# Deep learning theory lecture notes

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## Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preface</td>
<td>Basic setup: feedforward networks and test error decomposition</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Highlights</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Missing topics and references</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>Acknowledgements</td>
<td>7</td>
</tr>
<tr>
<td>1</td>
<td>Approximation: preface</td>
<td>7</td>
</tr>
<tr>
<td>1.1</td>
<td>Omitted topics</td>
<td>8</td>
</tr>
<tr>
<td>2</td>
<td>Classical approximations and “universal approximation”</td>
<td>8</td>
</tr>
<tr>
<td>2.1</td>
<td>Elementary folklore constructions</td>
<td>9</td>
</tr>
<tr>
<td>2.2</td>
<td>Universal approximation with a single hidden layer</td>
<td>11</td>
</tr>
<tr>
<td>3</td>
<td>Infinite-width Fourier representations and the Barron norm</td>
<td>14</td>
</tr>
<tr>
<td>4</td>
<td>Approximation near initialization and the Neural Tangent Kernel (NTK)</td>
<td>14</td>
</tr>
<tr>
<td>5</td>
<td>Benefits of depth</td>
<td>14</td>
</tr>
<tr>
<td>6</td>
<td>Optimization: preface</td>
<td>14</td>
</tr>
<tr>
<td>6.1</td>
<td>Omitted topics</td>
<td>15</td>
</tr>
<tr>
<td>7</td>
<td>Smooth objectives in ML</td>
<td>16</td>
</tr>
<tr>
<td>7.1</td>
<td>Convergence to stationary points</td>
<td>18</td>
</tr>
<tr>
<td>7.2</td>
<td>Convergence rate for smooth &amp; convex</td>
<td>20</td>
</tr>
<tr>
<td>8</td>
<td>Strong convexity</td>
<td>22</td>
</tr>
<tr>
<td>8.1</td>
<td>Rates when strongly convex and smooth</td>
<td>23</td>
</tr>
<tr>
<td>9</td>
<td>Stochastic gradients</td>
<td>26</td>
</tr>
<tr>
<td>10</td>
<td>NTK-based Gradient flow analysis on smooth shallow networks, following (Chizat and Bach 2019)</td>
<td>29</td>
</tr>
<tr>
<td>10.1</td>
<td>Proof of Theorem 10.1</td>
<td>33</td>
</tr>
<tr>
<td>11</td>
<td>Nonsmoothness, Clarke differentials, and positive homogeneity</td>
<td>37</td>
</tr>
<tr>
<td>11.1</td>
<td>Positive homogeneity</td>
<td>38</td>
</tr>
<tr>
<td>11.2</td>
<td>Positive homogeneity and the Clarke differential</td>
<td>39</td>
</tr>
</tbody>
</table>
Preface

Philosophy of these notes. Two key ideas determined what has been included so far.
1. I aim to provide simplified proofs over what appears in the literature, ideally reducing difficult things to something that fits in a single lecture.

2. I have primarily focused on a classical perspective of achieving a low test error for binary classification with IID data via standard (typically ReLU) feedforward networks.

Organization. Following the second point above, the classical view decomposes the test error into three parts.

1. Approximation (starts in section 1): given a classification problem, there exists a deep network which achieves low error over the distribution.

2. Optimization (starts in section 6): given a finite training set for a classification problem, there exist algorithms to find predictors with low training error and low complexity.

3. Generalization (starts in section 13): the gap between training and testing error is small for low complexity networks.

Remark 0.1 (weaknesses of this “classical” approach)

- Recent influential work suggests that the classical perspective is hopelessly loose, and has poor explanatory power (Neyshabur, Tomioka, and Srebro 2014; Zhang et al. 2017). Follow-ups highlight this looseness and its lack of correlation with good test error performance (Dziugaite and Roy 2017), and even suggest the basic approach is flawed (Nagarajan and Kolter 2019); please see section 13.1 for further discussion and references.

- The reasons for keeping with this approach here are as follows:

  1. It appears that all of these negative results consider the consequences of worst-case behavior in one of these three terms on the other two. Here instead we study how they inter-connect in a favorable way. A common theme is how they all work together with low complexity models on reasonable data.

  2. Even if the preceding point is overly optimistic at times, this decomposition still gives us a way to organize and categorize much of what is known in the field, and secondly these ideas will always be useful at least as tools in a broader picture.

Formatting.

- These notes use pandoc markdown with various extensions. A current html version is always at https://mjt.cs.illinois.edu/dlt/, and a current pdf version is always at https://mjt.cs.illinois.edu/dlt/index.pdf.

- Owing to my unfamiliarity with pandoc, there are still various formatting bugs.

- [ mjtΩ: Various todo notes are marked throughout the text like this.]

Feedback. I’m very eager to hear any and all feedback!

How to cite. Please consider using a format which makes the version clear:

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Basic setup: feedforward networks and test error decomposition

In his section we outline our basic setup, which can be summarized as follows:

1. We consider standard shallow and deep feedforward networks.
2. We study mainly binary classification in the supervised learning setup.
3. As above, we study an error decomposition into three parts.

Although this means we exclude many settings, as discussed above, much of the work in other settings uses tools from this most standard one.

Basic shallow network. Consider the mapping

\[ x \mapsto \sum_{j=1}^{m} a_j \sigma(w_j^T x + b_j). \]

- \( \sigma \) is the nonlinearity/activation/transfer. Typical choices: ReLU \( z \mapsto \max\{0, z\} \), sigmoid \( z \mapsto \frac{1}{1+\exp(-z)} \).
- \((a_j, w_j, b_j))_{j=1}^{m} \) are trainable parameters; varying them defines the function class. Sometimes in this shallow setting we freeze \((a_j)_{j=1}^{m}\), which gives a simple model that is still difficult to analyze (e.g., nonconvex).
- We can think of this as a directed graph of width \( m \): we have a hidden layer of \( m \) nodes, where the \( j \)th computes \( x \mapsto \sigma(w_j^T x + b_j) \).
- Define weight matrix \( W \in \mathbb{R}^{m \times d} \) and bias vector \( v \in \mathbb{R}^m \) as \( W_j = w_j^T \) and \( v_j := b_j \). The first layer computes \( h := \sigma(Wx + b) \in \mathbb{R}^m \) (\( \sigma \) applied coordinate-wise), the second computes \( h \mapsto a^T h \).

Basic deep network. Extending the matrix notation, given parameters \( w = (W_1, b_1, \ldots, W_L, b_L) \),

\[ f(x; w) := \sigma_L(W_L \sigma_{L-1}(\cdots W_2 \sigma_1(W_1 x + b_1) + b_2 \cdots) + b_L). \]  \hspace{1cm} (1)

- \( \sigma_j \) is now a multivariate mapping; in addition to coordinate-wise ReLU and sigmoid, we can do softmax \( z' \propto \exp(z) \), max-pooling (a few coordinates of input replaced with their maximum), attention layers, and many others.
- We can replace \( x \mapsto Wx + b \) with some compact representation while still preserving linearity, e.g., the standard implementation of a convolution layer. [ mjt\$: Maybe I will add the explicit formalisms somewhere? ].
- Often biases \((b_1, \ldots, b_L)\) are dropped; the handling of these biases can change many elements of the story.
- Typically \( \sigma_L \) is identity, so we refer to \( L \) as the number of affine layers, and \( L-1 \) the number of activation or hidden layers.
- Width now means the maximum output dimension of each activation. (For technical reasons, sometimes need to also take max or input dimension, or treat inputs as a fake layer.)
Once again we can describe the computation via an acyclic graph. Classically, the activations were univariate mappings applied coordinate-wise, and single rows of the weight matrix were composed with univariate activations to give a node. Now, however, activations are often multivariate (and in particular can not be written as identical univariate mappings, applied coordinate-wise), and for computation reasons we prefer not to break the matrices into vectors, giving a more general graph with each matrix and activation as its own node.

**Basic supervised learning setup; test error decomposition.**

- Given pairs $\mathcal{P} = \{(x_i, y_i)\}_{i=1}^n$ (training set), our job is to produce a mapping $x \mapsto y$ which performs well on future examples.
- If there is no relationship between past and future data, we can’t hope for much.
- The standard classical learning assumption is that both the training set, and future data, are drawn IID from some distribution on $(x, y)$.
- This IID assumption is not practical: it is not satisfied by real data. Even so, the analysis and algorithms here have many elements that carry over to more practical settings.

How do we define “performs well on future examples?”

- Given one $(x, y)$ and a prediction $\hat{y} = f(x)$, we suffer a loss $\ell(\hat{y}, y)$, e.g., logistic $\ln(1+\exp(-\hat{y}y))$, or squared $(\hat{y} - y)^2/2$.
- On a training set, we suffer empirical risk $\tilde{\mathcal{R}}(f) = \frac{1}{n} \sum_i \ell(f(x_i), y_i)$.
- For future (random!) data, we consider (population) risk $\mathcal{R}(f) = \mathbb{E} \ell(f(x), y) = \int \ell(f(x), y) d\mu(x, y)$.

“Performs well on future examples” becomes “minimize $\mathcal{R}(f)$.” We can decompose $\mathcal{R}(f)$ into three separate concerns: given a training algorithm’s choice $\hat{f}$ in some class of functions/predictors $\mathcal{F}$, as well as some reference solution $\bar{f} \in \mathcal{F}$,

$$\mathcal{R}(\hat{f}) = \mathcal{R}(\hat{f}) - \mathcal{R}(\bar{f})$$

**(generalization)**

$$+ \mathcal{R}(\bar{f}) - \mathcal{R}(\bar{f})$$

**(optimization)**

$$+ \mathcal{R}(\hat{f}) - \mathcal{R}(\bar{f})$$

**(concentration/generalization)**

$$+ \mathcal{R}(\bar{f})$$

**(approximation)**

These notes are organized into separately considering these three terms (treating “generalization” and “concentration/generalization” together).

**Remark 0.2 (sensitivity to complexity)** As discussed, we aim to circumvent the aforementioned pitfalls by working with notions of low complexity model which work well with all three parts. There is still very little understanding of the right way to measure complexity, however here are some informal comments.

- First suppose there exists a low complexity $\bar{f} \in \mathcal{F}$ so that the approximation term $\mathcal{R}(\bar{f})$ is small. Since the complexity is low, then the concentration/generalization term $\tilde{\mathcal{R}}(\bar{f}) - \mathcal{R}(\bar{f})$ is small.

- Since $\bar{f}$ has low complexity, then hopefully we can find $\hat{f}$ with not much larger complexity via an algorithm that balances the optimization term $\tilde{\mathcal{R}}(\hat{f}) - \tilde{\mathcal{R}}(\bar{f})$ with the complexity of $\hat{f}$; if $\hat{f}$ has low complexity, then the generalization term $\mathcal{R}(\hat{f}) - \tilde{\mathcal{R}}(\hat{f})$ will be small.
Remark 0.3 The two-argument form $\ell(\hat{y}, y)$ is versatile. We will most often consider binary classification $y \in \{\pm 1\}$, where we always use the product $\hat{y}y$, even for the squared loss:

$$[\hat{y} - y]^2 = [y(y\hat{y} - 1)]^2 = (y\hat{y} - 1)^2.$$ 

This also means binary classification networks have output dimension one, not two.

**Highlights**

Here are a few of the shortened and/or extended proofs in these notes.

1. **Approximation.**
   - (Section 2.2) Succinct universal approximation via Stone-Weierstrass.
   - (Section 3) Succinct Barron’s theorem (Fourier representation), with an explicit infinite width form.
   - (Section 5) Shorter depth separation proof.

2. **Optimization.**
   - (Section 10) Short re-proof of gradient flow convergence in the shallow NTK regime, due to (Chizat and Bach 2019).
   - (Section 12.3) Short proof that smooth margins are non-decreasing for homogeneous networks; originally due to (Lyu and Li 2019), this short proof is due to (Ji 2020).

3. **Generalization.**
   - (Section 18.2) Shortened “spectrally-normalized bound” proof (P. Bartlett, Foster, and Telgarsky 2017).
   - (Section 19.3) Shortened ReLU network VC dimension proof.

**Missing topics and references**

Due to the above philosophy, many topics are currently omitted. Over time I hope to fill the gaps. Here are some big omissions, hopefully resolved soon:

- **Architectures:**
  - Non-feedforward, e.g., recurrent (Siegelmann and Sontag 1994).
  - Specific feedforward architecture choices like convolutional layers and skip connections.
  - Continuous depth, for instance various neural ODE frameworks (Chen et al. 2018; Tzen and Raginsky 2019).

- **Other learning paradigms:**
  - Data augmentation, self-training, and distribution shift.
  - Unsupervised learning (e.g., GANs), Adversarial ML, RL.

Further omitted topics, in a bit more detail, are discussed separately for approximation (section 1.1), optimization (section 6.1), and generalization (section 13.1).
Acknowledgements

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1 Approximation: preface

As above, we wish to ensure that our predictors $\mathcal{F}$ (e.g., networks of a certain architecture) have some element $f \in \mathcal{F}$ which simultaneously has small $\mathcal{R}(f)$ and small complexity; we can re-interpret our notation and suppose $\mathcal{F}$ already is some constrained class of low-complexity predictors, and aim to make $\inf_{f \in \mathcal{F}} \mathcal{R}(f)$ small.

What is $\mathcal{F}$? In keeping with the earlier theme, it should be some convenient notion of “low complexity model”; but what is that?

1. **Models reached by gradient descent.** Since standard training methods are variants of simple first-order methods, it seems this might be a convenient candidate for $\mathcal{F}$ which is tight with practice. Unfortunately, firstly we only have understanding of these models very close to initialization and very late in training, whereas practice seems to lie somewhere between. Secondly, we can’t just make this our definition as it breaks things in the standard approach to generalization.

2. **Models of low norm,** where norm is typically measured layer-wise, and also typically the “origin” is initialization. This is the current most common setup, though it doesn’t seem to be able to capture the behavior of gradient descent that well, except perhaps when very close to initialization.

3. **All models of some fixed architecture,** meaning the weights can be arbitrary. This is the classical setup, and we’ll cover it here, but it can often seem loose or insensitive to data, and was a key part of the criticisms against the general learning-theoretic approach (Zhang et al. 2017). The math is still illuminating and still key parts can be used as tools in a more sensitive analysis, e.g., by compressing a model and then applying one of these results.

The standard classical setup (“all models of some fixed architecture”) is often stated with a goal of competing with all continuous functions:

$$\inf_{f \in \mathcal{F}} \mathcal{R}(f) \quad \text{versus} \quad \inf_{g \text{ continuous}} \mathcal{R}(g).$$

E.g.,

$$\sup_{g \text{ cont.}} \inf_{f \in \mathcal{F}} \mathcal{R}(f) - \mathcal{R}(g).$$
To simplify further, if $\ell$ is $\rho$-Lipschitz,
\[
\mathcal{R}(f) - \mathcal{R}(g) = \int (\ell(yf(x)) - \ell(yg(x))) \, d\mu(x, y)
\leq \int \rho |yf(x) - yg(x)| \, d\mu(x, y) = \rho \int |f(x) - g(x)| \, d\mu(x, y),
\]
and in particular we have reduced the approximation question to one about studying $\|f - g\|$ with function space norms.

**Remark 1.1** *(Is this too strenuous?)* Most of the classical work uses the uniform norm: $\|f - g\|_u = \sup_{x \in S} |f(x) - g(x)|$ where $S$ is some compact set, and compares against continuous functions. Unfortunately, already if the target is Lipschitz continuous, this means our function class needs complexity which scales exponentially with dimension (Luxburg and Bousquet 2004): this highlights the need for more refined target functions and approximation measures.

*(Lower bounds.)* The uniform norm has certain nice properties for proving upper bounds, but is it meaningful for a lower bound? Functions can be well-separated in uniform norm even if they are mostly the same: they just need one point of large difference. For this reason, $L_1$ norms, for instance $\int_{[0,1]^d} |f(x) - g(x)| \, dx$ are preferred for lower bounds.

**Remark 1.2** While norms have received much recent attention as a way to measure complexity, this idea is quite classical. For instance, a resurgence of interest in the 1990s led to the proof of many deep network VC dimension bounds, however very quickly it was highlighted (and proved) in (P. L. Bartlett 1996) that one has situations where the architecture (and connection cardinality) stays fixed (along with the VC dimension), yet the norms (and generalization properties) vary.

### 1.1 Omitted topics

- Full proofs for sobolev space approximation (Yarotsky 2016; Schmidt-Hieber 2017). [mjtΩ: Planning to add in Fall 2021!!!]
- Approximation of distributions and other settings.
- Approximation power of low-norm functions.

### 2 Classical approximations and “universal approximation”

We start with two types of standard approximation results, in the “classical” regime where we only care about the number of nodes and not the magnitude of the weights, and also the worst-case goal of competing with an arbitrary continuous function using some function space norm.

1. Elementary folklore results: univariate approximation with one hidden layer, and multivariate approximation with two hidden layers, just by stacking bricks. Latter use $L_1$ metric, which is disappointing.

2. Celebrated “universal approximation” result: fitting continuous functions over compact sets in uniform norm with a single hidden layer (Hornik, Stinchcombe, and White 1989).

There are weaknesses in these results (e.g., curse of dimension), and thus they are far from the practical picture. Still, they are very interesting and influential.
2.1 Elementary folklore constructions

We can handle the univariate case by gridding the line and taking steps appropriately.

**Proposition 2.1** Suppose \( g : \mathbb{R} \to \mathbb{R} \) is \( \rho \)-Lipschitz. For any \( \epsilon > 0 \), there exists a 2-layer network \( f \) with \( \left\lceil \frac{\epsilon}{\rho} \right\rceil \) threshold nodes \( z \mapsto 1[z \geq 0] \) so that \( \sup_{x \in [0,1]} |f(x) - g(x)| \leq \epsilon \).

**Proof.** Define \( m := \left\lceil \frac{\epsilon}{\rho} \right\rceil \) and \( x_i := (i-1)\epsilon/\rho \), and Construct \( \sum_i a_i 1[x - b_i] \) with
\[
a_1 = g(0), a_i = g(x_i) - g(x_{i-1}), b_i := x_i.
\]
Then for any \( x \in [0,1] \), pick the closest \( x_i \leq x \), and note
\[
|g(x) - f(x)| = |g(x) - f(x_i)| \leq |g(x) - g(x_i)| + |g(x_i) - f(x_i)|
\]
\[
= \rho(\epsilon/\rho) + |g(x_i) - g(x_0) - \sum_{j=2}^{i} (g(x_j) - g(x_{j-1})) 1[x_i \geq x_j]| = \epsilon.
\]

**Remark 2.1** This is standard, but we’ve lost something! We are paying for flat regions, which are a specialty of standard networks! A more careful proof only steps when it needs to and pays in total variation.

Now let’s handle the multivariate case. We will replicate the univariate approach: we will increment function values when the target function changes. In the univariate case, we could “localize” function modifications, but in the multivariate case by default we will modify an entire halfspace at once. To get around this, we use an additional layer.

**Remark 2.2**
- Note the problem is easy for finite point-sets: can reduce to univariate apx after projection onto a random line (homework 1?) But our goal is approximation over a distribution of points.
- We will not get any nice theorem that says, roughly: “the exact complexity of shallow approximation depends on this function of the first \( O(d) \) derivatives” (see also (Yarotsky 2016) for the deep case). This is part of why I like discussing the univariate case, where we have nice characterizations with total variation distance.

**Theorem 2.1** Let continuous \( g : \mathbb{R}^d \to \mathbb{R} \) and an \( \epsilon > 0 \) be given, and choose \( \delta > 0 \) so that \( \|x - x'\|_{\infty} \leq \delta \) implies \( |g(x) - g(x')| \leq \epsilon \). Then there exists a 3-layer network \( f \) with \( \Omega(\frac{1}{\delta^d}) \) ReLU with \( \int_{[0,1]^d} |f(x) - g(x)| dx \leq 2\epsilon \).

**Remark 2.3**
- Note the curse of dimension (exponential dependence on \( d \), which also appears in lower bounds (Luxburg and Bousquet 2004)). Note CIFAR has \( d = 3072 \). This issue is inherent in approximating arbitrary continuous functions, and makes this irrelevant in practice.
- Construction also has large weights and Lipschitz constant.
- Later in Theorem 2.3 ((Hornik, Stinchcombe, and White 1989)) we’ll give another approach that controls \( \sup_{x} |f(x) - g(x)| \) and uses only one activation layer, but it will not be a constructive proof, and trying to obtain estimates from it has all the preceding weaknesses as well.
The proof uses the following lemma (omitted in class), approximating continuous functions by piecewise constant functions.

**Lemma 2.1** Let $g, \delta, \epsilon$ be given as in the theorem. For any partition $\mathcal{P}$ of $[0, 1]^d$ into rectangles (products of intervals) $\mathcal{P} = (R_1, \ldots, R_N)$ with all side lengths not exceeding $\delta$, there exist scalars $(\alpha_1, \ldots, \alpha_N)$ so that

$$\sup_{x \in [0, 1]^d} |g(x) - h(x)|_u \leq \epsilon, \quad \text{where} \quad h = \sum_{i=1}^N \alpha_i \mathbf{1}_{R_i}.$$ 

**Proof.** Let partition $\mathcal{P} = (R_1, \ldots, R_N)$ be given, and for each $R_i$, pick some $x_i \in R_i$, and set $\alpha_i := g(x_i)$. Since each side length of each $R_i$ is at most $\delta$,

$$\sup_{x \in [0, 1]^d} |g(x) - h(x)| = \sup_{i \in \{1, \ldots, N\}} \sup_{x \in R_i} |g(x) - h(x)|$$

$$\leq \sup_{i \in \{1, \ldots, N\}} \sup_{x \in R_i} (|g(x) - g(x_i)| + |g(x_i) - h(x)|)$$

$$\leq \sup_{i \in \{1, \ldots, N\}} \sup_{x \in R_i} (\epsilon + |g(x_i) - \alpha_i|) = \epsilon.$$

**Proof of Theorem 2.1.** We will use the function $h = \sum_i \alpha_i \mathbf{1}_{R_i}$ from the earlier lemma. Specifically, we will use the first two layers to approximate $x \mapsto \mathbf{1}_{R_i}(x)$ for each $i$ using $O(d)$ nodes, and a final linear layer for the linear combination. Writing $\|f - g\|_1 = \int_{[0, 1]^d} |f(x) - g(x)| dx$ for convenience, since

$$\|f - g\|_1 \leq \|f - h\|_1 + \|h - g\|_1 \leq \epsilon + \|h - g\|_1,$$

and letting $g_i$ denote the approximation to $\mathbf{1}_{R_i}$,

$$\|h - g\|_1 = \left\| \sum_i \alpha_i (\mathbf{1}_{R_i} - g_i) \right\|_1 \leq \sum_i |\alpha_i| \cdot \|\mathbf{1}_{R_i} - g_i\|_1,$$

so it suffices to make $\|\mathbf{1}_{R_i} - g_i\|_1 \leq \sum_i |\alpha_i|$. (If $\sum_i |\alpha_i| = 0$, we can set $g$ to be the constant 0 network.)

Let’s do what we did in the univariate case, putting nodes where the function value changes. For each $R_i := \times_{j=1}^d [a_j, b_j]$, and any $\gamma > 0$, define

$$g_{\gamma,j}(z) := \sigma \left( \frac{z - (a_j - \gamma)}{\gamma} \right) - \sigma \left( \frac{z - a_j}{\gamma} \right) - \sigma \left( \frac{z - b_j}{\gamma} \right) + \sigma \left( \frac{z - (b_j + \gamma)}{\gamma} \right)$$

and $g_{\gamma}(x) := \sigma(\sum_j g_{\gamma,j}(x_j) - (d - 1))$ (adding the additional ReLU layer is the key step!), whereby

$$g_{\gamma}(x) = \begin{cases} 
1 & x \in R_i, \\
0 & x \not\in \times_{j=1}^d [a_j - \gamma, b_j + \gamma], \\
[0, 1] & \text{otherwise.}
\end{cases}$$

Since $g_{\gamma} \to \mathbf{1}_{R_i}$ pointwise, there exists $\gamma$ with $\|g_{\gamma} - \mathbf{1}_{R_i}\|_1 \leq \sum_i |\alpha_i|$.
Universal approximation with a single hidden layer

The proof of Theorem 2.1 developed \( g_\gamma \) such that \( g_\gamma(x) \approx 1 \) \( x \in \times_i [a_i, b_i] \).

If deep networks could multiply, we could do

\[
x \mapsto \prod_i 1_{x_i \in [a_i, b_i]}
\]

Can deep networks multiply and form a linear combination, all with a single hidden layer?

The answer will be yes, and we will use this to resolve the classical universal approximation question with a single hidden layer.

**Definition 2.1** A class of functions \( \mathcal{F} \) is a universal approximator over a compact set \( S \) if for every continuous function \( g \) and target accuracy \( \epsilon > 0 \), there exists \( f \in \mathcal{F} \) with

\[
\sup_{x \in S} |f(x) - g(x)| \leq \epsilon.
\]

**Remark 2.4** Typically we will take \( S = [0, 1]^d \). We can also consider more general universal approximation settings. Lastly, the compactness is in a sense necessary: as in the homework, consider approximating the \( \sin \) function with a finite-size ReLU network over all of \( \mathbb{R} \). Lastly, there are also ways to phrase universal approximation in terms of denseness, but we won’t use it here.

Consider unbounded width networks with one hidden layer:

\[
\mathcal{F}_{\sigma,d,m} := \mathcal{F}_{d,m} := \left\{ x \mapsto a^T \sigma(W x + b) : a \in \mathbb{R}^m, W \in \mathbb{R}^{m \times d}, b \in \mathbb{R}^m \right\},
\]

\[
\mathcal{F}_{\sigma,d} := \bigcup_{m \geq 0} \mathcal{F}_{\sigma,d,m}.
\]

**Remark 2.5** Classical perspective: \( \mathcal{F}_d \) is the linear span of functions computed by single-node networks \( (\mathcal{F}_{m,1} \text{ with } a_1 = 1) \).

Consider \( \sigma = \cos \). Since \( 2 \cos(y) \cos(z) = \cos(y + z) + \cos(y - z) \),

\[
2 \left[ \sum_{i=1}^m a_i \cos(w_i^T x + b_i) \right] \cdot \left[ \sum_{j=1}^n c_j \cos(u_j^T x + v_j) \right] = \sum_{i=1}^m \sum_{j=1}^n a_i c_j \cos((w_i + u_j)^T x + (b_i + v_j)) + \cos((w_i - u_j)^T x + (b_i - v_j)),
\]

thus \( f, g \in \mathcal{F}_{\cos,d} \implies fg \in \mathcal{F}_{\cos,d} \)!

This gives us “bumps” via

\[
x \mapsto \prod_{i=1}^d \cos(x_i)^r,
\]

and we can linearly combine bumps to get continuous functions.

Where does this leave us?
• **Polynomials** are also closed under addition and multiplication, and they are universal approximators (Weierstrass 1885). (“Universal apx” means apx continuous functions over compact sets for all $\epsilon > 0$.)

• An extension, the “Stone-Weierstrass theorem,” says “polynomial-like” classes of functions also work.

• These “polynomial-like” properties are satisfied by $\mathcal{F}_{\cos, d}$.

• Since $\cos$ can be typically approximated by $\mathcal{F}_{\sigma,1}$, we also have $\mathcal{F}_{\sigma, d} \approx \mathcal{F}_{\cos, d} \approx \text{cont.}$ (Probably formalized in homework 1.)

**Theorem 2.2 (Stone-Weierstrass; (Folland 1999, Theorem 4.45))** Let functions $\mathcal{F}$ be given as follows.

1. Each $f \in \mathcal{F}$ is continuous.
2. For every $x$, there exists $f \in \mathcal{F}$ with $f(x) \neq 0$.
3. For every $x \neq x'$ there exists $f \in \mathcal{F}$ with $f(x) \neq f(x')$ ($\mathcal{F}$ separates points).
4. $\mathcal{F}$ is closed under multiplication and vector space operations ($\mathcal{F}$ is an algebra).

Then for every continuous $g : \mathbb{R}^d \to \mathbb{R}$ and $\epsilon > 0$, there exists $f \in \mathcal{F}$ with $\|f - g\|_u \leq \epsilon$. ($\mathcal{F}$ is universal.)

**Remark 2.6**

• It is heavyweight, but a good tool to have.

• Proofs are not constructive, but seem to require size $\Omega(\frac{1}{\epsilon^d})$.

• Proofs are interesting:
  
  – We will revisit the standard one due to Bernstein, which picks a fine grid and interpolating polynomials that are well-behaved off the grid.

  – Weierstrass’s original proof convolved the target with a Gaussian, which makes it analytic, and also leads to good polynomial approximation.

• As a technical point, we could also approximately satisfy the properties, and apply the theorem to the closure of $\mathcal{F}$.

• The second and third conditions are necessary; if there exists $x$ so that $f(x) = 0$ for $f \in \mathcal{F}$, then we can’t approximate $g$ with $g(x) \neq 0$; if we can’t separate points $x \neq x'$, then we can’t approximate functions with $g(x) \neq g(x')$.

[ mjt@: I used to have some remark about different ways to prove weierstrass? put that back in?]

**Lemma 2.2 ((Hornik, Stinchcombe, and White 1989))** $\mathcal{F}_{\cos, d}$ is universal.
Proof. Let’s check the Stone-Weierstrass conditions:
1. Each \( f \in \mathcal{F}_{\cos,d} \) is continuous.
2. For each \( x \), \( \cos(0^\top x) = 1 \neq 0 \).
3. For each \( x \neq x' \), \( f(z) := \cos((z - x')^\top (x - x')/\|x - x'\|^2) \in \mathcal{F}_d \) satisfies
   \[
   f(x) = \cos(1) \neq \cos(0) = f(x').
   \]
4. \( \mathcal{F}_{\cos,d} \) is closed under products and vector space operations as before.

We can work it out even more easily for \( \mathcal{F}_{\exp,d} \).

Lemma 2.3 \( \mathcal{F}_{\exp,d} \) is universal.

Proof. Let’s check the Stone-Weierstrass conditions:
1. Each \( f \in \mathcal{F}_{\exp,d} \) is continuous.
2. For each \( x \), \( \exp(0^\top x) = 1 \neq 0 \).
3. For each \( x \neq x' \), \( f(z) := \exp((z - x')^\top (x - x')/\|x - x'\|^2) \in \mathcal{F}_d \) satisfies
   \[
   f(x) = \exp(1) \neq \exp(0) = f(x').
   \]
4. \( \mathcal{F}_{\exp,d} \) is closed under VS ops by construction; for products,
   \[
   \left( \sum_{i=1}^{n} r_i \exp(a_i^\top x) \right) \left( \sum_{j=1}^{m} s_j \exp(b_j^\top x) \right) = \sum_{i=1}^{m} \sum_{j=1}^{m} r_i s_j \exp((a + b)^\top x).
   \]

Now let’s handle arbitrary activations.

Theorem 2.3 (Hornik, Stinchcombe, and White 1989) Suppose \( \sigma : \mathbb{R} \to \mathbb{R} \) is continuous, and
\[
\lim_{z \to -\infty} \sigma(z) = 0, \quad \lim_{z \to +\infty} \sigma(z) = 1.
\]

Then \( \mathcal{F}_{\sigma,d} \) is universal.

Proof sketch. Given \( \epsilon > 0 \) and continuous \( g \), pick \( h \in \mathcal{F}_{\cos,d} \) (or \( \mathcal{F}_{\exp,d} \)) with \( \sup_{x \in [0,1]^d} |h(x) - g(x)| \leq \epsilon/2 \). To finish, replace all appearances of \( \cos \) with an element of \( \mathcal{F}_{\sigma,1} \). (Details in hw1.)

Remark 2.7

- Sometimes this condition on \( \sigma \) is called “sigmoidal.”
- ReLU is fine: use \( z \mapsto \sigma(z) - \sigma(z - 1) \) and split nodes.
- exp didn’t need bias in the proof, but this seems natural due to \( \exp(a^\top x + b) = e^b \cdot \exp(a^\top x) \).
  On the other hand, approximating exp with ReLU uses bias terms, so we don’t obtain a trick from exp to remove biases in general.
- Weakest conditions on \( \sigma \) (Leshno et al. 1993): universal apx iff not a polynomial.
- Carefully accounting within the proof seems to indicate curse of dimension again (size \( \Omega(1/\epsilon^d) \)), due for instance to expanding all terms in a product of \( d \) terms.

Remark 2.8 (other universal approximation proofs)
• (Cybenko 1989) Assume contradictorily you miss some functions. By duality, $0 = \int \sigma(a^T x - b) d\mu(x)$ for some signed measure $\mu$, all $(a,b)$. Using Fourier, can show this implies $\mu = 0$.

• (Leshno et al. 1993) If $\sigma$ a polynomial, …; else can (roughly) get derivatives and polynomials of all orders (we’ll have homework problems on this).

• (Barron 1993) Use inverse Fourier representation to construct an infinite-width network; we’ll cover this next. It can beat the worst-case curse of dimension!

• (Funahashi 1989) 

3 Infinite-width Fourier representations and the Barron norm

Under construction.

4 Approximation near initialization and the Neural Tangent Kernel (NTK)

Under construction.

5 Benefits of depth

Under construction.

6 Optimization: preface

Classically, the purpose of optimization is to approximately minimize (or maximize) an objective function $f$ over a domain $S$:

$$\min_{w \in S} f(w).$$

A core tension in the use of optimization in machine learning is that we would like to minimize the population risk $R(w) := \mathbb{E} \ell(Y f(X; w))$; however, we only have access to the empirical risk $\hat{R}(w) := n^{-1} \sum_i \ell(y_i, f(x_i; w))$.

As a result, when choosing a $w_t$, we not only care that $\hat{R}(w_t)$ is small, but also other good properties which may indicate $R(w_t)$ is small as well. Foremost amongst these are that $w_t$ has low norm, but there are other possibilities.

Outline.

• We will cover primarily first-order methods, namely gradient descent

$$w_{t+1} := w_t - \eta_t \nabla \hat{R}(w_t),$$

as well as the gradient flow

$$\frac{dw}{dt} = \dot{w}(t) = -\nabla \hat{R}(w(t)).$$
These dominate machine learning since:

- They have low per-iteration complexity (which can be reduced further with stochastic gradients); classical optimization developed many methods with higher per-iteration cost but a lower number of iterations, but the high accuracy these give is not important here since our true objective is unknown anyway.

- It seems they might have additional favorable properties; e.g., we will highlight the preference for low-norm solutions of first-order methods.

- First we’ll cover classical smooth and convex opt, including strong convexity and stochastic gradients.

Here our analysis differs from the literature by generally not requiring boundedness or existence of minima. Concretely, many proofs will use an arbitrary reference point \( z \) in place of an optimum \( \hat{w} \) (which may not exist); this arbitrary \( z \) will be used effectively in the margin maximization lectures.

- Then we will cover topics closer to deep learning, including gradient flow in a smooth shallow NTK case, and a few margin maximization cases, with a discussion of nonsmoothness.

**Remark 6.1**

- Even though our models are not convex (and \( \hat{R} \) is not convex in the parameters), our losses will always be convex.

- Analyzing gradient flow simplifies analyses, but in some cases it is difficult or completely unclear how to reproduce the same rates with gradient descent, and secondly it isn’t clear that they should have the same rates or convergence properties; in deep learning, for instance, the role of step size is not well-understood, whereas approximating gradient flow suggests small step sizes.

- A regularized ERM objective has the form \( w \mapsto \hat{R}(w) + P(w) \), where (for example) \( P(w) := \lambda \|w\|^2/2 \). We will not discuss these extensively, and we will similarly hardly discuss constrained optimization.

- A good introductory text on various optimization methods in machine learning is (Bubeck 2014); for more on convex optimization, see for instance (Nesterov 2003), and for more on convex analysis, see for instance (Bubeck 2014; Borwein and Lewis 2000).

[ mjt@: ...maybe I should always use \( \hat{R} \) or \( F \) for objectives]

### 6.1 Omitted topics

- **Mean-field perspective** (Chizat and Bach 2018; Mei, Montanari, and Nguyen 2018): as \( m \to \infty \), gradient descent mimics a Wasserstein flow on a distribution over nodes (random features). Many mean-field papers are in the 2-homogeneous case, whereas many NTK papers are in the 1-homogeneous case, which further complicates comparisons.

- **Landscape analysis.** (E.g., all local optima are global.)
  - Matrix completion: solve (under RIP)
    \[
    \min_{X \in \mathbb{R}^{d \times r}} \sum_{(i,j) \in S} (M_{i,j} - XX^\top)^2.
    \]
Recently it was shown that all local optima are global, and so gradient descent from random initialization suffices (Ge, Lee, and Ma 2016).

- For linear networks optimized with the squared loss, local optima are global, but there are bad saddle points (Kawaguchi 2016).
- Width $n$ suffices with general losses and networks (Nguyen and Hein 2017).
- [ There is also work on residual networks but I haven’t looked closely. ]

**Acceleration.** Consider gradient descent with momentum: $w_0$ arbitrary, and thereafter

$$v_{i+1} := w_i - \eta_i \nabla \mathcal{R}(w_i), \quad w_{i+1} := v_{i+1} + \gamma_i (v_{i+1} - v_i)$$

This sometimes seems to help in deep learning (even in stochastic case), but no one knows why (and opinions differ).

If set $\eta_i = 1/\beta$ and $\gamma_i = i/(i + 3)$ (constants matter) and $\mathcal{R}$ convex, $\mathcal{R}(w_i) - \inf_w \mathcal{R}(w) \leq \mathcal{O}(1/t^2)$ (“Nesterov’s accelerated method”). This rate is tight amongst algorithms outputting iterates in the span of gradients, under some assumptions people treat as standard.

**Escaping saddle points.** By adding noise to the gradient step, it is possible to exit saddle points (Jin et al. 2017). Some papers use this technique, though it is most useful in settings where all local minima (stationary points that are not saddles) are global minima.

**Beyond NTK.** A very limited amount of work studies nonlinear cases beyond what is possible with the NTK and/or highlighting ways in which the NTK does not capture the behavior of deep networks in practice, in particular showing sample complexity separations (Allen-Zhu and Li 2019; Daniely and Malach 2020; Ghorbani et al. 2020; Kamath, Montasser, and Srebro 2020; Yehudai and Shamir 2019, 2020).

**Benefits of depth for optimization.** Most of these works are either for shallow networks, or the analysis allows depth but degrades with increasing depth, in contrast with practical observations. A few works now are trying to show how depth can help optimization; one perspective is that sometimes it can accelerate convergence (Arora, Cohen, and Hazan 2018; Arora, Cohen, et al. 2018a).

**Other first-order optimizers,** e.g., Adam. There is recent work on these but afaik it doesn’t capture why these work well on many deep learning tasks.

**Further analysis of overparameterization.** Overparameterization makes many aspects of the optimization problem nicer, in particular in ways not investigated in these notes (Shamir 2018; S. Du and Hu 2019).

**Hardness of learning and explicit global solvers.** Even in simple cases, network training is NP-hard, but admits various types of approximation schemes (Goel et al. 2020; Diakonikolas et al. 2020).

### 7 Smooth objectives in ML

We say “$\mathcal{R}$ is $\beta$-smooth” to mean $\beta$-Lipschitz gradients:

$$\| \nabla \mathcal{R}(w) - \nabla \mathcal{R}(v) \| \leq \beta \| w - v \|.$$
(The math community says “smooth” for $C^\infty$.)

- We primarily invoke smoothness via the key inequality

$$\hat{R}(v) \leq \hat{R}(w) + \left\langle \nabla \hat{R}(w), v - w \right\rangle + \frac{\beta}{2} \|v - w\|^2.$$ 

In words: $f$ can be upper bounded with the convex quadratic

$$v \mapsto \frac{\beta}{2} \|v - w\|^2 + \left\langle \nabla \hat{R}(w), v - w \right\rangle + \hat{R}(w)$$

which shares tangent and function value with $\hat{R}$ at $w$. (The first definition also implies that we are lower bounded by concave quadratics.)

**Remark 7.1** Smoothness is trivially false for standard deep networks: the ReLU is not even differentiable. However, many interesting properties carry over, and many lines of research proceed by trying to make these properties carry over, so at the very least, it’s good to understand.

A key consequence: we can guarantee gradient descent does not increase the objective.

Consider gradient iteration $w' = w - \frac{1}{\beta} \nabla \hat{R}(w)$, then smoothness implies

$$\hat{R}(w') \leq \hat{R}(w) - \left\langle \nabla \hat{R}(w), \hat{R}(w)/\beta \right\rangle + \frac{1}{2\beta} \|\nabla \hat{R}(w)\|^2 = \hat{R}(w) - \frac{1}{2\beta} \|\nabla \hat{R}(w)\|^2,$$

and $\|\nabla \hat{R}(w)\|^2 \leq 2\beta(\hat{R}(w) - \hat{R}(w'))$.

With deep networks, we’ll produce similar bounds but in other ways.

As an exercise, let’s prove the earlier smoothness consequence. Considering the curve $t \mapsto \hat{R}(w + t(v - w))$ along $[0, 1]$,

$$\left| \hat{R}(v) - \hat{R}(w) - \left\langle \nabla \hat{R}(w), v - w \right\rangle \right|$$

$$= \left| \int_0^1 \left\langle \nabla \hat{R}(w + t(v - w)), v - w \right\rangle dt - \left\langle \nabla \hat{R}(w), v - w \right\rangle \right|$$

$$\leq \int_0^1 \left| \left\langle \nabla \hat{R}(w + t(v - w)) - \nabla \hat{R}(w), v - w \right\rangle \right| dt$$

$$\leq \int_0^1 \|\nabla \hat{R}(w + t(v - w)) - \nabla \hat{R}(w)\| \cdot \|v - w\| dt$$

$$\leq \int_0^1 t\beta\|v - w\|^2 dt$$

$$= \frac{\beta}{2} \|v - w\|^2.$$ 

**Example 7.1** Define $\hat{R}(w) := \frac{1}{2} \|Xw - y\|^2$, and note $\nabla \hat{R}(w) = X^T(Xw - y)$. For any $w, w'$,

$$\hat{R}(w') = \frac{1}{2} \|Xw' - Xw + Xw - y\|^2$$

$$= \frac{1}{2} \|Xw' - Xw\|^2 + \langle Xw' - Xw, Xw - y \rangle + \frac{1}{2} \|Xw - y\|^2$$

$$= \frac{1}{2} \|Xw' - Xw\|^2 + \left\langle w' - w, \hat{R}(w) \right\rangle + \hat{R}(w).$$
Since $\frac{\sigma_{\min}(X)}{2} \| w' - w \|^2 \leq \frac{1}{2} \| X w' - X w \|^2 \leq \frac{\sigma_{\max}(X)}{2} \| w' - w \|^2$, thus $\hat{R}$ is $\sigma_{\max}(X)$-smooth (and $\sigma_{\min}$-strongly-convex, as we’ll discuss).

The smoothness bound holds with equality if we use the seminorm $\| v \|_X = \| X v \|$. We’ll discuss smoothness wrt other norms in homework.

[ mjtm: I should use $L$ not $R$ since unnormalized.]

### 7.1 Convergence to stationary points

Consider first the gradient iteration

$$w' := w - \eta \nabla \hat{R}(w),$$

where $\eta \geq 0$ is the step size. When $f$ is $\beta$ smooth but not necessarily convex, the smoothness inequality directly gives

$$\hat{R}(w') \leq \hat{R}(w) + \langle \nabla \hat{R}(w), w' - w \rangle + \frac{\beta}{2} \| w' - w \|^2$$

$$= \hat{R}(w) - \eta \| \nabla \hat{R}(w) \|^2 + \frac{\beta \eta^2}{2} \| \nabla \hat{R}(w) \|^2$$

$$= \hat{R}(w) - \eta \left(1 - \frac{\beta \eta}{2}\right) \| \nabla \hat{R}(w) \|^2.$$

If we choose $\eta$ appropriately ($\eta \leq 2/\beta$) then: either we are near a critical point ($\nabla \hat{R}(w) \approx 0$), or we can decrease $f$.

Let’s refine our notation to tell iterates apart:

1. Let $w_0$ be given.
2. Recurse: $w_i := w_{i-1} - \eta_i \nabla \hat{R}(w_{i-1})$.

[ mjtm: am I consistent with indexing? is $\eta_i$ always with $w_{i-1}$?]

Rearranging our iteration inequality and summing over $i < t$,

$$\sum_{i<t} \eta_{i+1} \left(1 - \frac{\beta \eta_{i+1}}{2}\right) \| \nabla \hat{R}(w_i) \|^2 \leq \sum_{i<t} \left(\hat{R}(w_i) - \hat{R}(w_{i+1})\right)$$

$$= \hat{R}(w_0) - \hat{R}(w_t).$$

We can summarize these observations in the following theorem.

**Theorem 7.1** Let $(w_i)_{i \geq 0}$ be given by gradient descent on $\beta$-smooth $f$.

- If $\eta_{i+1} \in [0, 2/\beta]$, then $\hat{R}(w_{i+1}) \leq \hat{R}(w_i)$.
- If $\eta_i := \eta \in [0, 2/\beta]$ is constant across $i$,

$$\min_{i < t} \| \nabla \hat{R}(w) \|^2 \leq \frac{1}{t} \sum_{i<t} \| \nabla \hat{R}(w) \|^2 \leq \frac{2}{t \eta (2 - \eta \beta)} \left(\hat{R}(w_0) - \hat{R}(w_t)\right)$$

$$\leq \frac{2}{t \eta (2 - \eta \beta)} \left(\hat{R}(w_0) - \inf_w \hat{R}(w)\right).$$
Remark 7.2

- In lecture I set $\eta = \frac{1}{\beta}$, whereby
  \[
  \frac{2}{t\eta(2-\eta\beta)} = \frac{2\beta}{t}.
  \]

- We have no guarantee about the last iterate $\|\nabla \hat{R}(w_t)\|$; we may get near a flat region at some $i < t$, but thereafter bounce out. With a more involved proof, we can guarantee we bounce out (Lee et al. 2016), but there are cases where the time is exponential in dimension.

- This derivation is at the core of many papers with a “local optimization” (stationary point or local optimum) guarantee for gradient descent.

- The gradient iterate with step size $1/\beta$ is the result of minimizing the quadratic provided by smoothness:
  \[
  w - \frac{1}{\beta} \nabla \hat{R}(w) = \arg \min_{w'} \left( \hat{R}(w) + \langle \nabla \hat{R}(w), w' - w \rangle + \frac{\beta}{2} \|w' - w\|^2 \right)
  \]
  \[
  = \arg \min_{w'} \left( \langle \nabla \hat{R}(w), w' \rangle + \frac{\beta}{2} \|w' - w\|^2 \right).
  \]
  This relates to proximal descent and mirror descent generalizations of gradient descent.

- In $t$ iterations, we found a point $w$ with $\|\nabla \hat{R}(w)\| \leq \sqrt{2\beta/t}$. We can do better with Nesterov-Polyak cubic regularization: by choosing the next iterate according to
  \[
  \arg \min_{w'} \left( \hat{R}(w) + \langle \nabla \hat{R}(w), w' - w \rangle + \frac{1}{2} \langle \nabla^2 \hat{R}(w)(w' - w), w' - w \rangle + \frac{L}{6} \|w' - w\|^3 \right)
  \]
  where $\|\nabla^2 \hat{R}(x) - \nabla^2 \hat{R}(y)\| \leq L \|x - y\|$, then after $t$ iterations, some iterate $w_j$ with $j \leq t$ satisfies
  \[
  \|\nabla \hat{R}(w_j)\| \leq \mathcal{O}(1/\sqrt[3]{t}), \quad \lambda_{\min} \left( \nabla^2 \hat{R}(w_j) \right) \geq -\mathcal{O}(1/\sqrt[3]{t}).
  \]
  Note: it is not obvious that the above cubic can be solved efficiently, but indeed there are various ways. If we go up a few higher derivatives, it becomes NP-hard. Original used an eigenvalue solver for this cubic polynomial (Nesterov and Polyak 2006). Other approaches are given by (Carmon and Duchi 2018; Jin et al. 2017), amongst many others.

Gradient flow version. Using FTC, chain rule, and definition,
\[
\hat{R}(w(t)) - \hat{R}(w(0)) = \int_0^t \langle \nabla \hat{R}(w(s)), \dot{w}(s) \rangle \, ds
\]
\[
= -t \int_0^t \|\nabla \hat{R}(w(s))\| \, ds
\]
\[
\leq -t \inf_{s \in [0,t]} \|\nabla \hat{R}(w(s))\|^2,
\]
which can be summarized as follows.
**Theorem 7.2** For the gradient flow, 

\[
\inf_{s \in [0,t]} \| \nabla \hat{R}(w(s)) \|^2 \leq \frac{1}{t} \left( \hat{R}(w(0)) - \hat{R}(w(t)) \right).
\]

**Remark 7.3** GD: \( \min_{i<t} \| \nabla \hat{R}(w) \|^2 \leq \frac{2\beta}{t} \left( \hat{R}(w_0) - \hat{R}(w_t) \right) \).

- \( \beta \) is from step size.
- “2” is from the order smoothness term (avoided in GF).

### 7.2 Convergence rate for smooth & convex

**Theorem 7.3** Suppose \( \hat{R} \) is \( \beta \)-smooth and convex, and \( (w_i)_{i \geq 0} \) given by GD with \( \eta_i := 1/\beta \). Then for any \( z \),

\[
\hat{R}(w_i) - \hat{R}(z) \leq \frac{\beta}{2t} \left( \| w_0 - z \|^2 - \| w_t - z \|^2 \right).
\]

**Remark 7.4** We only invoke convexity via the inequality

\[
\hat{R}(w') \geq \hat{R}(w) + \langle \nabla \hat{R}(w), w' - w \rangle,
\]

meaning \( f \) lies above all tangents. [mjt@: I should give a summary of convexity characterizations. after all, I do it for strong convexity...]

The reference point \( z \) allows us to use this bound effectively when \( \hat{R} \) lacks an optimum, or simply when the optimum is very large. For an example of such an application of \( z \), see the margin maximization material.

**Proof.** By convexity and the earlier smoothness inequality \( \| \nabla \hat{R}(w) \|^2 \leq 2\beta(\hat{R}(w) - \hat{R}(w')) \),

\[
\| w' - z \|^2 = \| w - z \|^2 - \frac{2}{\beta} \langle \nabla \hat{R}(w), w - z \rangle + \frac{1}{\beta^2} \| \nabla \hat{R}(w) \|^2 \\
\leq \| w - z \|^2 + \frac{2}{\beta} (\hat{R}(z) - \hat{R}(w)) + \frac{2}{\beta} (\hat{R}(w) - \hat{R}(w')) \\
= \| w - z \|^2 + \frac{2}{\beta} (\hat{R}(z) - \hat{R}(w')).
\]

Rearranging and applying \( \sum_{i<t} \),

\[
\frac{2}{\beta} \sum_{i<t} (\hat{R}(w_{i+1}) - \hat{R}(z)) \leq \sum_{i<t} \left( \| w_i - z \|^2 - \| w_{i+1} - z \|^2 \right)
\]

The final bound follows by noting \( \hat{R}(w_i) \geq \hat{R}(w_i) \), and since the right hand side telescopes.

For GF, we use the same potential, but indeed start from the telescoping sum, which can be viewed
as a Riemann sum corresponding to the following application of FTC:

\[
\frac{1}{2} \|w(t) - z\|_2^2 - \frac{1}{2} \|w(0) - z\|_2^2 = \frac{1}{2} \int_0^t \frac{d}{ds} \|w(s) - z\|_2^2 ds
\]

\[
= \int_0^t \left\langle \frac{dw}{ds}, w(s) - z \right\rangle ds
\]

\[
\geq \int_0^t \left( \tilde{R}(w(s)) - \tilde{R}(z) \right) ds.
\]

**Theorem 7.4** For any \( z \in \mathbb{R}^d \), GF satisfies

\[
t \tilde{R}(w(t)) + \frac{1}{2} \|w(t) - z\|_2^2 \leq \int_0^t \tilde{R}(w(s)) + \frac{1}{2} \|w(0) - z\|_2^2
\]

\[
= t \tilde{R}(z) + \frac{1}{2} \|w(0) - z\|_2^2.
\]

**Remark 7.5 ("units" of GD and GF: \( t \) vs \( t/\beta \))** Here’s a back-of-the-envelope calculation to see why \( t \) becomes \( t/\beta \) and why they are really the same, and not a sloppiness of the analysis.

- Suppose \( \|\nabla \tilde{R}(w)\| \approx 1 \) for sake of illustration.
- The “distance traveled” by GD is

\[
\|w_t - w_0\| = \frac{1}{\beta} \sum_i \nabla \tilde{R}(w_i) \leq \sum_i \frac{1}{\beta} \|\nabla \tilde{R}(w_i)\| \approx \frac{t}{\beta}.
\]

- The “distance traveled” by GF is (via Jensen)

\[
\|w(t) - w(0)\| = \| \int_0^t \nabla \tilde{R}(w(s)) ds \| = \frac{1}{t} \int_0^t \| \nabla \tilde{R}(w(s)) \| ds \|
\]

\[
\leq \frac{1}{t} \int_0^t \| t \nabla \tilde{R}(w(s)) \| ds \approx t.
\]

**Remark 7.6 (potential functions)**

- For critical points, the potential was \( \tilde{R}(w(s)) \) (or arguably \( \|\nabla \tilde{R}(w(s))\|_2^2 \)).
- Here, the potential was \( \|w(s) - z\|_2^2 \). This particular choice is widespread in optimization. It is interesting since it is not part of the objective function; it’s some gradient descent magic?

We can use similar objective functions with deep learning, without smoothness (!).

**Remark 7.7 (rates)** Some rules of thumb (not comprehensive, and there are other ways).

- \( \frac{1}{t} \) is often a smoothness argument as above.
- \( \frac{1}{\sqrt{t}} \) uses Lipschitz (thus \( \|\nabla \tilde{R}\| = O(1) \)) in place of smoothness upper bound on \( \|\nabla \tilde{R}\| \).
- \( \frac{1}{t^2} \) uses “acceleration,” which is a fancy momentum inside the gradient.
- \( \exp(-O(t)) \) uses strong convexity (or other fine structure on \( \tilde{R} \)).
- Stochasticity changes some rates and what is possible, but there are multiple settings and inconsistent terminology.
Here is a sort of companion to Lipschitz gradients; a stronger condition than convexity which will grant much faster convergence rates. (Convexity references I recommend: (Hiriart-Urruty and Lemaréchal 2001; Borwein and Lewis 2000).)

Say that \( R \) is \( \lambda \)-strongly-convex (\( \lambda \)-sc) when
\[
R(w') \geq R(w) + \langle \nabla R(w), w' - w \rangle + \frac{\lambda}{2} \| w' - w \|^2.
\]

Some alternative definitions:

- When twice differentiable, \( \nabla^2 R \succeq \lambda I \) (\( \beta \)-smooth implies \( -\beta I \preceq \nabla^2 R \preceq \beta I \)).
- When differentiable, \( \langle \nabla R(w) - \nabla R(w'), w - w' \rangle \geq \lambda \| w - w' \|^2 \) (\( \beta \)-smooth gives \( \leq \beta \| w - w' \|^2 \)).
- \( \hat{R} \) is \( \lambda \)-sc iff \( \hat{R} - \| \cdot \|_2^2 / 2 \) is convex.
- Definitions in terms of subgradients and function values also exist.

**Example 8.1 (least squares)** Earlier we pointed out
\[
\frac{1}{2} \| Xw' - w' \|^2 =: \hat{R}(w') = \hat{R}(w) + \langle \nabla \hat{R}(w), w' - w \rangle + \frac{1}{2} \| Xw' - Xw \|^2
\]
and
\[
\sigma_{\text{min}}(X) \| w' - w \|^2 \leq \| Xw' - Xw \|^2 \leq \sigma_{\text{max}}(X) \| w' - w \|^2.
\]

The latter implies a smoothness upper bound we used, now we know the former implies strong convexity. (We can also say that both hold with equality using the special seminorm \( \| v \|_X = \| Xv \| \).) We can also verify these properties by noting \( \nabla^2 \hat{R} = X^T X \).

**Example 8.2 (regularization)** Define regularized risk \( \hat{R}_\lambda(w) := \hat{R}(w) + \lambda \| w \|^2 / 2 \).

If \( \hat{R} \) is convex, then \( \hat{R}_\lambda \) is \( \lambda \)-sc:

- A quick check is that if \( f \) is twice-differentiable, then \( \nabla^2 \hat{R}_\lambda = \nabla^2 \hat{R} + \lambda I \succeq 0 + \lambda I \).
- Alternatively, it also follows by summing the inequalities
\[
\hat{R}(w') \geq \hat{R}(w) + \langle \nabla \hat{R}(w), w' - w \rangle, \quad \lambda \| w' \|^2 / 2 = \lambda \| w \|^2 / 2 + \langle \lambda w, w' - w \rangle + \lambda \| w' - w \|^2 / 2.
\]

Another very useful property is that \( \lambda \)-sc gives a way to convert gradient norms to suboptimality.

**Lemma 8.1** Suppose \( \hat{R} \) is \( \lambda \)-sc. Then
\[
\forall w, \quad \hat{R}(w) - \inf_v \hat{R}(v) \leq \frac{1}{2\lambda} \| \nabla \hat{R}(w) \|^2.
\]

**Remark 8.1** Smoothness gave \( \frac{1}{2\beta} \| \nabla \hat{R}(w_i) \|^2 \leq \hat{R}(w_i) - \hat{R}(w_{i+1}) \).
Proof. Let \( w \) be given, and define the convex quadratic
\[
Q_w(v) := \tilde{R}(w) + \langle \nabla \tilde{R}(w), v - w \rangle + \frac{\lambda}{2} \|v - w\|^2,
\]
which attains its minimum at \( \tilde{v} := w - \nabla \tilde{R}(w) / \lambda \). By definition \( \lambda \)-sc,
\[
\inf_v \tilde{R}(v) \geq \inf_v Q_w(v) = Q_w(\tilde{v}) = \tilde{R}(w) - \frac{1}{2\lambda} \|\nabla \tilde{R}(w)\|^2.
\]

Remark 8.2 (stopping conditions) Say our goal is to find \( w \) so that \( f(w) - \inf_v \tilde{R}(v) \leq \epsilon \).
When do we stop gradient descent?

- The \( \lambda \)-sc case is easy: by the preceding lemma, we know that we can stop when \( \|\nabla \tilde{R}(w)\| \leq \sqrt{2\lambda} \epsilon \).
- Another easy case is when \( \inf_v \tilde{R}(v) \) is known, and we just watch \( \tilde{R}(w_t) \). E.g., in classification tasks, deep networks are expect to get 0. For things like deep RL, once again it becomes a problem.
- Many software packages use heuristics. Some people just run their methods as long as possible. In convex cases, sometimes we can compute duality gaps.

Remark 8.3 (Regularization and boundedness)
- Given \( \tilde{R}_\lambda(w) = \tilde{R}(w) + \lambda \|w\|^2/2 \) with \( \tilde{R} \geq 0 \), optimal point \( \tilde{w} \) satisfies
\[
\frac{\lambda}{2} \|\tilde{w}\|^2 \leq \tilde{R}_\lambda(\tilde{w}) \leq \tilde{R}_\lambda(0) = \tilde{R}(0),
\]
thus it suffices to search over bounded set \( \{w \in \mathbb{R}^p : \|w\|^2 \leq 2\tilde{R}(0)/\lambda\} \). This can often be plugged directly into generalization bounds.
- In deep learning, this style of regularization (“weight decay”) is indeed used, but it isn’t necessary for generalization, and is much smaller than what many generalization analyses suggest, and thus its overall role is unclear.

[mjt*: I should lemmas lemmas giving level set containment, and existence of minimizers.]

8.1 Rates when strongly convex and smooth

Theorem 8.1 Suppose \( f \) is \( \lambda \)-sc and \( \beta \)-smooth, and GD is run with step size \( 1/\beta \). Then a minimum \( \tilde{w} \) exists, and
\[
\tilde{R}(w_t) - \tilde{R}(\tilde{w}) \leq \left( \tilde{R}(w_0) - \tilde{R}(\tilde{w}) \right) \exp(-t\lambda/\beta),
\]
\[
\|w_t - \tilde{w}\|^2 \leq \|w_0 - \tilde{w}\|^2 \exp(-t\lambda/\beta).
\]
Proof. Using previously-proved Lemmas from smoothness and strong convexity,

\[
\begin{align*}
\hat{R}(w_{i+1}) - \hat{R}(\bar{w}) &\leq \hat{R}(w_i) - \hat{R}(\bar{w}) - \frac{\|\nabla \hat{R}(w_i)\|^2}{2\beta} \\
&\leq \hat{R}(w_i) - \hat{R}(\bar{w}) - \frac{2\lambda(\hat{R}(w_i) - \hat{R}(\bar{w}))}{2\beta} \\
&\leq \left(\hat{R}(w_i) - \hat{R}(\bar{w})\right) \left(1 - \lambda/\beta\right),
\end{align*}
\]

which gives the first bound by induction since

\[
\prod_{i<t} \left(1 - \lambda/\beta\right) \leq \prod_{i<t} \exp(-\lambda/\beta) = \exp(-t\lambda/\beta).
\]

For the second guarantee, expanding the square as usual,

\[
\begin{align*}
\|w' - \bar{w}\|^2 &= \|w - \bar{w}\|^2 + \frac{2}{\beta} \left(\langle \nabla \hat{R}(w), \bar{w} - w \rangle + \frac{1}{\beta^2} \|\nabla \hat{R}(w)\|^2\right) \\
&\leq \|w - \bar{w}\|^2 + \frac{2}{\beta} \left(\hat{R}(\bar{w}) - \hat{R}(w) - \frac{\lambda}{2} \|\bar{w} - w\|^2\right) \\
&\quad + \frac{1}{\beta^2} \left(2\beta(\hat{R}(w) - \hat{R}(w'))\right) \\
&= (1 - \lambda/\beta)\|w - \bar{w}\|^2 + \frac{2}{\beta} \left(\hat{R}(\bar{w}) - \hat{R}(w) + \hat{R}(w) - \hat{R}(w')\right) \\
&\leq (1 - \lambda/\beta)\|w - \bar{w}\|^2,
\end{align*}
\]

which gives the argument after a similar induction argument as before.

<table>
<thead>
<tr>
<th>Remark 8.4</th>
</tr>
</thead>
<tbody>
<tr>
<td>- $\beta/\lambda$ is sometimes called the condition number, based on linear system solvers, where it is $\sigma_{\text{max}}(X)/\sigma_{\text{min}}(X)$ as in least squares. Note that $\beta \geq \lambda$ and a good condition numbers improves these bounds.</td>
</tr>
<tr>
<td>- Setting the bounds to $\epsilon$, it takes a linear number of iterations to learn a linear number of bits of $\bar{w}$.</td>
</tr>
<tr>
<td>- Much of the analysis we’ve done goes through if the norm pair $(| \cdot |_2, | \cdot |<em>2)$ is replaced with $(| \cdot |, | \cdot |</em>*)$ where the latter dual norm is defined as</td>
</tr>
<tr>
<td>$|s|_* = \sup {\langle s, w \rangle : |w| \leq 1}$;</td>
</tr>
<tr>
<td>for instance, we can define $\beta$-smooth wrt $| \cdot |$ as</td>
</tr>
<tr>
<td>$|\nabla \hat{R}(w) - \nabla \hat{R}(w')|_* \leq \beta|w - w'|$.</td>
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</tbody>
</table>

Next let’s handle the gradient flow.

| Theorem 8.2 | If $\hat{R}$ is $\lambda$-sc, a minimum $\bar{w}$ exists, and the GF $w(t)$ satisfies |
|-------------| |
| $\|w(t) - \bar{w}\|^2 \leq \|w(0) - \bar{w}\|^2 \exp(-2\lambda t)$, |
| $\hat{R}(w(t)) - \hat{R}(\bar{w}) \leq \left(\hat{R}(w(0)) - \hat{R}(\bar{w})\right) \exp(-2t\lambda)$. |
Proof. By first-order optimality in the form $\nabla \bar{R}(\bar{w}) = 0$, then
\[
\frac{d}{dt} \frac{1}{2} \|w(t) - \bar{w}\|^2 = \langle w(t) - \bar{w}, \dot{w}(t) \rangle \\
= -\langle w(t) - \bar{w}, \nabla \bar{R}(w(t)) - \nabla \bar{R}(\bar{w}) \rangle \\
\leq -\lambda \|w(t) - \bar{w}\|^2.
\]

By Grönwall’s inequality, this implies
\[
\|w(t) - \bar{w}\|^2 \leq \|w(0) - \bar{w}\|^2 \exp\left(-\int_0^t 2\lambda ds\right) \\
\leq \|w(0) - \bar{w}\|^2 \exp(-2\lambda t),
\]
which establishes the guarantee on distances to initialization. For the objective function guarantee,
\[
\frac{d}{dt} (\bar{R}(w(t)) - \bar{R}(\bar{w})) = \langle \nabla \bar{R}(w(t)), \dot{w}(t) \rangle \\
= -\|\nabla \bar{R}(w(t))\|^2 \leq -2\lambda (\bar{R}(w(t)) - \bar{R}(\bar{w})).
\]

Grönwall’s inequality implies
\[
\bar{R}(w(t)) - \bar{R}(\bar{w}) \leq (\bar{R}(w(0)) - \bar{R}(\bar{w})) \exp(-2t\lambda).
\]

Remark 8.5 As in all other rates proved for GF and GD, time $t$ is replaced by “arc length units” $t/\beta$.

We have strayed a little from our goals by producing laborious proofs that not only separate the objective function and the distances, but also require minimizers. Interestingly, we can resolve this by changing the step size to a large (seemingly worse?) one.

Theorem 8.3 Suppose $\bar{R}$ is $\beta$-smooth and $\lambda$-sc, and a constant step size $\frac{2}{\beta + \lambda}$. Then, for any $z$,
\[
\bar{R}(w_t) - \bar{R}(z) + \lambda \frac{1}{2} \|w_t - z\|^2 \leq \left[\frac{\beta - \lambda}{\beta + \lambda}\right]^t \left(\bar{R}(w_0) - \bar{R}(z) + \lambda \frac{1}{2} \|w_0 - z\|^2\right).
\]

Proof. Homework problem ⋯.

Remark 8.6 (standard rates with strong convexity) Compared with standard proofs in the literature (Nesterov 2003, chap. 2), the preceding bound with step size $2/($ $\beta + \lambda$) is possibly loose: it seems possible to have a $2t$ and not just $t$ in the exponent, albeit after adjusting the other terms (and depending explicitly on minimizers). [ mjt#: I need to resolve what’s going on here⋯]

Moreover, another standard rate given in the literature is $1/t$ under just strong convexity (no smoothness); however, this requires a step size $\eta_t := (\lambda(i + 1))^{-1}$.

9 Stochastic gradients
Let’s generalize gradient descent, and consider the iteration

\[ w_{i+1} := w_i - \eta g_i, \]

where each \( g_i \) is merely some vector. If \( g_i := \nabla \mathcal{R}(w_i) \), then we have gradient descent, but in general we only approximate it. Later in this section, we’ll explain how to make \( g_i \) a “stochastic gradient.”

Our first step is to analyze this in our usual way with our favorite potential function, but accumulating a big error term:

\[
\|w_{i+1} - z\|^2 = \|w_i - \eta g_i - z\|^2
\]

\[
= \|w_i - z\|^2 - 2\eta_i \langle g_i, w_i - z \rangle + \eta_i^2 \|g_i\|^2
\]

\[
\leq \|w_i - z\|^2 + 2\eta \langle R(z) - R(w_i), w_i - z \rangle + \eta^2 \|g_i\|^2,
\]

which after rearrangement gives

\[
2\eta R(w_i) \leq 2\eta R(z) + \|w_i - z\|^2 - \|w_{i+1} - z\|^2 + 2\eta \epsilon_i + \eta^2 \|g_i\|^2,
\]

and applying \( \frac{1}{2\eta t} \sum_{i \leq t} \) to both sides gives

\[
\frac{1}{t} \sum_{i \leq t} R(w_i) \leq R(z) + \frac{\|w_0 - z\|^2 - \|w_t - z\|^2}{2\eta t} + \frac{1}{t} \sum_{i \leq t} \epsilon_i.
\]

The following lemma summarizes this derivation.

**Lemma 9.1** Suppose \( \mathcal{R} \) convex; set \( G := \max_i \|g_i\|_2 \), and \( \eta := \frac{c}{\sqrt{t}} \). For any \( z \),

\[
\mathcal{R}\left( \frac{1}{t} \sum_{i \leq t} w_i \right) \leq \frac{1}{t} \sum_{i \leq t} R(w_i) \leq R(z) + \frac{\|w_0 - z\|^2}{2ct} + \frac{cG^2}{2\sqrt{t}} + \frac{1}{t} \sum_{i \leq t} \epsilon_i.
\]

**Proof.** This follows from the earlier derivation after plugging in \( G, \eta = c/\sqrt{t} \), and applying Jensen’s inequality to the left hand side.

**Remark 9.1**

- We get a bound on the averaged iterate or a minimum iterate, but not the last iterate! (We’ll revisit this later.) Averaged iterates are often suggested in theory, but rare in applied classification (afaik), but I’ve heard of them used in deep RL; OTOH, averaging seems weird with nonconvexity?

- \( \eta = c/\sqrt{t} \) trades off between terms. If \( t \) not fixed in advance, can use \( \eta_i = c/\sqrt{1 + i} \), but I’d rather shorten lecture a little by avoiding the needed algebra with non-constant step sizes, and for deep learning at least this style seems to not work well.

- This analysis works fine with \( \nabla \mathcal{R}(w_i) \) replaced with subgradient \( s_i \in \partial \mathcal{R}(w_i) \).
• Suppose $\|\nabla R(w_i)\| \leq G$ and set $D := \max_i \|w_i - z\|$, then by Cauchy-Schwarz

$$\frac{1}{t} \sum_{i < t} \epsilon_i \leq \frac{1}{t} \sum_{i < t} \langle g_i - \nabla R(w_i), w_i - z \rangle \leq 2GD,$$

which does not go to 0 with $t$! Thus more structure needed on $\epsilon_i$, this worst-case argument is bad.

• This proof easily handles projection to convex closed sets: replace $w_i - \eta g_i$ with $\Pi_S(w_i - \eta g_i)$, and within the proof use the non-expansive property of $\Pi_S$. This can be used to ensure that $D$ up above is not too large. (We’ll return to this point.)

Now let us define the standard stochastic gradient oracle:

$$\mathbb{E}[g_i|w_{\leq i}] = \nabla R(w_i),$$

where $w_{\leq i}$ signifies all randomness in $(w_1, \ldots, w_i)$.

<table>
<thead>
<tr>
<th>Remark 9.2</th>
</tr>
</thead>
<tbody>
<tr>
<td>• We can’t use an unconditional expectation because gradient at $w_i$ should rely upon random variable $w_i$!</td>
</tr>
<tr>
<td>• One way to satisfy this: sample $(x, y)$, and set $g_i := l'(yf(x; w_i))y\nabla_w f(x; w_i)$; conditioned on $w_{\leq i}$, the only randomness is in $(x, y)$, and the conditional expectation is a gradient over the distribution!</td>
</tr>
</tbody>
</table>

Indeed, this setup allows the expectation to be nicely interpreted as an iterated integral over $(x_1, y_1)$, then $(x_2, y_2)$, and so on. The stochastic gradient $g_i$ depends on $(x_i, y_i)$ and $w_i$, but $w_i$ does not depend on $(x_i, y_i)$, rather on $((x_j, y_j))_{j=1}^{i-1}$.

• It’s standard to sample a minibatch and average the $g_i$ obtained from each, which ostensibly has the same conditional mean as $g_i$, but improved variance. It can be hard to analyze this.

• Stochastic minibatch gradient descent is standard for deep networks. However, there is a delicate interplay between step size, minibatch size, and number of training epochs (Shallue et al. 2018).

• Annoyingly, there are many different settings for stochastic gradient descent, but they refer to themselves in the same way and it requires a closer look to determine the precise setting.

• Previous slide suggested $(x, y)$ is a fresh sample from the distribution; in this case, we are doing stochastic gradient descent on the population directly!

• We can also resample the training set, in which case $R$ is our usual empirical risk, and now the randomness is under our control (randomized algorithm, not random data from nature). The “SVRG/SDCA/SAG/etc” papers are in this setting, as are some newer SGD papers. Since people typically do multiple passes over the time, perhaps this setting makes sense.

• There are many deep learning papers that claim SGD does miraculous things to the optimization process. Unfortunately, none of these seem to come with a compelling and general theoretical analysis. Personally I don’t know if SGD works further miracles (beyond computational benefits), but it’s certainly interesting!
Now let’s work towards our goal of showing that, with high probability, our stochastic gradient method does nearly as well as a regular gradient method. (We will not show any benefit to stochastic noise, other than computation!)

Our main tool is as follows.

**Theorem 9.1 (Azuma-Hoeffding)** Suppose \((Z_i)_{i=1}^n\) is a martingale difference sequence \((\mathbb{E}(Z_i | Z_{<i}) = 0)\) and \(\mathbb{E} |Z_i| \leq R\). Then with probability at least \(1 - \delta\),

\[
\sum_i Z_i \leq R \sqrt{2t \ln(1/\delta)}.
\]

**Proof omitted**, though we’ll sketch some approaches in a few weeks.

We will use this inequality to handle \(\sum_{i<t} \epsilon_i\). Firstly, we must show the desired expectations are zero. To start,

\[
\mathbb{E} \left[ \epsilon_i \bigg| w_{\leq i} \right] = \mathbb{E} \left[ \langle g_i - \nabla \mathcal{R}(w_i), z - w_i \rangle \bigg| w_{\leq i} \right]
= \langle \mathbb{E} \left[ g_i - \nabla \mathcal{R}(w_i) \bigg| w_{\leq i} \right], z - w_i \rangle
= \langle 0, z - w_i \rangle
= 0.
\]

Next, by Cauchy-Schwarz and the triangle inequality,

\[
\mathbb{E} |\epsilon_i| = \mathbb{E} \left| \langle g_i - \nabla \hat{R}(w_i), w_i - z \rangle \right| \leq \mathbb{E} \left( \|g_i\| + \|\nabla \hat{R}(w_i)\| \right) \|w_i - z\| \leq 2GD.
\]

Consequently, by Azuma-Hoeffding, with probability at least \(1 - \delta\),

\[
\sum_i \epsilon_i \leq 2GD \sqrt{2t \ln(1/\delta)}.
\]

Plugging this into the earlier approximate gradient lemma gives the following. [mjt\#: should give explicit cref]

**Lemma 9.2** Suppose \(\mathcal{R}\) convex; set \(G := \max_i \|g_i\|_2\), and \(\eta := \frac{1}{\sqrt{t}}, D \geq \max_i \|w_i - z\|\), and suppose \(g_i\) is a stochastic gradient at time \(i\). With probability at least \(1 - \delta\),

\[
\mathcal{R} \left( \frac{1}{t} \sum_{i<t} w_i \right) \leq \frac{1}{t} \sum_{i<t} \mathcal{R}(w_i)
\leq \mathcal{R}(z) + \frac{G^2}{2\sqrt{t}} + \frac{2DG \sqrt{2 \ln(1/\delta)}}{\sqrt{t}}.
\]

**Remark 9.3**

- If we tune \(\eta = c/\sqrt{t}\) here, we only get a \(DG\) term. [mjt\#: I should do it]
- We can ensure \(D\) is small by projecting to a small set each iteration. By the contractive property of projections, the analysis still goes through.
- By the tower property of conditional expectation, meaning \(\mathbb{E} = \mathbb{E}\mathbb{E}[: |w_{\leq i}]\), without Azuma-Hoeffding we easily get a bound on the expected average error:

\[
\mathbb{E} \left[ \frac{1}{t} \sum_{i<t} \mathcal{R}(w_i) \right] \leq \mathcal{R}(z) + \frac{\|w_0 - z\|^2}{2\sqrt{t}} + \frac{G^2}{2\sqrt{t}}.
\]
• If the preceding bound in expectation is sufficient, expected is enough, a more careful
analysis lets us use the last iterate (Shamir and Zhang 2013); AFAIK a high probability
version still doesn't exist.

• The Martingale structure is delicate: if we re-use even a single data-point, then we can’t
treat $\mathcal{R}$ as the population risk, but instead as the empirical risk. [mjt: and here my
notation is truly frustrating.]

• In practice, randomly sampling a permutation over the training data at the beginning of
each epoch is common; it can be hard to analyze.

• Why SGD in ML? In statistical problems, we shouldn’t expect test error better than
$\frac{1}{\sqrt{n}}$ or $\frac{1}{n}$ anyway, so we shouldn’t optimize to crazy accuracy. With SGD, the per-iteration
cost is low. Meanwhile, heavyweight solvers like Newton methods require a massive
per-iteration complexity, with the promise of crazy accuracy; but, again we don’t need
that crazy accuracy here. [mjt: summarize as “computation.”]

10 NTK-based Gradient flow analysis on smooth shallow networks, following (Chizat and Bach 2019)

Finally we will prove (rather than assert) that we can stay close to initialization long enough to
get a small risk with an analysis that is essentially convex, essentially following the NTK (Taylor
approximation).

• This proof is a simplification of one by Chizat and Bach (2019). There are enough differences
that it’s worth checking the original.

  – That paper highlights a “scaling phenomenon” as an explanation of the NTK. Essentially,
    increasing with always decreases initialization variance, and the paper argues this
    corresponds to “zooming in” on the Taylor expansion in function space, and flattening
    the dynamics.

  – This “scaling perspective” pervades much of the NTK literature and I recommend looking
    at (Chizat and Bach 2019) for further discussion; I do not discuss it much in this course
    or even in this proof, though I keep Chizat’s $\alpha > 0$ scale parameter.

• This proof comes after many earlier NTK analyses, e.g., (Jacot, Gabriel, and Hongler 2018;
Simon S. Du et al. 2018; Allen-Zhu, Li, and Liang 2018; Arora, Du, et al. 2019). I like the
proof by (Chizat and Bach 2019) very much and learned a lot from it; it was the most natural
for me to teach. OTOH, it is quite abstract, and we’ll need homework problems to boil it
down further.

Basic notation. For convenience, bake the training set into the predictor:

$$f(w) := \begin{bmatrix} f(x_1; w) \\ \vdots \\ f(x_n; w) \end{bmatrix} \in \mathbb{R}^n.$$ 

We’ll be considering squared loss regression:

$$\hat{\mathcal{R}}(\alpha f(w)) := \frac{1}{2}\|\alpha f(w) - y\|^2, \quad \hat{\mathcal{R}}_0 := \hat{\mathcal{R}}(\alpha f(w(0))),$$

29
where $\alpha > 0$ is a scale factor we’ll optimize later. [mjtodef: maybe I should use $\mathcal{L}$ not $\hat{\mathcal{L}}$ since unnormalized.]

We’ll consider gradient flow:

$$\dot{w}(t) := -\nabla_w \hat{\mathcal{L}}(\alpha f(w(t))) = -\alpha J_t^T \nabla \hat{\mathcal{L}}(\alpha f(w(t))),$$

where $J_t := J_w(t) := \begin{bmatrix} \nabla f(x_1; w(t))^T \\ \vdots \\ \nabla f(x_n; w(t))^T \end{bmatrix} \in \mathbb{R}^{n \times p}.$

We will also explicitly define and track a flow $u(t)$ over the tangent model; what we care about is $w(t)$, but we will show that indeed $u(t)$ and $w(t)$ stay close in this setting. (Note that $u(t)$ is \textit{not} needed for the analysis of $w(t)$.)

$$f_0(u) := f(w(0)) + J_0(u - w(0)).$$

$$\dot{u}(t) := -\nabla_u \hat{\mathcal{L}}(\alpha f_0(u(t))) = -\alpha J_0^T \nabla \hat{\mathcal{L}}(\alpha f_0(u(t))).$$

Both gradient flows have the same initial condition:

$$u(0) = w(0), \quad f_0(u(0)) = f_0(w(0)) = f(w(0)).$$

\textbf{Remark 10.1 (initialization, width, etc)}

- Notice that the setup so far doesn’t make any mention of width, neural networks, random initialization, etc! It’s all abstracted away! This is good and bad: the good is that it highlights the “scale” phenomenon, as $\alpha$ is the only concretely interpretable parameter here. On the downside, we need to do some work to get statements about width etc.

\textbf{Assumptions.}

$$\text{rank}(J_0) = n,$$

$$\sigma_{\min} := \sigma_{\min}(J_0) = \sqrt{\lambda_{\min}(J_0 J_0^T)} = \sqrt{\lambda_n(J_0 J_0^T)} > 0,$$

$$\sigma_{\max} := \sigma_{\max}(J_0) > 0,$$

$$\|J_w - J_v\| \leq \beta \|w - v\|.$$  \hfill (2)

\textbf{Remark 10.2 ($J_0 J_0^T$ has full rank, a “representation assumption”)} This is a “representation assumption” in an explicit sense: it implies the tangent model has exact solutions to the least squares problem, regardless of the choice of $y$, meaning the training error can always be made 0. In detail, consider the least squares problem solved by the tangent space:

$$\min_{u \in \mathbb{R}^p} \frac{1}{2} \|f_0(u) - y\|^2 = \min_{u \in \mathbb{R}^p} \frac{1}{2} \| J_0 u - y_0 \|^2,$$

where we have chosen $y_0 := y + J_0 w(0) - f(w(0))$ for convenience. The normal equations for this least squares problem are

$$J_0^T J_0 u = J_0^T y_0.$$

Let $J_0 = \sum_{i=1}^n s_i u_i v_i^T$ denote the SVD of $J_0$, which has $n$ terms by the rank assumption; the corresponding pseudoinverse is $J_0^T = \sum_{i=1}^n s_i^{-1} v_i u_i^T$. Multiplying both sides by $(J_0^T)^\dagger$,

$$J_0 u = (J_0^T)^\dagger J_0^T J_0 u = (J_0^T)^\dagger J_0^T y_0 = \left[ \sum_{i=1}^n u_i u_i^T \right] y_0 = y_0.$$
where the last step follows since $[\sum_i u_iu_i^T]$ is idempotent and full rank, and therefore the identity matrix. In particular, we can choose $\hat{u} = J_0^Ty_0$, then $J_0\hat{u} = [\sum_i u_iu_i^T]y_0 = y_0$, and in particular
\[
\frac{1}{2} \|f_0(\hat{u}) - y\|^2 = \frac{1}{2} \|J_0\hat{u} - y_0\|^2 = 0.
\]
As such, the full rank assumption is explicitly a representation assumption: we are forcing the tangent space least squares problem to always have solutions.

**Theorem 10.1 (see also (Theorem 3.2, Chizat and Bach 2019))** Assume $\alpha \geq \frac{\beta \sqrt{1152\sigma^2_{\max} R_0}}{\sigma_{\min}^2}$. Then
\[
\max \left\{ \mathcal{R}(\alpha f(w(t))), \mathcal{R}(\alpha f_0(u(t))) \right\} \leq \mathcal{R}_0 \exp(-t\alpha^2 \sigma_{\min}^2/2),
\]
\[
\max \{ \|w(t) - w(0)\|, \|u(t) - u(0)\| \} \leq \frac{3\sqrt{8\sigma^2_{\max} R_0}}{\alpha \sigma_{\min}^2}.
\]

**Remark 10.3 (shallow case)** To get a handle on the various abstract constants and what they mean, consider the shallow case, namely $f(x; w) = \sum_j s_j \sigma(w_j^T x)$, where $s_j \in \{\pm 1\}$ is not trained, and each $w_j$ is trained.

**Smoothness constant.** Let $X \in \mathbb{R}^{n \times d}$ be a matrix with the $n$ training inputs as rows, and suppose $\sigma$ is $\beta_0$-smooth. Then
\[
\|J_0 - J_\sigma\|^2 = \sum_{i,j} \|x_i\|^2 (\sigma'(w_j^T x_i) - \sigma'(v_j^T x_i))^2
\]
\[
\leq \sum_{i,j} \|x_i\|^2 4\beta_0^2 \|w_j - v_j\|^2
\]
\[
= \beta_0^2 \|X\|_F^2 \|w - v\|^2.
\]
Thus $\beta \leq \beta_0 \|X\|_F^2$ suffices, which we can ballpark as $\beta = \Theta(n)$.

**Singular values.** Now that we have an interpretation of the full rank assumption, ballpark the eigenvalues of $J_0J_0^T$. By definition,
\[
(J_0J_0^T)_{i,j} = \nabla f(x_i; w(0))^T \nabla f(x_j; w(0)).
\]
Holding $i$ fixed and letting $j$ vary, we can view the corresponding column of $(J_0J_0^T)$ as another feature representation, and rank($J_0$) = $n$ means none of these examples, in this feature representation, are linear combinations of others. This gives a concrete sense under which these eigenvalue assumptions are representation assumptions.

Now suppose each $w_j(0)$ is an iid copy of some random variable $v$. Then, by definition of $J_0$,
\[
\mathbb{E}_{w(0)} (J_0J_0^T)_{i,j} = \mathbb{E}_{w(0)} \nabla f(x_i; w(0))^T \nabla f(x_j; w(0)) = \mathbb{E}_{w(0)} \sum_k s_k^2 \sigma'(w_k(0)^T x_i) \sigma'(w_k(0)^T x_j) x_i^T x_j
\]
\[
= \mathbb{E}_v \sigma'(v^T x_i) \sigma'(v^T x_j) x_i^T x_j.
\]
In other words, it seems reasonable to expect $\sigma_{\min}$ and $\sigma_{\max}$ to scale with $\sqrt{m}$.
Initial risk $\tilde{R}_0$. Let’s consider two different random initializations.

In the first case, we use one of the fancy schemes we mentioned to force $f(w(0)) = 0$; e.g., we can make sure that $s_j$ is positive and negative an equal number of times, then sample $w_j$ for $s_j = +1$, and then make $w_j$ for $s_j = -1$ be the negation. With this choice, $\tilde{R}_0 = \|y\|^2/2 = \Theta(n)$.

On the other hand, if we do a general random initialization of both $s_j$ and $w_j$, then we can expect enough cancellation that, roughly, $f(x_i; w(0)) = \Theta(\sqrt{m})$ (assuming $w_j$’s variance is a constant and not depending on $m$: that would defeat the purpose of separating out the scale parameter $\alpha$). then $\|\alpha f(w(0))\|^2 = \Theta(\sqrt{m})$, and $\tilde{R}_0 = \Theta(\alpha^2 mn)$, and thus the lower bound condition on $\alpha$ will need to be checked carefully.

Combining all parameters. Again let’s split into two cases, based on the initialization as discussed immediately above.

- The case $\tilde{R}_0 = \Theta(\alpha^2 mn)$. Using $\beta = \Theta(n)$, the condition on $\alpha$ indeed has $\alpha$ on both sides, and becomes

$$\sigma^3_{\min} \geq \Omega(\beta \sigma_{\max} \sqrt{nm}) = \sigma_{\max} \Omega(\sqrt{nm^3}).$$

Since we said the singular values are of order $\sqrt{m}$, we get roughly $m^{3/2} \geq \sqrt{m^2 n^3}$, thus $m \geq n^3$.

Since the lower bound on $\alpha$ turned into a lower bound on $m$, let’s plug this $\tilde{R}_0$ into the rates to see how they simplify:

$$\max \left\{ \tilde{R}(\alpha f(w(t))), \tilde{R}(\alpha f_0(u(t))) \right\} \leq \tilde{R}_0 \exp \left(-\frac{t \alpha^2 \sigma^2_{\min}}{2}\right),$$

$$= \mathcal{O} \left( \frac{\alpha^2 nm \exp \left(-\frac{t \alpha^2 \sigma^2_{\min}}{2}\right)}{\sigma^2_{\min}} \right),$$

$$\max \{ \|w(t) - w(0)\|, \|u(t) - w(0)\| \} \leq \frac{3 \sqrt{8 \sigma^2_{\max} \tilde{R}_0}}{\alpha \sigma^2_{\min}}$$

$$= \mathcal{O} \left( \frac{\sqrt{\sigma^2_{\max} mn}}{\sigma^2_{\min}} \right).$$

In these inequalities, the distance to initialization is not affected by $\alpha$: this makes sense, as the key work needed by the gradient flow is to clear the initial noise so that $y$ can be fit exactly. Meanwhile, the empirical risk rate does depend on $\alpha$, and is dominated by the exponential term, suggesting that $\alpha$ should be made arbitrarily large. There is indeed a catch limiting the reasonable choices of $\alpha$, as will be pointed out shortly.

For now, to pick a value which makes the bounds more familiar, choose $\alpha = \hat{\alpha} := 1/\sigma_{\max}$, whereby additionally simplifying via $\sigma_{\min}$ and $\sigma_{\max}$ being $\Theta(\sqrt{m})$ gives

$$\max \left\{ \tilde{R}(\alpha f(w(t))), \tilde{R}(\alpha f_0(u(t))) \right\} = \mathcal{O} \left( \frac{\sigma^{-2}_{\max} nm \exp \left(-\frac{t \sigma^2_{\min}}{2\sigma^2_{\max}}\right)}{2\sigma^2_{\max}} \right),$$

$$= \mathcal{O} \left( \frac{n \exp \left(-\frac{t \sigma^2_{\min}}{2\sigma^2_{\max}}\right)}{2\sigma^2_{\max}} \right),$$

$$\max \{ \|w(t) - w(0)\|, \|u(t) - w(0)\| \} = \mathcal{O} \left( \frac{\sqrt{\sigma^2_{\max} mn}}{\sigma^2_{\min}} \right) = \mathcal{O} \left( \sqrt{n} \right).$$
Written this way, the empirical risk rate depends on the condition number $\sigma_{\text{max}}/\sigma_{\text{min}}$ of the NTK Gram matrix, which is reminiscent of the purely strongly convex and smooth analyses as in Theorem 8.1.

- **The case $\hat{R}_0 = \Theta(n)$**. Using $\beta = \Theta(n)$, the condition on $\alpha$ becomes

$$\alpha = \Omega \left( \frac{\beta \sqrt{\sigma_{\text{max}}^2 \hat{R}_0}}{\sigma_{\text{min}}^3} \right) = \Omega \left( \frac{\sigma_{\text{max}} n^{3/2}}{\sigma_{\text{min}}^3} \right).$$

We have removed the cancelation from the previous case, and are now constrained in our choice of $\alpha$; we can still set $\alpha := 1/\sigma_{\text{max}}$, which after using our estimate of $\sqrt{m}$ for $\sigma_{\text{min}}$ and $\sigma_{\text{max}}$ get a similar requirement $m = \Omega(n^3)$. More generally, we get $\alpha = \Omega(n^{3/2}/m)$, which means for large enough $m$ we can treat as close to $1/m$. [mjtΩ: Frederic Koehler points out that the first case can still look like $\hat{R}_0 = \Theta(\alpha^2 mn + n)$ and even $\Theta(n)$ when $\alpha$ is small; I need to update this story.]

**Possible values of $\alpha$.** The two preceding cases considered lower bounds on $\alpha$. In the case $\hat{R}_0 = \Theta(\alpha^2 mn)$, it even seemed that we can make $\alpha$ whatever we want; in either case, the time required to make $\hat{R}(\alpha f(w(t)))$ small will decrease as $\alpha$ increases, so why not simply make $\alpha$ arbitrarily large?

An issue occurs once we perform time discretization. Below, we will see that the smoothness of the model looks like $\alpha^2 \sigma_{\text{max}}^2$ near initialization; as such, a time discretization, using tools such as in Theorem 7.3, will require a step size roughly $1/(\alpha^2 \sigma_{\text{max}}^2)$, and in particular while we may increase $\alpha$ to force the gradient flow to seemingly converge faster, a smoothness-based time discretization will need the same number of steps.

As such, $\alpha = 1/\sigma_{\text{max}}$ seems a reasonable way to simplify many terms in this shallow setup, which translates into a familiar $1/\sqrt{m}$ NTK scaling.

### 10.1 Proof of Theorem 10.1

**Proof plan.**

- First we choose a fortuitous radius $B := \frac{\sigma_{\text{min}}}{2\beta}$, and seek to study the properties of weight vectors $w$ which are $B$-close to initialization:

$$\|w - w(0)\| \leq B;$$

This $B$ will be chosen to ensure $J_t$ and $J_0$ are close, amongst other things. Moreover, we choose a $T$ so that all $t \in [0,T]$ are in this good regime:

$$T := \inf \{ t \geq 0 : \|w(t) - w(0)\| > B \}.$$

- Now consider any $t \in [0,T]$. [mjtΩ: i should include explicit lemma pointers for each.]

  - First we show that if $J_tJ_t^T$ is positive definite, then we rapidly decrease risk, essentially following our old strong convexity proof.

  - Next, since the gradient of the least squares risk is the residual, then decreasing risk implies decreasing gradient norms, and in particular we can not travel far.
That is to say, in this setting, the original NTK paper, (Jacot, Gabriel, and Hongler 2018), had as its story

Suppose we can also prove this setting implies we stay close to initialization.

The evolution in prediction space is

\[
\frac{d}{dt} \alpha f(w(t)) = \alpha J_\alpha \dot{w}(t) = -\alpha^2 J_0 J_0^T \nabla \hat{R}(\alpha f(w(t))), \\
= -\alpha^2 J_0 J_0^T (\alpha f(w(t)) - y), \\
\]

\[
\frac{d}{dt} \alpha f_0(u(t)) = \frac{d}{dt} \alpha (f(w(0) + J_0(u(t) - w(0))) = \alpha J_0 \dot{u}(t) \\
= -\alpha^2 J_0 J_0^T \nabla \hat{R}(\alpha f_0(u(t))) \\
= -\alpha^2 J_0 J_0^T (\alpha f_0(u(t)) - y).
\]

The first one is complicated because we don’t know how \( J_t \) evolves.

But the second one can be written

\[
\frac{d}{dt} [\alpha f_0(u(t))] = -\alpha^2 (J_0 J_0^T) [\alpha f_0(u(t))] + \alpha^2 (J_0 J_0^T) y,
\]

which is a concave quadratic in the predictions \( \alpha f_0(u(t)) \).

Remark 10.5 The original NTK paper, (Jacot, Gabriel, and Hongler 2018), had as its story that GF follows a gradient in kernel space. Seeing the evolution of \( \alpha f_0(u(t)) \) makes this clear, as it is governed by \( J_0 J_0^T \), the Gram or kernel matrix!

Let’s fantasize a little and suppose \( (J_w J_w)^T \) is also positive semi-definite. Do we still have a nice convergence theory?

Lemma 10.1 Suppose \( \dot{z}(t) = -Q(t) \nabla \hat{R}(z(t)) \) and \( \lambda := \inf_{t \in [0, T]} \lambda_{\text{min}}(Q(t)) > 0. \) Then for any \( t \in [0, T], \)

\[
\hat{R}(z(t)) \leq \hat{R}(z(0)) \exp(-2t\lambda).
\]

Remark 10.6 A useful consequence is

\[
\|z(t) - y\| = \sqrt{2\hat{R}(z(t))} \leq \sqrt{2\hat{R}(z(0)) \exp(-2t\lambda)} = \|z(0) - y\| \exp(-t\lambda).
\]

Proof. Mostly just repeating our old strong convexity steps,

\[
\frac{d}{dt} \frac{1}{2} \|z(t) - y\|^2 = \langle -Q(t)(z(t) - y), z(t) - y \rangle \\
\leq -\lambda_{\text{min}}(Q(t)) \langle z(t) - y, z(t) - y \rangle \\
\leq -2\lambda \|z(t) - y\|^2 / 2,
\]

and Grönwall’s inequality completes the proof.

We can also prove this setting implies we stay close to initialization.
Lemma 10.2 Suppose $\dot{v}(t) = -S(t)^T \nabla \hat{R}(g(v(t)))$, where $S_t S_t^T = Q_t$, and $\lambda_t(Q_t) \in [\lambda, \lambda_1]$ for $[0, \tau]$. Then for $t \in [0, \tau]$,

$$
\|v(t) - v(0)\| \leq \frac{\sqrt{\lambda_1}}{\lambda} \|g(v(0)) - y\| \leq \frac{\sqrt{2\lambda_1 \hat{R}(g(v(0)))}}{\lambda}.
$$

Proof.

$$
\|v(t) - v(0)\| = \left\| \int_0^t \dot{v}(s) ds \right\| \leq \int_0^t \|\dot{v}(s)\| ds
$$

$$
\leq \sqrt{\lambda_1} \int_0^t \|g(v(s)) - y\| ds
\leq \sqrt{\lambda_1} \|g(v(0)) - y\| \int_0^t \exp(-s\lambda) ds
\leq \frac{\sqrt{\lambda_1}}{\lambda} \|g(v(0)) - y\| \leq \frac{\sqrt{2\lambda_1 \hat{R}(g(v(0)))}}{\lambda},
$$

where (\*) used +Lemma 10.1.

Where does this leave us?

We can apply the previous two lemmas to the tangent model $u(t)$, since for any $t \geq 0$,

$$
\dot{u}(t) = -\alpha J_0^T \nabla \hat{R}(\alpha f_0(u(t))), \quad \frac{d}{dt} \alpha f_0(u(t)) = -\alpha^2 (J_0 J_0^T) \nabla \hat{R}(\alpha f_0(u(t))).
$$

Thus since $Q_0 := \alpha^2 J_0 J_0^T$ satisfies $\lambda_t(Q_0) \in \alpha^2 [\sigma_{\min}^2, \sigma_{\max}^2]$,

$$
\hat{R}(\alpha f_0(u(t))) \leq \hat{R}_0 \exp(-2t\alpha^2 \sigma_{\min}^2),
$$

$$
\|u(t) - u(0)\| \leq \frac{\sqrt{2\sigma_{\max}^2 \hat{R}_0}}{\alpha \sigma_{\min}^2}.
$$

How about $w(t)$?

Let’s relate $(J_w J_w^T)$ to $(J_0 J_0^T)$.

Lemma 10.3 Suppose $\|w - w(0)\| \leq B = \frac{\sigma_{\min}}{2\beta}$. Then

$$
\sigma_{\min}(J_w) \geq \sigma_{\min} - \beta \|w - w(0)\|_2 \geq \frac{\sigma_{\min}}{2},
$$

$$
\sigma_{\max}(J_w) \leq \frac{3\sigma_{\max}}{2}.
$$

35
Proof. For the upper bound,
\[ \|J_w\| \leq \|J_0\| + \|J_w - J_0\| \leq \|J_0\| + \beta\|w - w(0)\| \leq \sigma_{\text{max}} + \beta B = \sigma_{\text{max}} + \frac{\sigma_{\text{min}}}{2}. \]

For the lower bound, given vector \( v \) define \( A_v := J_0 v \) and \( B_v := (J_w - J_0)^T v \), whereby
\[ \|A_v\| \geq \sigma_{\text{min}} \|v\|, \quad \|B_v\| \leq \|J_w - J_0\| \cdot \|v\| \leq \beta B \|v\|, \]
and thus
\[ \sigma_{\text{min}}(J_w)^2 = \min_{\|v\| = 1} v^T J_w J_w^T v \]
\[ = \min_{\|v\| = 1} ((J_0 + J_w - J_0)^T v)^T (J_0 + J_w - J_0)^T v \]
\[ = \min_{\|v\| = 1} \|A_v\|^2 + 2A_v^T B_v + \|B_v\|^2 \]
\[ \geq \min_{\|v\| = 1} \|A_v\|^2 - 2\|A_v\| \cdot \|B_v\| + \|B_v\|^2 \]
\[ = \min_{\|v\| = 1} (\|A_v\| - \|B_v\|)^2 \geq \min_{\|v\| = 1} (\sigma_{\text{min}} - \beta B)^2 \|v\|^2 = \left( \frac{\sigma_{\text{min}}}{2} \right)^2. \]

Using this, for \( t \in [0, T] \),
\[ \dot{w}(t) = -\alpha J_w^T \nabla \hat{R}(\alpha f(w(t))), \quad \frac{d}{dt} \alpha f(w(t)) = -\alpha^2 (J_w J_w^T) \nabla \hat{R}(\alpha f(w(t))). \]

Thus since \( Q_t := \alpha^2 J_t J_t^T \) satisfies \( \lambda_0(Q_t) \in [\alpha^2 \sigma_{\text{min}}^2/4, 9\alpha^2 \sigma_{\text{max}}^2/4] \),
\[ \hat{R}(\alpha f(w(t))) \leq \hat{R}_0 \exp(-t\alpha^2 \sigma_{\text{min}}^2/2), \]
\[ \|w(t) - w(0)\| \leq \frac{3\sqrt{8\sigma_{\text{max}}^2 \hat{R}_0}}{\alpha \sigma_{\text{min}}^2} =: B'. \]

It remains to show that \( T = \infty \). Invoke, for the first time, the assumed lower bound on \( \alpha \), namely
\[ \alpha \geq \frac{\beta \sqrt{1152\sigma_{\text{max}}^2 \hat{R}_0}}{\sigma_{\text{min}}^3}, \]
which by the above implies then \( B' \leq B' \). Suppose contradictorily that \( T < \infty \); since \( t \mapsto \|w(t) - w(0)\| \) is also continuous and starts from 0, and therefore \( \|w(T) - w(0)\| = B > 0 \) exactly. But due to the lower bound on \( \alpha \), we also have \( \|w(T) - w(0)\| \leq \frac{B}{2} < B', \) a contradiction.

This completes the proof.

Remark 10.7 (retrospective)

- On the downside, the proof is not only insensitive to benefits of \( w(t) \) over \( u(t) \), moreover the guarantees on \( w(t) \) are a degradation of those on \( u(t) \)! That is to say, this proof does not demonstrate any benefit to the nonlinear model over the linear one.

- Note that \( w(t) \) and \( u(t) \) are close by triangle inequality:
\[ \|w(t) - u(t)\| \leq \|w(t) - w(0)\| + \|u(t) - w(0)\|, \]
\[ \|\alpha f(w(t)) - \alpha f(u(t))\| \leq \|\alpha f(w(t)) - y\| + \|\alpha f(u(t)) - y\|. \]
Smoothness and differentiability do not in general hold for us (ReLU, max-pooling, hinge loss, etc.).

One relaxation of the gradient is the subdifferential set $\partial_s$ (whose elements are called subgradients), namely the set of tangents which lie below the predictor:

$$\partial_s R(w) := \{ s \in \mathbb{R}^p : \forall w' \cdot \widehat{R}(w') \geq \widehat{R}(w) + s^T(w' - w) \}.$$

- If $\widehat{R} : \mathbb{R}^d \to \mathbb{R}$ is convex, then $\partial_s \widehat{R}$ is nonempty everywhere.
- If $\nabla \widehat{R}$ exists and $\widehat{R}$ is convex, then $\partial_s \widehat{R}(w) = \{ \nabla \widehat{R}(w) \}$. [mjt: does this need some continuity on $\widehat{R}$? need to check and provide a reference.]
- Much of convex analysis and convex opt can use subgradients in place of gradients; cf. (Hiriart-Urruty and Lemaréchal 2001; Nesterov 2003). As an example from these notes, Lemma 9.1 can replace gradients with subgradients.

One fun application is a short proof of Jensen’s inequality.

**Lemma 11.1 (Jensen’s inequality)** Suppose random variable $X$ is supported on a set $S$, and $f$ is convex on $S$. Then $\mathbb{E} f(X) \geq f(\mathbb{E} X)$.

**Proof.** Choose any $s \in \partial_s f(\mathbb{E} X)$, and note

$$\mathbb{E} f(X) \geq \mathbb{E} [f(\mathbb{E} X) + s^T(X - \mathbb{E} X)] = f(\mathbb{E} X).$$

Typically, we lack convexity, and the subdifferential set is empty.

Our main formalism is the Clarke differential (Clarke et al. 1998):

$$\partial R(w) := \text{conv} \left( \{ s \in \mathbb{R}^p : \exists w_i \rightarrow w, \nabla \widehat{R}(w_i) \rightarrow s \} \right).$$

**Definition 11.1** $f$ is locally Lipschitz when for every point $x$, there exists a neighborhood $S \ni \{ x \}$ such that $f$ is Lipschitz when restricted to $S$.

Key properties:

- If $\widehat{R}$ is locally Lipschitz, $\partial R$ exists everywhere.
- If $\widehat{R}$ is convex, then $\partial R = \partial_s \widehat{R}$ everywhere. [mjt: need to check some continuity conditions and add a reference.]
- $\widehat{R}$ is continuously differentiable at $w$ iff $\partial R(w) = \{ \nabla R(w) \}$.

We can replace the gradient flow differential equation $\dot{w}(t) = -\nabla R(w(t))$ with a differential inclusion:

$$\dot{w}(t) \in -\partial R(w(t)) \quad \text{for a.e. } t \geq 0.$$
If $R$ satisfies some technical structural conditions, then the following nice properties hold; these properties are mostly taken from (Lemma 5.2, Theorem 5.8, Davis et al. 2018) (where the structural condition is $C^1$ Whitney stratifiability), which was slightly generalized in (Ji and Telgarsky 2020) under o-minimal definability; another alternative, followed in (Lyu and Li 2019), is to simply assume that a chain rule holds.

- **(Chain rule.)** For a.e. $t \geq 0$ and every $v \in \partial \hat{R}(w(t))$, then $\frac{d}{dt} \hat{R}(w(t)) = -\langle v, \dot{w}(t) \rangle$.
  
  This is the key strong property; since it holds for every element $v$ of the Clarke differential simultaneously, it implies the next property.

- **(Minimum norm path.)** For almost every $t \geq 0$, then $\dot{w}(t) = -\arg \min \{ \|v\| : v \in \partial \hat{R}(w(t)) \}$.

Consequently,

$$\hat{R}(w(t)) - \hat{R}(w(0)) = \int_0^t \frac{d}{ds} \hat{R}(w(s)) \, ds = -\int_0^t \min \{ \|v\|^2 : v \in \partial \hat{R}(w(s)) \} \, ds;$$

since the right hand side is nonpositive for all $t$, the flow never increases the objective.

This allows us to reprove our stationary point guarantee from an earlier lecture: since

$$\hat{R}(w(t)) - \hat{R}(w(0)) = -\int_0^t \min \{ \|v\|^2 : v \in \partial \hat{R}(w(s)) \} \, ds \leq -t \min \limits_{s \in [0,t]} \min \limits_{v \in -\partial \hat{R}(w(s))} \|v\|^2,$$

then just as before

$$\min \limits_{s \in [0,t]} \min \limits_{v \in -\partial \hat{R}(w(s))} \|v\|^2 \leq \frac{\hat{R}(w(t)) - \hat{R}(w(0))}{t},$$

thus for some time $s \in [0,t]$, we have an iterate $w(s)$ which is an approximate stationary point.

| **Remark 11.1** | Let’s go back to $\dot{w}(t) := \arg \min \{ \|v\| : v \in -\partial \hat{R}(w(t)) \}$, which we said will hold almost everywhere.

This is *not* satisfied by pytorch/tensorflow/jax/... (Kakade and Lee 2018) gives some bad examples, e.g.,

$$x \mapsto \sigma(\sigma(x)) - \sigma(-x)$$

with $\sigma$ the ReLU, evaluated at 0. (Kakade and Lee 2018) also give a randomized algorithm for finding good subdifferentials.

Does it matter? In the NTK regime, few activations change. In practice, many change, but it’s unclear what their effect is.

---

### 11.1 Positive homogeneity

Another tool we will use heavily outside convexity is *positive homogeneity*.

| **Definition 11.2** | $g$ is *positive homogeneous of degree $L$* when $g(\alpha x) = \alpha^L g(x)$ for $\alpha \geq 0$. (We will only consider continuous $g$, so $\alpha > 0$ suffices.)

| **Example 11.1** |

- Single ReLU: $\sigma(\alpha x) = \alpha \sigma(x)$.|

38
• Monomials of degree $L$ are positive homogeneous of degree $L$:

$$\prod_{i=1}^{d} (\alpha x_i)^{p_i} = \alpha \sum_{i} x_i^{p_i} = \alpha^L \prod_{i} x_i^{p_i}.$$

**Remark 11.2** The math community also has a notion of homogeneity without positivity; the monomial example above works with $\alpha < 0$. Homogeneity in math is often tied to polynomials and generalizations thereof.

**Example 11.2**

- A polynomial $p(x)$ is $L$-homogeneous when all monomials have the same degree; by the earlier calculation,

$$p(\alpha x) = \sum_{j=1}^{r} m_j(\alpha x) = \alpha^L \sum_{j=1}^{r} m_j(x).$$

The algebraic literature often discusses “homogeneous polynomials.”

- Norms are 1-homogeneous, meaning $\|\alpha x\| = \alpha \|x\|$ for $\alpha > 0$. But they moreover satisfy a stronger property $\|\alpha x\| = |\alpha| \cdot \|x\|$ when $\alpha < 0$. Also, $\ell_p$ norms are obtained by taking the root of a homogeneous polynomial, which in general changes the degree of a homogeneous function.

- Layers of a ReLU network are 1-homogeneous in the parameters for that layer:

$$f(x; (W_1, \ldots, \alpha W_i, \ldots, W_L))
= W_L \sigma(W_{L-1} \sigma(\ldots \alpha W_i \sigma(\ldots W_1 x \ldots) \ldots))
= \alpha W_L \sigma(W_{L-1} \sigma(\ldots W_i \sigma(\ldots W_1 x \ldots) \ldots))
= \alpha f(x; \alpha w).$$

The entire network is $L$-homogeneous in the full set of parameters:

$$f(x; \alpha w) = f(x; (\alpha W_1, \ldots, \alpha W_L))
= \alpha W_L \sigma(\alpha W_{L-1} \sigma(\ldots \sigma(\alpha W_1 x) \ldots))
= \alpha^L W_L \sigma(W_{L-1} \sigma(\ldots \sigma(W_1 x) \ldots))
= \alpha^L f(x; w).$$

What is the homogeneity as a function of the input?

- Homework will cover some nonsmooth architectures that are *not* positive homogeneous!

**11.2 Positive homogeneity and the Clarke differential**

Let’s work out an element of the Clarke differential for a ReLU network

$$x \mapsto W_L \sigma_{L-1}(\cdots W_2 \sigma_1(W_1 x)).$$

As a function of $x$, this mapping is 1-homogeneous and piecewise affine. As a function of $w = (W_L, \ldots, W_1)$, it is $L$-homogeneous and piecewise polynomial. The boundary regions form a set of (Lebesgue) measure zero (wrt to either weights or parameters).

Fixing $x$ and considering $w$, interior to each piece, the mapping is differentiable. Due to the definition of Clarke differential, it therefore suffices to compute the gradients in all adjacent pieces, and then take their convex hull.
Remark 11.3  Note that we are not forming the differential by choosing an arbitrary differential element for each ReLU: we are doing a more complicated region-based calculation. However, the former is what pytorch does.

So let’s return to considering some $w$ where are differentiable. Let $A_i$ be a diagonal matrix with activations of the output after layer $i$ on the diagonal:

$$A_i = \text{diag} (\sigma’(W_i \sigma(\ldots \sigma(W_1 x) \ldots))) ,$$

(note we’ve baked in $x$), and so $\sigma(r) = r \sigma’(r)$ implies layer $i$ outputs

$$x \mapsto A_i W_i \sigma(\ldots \sigma(W_1 x) \ldots) = A_i W_i A_{i-1} W_{i-1} \cdots A_1 W_1 x ,$$

and the network outputs

$$f(x; w) = W_L A_{L-1} W_{L-2} \cdots A_1 W_1 x .$$

and the gradient with respect to layer $i$ is

$$\frac{d}{dW_i} f(x; w) = (W_L A_{L-1} \cdots W_{i+1} A_i)^T (A_{i-1} W_{i-1} \cdots W_1 x)^T .$$

Additionally

$$\left\langle W_i, \frac{d}{dW_i} f(x; w) \right\rangle = \left\langle W_i, (W_L A_{L-1} \cdots W_{i+1} A_i)^T (A_{i-1} W_{i-1} \cdots W_1 x)^T \right\rangle$$

$$= \text{tr} (W_i^T (W_L A_{L-1} \cdots W_{i+1} A_i)^T (A_{i-1} W_{i-1} \cdots W_1 x)^T )$$

$$= \text{tr} ((W_L A_{L-1} \cdots W_{i+1} A_i)^T (W_i A_{i-1} W_{i-1} \cdots W_1 x)^T )$$

$$= \text{tr} (W_i A_{i-1} W_{i-1} \cdots W_{i+1} W_1 x)^T (W_L A_{L-1} \cdots W_{i+1} A_i)^T )$$

$$= \text{tr} (W_L A_{L-1} \cdots W_{i+1} A_i W_i A_{i-1} W_{i-1} \cdots W_1 x)$$

$$= f(x; w) ,$$

and

$$\left\langle W_i, \frac{d}{dW_i} f(x; w) \right\rangle = f(x; w) = \left\langle W_{i+1}, \frac{d}{dW_{i+1}} f(x; w) \right\rangle .$$

This calculation can in fact be made much more general (indeed with a simpler proof!).

Lemma 11.2  Suppose $f : \mathbb{R}^d \to \mathbb{R}$ is locally Lipschitz and $L$-positively homogeneous. For any $w \in \mathbb{R}^d$ and $s \in \partial f(w)$, 

$$\langle s, w \rangle = L f(w) .$$

Remark 11.4  This statement appears in various places (Lyu and Li 2019); the version here is somewhat more general, and appears in (Ji and Telgarsky 2020).
Proof. If \( w = 0 \), then \( \langle s, w \rangle = 0 = Lf(w) \) for every \( s \in \partial f(w) \), so consider the case \( w \neq 0 \). Let \( D \) denote those \( w \) where \( f \) is differentiable, and consider the case that \( w \in D \setminus \{0\} \). By the definition of gradient,
\[
\lim_{\delta \downarrow 0} \frac{f(w + \delta w) - f(w) - \langle \nabla f(w), \delta w \rangle}{\delta \|w\|} = 0,
\]
and by using homogeneity in the form \( f(w + \delta w) = (1 + \delta)^L f(w) \) (for any \( \delta > 0 \)), then
\[
0 = \lim_{\delta \downarrow 0} \frac{\left((1 + \delta)^L - 1\right) f(w) - \langle \nabla f(w), \delta w \rangle}{\delta} = -\langle \nabla f(w), w \rangle + \lim_{\delta \downarrow 0} f(w) (L + \mathcal{O}(\delta)),
\]
which implies \( \langle w, \nabla f(w) \rangle = Lf(w) \).
Now consider \( w \in \mathbb{R}^d \setminus D \setminus \{0\} \). For any sequence \( (w_i)_{i \geq 1} \) in \( D \) with \( \lim_i w_i = w \) for which there exists a limit \( s := \lim_i \nabla f(w_i) \), then
\[
\langle w, s \rangle = \lim_{i \to \infty} \langle w_i, \nabla f(w_i) \rangle = \lim_{i \to \infty} Lf(w_i) = Lf(w).
\]
Lastly, for any element \( s \in \partial f(w) \) written in the form \( s = \sum_i \alpha_i s_i \) where \( \alpha_i \geq 0 \) satisfy \( \sum_i \alpha_i = 1 \) and each \( s_i \) is a limit of a sequence of gradients as above, then
\[
\langle w, s \rangle = \left\langle w, \sum_i \alpha_i s_i \right\rangle = \sum_i \alpha_i \langle w, s_i \rangle = \sum_i \alpha_i Lf(w) = Lf(w).
\]

11.3 Norm preservation

If predictions are positive homogeneous with respect to each layer, then gradient flow preserves norms of layers.

Lemma 11.3 (Simon S. Du, Hu, and Lee (2018)) Suppose for \( \alpha > 0 \), \( f(x; (W_L, \ldots, \alpha W_i, \ldots, W_1)) = \alpha f(x; w) \) (predictions are 1-homogeneous in each layer). Then for every pair of layers \((i, j)\), the gradient flow maintains
\[
\frac{1}{2} \|W_i(t)\|^2 - \frac{1}{2} \|W_i(0)\|^2 = \frac{1}{2} \|W_j(t)\|^2 - \frac{1}{2} \|W_j(0)\|^2.
\]

Remark 11.5 We’ll assume a risk of the form \( \mathbb{E}_k \ell(y_k f(x_k; w)) \), but it holds more generally. We are also tacitly assuming we can invoke the chain rule, as discussed above.
Proof. Defining $\ell'_k(s) := y_k\ell'\left(y_k f(x_k; w(s))\right)$, and fixing a layer $i$,

$$\frac{1}{2} \|W_i(t)\|^2 - \frac{1}{2} \|W_i(0)\|^2 = \int_0^t \frac{d}{dt} \frac{1}{2} \|W_i(s)\|^2 ds$$

$$= \int_0^t \left\langle W_i(s), \dot{W}_i(s) \right\rangle ds$$

$$= \int_0^t \left\langle W_i(s), -\mathbb{E}_k \ell'_k(s) \frac{df(x_k; w)}{dW_i(s)} \right\rangle ds$$

$$= -\int_0^t \mathbb{E}_k \ell'_k(s) \left\langle W_i(s), \frac{df(x_k; w)}{dW_i(s)} \right\rangle ds$$

$$= -\int_0^t \mathbb{E}_k \ell'_k(s) f(x_k; w) ds.$$

This final expression does not depend on $i$, which gives the desired equality.

**Remark 11.6** One interesting application is to classification losses like $\exp(-z)$ and $\ln(1 + \exp(-z))$, where $R(w) \to 0$ implies $\min_k y_k f(x_k; w) \to \infty$.

This by itself implies $\|W_j\| \to \infty$ for some $j$; combined with norm preservation, $\min_j \|W_j\| \to \infty$.

[mjt: need to update this in light of the new material i’ve included?]

### 11.4 Smoothness inequality adapted to ReLU

Let’s consider: single hidden ReLU layer, only bottom trainable:

$$f(x; w) := \frac{1}{\sqrt{m}} \sum_j a_j \sigma(\langle x, w_j \rangle), \quad a_j \in \{\pm 1\}.$$

Let $W_s \in \mathbb{R}^{m \times d}$ denote parameters at time $s$, suppose $\|x\| \leq 1$.

$$\frac{df(x; W)}{dW} = \begin{bmatrix} a_1 x \sigma'(w_1^T x)/\sqrt{m} \\ \vdots \\ a_m x \sigma'(w_m^T x)/\sqrt{m} \end{bmatrix},$$

$$\left\| \frac{df(x; W)}{dW} \right\|_F^2 = \sum_j \left\| a_j x \sigma'(w_j^T x)/\sqrt{m} \right\|_2^2 \leq \frac{1}{m} \sum_j \|x\|_2^2 \leq 1.$$

We’ll use the logistic loss, whereby

$$\ell(z) = \ln(1 + \exp(-z)),$$

$$\ell'(z) = \frac{-\exp(-z)}{1 + \exp(-z)} \in (-1, 0),$$

$$\widehat{R}(W) := \frac{1}{n} \sum_k \ell(y_k f(x_k; W)).$$

A key fact (can be verified with derivatives) is

$$|\ell'(z)| = -\ell'(z) \leq \ell(z),$$

42
whereby

\[
\frac{d\hat{R}}{dW} = \frac{1}{n} \sum_k \ell'(y_k f(x_k; W)) y_k \nabla_W f(x_k W),
\]

\[
\left\| \frac{d\hat{R}}{dW} \right\|_F \leq \frac{1}{n} \sum_k |\ell'(y_k f(x_k; W))| \cdot \|y_k \nabla_W f(x_k W)\|_F \\
\leq \frac{1}{n} \sum_k |\ell'(y_k f(x_k; W))| \leq \min \left\{ 1, \hat{R}(W) \right\}.
\]

Now we can state a non-smooth, non-convex analog to Theorem 7.3

**Lemma 11.4** ((Lemma 2.6, Ji and Telgarsky 2019a)) If $\eta \leq 1$, for any $Z$,

\[
\|W_i - Z\|_F^2 + \eta \sum_{i < t} \hat{R}^{(i)}(W_i) \leq \|W_0 - Z\|_F^2 + 2\eta \sum_{i < t} \hat{R}^{(i)}(Z),
\]

where $\hat{R}^{(i)}(W) = \frac{1}{n} \sum_k \ell(y_k \langle W, \nabla f(x_k; W_i) \rangle)$.

**Remark 11.7**
- $\hat{R}^{(i)}(W_i) = \hat{R}(W_i)$.
- $\hat{R}^{(i)}(Z) \approx \hat{R}(Z)$ if $W_i$ and $Z$ have similar activations.
- (Ji and Telgarsky 2019a) uses this in a proof scheme like (Chizat and Bach 2019): consider those iterations where the activations are similar, and then prove it actually happens a lot. (Ji and Telgarsky 2019a), with additional work, can use this to prove low test error.

**Proof.** Using the squared distance potential as usual,

\[
\|W_{i+1} - Z\|_F^2 = \|W_i - Z\|_F^2 - 2\eta \left\langle \nabla \hat{R}(W_i), W_i - Z \right\rangle + \eta^2 \|\nabla \hat{R}(W_i)\|_F^2,
\]

where $\|\nabla \hat{R}(W_i)\|_F^2 \leq \|\nabla \hat{R}(W_i)\|_F \leq \hat{R}(W_i) = \hat{R}^{(i)}(W_i)$, and

\[
n \left\langle \nabla \hat{R}(W_i), Z - W_i \right\rangle = \sum_k y_k \ell'(y_k f(x_k; W_i)) \langle \nabla_W f(x_k; W_i), Z - W_i \rangle \\
= \sum_k \ell'(y_k f(x_k; W_i)) \langle \nabla_W f(x_k; W_i), Z - y_k f(x_k; W_i) \rangle \\
\leq \sum_k \left( \ell(y_k \langle \nabla_W f(x_k; W_i), Z \rangle) - \ell(y_k f(x_k; W_i)) \right) \\
= n \left( \hat{R}^{(i)}(Z) - \hat{R}^{(i)}(W_i) \right).
\]

Together,

\[
\|W_{i+1} - Z\|_F^2 \leq \|W_i - Z\|_F^2 + 2\eta \left( \hat{R}^{(i)}(Z) - \hat{R}^{(i)}(W_i) \right) + \eta \hat{R}_i(W_i);
\]

applying $\sum_{i < t}$ to both sides gives the bound.
**Margin maximization and implicit bias**

During 2015-2016, various works pointed out that deep networks generalize well, even though parameter norms are large, and there is no explicit generalization (Neyshabur, Tomioka, and Srebro 2014; Zhang et al. 2017). This prompted authors to study implicit bias of gradient descent, the first such result being an analysis of linear predictors with linearly separable data, showing that gradient descent on the cross-entropy loss is implicitly biased towards a maximum margin direction (Soudry, Hoffer, and Srebro 2017).

This in turn inspired many other works, handling other types of data, networks, and losses (Ji and Telgarsky 2019b, 2018, 2020; Gunasekar et al. 2018a; Lyu and Li 2019; Chizat and Bach 2020; Ji et al. 2020).

Margin maximization of first-order methods applied to exponentially-tailed losses was first proved for coordinate descent (Telgarsky 2013). The basic proof scheme there was pretty straightforward, and based on the similarity of the empirical risk (after the monotone transformation $\ln(\cdot)$) to $\ln \sum \exp$, itself similar to $\max(\cdot)$ and thus to margin maximization; we will use this connection as a basis for all proofs in this section (see also (Ji and Telgarsky 2019b; Gunasekar et al. 2018b)).

Throughout this section, fix training data $((x_i, y_i))_{i=1}^n$, define a (an unnormalized) margin mapping

$$m_i(w) := y_i f(x_i; w);$$

by this choice, we can also conveniently write an unnormalized risk $\mathcal{L}$:

$$\mathcal{L}(w) := \sum_i \ell(m_i(w)) = \sum_i \ell(y_i f(x_i; w)).$$

Throughout this section, we will always assume $f$ is locally-Lipschitz and $L$-homogeneous in $w$, which also means each $m_i$ is locally-Lipschitz and $L$-homogeneous.

We will also use the exponential loss $\ell(z) = \exp(-z)$. The results go through for similar losses.

**Remark 12.1 (generalization)** As hinted before, margin maximization is one way gradient descent prefers a solution which has a hope to generalize well, and not merely achieve low empirical risk. This low generalization error of large-margin predictors will appear explicitly later on in section 15.4.

**Remark 12.2 (implicit bias)** As mentioned above, the proofs here will show implicit margin maximization, which is enough to invoke the generalization theory in section 15.4. However, in certain cases it is valuable to moreover prove converges rates to the maximum margin direction. In the linear case, it is possible to convert a margin maximization rate to an implicit bias rate, however the rate degrades by a factor $\sqrt{\cdot}$ (Ji and Telgarsky 2019b); analyzing the implicit bias without degradation in the rate is more involved, and not treated here (Soudry, Hoffer, and Srebro 2017).

**Remark 12.3 (squared loss)** While the focus here is on losses with exponential tails and on bias towards the maximum margin direction, there are also many works (not further discussed here) which consider the squared loss (Gunasekar et al. 2017; Arora, Cohen, et al. 2018b, 2019).
We just said “maximum margin” and “separable data.” What do these mean? Consider a linear predictor, meaning \( x \mapsto \langle w, x \rangle \) for some \( w \in \mathbb{R}^d \). This \( w \) “separates the data” if \( y_i \) and \( \text{sgn}(\langle w, x_i \rangle) \) agree, which we can relax to the condition of strict separability, namely
\[
\min_i y_i \langle w, x_i \rangle > 0.
\]

It seems reasonable, or a nice inductive bias, if we are as far from 0 as possible:
\[
\max_{w \in \mathcal{W}} \min_i y_i \langle w, x_i \rangle > 0.
\]

The “?" indicates that we must somehow normalize or constrain, since otherwise, for separable data, this \( \max \) becomes a \( \sup \) and has value \( +\infty \).

**Definition 12.1** Data is linearly separable when there exists \( w \in \mathbb{R}^d \) so that \( \min_i y_i \langle w, x_i \rangle > 0 \). In this situation, the \((\ell_2)\) maximum margin predictor (which is unique!) is given by
\[
\bar{u} := \arg \max_{\|w\|=1} \min_i y_i \langle w, x_i \rangle,
\]
and the margin is \( \gamma := \min_i y_i \langle \bar{u}, x_i \rangle \).

**Remark 12.4** This concept has a long history. Margins first appeared in the classical perceptron analysis (Novikoff 1962), and maximum margin predictors were a guiding motivation for the SVM [mjt: need to add many more refs].

Consider now the general case of \( L \)-homogeneous predictors, where \( y_i \langle w, x_i \rangle \) is replaced by \( m_i(w) \).

**Proposition 12.1** Suppose \( f(x; w) \) is \( L \)-homogeneous in \( w \), \( \ell \) is the exponential loss, and there exists \( \hat{w} \) with \( \hat{R}(\hat{w}) < \ell(0)/n \). Then \( \inf_w \hat{R}(w) = 0 \), and the infimum is not attained.

**Proof.** Note
\[
\max_i \ell(-m_i(\hat{w})) \leq \sum_i \ell(-m_i(\hat{w})) = n \hat{R}(\hat{w}) < \ell(0),
\]
thus applying \( \ell^{-1} \) to both sides gives \( \min_i m_i(\hat{w}) > 0 \). Therefore
\[
0 \leq \inf_w \hat{R}(w) \leq \limsup_{c \to \infty} \hat{R}(cw) = \sum_i \limsup_{c \to \infty} \ell(-m_i(c\hat{w})) = \sum_i \limsup_{c \to \infty} \ell(-c^L m_i(\hat{w})) = 0.
\]

This seems to be problematic; how can we “find” an “optimum,” when solutions are off at infinity? Moreover, we do not even have unique directions, nor a way to tell different ones apart!

We can use margins, now appropriately generalized to the \( L \)-homogeneous case, to build towards a better-behaved objective function. First note that since
\[
\min_i m_i(w) = \|w\|^L \min_i m_i\left(\frac{w}{\|w\|}\right),
\]
we can compare different directions by normalizing the margin by \( \|w\|^L \). Moreover, again using the
exponential loss,
\[
\frac{\ell^{-1}(\mathcal{L}(w)) + \ln(n)}{\|w\|^L} = \frac{\ell^{-1}(\sum_i \ell(m_i(w))/n)}{\|w\|^L} \geq \frac{\min_i m_i(w)}{\|w\|^L} = \frac{\ell^{-1}(\max_i \ell(m_i(w)))}{\|w\|^L} \geq \frac{\ell^{-1}(\mathcal{L}(w))}{\|w\|^L}.
\]

(3)

This motivates the following definition.

**Definition 12.2** Say the data is $\vec{m}$-separable when there exists $w$ so that $\min_i m_i(w) > 0$. Define the margin, maximum margin, and smooth margin respectively as
\[
\gamma(w) := \frac{\min_i m_i(w)}{\|w\|^L}, \quad \bar{\gamma} := \max_{\|w\|=1} \gamma(w), \quad \tilde{\gamma}(w) := \frac{\ell^{-1}(\mathcal{L}(w))}{\|w\|^L}.
\]

(4)

**Remark 12.5** The terminology “smoothed margin” is natural for $L$-homogeneous predictors, but even so it seems to have only appeared recently in (Lyu and Li 2019). In the 1-homogeneous case, the smoothed margin appeared much earlier, indeed throughout the boosting literature (Schapire and Freund 2012).

**Remark 12.6 (multiclass margins)** There is also a natural notion of multiclass margin:
\[
\min_i \frac{f(x_i; w)_y - \max_{j \neq y} f(x_i; w)_j}{\|w\|^L}.
\]

The natural loss to consider in this setting is the cross-entropy loss.

The basic properties can be summarized as follows.

**Proposition 12.2** Suppose data is $\vec{m}$-separable. Then:
\begin{itemize}
  \item $\bar{\gamma} := \max_{\|w\| \leq 1} \gamma(w) > 0$ is well-defined (the maximum is attained).
  \item For any $w \neq 0$, For any $\hat{w}$ with $\bar{\gamma} = \gamma(\hat{w})$,
  \[
  \lim_{c \to \infty} \tilde{\gamma}(cw) = \gamma(w).
  \]
\end{itemize}

In particular, for $\hat{w}$ satisfying $\bar{\gamma} = \gamma(\hat{w})$, then $\lim_{c \to \infty} \tilde{\gamma}(c\hat{w}) = \bar{\gamma}$.

**Proof.** The first part follows by continuity of $m_i(w)$ and compactness of $\{w \in \mathbb{R}^p : \|w\| = 1\}$, and the second from eq. 4 and eq. 3.

**Remark 12.7** For the linear case, margins have a nice geometric interpretation. This is not currently true for the general homogeneous case: there is no known reasonable geometric characterization of large margin predictors even for simple settings.

12.2 Gradient flow maximizes margins of linear predictors

Let’s first see how far we can get in the linear case, using one of our earlier convex optimization tools, namely Theorem 7.4.
Lemma 12.1 Consider the linear case, with linearly separable data and the exponential loss, and \( \max_i \|x_i y_i\| \leq 1 \). Then
\[
\mathcal{L}(w_t) \leq \frac{1 + \ln(2tn\gamma^2)}{2t\gamma^2},
\]
\[
\|w_t\| \geq \ln(2tn\gamma^2) - \ln \left( 1 + \ln(2tn\gamma^2) \right).
\]

Remark 12.8 The intuition we will follow for the proof is: for every unit of norm, the (un-normalized) margin increases by at least \( \gamma \). Thus the margin bias affects the entire gradient descent process.

Later, when we study the \( L \)-homogeneous case, we are only able to show for every unit norm (to the power \( L \)), the (unnormalized) margin increases by at least the current margin, which implies nondecreasing, but not margin maximization.

Proof. By Theorem 7.4 with \( z = \ln(c)\bar{u}/\gamma \) for some \( c > 0 \),
\[
\mathcal{L}(w(t)) \leq \mathcal{L}(z) + \frac{1}{2t} \left( \|z\|^2 - \|w(t) - z\|^2 \right) \leq \sum_i \ell(m_i(z)) + \frac{\|z\|^2}{2t} \leq \sum_i \exp(-\ln(c)) + \frac{\ln(c)^2}{2t\gamma^2} = \frac{n}{c} + \frac{\ln(c)^2}{2t\gamma^2},
\]
and the first inequality follows from the choice \( c := 2tn\gamma^2 \). For the lower bound on \( \|w_t\| \), using the preceding inequality,
\[
\ell(\|w_t\|) \leq \min_i \ell(m_i(w_t)) \leq \frac{1}{n} \mathcal{L}(w_t) \leq \frac{1 + \ln(2tn\gamma^2)^2}{2tn\gamma^2},
\]
and the second inequality follows by applying \( \ell^{-1} \) to both sides.

This nicely shows that we decrease the risk to 0, but not that we maximize margins. For this, we need a more specialized analysis.

Theorem 12.1 Consider the linear case, with linearly separable data and the exponential loss, and \( \max_i \|x_i y_i\| \leq 1 \). Then
\[
\gamma(w_t) \geq \tilde{\gamma}(w_t) \geq \tilde{\gamma} - \frac{\ln n}{\ln t + \ln(2n\gamma^2) - 2\ln \ln(2tn\gamma^2)}.
\]
[ mjt: need to check some constants. also that denominator is hideous, maybe require slightly larger \( t \) to remove it?]
Proof. For convenience, define \( u(t) := \ell^{-1}(\mathcal{L}(w(t))) \) and \( v(t) := \|w(t)\| \), whereby
\[
\gamma(w(t)) = \frac{u(t)}{v(t)} = \frac{u(0)}{v(t)} + \int_0^t \frac{\dot{u}(s)}{v(t)} ds.
\]

Let’s start by lower bounding the second term. Since \( \ell' = -\ell \),
\[
\dot{u}(t) = \left\langle -\nabla \mathcal{L}(w(t)), \dot{w}(t) \right\rangle = \frac{\|\dot{w}(t)\|^2}{\mathcal{L}(w(t))},
\]
\[
\|\dot{w}(s)\| \geq \langle \dot{w}(s), \bar{u} \rangle = \left( -\sum_i x_i y_i \ell'(m_i(w(s))), \bar{u} \right) = \sum_i \ell(m_i(w(s))) (x_i y_i, \bar{u}) \geq \gamma \sum_i \ell(m_i(w(s))) = \gamma \mathcal{L}(w(s)),
\]
\[
v(t) = \|w(t) - w(0)\| = \left\| \int_0^t \dot{w}(s)ds \right\| \leq \int_0^t \|\dot{w}(s)\| ds,
\]
thus
\[
\frac{\int_0^t \dot{u}(s)ds}{v(t)} \geq \frac{\int_0^t \|\dot{w}(s)\|^2 ds}{v(t)} = \frac{\int_0^t \|\dot{w}(s)\| \|\dot{w}(s)\| ds}{v(t)} \geq \frac{\gamma \int_0^t \|\dot{w}(s)\| ds}{v(t)} = \gamma.
\]

For the first term \( u(0)/v(t) \), note \( \mathcal{L}(w(0)) = n \) and thus \( u(0) = -\ln n \), whereas by the lower bound on \( \|w(t)\| \) from Lemma 12.1,
\[
\frac{u(0)}{v(t)} = -\frac{\ln(n)}{\|w(t)\|} \geq \frac{-\ln(n)}{\ln(t) + \ln(2n\gamma^2) - 2\ln(2nt\gamma^2)}.
\]

Combining these inequalities gives the bound.

We are maximizing margins, but at a glacial rate of \( 1/\ln(t)! \)

To get some inspiration, notice that we keep running into \( \ell^{-1}(\mathcal{L}(w)) \) in all the analysis. Why don’t we just run gradient flow on this modified objective? In fact, the two gradient flows are the same!

**Remark 12.9 (time rescaling)** Let \( w(t) \) be given by gradient flow on \( \mathcal{L}(w(t)) \), and define a time rescaling \( h(t) \) via integration, namely so that \( \dot{h}(t) = 1/\mathcal{L}(w(h(t))) \). Then, by the substitution rule for integration,
\[
w(t) - w(0) = \int_0^t \dot{w}(s)ds = -\int_0^t \nabla \mathcal{L}(w(s))ds = \int_{h^{-1}([0,t])} \nabla \mathcal{L}(w(h(s)))|\dot{h}(s)|ds
\]
\[
= \int_{h^{-1}([0,t])} \frac{\nabla \mathcal{L}(w(h(s)))}{\mathcal{L}(w(h(s)))}ds = \int_{h^{-1}([0,t])} \nabla \ln \mathcal{L}(w(h(s)))ds
\]

As such, the gradient flow on \( \mathcal{L} \) and on \( \ell^{-1} \circ \mathcal{L} \) are the same, modulo a time rescaling. This perspective was first explicitly stated by Chizat and Bach (2020), though analyses using this rescaled time (and alternate flow characterization) existed before Lyu and Li (2019).

**Theorem 12.2 (time-rescaled flow)** Consider linear predictors with linearly separable data, and the logistic loss. Suppose \( \dot{\theta}(t) := \nabla_\theta \ell^{-1}(\mathcal{L}(\theta(t))) \). Then
\[
\gamma(\theta(t)) \geq \dot{\gamma}(\theta(t)) \geq \gamma - \frac{\ln n}{t\gamma^2 - \ln n}.
\]
**Proof.** We start as before: set \( u(t) := \ell^{-1}L(\theta(t)) \) and \( v(t) := \|\theta(t)\| \); then

\[
\tilde{\gamma}(t) = \frac{u(t)}{v(t)} = \frac{u(0)}{v(t)} + \int_0^t \frac{\dot{u}(s)}{v(t)} ds = -\frac{\ln n}{v(t)} + \int_0^t \frac{\dot{u}(s)}{v(t)} ds.
\]

Bounding these terms is now much simpler than for the regular gradient flow. Note

\[
\|\dot{\theta}(s)\| \geq \langle \nabla \ln L(\theta(s)), \tilde{u} \rangle = \sum_i \frac{\ell'\left(m_i(\theta(s))\right)}{L(\theta(s))} x_i y_i \geq \gamma \sum_i \frac{\ell'(m_i(\theta(s)))}{L(\theta(s))} = \gamma,
\]

\[
\tilde{u}(s) = \left\langle \nabla \ln L(\theta(s)), \dot{\theta}(s) \right\rangle = \|\dot{\theta}(s)\|^2,
\]

thus

\[
\ell^{-1}L(\theta(t)) = \ell^{-1}L(\theta(0)) + \int_0^t \frac{d}{ds} \ell^{-1}L(\theta(s)) ds \geq -\ln(n) + t\gamma^2,
\]

\[
\frac{\int_0^t \dot{u}(s) ds}{v(t)} \geq \frac{\gamma \int_0^t \|\dot{\theta}(s)\| ds}{v(t)} \geq \frac{\gamma \|\dot{\theta}(s)\| ds}{v(t)} = \gamma.
\]

On the other hand,

\[
\|\theta(t)\| \geq \|\theta(t)\| \gamma(\theta(t)) \geq \ell^{-1}L(\theta(t)) \geq t\gamma^2 - \ln(n).
\]

Together,

\[
\gamma(t) = \frac{u(t)}{v(t)} \geq \gamma - \frac{\ln(n)}{t\gamma^2 - \ln n}.
\]

**Remark 12.10** The preceding two proofs are simplified from (Ji and Telgarsky 2019b), but follow a general scheme from the (coordinate descent!) analysis in (Telgarsky 2013); this scheme was also followed in (Gunasekar et al. 2018b). The proof in (Soudry, Hoffer, and Srebro 2017) is different, and is based on an SVM analogy, since \( \tilde{\gamma} \to \gamma \).

Note also that the proofs here do not show \( w(t) \) converges to (the unique) maximum margin linear separator, which is easy to do with worse rates, and harder to do with good rates. However, large margins is sufficient for generalization in the linear case, as in section 15.4.

### 12.3 Smoothed margins are nondecreasing for homogeneous functions

In the nonlinear case, we do not have a general result, and instead only prove that smoothed margins are nondecreasing.

**Theorem 12.3 (originally from (Lyu and Li 2019), simplification due to (Ji 2020))**

Suppose there exists \( t_0 \) with \( \tilde{\gamma}(w(t_0)) > 0 \). Then \( t \mapsto \tilde{\gamma}(w(t)) \) is nondecreasing along \([t_0, \infty)\).
**Proof.** Write \( \bar{\gamma}_t := \bar{\gamma}(w(t)) = u_t/v_t, \) where

\[
u_t := \ell^{-1}(\mathcal{L}(w_t)), \quad v_t := \|w_t\|^L.
\]

By the quotient rule,

\[
\frac{d}{dt} \bar{\gamma}_t = \frac{\dot{u}_tv_t - \dot{v}_tu_t}{v_t^2}.
\]

Therefore it suffices to show \( v_t \neq 0 \) and that the numerator is positive.

First note a technical fact that since \( -\ell' = \ell > 0, \)

\[
\langle w, \dot{w} \rangle = \sum_j -\ell'(m_j(w)) \langle w, \nabla m_j(w) \rangle = L \sum_j -\ell'(m_j(w))m_j(w).
\]

\[
= L \sum_j -\ell'(m_j(w))\ell^{-1}(\ell(m_j(w))) \geq L \sum_j -\ell'(m_j(w))\ell^{-1}(\mathcal{L}(w))
\]

\[
= L\mathcal{L}(w)\ell^{-1}(\mathcal{L}(w)).
\]

Moreover,

\[
\dot{v}_t = \frac{d}{dt} \langle w_t, w_t \rangle^{L/2} = \frac{L}{2} \langle w_t, w_t \rangle^{L/2-1} 2 \langle w_t, \dot{w}_t \rangle = L\|w_t\|^{L-2} \langle w_t, \dot{w}_t \rangle.
\]

Consequently,

\[
\dot{v}_t = L\|w_t\|^{L-1} \left\langle \frac{w_t}{\|w_t\|}, \dot{w}_t \right\rangle \leq L\|w_t\|^{L-1} \sup_{\|v\| \leq 1} \langle v, \dot{w}_t \rangle = L\|w_t\|^{L-1}\|\dot{w}_t\|
\]

\[
\dot{v}_t \geq L^2\|w_t\|^{L-2} \mathcal{L}(w_t)\ell^{-1}(\mathcal{L}(w_t))
\]

For \( \dot{u}_t, \) using \( \ell(z) = \exp(-z) \) and the earlier lower bound on \( \langle w, \dot{w} \rangle, \) [ mj\text{\textcopyright} highlight this is a key step, differing from linear proof.]

\[
\dot{u}_t = -\frac{\mathcal{L}(w_t)\dot{w}_t}{\mathcal{L}(w_t)} = \frac{\|\dot{w}_t\|^2}{\mathcal{L}(w_t)} \geq \frac{\|\dot{w}_t\|}{\mathcal{L}(w_t)} \|\dot{w}_t, w_t\| \geq L\|w_t\|^{\ell^{-1}(\mathcal{L}(w_t))}.
\]

Therefore

\[
\dot{u}_tv_t - \dot{v}_tu_t \geq L\|w_t\|^{\ell^{-1}(\mathcal{L}(w_t))}\|w_t\|^L - L\|w_t\|^{L-1}\|\dot{w}_t\|\ell^{-1}(\mathcal{L}(w_t)) = 0.
\]

It remains to show \( v_t \) stays positive. \( v_0 > 0 \) since \( \mathcal{L}(w_t) < \ell(0)/n \leq \mathcal{L}(0). \) As before,

\[
\dot{v}_t \geq L^2\|w_t\|^{L-2} \mathcal{L}(w_t)\ell^{-1}(\mathcal{L}(w_t)) = L^2\|w_t\|^{2L-2} \mathcal{L}(w_t)\bar{\gamma}_t.
\]

Let \( T \) denote the first time where \( v_t = 0 \) for \( t \in [0, T) \), \( v_t > 0 \) and thus \( \bar{\gamma}_t \geq \bar{\gamma}_0 \) and \( \dot{v}_t > 0 \), meaning such a time \( T \) can not exist, and \( \dot{v}_t > 0 \) and \( (d/dt)\bar{\gamma}_t > 0 \).

**Remark 12.11** As mentioned, +Theorem 12.3 (originally from (Lyu and Li 2019), simplification due to (Ji 2020)) was originally presented in (Lyu and Li 2019), though this simplification is due to (Ji 2020), and its elements can be found throughout (Ji and Telgarsky 2020). The version in (Lyu and Li 2019) is significantly different, and makes heavy (and interesting) use of a polar decomposition of homogeneous functions and gradient flow on them.
For the case of an infinite-width 2-homogeneous network, assuming a number of convergence properties of the flow (which look technical, but are not “merely technical,” and indeed difficult to prove), margins are globally maximized (Chizat and Bach 2020).

13 Generalization: preface

The purpose of this generalization part is to bound the gap between testing and training error for standard (multilayer ReLU) deep networks via the classical uniform convergence tools, and also to present and develop these classical tools (based on Rademacher complexity).

These bounds are very loose, and there is extensive criticism now both of them and of the general approach, as will be discussed shortly (Neyshabur, Tomioka, and Srebro 2014; Zhang et al. 2017; Nagarajan and Kolter 2019; Dziugaite and Roy 2017); this work is ongoing and moving quickly and there are even already many responses to these criticisms (Negrea, Dziugaite, and Roy 2019; L. Zhou, Sutherland, and Srebro 2020; P. L. Bartlett and Long 2020).

13.1 Omitted topics

- Domain adaptation / covariate shift.
- Generalization properties of more architectures. One key omission is of convolution layers; for one generalization analysis, see (Long and Sedghi 2019).
- Other approaches and perspectives on generalization (possibly changing the basic definitions of “generalization”), for instance:
  - PAC-Bayes approaches (Dziugaite and Roy 2017). In the present notes, we only focus on uniform convergence bounds, which give high probability bounds between training and test error which hold simultaneously for every element of some class.
  
  By contrast, PAC-Bayes consider a distribution over predictors, and bound the expected gap between testing and training error for these predictors in terms of how close this distribution is to some prior distribution over the predictors.

  The looseness of the uniform-convergence bounds presented in these notes leads many authors to instead use them as explanatory tools, e.g., by studying their correlation with observed generalization. A correlation was claimed and presented in (P. Bartlett, Foster, and Telgarsky 2017), however it was on a single dataset and architecture. More extensive investigations have appeared recently (Jiang et al. 2020; Dziugaite et al. 2020), and highlight that while some bounds are correlated with generalization (or rather predictive of generalization) in some settings, there are other situations (e.g., large width) where no bound is correlated with observed generalization gaps.

  Compression-based approaches (Arora, Ge, et al. 2018), which bound the generalization of the network after applying some compression procedure, with no guarantees on the original network; that said, it is a promising approach, and there has been some effort to recover guarantees on the original network (Suzuki, Abe, and Nishimura 2019).

Another relevant work, from an explicitly PAC-Bayes perspective, is (W. Zhou et al. 2018). For further connections between PAC-Bayes methodology and compression, see
(Blum and Langford 2003), and for more on the concept of compression schemes, see for instance (Moran and Yehudayoff 2015).

- Double descent (Belkin et al. 2018; Belkin, Hsu, and Xu 2019; Hastie et al. 2019), and related “interpolating predictors.”

- Various omitted bounds in our uniform deviation framework:
  - (Wei and Ma 2019) give a bound which requires smooth activations; if we convert it to ReLU, it introduces a large factor which does not seem to improve over those presented here. That said, it is an interesting bound and approach. (There are a number of other bounds we don’t discuss since similarly they degrade for ReLU.)
  - (Golowich, Rakhlin, and Shamir 2018) have an additional bound over the one of theirs we present here: interestingly, it weakens the depends on $\sqrt{n}$ to $n^{1/4}$ or $n^{1/5}$ but in exchange vastly improves the dependence on norms in the numerator, and is a very interesting bound.

\section*{14 Concentration of measure}

- Concentration of measure studies how certain distribution families and operations on distributions lead to “clumping up” of probability mass. Examples we’ve seen:
  - Gaussians concentrate around the one-standard-deviation shell; we used this in NTK to say few activations change (so it’s concentrated away from 0, sometimes this is called “anti-concentration”).
  - Azuma-Hoeffding gave us control on the errors in SGD; note that we averaged together many errors before studying concentration!

- We’ll see in this section that concentration of measure allows us to handle the generalization gap of single predictors fixed in advance, but is insufficient to handle the output of training algorithms.

- We will be absurdly brief. Some other resources:
  - Martin Wainwright’s lecture notes (Wainwright 2015), now turned into a book (Wainwright 2019).
  - My learning theory class, as well as Maxim Raginsky’s.

\subsection*{14.1 sub-Gaussian random variables and Chernoff’s bounding technique}

Our main concentration tool will be the Chernoff bounding method, which works nicely with sub-Gaussian random variables.

\begin{definition}
Random variable $Z$ is sub-Gaussian with mean $\mu$ and variance proxy $\sigma^2$ when $\mathbb{E} e^{\lambda(Z-\mu)} \leq e^{\lambda^2 \sigma^2/2}$.
\end{definition}

\begin{example}
- Gaussian $\mathcal{N}(\mu, \sigma^2)$ is $(\mu, \sigma^2)$-sub-Gaussian.
\end{example}
• \( \Sigma_i Z_i \) is (\( \Sigma_i \mu, \|\sigma\|^2 \)) when \( Z_i \) is \((\mu_i, \sigma^2)\)-sub-Gaussian.

• If \( Z \in [a, b] \) a.s., then \( Z \) is \((\mathbb{E} Z, (b-a)^2/4)\)-sub-Gaussian (this is called the Hoeffding lemma \[ mj托: I should pick a proof.\]).

| Remark 14.1 | There is also “sub-exponential”; we will not use it but it is fundamental. |

Sometimes \( \mu \) is dropped from definition; in this case, one can study \( X - \mathbb{E} X \), and we’ll just say “\( \sigma^2\)-sub-Gaussian.”

\( \mathbb{E} \exp(\lambda Z) \) is the moment generating function of \( Z \); it has many nice properties, though we’ll only use it in a technical way.

Sub-Gaussian random variables will be useful to us due to their vanishing tail probabilities. This indeed is an equivalent way to define sub-Gaussian (see (Wainwright 2015)), but we’ll just prove implication. The first step is Markov’s inequality.

| Theorem 14.1 (Markov’s inequality) | For any nonnegative r.v. \( X \) and \( \epsilon > 0 \), |

\[
\Pr[X \geq \epsilon] \leq \frac{\mathbb{E} X}{\epsilon}.
\]

| Proof. | Apply \( \mathbb{E} \) to both sides of \( \epsilon 1[X \geq \epsilon] \leq X \). |

| Corollary 14.1 | For any nonnegative, nondecreasing \( f \geq 0 \) and \( f(\epsilon) > 0 \), |

\[
\Pr[X \geq \epsilon] \leq \frac{\mathbb{E} f(X)}{f(\epsilon)}.
\]

| Proof. | Note \( \Pr[X \geq \epsilon] \leq \Pr[f(X) \geq f(\epsilon)] \) and apply Markov. |

The Chernoff bounding technique is as follows. We can apply the proceeding corollary to the mapping \( t \mapsto \exp(tX) \) for all \( t > 0 \): supposing \( \mathbb{E} X = 0 \),

\[
\Pr[X \geq \epsilon] = \inf_{t \geq 0} \Pr[\exp(tX) \geq \exp(\epsilon t)] \leq \inf_{t \geq 0} \frac{\mathbb{E} \exp(tX)}{\exp(\epsilon t)}.
\]

Simplifying the RHS via sub-Gaussianity,

\[
\inf_{t > 0} \frac{\mathbb{E} \exp(tX)}{\exp(\epsilon t)} \leq \inf_{t > 0} \frac{\exp(t^2 \sigma^2/2)}{\exp(\epsilon t)} = \inf_{t > 0} \left( t^2 \sigma^2/2 - \epsilon t \right)
\]

\[
= \exp \left( \inf_{t > 0} \frac{t^2 \sigma^2/2 - \epsilon t}{\sigma^2} \right).
\]

The minimum of this convex quadratic is \( t := \frac{\epsilon}{2 \sigma^2} > 0 \), thus

\[
\Pr[X \geq \epsilon] = \inf_{t > 0} \frac{\mathbb{E} \exp(tX)}{\exp(\epsilon t)} \leq \exp \left( \inf_{t > 0} \frac{t^2 \sigma^2}{2 \sigma^2} - \epsilon t \right) = \exp \left( -\frac{\epsilon^2}{2 \sigma^2} \right). \tag{5}
\]

What if we apply this to an average of sub-Gaussian r.v.'s? (The point is: this starts to look like an empirical risk!)
Theorem 14.2 (Chernoff bound for subgaussian r.v.’s) Suppose \((X_1, \ldots, X_n)\) independent and respectively \(\sigma_i^2\)-subgaussian. Then
\[
\Pr \left[ \frac{1}{n} \sum_i X_i \geq \epsilon \right] \leq \exp \left( -\frac{n^2 \epsilon^2}{2 \sum_i \sigma_i^2} \right).
\]

In other words (“inversion form”), with probability \(\geq 1 - \delta\),
\[
\frac{1}{n} \sum_i \mathbb{E} X_i \leq \frac{1}{n} \sum_i X_i + \sqrt{\frac{2 \sum_i \sigma_i^2}{n^2} \ln \left( \frac{1}{\delta} \right)}.
\]

**Proof.** \(S_n := \sum_i X_i/n\) is \(\sigma^2\)-subgaussian with \(\sigma^2 = \sum_i \sigma_i^2/n^2\); plug this into the sub-Gaussian tail bound in eq. 5.

**Remark 14.2** (Gaussian sanity check.) Let’s go back to the case \(n = 1\). It’s possible to get a tighter tail for the Gaussian directly (see (Wainwright 2015)), but it only changes log factors in the “inversion form” of the bound. Note also the bound is neat for the Gaussian since it says the tail mass and density are of the same order (algebraically this makes sense, as with geometric series).

(“Inversion” form.) This form is how things are commonly presented in machine learning; think of \(\delta\) as “confidence”; \(\ln(1/\delta)\) term means adding more digits to the confidence (e.g., bound holds with probability 99.999\% means a linear increase in the term \(\ln(1/\delta)\)).

There are more sophisticated bounds (e.g., Bernstein, Freedman, McDiarmid) proved in similar ways, often considering a Martingale rather than IID r.v.s.

[ mjt: I should say something about necessary and sufficient, like convex lipschitz bounded vs lipschitz gaussian.]

[ mjt: maybe give heavy tail pointer? dunno.]

14.2 Hoeffding’s inequality and the need for uniform deviations

Let’s use what we’ve seen to bound misclassifications!

**Theorem 14.3 (Hoeffding inequality)** Given independent \((X_1, \ldots, X_n)\) with \(X_i \in [a_i, b_i]\) a.s.,
\[
\Pr \left[ \frac{1}{n} \sum_i (X_i - \mathbb{E} X_i) \geq \epsilon \right] \leq \exp \left( -\frac{2n^2 \epsilon^2}{\sum_i (b_i - a_i)^2} \right).
\]

**Proof.** Plug Hoeffding Lemma into sub-Gaussian Chernoff bound.

**Example 14.2** Fix classifier \(f\), sample \(((X_i, Y_i))_{i=1}^n\), and define \(Z_i := 1[f(X_i) \neq Y_i]\). With probability at least \(1 - \delta\),
\[
\Pr[f(X) \neq Y] - \frac{1}{n} \sum_{i=1}^n 1[f(x_i) = y_i] = \mathbb{E} Z_1 - \frac{1}{n} \sum_{i=1}^n Z_i \leq \sqrt{\frac{1}{2n} \ln \left( \frac{1}{\delta} \right)}.
\]

As in, test error is upper bounded by training error plus a term which goes \(\downarrow 0\) as \(n \to \infty\)!
Example 14.3 Classifier $f_n$ memorizes training data:

$$f_n(x) := \begin{cases} y_i & x = x_i \in (x_1, \ldots, x_n), \\ 17 & \text{otherwise.} \end{cases}$$

Consider two situations with $\Pr[Y = +1 | X = x] = 1$.

- Suppose marginal on $X$ has finite support. Eventually (large $n$), this support is memorized and $\mathcal{R}_z(f_n) = 0 = \mathcal{R}(f_n)$.
- Suppose marginal on $X$ is continuous. With probability 1, $\mathcal{R}_z(f_n) = 0$ but $\mathcal{R}(f_n) = 1$.

What broke Hoeffding’s inequality (and its proof) between these two examples?

- $f_n$ is a random variable depending on $S = ((x_i, y_i))_{i=1}^n$. Even if $((x_i, y_i))_{i=1}^n$ are independent, the new random variables $Z_i := 1[f_n(x_i) \neq y_i]$ are not.

This $f_n$ overfit: $\mathcal{R}(f_n)$ is small, but $\mathcal{R}(f_n)$ is large.

Possible fixes.

- Two samples: train on $S_1$, evaluate on $S_2$. But now we’re using less data, and run into the same issue if we evaluate multiple predictors on $S_2$.
- Restrict access to data within training algorithm: SGD does this, and has a specialized (martingale-based) deviation analysis.
- Uniform deviations: define a new r.v. controlling errors of all possible predictors $F$ the algorithm might output:

$$\left[ \sup_{f \in F} \mathcal{R}(f) - \mathcal{R}(f) \right].$$

This last one is the approach we’ll follow here. It can be adapted to data and algorithms by adapting $F$ (we’ll discuss this more shortly).

Remark 14.3 There are measure-theoretic issues with the uniform deviation approach, which we’ll omit here. Specifically, the most natural way to reason about

$$\left[ \sup_{f \in F} \mathcal{R}(f) - \mathcal{R}(f) \right]$$

is via uncountably intersections of events, which are not guaranteed to be within the $\sigma$-algebra. The easiest fix is to work with countable subfamilies, which will work for the standard ReLU networks we consider.

15 Rademacher complexity

As before we will apply a brute-force approach to controlling generalization over a function family $F$: we will simultaneously control generalization for all elements of the class by working with the random variable

$$\left[ \sup_{f \in F} \mathcal{R}(f) - \mathcal{R}(f) \right].$$

This is called “uniform deviations” because we prove a deviation bound that holds uniformly over all elements of $F$. 

55
Remark 15.1. The idea is that even though our algorithms output predictors which depend on data, we circumvent the independence issue by invoking a uniform bound on all elements of $\mathcal{F}$ before we see the algorithm’s output, and thus generalization is bounded for the algorithm output (and for everything else in the class). This is a brute-force approach because it potentially controls much more than is necessary.

On the other hand, we can adapt the approach to the output of the algorithm in various ways, as we will discuss after presenting the main Rademacher bound.

Example 15.1 (finite classes). As an example of what is possible, suppose we have $\mathcal{F} = (f_1, \ldots, f_k)$, meaning a finite function class $\mathcal{F}$ with $|\mathcal{F}| = k$. If we apply Hoeffding’s inequality to each element of $\mathcal{F}$ and then union bound, we get, with probability at least $1 - \delta$, for every $f \in \mathcal{F}$,

$$\Pr[f(X) \neq Y] - \hat{\Pr}[f(X) \neq Y] \leq \sqrt{\frac{\ln(k/\delta)}{2n}} \leq \sqrt{\frac{\ln |\mathcal{F}|}{2n}} + \sqrt{\frac{\ln(1/\delta)}{2n}}.$$ 

Rademacher complexity will give us a way to replace $\ln |\mathcal{F}|$ in the preceding finite class example with something non-trivial in the case $|\mathcal{F}| = \infty$.

Definition 15.1 (Rademacher complexity). Given a set of vectors $V \subseteq \mathbb{R}^n$, define the (un-normalized) Rademacher complexity as

$$\text{URad}(V) := E \sup_{u \in V} \langle \epsilon, u \rangle, \quad \text{Rad}(V) := \frac{1}{n} \text{URad}(V),$$

where $E$ is uniform over the corners of the hypercube over $\epsilon \in \{\pm 1\}^n$ (each coordinate $\epsilon_i$ is a Rademacher random variable, meaning $\Pr[\epsilon_i = +1] = \frac{1}{2} = \Pr[\epsilon_i = -1]$, and all coordinates are iid).

This definition can be applied to arbitrary elements of $\mathbb{R}^n$, and is useful outside machine learning. We will typically apply it to the behavior of a function class on $S = (z_i)_{i=1}^n$:

$$\mathcal{F}|_S := \{(f(x_1), \ldots, f(x_n)) : f \in \mathcal{F}\} \subseteq \mathbb{R}^n.$$ 

With this definition,

$$\text{URad}(\mathcal{F}|_S) = E \sup_{\epsilon \in \mathcal{F}|_S} \langle \epsilon, u \rangle = E \sup_{\epsilon \in \mathcal{F}} \sum_{i} \epsilon_i f(z_i).$$

Remark 15.2 (Loss classes). This looks like fitting random signs, but is not exactly that; often we apply it to the loss class: overloading notation,

$$\text{URad}(\ell \circ \mathcal{F})|_S = \text{URad} \{\ell(y_1f(x_1), \ldots, \ell(y_nf(x_n)) : f \in \mathcal{F}\}.$$ 

(Sanity checks.) We’d like $\text{URad}(V)$ to measure how “big” or “complicated” $V$ is. Here are a few basic checks:

1. $\text{URad}(\{u\}) = E \langle \epsilon, u \rangle = 0$; this seems desirable, as a $|V| = 1$ is simple.
2. More generally, $\text{URad}(V + \{u\}) = \text{URad}(V)$.
3. If $V \subseteq V'$, then $\text{URad}(V) \leq \text{URad}(V')$.
4. $\text{URad}(\{\pm 1\}^n) = E_{\epsilon} \epsilon^2 = n$; this also seems desirable, as $V$ is as big/complicated as possible (amongst bounded vectors).
5. \text{URad}\{(−1,\ldots,−1),(+1,\ldots,+1)\} = \mathbb{E}_ε |\sum_i ε_i| = Θ(\sqrt{n}). This also seems reasonable: |V| = 2 and it is not completely trivial.

\textbf{(URad vs Rad.)} I don’t know other texts or even papers which use \text{URad}, I only see \text{Rad}. I prefer \text{URad} for these reasons:

1. The 1/n is a nuisance while proving Rademacher complexity bounds.
2. When we connect Rademacher complexity to covering numbers, we need to change the norms to account for this 1/n.
3. It looks more like a regret quantity.

\textbf{(Absolute value version.)} The original definition of Rademacher complexity (P. L. Bartlett and Mendelson 2002), which still appears in many papers and books, is

\[ \text{URad}_{|!}(V) = \mathbb{E} \sup_{\epsilon \in V} |\langle \epsilon, u \rangle|. \]

Most texts now drop the absolute value. Here are my reasons:

1. \text{URad}_{|!} violates basic sanity checks: it is possible that \text{URad}_{|!}(\{u\}) ≠ 0 and more generally \text{URad}_{|!}(V + \{u\}) ≠ \text{URad}_{|!}(V), which violates my basic intuition about a “complexity measure.”
2. To obtain 1/n rates rather than 1/√n, the notion of local Rademacher complexity was introduced, which necessitated dropping the absolute value essentially due to the preceding sanity checks.
3. We can use \text{URad} to reason about \text{URad}_{|!}, since \text{URad}_{|!}(V) = \text{URad}(V ∪ −V).
4. While \text{URad}_{|!} is more convenient for certain operations, e.g., \text{URad}_{|!}(\bigcup_i V_i) ≤ \sum_i \text{URad}_{|!}(V_i), there are reasonable surrogates for \text{URad} (as below).

The following theorem shows indeed that we can use Rademacher complexity to replace the ln |F| term from the finite-class bound with something more general (we’ll treat the Rademacher complexity of finite classes shortly).

\textbf{Theorem 15.1} Let \( F \) be given with \( f(z) \in [a, b] \) a.s. \( \forall f \in F \).

1. With probability \( \geq 1 − \delta \),

\[ \sup_{f \in F} \mathbb{E} f(Z) − \frac{1}{n} \sum_i f(z_i) ≤ \mathbb{E}_{(z_i)_{i=1}^n} \left( \sup_{f \in F} \mathbb{E} f(z) − \frac{1}{n} \sum_i f(z_i) \right) + (b − a) \sqrt{\frac{\ln(1/\delta)}{2n}}. \]

2. With probability \( \geq 1 − \delta \),

\[ \mathbb{E}_{(z_i)_{i=1}^n} \text{URad}(F|S) ≤ \text{URad}(F|S) + (b − a) \sqrt{\frac{n \ln(1/\delta)}{2}}. \]

3. With probability \( \geq 1 − \delta \),

\[ \sup_{f \in F} \mathbb{E} f(Z) − \frac{1}{n} \sum_i f(z_i) ≤ \frac{2}{n} \text{URad}(F|S) + 3(b − a) \sqrt{\frac{\ln(2/\delta)}{2n}}. \]
Remark 15.3 To flip which side has an expectation and which side has an average of random variables, replace $\mathcal{F}$ with $-\mathcal{F} := \{-f : f \in \mathcal{F}\}$.

The proof of this bound has many interesting points and is spread out over the next few subsections. It has these basic steps:

1. The expected uniform deviations are upper bounded by the expected Rademacher complexity. This itself is done in two steps:
   1. The expected deviations are upper bounded by expected deviations between two finite samples. This is interesting since we could have reasonably defined generalization in terms of this latter quantity.
   2. These two-sample deviations are upper bounded by expected Rademacher complexity by introducing random signs.

2. We replace this difference in expectations with high probability bounds via a more powerful concentration inequality which we haven’t discussed, McDiarmid’s inequality.

15.1 Generalization without concentration; symmetrization

We’ll use further notation throughout this proof.

- $Z$ r.v.; e.g., $(x, y)$,
- $\mathcal{F}$ functions; e.g., $f(Z) = \ell(g(X), Y)$,
- $E_Z$ expectation over $Z$,
- $E_n$ expectation over $(Z_1, \ldots, Z_n)$,
- $E f = E f(Z)$,
- $\hat{E}_n f = \frac{1}{n} \sum_i f(Z_i)$.

In this notation, $R_\ell(g) = E \ell \circ g$ and $\hat{R}_\ell(g) = \hat{E} \ell \circ g$.

15.1.1 Symmetrization with a ghost sample

In this first step we’ll introduce another sample (“ghost sample”). Let $(Z'_1, \ldots, Z'_n)$ be another iid draw from $Z$; define $E'_n$ and $\hat{E}'_n$ analogously.

| Lemma 15.1 | $E_n \left( \sup_{f \in \mathcal{F}} E f - \hat{E}_n f \right) \leq E_n E'_n \left( \sup_{f \in \mathcal{F}} \hat{E}'_n f - \hat{E}_n f \right).$ |
Proof. Fix any $\epsilon > 0$ and apx max $f_\epsilon \in \mathcal{F}$; then

$$
\mathbb{E}\left( \sup_{f \in \mathcal{F}} \mathbb{E} f - \widehat{\mathbb{E}}_{n} f \right) \leq \mathbb{E}\left( \mathbb{E} f_\epsilon - \widehat{\mathbb{E}}_{n} f_\epsilon \right) + \epsilon \\
= \mathbb{E}\left( \frac{1}{n} \widehat{\mathbb{E}}_{n} f_\epsilon - \widehat{\mathbb{E}}_{n} f_\epsilon \right) + \epsilon \\
= \frac{1}{n} \mathbb{E} \left( \widehat{\mathbb{E}}_{n} f_\epsilon - \widehat{\mathbb{E}}_{n} f_\epsilon \right) + \epsilon \\
\leq \mathbb{E}\left( \sup_{f \in \mathcal{F}} \frac{1}{n} \widehat{\mathbb{E}}_{n} f - \widehat{\mathbb{E}}_{n} f \right) + \epsilon
$$

Result follows since $\epsilon > 0$ was arbitrary.

Remark 15.4 As above, in this section we are working only in expectation for now. In the subsequent section, we’ll get high probability bounds. But $\sup_{f \in \mathcal{F}} \mathbb{E} f - \mathbb{E}'_{n} f$ is a random variable; can describe it in many other ways too! (E.g., “asymptotic normality.”)

As mentioned before, the preceding lemma says we can instead work with two samples. Working with two samples could have been our starting point (and definition of generalization): by itself it is a meaningful and interpretable quantity!

15.1.2 Symmetrization with random signs

The second step swaps points between the two samples; a magic trick with random signs boils this down into Rademacher complexity.

Lemma 15.2 $\mathbb{E}_{n} \mathbb{E}'_{n} \sup_{f \in \mathcal{F}} \left( \widehat{\mathbb{E}}_{n} f - \widehat{\mathbb{E}}_{n} f \right) \leq \frac{2}{n} \mathbb{E}_{n} URad(\mathcal{F}|s)$. 

59
Proof. Fix a vector $\epsilon \in \{-1, +1\}^n$ and define a r.v. $(U_i, U'_i) := (Z_i, Z'_i)$ if $\epsilon = 1$ and $(U_i, U'_i) = (Z'_i, Z_i)$ if $\epsilon = -1$. Then
\[
\mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n^f f - \widehat{\mathbb{E}}_n f \right) = \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i (f(Z'_i) - f(Z_i)) \right) \right).
\]
Here’s the big trick: since $(Z_1, \ldots, Z_n, Z'_1, \ldots, Z'_n)$ and $(U_1, \ldots, U_n, U'_1, \ldots, U'_n)$ have same distribution, and $\epsilon$ arbitrary, then (with $\Pr[\epsilon_i = +1] = 1/2$ iid “Rademacher”)
\[
\mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n^f f - \widehat{\mathbb{E}}_n f \right) = \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (f(U'_i) - f(U_i)) \right) \right).
\]
Since similarly replacing $\epsilon_i$ and $-\epsilon_i$ doesn’t change $\mathbb{E}_\epsilon$,
\[
\mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n^f f - \widehat{\mathbb{E}}_n f \right) = \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (f(Z'_i) - f(Z_i)) \right) \right) \leq \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (f(Z'_i) - f'(Z_i)) \right) \right)
\]
\[
= \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (f(Z'_i)) \right) \right) + \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (-f'(Z_i)) \right) \right)
\]
\[
= 2 \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (f(Z_i)) \right) \right) = 2 \mathbb{E}_{\epsilon} \left( \sup_{f \in F} \mathbb{E}_n \left( \frac{1}{n} \sum_i \epsilon_i (f(Z_i)) \right) \right).
\]

15.2 Generalization with concentration

We controlled expected uniform deviations: $\mathbb{E}_n \sup_{f \in F} \mathbb{E} f - \widehat{\mathbb{E}}_n f$.

High probability bounds will follow via concentration inequalities.

Theorem 15.2 (McDiarmid) Suppose $F : \mathbb{R}^n \to \mathbb{R}$ satisfies “bounded differences”: $\forall i \in \{1, \ldots, n\} \exists c_i,$
\[
\sup_{z_1, \ldots, z_i, z'_i} |F(z_1, \ldots, z_i, \ldots, z_n) - F(z_1, \ldots, z'_i, \ldots, z_n)| \leq c_i.
\]

With $\Pr \geq 1 - \delta$,
\[
\mathbb{E}_n F(Z_1, \ldots, Z_n) \leq F(Z_1, \ldots, Z_n) + \sqrt{\frac{\sum_i c_i^2}{2} \ln(1/\delta)}.
\]
I’m omitting the proof. A standard way is via a *Martingale* variant of the Chernoff bounding method. The Martingale adds one point at a time, and sees how things grow.

Hoefding follows by setting $F′(Z) = \sum_i Z_i/n$ and verifying bounded differences $c_i := (b_i - a_i)/n$.

**Proof of Theorem 15.1.**

The third bullet item follows from the first two by union bounding. To prove the first two, it suffices to apply the earlier two lemmas on expectations and verify the quantities satisfy bounded differences with constant $(b - a)/n$ and $(b - a)$, respectively.

For the first quantity, for any $i$ and $(z_1, \ldots, z_n, z_i')$ and writing $z_j' := z_j$ for $z_j \neq z_i$ for convenience,

$$\left| \sup_{f \in \mathcal{F}} \mathbb{E} f - \hat{\mathbb{E}}_n f - \sup_{g \in \mathcal{F}} (\mathbb{E} g - \hat{\mathbb{E}}_n g) \right| = \left| \sup_{f \in \mathcal{F}} \mathbb{E} f - \hat{\mathbb{E}}_n f - \sup_{g \in \mathcal{F}} (\mathbb{E} g - \hat{\mathbb{E}}_n g + g(z_i) - g(z_i')) \right|$$

$$\leq \sup_{h \in \mathcal{F}} \left| \sup_{f \in \mathcal{F}} \mathbb{E} f - \hat{\mathbb{E}}_n f - \sup_{g \in \mathcal{F}} (\mathbb{E} g - \hat{\mathbb{E}}_n g + h(z_i)/n - h(z_i'))/n \right|$$

$$= \sup_{h \in \mathcal{F}} |h(z_i) - h(z_i')|/n$$

$$\leq \frac{b - a}{n}.$$

Using similar notation, and additionally writing $S$ and $S'$ for the two samples, for the Rademacher complexity,

$$\left| \text{URad}(\mathcal{F}|_S) - \text{URad}(\mathcal{F}|_{S'}) \right| = \left| \text{URad}(\mathcal{F}|_S) - \mathbb{E} \sup_{\mathcal{F}} \epsilon \sum_{i=1}^n \epsilon_i f(z_i) \right|$$

$$= \left| \text{URad}(\mathcal{F}|_S) - \mathbb{E} \sup_{\mathcal{F}} \epsilon \sum_{i=1}^n \epsilon_i f(z_i) - \epsilon_i f(z_i) + \epsilon_i f(z_i') \right|$$

$$\leq \sup_{h \in \mathcal{F}} \left| \text{URad}(\mathcal{F}|_S) - \mathbb{E} \sup_{\mathcal{F}} \epsilon \sum_{i=1}^n \epsilon_i f(z_i) - \epsilon_i h(z_i) + \epsilon_i h(z_i') \right|$$

$$\leq \sup_{h \in \mathcal{F}} \mathbb{E} |\epsilon_i h(z_i) + \epsilon_i h(z_i')| \leq (b - a).$$

**Example: basic logistic regression generalization analysis**

Let’s consider logistic regression with bounded weights:

$$\ell(yf(x)) := \ln(1 + \exp(-yf(x))),$$

$$|\ell'| \leq 1,$$

$$\mathcal{F} := \{ w \in \mathbb{R}^d : \|w\| \leq B \},$$

$$(\ell \circ \mathcal{F})|_S := \{(\ell(y_1w^T x_1), \ldots, \ell(y_nw^T x_n)) : \|w\| \leq B \},$$

$$\mathcal{R}_\ell(w) := \mathbb{E}\ell(Yw^TX),$$

$$\hat{\mathcal{R}}_\ell(w) := \frac{1}{n} \sum_i \ell(y_iw^T x_i).$$

The goal is to control $\mathcal{R}_\ell - \hat{\mathcal{R}}_\ell$ over $\mathcal{F}$ via the earlier theorem; our main effort is in controlling $\text{URad}((\ell \circ \mathcal{F})|_S)$.
This has two steps:

- "Peeling" off $\ell$.
- Rademacher complexity of linear predictors.

**Lemma 15.3** Let $\ell : \mathbb{R}^n \rightarrow \mathbb{R}^n$ be a vector of univariate $L$-lipschitz functions. Then $\operatorname{URad}(\ell \circ V) \leq L \cdot \operatorname{URad}(V)$.

**Proof.** The idea of the proof is to "de-symmetrize" and get a difference of coordinates to which we can apply the definition of $L$. To start,

\[
\operatorname{URad}(\ell \circ V) = \mathbb{E} \sup_{u \in V} \sum_{i} \epsilon_i \ell_i(u_i)
\]

\[
= \frac{1}{2} \mathbb{E} \sup_{u, w \in V} \left( \ell_1(u_1) - \ell_1(w_1) + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right)
\]

\[
\leq \frac{1}{2} \mathbb{E} \sup_{u, w \in V} \left( L |u_1 - w_1| + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right).
\]

To get rid of the absolute value, for any $\epsilon$, by considering swapping $u$ and $w$,

\[
\sup_{u, w \in V} \left( L |u_1 - w_1| + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right)
\]

\[
= \max \left\{ \sup_{u, w \in V} \left( L (u_1 - w_1) + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right), \right. \]

\[
\left. \sup_{u, w \in V} \left( L (w_1 - u_1) + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right) \right\}
\]

\[
= \sup_{u, w \in V} \left( L (u_1 - w_1) + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right).
\]

As such,

\[
\operatorname{URad}(\ell \circ V) \leq \frac{1}{2} \mathbb{E} \sup_{u, w \in V} \left( L |u_1 - w_1| + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right)
\]

\[
= \frac{1}{2} \mathbb{E} \sup_{u, w \in V} \left( L (u_1 - w_1) + \sum_{i=2}^{n} \epsilon_i (\ell_i(u_i) + \ell_i(w_i)) \right)
\]

\[
= \mathbb{E} \sup_{u \in V} \left[ L \epsilon_1 u_1 + \sum_{i=2}^{n} \epsilon_i \ell_i(u_i) \right].
\]

Repeating this procedure for the other coordinates gives the bound.

Revisiting our overloaded composition notation:

\[
(\ell \circ f) = ((x, y) \mapsto \ell(\ell(-yf(x)))),
\]

\[
\ell \circ F = \{ \ell \circ f : f \in F \}.
\]

**Corollary 15.1** Suppose $\ell$ is $L$-lipschitz and $\ell \circ F \in [a, b]$ a.s.. With probability $\geq 1 - \delta$, every
\( f \in F \) satisfies
\[
\mathcal{R}_\ell(f) \leq \tilde{\mathcal{R}}_\ell(f) + \frac{2L}{n} \text{URad}(\mathcal{F}|_S) + 3(b - a)\sqrt{\frac{\ln(2/\delta)}{2n}}.
\]

**Proof.** Use the lipschitz composition lemma with
\[
|\ell(-y_if(x_i) - \ell(-y_if'(x_i))| \leq L|y_if(x_i) + y_if'(x_i)|
\]
\[
\leq L|f(x_i) - f'(x_i)|.
\]

Now let’s handle step 2: Rademacher complexity of linear predictors (in \( \ell_2 \)).

**Theorem 15.3** Collect sample \( S := (x_1, \ldots, x_n) \) into rows of \( X \in \mathbb{R}^{n \times d} \).
\[
\text{URad}(\{x \mapsto \langle w, x \rangle : \|w\|_2 \leq B\}|_S) \leq B\|X\|_F.
\]

**Proof.** Fix any \( \epsilon \in \{-1, +1\}^n \). Then
\[
\sup_{\|w\| \leq B} \sum_i \epsilon_i \langle w, x_i \rangle = \sup_{\|w\| \leq B} \left\langle w, \sum_i \epsilon_i x_i \right\rangle = B \left\| \sum_i \epsilon_i x_i \right\|.
\]

We’ll bound this norm with Jensen’s inequality (only inequality in whole proof!):
\[
\mathbb{E} \left\| \sum_i \epsilon_i x_i \right\|^2 = \mathbb{E} \sum_i \epsilon_i x_i \|^2 \leq \mathbb{E} \left( \sum_i \| \epsilon_i x_i \|^2 + \sum_{i,j} \langle \epsilon_i x_i, \epsilon_j x_j \rangle \right) = \mathbb{E} \sum_i \| x_i \|^2 = \|X\|_F^2.
\]

**Remark 15.6** By Khinchine’s inequality, the preceding Rademacher complexity estimate is tight up to constants.

Let’s now return to the logistic regression example!

**Example 15.2 (logistic regression)** Suppose \( \|w\| \leq B \) and \( \|x_i\| \leq 1 \), and the loss is the 1-Lipschitz logistic loss \( \ell_{\log}(z) := \ln(1 + \exp(z)) \). Note \( \ell((\langle w, yx \rangle)) \geq 0 \) and \( \ell((\langle w, yx \rangle)) \leq \ln(2) + \langle w, yx \rangle \leq \ln(2) + B. \)

Combining the main Rademacher bound with the Lipschitz composition lemma and the Rademacher bound on linear predictors, with probability at least \( 1 - \delta \), every \( w \in \mathbb{R}^d \) with \( \|w\| \leq B \) satisfies
\[
\mathcal{R}_\ell(w) \leq \tilde{\mathcal{R}}_\ell(w) + \frac{2B\|X\|_F}{n} + 3(\ln(2) + B)\sqrt{\ln(2/\delta)/(2n)}
\]
\[
\leq \tilde{\mathcal{R}}_\ell(w) + \frac{2B\|X\|_F}{n} + 3(\ln(2) + B)\sqrt{\ln(2/\delta)/(2n)}
\]
\[
\leq \tilde{\mathcal{R}}_\ell(w) + \frac{2B + 3(B + \ln(2))\sqrt{\ln(2/\delta)/2}}{\sqrt{n}}.
\]
Remark 15.7 (Average case vs worst case.) Here we replaced \( \|X\|_F \) with the looser \( \sqrt{n} \).

This bound scales as the SGD logistic regression bound proved via Azuma, despite following a somewhat different route (Azuma and McDiarmid are both proved with Chernoff bounding method; the former approach involves no symmetrization, whereas the latter holds for more than the output of an algorithm).

It would be nice to have an “average Lipschitz” bound rather than “worst-case Lipschitz”; e.g., when working with neural networks and the ReLU, which seems it can kill off many inputs! But it’s not clear how to do this. Relatedly: regularizing the gradient is sometimes used in practice?

15.4 Margin bounds

In the logistic regression example, we peeled off the loss and bounded the Rademacher complexity of the predictors.

If most training labels are predicted not only accurately, but with a large margin, as in section 12, then we can further reduce the generalization bound.

Define \( \ell_{\gamma}(z) := \max\{0, \min\{1, 1 - z/\gamma\}\} \), \( \mathcal{R}_\gamma(f) := \mathbf{R}_{\ell_{\gamma}}(f) = \mathbb{E} \ell_{\gamma}(Yf(X)) \), and recall \( \mathcal{R}_z(f) = \Pr[f(X) \neq Y] \).

Theorem 15.4 For any margin \( \gamma > 0 \), with probability \( \geq 1 - \delta \), \( \forall f \in \mathcal{F} \),

\[
\mathcal{R}_z(f) \leq \mathcal{R}_\gamma(f) \leq \hat{\mathcal{R}}_\gamma(f) + \frac{2}{n\gamma} \text{URad}(\mathcal{F}) + 3\sqrt{\frac{\ln(2/\delta)}{2n}}.
\]

Proof. Since

\[
1[\text{sgn}(f(x)) \neq y] \leq 1[-f(x)y \geq 0] \leq \ell_{\gamma}(f(x)y),
\]

then \( \mathcal{R}_z(f) \leq \mathcal{R}_\gamma(f) \). The bound between \( \mathcal{R}_\gamma \) and \( \hat{\mathcal{R}}_\gamma \) follows from the fundamental Rademacher bound, and by peeling the \( \frac{1}{\gamma} \)-Lipschitz function \( \ell_{\gamma} \).

[ mjt: is that using per-example lipschitz? need to restate peeling? also, properly invoke peeling?]

Remark 15.8 (bibliographic notes) As a generalization notion, this was first introduced for 2-layer networks in (P. L. Bartlett 1996), and then carried to many other settings (SVM, boosting, ...)

There are many different proof schemes; another one uses sparsification (Schapire et al. 1997).

This approach is again being extensively used for deep networks, since it seems that while weight matrix norms grow indefinitely, the margins grow along with them (P. Bartlett, Foster, and Telgarsky 2017).

15.5 Finite class bounds

In our warm-up example of finite classes, our complexity term was \( \ln |\mathcal{F}| \). Here we will recover that, via Rademacher complexity. Moreover, the bound has a special form which will be useful in the later VC dimension and especially covering sections.
**Theorem 15.5 (Massart finite lemma)** \( \text{URad}(V) \leq \sup_{u \in V} \|u\|_2 \sqrt{2 \ln |V|} \).

**Remark 15.9** \( \ln |V| \) is what we expect from union bound.

The \( \| \cdot \|_2 \) geometry here is intrinsic here; I don’t know how to replace it with other norms without introducing looseness. This matters later when we encounter the Dudley Entropy integral.

We’ll prove this via a few lemmas.

**Lemma 15.4** If \((X_1, \ldots, X_n)\) are \(c^2\)-subgaussian, then \( \mathbb{E} \max_i X_i \leq c \sqrt{2 \ln(n)} \).

**Proof.**

\[
\mathbb{E} \max_i X_i = \inf_{t > 0} \mathbb{E} \frac{1}{t} \ln \max_i \exp(t X_i) \leq \inf_{t > 0} \mathbb{E} \frac{1}{t} \ln \sum_i \exp(t X_i) \\
\leq \inf_{t > 0} \frac{1}{t} \ln \sum_i \mathbb{E} \exp(t X_i) \leq \inf_{t > 0} \frac{1}{t} \ln \sum_i \exp(t^2 c^2 / 2) \\
= \inf_{t > 0} (\ln(n)/t + c^2 t/2)
\]

and plug in minimizer \( t = \sqrt{2 \ln(n)/c^2} \)

**Lemma 15.5** If \((X_1, \ldots, X_n)\) are \(c_i^2\)-subgaussian and independent, \( \sum_i X_i \) is \( \| \bar{c} \|_2^2 \)-subgaussian.

**Proof.** We did this in the concentration lecture, but here it is again:

\[
\mathbb{E} \exp(t \sum_i X_i) = \prod_i \mathbb{E} \exp(t X_i) \leq \prod_i \exp(t^2 c_i^2 / 2) = \exp(t^2 \| \bar{c} \|_2^2 / 2).
\]

**Proof of +Theorem 15.5 (Massart finite lemma).**

Let \( \epsilon \) be iid Rademacher and fix \( u \in V \). Define \( X_{u,i} := \epsilon_i u_i \) and \( X_u := \sum_i X_{u,i} \).

By Hoeffding lemma, \( X_{u,i} \) is \((u_i - u_i)^2/4 = u_i^2 - \) -subgaussian, thus (by Lemma) \( X_u \) is \( \|u\|_2^2 \)-subgaussian. Thus

\[
\text{URad}(V) = \mathbb{E} \max_{\epsilon \in V} \langle \epsilon, u \rangle = \mathbb{E} \max_{u \in V} X_u \leq \max_{u \in V} \|u\|_2 \sqrt{2 \ln |V|}.
\]

### 15.6 Weaknesses of Rademacher complexity

[mjt: not an exhaustive list…]

- The bounds we will prove shortly are all loose. To some extent, it was argued in (Neyshabur, Tomioka, and Srebro 2014; Zhang et al. 2017) and (Nagarajan and Kolter 2019) that this may be intrinsic to Rademacher complexity, though these arguments can be overturned in various settings (in the former, via \textit{a posteriori} bounds, e.g., as obtained via union bound; in the latter case, by considering a modified set of good predictors for the same problem); as such, that particular criticism is unclear. An alternative approach was highlighted in (Dziugaite and Roy 2017), however the bounds produced there are averages over some collection of predictors, and not directly comparable to the bounds here. Overall, though, many authors are investigating alternatives to the definition of generalization.
Looking outside the specific setting of neural network generalization, Rademacher complexity has been widely adopted since, to a great extent, it can cleanly re-prove many existing bounds, and moreover elements of Rademacher complexity proofs exist many decades prior to the coining of the term (P. L. Bartlett and Mendelson 2002). However, already in these settings, Rademacher complexity has extensive weaknesses.

- For many learning problems, extensive effort was put into fast or optimal learning rates, which often boiled down to replacing a $1/\sqrt{n}$ dependency with a $1/n$. While Local Rademacher Complexity is able to recover some of these bounds, it does not seem to recover all of them, and moreover the proofs are often very complicated.

- In many non-parametric learning settings, for example $k$-nearest-neighbor, the best bounds all use a direct analysis (Chaudhuri and Dasgupta 2014), and attempts to recover these analyses with Rademacher complexity have been unsuccessful.

- Closer to the investigation in these lecture notes, there are even cases where a direct Martingale analysis of SGD slightly beats the application of uniform convergence to the output of gradient descent, and similarly to the preceding case, attempts to close this gap have been unsuccessful (Ji and Telgarsky 2019a).

### 16 Two Rademacher complexity proofs for deep networks

We will give two bounds, obtained by inductively peeling off layers.

- One will depend on $\|W^T_i\|_{1,\infty}$. This bound has a pretty clean proof, and appeared in (P. L. Bartlett and Mendelson 2002).

- The other will depend on $\|W^T_i\|_F$, and is more recent (Golowich, Rakhlin, and Shamir 2018).

[ mjt: also i didn’t mention yet that the other proof techniques reduce to this one? ]

#### 16.1 First “layer peeling” proof: $(1, \infty)$ norm

**Theorem 16.1** Let $\rho$-Lipschitz activations $\sigma_i$ satisfy $\sigma_i(0) = 0$, and

$$\mathcal{F} := \{x \mapsto \sigma_L(W_L\sigma_L(\cdots \sigma_1(W_1x)\cdots)) : \|W^T_i\|_{1,\infty} \leq B\}$$

Then $\text{URad}(\mathcal{F}|_\mathcal{S}) \leq \|X\|_{2,\infty}(2\rho B)^L \sqrt{2\ln(d)}$.

**Remark 16.1** Notation $\|M\|_{b,c} = \|(\|M_1\|_b, \ldots, \|M_d\|_b\|_c$ means apply $b$-norm to columns, then $c$-norm to resulting vector.

Many newer bounds replace $\|W^T_i\|$ with a distance to initialization. (The NTK is one regime where this helps.) I don’t know how to use distance to initialize in the bounds in this section, but a later bound can handle it.

$(\rho B)^L$ is roughly a Lipschitz constant of the network according to $\infty$-norm bounded inputs. Ideally we’d have “average Lipschitz” not “worst case,” but we’re still far from that…

The factor $2^L$ is not good and the next section gives one technique to remove it.
We’ll prove this with an induction “peeling” off layers. This peeling will use the following lemma, which collects many standard Rademacher properties.

**Lemma 16.1**
1. $\text{URad}(V) \geq 0$.
2. $\text{URad}(cV + \{u\}) = |c| \text{URad}(V)$.
3. $\text{URad}(\text{conv}(V)) = \text{URad}(V)$.
4. Let $(V_i)_{i \geq 0}$ be given with $\sup_{u \in V_i} \langle u, \epsilon \rangle \geq 0 \forall \epsilon \in \{-1, +1\}^n$. (E.g., $V_i = -V_i$, or $0 \in V_i$.) Then $\text{URad}(\bigcup_i V_i) \leq \sum_i \text{URad}(V_i)$.
5. $\text{URad}(V) = \text{URad}(-V)$.

**Remark 16.2**
(3) is a mixed blessing: “Rademacher is insensitive to convex hulls,”
(4) is true for $\text{URad}_{|\cdot|}$ directly, where $\text{URad}_{|\cdot|}(V) = \mathbb{E}_\epsilon \sup_{u \in V} |\langle \epsilon, u \rangle|$ is the original definition of (unnormalized) Rademacher complexity: define $W_i := V_i \cup -V_i$, which satisfies the conditions, and note $(\bigcup_i V_i) \cup - (\bigcup_i V_i) = \bigcup_i W_i$. Since $\text{URad}_{|\cdot|}(V_i) = \text{URad}(W_i)$, then $\text{URad}_{|\cdot|}(\bigcup_i V_i) = \text{URad}(\bigcup_i W_i) \leq \sum_{i \geq 1} \text{URad}(W_i) = \sum_{i \geq 1} \text{URad}_{|\cdot|}(V_i)$. [mjt*: is this where I messed up and clipped an older Urada remark?]

(6) is important and we’ll do the proof and some implications in homework.
Proof of Lemma 16.1.

1. Fix any \( u_0 \in V \); then \( \mathbb{E}_\epsilon \sup_{u \in V} \langle \epsilon, v \rangle \geq \mathbb{E}_\epsilon \langle \epsilon, u_0 \rangle = 0 \).

2. Can get inequality with \(|c|\)-Lipschitz functions \( \ell_i(r) := c \cdot r + u_i \); for equality, note \( -\epsilon c \) and \( \epsilon c \) are same in distribution.

3. This follows since optimization over a polytope is achieved at a corner. In detail,

\[
\text{URad}(\text{conv}(V)) = \mathbb{E} \sup_{\epsilon} \sup_{u \in V} \sup_{\alpha \in \Delta_k} \langle \epsilon, \sum_j \alpha_j u_j \rangle \\
= \mathbb{E} \sup_{\epsilon} \sum_{k \geq 1} \sum_{j \in V} \sup_{u \in V} \langle \epsilon, u_j \rangle \\
= \mathbb{E} \left( \sup_{k \geq 1} \sum_{j \in V} \sup_{u \in V} \langle \epsilon, u \rangle \right) \\
= \text{URad}(V).
\]

4. Using the condition,

\[
\mathbb{E} \sup_{\epsilon} \sup_{u \in V} \langle \epsilon, u \rangle = \mathbb{E} \sup_{\epsilon} \sup_{i} \sup_{u \in V} \langle \epsilon, u \rangle \leq \mathbb{E} \sum_{i} \sup_{u \in V} \langle \epsilon, u \rangle \\
= \sum_{i \geq 1} \text{URad}(V_i).
\]

5. Since integrating over \( \epsilon \) is the same as integrating over \( -\epsilon \) (the two are equivalent distributions),

\[
\text{URad}(-V) = \mathbb{E} \sup_{\epsilon} \langle \epsilon, -u \rangle = \mathbb{E} \sup_{\epsilon} \langle -\epsilon, -u \rangle = \text{URad}(V).
\]
Proof of **Theorem 16.1.**

Let \( F_i \) denote functions computed by nodes in layer \( i \). It’ll be shown by induction that

\[
URad((F_i)|S) \leq \|X\|_{2,\infty}(2\rho B)^i \sqrt{2\ln(d)}.
\]

**Base case** \((i = 0)\): by the Massart finite lemma,

\[
URad((F_i)|S) = URad \left( \{x \mapsto x_j : j \in \{1, \ldots, d\} \} | S \right) \\
\leq \left( \max_j \| (x_1)_j, \ldots, (x_n)_j \|_2 \right) \sqrt{2\ln(d)} \\
= \|X\|_{2,\infty} \sqrt{2\ln d} = \|X\|_{2,\infty}(2\rho B)^0 \sqrt{2\ln d}.
\]

**Inductive step.** Since \(0 = \sigma((0,F(x))) \in F_{i+1}\), applying both Lipschitz peeling and the preceding multi-part lemma,

\[
URad((F_{i+1})|S) = URad \left( \{x \mapsto \sigma_{i+1}(\|W_{i+1}\|_1,\infty,g(x)) : g \in \text{conv}(-F_i \cup F_i) \} | S \right) \\
= \rho B \cdot URad \left( - (F_i)|S \cup (F_i)|S \right) \\
\leq 2\rho B \cdot URad \left( (F_i)|S \right) \\
\leq (2\rho B)^{i+1} \|X\|_{2,\infty} \sqrt{2\ln d}.
\]

**Remark 16.3** There are many related norm-based proofs now changing constants and also \((1,\infty)\); see for instance Neyshabur-Tomioka-Srebro, Bartlett-Foster-Telgarsky (we’ll cover this), Golowich-Rakhlin-Shamir (we’ll cover this), Barron-Klusowski.

The best lower bound is roughly what you get by writing a linear function as a deep network \(\tilde{\cdot}\).

The proof does not “coordinate” the behavior of adjacent layers in any way, and worst-cases what can happen.

### 16.2 Second “layer peeling” proof: Frobenius norm

**Theorem 16.2 (\((\text{Theorem 1, Golowich, Rakhlin, and Shamir 2018})\)** Let 1-Lipschitz positive homogeneous activation \(\sigma_i\) be given, and

\[
F := \{x \mapsto \sigma_L(W_L\sigma_{L-1}(\cdots \sigma_1(W_1x) \cdots )) : \|W_i\|_F \leq B \}.
\]

Then

\[
URad(F|S) \leq B^L\|X\|_F \left( 1 + \sqrt{2L\ln(2)} \right).
\]

**Remark 16.4** The criticisms of the previous layer peeling proof still apply, except we’ve removed \(2^L\).

The proof technique can also handle other matrix norms (though with some adjustment), bringing it closer to the previous layer peeling proof.
For an earlier version of this bound but including things like $2^L$, see Neyshabur-Tomioka-Srebro.

The main proof trick (to remove $2^L$) is to replace $\mathbb{E}_\epsilon$ with $\ln \mathbb{E}_\epsilon \exp$; the $2^L$ now appears inside the $\ln$.

To make this work, we need two calculations, which we’ll wrap up into lemmas.

- When we do “Lipschitz peeling,” we now have to deal with $\exp$ inside $\mathbb{E}_\epsilon$. Magically, things still work, but the proof is nastier, and we’ll not include it.
- That base case of the previous layer peeling could by handled by the Massart finite lemma; this time we end up with something of the form $\mathbb{E}_\epsilon \exp(t \|X^\epsilon\|_2)$.
- Comparing to the $\infty \to \infty$ operator norm (aka $(1, \infty)$) bound, let’s suppose $W \in \mathbb{R}^{m \times m}$ has row/node norm $\|W_j\|_2 \approx 1$, thus $\|W_j\|_1 \approx \sqrt{m}$, and

$$\|W\|_F \approx \sqrt{m} \approx \|W^T\|_{1,\infty},$$

so this bound only really improves on the previous one by removing $2^L$?

Here is our refined Lipschitz peeling bound, stated without proof.

**Lemma 16.2** *(Eq. 4.20, Ledoux and Talagrand 1991)* Let $\ell : \mathbb{R}^n \to \mathbb{R}^n$ be a vector of univariate $\rho$-lipschitz functions with $\ell_i(0) = 0$. Then

$$\mathbb{E} \exp \left( \sup_{u \in V} \sum_i \epsilon_i \ell_i(u_i) \right) \leq \mathbb{E} \exp \left( \rho \sup_{u \in V} \sum_i \epsilon_i u_i \right).$$

**Remark 16.5** With $\exp$ gone, our proof was pretty clean, but all proofs I know of this are more complicated case analyses. So I will not include a proof $\sim$.

The peeling proof will end with a term $\mathbb{E} \exp( t \|X^\epsilon\|_2)$, and we’ll optimize the $t$ to get the final bound. Consequently, we are proving $\|X^\epsilon\|$ is sub-Gaussian!

**Lemma 16.3** $\mathbb{E} \|X^\epsilon\|_2 \leq \|X\|_F$ and $\|X^\epsilon\|$ is $(\mathbb{E} \|X^\epsilon\|, \|X\|_F^2)$-sub-Gaussian.
Proof. Following the notation of (Wainwright 2015), define

\[ Y_k := \mathbb{E}[\|X^T \epsilon\|_2 | \epsilon_1, \ldots, \epsilon_k], \]
\[ D_k := Y_k - Y_{k-1}, \]

whereby \( Y_n - Y_0 = \sum_k D_k \). For the base case, as usual

\[ \mathbb{E}\|X^T \epsilon\|_2 \leq \sqrt{\mathbb{E}\|X^T \epsilon\|^2} = \sqrt{\sum_{j=1}^d \mathbb{E}(X^T_j \epsilon)^2} = \sqrt{\sum_{j=1}^d \|X_j\|^2} = \|X\|_F. \]

Supposing \( \epsilon \) and \( \epsilon' \) only differ on \( \epsilon_k \),

\[ \sup_{\epsilon_k} \|X^T \epsilon\| - \|X^T \epsilon'\|^2 \leq \sup_{\epsilon_k} \|X^T (\epsilon - \epsilon')\|^2 = \sup_{\epsilon_k} \sum_{j=1}^d (X^T_j (\epsilon - \epsilon'))^2 \]
\[ = \sup_{\epsilon_k} \sum_{j=1}^d (X_{k,j}(\epsilon_k - \epsilon'_k))^2 \leq 4\|X_{k,:}\|^2, \]

therefore by the (conditional) Hoeffding lemma, \( D_k \) is \( \|X_{k,:}\|^2 \)-sub-Gaussian, thus (Theorem 2.3, Wainwright 2015) grants \( \sum_k D_k \) is \( \sigma^2 \)-sub-Gaussian with \( \sigma^2 = \sum_k \|X_{k,:}\|^2 = \|X\|_F^2 \).

Remark 16.6 (pointed out by Ziwei Ji) Alternatively, we can use the Lipschitz-convex concentration bound for bounded random variables, and get a variance proxy of roughly \( \|X\|_2 \). Plugging this into the full peeling proof, we get an interesting bound

\[ B^L (\|X\|_F + \|X\|_2 \sqrt{128L \ln(2)}) \text{, thus dimension and depth don’t appear together.} \]
Proof of Theorem 16.2 ((Theorem 1, Golowich, Rakhlin, and Shamir 2018)). For convenience, let $X_i$ denote the output of layer $i$, meaning

$$X_0 = X \quad \text{and} \quad X_i := \sigma_i(X_{i-1}W_i^T).$$

Let $t > 0$ be a free parameter and let $w$ denote all parameters across all layers; the bulk of the proof will show (by induction on layers) that

$$\mathbb{E} \sup_w \exp(t^\top |X_i|) \leq \mathbb{E} 2^t \exp(tB^i |X_0|).$$

To see how to complete the proof from here, note by the earlier “base case lemma” (setting $\mu := \mathbb{E} |X_0|\epsilon_i$ for convenience) and Jensen’s inequality that

$$\text{URad}(F|S) = \mathbb{E} \sup_w \epsilon^\top X_L = \mathbb{E} \frac{1}{t} \ln \mathbb{E} \sup_w \exp(t\epsilon^\top X_L)$$

$$\leq \frac{1}{t} \ln \mathbb{E} \sup_w \exp(t|\epsilon^\top X_L|) \leq \frac{1}{t} \ln \mathbb{E} 2^t \exp(tB^L |\epsilon^\top X_0|)$$

$$\leq \frac{1}{t} \ln 2^t \exp(tB^L (|\epsilon^\top X_0| - \mu + \mu))$$

$$\leq \frac{1}{t} \ln \left[2^t \exp\left(t^2 B^{2L} |X|^2_F / 2 + tB^L \mu \right)\right]$$

$$\leq \frac{L \ln 2}{t} + \frac{tB^{2L} |X|^2_F}{2} + B^L |X|_F,$$

whereby the final bound follows with the minimizing choice

$$t := \sqrt{\frac{2L \ln(2)}{B^{2L} |X|^2_F}} \quad \Rightarrow \quad \text{URad}(F|S) \leq \sqrt{2 \ln(2) LB^{2L} |X|^2_F} + B^L |X|_F.$$

The main inequality is now proved via induction.

For convenience, define $\sigma := \sigma_i$ and $Y := X_{i-1}$ and $V := W_i$ and $\tilde{V}$ has $\ell_2$-normalized rows. By positive homogeneity and definition,

$$\sup_w |\epsilon^\top X_i|^2 = \sup_w \sum_j (\epsilon^\top \sigma(YV^T)_)^2$$

$$= \sup_w \sum_j (\epsilon^\top \sigma(YV^T)_j)^2$$

$$= \sup_w \sum_j (\epsilon^\top \sigma(||V_j||YV^T)_j)^2$$

$$= \sup_w \sum_j ||V_j||^2 (\epsilon^\top \sigma(YV^T)_j)^2.$$

The maximum over row norms is attained by placing all mass on a single row; thus, letting $u$ denote an arbitrary unit norm (column) vector, and finally applying the peeling lemma, and re-introducing the dropped terms, and closing with the IH,

$$\mathbb{E} \exp \left( t \sqrt{\sup_w |\epsilon^\top X_i|^2} \right) = \mathbb{E} \exp \left( t \sqrt{\sup_{w,u} B^2 (\epsilon^\top \sigma(Yu))^2} \right)$$

$$= \mathbb{E} \sup_{w,u} \exp \left( tB (\epsilon^\top \sigma(Yu)) \right)$$

$$\leq \mathbb{E} \sup_{w,u} \exp \left( tB (\epsilon^\top \sigma(Yu)) \right) - \exp \left( -tB (\epsilon^\top \sigma(Yu)) \right)$$

$$\leq \frac{1}{72} \mathbb{E} \sup_{w,u} \exp \left( tB (\epsilon^\top \sigma(Yu)) \right) + \mathbb{E} \sup_{w,u} \exp \left( -tB (\epsilon^\top \sigma(Yu)) \right)$$

$$= \mathbb{E} 2 \sup_{w,u} \exp \left( tB (\epsilon^\top \sigma(Yu)) \right)$$
Covering numbers

- Covering numbers are another way to do generalization. Covering numbers and Rademacher complexities are in some usual settings nearly tight with each other, though in these lectures we will only produce a way to upper bound Rademacher complexity with covering numbers.

- Covering numbers are a classical concept. The idea is we discretize or cover the function class with some finite collection of representative elements; in this way, it’s tight to the “totally bounded” definition of compact set. Their first use in a statistical context is due to (Kolmogorov and Tikhomirov 1959).

- [ mjt: i should discuss relating it to uniform convergence via rademacher, and how we have two ways, and neither is really tight, need chaining, and pointer to vershynin maybe.]

**Definition 17.1**  Given a set $U$, scale $\epsilon$, norm $\| \cdot \|$, $V \subseteq U$ is a (proper) cover when

$$\sup_{a \in U} \inf_{b \in V} \| a - b \| \leq \epsilon.$$  

Let $\mathcal{N}(U, \epsilon, \| \cdot \|)$ denote the covering number: the minimum cardinality (proper) cover.

**Remark 17.1**  “Improper” covers drop the requirement $V \subseteq U$. (We’ll come back to this.)

Most treatments define special norms with normalization $1/n$ baked in; we’ll use unnormalized Rademacher complexity and covering numbers.

Although the definition can handle directly covering functions $\mathcal{F}$, we get nice bounds by covering $\mathcal{F}|_S$, and conceptually it also becomes easier, just a vector (or matrix) covering problem with vector (and matrix) norms.

**17.1 Basic Rademacher-covering relationship**

**Theorem 17.1**  Given $U \subseteq \mathbb{R}^n$,

$$\text{URad}(U) \leq \inf_{\alpha > 0} \left( \alpha \sqrt{n} + \left( \sup_{a \in U} \| a \|_2 \right) \sqrt{2 \ln \mathcal{N}(U, \alpha, \| \cdot \|_2)} \right).$$

**Remark 17.2**  $\| \cdot \|_2$ comes from applying Massart. It’s unclear how to handle other norms without some technical slop.
\textbf{Proof.} Let $\alpha > 0$ be arbitrary, and suppose $\mathcal{N}(U, \alpha, \| \cdot \|_2) < \infty$ (otherwise bound holds trivially). Let $V$ denote a minimal cover, and $V(a)$ its closest element to $a \in U$. Then

\[
\text{URad}(U) = \mathbb{E} \sup_{a \in U} \langle \epsilon, a \rangle \\
= \mathbb{E} \sup_{a \in U} \langle \epsilon, a - V(a) + V(a) \rangle \\
= \mathbb{E} \sup_{a \in U} (\langle \epsilon, V(a) \rangle + \langle \epsilon, a - V(a) \rangle) \\
\leq \mathbb{E} \sup_{a \in U} (\langle \epsilon, V(a) \rangle + \| \epsilon \| \cdot \| a - V(a) \|) \\
\leq \text{URad}(V) + \alpha \sqrt{n} \\
\leq \sup_{b \in V} \| b \|_2 \sqrt{2 \ln |V|} + \alpha \sqrt{n} \\
\leq \sup_{a \in U} \| a \|_2 \sqrt{2 \ln |V|} + \alpha \sqrt{n},
\]

and the bound follows since $\alpha > 0$ was arbitrary.

\begin{remark}
The same proof handles improper covers with minor adjustment: for every $b \in V$, there must be $U(b) \in U$ with $\| b - U(v) \| \leq \alpha$ (otherwise, $b$ can be moved closer to $U$), thus

\[
\sup_{b \in V} \| b \|_2 \leq \sup_{b \in V} \| b - U(b) \|_2 + \| U(b) \|_2 \leq \alpha + \sup_{a \in U} \| a \|_2.
\]

To handle other norms, superficially we need two adjustments: Cauchy-Schwarz can be replaced with Hölder, but it’s unclear how to replace Massart without slop relating different norms.
\end{remark}

\section{Second Rademacher-covering relationship: Dudley’s entropy integral}

There is a classical proof that says that covering numbers and Rademacher complexities are roughly the same; the upper bound uses the Dudley entropy integral, and the lower bound uses a “Sudakov lower bound” which we will not include here.

\begin{itemize}
\item The Dudley entropy integral works at \textit{multiple scales}.
\begin{itemize}
\item Suppose we have covers $(V_N, V_{N-1}, \ldots)$ at scales $(\alpha_N, \alpha_N/2, \alpha_N/4, \ldots)$.
\item Given $a \in U$, choosing $V_i(a) := \arg \min_{b \in V_i} \| a - b \|$, 
\[ a = (a - V_N(a)) + (V_N(a) - V_{N-1}(a)) + (V_{N-1}(a) - V_{N-2}(a)) + \cdots. \]
\end{itemize}
\item We are thus rewriting $a$ as a sequence of \textbf{increments} at different scales.
\begin{itemize}
\item One way to think of it is as writing a number as its binary expansion 
\[ x = (0, b_1 b_2 b_3 \ldots) = \sum_{i \geq 1} \frac{(b_i b_{i+1} \ldots) - (0, b_{i+1} \ldots)}{2^i} = \sum_{i \geq 1} \frac{b_i}{2^i}. \]
\end{itemize}
\end{itemize}

In the Dudley entropy integral, we are covering these \textbf{increments} $b_i$, rather than the number $x$ directly.
One can cover increments via covering numbers for the base set, and that is why these basic covering numbers appear in the Dudley entropy integral. But internally, the argument really is about these increments.

[ mjt$: Seems this works with improper covers. I should check carefully and include it in the statement or a remark.]

[ mjt$: citation for dudley? to dudley lol?]

**Theorem 17.2 (Dudley)** Let $U \subseteq [-1, +1]^n$ be given with $0 \in U$.

\[
\text{URad}(U) \leq \inf_{N \in \mathbb{Z} \geq 1} \left( n \cdot 2^{-N+1} + 6\sqrt{n} \sum_{i=1}^{N-1} 2^{-i} \sqrt{\ln N(U, 2^{-i}\sqrt{n}, \|\cdot\|_2)} \right)
\leq \inf_{\alpha > 0} \left( 4\alpha \sqrt{n} + 12 \int_{\alpha}^{\sqrt{n}/2} \frac{\sqrt{\ln N(U, \beta, \|\cdot\|_2)} d\beta}{\beta} \right).
\]
Proof. We’ll do the discrete sum first. The integral follows by relating an integral to its Riemann sum.

- Let $N \geq 1$ be arbitrary.
- For $i \in \{1, \ldots, N\}$, define scales $\alpha_i := \sqrt{n}2^{1-i}$.
- Define cover $V_1 := \{0\}$; since $U \subseteq [-1, +1]^n$, this is a minimal cover at scale $\sqrt{n} = \alpha_1$.
- Let $V_i$ for $i \in \{2, \ldots, N\}$ denote any minimal cover at scale $\alpha_i$, meaning $|V_i| = \mathcal{N}(U, \alpha_i, \|\cdot\|_2)$.

Since $U \ni a = (a - V_N(a)) + \sum_{i=1}^{N-1} (V_{i+1}(a) - V_i(a)) + V_1(a)$,

\[
\text{URad}(U) = \mathbb{E} \sup_{a \in U} \langle \epsilon, a \rangle
\]

\[
= \mathbb{E} \sup_{a \in U} \left( \langle \epsilon, a - V_N(a) \rangle + \sum_{i=1}^{N-1} \langle \epsilon, V_{i+1}(a) - V_i(a) \rangle + \langle \epsilon, V_1(a) \rangle \right)
\]

\[
\leq \mathbb{E} \sup_{a \in U} \langle \epsilon, a - V_N(a) \rangle
\]

\[
+ \sum_{i=1}^{N-1} \mathbb{E} \sup_{a \in U} \langle \epsilon, V_{i+1} - V_i(a) \rangle
\]

\[
+ \mathbb{E} \sup_{a \in U} \langle \epsilon, V_1(a) \rangle.
\]

Let’s now control these terms separately. The first and last terms are easy:

\[
\mathbb{E} \sup_{a \in U} \epsilon V_1(a) = \mathbb{E} \langle \epsilon, 0 \rangle = 0,
\]

\[
\mathbb{E} \sup_{a \in U} \langle \epsilon, a - V_N(a) \rangle \leq \mathbb{E} \sup_{a \in U} \|\epsilon\| \|a - V_N(a)\| \leq \sqrt{n} \alpha_N = n 2^{1-N}.
\]

For the middle term, define increment class $W_i := \{V_{i+1}(a) - V_i(a) : a \in U\}$, whereby $|W_i| \leq |V_{i+1}| \cdot |V_i| \leq |V_{i+1}|^2$, and

\[
\mathbb{E} \sup_{a \in U} \langle \epsilon, V_{i+1}(a) - V_i(a) \rangle = \text{URad}(W_i)
\]

\[
\leq \left( \sup_{w \in W_i} \|w\|_2 \right) \sqrt{2 \ln |W_i|} \leq \left( \sup_{w \in W_i} \|w\|_2 \right) \sqrt{4 \ln |V_{i+1}|},
\]

\[
\sup_{w \in W_i} \|w\| \leq \sup_{a \in U} \|V_{i+1}\| + \|a - V_i(a)\| \leq \alpha_{i+1} + \alpha_i = 3 \alpha_{i+1}.
\]

Combining these bounds,

\[
\text{URad}(U) \leq n 2^{1-N} + 0 + \sum_{i=1}^{N} 6 \sqrt{n} 2^{-i} \sqrt{\ln \mathcal{N}(U, 2^{-i} \sqrt{n}, \|\cdot\|_2)}.
\]

$N \geq 1$ was arbitrary, so applying $\inf_{N \geq 1}$ gives the first bound. Since $\ln \mathcal{N}(U, \beta, \|\cdot\|_2)$ is nonincreasing in $\beta$, the integral upper bounds the Riemann sum:

\[
\text{URad}(U) \leq n 2^{1-N} + 6 \sum_{i=1}^{N-1} \alpha_{i+1} \sqrt{\ln \mathcal{N}(U, \alpha_{i+1}, \|\cdot\|_2)}
\]

\[
= n 2^{1-N} + 12 \sum_{i=1}^{N-1} (\alpha_{i+1} - \alpha_{i+2}) \sqrt{\ln \mathcal{N}(U, \alpha_{i+1}, \|\cdot\|_2)}
\]

\[
\leq \sqrt{n} \alpha_N + 12 \int_{\alpha_{N+1}}^{\alpha_2} \frac{76}{\alpha_{N+1} \sqrt{\ln \mathcal{N}(U, \alpha_{i+1}, \|\cdot\|_2)}} d\beta.
\]

To finish, pick $\alpha > 0$ and $N$ with
Remark 17.4  Tightness of Dudley: Sudakov’s lower bound says there exists a universal $C$ with

$$\text{URad}(U) \geq \frac{c}{\ln(n)} \sup_{\alpha > 0} \alpha \sqrt{\ln \mathcal{N}(U, \alpha \cdot \| \cdot \|)},$$

which implies $\text{URad}(U) = \Theta(\text{Dudley entropy integral})$. [mjt@: needs references, detail, explanation.]

Taking the notion of increments to heart and generalizing the proof gives the concept of chaining. One key question there is tightening the relationship with Rademacher complexity (shrinking constants and log factors in the above bound).

Another term for covering is “metric entropy.”

Recall once again that we drop the normalization $1/n$ from URad and the choice of norm when covering.

18  Two deep network covering number bounds

We will give two generalization bounds.

- The first will be for arbitrary Lipschitz functions, and will be horifically loose (exponential in dimension).
- The second will be, afaik, the tightest known bound for ReLU networks.

18.1 First covering number bound: Lipschitz functions

This bound is intended as a point of contrast with our deep network generalization bounds.

Theorem 18.1  Let data $S = (x_1, \ldots, x_n)$ be given with $R := \max_{i,j} \|x_i - x_j\|_\infty$. Let $\mathcal{F}$ denote all $\rho$-Lipschitz functions from $[-R + R]^{d} \rightarrow [-B, +B]$ (where Lipschitz is measured wrt $\| \cdot \|_\infty$). Then the improper covering number $\mathcal{N}$ satisfies

$$\ln \mathcal{N}(\mathcal{F}, \epsilon : \| \cdot \|_u) \leq \max \left\{ 0, \left\lceil \frac{4\rho(R + \epsilon)}{\epsilon} \right\rceil^d \ln \left\lfloor \frac{2B}{\epsilon} \right\rfloor \right\}.$$

Remark 18.1  Exponential in dimension!

Revisiting the “point of contrast” comment above, our deep network generalization bounds are polynomial and not exponential in dimension; consequently, we really are doing much better than simply treating the networks as arbitrary Lipschitz functions.
Proof.
- Suppose $B > \epsilon$, otherwise can use the trivial cover $\{x \mapsto 0\}$.
- Subdivide $[-R - \epsilon, +R + \epsilon]^d$ into $\left(\frac{4(R + \epsilon)\rho}{\epsilon}\right)^d$ cubes of side length $\frac{2\rho}{\epsilon}$; call this $U$.
- Subdivide $[-B, +B]$ into intervals of length $\epsilon$, thus $2B/\epsilon$ elements; call this $V$.
- Our candidate cover $\mathcal{G}$ is the set of all piecewise constant maps from $[-R - \epsilon, +R + \epsilon]^d$ to $[-B, +B]$ discretized according to $U$ and $V$, meaning

$$|\mathcal{G}| \leq \left[\frac{2B}{\epsilon}\right] \left[\frac{4(R + \epsilon)\rho}{\epsilon}\right]^d.$$

To show this is an improper cover, given $f \in \mathcal{F}$, choose $g \in \mathcal{G}$ by proceeding over each $C \in U$, and assigning $g|_C \in V$ to be the closest element to $f(x_C)$, where $x_C$ is the midpoint of $C$. Then

$$\|f - g\|_u = \sup_{C \in U} \sup_{x \in C} |f(x) - g(x)|$$

$$\leq \sup_{C \in U} \sup_{x \in C} (|f(x) - f(x_C)| + |f(x_C) - g(x)|)$$

$$\leq \sup_{C \in U} \sup_{x \in C} \left(\rho \|x - x_C\|_\infty + \frac{\epsilon}{2}\right)$$

$$\leq \sup_{C \in U} \sup_{x \in C} \left(\rho(\epsilon/(4\rho)) + \frac{\epsilon}{2}\right) \leq \epsilon$$

[ mjt*: hmm the proof used uniform norm... is it defined?]

### 18.2 “Spectrally-normalized” covering number bound

**Theorem 18.2 (P. Bartlett, Foster, and Telgarsky (2017))** Fix multivariate activations $(\sigma_i)_{i=1}^L$ with $\|\sigma\|_{\text{Lip}} =: \rho_i$ and $\sigma_i(0) = 0$, and data $X \in \mathbb{R}^{n \times d}$, and define

$$\mathcal{F}_n := \left\{ \sigma_L(W_L \sigma_{L-1} \cdots \sigma_1(W_1X^T) \cdots) : \|W_i\|_2 \leq s_i, \|W_i^T\|_{2,1} \leq b_i \right\},$$

and all matrix dimensions are at most $m$. Then

$$\ln \mathcal{N}(\mathcal{F}_n, \epsilon, \|\cdot\|_F) \leq \frac{\|X\|_F^2}{\epsilon^2} \prod_{j=1}^L \rho_j^2 s_j^2 \left( \sum_{i=1}^L \left( \frac{b_i}{s_i} \right)^{2/3} \right)^3 \ln(2m^2).$$

**Remark 18.2** Applying Dudley,

$$\text{URad}(\mathcal{F}_n) = \tilde{O} \left( \frac{\|X\|_F^2}{\epsilon^2} \prod_{j=1}^L \rho_j s_j \right) \cdot \left[ \sum_{i=1}^L \left( \frac{b_i}{s_i} \right)^{2/3} \right]^{3/2}.$$

[ mjt*: that’s annoying and should be included/performe rigorously.]

Proof uses $\|\sigma(M) - \sigma(M')\|_F \leq \|\sigma\|_{\text{Lip}} \cdot \|M - M'\|_F$; in particular, it allows multi-variate gates like max-pooling! See (P. Bartlett, Foster, and Telgarsky 2017) for $\|\sigma_i\|_{\text{Lip}}$ estimates.

This proof can be adjusted to handle “distance to initialization”; see (P. Bartlett, Foster, and Telgarsky 2017) and the notion “reference matrices.”
Let’s compare to our best “layer peeling” proof from before, which had $\prod_i \|W_i\|_F \lesssim m^{L/2} \prod_i \|W_i\|_2$. That proof assumed $\rho_i = 1$, so the comparison boils down to

$$m^{L/2} \left( \prod_i \|W_i\|_2 \right) \quad \text{vs.} \quad \left[ \sum_i \left( \frac{\|W_i^T\|_{2,1}^{2/3}}{\|W_i\|_2^{2/3}} \right)^{3/2} \right] \left( \prod_i \|W_i\|_2 \right),$$

where $L \leq \sum_i \left( \frac{\|W_i^T\|_{2,1}^{2/3}}{\|W_i\|_2^{2/3}} \right) \leq Lm^{2/3}$. So the bound is better but still leaves a lot to be desired, and is loose in practice.

It is not clear how to prove exactly this bound with Rademacher peeling, which is a little eerie (independent of whether this bound is good or not).

The proof, as with Rademacher peeling proofs, is an induction on layers, similarly one which does not “coordinate” the behavior of the layers; this is one source of looseness.

**Remark 18.3 (practical regularization schemes)** This bound suggests regularization based primarily on the Lipschitz constant of the network; similar ideas appeared in parallel applied work, both for classification problems (Cisse et al. 2017), and for GANs (Arjovsky, Chintala, and Bottou 2017).

**Remark 18.4 (another proof)** For an alternate proof a similar fact (albeit requiring univariate gates), see (Neyshabur, Bhojanapalli, and Srebro 2018).

The first step of the proof is a covering number for individual layers.

**Lemma 18.1**

$$\ln \mathcal{N}(\{WX^T : X \in \mathbb{R}^{m \times d}, \|W^T\|_{2,1} \leq b\}, \epsilon, \| \cdot \|_F) \leq \left[ \frac{\|X\|_F^2 b^2}{\epsilon^2} \right] \ln(2dm).$$
Proof. Let $W \in \mathbb{R}^{m \times d}$ be given with $\|W\|_{2,1} \leq r$. Define $s_{ij} := W_{ij}/|W_{ij}|$, and note

$$WX^T = \sum_{i,j} e_i e_j^T W_{ij} (Xe_j)^T = \sum_{i,j} e_i W_{ij} (Xe_j)^T = \sum_{i,j} \frac{|W_{ij}||Xe_j|_2}{r\|X\|_F} r\|X\|_F s_{ij} e_i (Xe_j)^T. $$

Note by Cauchy-Schwarz that

$$\sum_{i,j} q_{ij} \leq \frac{1}{r\|X\|_F} \sum_i \sqrt{\sum_j W_{ij}^2} \|X\|_F = \frac{\|WX^T\|_{2,1} \|X\|_F}{r\|X\|_F} \leq 1,$$

potentially with strict inequality, thus $q$ is not a probability vector, which we will want later. To remedy this, construct probability vector $p$ from $q$ by adding in, with equal weight, some $U_{ij}$ and its negation, so that the above summation form of $WX^T$ goes through equally with $p$ as with $q$.

Now define IID random variables $(V_1, \ldots, V_k)$, where

$$\Pr[V_l = U_{ij}] = p_{ij},$$

$$\mathbb{E} V_l = \sum_{i,j} p_{ij} U_{ij} = \sum_{i,j} q_{ij} U_{ij} = WX^T,$$

$$\|U_{ij}\| = \left(\frac{s_{ij} e_i (Xe_j)}{\|Xe_j\|_2} \right) \cdot r\|X\|_F = |s_{ij}| \cdot \|e_i\|_2 \cdot \frac{Xe_j}{\|Xe_j\|_2} \cdot r\|X\|_F = r\|X\|_F,$$

$$\mathbb{E} \|V_l\|^2 = \sum_{i,j} p_{ij} \|U_{ij}\|^2 \leq \sum_{i,j} p_{ij} r^2 \|X\|_F^2 = r^2 \|X\|_F^2.$$

By $\text{(lemma:maurey?)},$ there exist $(\hat{V}_1, \ldots, \hat{V}_k) \in S^k$ with

$$\left\|WX^T - \frac{1}{k} \sum_l \hat{V}_l\right\|^2 \leq \mathbb{E} \left\|V_l - \frac{1}{k} \sum_l V_l\right\|^2 \leq \frac{1}{k} \mathbb{E} \|V_l\|^2 \leq \frac{r^2 \|X\|_F^2}{k}.$$

Furthermore, the matrices $\hat{V}_l$ have the form

$$\frac{1}{k} \sum_l \hat{V}_l = \frac{1}{k} \sum_l \frac{s_{ij} e_i (Xe_j)^T}{\|Xe_j\|_2} = \left[\frac{1}{k} \sum_l \frac{s_{ij} e_i^T}{\|Xe_j\|_2}\right] X^T;$$

by this form, there are at most $(2nd)^k$ choices for $(\hat{V}_1, \ldots, \hat{V}_k)$.

**Lemma 18.2** Let $F_n$ be the same image vectors as in the theorem, and let per-layer tolerances $(\epsilon_1, \ldots, \epsilon_L)$ be given. then

$$\ln \mathcal{N}\left(F_n, \sum_{j=1}^L \rho_j \epsilon_j \prod_{k=j+1}^L \rho_k s_k, \|\cdot\|_F\right) \leq \sum_{i=1}^L \left[\frac{\|X\|_F^2 b^2_i \prod_{j<i} \rho_j^2 s^2_j}{\epsilon^2_i}\right] \ln(2m^2).$$
Proof. Let $X_i$ denote the output of layer $i$ of the network, using weights $(W_i, \ldots, W_1)$, meaning

$$X_0 := X \quad \text{and} \quad X_i := \sigma_i(X_{i-1}W_i^T).$$

The proof recursively constructs cover elements $\hat{X}_i$ and weights $\hat{W}_i$ for each layer with the following basic properties.

- Define $\hat{X}_0 := X_0$, and $\hat{X}_i := B_i\sigma_i(\hat{X}_{i-1}\hat{W}_i^T)$, where $B_i$ is the Frobenius-norm ball of radius $\|X\|_F \prod_{j<i} \rho_j s_j$.
- Due to the projection $B_i$, $\|\hat{X}_i\|_F \leq \|X\|_F \prod_{j<i} \rho_j s_j$. Similarly, using $\rho_i(0) = 0$, $\|X_i\|_F \leq \|X\|_F \prod_{j<i} \rho_j s_j$.
- Given $\hat{X}_{i-1}$, choose $\hat{W}_i$ via +Lemma 18.1 so that $\|\hat{X}_{i-1}W_i^T - \hat{X}_{i-1}\hat{W}_i^T\|_F \leq \epsilon_i$, whereby the corresponding covering number $N_i$ for this layer satisfies

$$\ln N_i \leq \left\lfloor \frac{\|\hat{X}_{i-1}\|_F^2 b_i^2}{\epsilon_i^2} \right\rfloor \ln(2m^2) \leq \left\lfloor \frac{\|X\|_F^2 b_i^2 \prod_{j<i} \rho_j^2 s_j^2}{\epsilon_i^2} \right\rfloor \ln(2m^2).$$

- Since each cover element $\hat{X}_i$ depends on the full tuple $(\hat{W}_i, \ldots, \hat{W}_1)$, the final cover is the product of the individual covers (not their union), and the final cover log cardinality is upper bounded by

$$\ln \prod_{i=1}^L N_i \leq \sum_{i=1}^L \left\lfloor \frac{\|X\|_F^2 b_i^2 \prod_{j<i} \rho_j^2 s_j^2}{\epsilon_i^2} \right\rfloor \ln(2m^2).$$

It remains to prove, by induction, an error guarantee

$$\|X_i - \hat{X}_i\|_F \leq \sum_{j=1}^i \rho_j \epsilon_j \prod_{k=j+1}^i \rho_k s_k.$$ 

The base case $\|X_0 - \hat{X}_0\|_F = \epsilon_0$ holds directly. For the inductive step, by the above ingredients and the triangle inequality,

$$\|X_i - \hat{X}_i\|_F \leq \rho_i \|X_{i-1}W_i^T - \hat{X}_{i-1}\hat{W}_i^T\|_F \leq \rho_i \|X_{i-1}W_i^T - \hat{X}_{i-1}W_i^T\|_F + \rho_i \|\hat{X}_{i-1}W_i^T - \hat{X}_{i-1}\hat{W}_i^T\|_F \leq \rho_i \epsilon_i + \rho_i \epsilon_i$$

$$\leq \rho_i \epsilon_i \left[ \sum_{j=1}^{i-1} \rho_j \epsilon_j \prod_{k=j+1}^{i-1} \rho_k s_k \right] + \rho_i \epsilon_i$$

$$= \sum_{j=1}^i \rho_j \epsilon_j \prod_{k=j+1}^i \rho_k s_k + \rho_i \epsilon_i$$

$$= \sum_{j=1}^i \rho_j \epsilon_j \prod_{k=j+1}^i \rho_k s_k.$$
Proof of Theorem 18.2 (P. Bartlett, Foster, and Telgarsky (2017)). By solving a Lagrangian (minimize cover size subject to total error $\leq \epsilon$), choose

$$\epsilon_i := \frac{\alpha_i \epsilon}{\rho_i \prod_{j>i} \rho_j s_j}, \quad \alpha_i := \frac{1}{\beta} \left( \frac{b_i}{s_i} \right)^{2/3}, \quad \beta := \sum_{i=1}^{L} \left( \frac{b_i}{s_i} \right)^{2/3}.$$ 

Invoking the induction lemma with these choices, the resulting cover error is

$$\sum_{i=1}^{L} \epsilon_i \rho_i \prod_{j>i} \rho_j s_j = \epsilon \sum_{j=1}^{L} \alpha_i = \epsilon.$$ 

and the main term of the cardinality (ignoring $\ln(2m^2)$) satisfies

$$\sum_{i=1}^{L} \frac{\|X\|^2 F}{\epsilon^2} \frac{b_i^2 \prod_{j<i} \rho_j^2 s_j^2}{\alpha_i^2 s_i^2} = \frac{\|X\|^2 F}{\epsilon^2} \sum_{i=1}^{L} \frac{b_i^2 \prod_{j=1}^{L} \rho_j^2 s_j^2}{\alpha_i^2 s_i^2} = \frac{\|X\|^2 F}{\epsilon^2} \sum_{i=1}^{L} \frac{\beta^2 b_i^{4/3}}{s_i^{2/3}} = \frac{\|X\|^2 F}{\epsilon^2} \left( \sum_{i=1}^{L} \left( \frac{b_i}{s_i} \right)^{2/3} \right)^3.$$ 

[mjt$: I should include the Lagrangian explicitly and also explicit Dudley.]

19 VC dimension

[mjt$: ok if VC dim is one section, covering numbers should be as well?]
[mjt$: remainder is copy/pasted from fall2018, was not taught in fall2019.]
[mjt$: should include in preamble various bounds not taught, and a comment that VC dim proofs are interesting and reveal structure not captured above.]

• VC dimension is an ancient generalization technique; essentially the quantity itself appears in the work of Kolmogorov, and was later rediscovered a few times, and named after Vapnik and Chervonenkis, whose used it for generalization.

• To prove generalization, we will upper bound Rademacher complexity with VC dimension; classical VC dimension generalization proofs include Rademacher averages.

• There is some huge ongoing battle of whether VC dimension is a good measure or not. I think the proofs are interesting and are sensitive to interesting properties of deep networks in ways not capture by many of the bounds we spent time on. Anyway, a discussion for another time...

  – As stated the bounds are worst-case-y; I feel they could be adapted into more average-case-y bounds, though this has not been done yet...

First, some definitions. First, the zero-one/classification risk/error:

$$R_{\alpha}(\text{sgn}(f)) = \Pr[\text{sgn}(f(X)) \neq Y], \quad \tilde{R}_{\alpha}(\text{sgn}(f)) = \frac{1}{n} \sum_{i=1}^{n} 1[\text{sgn}(f(x_i)) \neq y_i].$$
The earlier Rademacher bound will now have

$$\text{URad}\left(\{(x, y) \mapsto 1[\text{sgn}(f(x)) \neq y] : f \in \mathcal{F}\}_{|S|}\right).$$

This is at most $2^n$; we’ll reduce it to a combinatorial quantity:

$$\text{sgn}(U) := \{(\text{sgn}(u_1), \ldots, \text{sgn}(u_n)) : u \in V\},$$

$$\text{Sh}(\mathcal{F}|S) := \left|\text{sgn}(\mathcal{F}|S)\right|,$$

$$\text{Sh}(\mathcal{F}; n) := \sup_{S \in \mathcal{S}} \left|\text{sgn}(\mathcal{F}|S)\right|,$$

$$\text{VC}(\mathcal{F}) := \sup\left\{i \in \mathbb{Z}_{\geq 0} : \text{Sh}(\mathcal{F}; i) = 2^i\right\}.$$

**Remark 19.1**  Sh is “shatter coefficient,” VC is “VC dimension.”

Both quantities are criticized as being too tied to their worst case; bounds here depend on (empirical quantity!) $\text{URad}(\text{sgn}(\mathcal{F}|S))$, which can be better, but throws out the labels.

**Theorem 19.1** (“VC Theorem”) With probability at least $1 - \delta$, every $f \in \mathcal{F}$ satisfies

$$\mathcal{R}_z(\text{sgn}(f)) \leq \tilde{\mathcal{R}}_z(\text{sgn}(f)) + \frac{2}{n} \text{URad}(\text{sgn}(\mathcal{F}|S)) + 3\sqrt{\frac{\ln(2/\delta)}{2n}},$$

and

$$\text{URad}(\text{sgn}(\mathcal{F}|S)) \leq \sqrt{2n \ln \text{Sh}(\mathcal{F}|S)},$$

$$\ln \text{Sh}(\mathcal{F}|S) \leq \ln \text{Sh}(\mathcal{F}; n) \leq \text{VC}(\mathcal{F}) \ln(n + 1).$$

**Remark 19.2**  [ mjt: Say something like “Need Sh($\mathcal{F}|_s$) = o(n)” in order to learn” ?]

Minimizing $\tilde{\mathcal{R}}_z$ is NP-hard in many trivial cases, but those require noise and neural networks can often get $\tilde{\mathcal{R}}_z(\text{sgn}(f)) = 0$.

$\text{VC}(\mathcal{F}) < \infty$ suffices; many considered this a conceptual breakthrough, namely “learning is possible!”

The quantities (VC, Sh) appeared in prior work (not by V-C). Symmetrization apparently too, though I haven’t dug this up.

First step of proof: pull out the zero-one loss.

**Lemma 19.1**  $\text{URad}(\{(x, y) \mapsto 1[\text{sgn}(f(x)) \neq y] : f \in \mathcal{F}\}_{|S|}) \leq \text{URad}(\text{sgn}(\mathcal{F}|S)).$
**Proof.** For each $i$, define

$$\ell_i(z) := \max \left\{ 0, \min \left\{ 1, \frac{1 - y_i(2z - 1)}{2} \right\} \right\},$$

which is 1-Lipschitz, and satisfies

$$\ell_i(\text{sgn}(f(x_i))) = 1[\text{sgn}(f(x_i)) \neq y_i].$$

(Indeed, it is the linear interpolation.) Then

$$\text{URad}(\{(x, y) \mapsto 1[\text{sgn}(f(x)) \neq y] : f \in F\} |_S)$$

$$= \text{URad}(\{\ell_1(\text{sgn}(f(x_1))), \ldots, \ell_n(\text{sgn}(f(x_n))) : f \in F\} |_S)$$

$$= \text{URad}(\ell \circ \text{sgn}(F) |_S)$$

$$\leq \text{URad}(\text{sgn}(F) |_S).$$

[ mjtΩ: is that using the fancier per-coordinate vector-wise peeling again?] 

Plugging this into our Rademacher bound: w/ $\Pr \geq 1 - \delta$, $\forall f \in F$,

$$R_z(\text{sgn}(f)) \leq \hat{R}_z(\text{sgn}(f)) + \frac{2}{n} \text{URad}(\text{sgn}(F) |_S) + 3 \sqrt{\frac{\ln(2/\delta)}{2n}}.$$

The next step is to apply Massart’s finite lemma, giving

$$\text{URad}(\text{sgn}(F) |_S) \leq \sqrt{2n \text{Sh}(F |_S)}.$$

One last lemma remains for the proof.

[ mjtΩ: lol why mention warren. should be explicit and not passive-aggressive.]

**Lemma 19.2 (Sauer-Shelah? Vapnik-Chervonenkis? Warren? ...)** Let $F$ be given, and define $V := \text{VC}(F)$. Then

$$\text{Sh}(F; n) \leq \begin{cases} 2^n & \text{when } n \leq V, \\ \left(\frac{e^n}{V}\right)^V & \text{otherwise}. \end{cases}$$

Moreover, $\text{Sh}(F; n) \leq n^V + 1$.

**(Proof.** Omitted. Exists in many standard texts.)

[ mjtΩ: okay fine but i should give a reference, and eventually my own clean proof.]

**19.1 VC dimension of linear predictors**

**Theorem 19.2** Define $F := \{x \mapsto \text{sgn}(\langle a, x \rangle - b) : a \in \mathbb{R}^d, b \in \mathbb{R}\}$ (“linear classifiers”/“affine classifier”/ “linear threshold function (LTF)”). Then $\text{VC}(F) = d + 1.$
Remark 19.3  By Sauer-Shelah, $\text{Sh}(\mathcal{F}; n) \leq n^{d+1} + 1$. Anthony-Bartlett chapter 3 gives an exact equality; only changes constants of $\ln \text{VC}(\mathcal{F}; n)$.

Let’s compare to Rademacher:

$$\text{URad}(\text{sgn}(\mathcal{F}|_S)) \leq \sqrt{2nd \ln(n+1)},$$

$$\text{URad}(\{x \mapsto \langle w, x \rangle : \|w\| \leq R\}|_S) \leq R\|X_S\|_F,$$

where $\|X_S\|_F^2 = \sum_{x \in S} \|x\|_2^2 \leq n \cdot d \cdot \max_i, j \ x_{i,j}$. One is scale-sensitive (and suggests regularization schemes), other is scale-insensitive.

**Proof.** First let’s do the lower bound $\text{VC}(\mathcal{F}) \geq d + 1$.

- Suffices to show $\exists S := \{x_1, \ldots, x_{d+1}\}$ with $\text{Sh}(\mathcal{F}|_S) = 2^{d+1}$.
- Choose $S := \{e_1, \ldots, e_d, (0, \ldots, 0)\}$.

Given any $P \subseteq S$, define $(a, b)$ as

$$a_i := 2 \cdot 1[e_i \in P] - 1, \quad b := \frac{1}{2} - 1[0 \in P].$$

Then

$$\text{sgn}((a, e_i) - b) = \text{sgn}(21[e_i \in P] - 1 - b) = 21[e_i \in P] - 1,$$

$$\text{sgn}((a, 0) - b) = \text{sgn}(21[0 \in P] - 1/2) = 21[0 \in P] - 1,$$

meaning this affine classifier labels $S$ according to $P$, which was an arbitary subset.

Now let’s do the upper bound $\text{VC}(\mathcal{F}) < d + 2$.

- Consider any $S \subseteq \mathbb{R}^d$ with $|S| = d + 2$.
- By Radon’s Lemma (proved next), there exists a partition of $S$ into nonempty $(P, N)$ with $\text{conv}(P) \cap \text{conv}(N) \neq \emptyset$.
- Label $P$ as positive and $N$ as negative. Given any affine classifier, it can not be correct on all of $S$ (and thus $\text{VC}(\mathcal{F}) < d + 2$): either it is incorrect on some of $P$, or else it is correct on $P$, and thus has a piece of $\text{conv}(N)$ and thus $x \in N$ labeled positive.

[ mjt@: needs ref]

**Lemma 19.3 (Radon’s Lemma)**  Given $S \subseteq \mathbb{R}^d$ with $|S| = d + 2$, there exists a partition of $S$ into nonempty $(P, N)$ with $\text{conv}(P) \cap \text{conv}(S) \neq \emptyset$. 

85
Proof. Let $S = \{x_1, \ldots, x_{d+2}\}$ be given, and define $\{u_1, \ldots, u_{d+1}\}$ as $u_i := x_i - x_{d+2}$, which must be linearly dependent:

- Exist scalars $(\alpha_1, \ldots, \alpha_{d+1})$ and a $j$ with $\alpha_j := -1$ so that
  $$\sum_i \alpha_i u_i = -u_j + \sum_{i \neq j} \alpha_i u_i = 0;$$

- thus $x_j - x_{d+2} = \sum_{i \neq j} \alpha_i (x_i - x_{d+2})$ and $0 = \sum_{i<d+2} \alpha_i x_i - x_{d+2} \sum_{i<d+2} \alpha_i =: \sum_j \beta_j x_j$,

  where $\sum_j \beta_j = 0$ and not all $\beta_j$ are zero.

Set $P := \{i : \beta_i > 0\}$, $N := \{i : \beta_i \leq 0\}$; where neither set is empty.

Set $\beta := \sum_{i \in P} \beta_i - \sum_{i \in N} \beta_i > 0$.

Since $0 = \sum_i \beta_i x_i = \sum_{i \in P} \beta_i x_i + \sum_{i \in N} \beta_i x_i$, then

$$0 \beta = \sum_{i \in P} \beta_i x_i + \sum_{i \in N} \beta_i x_i$$

and the point $z := \sum_{i \in P} \beta_i x_i / \beta = \sum_{i \in N} \beta_i x_i / (-\beta)$ satisfies $z \in \text{conv}(P) \cap \text{conv}(N)$.

Remark 19.4 Generalizes Minsky-Papert “xor” construction.

Indeed, the first appearance I know of shattering/VC was in approximation theory, the papers of Warren and Shapiro, and perhaps it is somewhere in Kolmogorov’s old papers.

19.2 VC dimension of threshold networks

Consider iterating the previous construction, giving an “LTF network”: a neural network with activation $z \mapsto \mathbf{1}[z \geq 0]$.

We’ll analyze this by studying output of all nodes. To analyze this, we’ll study not just the outputs, but the behavior of all nodes.

| mjt@: another suggestion of definition in pandoc-numbering |

Definition.

- Given a sample $S$ of size $n$ and an LTF network with $m$ nodes (in any topologically sorted order), define activation matrix $A := \text{Act}(S; W := (a_1, \ldots, a_m))$ where $A_{ij}$ is the output of node $j$ on input $i$, with fixed network weights $W$.

- Let $\text{Act}(S; \mathcal{F})$ denote the set of activation matrices with architecture fixed and weights $W$ varying.

Remark 19.5 Since last column is the labeling, $|\text{Act}(S; \mathcal{F})| \geq \text{Sh}(\mathcal{F} | S)$.

Act seems a nice complexity measure, but it is hard to estimate given a single run of an algorithm (say, unlike a Lipschitz constant).

We’ll generalize Act to analyze ReLU networks.

Theorem 19.3 For any LTF architecture $\mathcal{F}$ with $p$ parameters,

$$\text{Sh}(\mathcal{F}; n) \leq |\text{Act}(S; \mathcal{F})| \leq (n + 1)^p.$$
Proof.

- Topologically sort nodes, let \((p_1, \ldots, p_m)\) denote numbers of respective numbers of parameters (thus \(\sum_i p_i = p\)).
- Proof will iteratively construct sets \((U_1, \ldots, U_m)\) where \(U_i\) partitions the weight space of nodes \(j \leq i\) so that, within each partition cell, the activation matrix does not vary.
- The proof will show, by induction, that \(|U_i| \leq (n+1)^{\sum_{j \leq i} p_j}\). This completes the proof of the first claim, since \(\text{Sh}(\mathcal{F}|S) \leq |\text{Act}(\mathcal{F}; S)| = |U_m|\).
- For convenience, define \(U_0 = \{\emptyset\}\), \(|U_0| = 1\); the base case is thus \(|U_0| = 1 = (n+1)^0\).

(Inductive step). Let \(j \geq 1\) be given; the proof will now construct \(U_{j+1}\) by refining the partition \(U_j\).

- Fix any cell \(C\) of \(U_j\); as these weights vary, the activation is fixed, thus the input to node \(j + 1\) is fixed for each \(x \in S\).
- Therefore, on this augmented set of \(n\) inputs (\(S\) with columns of activations appended to each example), there are \((n+1)^{p_{j+1}}\) possible outputs via Sauer-Shelah and the VC dimension of affine classifiers with \(p_{j+1}\) inputs.
- In other words, \(C\) can be refined into \((n+1)^{p_{j+1}}\) sets; since \(C\) was arbitrary,

\[
|U_{j+1}| = |U_j|(n+1)^{p_{j+1}} \leq (n+1)^{\sum_{l \leq j+1} p_l}.
\]

This completes the induction and establishes the Shattering number bound.

It remains to bound the VC dimension via this Shatter bound:

\[
\text{VC}(\mathcal{F}) < n
\]

\[
\iff \forall i \geq n \cdot \text{Sh}(\mathcal{F}; i) < 2^i
\]

\[
\iff \forall i \geq n \cdot (i+1)^p < 2^i
\]

\[
\iff \forall i \geq n \cdot p \ln(i+1) < i \ln 2
\]

\[
\iff \forall i \geq n \cdot p < \frac{i \ln(2)}{\ln(i+1)}
\]

\[
\iff p < \frac{n \ln(2)}{\ln(n+1)}
\]

If \(n = 6p \ln(p)\),

\[
\frac{n \ln(2)}{\ln(n+1)} \geq \frac{n \ln(2)}{\ln(2n)} = \frac{6p \ln(p) \ln(2)}{\ln 12 + \ln p + \ln \ln p}
\]

\[
\geq \frac{6p \ln p \ln 2}{3 \ln p} > p.
\]

Remark 19.6 Had to do handle \(\forall i \geq n\) since VC dimension is defined via sup; one can define funky \(\mathcal{F}\) where Sh is not monotonic in \(n\).

Lower bound is \(\Omega(p \ln m)\); see Anthony-Bartlett chapter 6 for a proof. This lower bound however is for a specific fixed architecture!

Other VC dimension bounds: ReLU networks have \(\tilde{O}(pL)\), sigmoid networks have \(\tilde{O}(p^2m^2)\), and there exists a convex-concave activation which is close to sigmoid but has VC dimension \(\infty\).

Matching lower bounds exist for ReLU, not for sigmoid; but even the “matching” lower bounds
are deceptive since they hold for a fixed architecture of a given number of parameters and layers.

### 19.3 VC dimension of ReLU networks

Today’s ReLU networks will predict with $x \mapsto A_L \sigma_{L-1} (A_{L-1} \cdots A_2 \sigma_1 (A_1 x + b_1) + b_2 \cdots + b_{L-1}) + b_L$, where $A_i \in \mathbb{R}^{d_i \times d_{i-1}}$ and $\sigma_i : \mathbb{R}^{d_i} \to \mathbb{R}^{d_i}$ applies the ReLU $z \mapsto \max\{0, z\}$ coordinate-wise.

Convenient notation: collect data as rows of matrix $X \in \mathbb{R}^{n \times d}$, and define

\[
X_0 := X^\top, \quad Z_0 := \text{all 1s matrix},
\]

\[
X_i := A_i (Z_{i-1} \odot X_{i-1}) + b_i 1_n^\top, \quad X_i := 1[X_i \geq 0],
\]

where $(Z_1, \ldots, Z_L)$ are the activation matrices.

[ mjJO: i should double check i have the tightest version? which is more sensitive to earlier layers? i should comment on that and the precise structure/meaning of the lower bounds?]

#### Theorem 19.4 ((Theorem 6, P. L. Bartlett et al. 2017))

Let fixed ReLU architecture $F$ be given with $p = \sum_{i=1}^L p_i$ parameters, $L$ layers, $m = \sum_{i=1}^L m_i$ nodes. Let examples $(x_1, \ldots, x_n)$ be given and collected into matrix $X$. There exists a partition $U_L$ of the parameter space satisfying:

- Fix any $C \in U_L$. As parameters vary across $C$, activations $(Z_1, \ldots, Z_L)$ are fixed.
- $\text{Sh}(F; n) \leq |\{Z_L(C) : C \in U_L\}| \leq |U_L| \leq (12nL)^{pL}$, where $Z_L(C)$ denotes the sign pattern in layer $L$ for $C \in U_L$.
- If $pL^2 \geq 72$, then $\text{VC}(F) \leq 6pL \ln(pL)$.

#### Remark 19.7 (on the proof)

As with LTF networks, the prove inductively constructs partitions of the weights up through layer $i$ so that the activations are fixed across all weights in each partition cell. Consider a fixed cell of the partition, whereby the activations are fixed zero-one matrices. As a function of the inputs, the ReLU network is now an affine function; as a function of the weights it is multilinear or rather a polynomial of degree $L$.

Consider again a fixed cell and some layer $i$; thus $\sigma(X_i) = Z_i \odot X_i$ is a matrix of polynomials of degree $i$ (in the weights). If we can upper bound the number of possible signs of $A_{i+1}(Z_i \odot X_i) + b_i 1_n^\top$, then we can refine our partition of weight space and recurse. For that we need a bound on sign patterns of polynomials, as on the next slide.

#### Theorem 19.5 (Warren ’68; see also Anthony-Bartlet Theorem 8.3)

Let $F$ denote functions $x \mapsto f(x; w)$ which are $r$-degree polynomials in $w \in \mathbb{R}^p$. If $n \geq p$, then $\text{Sh}(F; n) \leq 2 (2^{enr})^p$.

#### Remark 19.8

Proof is pretty intricate, and omitted. It relates the VC dimension of $F$ to the zero sets $Z_l := \{w \in \mathbb{R}^p : f(x; w) = 0\}$, which it controls with an application of Bezout’s Theorem. The zero-counting technique is also used to obtain an exact Shatter coefficient for affine classifiers.

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88
Proof (of ReLU VC bound).
We'll inductively construct partitions \((U_0, \ldots, U_L)\) where \(U_i\) partitions the parameters of layers \(j \leq i\) so that for any \(C \in U_i\), the activations \(Z_j\) in layer \(j \leq i\) are fixed for all parameter choices within \(C\) (thus let \(Z_j(C)\) denote these fixed activations).

The proof will proceed by induction, showing \(|U_i| \leq (12nL)^p_i\).

Base case \(i = 0\): then \(U_0 = \{\emptyset\}\), \(Z_0\) is all ones, and \(|U_0| = 1 \leq (12nL)^p_0\).

(Inductive step).

- Fix \(C \in S_i\) and \((Z_1, \ldots, Z_i) = (Z_1(C), \ldots, Z_i(C))\).
- Note \(X_{i+1} = A_{i+1}(Z_i \odot X_i) + b_i1_n^\top\) is polynomial (of degree \(i + 1\)) in the parameters since \((Z_1, \ldots, Z_i)\) are fixed.
- Therefore carving \(U_i\) into pieces according to \(Z_{i+1} = 1[X_{i+1} \geq 0]\) being fixed gives

  \[|\{1[X_{i+1} \geq 0] : \text{params} \in C\}| \leq Sh(i + 1 \text{ deg poly}; m_i \cdot n \text{ functions}) \leq 2 \left(2enm_{i+1} \sum_{j \leq i} p_j \right) \leq (12nL)^p_\ast\).

  [Technical comment: to apply the earlier shatter bound for polynomials, we needed \(n \cdot m_{i+1} \geq \sum_{j \leq i} p_j\); but if (even more simply) \(p \geq nm_{i+1}\), we can only have \(\leq 2^{nm_{i+1}} \leq 2^p\) activation matrices anyway, so the bound still holds.]  

- Therefore termcarving \(U_i\) into pieces according to \(Z_{i+1} = 1[X_{i+1} \geq 0]\) being fixed gives

  \[|U_{i+1}| \leq |U_i|(12nL)^p_i \leq (12nL)^p_{i+1}\).

This completes the induction and upper bounds the number of possible activation patterns and also the shatter coefficient.

It remains to upper bound the VC dimension via the Shattering bound. As with LTF networks,

\[
\text{VC}(F) < n \iff \forall i \geq n \cdot \text{Sh}(F; i) < 2^i
\]

\[
\iff \forall i \geq n \cdot (12iL)^p L < 2^i
\]

\[
\iff \forall i \geq n \cdot pL \ln(12iL) < i \ln 2
\]

\[
\iff \forall i \geq n \cdot pL < \frac{i \ln 2}{\ln(12iL)}
\]

\[
\iff pL < \frac{n \ln 2}{\ln(12nL)}
\]

If \(n = 6pL \ln(pL)\),

\[
\frac{n \ln 2}{\ln(12nL)} = \frac{6pL \ln(pL) \ln(2)}{\ln(72pL^2 \ln(pL))} = \frac{6pL \ln(pL) \ln(2)}{\ln(72) + \ln(pL^2) + \ln \ln(pL)}
\]

\[
\geq \frac{6pL \ln(pL) \ln(2)}{\ln(72) + \ln(pL^2) + \ln(pL) - 1} \geq \frac{6 \ln(pL) \ln(2)}{3 \ln(pL^2)}
\]

\[
= 2pL \ln 2 > pL.
\]

Remark 19.9 If ReLU is replaced with a degree \(r \geq 2\) piecewise polynomial activation, have \(r^i\)-degree polynomial in each cell of partition, and shatter coefficient upper bound scales with \(L^2\) not \(L\). The lower bound in this case still has \(L\) not \(L^2\); it’s not known where the looseness is.
Lower bounds are based on digit extraction, and for each pair \((p, L)\) require a fixed architecture.

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