Gradient-based Optimization

A short introduction to optimization in Deep Learning

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AGENDA

INTRODUCTION
  Motivation
  Probability framework
  Taylor approximation

GRADIENT DESCENT
  Gradient Descent
  Momentum
  Stochastic Gradient Descent

ADAPTATION AND PRECONDITIONING
  Adam
  Hessian
  Preconditioning
  Fisher Information Matrix

NATURAL GRADIENT
  Natural Gradient
  Riemannian manifold
  Empirical Fisher
  K-FAC

THOUGHTS
WHO AM I

Christian S. Perone

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Blog

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Section I

Introduction
**Motivation**

Mathematical optimization is the core of Machine Learning, without it we wouldn’t be able to find the needle in the haystack of the parameter space.
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- It materializes in Machine Learning by minimizing an objective function such as a divergence or any function that penalizes for mistakes of the model;
Motivation

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- It materializes in Machine Learning by minimizing an **objective function** such as a divergence or any function that penalizes for mistakes of the model;
- We will talk here about **local methods** that are characterized by the search of an optimal value within a neighboring set of parameter space;
Motivation

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- It materializes in Machine Learning by minimizing an **objective function** such as a divergence or any function that penalizes for mistakes of the model;
- We will talk here about **local methods** that are characterized by the search of an optimal value within a neighboring set of parameter space;
- We have a huge variety of methods that were recently developed, therefore **this talk is by far from being a comprehensive collection**. I will focus on **intuition and understanding**, instead of throwing algorithms.
Empirical Risk Minimization (ERM)

- On a supervised setting, we want to find a function or a model $f_\theta(\cdot)$ that describes the relationship between a random feature vector $\mathbf{x}$ and the label target vector $\mathbf{y}$. We assume a joint distribution $p_{\text{data}}(\mathbf{x}, \mathbf{y})$;
Empirical Risk Minimization (ERM)

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- We start by defining a loss function $L$, evaluated as $L(f_\theta(x), y)$ that gives us a penalization for the difference between predictions $f_\theta(x)$ and the true label $y$;
**Empirical Risk Minimization (ERM)**

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- We start by defining a loss function $L$, evaluated as $L(f_\theta(\mathbf{x}), \mathbf{y})$ that gives us a penalization for the difference between predictions $f_\theta(\mathbf{x})$ and the true label $\mathbf{y}$;
- Now, taking the expectation of the loss we have our risk $R$:

<table>
<thead>
<tr>
<th>Definition: Risk</th>
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\[
R(f) = \mathbb{E}_{\mathbf{x}, \mathbf{y} \sim p_{\text{data}}} [L(f_\theta(\mathbf{x}), \mathbf{y})] = \int L(f_\theta(\mathbf{x}), \mathbf{y}) \, dp_{\text{data}}(\mathbf{x}, \mathbf{y}),
\]

that we want to minimize.
Empirical Risk Minimization (ERM)

▸ However, we don’t know $p_{\text{data}}(x, y)$, we only have access to a sample training set $\mathcal{D} = (x_i, y_i) \sim p_{\text{data}}$;
Empirical Risk Minimization (ERM)

- However, we don’t know $p_{data}(x, y)$, we only have access to a sample training set $\mathcal{D} = (x_i, y_i) \sim p_{data}$;
- Therefore, we can approximate the risk with the empirical risk:

**Definition: Empirical Risk**

$$R_{emp}(f) = \frac{1}{n} \sum_{i=1}^{n} L(f_{\theta}(x_i), y_i)$$
Empirical Risk Minimization (ERM)

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Empirical Risk Minimization (ERM)

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Definition: Empirical Risk

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- The Empirical Risk Minimization (ERM) principle says that our learning algorithm should minimize the empirical risk;
- The MLE (Maximum Likelihood Estimation) can be posed as a special case of ERM where the loss function is the negative log-likelihood.
**Maximum Likelihood Estimation (MLE)**

Under the ERM framework we can describe the MLE cost function $J(\cdot)$ as:

$$J(\theta) = \mathbb{E}_{x,y \sim \hat{p}_{\text{data}}} - \log p_\theta(y | x)$$

where we define the cost as the expectation under the empirical distribution $\hat{p}_{\text{data}}$, as we only have access to a sample training set $\mathcal{D} = (x_i, y_i) \sim p_{\text{data}}$. 

We might be interested in let's say predicting a statistic of the distribution, such as the mean of $y$ using the predictor $f_\theta(x)$.

Our interest here in terms of optimization is:

$$\theta^* = \arg \min_\theta J(\theta),$$

where $\theta \in \mathbb{R}^n$. 

**Maximum Likelihood Estimation (MLE)**

Under the ERM framework we can describe the MLE cost function $J(\cdot)$ as:

$$J(\theta) = \mathbb{E}_{x,y \sim \hat{p}_{\text{data}}} \left[ -\log p_{\theta}(y \mid x) \right]$$

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- Our interest here in terms of optimization is:

$$\theta^* = \arg \min_{\theta} J(\theta), \text{ where } \theta \in \mathbb{R}^n$$
The global optimum
**Taylor approximation**

Let’s talk about a powerful calculus tool called *Taylor approximation*:

- Taylor approximation is based on the Taylor theorem¹:

\[
h(\theta) = f(\theta_0) + \nabla f(\theta_0)(\theta - \theta_0) + \frac{1}{2} \nabla^2 f(\theta_0)(\theta - \theta_0)^2,
\]

where we want an approximation of the function at the point \(\theta_0\);

---

¹Taylor’s theorem gives an approximation of a \(k\)-times differentiable function around a given point by a polynomial of degree \(k\). We’re using only up to second-order here.
Taylor approximation

Let's talk about a powerful calculus tool called *Taylor approximation*:

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\]

where \(h(\theta)\) is an approximation of the function at the point \(\theta_0\);

- This theorem is very powerful as it allows us to approximate any differentiable (and twice differentiable) function;

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- The \( \nabla^2 f(\cdot) \) is also called the Hessian, or \( H_f \). We will talk more about it later;

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- This theorem is very powerful as it allows us to approximate any differentiable (and twice differentiable) function;

- The \(\nabla^2 f(\cdot)\) is also called the Hessian, or \(\mathbf{H}_f\). We will talk more about it later;

- We will understand the deep connection of this approximation with Gradient Descent.

\(^1\)Taylor’s theorem gives an approximation of a \(k\)-times differentiable function around a given point by a polynomial of degree \(k\). We’re using only up to second-order here.
**Taylor approximation**

![Graph showing Taylor approximations of a function. The graph includes two curves: one for the first-order approximation (green dashed line) and one for the second-order approximation (red dashed line). The function values are plotted against the variable \( \theta \). The graph highlights the difference between first-order and second-order approximations around a specific point.]
**Taylor approximation**

![Graph showing Taylor approximation](image)

- **J(θ)**
- **First-order**
- **Second-order**
Taylors approximation
Taylor approximation

\[ J(\theta) \]

- First-order
- Second-order
Taylor approximation

- First-order
- Second-order
Taylor approximation
Taylor approximation
Taylor approximation
Taylor approximation
TaylOR approxIMATion
Taylor approximation

The graph illustrates the Taylor approximation of a function $J(\theta)$, showing the first-order and second-order approximations. The first-order approximation is represented by a dashed red line, while the second-order approximation is represented by a solid blue line. The graph highlights the difference in accuracy between the two approximations at different values of $\theta$. The x-axis represents $\theta$, ranging from -10.0 to 10.0, and the y-axis represents $J(\theta)$, ranging from -20 to 120.
Taylor approximation
Taylor approximation
Taylor approximation
Taylor approximation
Taylor approximation
Taylor approximation
Taylor approximation

![Graph showing Taylor approximation with two lines: one for First-order and another for Second-order. The graph plots J(θ) against θ. The First-order line is dashed green, and the Second-order line is dashed red. There is a point where the two lines meet, indicating the approximation.]
Taylor approximation
Taylor approximation
Taylor approximation in Jax

```python
from jax import grad

def taylor_first_order(θ, θ0):
    return f(θ0) + grad(f)(θ0)*(θ - θ0)

def taylor_second_order(θ, θ0):
    d1 = taylor_first_order(θ, θ0)
    d2 = 1./2. * grad(grad(f))(θ0) * (θ - a)**2
    return d1 + d2
```

Do not use greek symbols on your Python code, your colleagues will curse you.
### Taylor approximation in Jax

```python
from jax import grad

def taylor_first_order(θ, θ₀):
    return f(θ₀) + grad(f)(θ₀)*(θ - θ₀)

def taylor_second_order(θ, θ₀):
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    d2 = 1./2. * grad(grad(f))(θ₀) * (θ - a)**2
    return d1 + d2

>>> taylor_first_order(6.01, 6.0)
33.421864
>>> taylor_second_order(6.01, 6.0)
33.422104
>>> taylor_first_order(6.5, 6.0)
44.0067
>>> taylor_second_order(6.5, 6.0)
44.60597
```

---

Do not use greek symbols on your Python code, your colleagues will curse you.
Linear approximation plane

Let’s now think about that second-order term:

\[ h(\theta) = f(\theta_0) + \nabla f(\theta_0)(\theta - \theta_0) + \frac{1}{2} \nabla^2 f(\theta_0)(\theta - \theta_0)^2, \]

- first-order
- second-order

If we do a small step from \( \theta_0 \), what happens with the second-term?
Local approximation and second-order

Let’s now think about that second-order term:

\[ h(\theta) = f(\theta_0) + \nabla f(\theta_0)(\theta - \theta_0) + \frac{1}{2} \nabla^2 f(\theta_0)(\theta - \theta_0)^2, \]

If we do a small step from \( \theta_0 \), what happens with the second-term?
The steepest descent

- Even if \( f(\cdot) \) is very complex, **locally** it is simple, and we can use a simple function to approximate it, a linear function:

\[
h(\theta) \approx f(\theta_0) + \nabla f(\theta_0)(\theta - \theta_0)
\]

- This is also called *linearization*;
**The steepest descent**

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$$h(\theta) \approx f(\theta_0) + \nabla f(\theta_0)(\theta - \theta_0)$$

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- It is already apparent what we need now. How can we guarantee, locally, that we can always minimize the function (reduce the loss)?
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- We can just follow the slope (negative) of the approximation that is given by $-\nabla f(\theta_0)$;
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- It is already apparent what we need now. How can we guarantee, locally, that we can always minimize the function (reduce the loss) ?

- We can just follow the slope (negative) of the approximation that is given by $-\nabla f(\theta_0)$;

- No twice differentiability requirement, less computational resources;
Section II

Gradient Descent
Algorithm

The general gradient descent algorithm.

**Input:** initial weights $\theta^{(0)}$, iterations $T$, learning rate $\eta$

**Output:** final weights $\theta^{(T)}$

1. **for** $t = 0$ **to** $T - 1$
2. compute $\nabla L(\theta^{(t)})$
3. $\theta^{(t+1)} := \theta^{(t)} - \eta \nabla L(\theta^{(t)})$
4. **return** $\theta^{(T)}$
**Algorithm** The general gradient descent algorithm.

**Input:** initial weights $\theta^{(0)}$, iterations $T$, learning rate $\eta$

**Output:** final weights $\theta^{(T)}$

1. **for** $t = 0$ **to** $T - 1$
2. compute $\nabla L(\theta^{(t)})$
3. $\theta^{(t+1)} := \theta^{(t)} - \eta \nabla L(\theta^{(t)})$
4. **return** $\theta^{(T)}$

The important part here is the iterative rule:

$$\theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(\theta^{(t)})$$

How much we move
Gradient Descent - Loss surface
Gradient Descent - Loss surface
Gradient Descent - Loss surface
Gradient Descent - Loss surface
Gradient Descent - Loss surface
Gradient Descent - Loss surface
Learning Rate

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Learning Rate

\[\alpha = 0.06\]

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Learning Rate

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Learning Rate

\[ \alpha = 0.19 \]

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Learning Rate

\[ \alpha = 0.38 \]

\[ g(w) = 0.38 \]

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Learning Rate

\[ \alpha = 0.44 \]

\[ g(w) \]

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\[ \alpha = 0.50 \]

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Learning Rate

\[ \alpha = 0.56 \]

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**Learning Rate**

\[
g(w) = 0.63
\]

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Learning Rate

\[ \alpha = 0.69 \]

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Learning Rate

\[ g(w) = 0.75 \]

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Learning Rate

\[ \alpha = 0.81 \]

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\[ \alpha = 1.00 \]

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**Learning Rate**

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**High curvatures**

Gradient descent can suffer on some pathological curvatures and cause a lot of oscillations:

Momentum

Momentum is a method to damp out oscillations:

**Vanilla gradient descent:**

\[ \theta(t+1) = \theta(t) - \eta \nabla L(\theta(t)) \]
MOMENTUM

Momentum is a method to damp out oscillations:

**Vanilla gradient descent:**

\[ \theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(\theta^{(t)}) \]

**Momentum:**

\[ V^{(t+1)} = \beta V^{(t)} + \nabla L(\theta^{(t)}) \]

\[ \theta^{(t+1)} = \theta^{(t)} - \eta V^{(t+1)} \]

Momentum buffer
Momentum

Momentum is a method to damp out oscillations:

Vanilla gradient descent:

\[ \theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(\theta^{(t)}) \]

Momentum:

\[ V^{(t+1)} = \beta V^{(t)} + \nabla L(\theta^{(t)}) \]

\[ \theta^{(t+1)} = \theta^{(t)} - \eta V^{(t+1)} \]

Momentum works by acceleration and smoothing, it makes the trajectories to take more time to react to changes in the loss landscape;
Momentum

Momentum is a method to damp out oscillations:

**Vanilla gradient descent:**

$$\theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(\theta^{(t)})$$

**Momentum:**

$$V^{(t+1)} = \beta \underbrace{V^{(t)} + \nabla L(\theta^{(t)})}_{\text{Constant}}$$

$$\theta^{(t+1)} = \theta^{(t)} - \eta \underbrace{V^{(t+1)}}_{\text{Momentum buffer}}$$

- Momentum works by acceleration and smoothing, it makes the trajectories to take more time to react to changes in the loss landscape;
- Note that with $\beta = 0$ we recover vanilla Gradient descent;
**Momentum**

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Momentum

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Momentum

Pause for a quick demo from Lili Jiang, from:
https://github.com/lilipads/gradient_descent_viz
Stochastic Gradient Descent (SGD)

It turns out that we don’t quite need to compute the gradients $\nabla L(\theta)$ over the whole dataset at every iteration of Gradient descent:

$$\theta^{(t+1)} = \theta^{(t)} - \eta \nabla L_i(\theta^{(t)})$$

where we do random sampling (or not, we can stratify too, in practice it can lead to better results) of individual samples $i$ at every step.

---

3Robbins and Monro, “A Stochastic Approximation Method”, 1951
Stochastic Gradient Descent (SGD)

It turns out that we don’t quite need to compute the gradients $\nabla L(\theta)$ over the whole dataset at every iteration of Gradient descent:

$$\theta(t+1) = \theta(t) - \eta \nabla L_i(\theta(t))$$

where we do random sampling (or not, we can stratify too, in practice it can lead to better results) of individual samples $i$ at every step.

- Much more efficient (don’t have to compute gradient for entire dataset);
- Noise (can be beneficial);
- Lots of redundancy on real datasets;
- Highly correlation at early steps (similar gradients SGD vs GD);

SGD can be traced back to 1950s work on the Robbins–Monro algorithm.³

³Robbins and Monro, “A Stochastic Approximation Method”, 1951
Graphics Processing Unit (GPUs)

Most of the operations in Machine Learning ends up being lowered to GEMM (*General Matrix Multiplication*) and MAC (*Multiply–accumulate operation*) operations. To leverage these massively parallel engines, we need to provide enough data to take advantage of the parallelization potential.

MINI-BATCH SGD

That’s why using mini-batches instead of individual samples on SGD is a perfect marriage of having better gradient estimates together with improved parallelization:

\[
\tilde{\nabla} L(\theta(t)) = \frac{1}{|B|} \sum_{i \in B} \nabla L_i(\theta(t))
\]

\[\theta(t+1) = \theta(t) - \eta \tilde{\nabla} L(\theta(t))\]

Estimated gradients
**Mini-batch SGD**

That’s why using mini-batches instead of individual samples on SGD is a perfect marriage of having better gradient estimates together with improved parallelization:

\[
\tilde{\nabla} L(\theta^{(t)}) = \frac{1}{|B|} \sum_{i \in B} \nabla L_i(\theta^{(t)})
\]

Batch size

\[
\theta^{(t+1)} = \theta^{(t)} - \eta \quad \tilde{\nabla} L(\theta^{(t)})
\]

Estimated gradients

If we do random sampling, then:

\[
\mathbb{E}[\tilde{\nabla} L(\theta^{(t)})] = \nabla L(\theta)
\]

Unbiased estimate
Section III

Adaptation and Preconditioning
Adaptive Moment Estimation (Adam)

There are many adaptive methods, we will focus on one of the most frequently used in Deep Learning, the *Adaptive Moment Estimation* \(^4\), also called **Adam**.

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- Many of these algorithms are still not well understood, lots of folklore in the field;
- Will try to focus on building intuition from the original algorithm.

Adaptive Moment Estimation (Adam)

**Algorithm** \( g_t^2 = g_t \odot g_t \). Good defaults: \( \alpha = 0.001, \beta_1 = 0.9, \beta_2 = 0.999 \) and \( \epsilon = 10^{-8} \). \( \beta_1^t \) and \( \beta_2^t \) are \( \beta_1 \) and \( \beta_2 \) to the power \( t \).

**Require:** \( \beta_1, \beta_2 \in [0, 1) \): Exponential decay rates for the moment estimates

**Require:** \( f(\theta) \): Stochastic objective function with parameters \( \theta \)

**Require:** \( \theta_0 \): Initial parameter vector, \( \alpha \): Stepsize

\[
\begin{align*}
m_0 &\leftarrow 0 \text{ (Initialize 1st moment vector)} \\
v_0 &\leftarrow 0 \text{ (Initialize 2nd moment vector)} \\
t &\leftarrow 0 \text{ (Initialize timestep)} \\
\textbf{while} \ \theta_t \text{ not converged do} \\
\hspace{1em} t &\leftarrow t + 1 \\
g_t &\leftarrow \nabla_\theta f_t(\theta_{t-1}) \text{ (Get gradients w.r.t. stochastic objective at timestep } t) \\
m_t &\leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t \text{ (Update biased first moment estimate)} \\
v_t &\leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2 \text{ (Update biased second raw moment estimate)} \\
\widehat{m}_t &\leftarrow m_t/(1 - \beta_1^t) \text{ (Compute bias-corrected first moment estimate)} \\
\widehat{v}_t &\leftarrow v_t/(1 - \beta_2^t) \text{ (Compute bias-corrected second raw moment estimate)} \\
\theta_t &\leftarrow \theta_{t-1} - \alpha \cdot \widehat{m}_t/((\sqrt{\widehat{v}_t} + \epsilon)) \text{ (Update parameters)} \\
\textbf{end while} \\
\textbf{return} \ \theta_t \text{ (Resulting parameters)}
\]
Adaptive Moment Estimation (Adam)

Lots of things going on here, let’s focus on how moments are being computed and neglect bias correction and initialization:

\[ g_t \leftarrow \nabla_{\theta} f_t(\theta_{t-1}) \]
\[ m_t \leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t \]
\[ v_t \leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2 \]

And the parameter updates:

\[ \theta_t \leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \epsilon} \]
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And the parameter updates:

\[ \theta_t \leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \epsilon} \]

- Do you recognize \( m_t \) ?
- What happens when the uncentered variance grows ?
The good, the bad, and the Hessian

- The convergence rate of Gradient descent is deeply connected to the curvature of the landscape it is trying to optimize;
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▶ The Hessian matrix $H_f$ carries information about the curvature, therefore we usually use it understand problems or even make them better conditioned;
The good, the bad, and the Hessian

▶ The convergence rate of Gradient descent is deeply connected to the curvature of the landscape it is trying to optimize;
▶ The Hessian matrix $H_f$ carries information about the curvature, therefore we usually use it understand problems or even make them better conditioned;
▶ The $H_f$ is often very costly to compute for real-life problems, therefore much of the work rely on approximating it or computing information about it without having to materialize the entire matrix;
The $\mathbf{H}_f$ is a square matrix of 2nd-order partial derivatives. Let’s compute the $\mathbf{H}_f$ of $f(x, y) = x^2y + xy^3$, starting with first-order:

$$\frac{\partial f}{\partial x} = 2xy + y^3, \quad \frac{\partial f}{\partial y} = x^2 + 3xy^2$$

Note that the $\mathbf{H}_f$ can be constant and not depend on variables or depend only on some of them. We will see this case later.
**Hessian**

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**Second order**

$$\frac{\partial^2 f}{\partial x^2} = 2y \quad , \quad \frac{\partial^2 f}{\partial y \partial x} 2x + 3y^2 \quad , \quad \frac{\partial^2 f}{\partial x \partial y} = 2x + 3y^2 \quad , \quad \frac{\partial^2 f}{\partial y^2} = 6xy$$

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Second order

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Hessian

$$H_f = \begin{bmatrix} \frac{\partial^2 f}{\partial x^2} & \frac{\partial^2 f}{\partial y \partial x} \\ \frac{\partial^2 f}{\partial x \partial y} & \frac{\partial^2 f}{\partial y^2} \end{bmatrix} = \begin{bmatrix} 2y & 2x + 3y^2 \\ 2x + 3y^2 & 6xy \end{bmatrix}$$

Note that the $H_f$ can be constant and not depend on variables or depend only on some of them. We will see this case later.
Hessian Eigenvalues

All positive eigenvalues
(positive definite)
Hessian Eigenvalues

All positive eigenvalues
(positive definite)

All negative eigenvalues
(negative definite)
The **Condition Number**, also defined as $\kappa$, is the ratio of maximum and minimum eigenvalues ($\lambda_{\text{max}}$ and $\lambda_{\text{min}}$) of the Hessian $H_f$:

$$\kappa = \frac{\lambda_{\text{max}}}{\lambda_{\text{min}}}$$

- When $\kappa$ is high we say that the problem is ill-conditioned;
- **Steepest descent convergence rate is slow** for ill-conditioned problems;
- Let's understand it on a quadratic problem to gain intuition.
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**Condition Number**

\[ f(\theta) = \frac{1.0}{2.0}\theta_1 + \frac{2.0}{2.0}\theta_2 \]

\[ H_f \begin{bmatrix} 1.0 & 0.0 \\ 0.0 & 2.0 \end{bmatrix} \]

\[ \kappa = 2.00 \ (\lambda_{max} = 2.0, \lambda_{min} = 1.0) \]
**Condition Number**

\[ f(\theta) = \frac{1.5}{2.0} \theta_1 + \frac{2.0}{2.0} \theta_2 \]

\[ H_f = \begin{bmatrix} 1.5 & 0.0 \\ 0.0 & 2.0 \end{bmatrix} \]

\[ \kappa = 1.33 \ (\lambda_{max} = 2.0, \lambda_{min} = 1.5) \]
**Condition Number**

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**Condition Number**

\[ f(\theta) = \frac{2.5}{2.0} \theta_1 + \frac{2.0}{2.0} \theta_2 \]

\[ \kappa = 1.25 \ (\lambda_{max} = 2.5, \lambda_{min} = 2.0) \]
**Condition Number**

\[ f(\theta) = \frac{3.0}{2.0} \theta_1 + \frac{2.0}{2.0} \theta_2 \]

\[
\begin{bmatrix}
3.0 & 0.0 \\
0.0 & 2.0
\end{bmatrix}
\]

\[ \kappa = 1.50 \ (\lambda_{max} = 3.0, \lambda_{min} = 2.0) \]
**Condition Number**

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\[ \kappa = 1.75 \ (\lambda_{max} = 3.5, \lambda_{min} = 2.0) \]
Condition Number

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Hessian eigenvalue spectral density (ESD)


ResNet with depth 20 trained on Cifar-10. ResNet_{BN} is the ResNet without Batch Normalization and the ResNet_{Res} is without the residual connections. In \(^6\), they also show that the distribution seem to composed of two parts: the bulk around zero, and the edges scattered away from zero.

Preconditioning

From Adam’s Original Paper:

(…) Like natural gradient descent (NGD)\(^7\), Adam employs a **preconditioner** that adapts to the **geometry of the data**, since \(\hat{v}_t\) is an approximation to the **diagonal of the Fisher information matrix**\(^8\); (…) 

▶ Preconditioning can be viewed as a change in the geometry;

---

\(^7\) Amari, “Natural Gradient Works Efficiently in Learning”, 1998

\(^8\) Pascanu and Bengio, “Revisiting Natural Gradient for Deep Networks”, 2013
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- It can help with poorly conditioned problems;

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- Preconditioning can be viewed as a change in the geometry;
- It can help with poorly conditioned problems;
- We will talk about the Fisher Information Matrix (FIM) later;

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Preconditioning

\[ \theta^{(t+1)} = \theta^{(t)} - \eta \nabla L(\theta^{(t)}) \]
Preconditioning

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Gradients
Preconditioning

\[ \theta^{(t+1)} = \theta^{(t)} - \eta \left[ P \nabla L(\theta^{(t)}) \right] \]

- Preconditioner
- Gradients
Preconditioning

\[ \theta^{(t+1)} = \theta^{(t)} - \eta I \nabla L(\theta^{(t)}) \]

\( \theta \) (t+1) = \theta (t) - \eta \quad \text{Identity} \quad \nabla L(\theta (t)) \quad \text{Gradients}
Preconditioning

\[ \theta(t+1) = \theta(t) - \eta \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \nabla L(\theta(t)) \]

Identity

Gradients
Preconditioning

\[ \theta^{(t+1)} = \theta^{(t)} - H_{L}^{-1} \nabla L(\theta^{(t)}) \]

- Can be interpreted as an iterative minimization of the quadratic approximation, we’re using a 2nd-order term here, remember the Taylor approximation?

---

The superscript \( t \) was omitted from the \( H_{L}^{-1} \) for clarity.
Preconditioning

\[ \theta^{(t+1)} = \theta^{(t)} - (H_L + \lambda I)^{-1} \nabla L(\theta^{(t)}) \]

Damped Hessian \hspace{2cm} Gradients

The superscript \( t \) was omitted from the \( H_L \) for clarity.
Preconditioning

\[ f(\theta) = \frac{5.0}{2.0} \theta_1 + \frac{2.0}{2.0} \theta_2 \]

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Let’s think about what the preconditioner is doing in this situation, we have a point \( \theta \in \mathbb{R}^2 \) at \( \theta = (0.5, 0.5) \) and we have that:

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Let’s think about what the preconditioner is doing in this situation, we have a point \( \theta \in \mathbb{R}^2 \) at \( \theta = (0.5, 0.5) \) and we have that:

\[
\nabla f(\theta) = (2.5, 1.0)
\]

\[
H_f = \begin{bmatrix} 5.0 & 0.0 \\ 0.0 & 2.0 \end{bmatrix}
\]

\[
\theta - \mathbf{H}_L^{-1} \nabla f(\theta) = (0., 0.)
\]

\[
\theta - \nabla f(\theta) = (-2., -0.5)
\]
**Hessian as preconditioner**

\[ f(\theta) = \frac{1.0}{2.0} \theta_1 + \frac{2.0}{2.0} \theta_2 \]

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Hessians as preconditioner

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Difficulties of the Hessian preconditioning

▸ Using the Hessian as preconditioner is the basis of the Newton’s method;

---

11 Dauphin et al., “Identifying and attacking the saddle point problem in high-dimensional non-convex optimization”, 2014
12 Yao et al., PYHESSIAN: Neural networks through the lens of the hessian, 2019
13 Martens, Deep learning via Hessian-free optimization, 2010
Difficulties of the Hessian preconditioning

- Using the Hessian as preconditioner is the basis of the Newton’s method;
- Invariant to affine transformations;

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Difficulties of the Hessian preconditioning

- Using the Hessian as preconditioner is the basis of the Newton’s method;
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- However, a model with 23 million parameters (i.e. ResNet-50), what is the space complexity to store the $\mathbf{H}_f$ and the computational complexity to invert it?

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- Using the Hessian as preconditioner is the basis of the Newton’s method;
- Invariant to affine transformations;
- However, a model with 23 million parameters (i.e. ResNet-50), what is the space complexity to store the $H_f$ and the computational complexity to invert it?
- Difficult on non-convex problems, not always invertible, attracted by saddle points\(^\text{11}\);
- Among other reasons, you now understand all the efforts into Hessian approximations\(^\text{12}\), alternative curvature matrices and hessian-free optimization\(^\text{13}\).

\(^{11}\) Dauphin et al., “Identifying and attacking the saddle point problem in high-dimensional non-convex optimization”, 2014

\(^{12}\) Yao et al., PYHESSIAN: Neural networks through the lens of the hessian, 2019

\(^{13}\) Martens, Deep learning via Hessian-free optimization, 2010
**Saddle points**

![3D plot of saddle points](image)
Fisher Information Matrix (FIM)

Going back to the Adam’s article:

From Adam’s original paper:

(… ) Like natural gradient descent (NGD) \(^{14}\), Adam employs a **preconditioner** that adapts to the **geometry of the data**, since \(\hat{v}_t\) is an approximation to the **diagonal of the Fisher information matrix** \(^{15}\); (…)

▶ We now know what a preconditioner means;
▶ The missing ingredient now is the **Fisher Information Matrix** (also known as FIM).

---

\(^{14}\) Amari, “Natural Gradient Works Efficiently in Learning”, 1998

\(^{15}\) Pascanu and Bengio, “Revisiting Natural Gradient for Deep Networks”, 2013
**Fisher Information Matrix (FIM)**

The Fisher Information Matrix is the covariance of the score function (gradients of the log-likelihood function) with expectation over the model’s predictive distribution (pay attention to this detail).

**Definition:** Fisher Information Matrix

\[
F_\theta = \mathbb{E}_{y \sim p_\theta(y|x), \; x \sim p_{\text{data}}} \left[ \nabla_\theta \log p_\theta(y|x) \nabla_\theta \log p_\theta(y|x)^T \right]
\]

Where \( F_\theta \in \mathbb{R}^{n \times n} \).
**Fisher Information Matrix (FIM)**

The Fisher Information Matrix is the covariance of the score function (gradients of the log-likelihood function) with expectation over the *model’s predictive distribution* (pay attention to this detail).

**Definition: Fisher Information Matrix**

\[
\mathbf{F}_\theta = \mathbb{E}_{\begin{subarray}{l} y \sim p_\theta(y|x) \\ x \sim p_{\text{data}} \end{subarray}} \left[ \nabla_\theta \log p_\theta(y|x) \nabla_\theta \log p_\theta(y|x)^T \right]
\]

Where \( \mathbf{F}_\theta \in \mathbb{R}^{n \times n} \). We often approximate it using input samples (*y* is still from model’s predictive distribution), as we don’t have access to \( p_{\text{data}} \):

\[
\mathbf{F}_\theta = \frac{1}{N} \sum_{i=1}^{N} \nabla_\theta \log p_\theta(y|x_i) \nabla_\theta \log p_\theta(y|x_i)^T
\]
Kullback-Leibler divergence

$KL[P \parallel Q] = 5683.243$
Kullback-Leibler divergence

\[ KL[P \parallel Q] = 3488.456 \]
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Fisher Information Matrix (FIM)

▶ We can parametrize the same distribution family on many different ways;
▶ Moving in the parameter space using the Euclidean distance as a metric makes us tied to the particular parametrization;

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\[\text{For a full derivation please refer to: Ratliff, N. (2013). Information Geometry and Natural Gradients.}\]

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\(^{17}\)Martens, “New insights and perspectives on the natural gradient method”, 2014
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- We won’t be talking here, but the Fisher has a strong connection to the Hessian and the Generalized Gauss-Newton (GGN), please refer to$^{17}$ if you are interested.


$^{17}$Martens, “New insights and perspectives on the natural gradient method”, 2014
Section IV

Natural Gradient
Natural Gradient

When we do a preconditioning on Gradient descent using the Fisher, we have the Natural Gradient Descent: 

\[ \theta^{(t+1)} = \theta^{(t)} - \eta \mathbf{F}_{\theta}^{-1} \nabla L(\theta^{(t)}) \]

FIM Gradients

---

18 Amari, “Natural Gradient Works Efficiently in Learning”, 1998
19 Léon Bottou, Curtis, and Nocedal, Optimization methods for large-scale machine learning, 2018
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When we do a preconditioning on Gradient descent using the Fisher, we have the Natural Gradient Descent $^{18}$:

$$\theta^{(t+1)} = \theta^{(t)} - \eta F^{-1}_\theta \nabla L(\theta^{(t)})$$

- It converges much faster than ordinary Gradient descent;

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- It moves on the distribution space manifold, invariant with respect to all differentiable and invertible transformations $^{19}$;

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- Given that the FIM is the result of an outer-product, it is always PSD (positive semidefinite matrix);

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- Given that the FIM is the result of an outer-product, it is always PSD (positive semidefinite matrix);
- It is still a $n \times n$ matrix, that needs to be inverted;

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Natural Gradient

The natural gradient is connected to *information geometry*.

- In a **Euclidean space**, the shortest path between two points is always the straight line;

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The natural gradient is connected to information geometry\(^{20}\).

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- The metric tensor represents this curvature and can be different at different points;
- With the natural gradient, we are moving in this Riemannian manifold using the Fisher as the metric tensor;
- Parameters move more quickly along directions that have a small impact on the decision function, and more cautiously along directions that have a large impact\(^{21}\);

---

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\(^{21}\)Léon Bottou, Curtis, and Nocedal, Optimization methods for large-scale machine learning, 2018
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- A manifold is a collection of points, where locally (but not globally), is Euclidean;
- A metric induces an inner product on the tangent space at each point on the manifold;
- The metric on the statistical manifold is unique, it is an intrinsic geometry;
- In Euclidean space we don’t care because the metric is constant everywhere;

Empirical Fisher

There is a lot of confusion\(^{22}\) about the Fisher Information Matrix\(^{23}\).

▶ In some scenarios you will see people sampling \(y \sim p\text{data} \) too instead of sampling from the model’s predictive distribution \(y \sim p_{\theta}(y|x)\);

\(^{22}\)I blame evil people who omit expectation qualifiers about where \(y\) is coming from.

\(^{23}\)Kunstner, Balles, and Hennig, “Limitations of the empirical fisher approximation for natural gradient descent”, 2019
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- This is called the Empirical Fisher, Empirical FIM or just EF:

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- It turns out that Adam is using the Empirical Fisher, and to make things more confusing it is using the square root of it.

---

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\textsuperscript{23}Kunstner, Balles, and Hennig, “Limitations of the empirical fisher approximation for natural gradient descent”, 2019
Adam and the Natural Gradient Descent

Original Adam paper\textsuperscript{24} claims that Adam is an approximation to the natural gradient descent (diagonal of the FIM):

\[
g_t \leftarrow \nabla_{\theta} f_t(\theta_{t-1})
\]

\[
m_t \leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t
\]

\[
v_t \leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot \frac{g_t^2}{g_t}
\]

\[
\hat{m}_t \leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \epsilon}
\]

\textsuperscript{24}Kingma and Ba, “Adam: a Method for Stochastic Optimization”, 2015

\textsuperscript{25}Staib et al., “Escaping saddle points with adaptive gradient methods”, 2019
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  \theta_t &\leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \epsilon}
\end{align*}
\]

However, the approximation is only valid near optimality (why?). The exponent is also different, since Adam is taking square root, it doesn’t change direction of the descent (only stepsize)\textsuperscript{25}.

\textsuperscript{24}Kingma and Ba, “Adam: a Method for Stochastic Optimization”, 2015
\textsuperscript{25}Staib et al., “Escaping saddle points with adaptive gradient methods”, 2019
Empirical Fisher


- Vector fields of the gradients conditioned using the FIM vs using the EF are very different;
- Are they close to each other close to the minima?
Empirical Fisher

EF is a good approximation of the Fisher at the minimum if model is well-specified. Otherwise, even at the minimum and with a large amount of samples, it can be a very poor approximation.\(^{26}\)

Is EF just the non-central gradient covariance matrix, working as variance reduction instead of curvature adaptation?


The epsilon that might not be an epsilon

Many implementations use the epsilon to avoid division by zero:

\[
\begin{align*}
g_t & \leftarrow \nabla_{\theta} f_t(\theta_{t-1}) \\
m_t & \leftarrow \beta_1 \cdot m_{t-1} + (1 - \beta_1) \cdot g_t \\
v_t & \leftarrow \beta_2 \cdot v_{t-1} + (1 - \beta_2) \cdot g_t^2 \\
\tilde{\theta}_t & \leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \epsilon}
\end{align*}
\]

---

27 Choi et al., *On empirical comparisons of optimizers for deep learning*, 2019
The epsilon that might not be an epsilon

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\theta_t & \leftarrow \theta_{t-1} - \alpha \cdot \frac{\hat{m}_t}{\sqrt{\hat{v}_t} + \epsilon}
\end{align*}
\]

However, remember about the damping mechanism? The \( \epsilon \) can be seen as setting a trust region radius \(^{27}\).

\(^{27}\)Choi et al., *On empirical comparisons of optimizers for deep learning*, 2019
Fisher is a big Fisher

Computing the inverse of the diagonal Fisher is easy, but computing the inverse of the “full” Fisher $F^{-1}$ and the natural gradient $F^{-1}_\theta \nabla L(\theta(t))$, on networks with millions of parameters, is just intractable.

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▶ What about other structural approximations? We don’t want to lose all of the off-diagonal structure;

▶ However, there are certain goals that we should be ideally try to achieve: memory (remember we have $F \in \mathbb{R}^{n \times n}$) and computation (we want to have an efficient $F^{-1}$);

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▷ What about other structural approximations? We don’t want to lose all of the off-diagonal structure;

▷ However, there are certain goals that we should ideally try to achieve: memory (remember we have $\mathbf{F} \in \mathbb{R}^{n \times n}$) and computation (we want to have an efficient $\mathbf{F}^{-1}$);

▷ That is what Kronecker-Factored Approximate Curvature (K-FAC) $^{28}$ proposes, an structured approximation to natural gradient descent;

Kronecker product

\[
A \otimes B := \begin{pmatrix}
[A]_{1,1} B & \cdots & [A]_{1,n} B \\
\vdots & \ddots & \vdots \\
[A]_{m,1} B & \cdots & [A]_{m,n} B
\end{pmatrix} \in \mathbb{R}^{mn \times nb}
\]

\[
A \in \mathbb{R}^{m \times n}, B \in \mathbb{R}^{a \times b}: \text{Kronecker factors}
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A \in \mathbb{R}^{m \times n}, B \in \mathbb{R}^{a \times b} \text{: Kronecker factors}
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**Fisher approximation**

Fisher approximation

Fisher approximation

Kronecker Inversion

Kronecker product has a very interesting and critical property:

$$(A \otimes B)^{-1} = A^{-1} \otimes B^{-1}$$

This means that the inverse of the product is the same as the product of the inverse of the operands. And this gives us a critical performance speed-up because we just need to invert small factor matrices.
If you want to play with K-FAC on PyTorch, you can try using Backpack:

```python
from torch import nn
from backpack import backpack, extend
from backpack.extensions import KFAC
from backpack.utils.examples import load_one_batch_mnist
from backpack.utils import kroneckers

X, y = load_one_batch_mnist(batch_size=512)

model = nn.Sequential(
    nn.Flatten(),
    nn.Linear(784, 10)
)

lossfunc = nn.CrossEntropyLoss()

model = extend(model)
lossfunc = extend(lossfunc)

loss = lossfunc(model(X), y)

with backpack(KFAC(mc_samples=1)):
    loss.backward()
```

(named_params = dict(model.named_parameters()))
layer_weights = named_params['1.weight']
# layer_weights.grad = [10, 784]

kfac_f1, kfac_f2 = layer_weights.kfac
# kfac_f1 = [10, 10]
# kfac_f2 = [784, 784]

mat = kroneckers.two_kfacs_to_mat(kfac_f1, kfac_f2)
# mat = [7840, 7840]

---

29 Dangel, Kunstner, and Hennig, "BackPACK: Packing more into backprop", 2019
Kronecker Matrices

\[ \mathbf{A} \in \mathbb{R}^{10 \times 10} \quad \mathbf{B} \in \mathbb{R}^{784 \times 784} \quad \tilde{\mathbf{G}}(\theta) \in \mathbb{R}^{7840 \times 7840} \]

Note that the colormap of the \( \tilde{\mathbf{G}}(\theta) \) was changed for visualization purposes.
Some empirical results

Section V

Thoughts
BENCHMARKING OPTIMIZERS


Lines in gray (—, smoothed by cubic splines for visual guidance only) show the relative improvement for a certain tuning and schedule (compared to the one-shot tuning without schedule) for all 14 optimizers on all eight test problems. The median over all lines is plotted in orange (—) with the shaded area indicating the area between the 25th and 75th percentile. ³⁰

³⁰ Schmidt, Schneider, and Hennig, “Descending through a Crowded Valley – Benchmarking Deep Learning Optimizers”, 2020
To think

▶ Do we really need normalization techniques (i.e. Batch Normalization) if we can come up with optimization methods that embed invariant properties?
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- What are the barriers to the use of second-order or approximately second-order methods? Are we going to see more software support?
- Are we driving towards more hyper-parameters or more robust methods?
- What are properties of the different solutions that different optimization methods can achieve?
Q&A
References I


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References IV


