FYS-STK 3155 Project 3 Erlend Abrahamsen

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Abstract

Partial differential equations are very important and have wide applications, one example is the use of Navier-Stokes equations for weather predictions. Much study and improvement of solvers for these are crucial. In this project we study deep neural networks (DNN) for solving a version of the heat equation as well as a rather ambitious ODE approach for finding eigenvalues. For comparison of the heat equation results, we also solve it analytically, and with the numerical finite differences method. Results showed that the DNN was efficient and had decent accuracy while finite differences had great accuracy but with poor computational efficiency. For eigenvalues the predicted eigenvalues had chaotic behavior and oscillated between many of the true eigenvalues. The model had sometimes good results but lacked stability.

1 Introduction

For the source code (folder Source_code) and the results (folder Results) go to my Github: https://github.com/ErlendAbrahamsen/FYS-STK3155/tree/master/Project3

PDE's are widely used for explaining real physical problems and are generally approximated using numerical methods. An example of great interest is modeling of fluid dynamics. More computational efficiency is much needed in this area, and with a lot recent success of Neural networks, it is interesting to investigate a deep learning approach on PDE's. For more motivation see the recent paper Deep Neural Networks Motivated by Partial Differential Equations.

In this project we explore the finite differences scheme and a feed forward Neural network based on Tensorflow functionality for solving the 1d heat equation given some nice initial condition.

In the article Computers and Mathematics with Applications (Reference 2. Authors Yi, Fu and Jin Tang) they have an interesting approach of predicting eigenvectors/eigenvalues by solving a vector ODE with a recurrent neural network. Eigenvalues are essential in linear algebra, and we will also attempt their method with a feed forward network.

Outline [Introduction, Methods, Results and discussion, Summary] (I will reference to source code/project part where relevant!)

1.1 The 1d heat equation (Part a)

The general heat equation is $\frac{\partial u}{\partial t} = \alpha \left[\frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} + \frac{\partial^2 u}{\partial z^2} \right]$, where u(x,y,z,t) is the temperature of a medium at position (x,y,z) in time t. The α constant is thermal diffusivity of the medium. We look at a 1d case with $\alpha = 1$.

For more explanation of units and physical interpretation see The Heat Equation (Ref. 5).

Given a 1d rod of length L = 1, from position x = 0 til x = 1, we want to find the temperature u(x,t) at any position in time (x,t).

We set the initial heat distribution for time t = 0 as $u(x, 0) = sin(\pi x)$ and constant zero temperatures at endpoints u(0, t) = u(L, t) = 0.

Heat equation with chosen conditions:

$$u_t(x,t) = u_{xx}(x,t), t > 0, x \in [0,1]$$

or
$$\frac{\partial u(x,t)}{\partial t} = \frac{\partial^2 u(x,t)}{\partial x^2}$$

$$u(x,0) = sin(\pi x), 0 \le x \le 1$$
 (Initial conditions)

$$u(0,t)=u(1,t)=0,\,t\geq 0$$
 (Boundary conditions)

The solution is $u(x,t) = e^{-\pi^2 t} \sin(\pi x)$ which we show in next section.

2 Methods

2.1 Analytic solution of 1d heat pde (Part a)

For the solution we use the pde technique separation of variables. Also note that for most initial conditions infinite Fourier series are computed. This is not needed for our simplified version and we avoid extra errors for lack of terms used.

Sep. of var.: Assume u(x,t) = X(x)T(t) for some $X,T: R \to R$.

Plugging this into $u_t = u_{xx}$ and ordering terms gives $\frac{T'(t)}{T(t)} = \frac{X''(x)}{X(x)}$.

This has to be some constant $\lambda \in R$, since t and x are independent variables. Further we get two ODE's $T'(t) - \lambda T(t) = 0$ and $X''(x) - \lambda X(x) = 0$. These are easy to solve and have some different forms dependent on λ .

(Case 1. $\lambda = 0$)

Using boundary conditions yield

0 = u(0,t) = X(0)T(t) = u(1,t) = X(1)T(t) which implies X(x) = 0 since X(x) = a + bx. We are looking for non-zero solutions, thus $\lambda = 0$ is excluded.

(Case 2. $\lambda > 0$)

Again from boundary conditions we have X(0) = X(1) = 0 implying a = -b and $ae^{\sqrt{\lambda}x} - ae^{\sqrt{\lambda}x} = 0$, since $X(x) = ae^{\sqrt{\lambda}x} + be^{-\sqrt{\lambda}x}$, giving X(x) = 0. $\lambda > 0$ is also excluded.

(Case 3 $\lambda < 0$)

 $X(x) = a\cos(\sqrt{-\lambda}x) + b\sin(\sqrt{-\lambda}x)$ and $T(t) = ce^{\lambda t}$, giving a = X(0) = 0, and $X(1) = b\sin(\sqrt{-\lambda}x)$ yielding $\sqrt{-\lambda} = \pi n$, $n \in N_1$. This we can solve for a non-zero solution.

Putting it all together we have now found the particular solutions $u_n = b_n e^{\lambda t} \sin(\sqrt{-\lambda}x) = b_n e^{-(\pi n)^2 t} \sin(\pi n x)$, giving the general solution $u = \sum_{n=1}^{\infty} u_n$ by super position principle.

 $u(x,0) = \sin(\pi x) = \sum_{n=1}^{\infty} b_n \sin(\pi n x)$, so $b_1 = 1$ and $b_{n\neq 1} = 0$ giving our final general solution.

$$u(x,t) = e^{-\pi^2 t} \sin(\pi x)$$
 for $t \ge 0, x \in [0,1]$.

Verifying the solution:

$$u_t(x,t) = -\pi^2 e^{-\pi^2 t} \sin(\pi x)$$

$$u_{xx}(x,t) = \frac{d}{dx}\pi e^{-\pi^2 t}\cos(\pi x) = -\pi^2 e^{-\pi^2 t}\sin(\pi x) = u_t(x,t)$$

$$u(x,0) = \sin(\pi x), \ u(0,t) = u(1,t) = 0$$

2.2 Numerical approximation with finite differences (Part b)

For deriving derivative approximations of u_t and u_{xx} we make use of 1. and 2. order Taylor expansions. For notation simplicity let $f: R \to R$ be a sufficient continuous function.

1. order Taylor expansion gives $f(x + \Delta x) = f(x) + f'(x)\Delta x + O(\Delta x^2)$

So $f'(x) \approx \frac{f(x+\Delta x)-f(x)}{\Delta x}$ with truncation error $O(\Delta x)$. (Forwards difference)

2. order Taylor expansions of $f(x + \Delta x)$ and $f(x - \Delta x)$ yields

(i)
$$f(x + \Delta x) = f(x) + f'(x)\Delta x + \frac{1}{2}f''(x)\Delta x^2 + O(\Delta x^3)$$

(ii)
$$f(x - \Delta x) = f(x) - f'(x)\Delta x + \frac{1}{2}f''(x)\Delta x^2 - O(\Delta x^3)$$

Adding (i), (ii) and solving for f''(x) yields

 $f''(x) \approx \frac{f(x+\Delta x)-2f(x)+f(x-\Delta x)}{\Delta x^2}$ with truncation error $O(\Delta x^2)$. (Central difference)

We use this to set up the forward time, centered space, discrete approximation:

Using the derivative approximations on $\frac{\partial u(x,t)}{\partial t} = \frac{\partial^2 u(x,t)}{\partial x^2}$ we let $u(x_i,t_j) = u_i^j$:

$$\frac{u_{i}^{j+1}-u_{i}^{j}}{\Delta t}=\frac{u_{i+1}^{j}-2u_{i}^{j}+u_{i-1}^{j}}{\Delta x^{2}}$$

$$u_i^{j+1}=u_i^j+\frac{\Delta t}{\Delta x^2}(u_{i+1}^j-2u_i^j+u_{i-1}^j),$$
 let the ratio $\frac{\Delta t}{\Delta x^2}=d$

$$u_i^{j+1} = du_{i+1}^j + (1-2d)u_i^j + du_{i-1}^j, i = 1, ..., n-1, j = 0, 1, ..., m-1$$

With conditions:

$$u_i^0 = \sin(\pi x_i), i = 1, ..., n - 1$$

$$u_0^j = u_n^j = 0, j = 0, 1, ..., m$$

We choose $d = \Delta t/\Delta x^2 \leq 1/2$, as it ensures numerical stability and set up the grids $x_i = \Delta x \cdot i \in [0, 1]$, $t_j = \Delta t \cdot j \in [0, t_{stop}]$ and iterate explicitly with double loop.

The time complexity of our algorithm becomes

$$O(len(x) \cdot len(t)) = O(\frac{1}{\Delta x} \cdot \frac{t_{stop}}{d\Delta x^2}) = O(1/\Delta x^3).$$

(See finit_diff() in Part_b.py for Python implementation)

2.3 Neural Network learning PDE solution (Part c)

We will utilize the flexibility of Tensorflow (API Documentation Ref. 7) to explore DNN's with an appropriate defined cost function. By the universal approximation theorem we know that a simple neural network can approximate any sufficient continuous function.

We let the approximation of PDE solution u(x,t) be $\hat{u} = N(x,t)$, where $N = a^L$ is the DNN output of the final layer L.

The activation at each layer l=1,...,L is $a^l=f(a^{l-1}W^l+b^l)$ for some activation function $f\in R$ and input $a^0=[x'\quad t']\in R^{(n\times 2)}$.

Dimensions:

 W^l : $(2 \times m)$, $(m \times m)$, ..., $(m \times m)$, $(m \times 1)$ for l = 1, 2, ..., L - 1 (m is num. hidden neurons).

$$b^{l}$$
: $(n \times 2)$, $(n \times m)$, ..., $(n \times m)$, $(n \times 1)$ for $l = 1, 2, ..., L - 1$

$$a^l$$
: $(n \times 2)$, $(n \times m)$, ..., $(n \times m)$, $(n \times 1)$ for $l = 0, 1, ..., L - 1, L$ (n is length of x, t meshgrid)

Clearly we want $a^L \in [0,1]$ since $u(x,t) \in [0,1]$. A possibility for this is using $sigmoid(t) = \frac{1}{1+e^{-t}}$ as the output function, else one should think of good candidates and test them.

For optimizing the weights W^l and biases b^l we utilize Tensorflow's Adam optimizer, which uses variable learning rate and usually performs better than SGD (stochastic gradient descent).

For further explanations on optimization with SGD and back propagation see section 2.2 in project 2 (Ref. 6).

Also for additional information on Adam optimizer visit Adam - latest trends in deep learning optimization (Ref. 4).

(HeatLearner class in Part_c.py for DNN implementation)

2.3.1 Defining cost or loss for PDE problem

We need our loss/cost to catch properties and conditions:

$$u_t - u_{xx} = 0$$
 (i)

$$u(x,0) - \sin(\pi x) = 0$$
 (ii)

$$u(0,t) = 0$$
 (iii)

$$u(1,t) = 0$$
 (iv).

A good suggestion would be

$$L(x,t) =$$

$$MSE[\hat{u}_t, \hat{u}_{xx}] + MSE[\hat{u}(x, 0), sin(\pi x)] + MSE[\hat{u}(0, t), 0] + MSE[\hat{u}(1, t), 0]$$

This punishes each property (i-iv) equally for being non-zero and the DNN learns them when $L \to 0$.

We use this loss function and implement it as loss() in HeatLearner class.

2.4 Eigenvalue learning with Neural network (Part d)

From the paper on eigenvalue computations (Ref. 2) they proposed solving a vector differential equation with an artificial recurrent neural network for finding the eigenvectors/eigenvalues of a symmetric $n \times n$ matrix.

Let A be a $n \times n$ real symmetric matrix and $x(t) = (x_1(t), ..., x_n(t))^T \in \mathbb{R}^n$. The proposed ODE:

(1)
$$x'(t) = f(x(t)) - x(t)$$
, where $f(x) = [x^T x A + (1 - x^T A x)I]x$.

For a representation of the ODE solution see Theorem 2, section 3 in the paper (Ref. 2).

Theorem 3 in the paper states that each solution of (1) starting from any non-zero point $x(0) \in \mathbb{R}^n$ will converge to an eigenvector $v = \lim_{t \to \infty} x(t)$ of A (Convergence Analysis, section 4).

We find the corresponding eigenvalue $\lambda = \frac{v^T \lambda v}{v^T v} = \frac{v^T A v}{v^T v}$.

Let $\hat{x}(t) = N(x_0, t)$ for some initial guess $x_0 \in \mathbb{R}^n$ where N is the NN output.

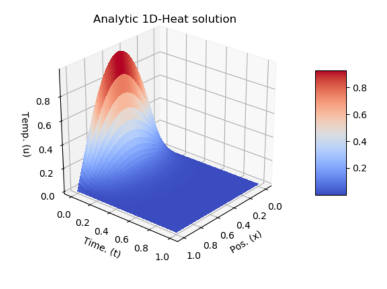
We set up the loss function $L(t) = mse[\hat{x}'(t), f(\hat{x}) - \hat{x}]$ since $\hat{x}'(t) \to f(\hat{x}) - \hat{x}$ when $L \to 0$.

In the implementation we reuse the PDE solver class HeatLearner. (See $Part_d$ for implementation)

3 Results and discussion

3.1 Finite differences (Part b)

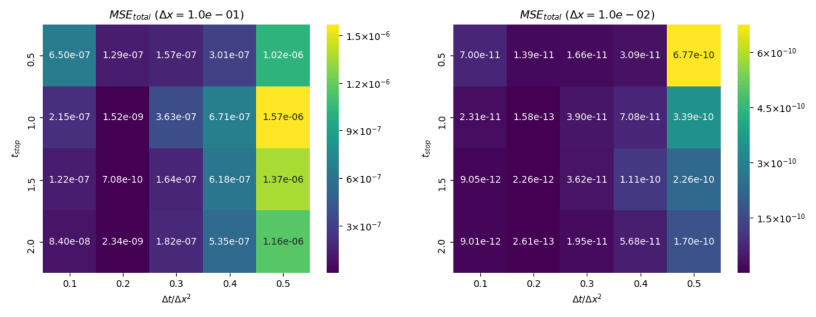
The analytic solution $u(x,t) = e^{-\pi^2 t} \sin(\pi x), x, t \in [0,1]$:



Notice the quick temperature drop with respect to time caused by the exponential term.

For using the finit-diff. scheme we need the ratio $\Delta t/\Delta x^2 \leq 1/2$, else unstable results with massive erros occur.

Tuning the model we create a heat-map, $MSE_{total} = MSE(u, \hat{u})$, for different time ranges $t \in [0, t_{stop}]$ and ratios $\Delta t/\Delta x^2$. We wont bother looking at t > 2 because of the exponential decay.

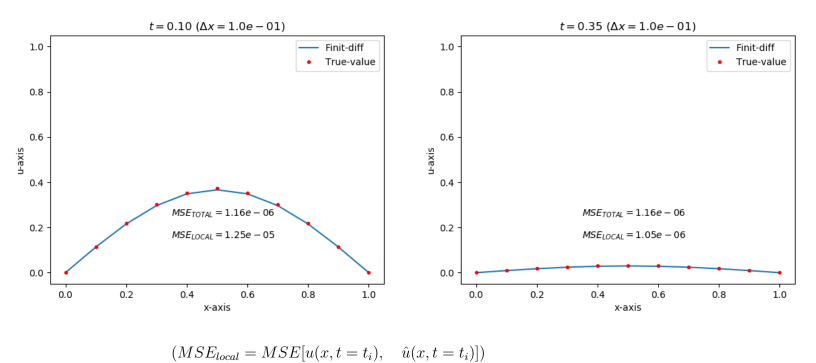


For fixed $(\Delta x)_1 = 10^{-1}$ and $(\Delta x)_2 = 10^{-2}$ we notice over all great results error-wise. The best ratio with respect to error is 0.2. Smaller ratios gives increased error.

Ratio at 0.5 is sufficient for both $(\Delta x)_1$ and $(\Delta x)_2$ at $t_{stop} = 2$, since MSE errors are respectively $1.16 \cdot 10^{-6}$ and $1.70 \cdot 10^{-10}$.

The model is not very flexible with respect to grid selection because of the stability criterion. Decreasing Δx is very computationally expensive and we might get a smaller Δt then necessary, which is a waste of time. The algorithm scales poorly with $O(1/\Delta x^3)$ time complexity, derived in section 2.2.

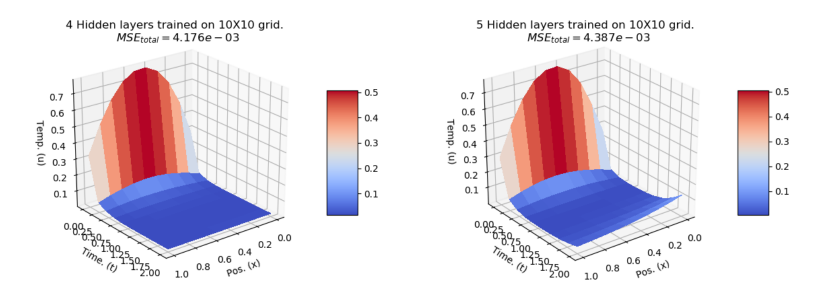
We look at the cheapest model $((\Delta x)_1 \text{ with } Ratio = 0.5)$ at two fixed times $t_1 = 0.1$ and $t_2 = 0.35$:



At $t_1=0.1$ when \hat{u} is significantly curved, the local 'sine term' error is greater than the total error. When $t>t_2=0.35$ \hat{u} is closer to the stationary state and $MSE_{total}>MSE_{local}\rightarrow 0$.

3.2 DNN on heat equation (Part c)

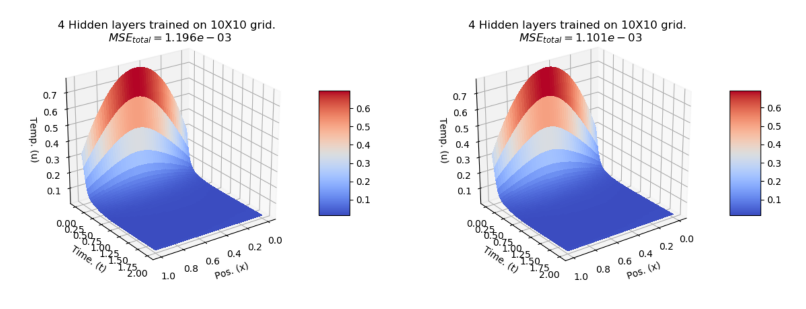
Trying Relu variations with some linear output function $f \in [0,1]$ doesn't seem to do well with sine curves. It learns the exponential decay but along the x-axis we get very pointy linear parts. We end up using Sigmoid as hidden activation function and output function. The DNN output is then in value range [0,1] which we want.



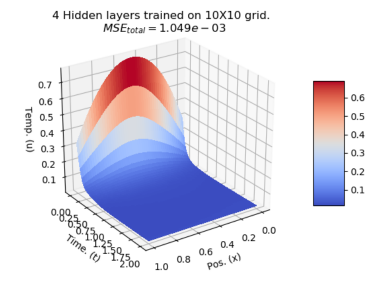
 $(10 \times 10 \text{ prediction})$

Adding more than 4 layers makes the model worse and at 5 hidden layers we see that \hat{u} starts to curve again near t=2 (Right figure). In the final model we end up with 4 hidden layers with 130 neurons each. The best we do are errors of magnitude 10^{-3} which is decent but not as good as 10^{-6} with finite-differences. The upside is that we can do fast training on a small grid and get relatively god prediction on much larger untrained grids, hence a big potential in computational efficiency gains. Training time for 10×10 grid was ≈ 1.3 seconds.

$(100 \times 100 \text{ grid prediction}) (200 \times 200 \text{ grid prediction})$

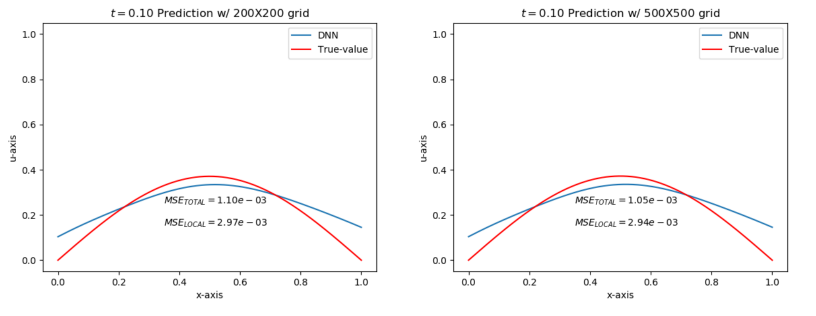


 $(500 \times 500 \text{ grid prediction})$



We see that the errors are very slightly decreasing for the grids $\{100^2, 200^2, 500^2\}$. The errors are $MSE_{total} = \{1.20 \cdot 10^{-3}, 1.10 \cdot 10^{-3}, 1.05 \cdot 10^{-3}\}$ respectively, with efficient prediction computations of course. The prediction method $u_hat()$ uses one loop over the number of layers and computes a matrix product, all the matrices are stored in memory.

Let's also look at the local sine errors for fixed times:



At t = 0.1 the cheapest finit-diff. model had MSE_{local} of magnitude 10^{-5} , here we have MSE_{local} of magnitude 10^{-3} . The \hat{u} in x end points are a bit bad and the model is averaging the initial sine condition with the boundary conditions.

[Initial: $u(x,0) = sin(\pi x), x \neq \{0,1\}$ Boundary: u(0,t) = u(1,t) = 0]

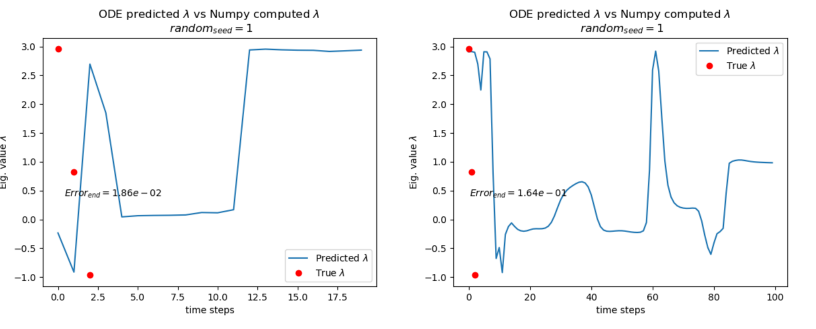
3.3 DNN for eigenvalues (Part d)

We look at different pseudo random symmetrical matrices using numpy seeds. Because of very unreliable convergence results we change to SGD optimizer attempting to get some control by setting a learning rate. We ended up with tanh as activation's and an architecture of 4 hidden layers with 50 neurons each. Note that $a^l = tanh(z^l)$ is fine for eigenvector elements in [0, 1], which is what was generated from the random matrices.

We managed sometimes to get absolute error $|\lambda_{predicted} - \lambda_{numpy}|$ of magnitude $\leq 10^{-2}$. This is a bit poor, but the overall results are still rather interesting. As you will see in the plots the predictions jump chaotically between multiple eigenvalues.

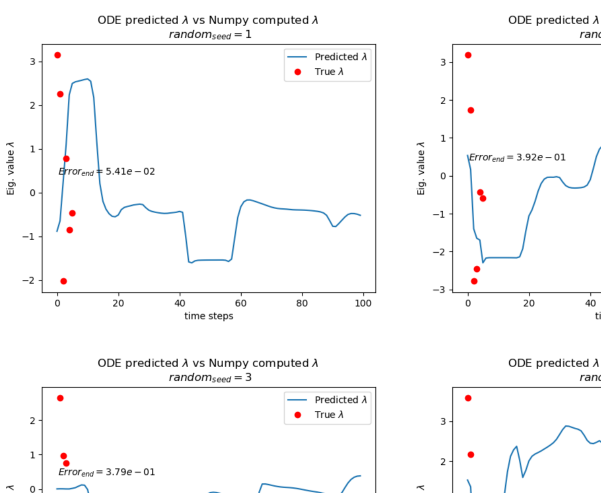
We test with matrices up to 6×6 . $Error_{end}$ is absolute error between nearest eigenvalue and the last predicted one. See plots in the next two pages.

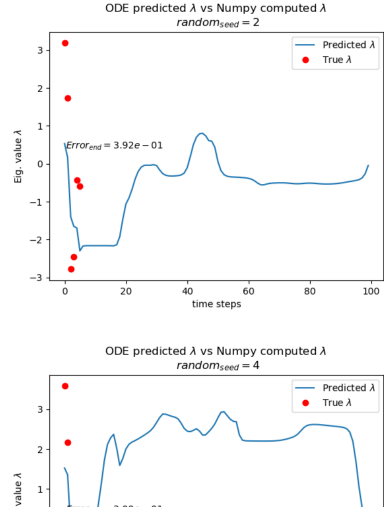
We seem to get the best results with 3×3 (or less) matrices:

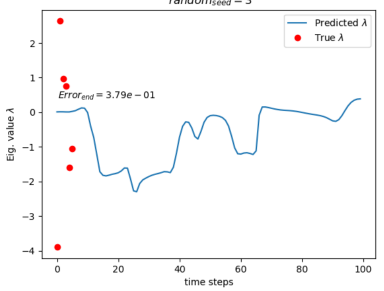


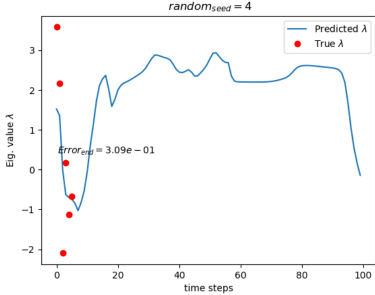
In the first plot, with fewer time steps, we get decent end error for the largest eigenvalue. The first/smallest peak is pretty close to the smallest λ_{true} , but for the middle λ_{true} the model is way off. In the second plot, with more time steps, it gets a little more chaotic and most of the peeks seem to be somewhat close to the true eigenvalues. If we use first peak, smallest peak and end prediction we are 'close' to all eigenvalues, but again we would want much higher precision.

Looking at 6×6 matrices:









Not really any convergence, but again some of the peaks goes toward different eigenvalues. Could be interesting spending some more time on this and improve. This approach on eigenvalues was motivated by the paper by Yi et al (Ref. 2) where they made a very successful model for finding the largest/smallest eigenvalue with a efficient recurrent neural network. Here we attempted to find every eigenvalue and it worked decent for 3×3 matrices if we pick certain stationary points/peaks. For greater dimensions the results got worse.

3.4 Unit test of HeatLearner class

A simple test was set up with u(x,t) properties $u_x = cos(x)$ and u(0,0) = 0.

We test for $MSE[u(x,0),\hat{u}(x,0)] < tol$ with $u(x,t)=sin(x),\,tol=10^{-3}$ using one hidden layer in the network.

 $(See unit_test.py)$

4 Summary (Part e)

The findings from the heat equation were overall that DNN's are more flexible and can be much faster on larger grids which can make it a very important part of solving large scale real world problems. For the finite-differences it was very precise on the 1d heat eq. but it is not flexible and scales badly for larger problems.

Finite-difference worked very well error wise. Cheapest model had $MSE_{total} = 1.16 \cdot 10^{-6}$, but with poorly scaling at $O(1/\Delta x^3)$ time complexity.

With the DNN model we trained it on a small 10×10 grid and managed errors MSE_{total} of magnitude 10^{-3} for grids 100×100 and up to much larger ones. The efficiency gains lays in minimizing the sub-sets used for training, while having sufficient out of sample precision.

Our implementation of the eigenvalue solver had interesting plots but poor convergence. We managed sometimes absolute errors of magnitude 10^{-2} to 10^{-4} . In Yi et al's paper (Ref. 2) they had pretty good results with symmetrical matrices with ability to achieve high computing performance since their recurrent model could parallel process asynchronously. In section 5 they did simulations on different random symmetrical matrices and found the biggest/smallest eigenvalues efficiently with precision 10^{-4} and stable convergence.

References

- 1. Lecture note on ODE in Tensorflow: https://github.com/krisbhei/DEnet/blob/master/DNN_Diffeq/example_ode_poisson.ipynb
- 2. Paper on Eigenvalues with Neural Networks (Yi, Fu and Jin Tang): https://www.sciencedirect.com/science/article/pii/S0898122104901101
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