Neural Network

Aiming to identify the main mathematical research threads, we first have to understand how a deep neural network is used for a given application setting.

**Step 1 (Train-test split of the dataset):** We assume that we are given samples $(x^{(i)}, y^{(i)})_{i=1}^m$ of inputs and outputs. The task of the deep neural network is then to identify the relation between those. For instance, in a classification problem, each output $y^{(i)}$ is considered to be the label of the respective class to which the input $x^{(i)}$ belongs. One can also take the viewpoint that $(x^{(i)}, y^{(i)})_{i=1}^m$ arises as samples from a function such as $g : \mathcal{M} \to \{1, 2, \ldots, K\}$, where $\mathcal{M}$ might be a lower-dimensional manifold of $\mathbb{R}^d$, in the sense of $y^{(i)} = g(x^{(i)})$ for all $i = 1, \ldots, m$.

The set $(x^{(i)}, y^{(i)})_{i=1}^m$ is then split into a training data set $(x^{(i)}, y^{(i)})_{i=1}^m$ and a test data set $(x^{(i)}, y^{(i)})_{i=m+1}^\tilde{m}$. The training data set is—as the name indicates—used for training, whereas the test data set will later on solely be exploited for testing the performance of the trained network. We emphasize that the neural network is not exposed to the test data set during the entire training process.

**Step 2 (Choice of architecture):** For preparation of the learning algorithm, the architecture of the neural network needs to be decided upon, which means the number of layers $L$, the number of neurons in each layer $(N_\ell)_{\ell=1}^L$, and the activation function $\rho$ have to be selected. It is known that a fully connected neural network is often difficult to train, hence, in addition, one typically preselects certain entries of the weight matrices $(W^{(ℓ)})_{\ell=1}^L$ to already be set to zero at this point.

For later purposes, we define the selected class of deep neural networks by $\mathcal{N}_\theta$ with $\theta$ encoding this chosen architecture.

**Step 3 (Training):** The next step is the actual training process, which consists of learning the affine-linear functions $(T_\ell)_{\ell=1}^L = (W^{(\ell)} + b^{(\ell)})_{\ell=1}^L$. This is accomplished by minimizing the empirical risk

$$\hat{\mathcal{R}}(\Phi_{(W^{(\ell)}, b^{(\ell)})}) := \frac{1}{m} \sum_{i=1}^m \left(\Phi_{(W^{(\ell)}, b^{(\ell)})}(x^{(i)}) - y^{(i)}\right)^2. \quad (2.1)$$

A more general form of the optimization problem is

$$\min_{(W^{(\ell)}, b^{(\ell)})_{\ell=1}^L} \sum_{i=1}^m \mathcal{L}(\Phi_{(W^{(\ell)}, b^{(\ell)})}(x_i), y^{(i)}) + \lambda \mathcal{P}((W^{(\ell)}, b^{(\ell)})_{\ell=1}^L), \quad (2.2)$$

where $\mathcal{L}$ is the loss function and $\mathcal{P}$ is a penalty function.
where $\mathcal{L}$ is a loss function to determine a measure of closeness between the network evaluated in the training samples and the (known) values $y^{(i)}$ and where $\mathcal{P}$ is a penalty/regularization term to impose additional constraints on the weight matrices and bias vectors.

One common algorithmic approach is gradient descent. Since, however, $m$ is typically very large, this is computationally not feasible. This problem is circumvented by randomly selecting only a few gradients in each iteration, assuming that they constitute a reasonable average, which is coined stochastic gradient descent.

Solving the optimization problem then yields a network $\Phi_{(W^{(i)},b^{(i)})} : \mathbb{R}^d \rightarrow \mathbb{R}^{N_L}$, where

$$\Phi_{(W^{(i)},b^{(i)})}(x) = T_L \rho(T_{L-1} \rho(\ldots \rho(T_1(x)))).$$ 

Step 4 (Testing): Finally, the performance (often also called generalization ability) of the trained neural network is tested using the test data set $(x^{(i)}, y^{(i)})_{i=m+1}^\tilde{m}$ by analyzing whether

$$\Phi_{(W^{(i)},b^{(i)})}(x^{(i)}) \approx y^{(i)}, \quad \text{for all } i = m + 1, \ldots, \tilde{m}.$$ 

## 2.3 Relation to a Statistical Learning Problem

From the procedure above, we can already identify the selection of architecture, the optimization problem, and the generalization ability as the key research directions for mathematical foundations of deep neural networks. Considering the entire learning process of a deep neural network as a statistical learning problem reveals those three research directions as indeed the natural ones for analyzing the overall error.

For this, let us assume that there exists a function $g : \mathbb{R}^d \rightarrow \mathbb{R}$ such that the training data $(x^{(i)}, y^{(i)})_{i=1}^m$ is of the form $(x^{(i)}, g(x^{(i)}))_{i=1}^m$ and $x^{(i)} \in [0,1]^d$ for all $i = 1, \ldots, m$. A typical continuum viewpoint to measure success of the training is to consider the risk of a function $f : \mathbb{R}^d \rightarrow \mathbb{R}$ given by

$$\mathcal{R}(f) := \int_{[0,1]^d} (f(x) - g(x))^2 dx, \quad (2.3)$$

where we used the $L^2$-norm to measure the distance between $f$ and $g$. The error between the trained deep neural network $\Phi^0(=\Phi_{(W^{(i)},b^{(i)})}) \in \mathcal{NN}_\theta$ and the optimal function $g$ can then be estimated by

$$\mathcal{R}(\Phi^0) \leq \underbrace{\mathcal{R}(\Phi^0) - \inf_{\Phi \in \mathcal{NN}_\theta} \mathcal{R}(\Phi)}_{\text{Optimization error}} + 2 \sup_{\Phi \in \mathcal{NN}_\theta} |\mathcal{R}(\Phi) - \mathcal{R}(\Phi)| + \inf_{\Phi \in \mathcal{NN}_\theta} \mathcal{R}(\Phi), \quad (2.4)$$

These considerations lead to the main research threads described in the following subsection.

## 2.4 Main Research Threads

We can identify two conceptually different research threads, the first one being focused on developing mathematical foundations of artificial intelligence and the second one aiming to use methodologies from artificial intelligence to solve mathematical problems. It is intriguing to see how both have already led to some extent to a paradigm shift in some mathematical research areas, most prominently the area of numerical analysis.

### 2.4.1 Mathematical Foundations for Artificial Intelligence

Following up on the discussion in Subsection 2.3, we can identify three research directions which are related to the three types of errors which one needs to control in order to estimate the overall error of the entire training process.

- **Expressivity.** This direction aims to derive a general understanding whether and to which extent aspects of a neural network architecture affect the best case performance of deep neural networks. More precisely,
the goal is to analyze the approximation error \( \inf_{\Phi \in \mathcal{N}_\theta} \mathcal{R}(\Phi) \) from (2.4), which estimates the approximation accuracy when approximating \( g \) by the hypothesis class \( \mathcal{N}_\theta \) of deep neural networks of a particular architecture. Typical methods for approaching this problem are from applied harmonic analysis and approximation theory.

- **Learning/Optimization.** The main goal of this direction is the analysis of the training algorithm such as stochastic gradient descent, in particular, asking why it usually converges to suitable local minima even though the problem itself is highly non-convex. This requires the analysis of the optimization error, which is \( \mathcal{R}(\Phi^0) - \inf_{\Phi \in \mathcal{N}_\theta} \mathcal{R}(\Phi) \) (cf. (2.4)) and which measures the accuracy with which the learnt neural network \( \Phi^0 \) minimizes the empirical risk \( (2.1), (2.2) \). Key methodologies for attacking such problems come from the areas of algebraic/differential geometry, optimal control, and optimization.

- **Generalization.** This direction aims to derive an understanding of the out-of-sample error, namely, \( \sup_{\Phi \in \mathcal{N}_\theta} |\mathcal{R}(\Phi) - \mathcal{R}(\Phi)| \) from (2.4), which measures the distance of the empirical risk \( (2.1), (2.2) \) and the actual risk \( (2.3) \). Predominantly, learning theory, probability theory, and statistics provide the required methods for this research thread.

A very exciting and highly relevant new research direction has recently emerged, coined explainability. At present, it is from the standpoint of mathematical foundations still a wide open field.

- **Explainability.** This direction considers deep neural networks, which are already trained, but no knowledge about the training is available; a situation one encounters numerous times in practice. The goal is then to derive a deep understanding of how a given trained deep neural network reaches decisions in the sense of which features of the input data are crucial for a decision. The range of required approaches is quite broad, including areas such as information theory or uncertainty quantification.

### 2.4.2 Artificial Intelligence for Mathematical Problems

Methods of artificial intelligence have also turned out to be extremely effective for mathematical problem settings. In fact, the area of inverse problems, in particular, in imaging sciences, has already undergone a profound paradigm shift. And the area of numerical analysis of partial differential equations seems to soon follow the same path, at least in the very high dimensional regime.

Let us briefly characterize those two research threads similar to the previous subsection on mathematical foundations of artificial intelligence.

- **Inverse Problems.** Research in this direction aims to improve classical model-based approaches to solve inverse problems by exploiting methods of artificial intelligence. In order to not neglect domain knowledge such as the physics of the problem, current approaches aim to take the best out of both worlds in the sense of optimally combining model- and data-driven approaches. This research direction requires a variety of techniques, foremost from areas such as imaging science, inverse problems, and microlocal analysis, to name a few.

- **Partial Differential Equations.** Similar to the area of inverse problems, here the goal is to improve classical solvers of partial differential equations by using ideas from artificial intelligence. A particular focus is on high dimensional problems in the sense of aiming to beat the curse of dimensionality. This direction obviously requires methods from areas such as numerical mathematics and partial differential equations.

### 3 Mathematical Foundations for Artificial Intelligence

This section shall serve as an introduction into the main research threads aiming to develop a mathematical foundation for artificial intelligence. We will introduce the problem settings, showcase some exemplary results, and discuss open problems.
3.1 Expressivity

Expressivity is maybe the richest area at present in terms of mathematical results. The general question can be phrased as follows: Given a function class/space $C$ and a class of deep neural networks $\mathcal{NN}_\theta$, how does the approximation accuracy when approximating elements of $C$ by networks $\Phi \in \mathcal{NN}_\theta$ relate to the complexity of such $\Phi$? Making this precise thus requires the introduction of a complexity measure for deep neural networks. In the sequel, we will choose the canonical one, which is the complexity in terms of memory requirements. Notice though that certainly various other complexity measures exist. Further, recall that the $\| \cdot \|_0$-"norm" counts the number of non-zero components.

Definition 3.1. Retaining the same notation for deep neural networks as in Definition 2.2, the complexity $C(\Phi)$ of a deep neural network $\Phi$ is defined by

$$C(\Phi) := \sum_{\ell=1}^{L} \left( \| W^{(\ell)} \|_0 + \| b^{(\ell)} \|_0 \right).$$

The most well-known—and maybe even the first—result on expressivity is the universal approximation theorem [8; 13]. It states that each continuous function on a compact domain can be approximated up to an arbitrary accuracy by a shallow neural network.

Theorem 3.2. Let $d \in \mathbb{N}$, $K \subset \mathbb{R}^d$ compact, $f : K \to \mathbb{R}$ continuous, $\rho : \mathbb{R} \to \mathbb{R}$ continuous and not a polynomial. Then, for each $\epsilon > 0$, there exist $N \in \mathbb{N}$ and $a_k, b_k \in \mathbb{R}, w_k \in \mathbb{R}^d, 1 \leq k \leq N$, such that

$$\| f - \sum_{k=1}^{N} a_k \rho(\langle w_k, \cdot \rangle - b_k) \|_\infty \leq \epsilon.$$

While this is certainly an interesting result, it is not satisfactory in several regards: It does not give bounds on the complexity of the approximating neural network and also does not explain why depth is so important. A particularly intriguing example for a result, which considers complexity and also targets a more sophisticated function space, was derived in [31].

Theorem 3.3. For all $f \in C^s([0, 1]^d)$ and $\rho(x) = \max\{0, x\}$, i.e., the ReLU, there exist neural networks $(\Phi_n)_{n \in \mathbb{N}}$ with the number of layers of $\Phi_n$ being approximately of the order of $\log(n)$ such that

$$\| f - \Phi_n \|_\infty \lesssim C(\Phi_n)^{-\gamma} \to 0 \quad \text{as} \quad n \to \infty.$$

This result provides a beautiful connection between approximation accuracy and complexity of the approximating neural network, and also to some extent takes the depth of the network into account. However, to derive a result on optimal approximations, we first require a lower bound. The so-called VC-dimension (Vapnik-Chervonenkis-dimension) (see also (3.2)) was for a long time the main method for achieving such lower bounds. We will recall here a newer result from [7] in terms of the optimal exponent $\gamma^*(\mathcal{C})$ from information theory to measure the complexity of $\mathcal{C} \subset L^2(\mathbb{R}^d)$. Notice that we will only state the essence of this result without all technicalities.

Theorem 3.4. Let $d \in \mathbb{N}$, $\rho : \mathbb{R} \to \mathbb{R}$, and let $\mathcal{C} \subset L^2(\mathbb{R}^d)$. Further, let

$$\text{Learn} : (0, 1) \times \mathcal{C} \to \mathcal{NN}_\theta$$

satisfy that, for each $f \in \mathcal{C}$ and $0 < \epsilon < 1$,

$$\sup_{f \in \mathcal{C}} \| f - \text{Learn}(\epsilon, f) \|_2 \leq \epsilon.$$

Then, for all $\gamma < \gamma^*(\mathcal{C})$,

$$c^\gamma \sup_{f \in \mathcal{C}} C(\text{Learn}(\epsilon, f)) \to \infty, \quad \text{as} \quad \epsilon \to 0.$$
Let Theorem 3.8. from [7] in addition proves that the bound in Theorem 3.4 is sharp. 

\[ \text{N mimics best and in fact as all affine systems. Even more, the construction in the proof of suitable neural networks, which } \]

introduced in [9].

\[ \text{fronts in the solution of transport-dominated equations, the following suitable model class of functions was } \]

Definition 3.6.

\[ \text{SH scaling, i.e., } \]

\[ A_{2^j} = \begin{pmatrix} 2^j & 0 \\ 0 & 2^{j/2} \end{pmatrix}, \quad j \in \mathbb{Z} \]

and \[ \tilde{A}_{2^j} = \text{diag}(2^{j/2}, 2^j) \] as well as changing the orientation via shearing defined by \[ S_k = \begin{pmatrix} 1 & k \\ 0 & 1 \end{pmatrix}, \quad k \in \mathbb{Z}. \]

(Cone-adapted) discrete shearlet systems can then be defined as follows, cf. [17]. A faithful implementation of the shearlet transform as a 2D and 3D (parallelized) fast shearlet transform can be found at www.ShearLab.org.

Definition 3.5. The (cone-adapted) discrete shearlet system \( \mathcal{SH}(\phi, \psi, \hat{\psi}) \) generated by \( \phi \in L^2(\mathbb{R}^2) \) and \( \psi, \hat{\psi} \in L^2(\mathbb{R}^2) \) is the union of

\[ \{ \phi(\cdot - m) : m \in \mathbb{Z}^2 \}, \]

\[ \{ 2^{3j/4} \psi(S_k A_{2^j} \cdot - m) : j \geq 0, |k| \leq |2^{j/2}|, m \in \mathbb{Z}^2 \}, \]

\[ \{ 2^{3j/4} \hat{\psi}(S_k^T \tilde{A}_{2^j} \cdot - m) : j \geq 0, |k| \leq |2^{j/2}|, m \in \mathbb{Z}^2 \}. \]

Since multivariate problems are typically governed by anisotropic features such as edges in images or shock fronts in the solution of transport-dominated equations, the following suitable model class of functions was introduced in [9].

Definition 3.6. The set of cartoon-like functions \( \mathcal{E}^2(\mathbb{R}^2) \) is defined by

\[ \mathcal{E}^2(\mathbb{R}^2) = \{ f \in L^2(\mathbb{R}^2) : f = f_0 + f_1 \cdot \chi_B \}, \]

where \( \emptyset \neq B \subset [0,1]^2 \) is simply connected with a \( C^2 \)-curve with bounded curvature as its boundary, and \( f_i \in C^2(\mathbb{R}^2) \) with \( \text{supp } f_i \subseteq [0,1]^2 \) and \( \| f_i \|_{C^2} \leq 1, \quad i = 0,1 \).

While wavelets are deficient in optimally approximating cartoon-like functions due to their isotropic structure, shearlets provide an optimal (sparse) approximation rate up to a log-factor. The following statement is taken from [17], where also the precise hypotheses can be found. Notice that the justification for optimality is a benchmark result from [9].

Theorem 3.7. Let \( \phi, \psi, \hat{\psi} \in L^2(\mathbb{R}^2) \) be compactly supported, and let \( \hat{\psi}, \hat{\psi} \) satisfy certain decay conditions. Then \( \mathcal{SH}(\phi, \psi, \hat{\psi}) \) provides an optimally sparse approximation of \( f \in \mathcal{E}^2(\mathbb{R}^2) \), i.e.,

\[ \| f - f_N \|_2 \lesssim N^{-1}(\log N)^{3/2} \quad \text{as } N \to \infty. \]

One can now use Theorem 3.4 to show that indeed deep neural networks are as good approximators as shearlets and in fact as all affine systems. Even more, the construction in the proof of suitable neural networks, which mimics best \( N \)-term approximations, also leads to memory-optimal neural networks. The resulting statement from [7] in addition proves that the bound in Theorem 3.4 is sharp.

Theorem 3.8. Let \( \rho \) be a suitably chosen activation function, and let \( \epsilon > 0 \). Then, for all \( f \in \mathcal{E}^2(\mathbb{R}^2) \) and \( N \in \mathbb{N} \), there exists a neural network \( \Phi \) with complexity \( O(N) \) and activation function \( \rho \) with

\[ \| f - \Phi \|_2 \lesssim N^{-1+\epsilon} \to 0 \quad \text{as } N \to \infty. \]
Summarizing, one can conclude that deep neural networks achieve optimal approximation properties of all affine systems combined.

Let us finally mention that lately a very different viewpoint of expressivity was introduced in [21] according to so-called trajectory lengths. The standpoint taken in this work is to measure expressivity in terms of changes of the expected length of a (non-constant) curve in the input space as it propagates through layers of a neural network.

3.2 Optimization

This area aims to analyze optimization algorithms, which solve the (learning) problem in (2.1), or, more generally, (2.2). A common approach is gradient descent, since the gradient of the loss function (or optimized functional) with respect to the weight matrices and biases, i.e., the parameters of the network, can be computed exactly. This is done via backpropagation [27], which is in a certain sense merely an efficient application of the chain rule. However, since the number of training samples is typically in the millions, it is computationally infeasible to compute the gradient on each sample. Therefore, in each iteration only one or several (a batch) randomly selected gradients are computed, leading to the algorithm of stochastic gradient descent [25].

In convex settings, guarantees for convergence of stochastic gradient descent do exist. However, in the neural network setting, the optimization problem is non-convex, which makes it—even when using a non-random version of gradient descent—very hard to analyze. Including randomness adds another level of difficulty as is depicted in Figure 2, where the two algorithms reach different (local) minima.

![Gradient descent versus stochastic gradient descent](image)

Figure 2: Gradient descent versus stochastic gradient descent [6]

This area is by far less explored than expressivity. Most current results focus on shallow neural networks, and for a survey, we refer to [6].

3.3 Generalization

This research direction is perhaps the least explored and maybe also the most difficult one, sometimes called the “holy grail” of understanding deep neural networks. It targets the out-of-sample error

\[
\sup_{\Phi \in \mathcal{NN}_0} |\mathcal{R}(\Phi) - \hat{\mathcal{R}}(\Phi)|
\]  

(3.1)

as described in Subsection 2.4.1.

One of the mysteries of deep neural networks is the observation that highly overparameterized deep neural networks in the sense of high complexity of the network do not overfit with overfitting referring to the problem of fitting the training data too tightly and consequently endangering correct classification of new data. An illustration of the phenomenon of overfitting can be found in Figure 3.
Let us now analyze the generalization error in (3.1) in a bit more depth. For a large number $m$ of training samples the law of large numbers tells us that with high probability $\hat{R}(\Phi) \approx R(\Phi)$ for each neural network $\Phi \in \mathcal{NN}_\theta$. Bounding the complexity of the hypothesis class $\mathcal{NN}_\theta$ by the VC-dimension, the generalization error can be bounded with probability $1 - \delta$ by

$$\sqrt{\frac{\text{VCdim}(\mathcal{NN}_\theta) + \log(1/\delta)}{m}}.$$  \hfill (3.2)

For classes of highly over-parametrized neural networks, i.e., where $\text{VCdim}(\mathcal{NN}_\theta)$ is very large, we need an enormous amount of training data to keep the generalization error under control. It is thus more than surprising that numerical experiments show the phenomenon of a so-called double descent curve [5]. More precisely, the test error was found to decrease after passing the interpolation point, followed by an increase consistent with statistical learning theory (see Figure 4).

3.4 Explainability

The area of explainability aims to “open the black box” of deep neural networks in the sense as to explain decisions of trained neural networks. These explanations typically consist of providing relevance scores for features of the input data. Most approaches focus on the task of image classification and provide relevance scores for each pixel of the input image. One can roughly categorize the different types of approaches into gradient-based methods [28], propagation of activations in neurons [4], surrogate models [24], and game-theoretic approaches [19].

We would now like to describe in more detail an approach which is based on information theory and also allows an extension to different modalities such as audio data as well as analyzing the relevance of higher-level features; for a survey paper, we refer to [15]. This rate-distortion explanation (RDE) framework was introduced in 2019 and later extended by applying RDE to non-canonical input representations.

Let now $\Phi : \mathbb{R}^d \to \mathbb{R}^n$ be a trained neural network, and $x \in \mathbb{R}^d$. The goal of RDE is to provide an explanation for the decision $\Phi(x)$ in terms of a sparse mask $s \in \{0, 1\}^d$ which highlights the crucial input features of $x$. This
mask is determined by the following optimization problem:

$$\min_{s \in \{0, 1\}^d} \mathbb{E}_{v \sim \mathcal{V}} d(\Phi(x), \Phi((x \odot s + (1 - s) \odot v))) \text{ subject to } \|s\|_0 \leq \ell,$$

where $\odot$ denotes the Hadamard product, $d$ is a measure of distortion such as the $\ell_2$-distance, $\mathcal{V}$ is a distribution over input perturbations $v \in \mathbb{R}^d$, and $\ell \in \{1,\ldots,d\}$ is a given sparsity level for the explanation mask $s$. The key idea is that a solution $s^*$ is a mask marking few components of the input $x$ which are sufficient to approximately retain the decision $\Phi(x)$. This viewpoint reveals the relation to rate-distortion theory, which normally focuses on lossy compression of data.

Since it is computationally infeasible to compute such a minimizer (see [30]), a relaxed optimization problem providing continuous masks $s \in [0, 1]^d$ is used in practice:

$$\min_{s \in [0, 1]^d} \mathbb{E}_{v \sim \mathcal{V}} d(\Phi(x), \Phi((x \odot s + (1 - s) \odot v))) + \lambda \|s\|_1,$$

where $\lambda > 0$ determines the sparsity level of the mask. The minimizer now assigns each component $x_i$ of the input—in case of images each pixel—a relevance score $s_i \in [0, 1]$. This is typically referred to as Pixel RDE.

Extensions of the RDE-framework allow the incorporation of different distributions $\mathcal{V}$ better adapted to data distributions. Another recent improvement was the assignment of relevance scores to higher-level features such as arising from a wavelet decomposition, which ultimately led to the approach CartoonX. An example of Pixel RDE versus CartoonX, which also shows the ability of the higher-level explanations of CartoonX to give insights into what the neural network saw when misclassifying an image, is depicted in Figure 5.

![Pixel RDE versus CartoonX](image)

**Figure 5:** Pixel RDE versus CartoonX for analyzing misclassifications of a deep neural network

## 4 Artificial Intelligence for Mathematical Problems

We now turn to the research direction of artificial intelligence for mathematical problems, with the two most prominent problems being inverse problems and partial differential equations. As before, we will introduce the problem settings, showcase some exemplary results, and also discuss open problems.

### 4.1 Inverse Problems

Methods of artificial intelligence, in particular, deep neural networks have a tremendous impact on the area of inverse problems, as already indicated before. One current major trend is to optimally combine classical solvers
with deep learning in the sense of taking the best out of the model- and data-world.

To introduce such results, we start by recalling some basics about solvers of inverse problems. For this, assume that we are given an (ill-posed) inverse problem

$$Kf = g,$$

(4.1)

where \(K: X \rightarrow Y\) is an operator and \(X\) and \(Y\) are, for instance, Hilbert spaces. Drawing from the area of imaging science, examples include denoising, deblurring, or inpainting (recovery of missing parts of an image). Most classical solvers are of the form (which includes Tikhonov regularization)

$$f^\alpha := \arg\min_f \left[ \|Kf - g\|^2 + \alpha \cdot \mathcal{P}(f) \right],$$

where \(\mathcal{P}: X \rightarrow \mathbb{R}\) and \(f^\alpha \in X, \alpha > 0\) is an approximate solution of the inverse problem (4.1). One very popular and widely applicable special case is sparse regularization, where \(\mathcal{P}\) is chosen by

$$\mathcal{P}(f) := \|\langle f, \varphi_i \rangle\|_1$$

and \((\varphi_i)_{i \in I}\) is a suitably selected orthonormal basis or a frame for \(X\).

We now turn to deep learning approaches to solve inverse problems, which might be categorized into three classes:

- **Supervised approaches.** An ad-hoc approach in this regime is given in [14], which first applies a classical solver followed by a neural network to remove reconstruction artifacts. More sophisticated approaches typically replace parts of the classical solver by a custom-build neural network [26] or a network specifically trained for this task [1].

- **Semi-supervised approaches.** These approaches encode the regularization as a neural network with an example being adversarial regularizers [20].

- **Unsupervised approaches.** A representative of this type of approaches is the technique of deep image prior [29]. This method interestingly shows that the structure of a generator network is sufficient to capture necessary statistics of the data prior to any type of learning.

Aiming to illustrate the superiority of approaches from artificial intelligence for inverse problems, we will now focus on the inverse problem of computed tomography (CT) from medical imaging. The forward operator \(K\) in this setting is the Radon transform, defined by

$$\mathcal{R}f(\phi, s) = \int_{L(\phi, s)} f(x)dS(x),$$

where \(L(\phi, s) = \{x \in \mathbb{R}^2: x_1 \cos(\phi) + x_2 \sin(\phi) = s\}, \phi \in [-\pi/2, \pi/2),\) and \(s \in \mathbb{R}\). Often, only parts of the so-called sinogram \(\mathcal{R}f\) can be acquired due to physical constraints as in, for instance, electron tomography. The resulting, more difficult problem is termed limited-angle CT. One should notice that this problem is even harder than the problem of low-dose CT, where not an entire block of measurements is missing, but the angular component is “only” undersampled.

The most prominent features in images \(f\) are edge structures. This is also due to the fact that the human visual system reacts most strongly to those. These structures in turn can be accurately modeled by microlocal analysis, in particular, by the notion of wavefront sets \(WF(f) \subseteq \mathbb{R}^2 \times \mathbb{S}\), which—coarsely speaking—consist of singularities together with their direction. Basing in this sense the application of a deep neural network on microlocal considerations, in particular, also using a deep learning-based wavefront set detector [2] in the regularization term, the reconstruction performance significantly outperforms classical solvers such as sparse regularization with shearlets (see Figure 6 we also refer to [3] for details). Notice that this approach is of a
hybrid type and takes the best out of both worlds in the sense of combining model- and artificial intelligence-based approaches.

Finally, the deep learning-based wavefront set extraction itself is yet another evidence of the improvements on the state-of-the-art now possible by artificial intelligence. Figure 7 shows a classical result from [23], whereas [2] uses the shearlet transform as a coarse edge detector, which is subsequently combined with a deep neural network.

4.2 Partial Differential Equations

The second main range of mathematical problem settings, where methods from artificial intelligence are very successfully applied to, are partial differential equations. Although the benefit of such approaches was not initially clear, both theoretical and numerical results show their superiority in high-dimensional regimes.

The most common approach aims to approximate the solution of a partial differential equation by a deep neural network, which is trained according to this task by incorporating the partial differential equation into the loss function. More precisely, given a partial differential equation \( L(u) = f \), we train a neural network \( \Phi \) such that

\[
L(\Phi) \approx f.
\]

Since 2017, research in this general direction has significantly accelerated. Some of the highlights are the Deep Ritz Method [10] and Physics Informed Neural Networks [22], or a very general approach for high-dimensional parabolic partial differential equations [12].

One should note that most theoretical results in this regime are of an expressivity type and also study the phenomenon whether and to which extent deep neural networks are able to beat the curse of dimensionality. In the sequel, we briefly discuss one such result as an example. In addition, notice that there already exist contributions—though very few—which analyze learning and generalization aspects.
Let $L(u_y, y) = f_y$ denote a parametric partial differential equation with $y$ being a parameter from a high-dimensional parameter space $\mathcal{Y} \subseteq \mathbb{R}^p$ and $u_y$ the associated solution in a Hilbert space $\mathcal{H}$. After a high-fidelity discretization, let $b_y(u^h_y, v) = f_y(v)$ be the associated variational form with $u^h_y$ now belonging to the associated high-dimensional space $U^h$, where we set $D := \dim(U^h)$. We moreover denote the coefficient vector of $u^h_y$ with respect to a suitable basis of $U^h$ by $u^h_y$. Of key importance in this area is the parametric map given by

$$\mathbb{R}^p \supseteq \mathcal{Y} \ni y \mapsto u^h_y \in \mathbb{R}^D$$

such that $b_y(u^h_y, v) = f_y(v)$ for all $v$, which in multi-query situations such as complex design problems needs to be solved several times. If $p$ is very large, the curse of dimensionality could lead to an exponential computational cost.

We now aim to analyze whether the parametric map can be solved by a deep neural network, which would provide a very efficient and flexible method, hopefully also circumventing the curse of dimensionality in an automatic manner. From an expressivity viewpoint, one might ask whether, for each $\epsilon > 0$, there does exist a neural network $\Phi$ such that

$$\|\Phi(y) - u^h_y\| \leq \epsilon \quad \text{for all } y \in \mathcal{Y}.$$  (4.2)

The ability of this approach to tackle the curse of dimensionality can then be studied by analyzing how the complexity of $\Phi$ depends on $p$ and $D$. A result of this type was proven in [18], the essence of which we now recall.

**Theorem 4.1.** There exists a neural network $\Phi$ which approximates the parametric map, i.e., which satisfies (4.2), and its dependence on $C(\Phi)$ on $p$ and $D$ can be (polynomially) controlled.

Analyzing the learning procedure and the generalization ability of the neural network in this setting is currently out of reach. Aiming to still determine whether a trained neural networks does not suffer from the curse of dimensionality as well, in [11] extensive numerical experiments were performed, which indicates that indeed the curse of dimensionality is also beaten in practice.

## 5 Conclusion: Seven Mathematical Key Problems

Let us conclude with seven mathematical key problems of artificial intelligence as they were stated in [6]. Those constitute the main obstacles in *Mathematical Foundations for Artificial Intelligence* with its subfields expressivity, optimization, generalization, and explainability as well as in *Artificial Intelligence for Mathematical Problems* which focusses on the application to inverse problems and partial differential equations.

1. What is the role of depth?
2. Which aspects of a neural network architecture affect the performance of deep learning?
3. Why does stochastic gradient descent converge to good local minima despite the non-convexity of the problem?
4. Why do large neural networks not overfit?
5. Why do neural networks perform well in very high-dimensional environments?
6. Which features of data are learned by deep architectures?
7. Are neural networks capable of replacing highly specialized numerical algorithms in natural sciences?

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