# 9 Inverse Problems & Deep Learning: Basic Methodology

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Deep Learning in Computational Mechanics – an introductory course,

Herrmann et al. 2025





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## 9 Inverse Problems

Given a parametrized non-linear partial differential equations of the form

$$\frac{\partial^a u}{\partial^a t} + \mathcal{N}(u; \lambda) = 0, \qquad x \in \Omega, \qquad t \in \mathcal{T}$$

Inverse problems consider the identification of the differential equation given a (partial) solution u

- in the form of the non-linear differential operator  $\mathcal{N}(u; \lambda)$
- or/and the coefficients  $\lambda$
- or/and a order of derivative in time t

#### Examples of inverse problems

- X-ray computed tomography: geometry reconstruction using the attenuation of the x-rays
- Calculation of the earth's density from measurements of the variation in the gravity field
- Flaw identification through the disturbance of ultrasonic pulses
- Topology optimization

Inverse problems are often ill-posed. Ill-posed means that the solution is either not unique or a small variation of the input causes a large variation of the output.

Non-unique solution: If the answer is 42 then it could have been composed of 41+1 or 40+2 or....

# 9.1 Basic Methodology

Three main methodologies in deep learning

Physics-Informed Neural Networks (e.g. the elastic bar discussed in Chapter 5)

- Minimization of the residual of the partial differential equation
- Sub-method of physics-informed learning (see Chapter 10)

#### **Iterative Forward Solvers**

- Minimization of the residual between measurement data and the solution to the differential equation in an alternating fashion
- Enforcement of physical laws by use of classical methods for the solution of differential equations

#### Data-driven solvers

- Minimization of the residual between predictions and labelled data
- Physics is "learnt" by NN

Degree of enforcement of the underlying physics decreases from top to bottom

# 9.1.1 Physics-Informed Neural Networks

In general the network provides a prediction  $\widehat{X}$  given an input X

Inputs X can be, e.g.

- coordinates x, y
- solutions of PDEs u
- solutions of PDEs at previous time-steps  $u_{t-1}$
- physical parameters  $\lambda$

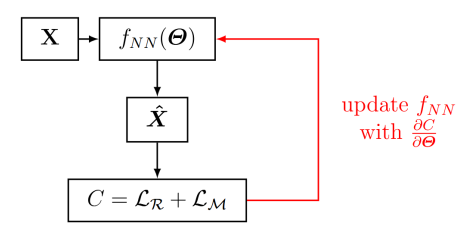
Predictions  $\widehat{X}$  can be

- solutions u
- inverse quantities  $\lambda$
- the non-linear differential operators  $\mathcal{N}(u;\lambda)$ .

The cost function is composed of

$$\mathcal{L}_M = \frac{1}{n} \sum_{i=1}^{m_M} (\tilde{u}_i - \hat{u}_i)^2$$
 ,

repeat for number of epochs



$$\mathcal{L}_{M} = \frac{1}{n} \sum_{i=1}^{m_{M}} (\tilde{u}_{i} - \hat{u}_{i})^{2}, \qquad \qquad \mathcal{L}_{R} = \frac{1}{m_{R}} \sum_{i=1}^{m_{R}} \left( \frac{\partial \hat{u}_{i}}{\partial t} + \widehat{\mathcal{N}} \left[ \hat{u}_{i}; \hat{\lambda}_{i} \right] \right)^{2}$$

### 9.1.2 Iterative Forward Solvers

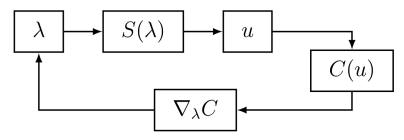
In general the network provides a prediction  $\widehat{X} \setminus \widehat{u}$  without the solution to the PDE given an input X

- The prediction  $\widehat{X} \setminus \widehat{u}$  is the used to solve the forward problem with a conventional approach yielding  $\widehat{u}$
- Physical laws are indirectly enforced by the forward solver
- The predicted solution  $\hat{u}$  is used to compute the measurement loss as cost function

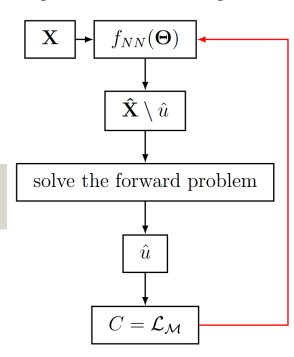
$$C = \mathcal{L}_M = \frac{1}{n} \sum_{i=1}^{m_M} (\tilde{u}_i - \hat{u}_i)^2$$

Only finds a single solution

Without a neural network  $f_{NN}(\Theta)$  the scheme is equivalent to standard gradient-based optimization:  $\underline{\lambda} = \widehat{X} \setminus \widehat{u}$ ,  $S(\lambda)$  is the forward solver But what is the point of the neural network?



repeat for number of epochs



update  $f_{NN}$  with  $\frac{\partial C}{\partial \mathbf{Q}}$ 

C(u)

## 9.1.2.1 Neural Network Ansatz in Iterative Solvers

#### Standard gradient-based optimizer

• Parametrization of  $\lambda$  with classical ansatz functions (e.g., FEM)

$$\hat{\lambda}(x) \approx \sum_{i} \lambda_{i} N_{i}(x)$$

- Where  $\lambda_i$  are the coefficients and  $N_i(x)$  the shape functions
- Given a set of coefficients  $\lambda_i$  the forward solution u is computed with the forward solver  $S(\lambda)$
- The quality of  $\lambda_i$  is assessed by the cost function C(u)
- The derivative  $\nabla_{\lambda} C$  is used to update the set of coefficients  $\lambda_i$

$$\lambda_i^{n+1} = \lambda_i^n - \alpha \nabla_{\lambda} C$$

#### **Iterative Forward Solver with Neural Network Ansatz**

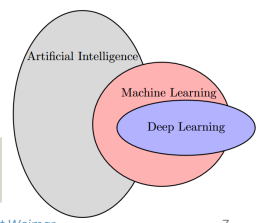
• Instead of relying on a classical parametrization of  $\lambda$ , a neural network is employed

$$\hat{\lambda}(x) \approx \lambda_{NN}(x; \mathbf{\Theta})$$

The coefficients are thus the neural network parameters Θ

As optimizing the neural network retrieves the parameters  $\Theta$  for <u>a single solution and cannot be applied to a different problem</u>, the procedure is **not machine learning**.

However, as a neural network is still being optimized, it can still be considered deep learning.



 $\nabla_{\lambda}C$ 

## 9.1.2.2 Motivation for a Neural Network Ansatz

On neural networks for generating better local optima in topology optimization, Herrmann et al. 2024

Consider the Rosenbrock function as toy example

$$g(y_1, y_2) = (1 - y_1)^2 + 100(y_2 - y_1^2)^2$$

Goal is to find the optimum from an initial guess  $y_1^{(0)}$ ,  $y_2^{(0)}$ 

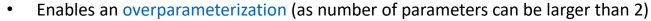
Standard gradient-based optimizer optimizes design variables  $y_1$ ,  $y_2$  directly

as optimization is performed directly, we call this linear ansatz

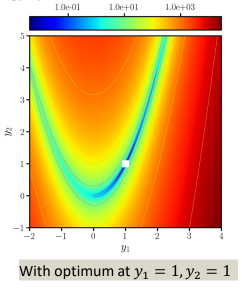
Iterative Forward Solver with Neural Network Ansatz parametrizes  $y_1$ ,  $y_2$  with a neural network

$$\hat{\lambda} = \begin{pmatrix} \hat{y}_1 \\ \hat{y}_2 \end{pmatrix} = f_{NN}(\boldsymbol{\xi}; \boldsymbol{\Theta})$$

Such that the parameters  $\Theta$  become the design variables (instead of  $y_1, y_2$ )

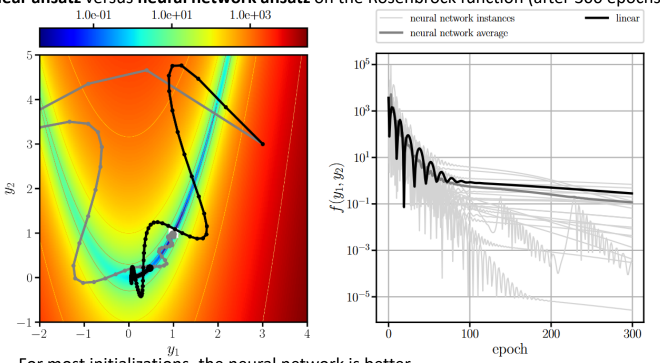


- ξ is kept constant throughout the optimization (it is an artifact of the neural network architecture, which requires an input)
- $\xi$  is typically set via random noise



## 9.1.2.2 Motivation for a Neural Network Ansatz

**Linear ansatz** versus **neural network ansatz** on the Rosenbrock function (after 300 epochs)



Further improvements are possible by reintegrating machine learning via transfer learning (see 9.2.5.1)

- For most initializations, the neural network is better
- Advantage is only possible when using the Adam optimizer for both parametrizations

### **Exercises**

- E.31 Neural Network Ansatz on Optimization Benchmarks (C)
  - Compare a neural network ansatz with a linear ansatz on four basic optimization benchmarks (Rosenbrock, Rastrigrin, Ackley, Levy) using different optimizers (gradient descent with momentum, AdaGrad, RMSprop, Adam, and L-BFGS).

### 9.1.3 Data-Driven Solvers

In data-driven solvers, a labelled data-set of a physical problem is known as

 $(\widetilde{X},\widetilde{y})$ 

- The neural network predicts the label  $\widehat{y}$  given an input X
- This process is repeated over the whole dataset
- The cost function is the error between the predictions and the labelled data

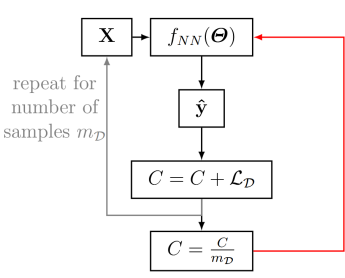
$$C = \frac{1}{m_D} \sum_{i=1}^{m_D} (\widetilde{\mathbf{y}}_i - \widehat{\mathbf{y}}_i)^2$$

Data driven solvers...

- require an off-line training phase ("expensive")
- learn multiple solutions
- deliver fast predictions in the on-line phase ("cheap")
- require a very large amount of data
- Do not enforce laws of physics. Physics is "observed and learnt"

Conceptually identical with the surrogate model for learning strain distributions in Chapter 3

repeat for number of epochs



update  $f_{NN}$  with  $\frac{\partial C}{\partial \Omega}$ 

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