Mathematics of Data: From Theory to Computation

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Lecture 9: Generalization in deep learning

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Outline

This lecture :

- The classical trade-off between model complexity and risk
- Generalization bounds via uniform convergence
- The generalization mystery in deep learning
- Implicit regularization of optimization algorithms
- ▶ Double descent curves: Generalization bounds via bias-variance decomposition
- Generalization bounds based on algorithmic stability
- Boosting

Next lecture :

Optimization in Deep Learning

Understanding the trade-off between model complexity and expected risk



Models

Let $[\mathcal{X}_i : i = 1, ...]$ be a nested sequence of parameter domain, i.e., $\mathcal{X}_i \subseteq \mathcal{X}_{i+1}$. For example, let \mathcal{X}_i = neural networks with *i* neurons.

1. $R_n(\mathbf{x}_i^{\star}) = \min_{\mathbf{x} \in \mathcal{X}_i} R_n(\mathbf{x})$: ERM solution over \mathcal{X}_i

- 2. $R(\mathbf{x}_{i}^{\star})$: True risk of the ERM solution over \mathcal{X}_{i}
- 3. $\sup_{\mathbf{x} \in \mathcal{X}_i} |R(\mathbf{x}) R_n(\mathbf{x})|$: Worst-case Generalization error of \mathcal{X}_i

Practical performance of the ERM estimator

$$R(\mathbf{x}_{i}^{\star}) \leq \min_{\mathbf{x} \in \mathcal{X}_{i}} R_{n}(\mathbf{x}) + \sup_{\mathbf{x} \in \mathcal{X}_{i}} |R(\mathbf{x}) - R_{n}(\mathbf{x})|$$
(1)

As we increase the index $i \rightarrow i+1$ of the parameter domain, i.e., we choose a larger (more complex) model

- 1. The minimum empirical risk decreases: $\min_{\mathbf{x} \in \mathcal{X}_i} R_n(\mathbf{x}) \geq \min_{\mathbf{x} \in \mathcal{X}_{i+1}} R_n(\mathbf{x})$.
- 2. The generalization error increases: $\sup_{\mathbf{x}\in\mathcal{X}_i}|R(\mathbf{x}) R_n(\mathbf{x})| \leq \sup_{\mathbf{x}\in\mathcal{X}_{i+1}}|R(\mathbf{x}) R_n(\mathbf{x})|.$
- 3. What happens with the true risk $R(\mathbf{x}_i^{\star})$?



Peeling the onion



Models

Let $d(\cdot, \cdot) : \mathcal{H}^{\circ} \times \mathcal{H}^{\circ} \to \mathbb{R}^+$ be a metric in an extended function space \mathcal{H}° that includes \mathcal{H} ; i.e., $\mathcal{H} \subseteq \mathcal{H}^{\circ}$. Let

- $1.\ h^\circ \in \mathcal{H}^\circ$ be the true, expected risk minimizing model
- 2. $h^{\natural} \in \mathcal{H}$ be the solution under the assumed function class $\mathcal{H} \subseteq \mathcal{H}^{\circ}$
- 3. $h^{\star} \in \mathcal{H}$ be the estimator solution
- 4. $h^t \in \mathcal{H}$ be the numerical approximation of the algorithm at time t

Practical performance

$$\underbrace{d(h^t, h^\circ)}_{\overline{e}(t, n)} \leq \underbrace{d(h^t, h^\star)}_{\text{optimization error}} + \underbrace{d(h^\star, h^\natural)}_{\text{statistical error}} + \underbrace{d(h^\natural, h^\circ)}_{\text{model error}}$$

where $\bar{\varepsilon}(t,n)$ denotes the total error of the Learning Machine. We can try to

- $1. \,$ reduce the optimization error with computation
- 2. reduce the statistical error with more data samples, with better estimators, and with prior information
- 3. reduce the model error with flexible or universal representations

The classical trade-off between model complexity and risk



Figure: Bias-variance trade-off [19].

Occam's Razor: Simple is better than complex.



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The dangers of complex function classes: sévère (cevher) overfitting



Figure: Training over a complex function class can lead to overfitting.



The dangers of complex function classes: sévère (cevher) overfitting



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Figure: Training over a complex function class can lead to overfitting.



Estimation of parameters vs estimation of risk



Nomenclature

$R_n(\cdot)$	training error
$R(\cdot)$	test error
$R(\mathbf{x}^{\natural}) - R(\mathbf{x}^{\circ})$	modeling error
$R(\mathbf{x}^{\star}) - R(\mathbf{x}^{\natural})$	excess risk
$\sup_{\mathbf{x}\in\mathcal{X}} R(\mathbf{x})-R_n(\mathbf{x}) $	generalization error
$R_n(\mathbf{x}^t) - R_n(\mathbf{x}^\star)$	optimization error

$\mathcal{X} \to \mathcal{X}^{\circ}$ $n\uparrow$ Training error Excess risk Generalization error Modeling error Time

 $R_n(h_{\mathbf{x}}) = \frac{1}{n} \sum_{i=1}^n L(h_{\mathbf{x}}(\mathbf{a}_i), b_i)$ be the empirical estimate. Let $\mathcal{X} \subseteq \mathcal{X}^\circ$ be parameter domains, where \mathcal{X} is known. Define

1. $\mathbf{x}^{\circ} \in \arg \min_{\mathbf{x} \in \mathcal{X}^{\circ}} R(h_{\mathbf{x}})$: true minimum risk model 2. $\mathbf{x}^{\natural} \in \operatorname{arg\,min}_{\mathbf{x} \in \mathcal{X}} R(h_{\mathbf{x}})$: assumed minimum risk model

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Recall the general setting

Let $R(h_{\mathbf{x}}) = \mathbb{E}L(h_{\mathbf{x}}(\mathbf{a}), b)$ be the risk function and

3. $\mathbf{x}^* \in \operatorname{arg\,min}_{\mathbf{x} \in \mathcal{X}} R_n(h_{\mathbf{x}})$: ERM solution 4. \mathbf{x}^t : numerical approximation of \mathbf{x}^* at time t

What theoretical challenges in Deep Learning will we study?



Models

- Let $\mathcal{X} \subseteq \mathcal{X}^\circ$ be parameter domains, where \mathcal{X} is known. Define
- 1. $\mathbf{x}^{\circ} \in \arg\min_{\mathbf{x} \in \mathcal{X}^{\circ}} R(h_{\mathbf{x}})$: true minimum risk model
- 2. $\mathbf{x}^{\natural} \in \operatorname{arg\,min}_{\mathbf{x} \in \mathcal{X}} R(h_{\mathbf{x}})$: assumed minimum risk model
- 3. $\mathbf{x}^{\star} \in \operatorname{arg\,min}_{\mathbf{x} \in \mathcal{X}} R_n(h_{\mathbf{x}})$: ERM solution
- 4. \mathbf{x}^t : numerical approximation of \mathbf{x}^* at time t

Practical performance in Deep Learning



where $\bar{e}(t, n)$ denotes the total error of the Learning Machine. In Deep Learning applications

- 1. Optimization error is almost zero, in spite of non-convexity. \Rightarrow lecture 10
- 2. Generalization error is usually small, but theory is lacking. \Rightarrow lecture 9 (this one)
- 3. Large architectures + inductive bias might lead to small model error.

Generalization error bounds and Rademacher Complexity

Goal: Obtain generalization bounds for multi-layer, fully-connected neural networks

 \circ We want to find high-probability upper bounds for the quantity

$$\sup_{\mathbf{x}\in\mathcal{X}}|R(\mathbf{x})-R_n(\mathbf{x})|$$

• Need a notion of *complexity* to derive generalization bounds for infinite classes of functions

Definition (Rademacher Complexity [10])

Let $A = \{\mathbf{a}_1, \dots, \mathbf{a}_n\} \subseteq \mathbb{R}^p$ and let $\{v_i : i = 1, \dots, n\}$ be independent Rademacher random variables i.e., taking values uniformly in $\{-1, +1\}$ (coin flip). Let \mathcal{H} be a class of functions of the form $h : \mathbb{R}^p \to \mathbb{R}$. The Rademacher complexity of \mathcal{H} with respect to A is defined as:

$$\mathcal{R}_A(\mathcal{H}) \coloneqq \mathbb{E}_v \sup_{h \in \mathcal{H}} \frac{1}{n} \sum_{i=1}^n v_i h(\mathbf{a}_i).$$

Remarks: $\circ \mathcal{R}_A(\mathcal{H})$ measures how well we fit random (±1) with the output of an element of \mathcal{H} on the set A. \circ The derivation of Rademacher Complexity for specific function classes are in the appendix.

Fundamental theorem about the Rademacher Complexity

Theorem (See Theorem 3.3 and 5.8 in [33])

Suppose that the loss function has the form $L(h_{\mathbf{x}}(\mathbf{a}), b) = \phi(b \cdot h_{\mathbf{x}}(\mathbf{a}))$ for a 1-Lipschitz function $\phi : \mathbb{R} \to \mathbb{R}$.

Let $\mathcal{H}_{\mathcal{X}} := \{h_{\mathbf{x}} : \mathbf{x} \in \mathcal{X}\}$ be a class of parametric functions $h_{\mathbf{x}} : \mathbb{R}^p \to \mathbb{R}$. For any $\delta > 0$, with probability at least $1 - \delta$ over the draw of an *i.i.d.* sample $\{(\mathbf{a}_i, b_i)\}_{i=1}^n$, letting $A = (\mathbf{a}_1, \ldots, \mathbf{a}_n)$, the following holds:

$$\sup_{\mathbf{x}\in\mathcal{X}} |R_n(\mathbf{x}) - R(\mathbf{x})| \le 2\mathbb{E}_A \mathcal{R}_A(\mathcal{H}_{\mathcal{X}}) + \sqrt{\frac{\ln(2/\delta)}{2n}},$$
$$\sup_{\mathbf{x}\in\mathcal{X}} |R_n(\mathbf{x}) - R(\mathbf{x})| \le 2\mathcal{R}_A(\mathcal{H}_{\mathcal{X}}) + 3\sqrt{\frac{\ln(4/\delta)}{2n}}.$$

Assumption is true for common losses

$$\blacktriangleright L(h_{\mathbf{x}}(\mathbf{a}), b) = \log(1 + \exp(-b \cdot h_{\mathbf{x}}(\mathbf{a}))) \Rightarrow \phi(z) := \log(1 + \exp(z)) \text{ (logistic loss)}$$

 $L(h_{\mathbf{x}}(\mathbf{a}), b) = \max(0, 1 - b \cdot h_{\mathbf{x}}(\mathbf{a})) \Rightarrow \phi(z) := \max(0, 1 - z) \text{ (hinge loss)}$

The complexity vs risk trade-off in practice (I)



Figure: Training (empirical) and test (true) error for one-hidden-layer networks of increasing width, trained with SGD [38].

Empirical error becomes zero for a wide enough network. What should happen for even wider networks?

The complexity vs risk trade-off in practice (II)



Figure: Training (empirical) and test (true) error for one-hidden-layer networks of increasing width, trained with SGD [38].

Test error continues to go down even if we keep incresing the complexity of the model!

How well do complexity measures correlate with generalization?

name	definition	$correlation^1$
Frobenius distance to initialization [35]	$\sum_{i=1}^d \ \mathbf{X}_i - \mathbf{X}_i^0\ _F^2$	-0.263
Spectral complexity ² [8]	$\prod_{i=1}^{d} \ \mathbf{X}_{i}\ \left(\sum_{i=1}^{d} rac{\ \mathbf{X}_{i}\ _{2,1}^{3/2}}{\ \mathbf{X}_{i}\ ^{3/2}} ight)^{2/3}$	-0.537
Parameter Frobenius norm	$\sum_{i=1}^d \ \mathbf{X}_i\ _F^2$	0.073
Fisher-Rao [29]	$\frac{(d+1)^2}{n} \sum_{i=1}^n \langle \mathbf{x}, \nabla_{\mathbf{x}} \ell(h_{\mathbf{x}}(\mathbf{a}_i), b_i) \rangle$	0.078
Path-norm [39]	$\sum_{(i_0,\ldots,i_d)}\prod_{j=1}^d \left(\mathbf{X}_{i_j,i_{j-1}} ight)^2$	0.373

Table: Complexity measures compared in the empirical study [27], and their correlation with generalization

Complexity measures are still far from explaining generalization in Deep Learning!

A more recent evaluation of many complexity measures is available [18].

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¹Kendall's rank correlation coefficient

²The definition in [27] differs slightly

The benefits of overparametrization

Overparameterization: #model parameters $\gg \#$ training data



Figure: Overparametrization leads to benign overfitting.



The generalization mystery in deep learning

UNDERSTANDING DEEP LEARNING REQUIRES RE-THINKING GENERALIZATION

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Abstract

Despite their massive size, successful deep artificial neural networks can exhibit a remarkably small difference between training and test performance. Conventional wisdom attributes small generalization error either to properties of the model family, or to the regularization techniques used during training.

Through extensive systematic experiments, we show how these traditional approaches fail to explain why large neural networks generalize well in practice. Specifically, our experiments establish that state-of-the-ard convolutional networks for image classification trained with stochastic gradient methods easily fit a random labeling of the training data. This phenomenon is qualitatively unaffected by explicit regularization, and occurs even if we replace the true images by completely unstructured random noise. We corroborate these experimental findings with a theoretical construction chuving that it may have the nature attracts of the structure of the stru

A gap between theory and practice

 \circ In practice, simple algorithms like SGD can train neural networks to zero error and achieve low test error.

 \circ This happens even for large and complex neural network architectures.

 Complexity measures like the Rademacher complexity suggest the opposite behaviour (overfitting)



Multiple global minimizers of the empirical risk



 \circ The global minimum is R_n^{\star}



Multiple global minimizers of the empirical risk



 \circ The global minimum is R_n^\star , but many parameters can attain such value.



Multiple global minimizers of the empirical risk



 \circ The global minimum is R_n^{\star} , but many parameters can attain such value.

 \circ Each minimizer of the empirical risk might have a different true risk.

*From multiple global minimizers to single global minimizer

 \circ Under some settings, the training objective of deep ReLU is almost convex and semi-smooth [2].³

o In such settings, the training behavior of NNs are close to training with kernel methods (see supp. material).

 \circ Define feature mapping $\mathbf{a} \mapsto \frac{\partial h}{\partial \mathbf{x}}(\mathbf{a}, \mathbf{x}_0)$, the (empirical) neural tangent kernel (NTK) [26] is defined as

 $K(\mathbf{a}_i, \mathbf{a}_j) := \langle \nabla_{\mathbf{x}} h(\mathbf{a}_i, \mathbf{x}), \nabla_{\mathbf{x}} h(\mathbf{a}_j, \mathbf{x}) \rangle, \forall i, j \in [n].$

 $^{^{3}}$ Classical smoothness only has a second-order term, but semi-smoothness also has an extra first-order term that is smaller for a larger width.



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Training dynamics [26]

Under the squared loss, the dynamics of $h(\mathbf{a}, \mathbf{x})$ is equivalent to kernel regression

$$\dot{h}(\mathbf{a}, \mathbf{x}(t)) = \nabla_{\mathbf{x}} h(\mathbf{a}, \mathbf{x}) \dot{\mathbf{x}}(t) = K_{\infty}(\mathbf{a}, \mathbf{a}_i) (h(\mathbf{a}, \mathbf{x}(t)) - b),$$

where, under proper initialization and large enough width, we have

$$K_{\infty} := \lim_{\text{width} \to \infty} K_{\mathbf{x}(0)}(\mathbf{a}_i, \mathbf{a}_j) = \mathbb{E}_{\mathbf{x}}[K_{\mathbf{x}(0)}(\mathbf{a}_i, \mathbf{a}_j)].$$

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where, under proper initialization and large enough width, we have

$$K_{\infty} := \lim_{\text{width}\to\infty} K_{\mathbf{x}(0)}(\mathbf{a}_i, \mathbf{a}_j) = \mathbb{E}_{\mathbf{x}}[K_{\mathbf{x}(0)}(\mathbf{a}_i, \mathbf{a}_j)].$$

Remarks: • NTK stays unchanged during training

- General loss functions: equivalence between infinite NNs and kernel methods [15]
 - ▶ e.g., NN trained by soft margin loss vs. SVM trained by subgradient descent

³Classical smoothness only has a second-order term, but semi-smoothness also has an extra first-order term that is smaller for a larger width.



*Optimization and generalization by NTK

Theorem (optimization and generalization [2, 13])

For a DNN with a large enough width trained by (S)GD on $\{(\mathbf{a}_i, b_i)\}_{i=1}^n$, under proper data assumptions and step-size η , we have

global convergence

 $L(\mathbf{x}(t)) \leq [1 - \eta \lambda_{\min}(K_{\infty})]^{t} L(\mathbf{x}(0))$ whp,

where $\lambda_{\min}(K_{\infty})$ is the minimum eigenvalue of K_{∞} .

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generalization guarantee

$$\mathcal{R}(h_{\mathbf{x}(t)}) \lesssim \mathcal{O}\left(\sqrt{\frac{\mathbf{b}^\top K_\infty^{-1} \mathbf{b}}{n}}\right) + \mathcal{O}\left(\frac{1}{\sqrt{n}}\right) \textit{whp}.$$

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Remarks: • The minimum eigenvalue of NTK plays an important role!

- robustness: generate adversarial examples [49]
- image denoising [48]
- neural architecture search in a "train-free" fashion [53, 14]
- \circ Under proper assumptions, we have $\lambda_{\min}(K_{\infty})=\Omega(p)$ [41] for the input dimension p

Not all global minimizers are the same

Consider a simple 2D classification task, and train a neural network with fixed step-size SGD.
The plots below correspond to two different global minimizers:



SGD almost never lands on the global minimum on the right! Why?

Understanding the implicit bias of optimization algorithms

• SGD seems to be *biased* towards good global minimizers (low true risk).

• Some optimization algorithms have an implicit bias towards certain kinds of global minimizers.

• Can we characterize this implicit bias?



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- Some optimization algorithms have an implicit bias towards certain kinds of global minimizers.
- Can we characterize this implicit bias?

Definition (Algorithm)

We will refer to a function (deterministic or randomized) $\mathscr{A} : \mathcal{Z} \to \mathcal{X}$, mapping $Z \mapsto \mathscr{A}_Z$ as an *algorithm* with *input* $Z \in \mathcal{Z}$ and *output* $\mathscr{A}_Z \in \mathcal{X}$.

Example: Gradient Descent Algorithm

We denote $GD_{(T,\alpha,\mathbf{x}^0,\nabla f)} := T$ -steps of GD with stepsize α , starting from \mathbf{x}^0 , using gradient ∇f .



What is implicit regularization?

Definition (Implicit Regularization of a Deterministic Algorithm)

Consider a minimization problem

$$F^{\star} = \min_{\mathbf{x} \in \mathcal{X}} F(\mathbf{x}) \tag{2}$$

and let \mathscr{A} be a deterministic algorithm with input $Z \in \mathcal{Z}$ and output $\mathscr{A}_Z \in \mathcal{X}$.

We say that \mathscr{A} solves problem (2) and has *implicit regularization* $H: \mathcal{X} \times \mathcal{Z} \to \mathbb{R}$ if

 $\mathscr{A}_Z \in \underset{F(\mathbf{x})=F^{\star}}{\operatorname{arg\,min}} H(\mathbf{x}, Z).$

Given the input $Z \in \mathcal{Z}$, the algorithm outputs a global minimizer of F that, additionally, minimizes $H(\cdot, Z)$.



Implicit bias of gradient descent for linear regression

o Consider for example an underdetermined linear system

$$\mathbf{A}\mathbf{x} = \mathbf{b}, \quad \text{with } \mathbf{A} \in \mathbb{R}^{n \times p}, \quad n < p$$

• If a solution exists (i.e., $\mathbf{b} \in \mathbf{colspan}(\mathbf{A})$), then there is an *infinite number of solutions* to this system.

Finding a solution

To find a valid \mathbf{x} , we could apply one of the optimization algorithms seen in class to the convex problem

$$\operatorname*{arg\,min}_{\mathbf{x}\in\mathbb{R}^p}\frac{1}{2}\|\mathbf{A}\mathbf{x}-\mathbf{b}\|_2^2$$

Among all the possible solutions, which one will the algorithm converge to ?

Same problem and same initialization vs different algorithms and different solutions

Consider the following simple 2D example :

$$\begin{bmatrix} 1 & 2 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \end{bmatrix} = 5$$



Different Solutions

Gradient Descent and AdaGrad converge to *different* points on the line.



Implicit bias of gradient descent for linear regression

 \circ Gradient descent seems to converge to the closest one in terms of $\ell_2\text{-norm.}$

Theorem (Implicit bias of Gradient Descent [20])

For the underdetermined, realizable linear system

$$F^{\star} = \min_{\mathbf{x} \in \mathcal{X}} F(\mathbf{x}) = \frac{1}{2} \|\mathbf{A}\mathbf{x} - \mathbf{b}\|_{2}^{2}$$

the gradient descent algorithm $GD_{(T,\alpha,\mathbf{x}^0,\nabla F)}$, for $T = \infty$ and for any $\mathbf{x}^0 \in \mathbb{R}^p$, and valid step-size α , has implicit bias $H(\mathbf{x}) = \|\mathbf{x} - \mathbf{x}^0\|_2$, i.e.,

$$GD_{(T=\infty,\alpha,\mathbf{x}^0,\nabla F)} = \operatorname*{arg\,min}_{F(\mathbf{x})=F^{\star}} \|\mathbf{x}-\mathbf{x}_0\|_2.$$

Remark: • The theorem also holds for stochastic gradient descent, see [3].

Same problem and same initialization vs different algorithms and different solutions



Proof : For simplicity, take $\mathbf{x}_0 = 0$.

- The gradient of F is $\mathbf{A}^T (\mathbf{A}\mathbf{x} \mathbf{b})$.
- This implies that $\forall \mathbf{x}, \nabla f(\mathbf{x}) \in \mathbf{colspan}(\mathbf{A}^T)$.

GD iterates stay in the rowspan

Gradient Descent is therefore constrained to the space

 $\mathbf{colspan}(\mathbf{A}^T) = \mathbf{rowspan}(\mathbf{A})$

So its limit point at $T = \infty$ is in rowspan(A).

Note that because of the preconditionning, AdaGrad can get out of the rowspan(A).

Same problem and same initialization vs different algorithms and different solutions



Proof (continued):

The minimum norm solution

$$\hat{\mathbf{x}}_{\mathsf{candidate}} = \mathop{\arg\min}_{\mathbf{x}:\mathbf{Ax}=\mathbf{b}} \|\mathbf{x}\|_2^2$$

is also in $\mathbf{rowspan}(\mathbf{A})$.

- So both $\hat{\mathbf{x}}_{candidate}$ and the limit point of GD are solutions of $\mathbf{A}\mathbf{x} = \mathbf{b}$ that are in the $\mathbf{rowspan}(\mathbf{A})$
- ► Since nullA ∩ rowspan(A) = {0}, there can only be one solution in the rowspan(A), so

$$\mathbf{x}_{\mathsf{GD}}^{\star} = \hat{\mathbf{x}}_{\mathsf{candidate}}$$
Implicit bias for linear models

 \circ We can extend this analysis to linear models:

$$\operatorname*{arg\,min}_{\mathbf{x}\in\mathbb{R}^p} F(\mathbf{x}) := \sum_{i=1}^n L(\langle \mathbf{x}, \mathbf{a}_i \rangle, b_i).$$

• If the observations are realizable and there are many global minima $\mathbf{Glob} = \{\mathbf{x} : F(\mathbf{x}) = 0\}$, then

Theorem (Implicit Bias of Gradient Descent [20])

If the loss L is convex and has a unique (attained) minimum, then the iterates \mathbf{x}^t of Gradient Descent converge to the global minimum that is closest to initialization \mathbf{x}_0 in the ℓ_2 -distance :

$$\mathbf{x}^t \xrightarrow[t \to \infty]{} \arg\min_{\mathbf{x} \in \mathbf{Glob}} \|\mathbf{x} - \mathbf{x}_0\|_2$$

Proof: (Sketch) The assumption on *L* implies the problem reduces to a linear system: If x is a global minimum, we must have $\langle \mathbf{x}, \mathbf{a}_i \rangle = b_i$ for all $i \in \{1, ..., n\}$. We can recycle the results we have just seen. **Remarks:** \circ Implicit bias for wide two-layer neural networks [16] can be found in supplementary material.

The double descent phenomenon

• A failure of conventional wisdom



Figure: The classical U-shaped risk curve vs. double-descent risk curve. source: [11].

- \blacktriangleright classical large-sample limit setting: $n \rightarrow \infty$ under fixed p
- \blacktriangleright high dimensional setting: n and p comparably large

Double descent curve in practice (I)

- Typical examples:
 - linear/nonlinear regression [24]
 - ▶ random features, random forest, and shallow neural networks [11]



Figure: Experiments on MNIST. Source: [11].

Double descent curve in practice (II)



Figure: Left: Train and test error as a function of model size, for ResNet18s of varying width on CIFAR-10 with 15% label noise. Right: Test error, shown for varying train epochs. source: [36].

Double descent curve in practice (III)



Figure: Left: The double descent phenomenon, where the number of parameters is used as the model complexity. Middle: The norm of the learned model is peaked around $n \approx p$. Right: The test error against the norm of the learnt model. The color bar indicate the number of parameters and the arrows indicates the direction of increasing model size. Their relationship are closer to the convention wisdom than to a double descent. source: [40]. This is the same setting as in Section 5.2 of [37].

Underparametrized regime





Figure: Low generalization but high empirical error

Figure: Sweet spot for the model complexity



Interpolation threshold



Figure: The unique degree 19 polynomial that can fit 20 samples.



Benign overfitting in the over-parametrized regime



Figure: A degree 200 polynomial that can harmlessly fits noisy 20 points.



Figure: Double descent for polynomial fits

Benign Overfitting [9]: good prediction with zero training error

- Statistical wisdom: a predictor should not fit too well.
- deep networks fit perfectly on noisy data and generalize well on test data.



Double descent in 1998: AdaBoost

Definition (Informal [45])

"Boosting solves hard machine learning problems by forming a very smart committee of grossly incompetent but carefully selected members."

AdaBoost

- 1. Initialize the observation weights $w_i = 1/N$, i = 1, 2, ..., N
- 2. For t = 1 to T:
 - 2.1 Fit a classifier $h_{\mathbf{x},t}(\mathbf{a})$ to the training data using weights w_i .
 - 2.2 Compute

$$\mathsf{err}_t = \frac{\sum_{i=1}^N w_i I(\mathbf{b}_i \neq h_{\mathbf{x},t}(\mathbf{a}_i))}{\sum_{i=1}^N w_i}$$

2.3 Compute
$$\alpha_t = \log((1 - \operatorname{err}_t)/\operatorname{err}_t)$$
.
2.4 Set $w_i \leftarrow w_i \cdot \exp[\alpha_t \cdot I(\mathbf{b}_i \neq h_{\mathbf{x},t}(\mathbf{a}_i))], i = 1, 2, \dots, N$.

3. Output
$$h(\mathbf{a}) = \left[\sum_{t=1}^{T} \alpha_t h_{\mathbf{x},t}(\mathbf{a})\right]$$
.

Remarks:

 $\,\circ\,$ At each round, the weights are updated so the weak learner focuses on the hard examples.

 \circ The more iterations are run, the more complex the output function becomes (e.g., overfitting).



AdaBoost with large number of rounds





Figure: AdaBoost on letters dataset [7]. Test error keeps improving even after 0 training error is reached.

Margin theory [7]

The *margin* is a measure of confidence in the prediction. Boosting can be shown to increase the margin at each round.

Wrap up!

 \circ The visualizations can be deceiving to understand the high-dimensional behavior

 \circ Are we really in the interpolation regime in machine learning?

Theorem (Probability of interpolation [6])

Given a p-dimensional dataset $\mathcal{A}_n = \{\mathbf{a}_1, \dots, \mathbf{a}_n\}$ with i.i.d. samples, where $\mathbf{a}_i \sim \mathcal{N}(0, I)$ for all $i = 1, \dots, n$, the probability that a new sample $\mathbf{a} \sim \mathcal{N}(0, I)$ is in the interpolation regime (i.e., within the convex hull of \mathcal{A}_n) has the following limiting behavior

$$\lim_{p \to \infty} p(\mathbf{a} \in \textit{ConvexHull}(\mathcal{A}_n)) = \begin{cases} 1 & \text{if } n > 2^{p/2}/p; \\ 0 & \text{if } n < 2^{p/2}/p. \end{cases}$$

 \circ We are most likely in the extrapolation regime [5]

*Concentration inequality

o Main tool for generalization bound: concentration inequalities!

Measure of how far is an empirical average from the true mean

Theorem (Hoeffding's Inequality [33])

Let Y_1, \ldots, Y_n be i.i.d. random variables with Y_i taking values in the interval $[a_i, b_i] \subseteq \mathbb{R}$ for all $i = 1, \ldots, n$. Let $S_n := \frac{1}{n} \sum_{i=1}^n Y_i$. It holds that

$$\mathbb{P}(|S_n - E[S_n]| > t) \le 2 \exp\left(-\frac{2n^2t^2}{\sum_{i=1}^n (b_i - a_i)^2}\right)$$

*Generalization bound for a singleton

Lemma

For i = 1, ..., n, let $(\mathbf{a}_i, b_i) \in \mathbb{R}^p \times \{-1, 1\}$ be independent random variables and $h_{\mathbf{x}} : \mathbb{R}^p \to \mathbb{R}$ be a function parametrized by $\mathbf{x} \in \mathcal{X}$. Let $\mathcal{X} = \{\mathbf{x}_0\}$ and $L(h_{\mathbf{x}}(\mathbf{a}), b) = \{sign(h_{\mathbf{x}}(\mathbf{a})) \neq b\}$ be the 0-1 loss. With probability at least $1 - \delta$, we have that

$$\sup_{\mathbf{x}\in\mathcal{X}} |R(\mathbf{x}) - R_n(\mathbf{x})| = |R(\mathbf{x}_0) - R_n(\mathbf{x}_0)| \le \sqrt{\frac{\ln(2/\delta)}{2n}}.$$

Proof.

lions@epf

Note that $E[\frac{1}{n}\sum_{i=1}^{n}L(h_{\mathbf{x}_{0}}(\mathbf{a}_{i}),b_{i})] = R(\mathbf{x}_{0})$, the expected risk of the parameter \mathbf{x}_{0} . Moreover $L(h_{\mathbf{x}_{0}}(\mathbf{a}_{i}),b_{i}) \in [0,1]$. We can use Hoeffding's inequality and obtain

$$\mathbb{P}(|R_n(\mathbf{x}_0) - R(\mathbf{x}_0)| > t) = \mathbb{P}\left(\left| \frac{1}{n} \sum_{i=1}^n L_i(h_{\mathbf{x}_0}(\mathbf{a}_i), b_i) - R(\mathbf{x}_0) \right| > t \right) \le 2 \exp\left(-2nt^2 \right)$$

Setting $\delta:=2\exp\left(-2nt^2\right)$ we have that $t=\sqrt{rac{\ln 2/\delta}{2n}}$, thus obtaining the result.

Mathematics of Data | Prof. Volkan Cevher, volkan.cevher@epfl.ch Sli

*Generalization bound for finite sets

Lemma

For i = 1, ..., n, let $(\mathbf{a}_i, b_i) \in \mathbb{R}^p \times \{-1, 1\}$ be independent random variables and $h_{\mathbf{x}} : \mathbb{R}^p \to \mathbb{R}$ be a function parametrized by $\mathbf{x} \in \mathcal{X}$. Let \mathcal{X} be a finite set and $L(h_{\mathbf{x}}(\mathbf{a}), b) = \{sign(h_{\mathbf{x}}(\mathbf{a})) \neq b\}$ be the 0-1 loss. With probability at least $1 - \delta$, we have that

$$\sup_{\mathbf{x}\in\mathcal{X}}|R(\mathbf{x})-R_n(\mathbf{x})| \leq \sqrt{\frac{\ln|\mathcal{X}|+\ln(2/\delta)}{2n}}.$$

Proof.

lions@epf

Let $\mathcal{X} = \{\mathbf{x}_1, \dots, \mathbf{x}_{|\mathcal{X}|}\}$. We can use a union bound and the analysis of the singleton case to obtain:

$$\mathbb{P}(\exists j: |R_n(\mathbf{x}_j) - R(\mathbf{x}_j)| > t) \le \sum_{j=1}^{|\mathcal{X}|} \mathbb{P}(|R_n(\mathbf{x}_j) - R(\mathbf{x}_j)| > t) = 2|\mathcal{X}| \exp\left(-2nt^2\right)$$

Setting $\delta := 2|\mathcal{X}| \exp\left(-2nt^2\right)$, we have that $t = \sqrt{\frac{\ln|\mathcal{X}| + \ln \frac{2}{\delta}}{2n}}$, thus obtaining the result.

*Visualizing Rademacher complexity

X X X X X X X X X -1 -1 -1 -1 -1 -1 -1 -1

Figure: Rademacher complexity measures correlation with random signs



*Visualizing Rademacher complexity







*Computing the Rademacher complexity of linear functions

Theorem

Let $\mathcal{X} := \{\mathbf{x} \in \mathbb{R}^p : \|\mathbf{x}\|_2 \leq \lambda\}$ and let $\mathcal{H}_{\mathcal{X}}$ be the class of functions of the form $h_{\mathbf{x}} : \mathbb{R}^p \to \mathbb{R}, h_{\mathbf{x}}(\mathbf{a}) = \langle \mathbf{x}, \mathbf{a} \rangle$, for some $\mathbf{x} \in \mathcal{X}$. Let $A = {\mathbf{a}_1, \dots, \mathbf{a}_n} \subseteq \mathbb{R}^p$ such that $\max_{i=1,\dots,n} \|\mathbf{a}_i\| \leq M$. It holds that $\mathcal{R}_A(\mathcal{H}_{\mathcal{X}}) \leq \lambda M / \sqrt{n}.$

Proof.

$$\begin{aligned} \mathcal{R}_{A}(\mathcal{H}_{\mathcal{X}}) &= \mathbf{E} \sup_{\|\mathbf{x}\|_{2} \leq \lambda} \frac{1}{n} \sum_{i=1}^{n} v_{i} \langle \mathbf{x}, \mathbf{a} \rangle \\ &= \mathbf{E} \sup_{\|\mathbf{x}\|_{2} \leq \lambda} \frac{1}{n} \left\langle \mathbf{x}, \sum_{i=1}^{n} v_{i} \mathbf{a} \right\rangle \\ &\leq \frac{1}{n} \lambda \mathbf{E} \left\| \sum_{i=1}^{n} v_{i} \mathbf{a}_{i} \right\|_{2} \end{aligned} \qquad \Rightarrow \mathcal{R}_{A}(\mathcal{H}_{\mathcal{X}}) \leq \frac{1}{n} \lambda \left(\mathbf{E} \sum_{i=1}^{n} \|v_{i} \mathbf{a}_{i}\|_{2}^{2} \right)^{1/2}$$
(Jensen)
$$&\leq \frac{1}{n} \lambda \left(\sum_{i=1}^{n} \|u_{i}\|_{2}^{2} \right)^{1/2} \\ &\leq \lambda M / \sqrt{n} \end{aligned}$$

*Rademacher complexity estimates of fully connected Neural Networks

Notation

For a matrix $\mathbf{X} \in \mathbb{R}^{n,m}$, $\|\mathbf{X}\|$ denotes its spectral norm. Let $\mathbf{X}_{:,k}$ be the k-th column of \mathbf{X} . We define

$$\|\mathbf{X}\|_{2,1} = \|(\|\mathbf{X}_{:,1}\|_{2}, \dots, \|\mathbf{X}_{:,m}\|_{2})\|_{1}.$$
(3)

Theorem (Spectral bound [8])

For positive integers $p_0, p_1, \ldots, p_d = 1$, and positive reals $\lambda_1, \ldots, \lambda_d$ and ν_1, \ldots, ν_d , define the set

$$\mathcal{X} := \{ (\mathbf{X}_1, \dots, \mathbf{X}_d) : \mathbf{X}_i \in \mathbb{R}^{p_i \times p_{i-1}}, \|\mathbf{X}_i\| \le \lambda_i, \|\mathbf{X}_i^T\|_{2,1} \le \nu_i \}$$

Let $H_{\mathcal{X}}$ be the class of neural networks $h_{\mathbf{x}} : \mathbb{R}^p \to \mathbb{R}$, $h_{\mathbf{x}} = \mathbf{X}_d \circ \sigma \circ \ldots \circ \sigma \circ \mathbf{X}_1$ where $\mathbf{x} = (\mathbf{X}_1, \ldots, \mathbf{X}_d) \in \mathcal{X}$. Suppose that σ is 1-Lipschitz. Let $A = \{\mathbf{a}_1, \ldots, \mathbf{a}_n\} \subseteq \mathbb{R}^p$, $M := \max_{i=1,\ldots,n} \|\mathbf{a}_i\|$ and $W := \max\{p_i : i = 0, \ldots, d\}$.

The Rademacher complexity of $\mathcal{H}_{\mathcal{X}}$ with respect to A is bounded as

$$\mathcal{R}_A(\mathcal{H}_{\mathcal{X}}) = \mathcal{O}\left(\frac{\log(W)M}{\sqrt{n}} \prod_{i=1}^d \lambda_i \left(\sum_{j=1}^d \frac{\nu_j^{2/3}}{\lambda_j^{2/3}}\right)^{3/2}\right).$$
(4)

*Implicit bias for linearly separable datasets

For linearly separable datasets, we know of an algorithm capable of finding a separating hyperplane.
It maximizes the *margin* (i.e., distance between the boundary and the nearest training-data point).



Hard-margin Support Vector Machines

The hard margin Support Vector Machine solves the following optimization problem :

 $\underset{\mathbf{x}\in\mathbb{R}^{p}}{\arg\min}\|\mathbf{x}\|_{2} \quad \text{ subject to } y_{i}\langle\mathbf{x},\mathbf{a}_{i}\rangle\geq 1.$

It finds a hyperplane that maximizes the margin. It does so *by design*.

*Implicit bias for linearly separable datasets

 \circ What happens if we do not explicitly enforce margin maximization?

Theorem (Implicit Bias of Gradient Descent on Separable Data [47, 20])

For the logistic loss (and some other strictly monotonically decreasing losses) and for linearly separable datasets, the direction of the iterates x^t of Gradient Descent for any initialization converges to the hard-margin SVM direction:

$$\frac{\mathbf{x}^{t}}{\|\mathbf{x}^{t}\|_{2}} \xrightarrow[t \to \infty]{} \frac{\mathbf{x}_{SVM}^{\star}}{\|\mathbf{x}_{SVM}^{\star}\|_{2}} \quad \textit{where } \mathbf{x}_{SVM}^{\star} = \left\{ \underset{\mathbf{x} \in \mathbb{R}^{p}}{\arg\min} \|\mathbf{x}\|_{2} \quad \textit{subject to } y_{i} \langle \mathbf{x}, \mathbf{a}_{i} \rangle \geq 1 \right\}$$

Remarks: • Here, without explicit instructions, gradient descent maximizes the margin.

• The rate of this convergence is $O\left(\frac{1}{\log t}\right)$.

*Implicit bias for linearly separable datasets

• A similar result can be established for stochastic gradient descent for the logistic loss on separable datasets.

Theorem (Implicit Bias of *Stochastic* Gradient Descent on Separable Data [34])

The direction of the iterates \mathbf{x}^t of Stochastic Gradient Descent for any initialization and for a small enough fixed step-size, converges almost surely to the hard-margin SVM direction:

$$\left\| \frac{\mathbf{x}^t}{\|\mathbf{x}^t\|_2} - \frac{\mathbf{x}^{\star}_{SVM}}{\|\mathbf{x}^{\star}_{SVM}\|_2} \right\|_2 = O\left(\frac{1}{\log t}\right)$$

 Remarks:
 • This result is particularly interesting as it establishes convergence of fixed step-size SGD.

 • Both SGD and GD have the same implicit bias towards maximizing margins.

*Implicit bias for non-convex objectives

• Characterizing implicit bias of stochastic gradient descent for non-convex objectives is an active research area.

• Some papers study deep matrix factorization as a first step towards getting results for neural networks.

Deep Matrix Factorization

Deep matrix factorization consists of parametrizing a matrix \mathbf{M} as a product of N matrices:

$$\mathbf{M} = \mathbf{X}_N \mathbf{X}_{N-1} \dots \mathbf{X}_1$$

which can be understood as parametrizing M by a depth N "linear neural network," i.e., a neural network with no activations and with weight matrices \mathbf{X} .

*Implicit bias for deep matrix completion

• The matrix completion problem consists of filling the missing entries of a partially observed matrix.

• The deep matrix factorization approach consists of solving the following problem with gradient descent:

$$\operatorname*{arg\,min}_{\mathbf{X}_N,\mathbf{X}_{N-1}\ldots,\mathbf{X}_1} \sum_{(i,j)\in\Omega} ([\mathbf{X}_N\mathbf{X}_{N-1}\ldots\mathbf{X}_1]_{i,j} - b_{i,j})^2.$$

o It was conjectured in 2017 [21] that gradient descent was biased towards solutions with small nuclear norm.

Theorem (Implicit Regularization May Not Be Explainable by Norms (2020) [44])

For deep matrix completion the implicit bias can not be expressed as a function of a norm or semi-norm.



*Implicit bias for wide two-layer neural networks

• Assume a wide two-layer neural network $h_{\mathbf{x}}(\mathbf{a}) = \frac{1}{m} \sum_{i=1}^{m} \sigma(\langle \mathbf{x}_i, \mathbf{a} \rangle)$, where m is the width

 \circ An integral representation parameterized with a probability measure ν is given by

$$h_
u(\mathbf{a}) = \int_{\mathbb{R}^p} \sigma(\langle \mathbf{x}, \mathbf{a}
angle) \mathrm{d}
u(\mathbf{x}) \, .$$

Theorem (Implicit bias of gradient flow on two-layer neural networks [16])

Under proper initialization and technical conditions (in particular, of convergence), the output of the gradient flow h_{ν_t} under a proper normalization scheme converges to a certain max-margin classifier.

Remarks: • Gradient flow is the continuous limit of gradient descent [46].

- Fixing the hidden layer (i.e., random features) leads to a max-margin classifier in RKHS [16].
- \circ Other extensions of implicit bias of SGD depend on different models or settings:
 - overparameterized least squares [51], diagonal linear networks [42], stochastic differential equations [28].
 - multi-pass SGD [54], diferent noise types [12, 22], different momentum types [50].

*Implicit bias for wide two-layer neural networks

• Assume that we have a wide two-layer neural network $h_{\mathbf{x}}(\mathbf{a}) = \frac{1}{m} \sum_{i=1}^{m} \sigma(\langle \mathbf{x}_i, \mathbf{a} \rangle)$ • An integral representation parameterized with a probability measure ν

$$h_{
u}(\mathbf{a}_i) = \int_{\mathbb{R}^p} \sigma(\langle \mathbf{x}, \mathbf{a}_i
angle) \mathrm{d}
u(\mathbf{x}) \, ,$$

 $\circ \ \nu \in \mathcal{P}_2(\mathbb{R}^{d+2}) \text{ in the set of probability measures with finite second moment}$ $\circ \text{ the variation norm: } \|h\|_{\mathcal{F}_1} = \min_{\nu \in \mathcal{P}_2(\mathbb{R}^{d+2})} \left\{ \frac{1}{2} \int \|\mathbf{x}\|_2^2 \, \mathrm{d}\nu(\mathbf{x}); \quad h_\nu(\mathbf{a}_i) = \int \sigma(\langle \mathbf{x}, \mathbf{a}_i \rangle) \mathrm{d}\nu(\mathbf{x}) \right\}$

Theorem (Implicit Bias of wide two-layer Neural Networks [16])

Assume that $\nu_0 = \mathcal{U}_{\mathbb{S}^d} \otimes \mathcal{U}_{\{-1,1\}}$, the training set is consistent $([\mathbf{a}_i = \mathbf{a}_j] \Longrightarrow [b_i = b_j])$ and technical conditions (in particular, of convergence). Then $h_{\nu_t} / \|h_{\nu_t}\|_{\mathcal{F}_1}$ trained by an exponential tail loss converges to the \mathcal{F}_1 -max-margin classifier, i.e. it solves

$$\max_{\|h\|_{\mathcal{F}_1} \le 1} \min_{i \in [n]} b_i h(\mathbf{a}_i),$$

• Gradient flow is the continuous limit of gradient descent [46].

• Fixing the hidden layer (i.e., random features) leads to a max-margin classifier in RKHS [16].



*Example: Benign overfitting of DNNs on binary classification [52]

Problem setting: linear signal with label noise

- \blacktriangleright clean data distribution $(\tilde{\mathbf{a}}, \tilde{b}) \sim \tilde{\rho}$ $\circ \tilde{b} \sim \{\pm 1, -1\}, \tilde{\mathbf{a}} = \mathbf{z} + \tilde{b}\mu$ $\circ \mu$ -separated, 1-subgaussian, log-concave distributions in \mathbb{R}^d
- under a noise rate η , marginal distribution is the same: $\rho_A = \tilde{\rho}_A$ over A with $d_{\text{TV}}(\rho, \tilde{\rho}) \leq \eta$
- ▶ labels are flipped with probability $\eta(\mathbf{a})$: $\Pr[b(\mathbf{a}) = \tilde{b}] = 1 \eta$ and $\Pr[b(\mathbf{a}) \neq \tilde{b}] = \eta$
- DNNs with ReLU trained by gradient descent under the logistic loss

Theorem (Binary classification)

Under the above setting and assumptions, after t steps, DNNs can obtain the Bayes-optimal test error

$$\mathbb{P}_{(\mathbf{a},b)\sim\rho}(b\neq \operatorname{sgn}(h(\mathbf{a};\mathbf{X}^{(t)}))) \leq \eta + \exp\left(-\lambda\Theta\left(\frac{t\alpha(1-2\eta)}{\operatorname{Lip}_{h(\mathbf{a};\mathbf{X}^{(t)})}}\right)^2\right), w.h.p,$$

where α is the step size and η is the label flip rate.

Remarks: o smaller Lipchitz constant, faster convergence rate Lipschitz constant used for generalization NTK initialization: lazy training regime



*From neural networks to random features model [25, 43]

1-hidden-layer neural network with m neurons (fully-connected architecture):

Let $\mathbf{X}_1 \in \mathbb{R}^{m imes p}$, $\mathbf{a} \in \mathbb{R}^p$, $\mathbf{X}_2 \in \mathbb{R}^m$, and $\mu_2 \in \mathbb{R}$



- X₁: Gaussian initialization and then fixed
- ► X₂: to be learned
- over-parameterized model: #neurons m > #training data n

*Double descent: random features model (I)

 \circ high dimensions: #training data n, #neurons m, feature dimension p are comparably large



Figure: Test MSE, Bias, and Variance of RF regression as a function of the ratio m/n on MNIST data set (digit 3 vs. 7) for p = 784 and n = 600 across the Gaussian kernel. Source: [31].

- ▶ random features regression solved by SGD: interplay between excess risk and optimization
- bias variance decomposition for understanding multiple randomness sources
- monotonic decreasing bias and unimodal variance \Rightarrow double descent

*Double descent: random features model (II)

Algorithm	data assumption	solution type	Result on risk curve
[24]	Gaussian	closed-form	variance 🦯 📐
[32]	i.i.d on sphere	closed-form	variance, bias 🥕 🔪
[17]	Gaussian	closed-form	refined decomposition on variance
[1]	Gaussian	closed-form	fully decomposition on variance
[30]	general	closed-form	\nearrow
[4]	Gaussian	GD	variance 🦯 📐
[31]	sub-Gaussian	SGD	variance 🦯 📐, bias 📐

Table: Comparison of representative random features on double descent.

 $\circ\,$ multiple randomness sources: data sampling, label noise, initialization

 $\circ\,$ phase transition due to non-monotonic variance

*Peeling the onion (risk minimization setting) - Decomposition details

$$R(\mathbf{x}^{t}) - R(\mathbf{x}^{\natural}) = R(\mathbf{x}^{t}) - R_{n}(\mathbf{x}^{t}) + R_{n}(\mathbf{x}^{t}) - R_{n}(\mathbf{x}^{\star}) + \underbrace{R_{n}(\mathbf{x}^{\star}) - R_{n}(\mathbf{x}^{\natural})}_{\leq 0} + R_{n}(\mathbf{x}^{\natural}) - R(\mathbf{x}^{\natural})$$
$$\leq R_{n}(\mathbf{x}^{t}) - R_{n}(\mathbf{x}^{\star}) + \underbrace{R(\mathbf{x}^{t}) - R_{n}(\mathbf{x}^{t}) + R_{n}(\mathbf{x}^{\natural}) - R(\mathbf{x}^{\natural})}_{2 \sup_{\mathbf{x} \in \mathcal{X}} |R_{n}(\mathbf{x}) - R(\mathbf{x})|}$$

$$R(\mathbf{x}^{t}) - R(\mathbf{x}^{\circ}) = R(\mathbf{x}^{t}) - R(\mathbf{x}^{\natural}) + R(\mathbf{x}^{\natural}) - R(\mathbf{x}^{\circ})$$

$$\leq R_{n}(\mathbf{x}^{t}) - R_{n}(\mathbf{x}^{\star}) + 2 \sup_{\mathbf{x} \in \mathcal{X}} |R_{n}(\mathbf{x}) - R(\mathbf{x})| + R(\mathbf{x}^{\natural}) - R(\mathbf{x}^{\circ})$$



*Generalization bounds based on uniform stability — definitions

Definition (Empirical Risk on a set)

Let $S := [(\mathbf{a}_1, b_1), \dots, (\mathbf{a}_n, b_n)]$ be an i.i.d. sample drawn from a distribution on $\mathcal{A} \times \mathcal{B}$. Let $L : \mathcal{B} \times \mathcal{B} \to \mathbb{R}$ be a loss function and \mathcal{H} be a class of functions $h : \mathcal{A} \to \mathcal{B}$. The empirical risk of $h \in \mathcal{H}$ on the set S is defined as:

$$R_S(h) := \frac{1}{n} \sum_{i=1}^n L(h(\mathbf{a}_i), b_i)$$

(Almost) same definition as before. Makes explicit the dependence on the set S.

Definition (Expected Generalization Error)

Let $\mathscr{A} : \mathscr{Z} \to \mathcal{H}$ be a randomized algorithm that takes as input a finite sample S of arbitrary size, and outputs a function $\mathscr{A}_S \in \mathcal{H}$. Suppose that $S = [(\mathbf{a}_1, b_1), \dots, (\mathbf{a}_n, b_n)]$ is an i.i.d. sample form probability distribution on $\mathcal{A} \times \mathcal{B}$. The expected generalization error on a sample of size n is the value

$$\mathbb{E}[R_S(\mathscr{A}_S) - R(\mathscr{A}_S))]$$

the expectation is taken with respect to the draw of the sample S and the randomness of $\mathscr{A}.$



*Generalization bounds based on uniform stability — Fundamental theorem (I)

Theorem (Hardt et al. 2016 [23])

Let A be uniformly stable with stability $(\beta_n)_{n\geq 1}$, then for a random i.i.d. sample S of size n, the expected generalization error is bounded as follows

$$\mathbb{E}[|R_S(\mathscr{A}_S) - R(\mathscr{A}_S))|] \le \beta_n$$

Proof.

Let $S = [(\mathbf{a}_1, b_1), \dots, (\mathbf{a}_n, b_n)]$ and $S' = [(\mathbf{a}'_1, b'_1), \dots, (\mathbf{a}'_n, b_n)]$ be two i.i.d. samples of size n. Denote

$$S^{(i)} := [(\mathbf{a}_1, b_1), \dots, (\mathbf{a}_{i-1}, b_{i-1}), (\mathbf{a}'_i, b'_i), (\mathbf{a}_{i+1}, b_{i+1}), \dots, (\mathbf{a}_n, b_n)]$$

the sample that results from replacing (\mathbf{a}_i, b_i) by (\mathbf{a}'_i, b'_i) in S.

$$\mathbb{E}[R_S(\mathscr{A}_S)] = \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_S(\mathbf{a}_i), b_i)\right] = \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_{S(i)}(\mathbf{a}'_i), b'_i)\right]$$
$$= \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_{S(i)}(\mathbf{a}'_i), b'_i) - \frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_S(\mathbf{a}'_i), b'_i)\right] + \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_S(\mathbf{a}'_i), b'_i)\right]$$



*Generalization bounds based on uniform stability — Fundamental theorem (II)

Proof. (continued).

We have

$$\mathbb{E}[R_S(\mathscr{A}_S)] = \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_{S(i)}(\mathbf{a}'_i), b'_i) - \frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_S(\mathbf{a}'_i), b'_i)\right] + \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^n L(\mathscr{A}_S(\mathbf{a}'_i), b'_i)\right]$$

Note that S and $S^{(i)}$ only differ in one sample: uniform stability allows bounding the first term as:

$$= \mathbb{E}\left[\frac{1}{n}\sum_{i=1}^{n}L(\mathscr{A}_{S(i)}(\mathbf{a}'_{i}), b'_{i}) - \frac{1}{n}\sum_{i=1}^{n}L(\mathscr{A}_{S}(\mathbf{a}'_{i}), b'_{i})\right] = \frac{1}{n}\sum_{i=1}^{n}\mathbb{E}\left[L(\mathscr{A}_{S(i)}(\mathbf{a}'_{i}), b'_{i}) - L(\mathscr{A}_{S}(\mathbf{a}'_{i}), b'_{i})\right] \leq \beta_{n}$$

Finally note that because the samples (\mathbf{a}_i, b_i) are independent of S we have:

$$\mathbb{E}\left[\frac{1}{n}\sum_{i=1}^{n}L(\mathscr{A}_{S}(\mathbf{a}_{i}'),b_{i}')\right]=R(\mathscr{A}_{S})$$

analogously we can show $\mathbb{E}\left[R(\mathscr{A}_S)-R_S(\mathscr{A}_S)\right]\leq\beta_n.$

*Alternatives to complexity-based generalization bounds

 \circ So far we have seen that complexity based generalization bounds:

- characterize worst-case scenario
- not tight in practice
- disregard the effect of the optimization algorithm

Can we understand generalization as a property of an optimization algorithm?



*Alternatives to complexity-based generalization bounds

 \circ So far we have seen that complexity based generalization bounds:

- characterize worst-case scenario
- not tight in practice
- disregard the effect of the optimization algorithm

Can we understand generalization as a property of an optimization algorithm? YES!



*Formal definition of stability (I)

Definition (Uniform Stability [23])

Let $\mathscr{A}: \mathcal{Z} \to \mathcal{H}$ be a randomized algorithm with input a finite sample S, and output a function $\mathscr{A}_S \in \mathcal{H}$.

The algorithm \mathscr{A} has uniform stability $(\beta_n)_{n\geq 1}$ with respect to the loss function L if for all subsets $S, S' \subseteq \mathcal{A} \times \mathcal{B}$ such that |S| = |S'| = n and \overline{S} and S' differ in at most one sample:

$$\sup_{\mathbf{a},b)\in\mathcal{A}\times\mathcal{B}} \mathbb{E}|L(\mathscr{A}_{S}(\mathbf{a}),b) - L(\mathscr{A}_{S'}(\mathbf{a}),b)| \leq \beta_{n}$$

The expectation is taken with respect to the randomness in the algorithm \mathscr{A} .

Misnomer: Lower stability (small values of β_n) means the difference in the output of the algorithm is smaller.
*Formal definition of stability (II)



Figure: Algorithm \mathscr{B} is less stable than algorithm \mathscr{A} .

*The stability of SGD

• Let $h_{\mathbf{x}} \in \mathcal{H}_{\mathcal{X}}$ be an element of a parametric function class. Consider the ERM optimization objective:

$$f(\mathbf{x}) := \frac{1}{n} \sum_{i=1}^{n} f_i(\mathbf{x}), \qquad f_i(\mathbf{x}) := L(h_{\mathbf{x}}(\mathbf{a}_i), b_i).$$

• The SGD iterates for t = 0, ..., T are $\mathbf{x}_{t+1} = \mathbf{x}_t - \alpha_t \nabla_{\mathbf{x}} f_i(\mathbf{x}_t)$, for $i \sim \text{Unif}[n]$.

Algorithm	Assumptions on f_i	Stability
SGD	convex, $L\text{-smooth},\ \beta\text{-Lipschitz},\ \alpha_t \leq 2/L$	$\frac{\beta^2}{n} \sum_{\substack{t=0\\ c^2}}^T \alpha_t$
SGD	$\mu\text{-str}$ convex, $L\text{-smooth}$, $\beta\text{-Lipschitz}$, $lpha_t\leq 2/L$	$\frac{\beta^2}{n\mu}$
SGD	$\mu ext{-str convex}, ext{ L-smooth, eta-Lipschitz, $lpha_t=rac{1}{\mu t}$}$	$\frac{\beta^2 + L\rho}{n\mu}$
SGD avg. iterate	convex, L -smooth, β -Lipschitz	$\frac{\beta^2 T}{\pi I}$
SGD	non-convex, L -smooth, β -Lipschitz, $\alpha_t=1/t$	$\frac{1+1/\beta}{n} \beta^{\frac{2}{L+1}} T^{\frac{L}{L+1}}$

Table: Summary of stability upper bounds for different assumptions on the objective function [23]

*Effect of the number of iterations on the stability of SGD and the generalization error



Figure: Normalized parameter distance between two networks trained on two datasets S, S' differing only in one sample, training error, test error and generalization error (0-1 loss) on CIFAR10 [23].

o Parameter distance is a stronger notion than stability.

 \circ More iterations \Rightarrow Parameter distance increases (we expect stability to increase).

 \circ Generalization error follows the same behavior as the parameter distance (proxy for stability).

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