Generalizaiton and Regularizaiton

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## Outline

Statistical Learning Theory

# 2 Regularization

Hyperparameter Tune



# Recap: Optimization in Neural Networks

### Training Process:

- MLP are parameterized function  $f_{\theta}$ , where  $\theta = \{ W^{\ell}, b^{\ell} \}$
- The training process involves solving an optimization problem with respect to  $\theta$ :

$$\min_{\boldsymbol{\theta}} \quad \mathcal{L}(\boldsymbol{\theta}) = \frac{1}{n} \sum_{i=1}^{n} \ell(f_{\boldsymbol{\theta}}(\boldsymbol{x}_i), \boldsymbol{y}_i)$$

where  $\ell$  is a loss function and  $\mathcal{S} := \{ \pmb{x}_i, \pmb{y}_i \}_{i=1}^{\ell}$  is a training set.

• One commonly used method is called gradient descent:

$$\boldsymbol{\theta}^+ = \boldsymbol{\theta} - \eta \nabla \mathcal{L}(\boldsymbol{\theta})$$

where  $\eta > 0$  is a **learning rate**.

#### **Convergence Issues:**

- Small  $\eta$  leads to slow convergence but large  $\eta$  cause oscillations or divergence.
- DNN loss landscapes are highly complex, exhibiting large and varying condition numbers  $\kappa$
- Ill-conditioned loss landscapes cause zig-zag patterns in gradient descent.
- Unstable information propagation in DNNs can result in vanishing or exploding gradients.

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# Recap: Advanced Optimizers

### Improving Optimizations:

- Averaging gradients (or with momentum) helps smooth the descent direction.
- A larger  $\eta$  is used in GD with momentum, but training also exhibits **damping** effects in the loss.
- Adaptive methods like RMSProp rescale gradients to maintain consistent update magnitudes.
- Adaptive optimizers provide an adaptive learning rate for each gradient coordinate.
- SGD with mini-batch improves computational efficiency by using small data subsets.

### Questions

- What are common activation functions beyond sigmoid and ReLU?
- How should I choose learning rate, width, and depth for my network?
- Does gradient descent always converge? How can I speed up training?
- Does good training performance guarantee good test performance?

## Outline

Statistical Learning Theory

### 2 Regularization

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Overparameterization

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Gaussian Mixture Model			

- Assume the output y follows a discrete uniform distribution over  $\{0,1\}$ , meaning  $y \sim \mathcal{U}\{0,1\}$ .
- For each value of y, the input x follows a *Gaussian distribution*:
  - When y = 0, x follows  $x|y = 0 \sim \mathcal{N}(\mu_1, \sigma_1^2)$ , e.g.,  $\mu_1 = 1$  and  $\sigma_1 = 1$
  - When y=1, x follows  $x|y=1 \sim \mathcal{N}(\mu_2, \sigma_2^2)$ , e.g.,  $\mu_2=2$  and  $\sigma_2=2$



- This setup defines a (binary) Gaussian Mixture Model (GMM).
- Both x and y are random variables, with a joint distribution denoted as  $\mathcal{D}$ , i.e.,  $(x, y) \sim \mathcal{D}$ .

# Statistical Learning Theory (SLT)

- Assume the data (x,y) is drawn from an underlying joint distribution  $\mathcal{D}$ , i.e.,  $(x,y) \sim \mathcal{D}$ .
- The goal of learning is to find a (parameterized) function f such that:

# $f(x)\approx y$

for "most" (x, y) pairs in a probabilistic sense.

• The expected risk of f is defined as:

$$R(f) := \mathbb{E}_{(x,y)\sim\mathcal{D}}[f(x) - y]^2,$$

where we use the squared loss to measure the difference between f(x) and y.

- In practice, the distribution  $\mathcal{D}$  is **unknown**.
- Instead, we collect a random training sample  $S := \{(x_i, y_i)\}_{i=1}^n$  and compute the empirical risk or training error:

$$R_S(f) := \frac{1}{n} \sum_{i=1}^n [f(x_i) - y_i]^2$$

• By the law of large numbers, we have:

$$R_S(f) \longrightarrow R(f)$$
 as  $n \to \infty$ .

# Example of Expected and Empirical Risk using GMM

Suppose (x, y) follows GMM, and the function  $f(x) = \theta x$ , *i.e.*, parameterized linear function.

 $\bullet\,$  The expected risk R(f) is given by

$$\begin{split} R(f) = & \mathbb{E}_{(x,y)\sim\mathcal{D}}\ell(f(x),y) \\ = & \int \left[f(x) - y\right]^2 p(x,y) dx dy = \int \left[f(x) - y\right]^2 p(x|y) p(y) dx dy \\ = & \frac{1}{2} \int \left[f(x)\right]^2 p(x|y=0) dx + \frac{1}{2} \int \left[f(x) - 1\right]^2 p(x|y=1) dx \\ = & \frac{1}{2} \int \left[\theta x\right]^2 \cdot \mathcal{N}(x;\mu_1,\sigma_1^2) dx + \frac{1}{2} \int \left[\theta x - 1\right]^2 \cdot \mathcal{N}(x;\mu_2,\sigma_1^2) dx \\ & \triangleq R(\theta), \end{split}$$

where p(x,y) is the joint density, and  $\mathcal{N}(x;\mu,\sigma^2)$  is the Gaussian density defined by

$$\mathcal{N}(x;\mu,\sigma^2) = \frac{1}{\sigma\sqrt{2\pi}} e^{-(x-\mu)^2/2\sigma^2}$$

• The empirical risk  $R_S(f)$  over a training sample is given by

$$R_S(f) = \frac{1}{n} \sum_{i=1}^n [\theta x_i - y_i]^2 \triangleq R_S(\theta).$$

Statistical Learning Theory	Regularization	Hyperparameter Tune	Overparameterization
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Hypothesis Class			

In practice, we cannot evaluate all possible functions f. Instead, we restrict our search to a family of functions called a **hypothesis class**  $\mathcal{H}$ . Each function  $h \in \mathcal{H}$  is called a **hypothesis**.

• The collection of all linear models or the collection of all two-layer neural networks:

$$\mathcal{H}_1 = \{h : h(\mathbf{x}) = \mathbf{w}^\top \mathbf{x}\}$$
$$\mathcal{H}_2 = \{h : h(\mathbf{x}) = \mathbf{v}^\top \phi(\mathbf{W}\mathbf{x})\}$$

• A learning algorithm aims to find the best hypothesis  $h \in \mathcal{H}$  that minimizes the **expected risk**:

$$f_{\mathcal{H}} := \operatorname*{argmin}_{f \in \mathcal{H}} R(f).$$

- The difference  $||f^* f_H||$  is called the **approximation error**, where  $f^*$  is the ground true function.
- The Universal Approximation Theorem (UAT) implies  $||f^* f_{\mathcal{H}}|| \approx 0$  if  $\mathcal{H} = \mathcal{H}_2$ .



Statistical Learning Theory	Regularization	Hyperparameter Tune	Overparameterization
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Decomposition of Expected F	Risk		

• Given a learned hypothesis  $f_S$  from a sample S, the expected risk of  $f_S$  can be decomposed as:



• The generalization error is the difference between the expected risk and the empirical risk.



• In practice, the generalization error is estimated using the test error on an independent test set.

Statistical Learning Theory	Regularization	Hyperparameter Tune	Overparameterization
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Bounding the Generalization	Error		

• The generalization error can be upper bounded by the complexity of the hypothesis class:

 $\sup_{h\in\mathcal{H}}|R(h)-R_S(h)|\leq \text{Complexity Term},$ 

where the "Complexity Term" quantifies how flexible or complex the hypothesis class  ${\cal H}$  is.

• One commonly used complexity measure is the (empirical) Rademacher complexity:

$$\mathfrak{R}_{S}(\mathcal{H}) := \mathbb{E}_{\sigma_{i} \sim \mathcal{U}\{-1,1\}} \left[ \min_{h \in \mathcal{H}} \frac{1}{n} \sum_{i=1}^{n} \ell(h(x_{i}), \sigma_{i}) \right],$$

where  $\ell(h(x), \sigma) = \sigma h(x)$  and  $\sigma_i \in \{-1, 1\}$  are i.i.d. Rademacher random variables (uniformly distributed), *i.e.*,  $\sigma \sim \mathcal{U}\{-1, 1\}$ , and the expectation is taken over these random labels.

- Rademacher complexity measures the ability of the hypothesis class to fit **random noise** (i.e., how well the hypothesis class can fit random labels).
- Using model complexity, we can derive the following generalization bound:

$$R(f_S) \le R_S(f_S) + \Re_S(\mathcal{H}) + \tilde{\mathcal{O}}(n^{-1}),$$

where the expected risk is upper bounded by the training error and the complexity of the model.

## Example: Complexity of Linear Models

Let  $S \subseteq \{ \boldsymbol{x} : \| \boldsymbol{x} \| \leq R \}$  be a random sample, and consider  $\mathcal{H}_1 := \{ h : h(\boldsymbol{x}) = \boldsymbol{w}^\top \boldsymbol{x}, \| \boldsymbol{w} \| \leq \Lambda \}.$ 

• The (empirical) Rademacher complexity  $\mathfrak{R}_{\mathrm{S}}(\mathcal{H}_1)$  is given by

$$\begin{aligned} \mathfrak{R}_{S}(\mathcal{H}_{1}) = & \mathbb{E}_{\sigma_{i}} \left[ \min_{h \in \mathcal{H}_{1}} \frac{1}{n} \sum_{i=1}^{n} \ell(h(x_{i}), \sigma_{i}) \right] = \mathbb{E}_{\sigma_{i}} \left[ \min_{\|\boldsymbol{w}\| \leq \lambda} \frac{1}{n} \sum_{i=1}^{n} \sigma_{i} \boldsymbol{w}^{\top} \boldsymbol{x}_{i} \right] \\ \leq & \frac{\Lambda}{n} \mathbb{E}_{\sigma_{i}} \left[ \left\| \sum_{i=1}^{n} \sigma_{i} \boldsymbol{x}_{i} \right\| \right] \leq \frac{\Lambda}{n} \left[ \mathbb{E}_{\sigma_{i}} \left\| \sum_{i=1}^{n} \sigma_{i} \boldsymbol{x}_{i} \right\|^{2} \right]^{1/2} \\ \leq & \frac{\Lambda}{n} \sqrt{nR^{2}} = \sqrt{\frac{R^{2}\Lambda^{2}}{n}}, \end{aligned}$$

where we use the Cauchy-Schwartz and Jensen's inequalities.

• As a result, the generalization error for linear models satisfies (with high probability):

$$R(h_S) \le R_S(h_S) + \sqrt{\frac{R^2 \Lambda^2}{n}} + \tilde{\mathcal{O}}(n^{-1}).$$

• More data improves the empirical risk  $R_S$  as an **approximation** of the expected risk R, reducing overfitting, but overall performance still depends on minimizing  $R_S$ .

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### Model Complexity Trade-Off

The expected risk  $R(f_S)$  is upper bounded by the training error and the model complexity:

 $R(f_S) \le R_S(f_S) + \mathfrak{R}_S(\mathcal{H}) + \tilde{\mathcal{O}}(n^{-1}).$ 



### Key Insights on Generalization Bound

- If the model is too simple, it may fail to fit the training data well. This is known as **underfitting**.
- Conversely, if the model is highly flexible, it may achieve low training error, but perform poorly on unseen data. This is known as **overfitting**.
- The goal is to find a "sweet spot" balancing underfitting and overfitting to minimize the overall expected risk.

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Optimal Hypothesis $f^*$			

**Claim**:  $f^*(x) = \mathbb{E}[y|x]$  is the **optimal hypothesis** that minimizes the expected risk.

#### Proof.

For any function f, we can decompose the expected risk as follows:

$$R(f) = \mathbb{E}(f - y)^2 = \mathbb{E}(f - f^* + f^* - y)^2$$
  
=  $\mathbb{E}(f - f^*)^2 + 2\mathbb{E}(f - f^*)(f^* - y) + \mathbb{E}(f^* - y)^2$   
=  $\mathbb{E}(f - f^*)^2 + \mathbb{E}(f^* - y)^2$   
 $\geq \mathbb{E}(f^* - y)^2$   
=  $R(f^*)$ 

where the cross term  $\mathbb{E}(f - f^*)(f^* - y) = 0$ , because  $f^*(x) = \mathbb{E}[y|x]$ .

- This is another existence result.
- The optimal hypothesis  $f^*$  is not directly accessible unless we know the joint distribution  $\mathcal{D}$ .
- Generally, we may have  $R(f^*) \neq 0$ . For example, consider  $y = \theta x + \varepsilon$ , where  $\varepsilon \sim \mathcal{N}(0, \sigma^2)$

$$f^*(x) = \mathbb{E}[y|x] = \mathbb{E}[\theta x + \varepsilon | x] = \theta x$$
  

$$R(f^*) = \mathbb{E}_x[f^*(x) - y]^2 = \mathbb{E}_x[\theta x - (\theta x + \varepsilon)]^2 = \sigma^2 \implies \text{ irreducible error.}$$

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- Bias-Variance Decomposition of Expected Risk
  - The learned function  $f_S$  depends on the random sample S, making  $f_S$  a random variable.
  - Hence, the expected risk  $R(f_S)$  is also random, and it varies across different random samples S.
  - To capture this variability, we consider the expectation of the  $R(f_S)$  over all possible samples S, i.e.,  $\mathbb{E}_S[R(f_S)]$ .
  - Let  $\overline{f} := \mathbb{E}_S[f_S]$ , the expected or average hypothesis over all random samples S.
  - Using  $\bar{f}$ , we can decompose  $\mathbb{E}_S[R(f_S)]$  as follows:

$$\begin{split} \mathbb{E}_{S}[R(f_{S})] = & \mathbb{E}_{S} \mathbb{E}_{(x,y)\sim\mathcal{D}}[f_{S}(x) - y]^{2} \\ = & \mathbb{E}_{S} \mathbb{E}_{\mathcal{D}}[f_{S} - f^{*})]^{2} + R(f^{*}) \\ = & \mathbb{E}_{S} \mathbb{E}_{\mathcal{D}}[f_{S} - \bar{f} + \bar{f} - f^{*}]^{2} + R(f^{*}) \\ = & \mathbb{E}_{S} \mathbb{E}_{\mathcal{D}}\Big[(f_{S} - \bar{f})^{2} + (\bar{f} - f^{*})^{2}\Big] + R(f^{*}) \\ = & \underbrace{\mathbb{E}_{S}(f_{S} - \bar{f})^{2}}_{\text{Variance term}} + \underbrace{\mathbb{E}_{\mathcal{D}}(\bar{f} - f^{*})^{2}}_{\text{Bias term}} + \underbrace{R(f^{*})}_{\text{irreducible}} \end{split}$$

where the cross term  $\mathbb{E}_{S,\mathcal{D}}(f_S - \mathbb{E}_S[f_S])(\mathbb{E}_S(f_S) - f^*) = 0$  cancels out.

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### **Bias-Variance Trade-Off**

The expected risk  $\mathbb{E}_S[R(f_S)]$  can be broken down into three parts:

- Squared Bias:  $\mathbb{E}_{\mathcal{D}}[(f^* \mathbb{E}_S(f_S))^2]$  measures the error from approximating the optimal function  $f^*$  with the learned model  $f_S$ . It reflects the error caused by using a simple model that cannot capture all the data patterns.
- Variance:  $Var(f_S) = \mathbb{E}_S[(f_S \mathbb{E}_S(f_S))^2]$  measures how much the learned function  $f_S$  varies with different training samples. It represents the error due to the model's sensitivity to fluctuations in the random training sample S.
- Irreducible Error:  $R(f^*)$  represents the inherent noise in the data, which no model can eliminate. It is the error we cannot reduce.



- **High bias, low variance**: Simple models (e.g., linear models) have *low variance* since they are less sensitive to training data, but have *high bias* because they are too simple to capture all patterns in the data.
- Low bias, high variance: Complex models (e.g., polynomial model) have *low bias* as they can model complex relations, but *high variance* due to overfitting to the training data.

Statistical Learning Theory	Regularization	Hyperparameter Tune	Overparameterization
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Summary of Statistica	l Learning Theory		

• The goal is to find a hypothesis f within a hypothesis class  $\mathcal{H}$  that minimizes the **expected risk**:

$$R(f) = \mathbb{E}_{(x,y)\sim\mathcal{D}}\left[\left(f(x) - y\right)^2\right].$$

• Since the underlying distribution  $\mathcal{D}$  is **unknown**, we approximate f by minimizing the **empirical** risk based on a random training sample S:

$$R_S(f) = \frac{1}{n} \sum_{i=1}^n (f(x_i) - y_i)^2.$$

• Using model complexity  $\mathfrak{R}_{\mathrm{S}}(\mathcal{H})$ , the expected risk is upper bound as:

$$R(f_S) \le R_S(f_S) + \mathfrak{R}_S(\mathcal{H}) + \tilde{\mathcal{O}}(n^{-1}),$$

- By considering variations across different random training samples S, the expected risk  $\mathbb{E}_S[R(f_S)]$  can be decomposed into three components: bias, variance, and irreducible error:
  - High bias, low variance: Simple models underfit and miss important patterns in the data.
  - Low bias, high variance: Complex models overfit and perform poorly on unseen data.
- Find the "sweet spot" between underfitting and overfitting to minimize the overall expected risk.

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# Outline

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# DNNs Can Fit Random Labels and Random Data



### Key Observation

- Label corruption: Replace true label with random label
- **Shuffled pixels**: The pixels of each image are rearranged using a *fixed* random permutation
- **Random pixels**: Each image has a *unique* random arrangement of pixels).
- Gaussian: The pixels in images are replaced with random Gaussian *noise*.
- Average loss: *Training error* using the cross-entropy loss

DNNs can perfectly fit random labels or data, achieving **zero training error** even on completely unstructured inputs.

Zhang et al. "Understanding deep learning requires rethinking generalization" ICLR 2017.

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# Weight Decay

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Weight Decay			

• Regularization typically involves adding an extra term, called the regularizer, to the training loss:

$$\mathcal{L}_{\lambda}(oldsymbol{ heta}) := \mathcal{L}(oldsymbol{ heta}) + rac{\lambda}{2} \|oldsymbol{ heta}\|^2,$$

where  $\lambda > 0$  is the **regularization hyperparameter**, and  $\|\cdot\|$  is the Euclidean norm.

• In deep learning, this regularization is known as weight decay because gradient descent on the regularized loss automatically shrinks (or decays) parameter  $\theta$  by the factor  $(1 - \eta\lambda)$ :

$$\begin{split} \boldsymbol{\theta}^{+} = & \boldsymbol{\theta} - \eta \nabla_{\boldsymbol{\theta}} \mathcal{L}_{\lambda}(\boldsymbol{\theta}) = \boldsymbol{\theta} - \eta \left[ \nabla_{\boldsymbol{\theta}} \mathcal{L}(\boldsymbol{\theta}) + \lambda \boldsymbol{\theta} \right] \\ = \underbrace{(1 - \eta \lambda)}_{\text{decaving weights}} \boldsymbol{\theta} - \eta \nabla_{\boldsymbol{\theta}} \mathcal{L}(\boldsymbol{\theta}). \end{split}$$

• However,  $\theta$  does **not** shrink to zero, as it must maintain a certain value to minimize the cost  $\mathcal{L}(\theta)$ .

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Interpretation: Sparsity			

• The regularized optimization can be reformulated as:

$$\min_{\boldsymbol{\theta}} \mathcal{L}(\boldsymbol{\theta}), \quad \text{s.t.} \quad \|\boldsymbol{\theta}\| \leq C_{\lambda},$$

where  $C_{\lambda} > 0$  is a constant that depends on  $\lambda$ .

- In deep learning,  $\theta$  is called **sparse** if most parameters are zero or close to zero (i.e.,  $\theta_i \approx 0$ ).
- Sparse  $\theta$  reduces the flexibility and complexity of the DNN, leading to a simpler model.



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Interpretation: Linearity			

Consider a simple two-layer neural network:

$$f_{\boldsymbol{\theta}}(x) = \sum_{i=1}^{n} v_i \phi(w_i x),$$

where  $x \in \mathbb{R}$  is a scalar and  $\phi(\cdot)$  is tanh.

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• When  $w_i \approx 0$ , then  $w_i x \approx 0$ , and the network operates near the linear region of tanh:



$$v_i\phi(w_ix)\approx v_i(w_ix)\approx (v_iw_i)x=u_ix \quad \Longrightarrow \quad \text{a linear model},$$
 where  $u_i:=v_iw_i.$  If  $v_i\approx 0,$  then

 $v_i\phi(w_ix)\approx 0,$ 

indicating fewer neurons are used.

Interpretation: Stability			
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A learning algorithm is stable if small changes to its input do not result in large changes to its output.

• Consider the same two-layer neural network:

$$f_{\boldsymbol{\theta}}(x) = \sum_{i=1}^{n} v_i \phi(w_i x).$$

• The derivative of  $f_{\theta}$  with respect to the **input** x is:

$$\nabla_x f_{\theta}(x) = \sum_{i=1}^n v_i \phi'(w_i x) w_i.$$

- If either  $v_i$  or  $w_i$  is small, then  $\nabla_x f_{\theta}(x)$  is small.
- Hence, DNNs with sparse parameters are generally more stable than those with dense parameters.

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# Dropout

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Dropout Regularization			
Recall the forward prop	pagation:		

$$\boldsymbol{z}^{\ell} = \boldsymbol{W}^{\ell} \boldsymbol{x}^{\ell-1}, \quad \boldsymbol{x}^{\ell} = \phi(\boldsymbol{z}^{\ell}).$$

• During training, each neuron is randomly dropped with probability p (a hyperparameter):

$$oldsymbol{z}^\ell = oldsymbol{W}^\ell \left(oldsymbol{r}^\ell \odot oldsymbol{x}^{\ell-1}
ight), \quad oldsymbol{x}^\ell = \phi(oldsymbol{z}^\ell),$$

where  $r_i^{\ell} \stackrel{\textit{i.i.d.}}{\sim}$  Bernoulli(p) and  $\odot$  is element-wise product.



- The gradient update applies only to a thinned subbnet of the network.
- At test time, dropout is turned off, and weights are scaled by p to respect the dropout probability:

$$\boldsymbol{z}^{\ell} = p \boldsymbol{W}^{\ell} \boldsymbol{x}^{\ell-1}.$$

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- By randomly dropping units, a different thinned subnet is trained at each gradient descent step.
- With n neurons in the full network, we are effectively training  $2^n$  different subnets simultaneously that all share the same weights.
- At test time, the output is an ensemble prediction, aggregating the contributions of all subnets.



### Key Insight

Dropout ensures that **no** single neuron or small group of neurons can dominate the prediction. By **spreading** the responsibility across all units, it improves model robustness to the input change and prevents overfitting.

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# **Stochastic Weight Averaging**

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Trajectories of SGD			





- SGD oscillates around the periphery of high-performing solutions, and averaging SGD iterates improves test performance.
- SGD trajectories resemble a high-dimensional Gaussian-like distribution, with most of the mass concentrated in a **thin shell**.

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Averaging Weights for	Better Pertormance		

• Averaging SGD iterates leads to improved generalization:

$$ar{m{w}} = rac{1}{k}\sum_{i=1}^km{w}^i$$

• Averaging weights approximates ensembling predictions via linearization (if the weights are close):

$$\frac{1}{k}\sum_{i=1}^{k}f(\boldsymbol{w}^{i})\approx f\left(\frac{1}{k}\sum_{i=1}^{k}\boldsymbol{w}^{i}\right)=f(\bar{\boldsymbol{w}})$$

• Moving average formulation:

$$\begin{split} \boldsymbol{w}^{k+1} &= \boldsymbol{w}^k - \eta \nabla \mathcal{L}(\boldsymbol{w}^k) \\ \boldsymbol{w}^{k+1}_{\text{swa}} &= (1 - \beta^k) \boldsymbol{w}^k_{\text{swa}} + \beta^k \boldsymbol{w}^{k+1} \end{split}$$

where  $\beta^k = \frac{k}{k+1}$  or  $\beta^k = \beta \in (0,1).$ 



- DNNs can fit random labels and data, achieving zero training error.
- Weight decay controls large weights, promoting sparsity, linearity, and stability.
- During training, dropout randomly drops units, effectively training an **exponential number** of **thinned subnets** simultaneously.
- At test time, the output is an ensemble prediction, aggregating contributions from all subnets.
- SGD oscillates near the **boundary** of local minima, while SWA finds a **centralized** solution in a flatter region.
- SWA approximates ensemble predictions through linearization.

# Outline

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Overparameterization

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Recap: Hyperparameters in	Neural Networks		
Statistical Learning Theory	Regularization	Hyperparameter Tune	Overparameterization
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The training process involves several key hyperparameters:

- Loss Function  $\ell(\cdot, \cdot)$ : Square loss, cross-entropy loss, hinge loss
- Activation Function  $\phi(\cdot)$ : Step, sigmoid, ReLU, tanh, GELU
- Optimizer: SGD, Momentum, RMSProp, Adam, AdamW
- Learning Rate  $(\eta)$ , Batch Size (b), Epochs
- Network Type: MLPs, CNNs, RNNs, Transformers, GNNs
- Width and Depth
- Layers: Normalization, pooling, dropout, softmax
- Otherwise: Initialization (Xavier, He),  $\ell_2$ -regularization, gradient clipping, early stop

### Key Difference: Hyperparameters vs. Trainable Parameters

- Hyperparameters are not trainable. Unlike weights and biases, they need to be tuned.
- Proper tuning is essential for faster convergence during training and achieving good generalization performance.

# Validation Set

- Split the dataset into three parts: training set, validation set, and test set.
- Build the model using the training set.
- Optimize or tune hyperparameters on the validation set.
- After tuning, evaluate the final model on the test set.
- Suggested split ratios:
  - For datasets between 100 and 1,000,000 samples: 60/20/20.
  - For datasets larger than 1,000,000 samples: 98/1/1.
- Ensure the validation and test sets come from the same distribution.
  - Example: Training and validation images from the web, but test images from user cell phones can cause a mismatch.

# Tuning Process

- Grid Search: Systematically explores a predefined set of hyperparameters; comprehensive but expensive
- Random Search: Randomly sample hyperparameters; more efficient than grid search when some hyperparameters are less important.
- Hyperband/Successive Halving: Dynamically allocate resources and discard poor configurations early, ideal for deep networks with long training times.
- Start with coarse tuning, then refine gradually.
- $\bullet$  Use  $\log$  scale for hyperparameter search when appropriate, e.g., learning rate  $\eta$  and smoothing factors  $\beta$
- Leverage parallelization to run multiple experiments simultaneously to accelerate the search.

Hyperparameter Tune

### Input Normalization

• Normalize the inputs using training set:

$$oldsymbol{\mu} = rac{1}{n}\sum_{i=1}^noldsymbol{x}_i, \quad oldsymbol{ar{x}}_i = oldsymbol{x}_i - oldsymbol{\mu}, \quad oldsymbol{\sigma}^2 = rac{1}{n}\sum_{i=1}^noldsymbol{ar{x}}_i^2, \quad oldsymbol{\hat{x}}_i = oldsymbol{ar{x}}_i/oldsymbol{\sigma},$$

where all operations are taken element-wise.

- Consider a binary classification problem using linear model:  $f_{\theta}(x) = w_1 x_1 + w_2 x_2$ 
  - if  $x_1 = \mathcal{O}(100)$  and  $x_2 = \mathcal{O}(1)$ , to have output  $f_{\theta} = \mathcal{O}(1)$ , we must have  $w_1 = \mathcal{O}\left(\frac{1}{100}\right)$  and  $w_2 = \mathcal{O}(1)$ .
  - After normalization,  $\bar{x}_1 = \mathcal{O}(1)$  and  $\bar{x}_2 = \mathcal{O}(1)$ , so we have  $w_1 = \mathcal{O}(1)$  and  $w_2 = \mathcal{O}(1)$ .



• At test time, apply  $\mu$  and  $\sigma$  from training to test set.

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# Learning Rate Decay

- $\bullet$  Recall that an  ${\bf epoch}\;k$  is one pass through all  ${\bf mini-batches}$  in SGD
- Instead of using a fixed learning rate, one can consider using learning rate decay

$$\eta_k = \frac{\eta}{k}, \qquad \eta_k = \frac{\eta}{\sqrt{k}}, \qquad \eta_k = (0.95)^k \eta$$

# Bag of Tips

### Learning Rate $\eta$ :

- Log-scale search:  $10^{-5} \sim 10^{-1}$ .
- Learning rate schedules: Linearly warm up, then decay periodically for smooth convergence.
- Early stopping: Monitor loss curves to detect divergence.

### Batch Size b:

- Small batches (e.g.,  $16 \sim 128$ ) generalize better, but noisy gradient.
- Large batches (e.g.,  $256 \sim 4096$ ) converge faster but may require higher learning rates.
- Rule of Thumb: Use the largest batch size that fits in memory, then tune;  $\eta' = \eta \times \frac{b'}{b}$ .

### Weight Decay:

- Log-scale search:  $10^{-5} \sim 10^{-3}$ .
- For Adam: Use AdamW instead of standard weight decay.

$$oldsymbol{w} \leftarrow oldsymbol{w} - \eta rac{oldsymbol{v}}{\sqrt{oldsymbol{s}+oldsymbol{arepsilon}}} - \eta \lambda oldsymbol{w} \quad \Longrightarrow \quad oldsymbol{w} \leftarrow oldsymbol{w} - \eta rac{oldsymbol{v}}{\sqrt{oldsymbol{s}+oldsymbol{arepsilon}}} - \lambda oldsymbol{w}$$

where weight decay is scaled by the small  $\eta$  in Adam, reducing the regularization effect.

• If validation loss diverges while training loss improves, increase weight decay.

### Dropout:

- Start with  $0.2 \sim 0.5$  for input layers,  $0.5 \sim 0.8$  for hidden layers.
- $\bullet$  Combine dropout with  $\ell_2\text{-}\mathsf{regularization}$  but avoid using it with Batch normalization.

### **Optimizers:**

- SGD+Momentum: More stable than vanilla SGD.
- Adam works well for most tasks with default values  $\beta_1=0.9$  and  $\beta_2=0.999$
- Use AdamW for better weight decay handling.
- **RMSProp**: Useful for RNNs and reinforcement learning.

### Network Architecture (Depth and Width):

- Start simple and gradually increase the complexity
- More layers (depth) improve feature extraction, using skip connections if too deep
- More neurons (width) increase capacity and stabilize training

### **Activation Functions:**

- ReLU: Standard choice for DNNs.
- Leaky ReLU: Fixes dying ReLU problem ( $\alpha = 0.01$ ).
- GELU: Used in Transformers.
- Swish: Works well in CNNs.

# Outline

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Hyperparameter Tune 00000000 Overparameterization 0000

### Overparameterization

- A deep neural network (DNN) is said to be **overparameterized** when the number of neurons or parameters is much larger than the number of training samples.
- This might seem counterintuitive, but it has been found to be surprisingly beneficial in practice.



Behnam, et al. "Towards Understanding the Role of Over-Parametrization in Generalization of Neural Networks," ICLR 2019 🚽 🔿 🖉

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### Double Descent

Overparameterized neural networks can perfectly fit or interpolate the training data.

ullet Mathematically, there exists a set of parameters  ${m heta}$  such that

$$f_{\theta}(x_i) = y_i, \quad \forall i \in [n].$$
(1)

- Overparameterization implies there are infinitely many interpolation solutions.
- Some interpolation solutions generalize much better than those in the *underparameterized* regime. This phenomenon is called **double descent**.



Hyperparameter Tune 00000000

# Implicit Regularization

- It is important to understand that different global minima lead to varying test performances.
- A flat minimum typically results in better generalization than a sharp minimum.
- Different optimizers may converge to different minima, each with different generalization outcomes. This is known as **implicit regularization**.
- Thus, even if your current optimizer achieves low training error, tuning or adjusting it may still be necessary to achieve better test performance.

