Towards Understanding the Regularization of Adversarial Robustness on Neural Networks

Yuxin Wen * 1 2 Shuai Li * 1 Kui Jia 1 2

Abstract

The problem of adversarial examples has shown that modern Neural Network (NN) models could be rather fragile. Among the more established techniques to solve the problem, one is to require the model to be ϵ -adversarially robust (AR); that is, to require the model not to change predicted labels when any given input examples are perturbed within a certain range. However, it is observed that such methods would lead to standard performance degradation, i.e., the degradation on natural examples. In this work, we study the degradation through the regularization perspective. We identify quantities from generalization analysis of NNs; with the identified quantities we empirically find that AR is achieved by regularizing/biasing NNs towards less confident solutions by making the changes in the feature space (induced by changes in the instance space) of most layers smoother uniformly in all directions; so to a certain extent, it prevents sudden change in prediction w.r.t. perturbations. However, the end result of such smoothing concentrates samples around decision boundaries, resulting in less confident solutions, and leads to worse standard performance. Our studies suggest that one might consider ways that build AR into NNs in a gentler way to avoid the problematic regularization.

1. Introduction

Despite the remarkable performance (Krizhevsky et al., 2012) of Deep Neural Networks (NNs), they are found to be rather fragile and easily fooled by adversarial examples

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(Szegedy et al., 2014). More intriguingly, these adversarial examples are generated by adding imperceptible noise to normal examples, and thus are indistinguishable for humans. NNs that are more robust to adversarial examples tend to have lower standard accuracy (Su et al., 2018), i.e., the accuracy measured on natural examples. The trade-off between robustness and accuracy has been empirically observed in many works (Fawzi et al., 2018; Kurakin et al., 2017; Madry et al., 2018; Tsipras et al., 2019), and has been theoretically analyzed under the context of simple models, e.g., linear models (Tsipras et al., 2019), quadratic models (Fawzi et al., 2018), but it is not clear whether the analysis generalizes to NNs. For example, Tsipras et al. (2019) show that for linear models, if examples are close to decision boundaries, robustness provably conflicts with accuracy, though the proof seems unlikely to generalize to NNs. Arguably, the most widely used remedy is developed to require NNs to be ϵ adversarially robust (AR), e.g., via Adversarial Training (Madry et al., 2018), Lipschitz-Margin Training (Tsuzuku et al., 2018); that is, they require the model not to change predicted labels when any given input examples are perturbed within a certain range. In practice, such AR methods are found to lead to worse performance measured in standard classification accuracy. Alternatives to build AR into NNs are also being developed. For instance, Zhang et al. (2019) show that a gap exists between surrogate risk gap and 0-1 risk gap if many examples are close to decision boundaries, and better robustness can be achieved by pushing examples away from decision boundaries. But pushing examples away again degrades NN performance in their experiments. But they are yet to be widely adopted by the community.

We investigate how adversarial robustness built into NNs by the arguably most established method, i.e., Adversarial Training (Madry et al., 2018), influences the behaviors of NNs to make them more robust but have lower performance through the lens of regularization. In an earlier time (Szegedy et al., 2014), adversarial training has been suggested as a form of regularization: it augments the training of NNs with adversarial examples, and thus might improve the generalization of the end models. Note that such a *hard requirement* that the adversarial examples need to be classified correctly is different from the methods that increase

^{*}Equal contribution ¹School of Electronic and Information Engineering, South China University of Technology, Guangzhou, Guangdong 510640, China ²Pazhou Lab, Guangzhou, 510335, China. Correspondence to: Shuai Li lishuai918@gmail.com>, Yuxin Wen <wen.yuxin@mail.scut.edu.cn>, Kui Jia <kuijia@scut.edu.cn>.

adversarial robustness by adding a soft penalty term to the risk function employed by Lyu et al. (2015) and Miyato et al. (2018), or a penalty term through curvature reduction (Moosavi-Dezfooli et al., 2019), or local linearization (Qin et al., 2019) (more discussion in appendix A). In these works, regularization is explicitly enforced by a penalty term, while in adversarial training, it is not clear that how training with augmented adversarial examples regularizes NNs. For example, if adversarial training does work as a regularizer, how does a possible improvement in generalization by using more data end up degrading performance? Even such a basic problem does not have a clear answer. To understand the regularization effects of AR on NNs, we go beyond simple linear or quadratic models and undertake a comprehensive generalization analysis of AR by establishing a rigorous generalization bound on NNs, and carrying out a series of empirical studies theoretically guided by the bound.

Technically, improved generalization implies the reduction in gap between training errors and test errors. Regularization achieves the gap reduction by reducing the size of the hypothesis space, which reduces the variance, but meanwhile increases the bias of prediction made — a constant classifier can have zero generalization errors, but also have low test performance. Thus, when a hypothesis space is improperly reduced, another possible outcome is biased poorly performing models with reduced generalization gaps.

Key results. Through a series of theoretically motivated experiments, we find that AR is achieved by regularizing/biasing NNs towards less confident solutions by making the changes in the feature space of most layers (which are induced by changes in the instance space) smoother uniformly in all directions; so to a certain extent, it prevents sudden change in prediction w.r.t. perturbations. However, the end result of such smoothing concentrates examples around decision boundaries and leads to worse standard performance. We elaborate the above statement in details shortly in section 1.1.

Implications. We conjecture that the improper reduction comes from the indistinguishability of the change induced in the intermediate layers of NNs by adversarial noise and that by inter-class difference. To guarantee AR, NNs are asked to smoothe out difference uniformly in all directions in a high dimensional space, and thus are biased towards less confident solutions that make similar/concentrated predictions. We leave the investigation of the conjecture as future works.

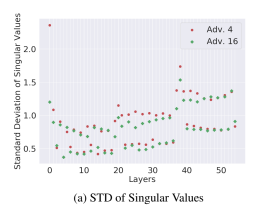
1.1. AR leads to less confident NNs with more indecisive misclassifications

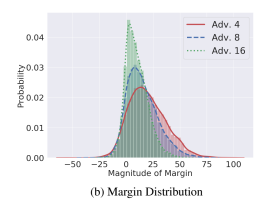
This section elaborates the *key results* we briefly present previously.

AR reduces the perturbations in the activation/outputs — the perturbations that are induced by perturbations in the inputs fed into the layer — of most layers. Through a series of theoretically motivated experiments, the results prompt us to look at the singular value distributions of the weight matrix of each layer of the NNs. Shown in fig. 1a, we find that overall the standard deviation (STD) of singular values associated with a layer of the NN trained with lower AR strength 4 is larger than that of the NN with higher AR strength 16 1 — the green dots are mostly below the red dots. Note that given a matrix W and an example x, singular values of W determine how the norm ||Wx|| is changed comparing with ||x||. More specifically, let $\sigma_{\min}, \sigma_{\max}$ be the minimal and maximal singular values. If x is not in the null space of W, then we have $||Wx|| \in [\sigma_{\min}||x||, \sigma_{\max}||x||]$, where $||\cdot||$ denotes 2-norm. This applies to norm $||\delta x||$ of a perturbation as well; that is, given possible changes δx of x of the same norm $||\delta x|| = c$, where c is a constant, the variance of $\sigma(\mathbf{W})$ roughly determines the variance of $||W\delta x||$, where $\sigma(W)$ denotes all singular values $\{\sigma_i\}$ of W. In more details, note that by SVD decomposition, $W\delta x = \sum_{i} \sigma_{i} u_{i} v_{i}^{T} \delta x$, thus σ_{i} determines how the component $v_{i}^{T} \delta x$ in the direction of v_{i} is amplified. To see an example, suppose that $\sigma_{\min} = \sigma_{\max} = \sigma_0$, then the variance of $\sigma(\mathbf{W})$ is zero, and $||\mathbf{W}\delta \mathbf{x}|| = \sigma_0 ||\delta \mathbf{x}||$. In this case, the variance of $||W\delta x||$ (given an ensemble of perturbations δx of the same norm c) is zero as well. The conclusion holds as well for ReLU($W\delta x$), where Where is a weight matrix of a layer of a NN, and ReLU denotes Rectifier Linear Unit activation function (proved by applying Cauchy interlacing law by row deletion (Chafai) in lemma 3.1). Consequently, by reducing the variance of singular values of weight matrix of a layer of the NN, AR reduces the variance of the norms of layer activations, or informally, perturbations in the activations, induced by input perturbations.

The perturbation reduction in activations concentrates examples, and it empirically concentrates them around decision boundaries; that is, predictions are less confident. The reduced variance implies that the outputs of each layer of the NN are more concentrated, but it does not tell where they are concentrated. Note that in the previous paragraph, the variance relationship discussed between $||\mathbf{W}\delta x||$ and $||\delta x||$ equally applies to $||\mathbf{W}x||$ and ||x||, where x is an actual example instead of perturbations. Thus, to find out the concentration of perturbations, we can look at the concentration of samples. Technically, we look at margins of examples. In a multi-class setting, suppose a NN computes a score function $f: \mathbb{R}^d \to \mathbb{R}^L$, where L is the number of classes; a way

 $^{^{1}}$ The AR strength is characterized by the maximally allowed l_{∞} norm of adversarial examples that are used to train the NNs — we use adversarial training (Madry et al., 2018) to build adversarial robustness into NNs. Details can be found in appendix B.1





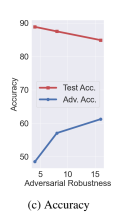


Figure 1. Experiment results on ResNet56 (He et al., 2016) trained on the CIFAR10 dataset. For the details of the experiments, refer to section 4. (a) The standard deviation of singular values of each layer of NNs with adversarial robustness (AR) strength 4, 16 (AR strength 8 is dropped for clarity of the plot). To emphasize, the x-axis is the layer index — overall 56 layers are involved. (b) The probability distribution of margins of NNs with AR strength 4, 8, 16. (c) The standard and adversarial accuracy of NNs with AR 4, 8, 16.

to convert this to a classifier is to select the output coordinate with the largest magnitude, meaning $x \mapsto \arg \max_i f_i(x)$. The confidence of such a classifier could be quantified by margins. It measures the gap between the output for the correct label and other labels, meaning $f_y(x) - \max_{i \neq y} f_i(x)$. Margin piece-wise linearly depends on the scores, thus the variance of margins is also in a piece-wise linear relationship with the variance of the scores, which are computed linearly from the activation of a NN layer. Thus, the consequence of concentration of activation discussed in the previous paragraph can be observed in the distribution of margins. More details of the connection between singular values and margins are discussed in section 4.2.2, after we present lemma 3.1. A zero margin implies that a classifier has equal propensity to classify an example to two classes, and the example is on the decision boundary. We plot the margin distribution of the test set of CIFAR10 in fig. 1b, and find that margins are increasingly concentrated around zero — that is, the decision boundaries — as AR strength grows.

The sample concentration around decision boundaries smoothes sudden changes induced perturbations, but also increases indecisive misclassifications. The concentration of test set margins implies that the induced change in margins by the perturbation in the instance space is reduced by AR. Given two examples x, x' from the test set, $\delta x = x - x'$ can be taken as a significant perturbation that changes the example x to x'. The concentration of overall margins implies the change induced by δx is smaller statistically in NNs with higher AR strength. Thus, for an adversarial perturbation applied on x, statistically the change of margins is smaller as well — experimentally it is reflected in the increased adversarial robustness of the network, as shown in the increasing curve in fig. 1c. That is, the sudden changes of margins originally induced by adversarial perturbations

are *smoothed* (to change slowly). However, the *cost* of such smoothness is lower confidence in prediction, and more test examples are slightly/indecisively moved to the wrong sides of the decision boundaries — incurring lower accuracy, as shown in the decreasing curve in fig. 1c.

Lastly, we note that experiments in this section are used to illustrate our main arguments in this section. Further consistent quality results are reported in section 4 by conducting experiments on CIFAR10/100 and Tiny-ImageNet with networks of varied capacity. And more corroborative experiment results are presented in the appendices, and outlined in section 1.2.

1.2. Outline and contributions

This work carries out generalization analysis on NNs with AR. The quantities in the previous section are identified by the generalization errors (GE) upper bound we establish at theorem 3.1, which characterizes the regularization of AR on NNs. The key result is obtained at the *end* of a series of analysis, thus we present the outline of the analysis here.

Outline. After presenting some preliminaries in section 2, we proceed to analyze the regularization of AR on NNs, and establish a GE upper bound in section 3. The bound prompts us to look at the GE gaps in experiments. In section 4.1, we find that for NNs trained with higher AR strength, the surrogate risk gaps (GE gaps) decrease for a range of datasets, i.e., CIFAR10/100 and Tiny-ImageNet. It implies AR effectively regularizes NNs. We then study the finer behavior change of NNs that might lead to such a gap reduction. Again, we follow the guidance of theorem 3.1. We look at the margins in section 4.2.1, then at the singular value distribution in section 4.2.2, and discover the main results described in section 1.1. More corroborative experiments are present

in appendix B.4 and appendix B.6 to show that such phenomenon exists in a broad range of NNs with varied capacity and adversarial training techniques. More complementary results are present in appendix B.3 to explain some seemingly abnormal observations, and in appendix B.5 to quantitatively demonstrate the smoothing effects of AR discussed in section 1.1. Related works are present in appendix A.

Contributions. Overall, the core contribution in this work is to show that adversarial robustness (AR) regularizes NNs in a way that hurts its capacity to learn to perform in test. More specifically:

- We establish a generalization error (GE) bound that characterizes the regularization of AR on NNs. The bound connects *margin* with adversarial robustness radius ϵ via *singular values of weight matrices* of NNs, thus suggesting the two quantities that guide us to investigate the regularization effects of AR empirically.
- Our empirical analysis tells that AR effectively regularizes NNs to reduce the GE gaps. To understand how reduced GE gaps turns out to degrade test performance, we study variance of singular values of layer-wise weight matrices of NNs and distributions of margins of samples, when different strength of AR are applied on NNs.
- The study shows that AR is achieved by regularizing/biasing NNs towards less confident solutions by making the changes in the feature space of most layers (which are induced by changes in the instance space) smoother uniformly in all directions; so to a certain extent, it prevents sudden change in prediction w.r.t. perturbations. However, the end result of such smoothing concentrates samples around decision boundaries and leads to worse standard performance.

2. Preliminaries

Assume an instance space $\mathcal{Z} = \mathcal{X} \times \mathcal{Y}$, where \mathcal{X} is the space of input data, and \mathcal{Y} is the label space. Z := (X,Y) are the random variables with an unknown distribution μ , from which we draw samples. We use $S_m = \{z_i = (\boldsymbol{x}_i, y_i)\}_{i=1}^m$ to denote the training set of size m whose examples are drawn independently and identically distributed (i.i.d.) by sampling Z. Given a loss function l, the goal of learning is to identify a function $T: \mathcal{X} \mapsto \mathcal{Y}$ in a hypothesis space (a class \mathcal{T} of functions) that minimizes the expected risk

$$R(l \circ T) = \mathbb{E}_{Z \sim \mu} [l (T(X), Y)],$$

Since μ is unknown, the observable quantity serving as the proxy to the expected risk R is the empirical risk

$$R_m(l \circ T) = \frac{1}{m} \sum_{i=1}^m l(T(\boldsymbol{x}_i), y_i).$$

Our goal is to study the discrepancy between R and R_m , which is termed as generalization error — it is also some-

times termed as generalization gap in the literature

$$GE(l \circ T) = |R(l \circ T) - R_m(l \circ T)|. \tag{1}$$

A NN is a map that takes an input x from the space \mathcal{X} , and builds its output by recursively applying a linear map W_i followed by a pointwise non-linearity g:

$$x_i = g(\boldsymbol{W}_i \boldsymbol{x}_{i-1}),$$

where i indexes the times of recursion, which is denoted as a layer in the community, $i=1,\ldots,L$, $x_0=x$, and g denotes the activation function. which is restricted to Rectifier Linear Unit (ReLU) (Glorot et al., 2011) or max pooling operator (Bécigneul, 2017) in this paper. To compactly summarize the operation of T, we denote

$$Tx = g(\mathbf{W}_L g(\mathbf{W}_{L-1} \dots g(\mathbf{W}_1 \mathbf{x}))). \tag{2}$$

Definition 1 (Covering number). Given a metric space (S, ρ) , and a subset $\tilde{S} \subset S$, we say that a subset \hat{S} of \tilde{S} is a ϵ -cover of \tilde{S} , if $\forall \tilde{s} \in \tilde{S}$, $\exists \hat{s} \in \hat{S}$ such that $\rho(\tilde{s}, \hat{s}) \leq \epsilon$. The ϵ -covering number of \tilde{S} is

$$\mathcal{N}_{\epsilon}(\tilde{\mathcal{S}}, \rho) = \min\{|\hat{\mathcal{S}}| : \hat{\mathcal{S}} \text{ is an } \epsilon\text{-covering of } \tilde{\mathcal{S}}\}.$$

Various notions of adversarial robustness have been studied in existing works (Madry et al., 2018; Tsipras et al., 2019; Zhang et al., 2019). They are conceptually similar; in this work, we formalize its definition to make clear the object for study.

Definition 2 $((\rho, \epsilon)$ -adversarial robustness). Given a multiclass classifier $f: \mathcal{X} \to \mathbb{R}^L$, and a metric ρ on \mathcal{X} , where L is the number of classes, f is said to be adversarially robust w.r.t. adversarial perturbation of strength ϵ , if there exists an $\epsilon > 0$ such that $\forall z = (x, y) \in \mathcal{Z}$ and $\delta x \in \{\rho(\delta x) \leq \epsilon\}$, we have

$$f_{\hat{y}}(\boldsymbol{x} + \delta \boldsymbol{x}) - f_i(\boldsymbol{x} + \delta \boldsymbol{x}) \ge 0,$$

where $\hat{y} = \arg\max_{j} f_{j}(\boldsymbol{x})$ and $i \neq \hat{y} \in \mathcal{Y}$. ϵ is called **adversarial robustness radius**. When the metric used is clear, we also refer (ρ, ϵ) -adversarial robustness as ϵ -adversarial robustness.

Note that the definition is an *example-wise* one; that is, it requires each example to have a guarding area, in which all examples are of the same class. Also note that the robustness is w.r.t. the predicted class, since ground-truth label is unknown for a \boldsymbol{x} in test.

We characterize the GE with ramp risk, which is a typical risk to undertake theoretical analysis (Bartlett et al., 2017; Neyshabur et al., 2018b).

Definition 3 (Margin Operator). A margin operator $\mathcal{M}: \mathbb{R}^L \times \{1,\dots,L\} \to \mathbb{R}$ is defined as

$$\mathcal{M}(\boldsymbol{s}, y) := s_y - \max_{i \neq y} s_i$$

Definition 4 (Ramp Loss). The ramp loss $l_{\gamma} : \mathbb{R} \to \mathbb{R}^+$ is defined as

$$l_{\gamma}(r) := \begin{cases} 0 & r < -\gamma \\ 1 + r/\gamma & r \in [-\gamma, 0] \\ 1 & r > 0 \end{cases}$$

Definition 5 (Ramp Risk). *Given a classifier* f, ramp risk is the risk defined as

$$R_{\gamma}(f) := \mathbb{E}(l_{\gamma}(-\mathcal{M}(f(X),Y))),$$

where X, Y are random variables in the instance space \mathcal{Z} previously.

We will use a different notion of margin in theorem 3.1, and formalize its definition as follows. We reserve the unqualified word "margin" specifically for the margin discussed previously — the output of margin operator for classification. We call this margin to-be-introduced *instance-space margin (IM)*.

Definition 6 (Smallest Instance-space Margin). Given an element $z = (x, y) \in \mathcal{Z}$, let v(x) be the distance from x to its closest point on the decision boundary, i.e., the instance-space margin (IM) of example x. Given a covering set \hat{S} of \mathcal{Z} , let

$$v_{\min} = \min_{\boldsymbol{x} \in \{\boldsymbol{x} \in \mathcal{X} | \exists \boldsymbol{x}' \in \hat{S}_m, ||\boldsymbol{x} - \boldsymbol{x}'||_2 \le \epsilon\}} v(\boldsymbol{x}), \quad (3)$$

where $\hat{S}_m := \{ \boldsymbol{x}' \in \hat{S} | \exists \boldsymbol{x}_i \in S_m, ||\boldsymbol{x}_i - \boldsymbol{x}'||_2 \leq \epsilon \}$. v_{\min} is the smallest instance-space margin of elements in the covering balls that contain training examples.

3. Theoretical instruments for empirical studies on AR

In this section, we rigorously establish the bound mentioned in the introduction. We study the map T defined in section 2 as a NN (though technically, T now is a map from \mathcal{X} to \mathbb{R}^L , instead of to \mathcal{Y} , such an abuse of notation should be clear in the context). To begin with, we introduce an assumption, before we state the generalization error bound guaranteed by adversarial robustness.

Assumption 3.1 (Monotony). Given a point $x \in \mathcal{X}$, let x' be the point on the decision boundary of a NN T that is closest to x. Then, for all x'' on the line segment x + t(x' - x), $t \in [0, 1]$, the margin $\mathcal{M}(Tx'', y)$ decreases monotonously.

The assumption is a regularity condition on the classifier that rules out undesired oscillation between x and x'. To see how, notice that the margin defined in definition 3 reflects how confident the decision is made. Since x' is on the decision boundary, it means the classifier is unsure how it

should be classified. Thus, when the difference x' - x is gradually added to x, ideally we want the confidence that we have on classifying x to decrease in a consistent way to reflect the uncertainty.

Theorem 3.1. Let T denote a NN with ReLU and MaxPooling nonlinear activation functions (the definition is put at eq. (2) for readers' convenience), l_{γ} the ramp loss defined at definition 4, and Z the instance space assumed in section 3. Assume that Z is a k-dimensional regular manifold that accepts an ϵ -covering with covering number $(\frac{C_{\chi}}{\epsilon})^k$, and assumption assumption 3.1 holds. If T is ϵ_0 -adversarially robust (defined at definition 2), $\epsilon \leq \epsilon_0$, and denote v_{\min} the smallest IM margin in the covering balls that contain training examples (defined at definition 6), σ_{\min}^i the smallest singular values of weight matrices W_i , i = 1, ..., L-1of a NN, $\{w_i\}_{i=1,...,|\mathcal{Y}|}$ the set of vectors made up with ith rows of \boldsymbol{W}_L (the last layer's weight matrix), then given an i.i.d. training sample $S_m = \{z_i = (\boldsymbol{x}_i, y_i)\}_{i=1}^m$ drawn from \mathcal{Z} , its generalization error $GE(l \circ T)$ (defined at eq. (1)) satisfies that, for any $\eta > 0$, with probability at least $1 - \eta$

$$GE(l_{\gamma} \circ T) \le \max\{0, 1 - \frac{u_{\min}}{\gamma}\} + \sqrt{\frac{2\log(2)C_{\mathcal{X}}^{k}}{\epsilon^{k}m} + \frac{2\log(1/\eta)}{m}}$$
(4)

where

$$u_{\min} = \min_{y,\hat{y} \in \mathcal{Y}, y \neq \hat{y}} ||\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}}||_2 \prod_{i=1}^{L-1} \sigma_{\min}^i v_{\min}$$
 (5)

is a lower bound of margins of examples in covering balls that contain training samples.

The proof of theorem 3.1 is in appendix C. The bound identifies quantities that would be studied experimentally in section 4 to understand the regularization of AR on NNs. The first term in eq. (4) in theorem 3.1 suggests that quantities related to the lower bound of margin u_{\min} might be useful to study how ϵ -adversarial robustness (ϵ -AR) regularizes NNs. However, ϵ -AR is guaranteed in the instance space that determines the smallest instance-space margin v_{\min} . To relate GE bound with ϵ -AR, we characterize in eq. (5) the relationship between margin with IM, via smallest singular values of NNs' weight matrices, suggesting that quantities related to singular values of NNs' weight matrices might be useful to study how AR regularizes NNs as well. An illustration on how AR could influence generalization of NNs through IM is also given in fig. 2a. The rightmost term in eq. (4) is a standard term in robust framework (Xu & Mannor, 2012) in learning theory, and is not very relevant to the discussion. The remaining of this paper are empirical studies that are based on the quantities, e.g., margin distributions and singular values of NNs' weight matrices, that are related to the identified quantities, i.e., u_{\min} , σ_{\min}^i .

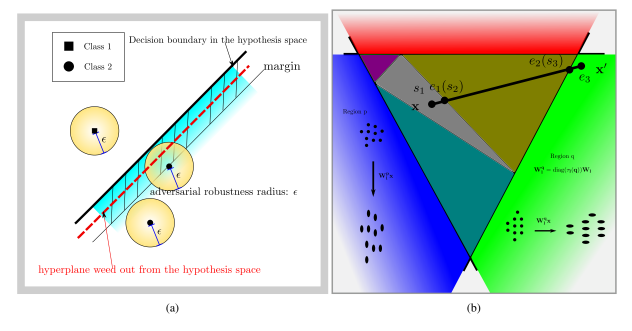


Figure 2. (a) Illustration of the regularization effect of adversarial robustness. If a NN T is ϵ -adversarially robust, for a given example \boldsymbol{x} (drawn as filled squares or circles) and points \boldsymbol{x}' in the yellow ball $\{\boldsymbol{x}' \mid \rho(\boldsymbol{x},\boldsymbol{x}') \leq \epsilon\}$ around \boldsymbol{x} , the predicted labels of $\boldsymbol{x},\boldsymbol{x}'$ should be the same, and the loss variation is potentially bigger as \boldsymbol{x}' moves from the center to the edge, as shown as intenser yellow color at the edge of a ball. Collectively, the adversarial robustness of each example requires an instance-space margin (IM) to exist for the decision boundary, shown as the shaded cyan margin. As normally known, margin is related to generalization ability that shrinks the hypothesis space. In this case, the IM required by adversarial robustness would weed out hypotheses that do not have an adequate IM, such as the red dashed line shown in the illustration. (b) Illustration of lemma 3.1. Given a NN with ReLU activation function, the feature map I_l at layer l is divided into regions where $I_l(\boldsymbol{x})$ is piecewise linear w.r.t. \boldsymbol{x} . The induced linear map \boldsymbol{W}_1^q is given by diag $(\tau_1(q))\boldsymbol{W}_1$, where diag $(\tau_l(q))$ is a diagonal matrix whose diagonal entries are given by a vector $\tau_1(q)$ that has 0-1 values. For example, in region p, $I_1 = \boldsymbol{W}_1^p \boldsymbol{x}$ and distance between instances \boldsymbol{x} are vertical elongated, while in region q, $I_1 = \boldsymbol{W}_1^q \boldsymbol{x}$ and distance are horizontally elongated. Thus given $\boldsymbol{x}, \boldsymbol{x}'$, the difference $||I_l(\boldsymbol{x}) - I_l(\boldsymbol{x}')||$ between $I_l(\boldsymbol{x})$ and $I_l(\boldsymbol{x}')$ is the length of the transformed line segment $\boldsymbol{x} - \boldsymbol{x}'$ drawn, of which each segment is linearly transformed in a different way.

These studies aim to illuminate with empirical evidence on the phenomena that AR regularizes NNs, reduces GE gaps, but degrades test performance. ²

Before turning into empirical study, we further present a lemma to illustrate the relation characterized in eq. (5) without the need to jump into proof of theorem 3.1. It would motivate our experiments later in section 4.2.2. We state the following lemma that relates distances between elements in the instance space with those in the feature space of any

intermediate network layers.

Lemma 3.1. Given two instances $x, x' \in \mathcal{X}$, let $I_l(x)$ be the activation $g(\mathbf{W}_l g(\mathbf{W}_{l-1} \ldots g(\mathbf{W}_1 x)))$ at layer l of x, then there exist $n \in \mathbb{N}$ sets of matrices $\{\mathbf{W}_i^{q_j}\}_{i=1...l}, j=1...n$, that each of the matrix $\mathbf{W}_i^{q_j}$ is obtained by setting some rows of \mathbf{W}_i to zero, and $\{q_j\}_{j=1...n}$ are arbitrary distinctive symbols indexed by j that index $\mathbf{W}_i^{q_j}$, such that

$$||oldsymbol{I}_l(oldsymbol{x})-oldsymbol{I}_l(oldsymbol{x}')|| = \sum_{j=1}^n \int_{s_j}^{e_j} \left| \left| \prod_{i=1}^l oldsymbol{W}_i^{q_j} dt (oldsymbol{x} - oldsymbol{x}')
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where $s_1 = 0$, $s_{j+1} = e_j$, $e_n = 1$, s_j , $e_j \in [0,1]$ — each $[s_j, e_j]$ is a segment in the line segment parameterized by t that connects x and x'.

Its proof is in appendix C, and an illustration is given in fig. 2b. Essentially, it states that difference in the feature space of a NN, induced by the difference between elements in the instance space, is a summation of the norms of the linear transformation $(\prod_{i=1}^{l} \boldsymbol{W}_{i}^{q_{j}})$ applied on segments of the line segment that connects $\boldsymbol{x}, \boldsymbol{x}'$ in the instance space.

²Note that in the previous paragraph, though we identifies quantities u_{\min} and σ_{\min}^i related to the upper bound of GE, the quantities we actually would study empirically are *margin distribution* and all *singular values* that characterize the GE of all samples, not just the extreme case (upper bound). The analytic characterization of the GE of all samples is not possible since we do not have enough information (we do not know the true distribution of samples). That's why to arrive at close-form analytic characterization of GE, we resort to the extreme non-asymptotic large-sample behaviors. *The analytic form is a neat way to present how relevant quantities influence GE.* In the rest of the paper, we would carry on empirical study on the distributions of margins and singular values to investigate AR's influence on GE of all samples.

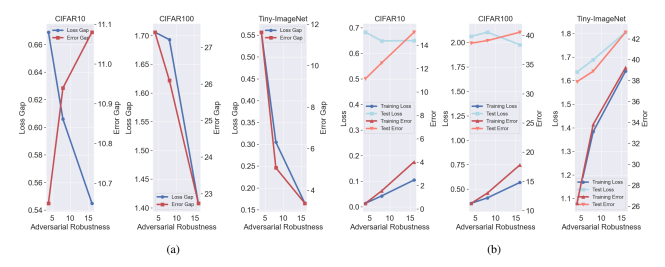


Figure 3. Experiment results on CIFAR10/100, and Tiny-ImageNet. The unit of x-axis is the adversarial robustness (AR) strength of NNs, c.f. the beginning of section 4. (a) Plots of loss gap (and error rate gap) between training and test datasets v.s. AR strength. (b) Plots of losses (and error rates) on training and test datasets v.s. AR strength.

Since $\boldsymbol{W}_i^{q_j}$ is obtained by setting rows of \boldsymbol{W}_i to zero, the singular values of these induced matrices are intimately related to weight matrices \boldsymbol{W}_i of NN by Cauchy interlacing law by row deletion (Chafai). Since the margin of an example \boldsymbol{x} is a linear transform of the difference between $I_{L-1}(\boldsymbol{x})$ and the $I_{L-1}(\boldsymbol{x}')$ of an element \boldsymbol{x}' on the decision boundary, singular values of $\{W_i\}_{i=1...L-1}$ determine the amplification/shrinkage of the IM $\boldsymbol{x}-\boldsymbol{x}'$.

4. Empirical studies on regularization of adversarial robustness

In this section, guided by theorem 3.1, we undertake empirical studies to explore AR's regularization effects on NNs. We first investigate the behaviors of off-the-shelf architectures of fixed capacity on various datasets in section 4.1 and section 4.2. More corroborative controlled studies that explore the regularization effects of AR on NNs with varied capacity are present in appendix B.3.

4.1. Adversarial robustness effectively regularizes NNs on various datasets

This section aims to explore whether AR can effectively reduce generalization errors — more specifically, the surrogate risk gaps. We use adversarial training (Madry et al., 2018) to build adversarial robustness into NNs. The AR strength is characterized by the maximally allowed l_{∞} norm of adversarial examples that are used to train the NNs. Details on the technique to build adversarial robustness into NNs is given in appendix B.1.

Our experiments are conducted on CIFAR10, CIFAR100, and Tiny-ImageNet (ImageNet, 2018) that represent learn-

ing tasks of increased difficulties. We use ResNet-56 and ResNet-110 (He et al., 2016) for CIFAR10/100, and Wide ResNet (WRN-50-2-bottleneck) (Zagoruyko & Komodakis, 2016) for Tiny-ImageNet (ImageNet, 2018). These networks are trained with increasing AR strength. Results are plotted in fig. 3.

Regularization of AR on NNs. We observe in fig. 3a (shown as blue lines marked by circles) that GE gaps (the gaps between training and test losses) decrease as strength of AR increase; we also observe in fig. 3a that training losses increase as AR strength increase; these results (and more results in subsequent fig. 6) imply that AR does regularize training of NNs by reducing their capacities to fit training samples. Interestingly, in the CIFAR 10/100 results in fig. 3b, the test losses show a decreasing trend even when test error rates increase. It suggests that the network actually performs better measured in test loss as contrast to the performance measured in test error rates. This phenomenon results from that less confident wrong predictions are made by NNs thanks to adversarial training, which will be explained in details in section 4.2, when we carry on finer analysis. We note that on Tiny-ImageNet, the test loss does not decrease as those on CIFAR10/100. It is likely because the task is considerably harder, and regularization hurts NNs even measured in test loss.

Trade-off between regularization of AR and test error rates. The error rate curves in fig. 3b also tell that the end result of AR regularization leads to biased-performing NNs that achieve degraded test performance. These results are consistent across datasets and networks.

Seemingly abnormal phenomenon. An seemingly abnormal phenomenon in CIFAR10 observed in fig. 3a is that the error rate gap actually increases. It results from the same underlying behaviors of NNs, which we would introduce in section 4.2, and an overfitting phenomenon that AR cannot control. Since it would be a digress to explain, it is put in appendix B.3.

We finally note that the adversarial robustness training reproduced is relevant, of which the defense effect is comparable with existing works. One may refer to fig. 12 in appendix D.2 for the details. We can see from it that similar adversarial robustness to Madry et al. (2018) and Li et al. (2018) is achieved for CIFAR10/100, Tiny-ImageNet in the NNs we reproduce.

4.2. Refined analysis through margins and singular values

The experiments in the previous sections confirm that AR reduces GE, but decreases accuracy. We study the underlying behaviors of NNs to analyze what have led to it here. More specifically, we show that adversarial training implements ϵ -adversarial robustness by making NNs biased towards less confident solutions; that is, the key finding we present in section 1.1 that explains both the prevented sudden change in prediction w.r.t. sample perturbation (i.e., the achieved AR), and the reduced test accuracy.

4.2.1. MARGINS THAT CONCENTRATE MORE AROUND ZERO LEAD TO REDUCED GE GAP

To study how GE gaps are reduced, theorem 3.1 suggests we first look at the margins of examples — a lower bound of margins is $u_{\rm min}$ in eq. (5). The analysis on margins has been a widely used tool in learning theory (Bartlett et al., 2017). It reflects the confidence that a classifier has on an example, which after being transformed by a loss function, is the surrogate loss. Thus, the loss difference between examples are intuitively reflected in the difference in confidence characterized by margins. To study how AR influences generalization of NNs, distributions of samples which are obtained by training ResNet-56 on CIFAR10 and CIFAR100 with increased AR strength (the same setting as for fig. 3). Applying the same network of ResNet-56 respectively on CIFAR-10 and CIFAR-100 of different learning difficulties creates learning settings of larger- and smaller-capacity NNs.

Concentration and reduced accuracy. In fig. 4, we can see that in both CIFAR10/100, the distributions of margins become more concentrated around zero as AR grows. The concentration moves the mode of margin distribution towards zero and more examples slightly across the decision boundaries, where the margins are zero, which explains the

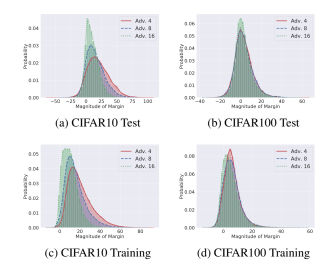


Figure 4. Margin distributions of NNs with AR strength 4, 8, 16 on Training and Test sets of CIFAR10/100.

reduced accuracy 3.

Concentration and reduced loss/GE gap. The concentration has different consequences on training and test losses. Before describing the consequences, to directly relate the concentration to loss gap, we further introduce estimated probabilities of examples. This is because though we use ramp loss in theoretical analysis, in the experiments, we explore the behaviors of more practically used cross entropy loss. The loss maps one-to-one to estimated probability, but not to margin, though they both serve as a measure of confidence. Suppose p(x) is the output of the softmax function of dimension L (L is the number of target classes), and y is the target label. The estimated probability of x would be the y-th dimension of (p(x)), i.e., $(p(x))_y$. On the training sets, since the NNs are optimized to perform well on the sets, only a tiny fraction of them are classified wrongly. To concentrate the margin distribution more around zero, is to

³We remark a possibly confusing phenomenon here about the margin. The bound eq. (4) might give the impression that a smaller margin might lead to a larger generalization error, while the empirical study instead shows that the NNs with a smaller margin have a smaller generalization error. The hypothesized confusion is a misunderstanding of the generalization bound analysis. The upper bound is a worst case analysis of GE. However, in practice, the interesting object is the average gap between the training losses and the test losses, i.e., the GE. Unfortunately, the average gap cannot be analyzed analytically (cf. footnote 2). Thus, we, and also the statistical learning community, resort to worst case analysis to find an upper bound on GE to identify quantities that might influence GE. In this case, the phenomenon suggests that the bound might be loose, though this is a problem that plagues the statistical learning community (Nagarajan & Kolter, 2019). But our focus in this work is not to derive tight bounds, or reach definite conclusions from bounds alone, but to guide experiments with the bound.

make almost all of predictions that are correct less confident. Thus, a higher expected training loss ensues. On the test sets, the estimated probabilities of the target class concentrate more around middle values, resulting from lower confidence/margins in predictions made by NNs, as shown in fig. 5a (but the majority of values are still at the ends). Note that wrong predictions away from decision boundaries (with large negative margins) map to large loss values in the surrogate loss function. Thus, though NNs with larger AR strength have lower accuracy, they give more predictions whose estimated probabilities are at the middle (compared with NNs with smaller AR strength). These predictions, even if relatively more of them are wrong, maps to smaller loss values, as shown in fig. 5b, where we plot the histogram of loss values of test samples. In the end, it results in expected test losses that are lower, or increase in a lower rate than the training losses on CIFAR10/100, Tiny-ImageNet, as shown in fig. 3b. The reduced GE gap results from the increased training losses, and decreased or less increased test losses.

4.2.2. AR MAKES NNS SMOOTHE PREDICTIONS W.R.T. INPUT PERTURABTIONS IN ALL DIRECTIONS

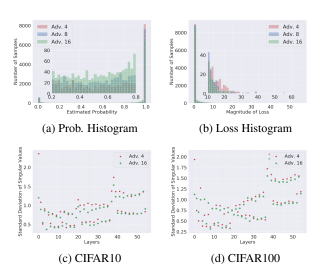


Figure 5. (a)(b) are histograms of estimated probabilities and losses of the test set sample of NNs trained with AR strength 4, 8, 16. We plot a subplot of a narrower range inside the plot of the full range to better show the histograms of examples that are around the middle values induced by AR. (c)(d) are standard deviations of singular values of weight matrices of NNs at each layer trained on CIFAR10/100 with AR strength 4, 16. The AR strength 8 is dropped for clarity.

The observation in section 4.2.1 shows that AR make NNs just *less confident* by reducing the variance of predictions made and concentrate margins more around zero. In this section, we study the *underlying factors* of AR that make NNs become less confident.

To begin with, we show that the singular values of the weight matrix of each layer determine the perturbation in margins of samples induced by perturbations in the instance space. Such a connection between singular values and the perturbation of outputs of a single layer, i.e., ReLU($W\delta x$), has been discussed in section 1.1. In the following, with lemma 3.1, we describe how the relatively more complex connection between margins and singular values of each weight matrix of layers of NNs holds. Observe that margins are obtained by applying a piece-wise linear mapping (c.f. the margin operator in definition 3) to the activation of the last layer of a NN. It implies the perturbations in activation of the last layer induce changes in margins in a piece-wise linear way. Meanwhile, the perturbation in the activation of the last layer (induced by perturbation in the instance space) is determined by the weight matrix's singular values of each layer of NNs. More specifically, this is explained as follows. Lemma 3.1 shows that the perturbation δI induced by δx , is given by $\sum_{j=1}^{n} \int_{s_{j}}^{e_{j}} \left| \left| \prod_{i=1}^{l} \boldsymbol{W}_{i}^{q_{j}} \delta \boldsymbol{x} dt \right| \right|$. Note that for each i, $\boldsymbol{W}_{i}^{q_{i}}$ is a matrix. By Cauchy interlacing law by row deletion (Chafai), the singular values of W_i , the weight matrix of layer i, determine the singular values of $W_i^{q_j}$. Thus, suppose l = 1, we have the change (measured in norm) induced by perturbation as $\sum_{j=1}^{n} \int_{s_{j}}^{e_{j}} || \boldsymbol{W}_{1}^{q_{j}} \delta \boldsymbol{x} dt ||$. The singular values of W_1 would determine the variance (of norms) of activation perturbations induced by perturbations δx , similarly as explained in section 1.1 except that the norm perturbation now is obtained by a summation of n terms $||\mathbf{W}_{1}^{q_{j}} \delta \mathbf{x} dt||$ (each of which is the exact form discussed in section 1.1) weighted by $1/(e_j - s_j)$. Similarly, for the case where $l = 2 \dots L - 1$, the singular values of W_l determine the variance of perturbations in the output of layer l that induced by the perturbations in the output of the previous layer (the input to layer l), i.e., layer l-1. Consequently, we choose to study these singular values.

We show the standard deviation of singular values of each layer of ResNet56 trained on CIFAR10/100 earlier in fig. 5c fig. 5d. Overall, the standard deviation of singular values associated with a layer of the NN trained with AR strength 4 is mostly larger than that of the NN with AR strength 16. The STD reduction in CIFAR100 is relatively smaller than CIFAR10, since as observed in fig. 4b, the AR induced concentration effect of margin distributions is also relatively less obvious than that in fig. 4a. More quantitative analysis is given in appendix B.2. This leads us to our *key results* described in section 1.1.

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Appendices

A. Related works

Generalization and robustness. Robustness in machine learning models is a large field. We review some more works that analyze robustness from the statistical perspective. The majority of works that study adversarial robustness from the generalization perspective study the generalization behaviors of machine learning models under adversarial risk. The works that study adversarial risk include Attias et al. (2018); Schmidt et al. (2018); Cullina et al. (2018); Yin & Bartlett (2018); Khim & Loh (2018); Sinha et al. (2018). The bounds obtained under the setting of adversarial risk characterize the risk gap introduced by adversarial examples. Thus, it is intuitive that a larger risk gap would be obtained for a larger allowed perturbation limit ϵ , which is roughly among the conclusions obtained in those bounds. That is to say, the conclusion normally leads to a larger generalization error as an algorithm is asked to handle more adversarial examples, for that it focuses on characterizing the error of adversarial examples, not that of natural examples. However, adversarial risk is not our focus. In this paper, we study when a classifier needs to accommodate adversarial examples, what is the influence that the accommodation has on generalization behaviors of empirical risk of natural data.

Hard and soft adversarial robust regularization. We study the behaviors of NNs that are trained in the way that adversarial examples are required to be classified correctly. We note that the adversarial robustness required can also be built in NNs in a soft way by adding a penalty term in the risk function. Relevant works includes Lyu et al. (2015) and Miyato et al. (2018). This line of works is not our subject of investigation. They focus on increasing test performance instead of defense performance. The focus of our works is to study the behaviors that lead to standard performance degradation when a network is trained to has a reasonable defense ability to adversarial examples. For example, a 50% accuracy on adversarial examples generated by PGD methods (Madry et al., 2018) in fig. 12 is a defense ability that can serve as a baseline for a reasonable defense performance. It is natural that in the setting where the requirement to defend against adversarial examples is dropped, the regularization can be weakened (added as a penalty term) to only aim to improve the test performance of the network. In this case, no performance degradation would occur, but the defense performance is also poor.

Explicit regularization that increases robustness of NNs by imposing smoothness through a penalty term. The smoothing effect of adversarial training on the loss surface has been observed in contemporary works (Moosavi-Dezfooli et al., 2019; Qin et al., 2019). And based on such an observation, explicit regularization is formulated by adding

a penalty term to the risk function to increase NNs' robustness. The message of this work is different from the insights of (Moosavi-Dezfooli et al., 2019; Qin et al., 2019) related to regularization. They (Moosavi-Dezfooli et al., 2019; Qin et al., 2019) show that if the output of NNs is explicitly smoothed through a penalty term thorough curvature regularization (Moosavi-Dezfooli et al., 2019), or local linearization (Qin et al., 2019), then a certain degree of adversarial robustness (AR) can be achieved. The penalty term works as a regularizer because it is explicitly formulated that way. It is not clear whether adversarial training, which is a different and arguably the most widely used technique, has the effect of a regularizer. This is the issue that is investigated in this work, and we show that adversarial training effectively regularizes NNs, which is not clear previously. In addition, this work has shown that adversarial training has a smoothing effect on features of all layers, instead of just the loss surface. Such a fine-grained analysis is possible because of the theoretical instruments developed in this work, and is absent previously.

B. Further empirical studies on adversarial robustness

B.1. Technique to build adversarial robustness

To begin with, we describe the technique that we use to build AR into NNs. As mentioned in the caption of fig. 1, we choose arguably the most well received technique, i.e., the adversarial training method (Madry et al., 2018). Specifically, we use l_{∞} -PGD (Madry et al., 2018) untargeted attack adversary, which creates an adversarial example by performing projected gradient descent starting from a random perturbation around a natural example. Then, NNs are trained with adversarial examples. NNs with different AR strength are obtained by training them with increasingly stronger adversarial examples. The adversarial strength of adversarial examples is measured in the l_{∞} norm of the perturbation applied to examples. l_{∞} -norm is rescaled to the range of 0-255 to present perturbations applied to different datasets in a comparable way; that means in fig. 1 fig. 3 fig. 4 fig. 5 and fig. 6 fig. 12, AR is measured in this scale. We use 10 steps of size 2/255 and maximum of = [4/255], 8/255, 16/255] respectively for different defensive strength in experiments. For example, a NN with AR strength 8 is a NN trained with adversarial examples generated by perturbations whose l_{∞} norm are at most 8. Lastly, we note that although adversarial training could not precisely guarantee an adversarial robustness radius of ϵ , a larger l_{∞} norm used in training would make NNs adversarially robust in a larger ball around examples. Thus, though the precise adversarial robustness radius is not known, we know that we are making NNs adversarially robust w.r.t. a larger ϵ . Consequently, it enables us to study the influence of ϵ -AR

on NNs by studying NNs trained with increasing l_{∞} norm.

B.2. Quantitative analysis of variance reduction in singular values

Here, we provide more quantitative analysis on fig. 5c and fig. 5d, as noted previously in section 4.2.2.

Quantitatively, we can look at the accumulated standard deviation (STD) difference in all layers. We separate the layers into two group: the group that the STD (denoted σ_i^4) of singular values of layer i (of the NN trained) with AR strength 4 that is larger than that (denoted σ_i^{16}) of AR strength 16; and the group that is smaller. In CIFAR10, for the first group, the summation of the difference/increments of STD of the two networks $(\sum_i \sigma_i^4 - \sigma_i^{16})$ is 4.7465, and the average is 0.1158. For the second groups, the summation $(\sum_i \sigma_i^{16} - \sigma_i^4)$ is 0.4372, and the average is 0.0312. In CIFAR100, the summation of the first group is 3.7511, and the average is 0.09618; the summation of the second group is 0.4372, and the average is 0.1103. The quantitative comparison shows that the accumulated STD decrease in layers that have their singular value STDs decreased (comparing STD of the NN with AR strength 16 with STD of the NN with AR strength 4) is a magnitude larger the accumulated STD increase in the layers that have their singular value STDs increased. The magnitude difference is significant since the STDs of singular values of most layers are around

B.3. Discrepancy between trends of loss and error rate gaps in large capacity NNs

In section 4.1, we have noted an inconsistent behaviors of CIFAR10, compared with that of CIFAR100 and Tiny-ImageNet: the error gap reduces for CIFAR100 and Tiny-ImageNet, but increases for CIFAR10. It might suggest that AR does not effectively regularize NNs in the case of CIFAR10. However, we show in this section that the abnormal behaviors of CIFAR10 are derived from the same margin concentration phenomenon observed in section 4.2.1 due to capacity difference, and compared with the error gaps, the GE/loss gaps are more faithfully representatives of the generalization ability of the NNs. Thus, the seemingly abnormal phenomenon corroborate, not contradict, the *key results* present in section 1.

Using CIFAR10 and CIFAR100 as examples and evidence in the previous sections, we explain how the discrepancy emerges from AR's influence on margin distributions of the same network trained on tasks with different difficulties. Further evidence that the discrepancy arises from capacity difference would be shown at appendix B.3, where we run experiments to investigate GE gap of NNs with varied capacities on the same task/dataset.

- 1. On CIFAR10, the margin distribution of training sets not only concentrate more around zero, but also skews towards zero. As shown in the margin distribution on training sets of CIFAR10 in fig. 4c, we find that the large error gap is caused by the high training accuracy that is achieved with a high concentration of training samples just slightly beyond the decision boundary. This phenomenon does not happen in CIFAR100. Comparing margin distribution on the test set in fig. 4(a) in fig. 4a, the margin distribution on the training set in fig. 4c is highly skewed, i.e., asymmetrically distributed w.r.t. mean. While the margin distributions of CIFAR100 training set in fig. 4d is clearly less skewed, and looks much more like a normal distribution, as that of the margin distribution on the test set.
- 2. The high skewness results from the fact that the NN trained on CIFAR10 is of large enough capacity to overfit the training set. As known, CIFAR100 is a more difficult task w.r.t. CIFAR10 with more classes and less training examples in each class. Thus, relatively, even the same ResNet56 network is used, the capacity of the network trained on CIFAR10 is larger than the one trained on CIFAR100. Recall that NNs have a remarkable ability to overfit training samples (Zhang et al., 2016). And note that though AR requires in a ball around an example, the examples in the ball should be of the same class, since the ball is supposed only to include imperceptible perturbation to the example, few of the training samples are likely in the same ball. Thus, the ability to overfit the training set is not regularized by AR: if NNs can overfit all training samples, it can still overfit some more examples that are almost imperceptibly different. For CI-FAR10, since NNs have enough capacity, the NN simply overfits the training set.
- 3. However, as shown in the observed overfitting phenomenon in fig. 4c, the high training accuracy is made up of correct predictions with relatively lower confidence (compared with NNs with lower AR), which is bad and not characterized by the error rate; and the low test accuracy are made up of wrong predictions with relatively lower confidence as well (as explain in section 4.2.1), which is good, and not characterized by error rate as well. Thus, the error gap in this case does not characterize the generalization ability (measured in term of prediction confidence) of NNs well, while the GE gap more faithfully characterizes the generalization ability, and show that AR effectively regularizes NNs. In the end, AR still leads to biased poorly performing solutions since the overfitting in training set does not prevent the test margin distribution concentrating more around zero, which leads to higher test errors of CIFAR10 as shown in fig. 3b. It further suggests that the damage AR done to the hypothesis space is not recovered by increasing capacity, however the ability of NNs to fit arbitrary labels

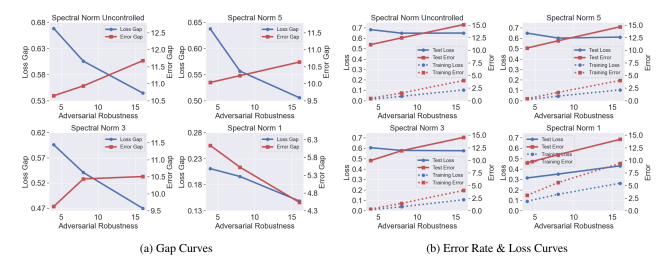


Figure 6. The four plots from upper left to lower bottom (in each subfigure) are NNs with increasingly smaller spectral complexity, where "Spectral Norm 1" means for each weight matrix of the NN, its spectral norm is at most 1. (a) Plots of training/test loss gap (and error gap) against adversarial robustness strength. (b) Training/test losses and error rates against increased strength of adversarial robustness.

is not hampered by AR.

B.4. Further evidence of regularization effects on NNs with varied capacity

In previous sections, we observe AR consistently effectively regularizes NNs; meanwhile, we also observe that in the case where a NN has a large capacity, it can spuriously overfit training samples and lead to an increased error gap. In this section, we present additional results by applying AR to networks of varied capacities. The effects of adversarial training on a larger NNs, i.e., ResNet 110 is given in appendix B.4.1. Then, AR applied on NNs with controlled capacities through spectral normalization is given in appendix B.4.2. This is to ensure that our observations and analysis in previous sections exist not just at some singular points, but also in a continuous area in the hypothesis space.

B.4.1. REGULARIZATION EFFECTS ON NNs WITH LARGER CAPACITY

To preliminarily validate that the regularization effects observed in section 4.1 manifest in NNs with varied capacities, we investigate the regularization effects of AR on a larger NNs, i.e., ResNet 110. The results are shown in fig. 7. The observed phenomenon is the same with that of ResNet56 presented in section 4.1, and thus corroborates our results.

B.4.2. REGULARIZATION EFFECTS ON NNS WITH CONTROLLED CAPACITIES

To control capacities of NNs quantitatively, we choose the measure based on spectral norm (Bartlett et al., 2017; Neyshabur et al., 2018a). In spectral norm based capacity measure bound (Bartlett et al., 2017; Neyshabur et al., 2018a), the NN capacity is normally proportional to a quantity called spectral complexity (SC), which is defined as follows.

Definition 7 (Spectral Complexity). *Spectral complexity* SC(T) of a NN T is the multiplication of spectral norms of weight matrices of layers in a NN.

$$SC(T) = \prod_{i=1}^{L} ||\boldsymbol{W}_i||_2$$

where $\{W_i\}_{i=1...L}$ denotes weight matrices of layers of the NN.

To control SC, we apply the spectral normalization (SN) (Sedghi et al., 2018) on NNs. The technique renormalizes the spectral norms of the weight matrices of a NN to a designated value after certain iterations. We carry out the normalization at the end of each epoch.

We train ResNet56 with increasingly strong AR and with increasingly strong spectral normalization. The results are shown in fig. 6.

As can be seen, as the capacity of NNs decreases (from upper left to bottom right in each sub-figure), the error gap between training and test gradually changes from an increasing trend to a decreasing trend, while the loss gap keeps a consistent decreasing trend. It suggests that the overfitting phenomenon is gradually prevented by another regularization techniques, i.e., the spectral normalization. As a result, the regularization effect of AR starts to emerge even in the error gap, which previously manifests only in the loss gap. The other curves corroborate our previous observations and analysis as well.

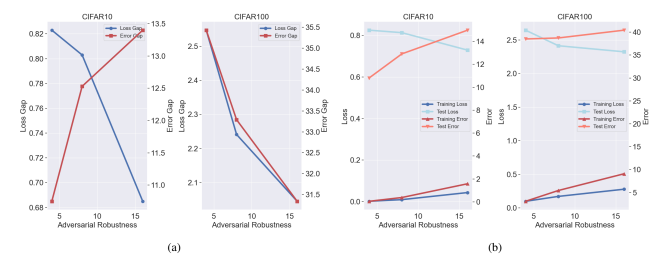


Figure 7. Experiment results on CIFAR10/100 with ResNet-110 (He et al., 2016). The unit of x-axis is the adversarial robustness (AR) strength of NNs, c.f. the beginning of section 4. (a) Plots of loss gap (and error rate gap) between training and test datasets v.s. AR strength. (b) Plots of losses (and error rates) on training and test datasets v.s. AR strength.

B.5. Further evidence on the smoothing effect of adversarial robustness

We quantitatively measure the smoothing effect around examples here by measuring the average maximal loss change/variation induced by the perturbation (of a fixed infinity norm) applied on examples. We found that the loss variation decreases as networks become increasingly adversarially robust. Note that the loss of an example is a proxy to the confidence of the example — it is the logarithm of the estimated probability (a characterization of confidence) of the NN classifier.

For a given maximal perturbation range characterized by the infinity norm, we generate adversarial examples within that norm for all test samples. For each example, the maximal loss variation/change of the adversarial example w.r.t. the natural example is computed for networks with different adversarial strength. To obtain statistical behaviors, we compute the average and standard deviation of such maxima of all test samples. The results are shown in fig. 8. The exact data can be found in table 1.

We can see that the average loss variation decreases with adversarial robustness. The standard deviation decreases with network adversarial robustness as well. The phenomenon that the standard deviation is comparably large with the mean might need some explanation. This is because different examples have different losses, thus the loss varies in relatively different regimens — the more wrongly classified examples vary in a larger magnitude, and vice versa for more correctly classified examples. This phenomenon leads to the large standard deviation of the loss variation.

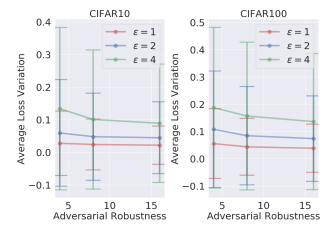


Figure 8. Average maximal loss variation induced by adversarial examples in networks with increasing adversarial robustness. The experiments are carried on CIFAR10/100. ϵ represents the maximal perturbation can be applied on natural test examples to generate adversarial examples. It is measured in the infinity norm. The larger the ϵ , the stronger the perturbation is. The error bars represent standard deviation.

B.6. Further experiments on using FGSM in adversarial training to build adversarial robustness

We explain the choice of PGD as the representative of adversarial training techniques here. Various adversarial training methods are variant algorithms that compute first order approximation to the point around the input example that minimizes the label class confidence. The difference is how close the approximation is. Recent works on adversarial examples exclusively only use PGD in experiments

	Attack Strength	Defensive Strength			
Dataset	Attack Stiength	4	8	16	
CIFAR10	$ \epsilon = 1 $ $ \epsilon = 2 $ $ \epsilon = 4 $	0.0273 ± 0.0989 0.0590 ± 0.1637 0.1337 ± 0.2494	$\begin{array}{c} 0.0236 \pm 0.0778 \\ 0.0477 \pm 0.1337 \\ 0.1006 \pm 0.2137 \end{array}$	$\begin{array}{c} 0.0215 \pm 0.0588 \\ 0.0443 \pm 0.1102 \\ 0.0888 \pm 0.1816 \end{array}$	
CIFAR100	$egin{aligned} \epsilon &= 1 \\ \epsilon &= 2 \\ \epsilon &= 4 \end{aligned}$	0.0550 ± 0.1276 0.1072 ± 0.2138 0.1868 ± 0.2946	0.0430 ± 0.1043 0.0839 ± 0.1802 0.1563 ± 0.2712	0.0379 ± 0.0886 0.0732 ± 0.1568 0.1355 ± 0.2494	

Table 1. Data of the smoothing effect of PGD adversarial training in fig. 8.

(Kannan et al., 2018; Schmidt et al., 2018; Xie et al., 2019; Ilyas et al., 2019; Wang & Zhang, 2019). It is also a very strong multi-step attack method that improves over many of its antecedents: NNs trained by FGSM could have no defense ability to adversarial examples generated by PGD, as shown in Table 5 in Madry et al. (2018); multi-step methods prevent the pitfalls of adversarial training with single-step methods that admit a degenerate global minimum (Tramèr et al., 2017). Thus, we believe the observations in this work is representative for various adversarial training techniques. Yet, even in the worst case, this work at least makes a first step to understand a representative approach of the approximation.

To corroborate the analysis, we also use FGSM (Goodfellow et al., 2015) in the adversarial training to build adversarial robustness into NNs. The results are consistent with the results obtained using PGD. The experiments are carried on CIFAR10/100. We present key plots that support the results obtained in the main content here. All the setting are same with that described in appendix B.1 of PGD, except that we replace PGD with FGSM.

Adversarial robustness reduces generalization gap and standard test performance. In section 4.1, we find that NNs with stronger adversarial robustness tend to have smaller loss/generalization gap between training and test sets. Consistent phenomenon has been observed in networks adversarially trained with FGSM on CIFAR10/100, as shown in fig. 9a. Consistent standard test performance degradation has been observed in adversarially trained with FGSM on CIFAR10/100 as well, as shown in fig. 9b. The exact data can be found in table 2.

Adversarial robustness concentrates examples around decision boundaries. In section 4.2.1, we find that the distributions of margins become more concentrated around zero as AR grows. The phenomenon has been observed consistently in networks adversarially trained with FGSM on CIFAR10/100, as shown in fig. 10. Phenomenon in fig. 5a and fig. 5b are also reproduced consistently in fig. 11b and fig. 11a. Please refer to section 4.2.1 for the analysis of the

results. Here we mainly present counterparts of the results analyzed there.

Adversarial robustness reduces the standard deviation of singular values of weight matrices in the network. In section 4.2.2, we find that for NNs with stronger adversarial robustness, the standard deviation of singular values of weight matrices is smaller in most layers. The phenomenon has been consistently observed in NNs trained with FGSM on CIFAR10/100, as shown in fig. 11c and fig. 11d. Please refer to section 1.1 and section 4.2.2 for the analysis of the results. Here we mainly present counterparts of the results analyzed there.

In conclusion, all key empirical results have been consistently observed in NNs trained with FGSM.

Table 2. Data of fig. 9

Dataset		Defensive Strength 4 8 16			
	Test Acc.	89.32	86.67	82.83	
GTE 1 5 1 0	Trn Loss	0.038	0.086	0.252	
CIFAR10	Test Loss	0.467	0.495	0.637	
	Δ Loss	0.429	0.409	0.385	
	Test Acc.	62.01	59.78	56.30	
	Trn Loss	0.469	0.656	0.822	
CIFAR100	Test Loss	1.776	1.723	1.797	
	Δ Loss	1.307	1.067	0.975	

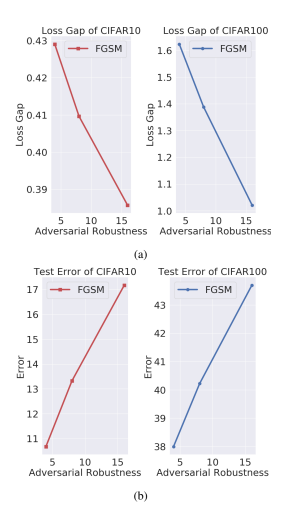


Figure 9. Experiment results on CIFAR10/100. The network is ResNet-56 (He et al., 2016). The unit of x-axis is the adversarial robustness (AR) strength of NNs, c.f. the beginning of section 4. (a) Plots of loss gap between training and test datasets v.s. AR strength. (b) Plots of error rates on training and test datasets v.s. AR strength.

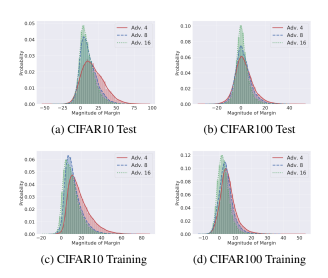


Figure 10. Margin distributions of NNs with AR strength 4, 8, 16 on Training and Test sets of CIFAR10/100.

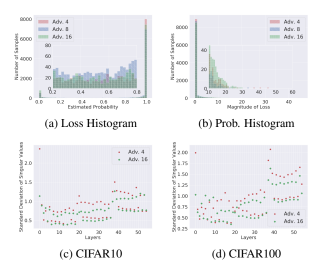


Figure 11. (a)(b) are histograms of estimated probabilities and losses respectively of the test set sample of NNs trained AR strength 4, 8, 16. We plot a subplot of a narrower range inside the plot of the full range to show the histograms of examples that are around the middle values to show the change induced by AR that induces more middle valued confidence predictions. (c)(d) are standard deviations of singular values of weight matrices of NNs at each layer trained on CIFAR10/100 with AR strength 4, 16. The AR strength 8 is dropped for clarity.

C. Proof of theorem 3.1

C.1. Algorithmic Robustness Framework

In order to characterize the bound to the GE, we build on the *algorithmic robustness* framework (Xu & Mannor, 2012).

We introduce the framework below.

Definition 8 $((K, \epsilon(\cdot))\text{-robust})$. An algorithm is $(K, \epsilon(\cdot))$ robust, for $K \in \mathbb{N}$ and $\epsilon(\cdot) : \mathcal{Z}^m \mapsto \mathbb{R}$, if \mathcal{Z} can be partitioned into K disjoint sets, denoted by $\mathcal{C} = \{C_k\}_{k=1}^K$, such that the following holds for all $s_i = (x_i, y_i) \in S_m, z = (x, y) \in \mathcal{Z}, C_k \in \mathcal{C}$:

$$\forall s_i = (\boldsymbol{x}_i, y_i) \in C_k, \forall z = (\boldsymbol{x}, y) \in C_k$$

$$\implies |l(f(\boldsymbol{x}_i), y_i) - l(f(\boldsymbol{x}), y)| \le \epsilon(S_m).$$

The gist of the definition is to constrain the variation of loss values on test examples w.r.t. those of training ones through local property of the algorithmically learned function f. Intuitively, if $s \in S_m$ and $z \in \mathcal{Z}$ are "close" (e.g., in the same partition C_k), their loss should also be close, due to the intrinsic constraint imposed by f.

For any algorithm that is robust, Xu & Mannor (Xu & Mannor, 2012) proves

Theorem C.1 (Xu & Mannor (Xu & Mannor, 2012)). *If* a learning algorithm is $(K, \epsilon(\cdot))$ -robust and \mathcal{L} is bounded, a.k.a. $\mathcal{L}(f(x), y) \leq M \ \forall z \in \mathcal{Z}$, for any $\eta > 0$, with probability at least $1 - \eta$ we have

$$GE(f_{S_m}) \le \epsilon(S_m) + M\sqrt{\frac{2K\log(2) + 2\log(1/\eta)}{m}}.$$
 (6)

To control the first term, an approach is to constrain the variation of the loss function. Covering number (Shalev-Shwartz & Ben-David, (Shalev-Shwartz & Ben-David, 2014), Chapter 27) provides a way to characterize the variation of the loss function, and conceptually realizes the actual number K of disjoint partitions.

For any regular k-dimensional manifold embedded in space equipped with a metric ρ , e.g., the image data embedded in $L^2(\mathbb{R}^2)$, the square integrable function space defined on \mathbb{R}^2 , it has a covering number $\mathcal{N}(\mathcal{X};\rho,\epsilon)$ of $(C_{\mathcal{X}}/\epsilon)^k$ (Verma, 2013), where $C_{\mathcal{X}}$ is a constant that captures its "intrinsic" properties, and ϵ is the radius of the covering ball. When we calculate the GE bound of NNs, we would assume the data space is a k-dimensional regular manifold that accepts a covering.

Adversarial robustness makes NNs a $(K, \epsilon(\cdot))$ -robust algorithm, and is able to control the variation of loss values on test examples. Building on covering number and theorem C.1, we are able to prove theorem 3.1.

C.2. Proof

Proof of lemma 3.1. By theorem 3 in Sokolic et al. (2017), we have

$$||I_l(\boldsymbol{x}) - I_l(\boldsymbol{x}')|| = \left| \left| \int_0^1 \boldsymbol{J}(\boldsymbol{x} - t(\boldsymbol{x}' - \boldsymbol{x})) dt(\boldsymbol{x} - \boldsymbol{x}') \right| \right|$$
(7)

where J(x) denotes the Jacobian of $I_l(x)$ at x.

By lemma 3.2 in Jia et al. (2019), when we only have max pooling layers and ReLU as nonlinear layer in NNs, J(x) is a linear operator at a local region around x. For terminology concerning regions, we follow the definitions in Jia et al. (2019). More specifically, we have

$$\boldsymbol{J}(\boldsymbol{x}) = \prod_{i=1}^l \boldsymbol{W}_i^{\boldsymbol{x}}$$

where \boldsymbol{W}_{i}^{x} is the linear mapping (matrix) induced by $\boldsymbol{J}(x)$ at \boldsymbol{x} . It is a matrix obtained by selectively setting certain rows of \boldsymbol{W}_{i} to zero. For the more concrete form of \boldsymbol{W}_{i}^{x} , refer to lemma 3.2 in Jia et al. (2019). In Jia et al. (2019), it is noted as \boldsymbol{W}_{i}^{q} , where q is a region where \boldsymbol{x} is in.

Suppose that from x to x', the line segment x - x' passes through regions $\{q_i\}_{i=1,\dots,n}$. The line segment is illustrated in fig. 2b as the boldest black line segment at the upper half of the figure. In the illustration, x - x' passes through three regions, colored coded as gray, dark yellow, light blue respectively. The line segment is divided into three subsegments. Suppose l(t) = x + t(x' - x). Then the three sub-segments can be represented by l(t) as $l(s_1)$ to $l(e_1)$, $l(s_2)$ to $l(e_2)$, and $l(s_3)$ to $l(e_3)$ respectively, as noted on the line segment in the illustration. Originally, the range of the integration in eq. (7) is from 0 to 1, representing the integration on the line segment l(0) to l(1) in the instance space. Now, since for each of these regions trespassed by the line segment, the Jacobian J(x) is a linear operator, denoted as $W_i^{q_j}$, the integration in eq. (7) from 0 to 1 can be decomposed as a summation of integration on segments $l(s_1)$ to $l(e_1)$ etc. In each of these integration, the Jacobian J(x) is the multiplication of linear matrices $W_i^{q_j}$, i.e., $\prod_{i=1}^{l} W_{i}^{q_{i}}$. Thus, eq. (7) can be written as

$$\sum_{i=1}^n \int_{s_j}^{e_j} \left| \left| \prod_{i=1}^l oldsymbol{W}_i^{q_j} dt(oldsymbol{x} - oldsymbol{x}')
ight|
ight|$$

where s_j, e_j denotes the start and end of the segment $[s_j, e_j] \subset [0, 1]$ of the segment [0, 1] that passes through the region q_j .

In the cases that a linear operator is applied on the feature map $I_l(x)$ without any activation function, we can also

obtain a similar conclusion. Actually, such cases are just degenerated cases of feature maps that have activation functions.

Corollary C.1. Given two elements x, x', and $I_l(x) = W_l g(W_{l-1} \dots g(W_1 x))$, we have

$$||I_l(oldsymbol{x}) - I_l(oldsymbol{x}')|| = \sum_{j=1}^n \int_{s_j}^{e_j} \left| \left| oldsymbol{W}_l \prod_{i=1}^{l-1} oldsymbol{W}_i^{q_j} dt (oldsymbol{x} - oldsymbol{x}')
ight|
ight|$$

where symbols are defined similar as in Proof of lemma 3.1.

Now, we are ready to prove theorem C.1.

Proof of theorem C.1. Similar with the proof of theorem C.1, we partition space $\mathcal Z$ into the ϵ -cover of $\mathcal Z$, which by assumption is a k-dimension manifold. Its covering number is upper bounded by $C_{\mathcal X}^k/\epsilon^k$, denoting $K=C_{\mathcal X}^k/\epsilon^k$, and $\hat C_i$ the ith covering ball. For how the covering ball is obtained from the ϵ -cover, refer to theorem 6 in Xu & Mannor (2012). We study the constraint/regularization that adversarial robustness imposes on the variation of the loss function. Since we only have ϵ -adversarial robustness, the radius of the covering balls is at most ϵ — this is why we use the same symbol. Beyond ϵ , adversarial robustness does not give information on the possible variation anymore. Let T' denotes the NN without the last layer.

First, we analyze the risk change in a covering ball C_i . The analysis is divided into two cases: 1 all training samples in C_i are classified correctly; 2) all training samples in C_i are classified wrong. Note that no other cases exist, for that the radius of C_i is restricted to be ϵ , and we work on ϵ -adversarial robust classifiers. It guarantees that all samples in a ball are classified as the same class. Thus, either all training samples are all classified correctly, or wrongly.

We first study case 1). Given any example $z = (x, y) \in C_i$, let $\hat{y} = \arg \max_{i \neq y} w_i^T T' x$. Its ramp loss is

$$l_{\gamma}(\boldsymbol{x}, y) = \max\{0, 1 - \frac{1}{\gamma}(\boldsymbol{w}_{y} - \boldsymbol{w}_{\hat{y}})^{T} T' \boldsymbol{x}\}.$$

Note that within C_i , $(\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T T' \boldsymbol{x} \geq 0$, thus $l_{\gamma}(\boldsymbol{x}, y)$ is mostly 1, and we would not reach the region where r > 0 in definition 4. Let $u(\boldsymbol{x}) := (\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T T' \boldsymbol{x}$, and $u_{\min}^i = \min_{\forall \boldsymbol{x} \in C_i} u(\boldsymbol{x})$. We have

$$l_{\gamma}(\boldsymbol{x}, y) \le \max\{0, 1 - \frac{u_{\min}^{i}}{\gamma}\} \le \max\{0, 1 - \frac{u_{\min}}{\gamma}\},$$

where u_{\min} denotes the smallest margin among all partitions

The inequality above shows adversarial robustness requires that T'x should vary slowly enough, so that in the worst case, the loss variation within the adversarial radius should

satisfy the above inequality. The observation leads to the constraint on the loss difference $\epsilon(\cdot)$ defined earlier in definition 8 in the following.

Given any training example $z := (x, y) \in C_i$, and any element $z' := (x', y') \in C_i$, where C_i is the covering ball that covers x, we have

$$|l_{\gamma}(\boldsymbol{x}, y) - l_{\gamma}(\boldsymbol{x}', y')|$$

$$= |\max\{0, 1 - \frac{u(\boldsymbol{x})}{\gamma}\} - \max\{0, 1 - \frac{u(\boldsymbol{x}')}{\gamma}\}|$$

$$\leq \max\{0, 1 - \frac{u_{\min}}{\gamma}\}.$$
(8)

Now we relate the margin to the margin in the instance space.

Given $z := (x, y) \in \mathcal{Z}$, and z', of which x' is the closest points to x (measured in Euclidean norm) on the decision boundary, we can derive the inequality below.

$$u(\mathbf{x}) = u(\mathbf{x}) - u(\mathbf{x}')$$

$$= \int_{0}^{1} \mathbf{J}(\mathbf{x} - t(\mathbf{x} - \mathbf{x}')) dt(\mathbf{x} - \mathbf{x}')$$

$$= \int_{0}^{1} (\mathbf{w}_{y} - \mathbf{w}_{\hat{y}})^{T} \prod_{i=1}^{L-1} \mathbf{W}_{i}^{\mathbf{x} - t(\mathbf{x} - \mathbf{x}')} dt(\mathbf{x} - \mathbf{x}')$$

$$= \int_{0}^{1} \left| (\mathbf{w}_{y} - \mathbf{w}_{\hat{y}})^{T} \prod_{i=1}^{L-1} \mathbf{W}_{i}^{\mathbf{x} - t(\mathbf{x} - \mathbf{x}')} dt(\mathbf{x} - \mathbf{x}') \right|$$

$$= \sum_{j=1}^{n} \int_{s_{j}}^{e_{j}} \left| (\mathbf{w}_{y} - \mathbf{w}_{\hat{y}})^{T} \prod_{i=1}^{L-1} \mathbf{W}_{i}^{q_{j}} dt(\mathbf{x} - \mathbf{x}') \right|$$

$$\geq \min_{y, \hat{y} \in \mathcal{Y}, y \neq \hat{y}} ||\mathbf{w}_{y} - \mathbf{w}_{\hat{y}}||_{2} \prod_{i=1}^{L-1} \sigma_{\min}^{i} ||\mathbf{x} - \mathbf{x}'||_{2} \int_{0}^{1} dt$$

$$\geq \min_{y, \hat{y} \in \mathcal{Y}, y \neq \hat{y}} ||\mathbf{w}_{y} - \mathbf{w}_{\hat{y}}||_{2} \prod_{i=1}^{L-1} \sigma_{\min}^{i} ||\mathbf{x} - \mathbf{x}'||_{2}$$

$$(13)$$

where J(x) denotes the Jacobian of $I_l(x)$ at x. eq. (10) can be reached by theorem 3 in Sokolic et al. (2017). eq. (11) can be reached because $(w_y - w_{\hat{y}})W_i^{x-t(x-x')}(x-x')$ is the actually classification score u(x), u(x') difference between x, x', and by assumptions assumption 3.1, they are positive throughout. eq. (12) is reached due to corollary C.1 — in this case, the matrix W_l in corollary C.1 is of rank one.

To arrive from eq. (12) to eq. (13), we observe that x' is the closest point to x on the decision boundary. Being the closest means $x - x' \perp \mathcal{N}((w_y - w_{\hat{y}})T')$. If the difference x' - x satisfies $x - x' \not\perp \mathcal{N}(T')$, we can always remove

the part in the $\mathcal{N}(T')$, which would identify a point that is closer to x, but still on the decision boundary, which would be a contradiction. Then if x-x' is orthogonal to the null space, we can bound the norm using the least singular values. We develop the informal reasoning above formally in the following.

Similarly in lemma 3.4 in Jia et al. (2019), by Cauchy interlacing law by row deletion, assuming $\boldsymbol{x} \perp \mathcal{N}(\prod_{i=1}^{L-1} \boldsymbol{W}_i^{q_j})$ (\mathcal{N} denotes the null space; the math statement means \boldsymbol{x} is orthogonal to the null space of $\boldsymbol{J}(\boldsymbol{x})$), we have

$$||\prod_{i=1}^{L-1} \boldsymbol{W}_{i}^{q_{j}} \boldsymbol{x}||_{2} \ge \prod_{i=1}^{L-1} \sigma_{\min}^{i} ||\boldsymbol{x}||_{2}$$
 (14)

where σ_{\min}^i is the smallest singular value of \boldsymbol{W}_i . Then conclusion holds as well for multiplication of matrices $\prod_{i=1}^{L-1} \boldsymbol{W}_i^{q_j}$, since the multiplication of matrices are also a matrix.

Notice that in each integral in eq. (12), we are integrating over constant. Thus, we have it equates to

$$\sum_{j=1}^{n} (e_j - s_j) \left| (\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T \prod_{i=1}^{L-1} \boldsymbol{W}_i^{q_j} (\boldsymbol{x} - \boldsymbol{x}') \right|.$$

Now we show that in each operand, $\boldsymbol{x} - \boldsymbol{x}' \perp \mathcal{N}((\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T \prod_{i=1}^{L-1} \boldsymbol{W}_i^{q_j})$. Denote T_{q_j} as $\mathcal{N}((\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T \prod_{i=1}^{L-1} \boldsymbol{W}_i^{q_j})$. Suppose that it does not hold. Then we can decompose $\boldsymbol{x} - \boldsymbol{x}'$ into two components $\boldsymbol{\Delta}_1, \boldsymbol{\Delta}_2$, where $\boldsymbol{\Delta}_1 \perp T_{q_j}, \boldsymbol{\Delta}_2 \not\perp T_{q_j}$. We can find a new point $\boldsymbol{x}'' = \boldsymbol{x} + \boldsymbol{\Delta}_1$ that is on the boundary. However, in this case

$$||x - x''||_2 = ||\Delta_1||_2 \le ||\Delta_1||_2 + ||\Delta_2||_2 = ||x - x'||_2$$

Recall that x' is the closest point to x on the decision boundary. This leads to a contradiction. Repeat this argument for all $j=1,\ldots,n$, then we have x-x' be orthogonal to all $\mathcal{N}(T_{q_j})$. Thus, by the inequality eq. (14) earlier, we can arrive at eq. (13) — notice that $w_y-w_{\hat{y}}$ is a matrix with one column, thus also satisfies the above reasoning.

Through the above inequality, we can transfer the margin to margin in the instance space. Let $v(\boldsymbol{x})$ be the shortest distance in $||\cdot||_2$ norm from an element $\boldsymbol{x} \in \mathcal{X}$ to the decision boundary. For a covering ball C_i , let v_{\min}^i be $\min_{\boldsymbol{x} \in C_i} v(\boldsymbol{x})$. Let v_{\min} be the smallest v_{\min}^i among all covering balls C_i that contain at least a training example. We have that

$$u_{\min} \geq \min_{y,\hat{y} \in \mathcal{Y}, y \neq \hat{y}} || \boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}} ||_2 \prod_{i=1}^{L-1} \sigma_{\min}^i v_{\min}$$

Consequently, we can obtain an upper bound of eq. (8)

parameterized on v_{\min} , as follows

$$\begin{split} \max\{0, 1 - \frac{u_{\min}}{\gamma}\} &\leq \max\{0, \\ 1 - \frac{\min_{y, \hat{y} \in \mathcal{Y}, y \neq \hat{y}} ||\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}}||_2 \prod_{i=1}^{L-1} \sigma_{\min}^i v_{\min}}{\gamma}\}. \end{split}$$

Notice that only because ϵ_0 -adversarial robustness, we can guarantee that v_{\min} is non-zero, thus the bound is influenced by AR.

Then, we study case 2), in which all training samples $z \in C_i$ are classified wrong. In this case, for all $z \in C_i$, the \hat{y} given by $\hat{y} = \arg\max_{i \neq y} \boldsymbol{w}_i^T T' \boldsymbol{x}$ in the margin operator is the same, for that \hat{y} is the wrongly classified class. Its ramp loss is

$$l_{\gamma}(\boldsymbol{x}, y) = \max\{0, 1 - \frac{1}{\gamma}(\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T T' \boldsymbol{x}\}.$$

Note that in the case 1), it is the y that stays fixed, while \hat{y} may differ from example to example; while in the case 2), it is the \hat{y} stays fixed, while y may differ.

Similarly, within C_i as required by adversarial robustness, $(\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T T' \boldsymbol{x} \leq 0$, thus we always have $1 - \frac{1}{\gamma} (\boldsymbol{w}_y - \boldsymbol{w}_{\hat{y}})^T T' \boldsymbol{x} \geq 1$, implying

$$l_{\gamma}(\boldsymbol{x},y)=1.$$

Thus, $\forall z = (\boldsymbol{x}, y), z' = (\boldsymbol{x}', y') \in C_i$

$$|l_{\gamma}(\mathbf{x}, y) - l_{\gamma}(\mathbf{x}', y')| = 0.$$
 (15)

Since only these two cases are possible, by eq. (8) and eq. (15), we have $\forall z, z' \in C_i$

$$|l_{\gamma}(\boldsymbol{x}, y) - l_{\gamma}(\boldsymbol{x}', y')| \le \max\{0, 1 - \frac{u_{\min}}{\gamma}\}.$$
 (16)

The rest follows the standard proof in algorithmic robust framework.

Let N_i be the set of index of points of examples that fall into C_i . Note that $(|N_i|)_{i=1...K}$ is an IDD multimonial random

variable with parameters m and $(|\mu(C_i)|)_{i=1...K}$. Then

D. Implementation Details

We summarize the details of the experiments in this section. The experiments are run with PyTorch (Pfeiffer, 2017).

$$|R(l \circ T) - R_{m}(l \circ T)|$$

$$= |\sum_{i=1}^{K} \mathbb{E}_{Z \sim \mu}[l(TX, Y)]\mu(C_{i}) - \frac{1}{m} \sum_{i=1}^{m} l(Tx_{i}, y_{i})|$$

$$\leq |\sum_{i=1}^{K} \mathbb{E}_{Z \sim \mu}[l(TX, Y)] \frac{|N_{i}|}{m} - \frac{1}{m} \sum_{i=1}^{m} l(Tx_{i}, y_{i})|$$

$$+ |\sum_{i=1}^{K} \mathbb{E}_{Z \sim \mu}[l(TX, Y)]\mu(C_{i}) - \sum_{i=1}^{K} \mathbb{E}_{Z \sim \mu}[l(TX, Y)] \frac{|N_{i}|}{m}|$$

$$\leq |\frac{1}{m} \sum_{i=1}^{K} \sum_{j \in N_{i}} \max_{z \in C_{i}} |l(Tx, y) - l(Tx_{j}, y_{j})|$$
(17)

$$+ |\max_{z \in \mathcal{Z}} |l(Tx, y)| \sum_{i=1}^{K} |\frac{|N_i|}{m} - \mu(C_i)||.$$
 (18)

Remember that z = (x, y).

By eq. (16) we have eq. (17) is equal or less than $\max\{0,2(1-\frac{u_{\min}}{\gamma})\}$. By Breteganolle-Huber-Carol inequality, eq. (18) is less or equal to $\sqrt{\frac{\log(2)2^{k+1}C_{\mathcal{X}}^k}{\gamma^k m}+\frac{2\log(1/\eta)}{m}}.$

The proof is finished.

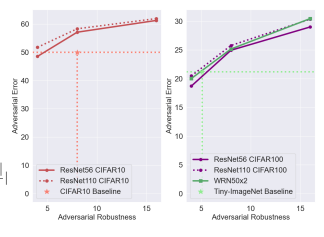


Figure 12. The plot of accuracy on adversarial examples v.s. adversarial defense strength built in NNs. The dotted line of which the intersections are marked by stars are adversarial accuracy in Madry et al. (2018) (CIFAR10), in Li et al. (2018) (Tiny ImageNet) under similar adversarial attack strength.

D.1. Datasets

CIFAR10/100. Each CIFAR dataset consists of 50,000 training data and 10,000 test data. CIFAR-10 and CIFAR-100 have 10 and 100 classes respectively. Our data augmentation follows the standard manner in Lee et al. (2015): during training, we zero-pad 4 pixels along each image side, and sample a 32×32 region cropped from the padded image or its horizontal flip; during testing, we use the original non-padded image.

Tiny-ImageNet. Tiny-ImageNet is a subset of ImageNet dataset, which contains 200 classes rather than 1,000 classes. Each class has 500 training images and 50 validation images. Images in the Tiny-ImageNet dataset are of 64×64 pixels, as opposed to 256×256 in the full ImageNet set. The data augmentation is straightforward: an input image is 56×56 randomly cropped from a resized image using the scale, aspect ratio augmentation as well as scale jittering. A single 56×56 cropped image is used for testing.

D.2. Experiments in section 4.1

CIFAR10/100 Models and Training. The models for CIFAR10/100 are the same as the ones in appendix B.3, except that we do not use spectral normalization anymore. CIFAR100 has 100 output neurons instead of 10.

Tiny-ImageNet Model. For Tiny ImageNet dataset, we use

Table 3. Raw data of CIFAR10 dataset for plots in fig. 3, fig. 7 and fig. 12.

Method		Defensive Strength			
Method		4	8	16	
	Trn Acc.	99.51	98.45	95.97	
	Test Acc.	88.86	87.51	84.89	
ResNet-56	$\Delta Acc.$	10.65	10.94	11.08	
+ Adv Trn	Trn Loss	0.014	0.043	0.105	
T Auv IIII	Test Loss	0.683	0.649	0.650	
	$\Delta { m Loss}$	0.669	0.606	0.545	
	PGD	65.92	65.24	72.16	
	Trn Acc.	99.95	99.62	98.42	
	Test Acc.	89.20	87.09	85.02	
ResNet-110	$\Delta Acc.$	10.75	12.53	13.40	
+ Adv Trn	Trn Loss	0.002	0.010	0.044	
T AUV IIII	Test Loss	0.825	0.813	0.729	
	ΔLoss	0.823	0.803	0.685	
	PGD	58.02	66.94	72.40	

50-layered wide residual networks with 4 groups of residual layers and [3,4,6,3] bottleneck residual units for each group respectively. The 3×3 filter of the bottleneck residual units have $[64\times k,128\times k,256\times k,512\times k]$ feature maps with the widen factor k=2 as mentioned in Zagoruyko & Komodakis (2016). We replace the first 7×7 convolution layer with 3×3 filters with stride 1 and padding 1. The max pooling layer after the first convolutional layer is also removed to fit the 56×56 input size. Batch normalization layers are retained for this dataset. The weights of convolution layers for Tiny ImageNet are initialized with Xavier uniform (Glorot & Bengio, 2010). Again, all dropout layers are omitted.

Tiny-ImageNet Training. The experiments on the Tiny-ImageNet dataset are based on a mini-batch size of 256 for 90 epochs. The initial learning rate is set to be 0.1 and decayed at 10 at 30 and 60 epochs respectively. All experiments are trained on the training set with stochastic gradient descent with the momentum of 0.9.

Results. The data for fig. 3 and fig. 7 are given in table 3, table 4 and table 5. More specifically, the data on CIFAR10 are given in table 3. The result on CIFAR100 are given in table 4. The result on Tiny-ImageNet are given in table 5.

Adversarial Robustness Attack Method. The adversarial accuracy is evaluated against l_{∞} -PGD (Madry et al., 2018) untargeted attack adversary, which is one of the strongest white-box attack methods. When considering adversarial attack, they usually train and evaluate against the same perturbation. And for our tasks, we only use the moderate adversaries that generated by 10 iterations with steps of size 2 and maximum of 8. When evaluating adversarial robustness, we only consider clean examples classified correctly originally, and calculate the accuracy of the adversarial ex-

Table 4. Raw data of CIFAR100 dataset for the plot in fig. 3, fig. 7 and fig. 12.

Method		Defensive Strength		
Method		4	8	16
	Trn Acc.	88.73	86.97	82.17
	Test Acc.	61.31	60.87	59.43
ResNet-56	$\Delta Acc.$	27.42	26.10	22.74
+ Adv Trn	Trn Loss	0.357	0.413	0.570
+ Auv IIII	Test Loss	2.063	2.106	1.978
	$\Delta { m Loss}$	1.706	1.693	1.408
	PGD	30.52	40.99	48.81
	Trn Acc.	96.91	94.55	90.90
	Test Acc.	61.48	61.26	59.56
ResNet-110	$\Delta Acc.$	35.43	33.29	31.34
+ Adv Trn	Trn Loss	0.098	0.171	0.278
+ Auv IIII	Test Loss	2.645	2.413	2.323
	$\Delta { m Loss}$	2.547	2.241	2.045
	PGD	33.33	42.08	50.99

Table 5. Raw data of Tiny-ImageNet dataset for the plot in fig. 3, fig. 7 and fig. 12.

Method		Defensive Strength				
Method		0	4	8	16	
	Trn Acc.	79.12	73.71	66.17	60.73	
	Test Acc.	63.43	62.09	61.09	57.36	
Wide ResNet	$\Delta Acc.$	15.69	11.62	5.08	3.37	
+ Adv Trn	Trn Loss	0.874	1.080	1.384	1.641	
+ Auv IIII	Test Loss	1.561	1.637	1.689	1.806	
	$\Delta { m Loss}$	0.687	0.557	0.305	0.165	
	PGD	0.00	32.26	41.20	53.12	

amples generated from them that are still correctly classified. The adversarial accuracy is given in table 3 table 4 table 5, the row named "PGD", and plotted in fig. 12.

D.3. Experiments in appendix B.3

Models. We use ResNet-type networks (Zhang et al., 2018). Given that we need to isolate factors that influence spectral complexity, we use ResNet without additional batch normalization (BN) layers. To train ResNet without BN, we rely on the fixup initialization proposed in Zhang et al. (2018). The scalar layers in Zhang et al. (2018) are also omitted, since it changes spectral norms of layers. Dropout layers are omitted as well. Following Sedghi et al. (2018), we clip the spectral norm every epoch rather than every iteration.

Training. The experiments on CIFAR10 datasets are based on a mini-batch size of 256 for 200 epochs. The learning rate starts at 0.05, and is divided by 10 at 100 and 150 epochs respectively. All experiments are trained on training set with stochastic gradient descent based on the momentum of 0.9.

Table 6. Raw data for fig. 6. SP denotes spectral norm.

Strength of		Defensive Strength		
Spectral Normalization		4	8	16
	Trn Acc.	96.91	94.38	90.58
	Test Acc.	90.47	88.87	85.82
	$\Delta Acc.$	6.44	5.51	4.76
SP 1	Trn Loss	0.092	0.159	0.265
	Test Loss	0.316	0.353	0.432
	ΔLoss	0.224	0.194	0.168
	PGD	57.93	69.98	75.98
	Trn Acc.	99.65	98.51	95.94
	Test Acc.	90.02	88.07	85.43
	$\Delta Acc.$	9.63	10.44	10.51
SP 3	Trn Loss	0.010	0.039	0.107
	Test Loss	0.606	0.580	0.577
	$\Delta { m Loss}$	0.596	0.541	0.470
	PGD	56.83	67.73	73.41
	Trn Acc.	99.57	98.33	95.96
	Test Acc.	89.53	88.09	85.32
	$\Delta Acc.$	10.04	10.24	10.64
SP 5	Trn Loss	0.012	0.045	0.105
	Test Loss	0.649	0.602	0.611
	$\Delta { m Loss}$	0.638	0.557	0.506
	PGD	54.91	65.96	72.37
	Trn Acc.	99.51	98.45	95.97
	Test Acc.	88.86	87.51	84.89
	$\Delta Acc.$	10.65	10.94	11.08
SP Uncontrolled	Trn Loss	0.014	0.043	0.105
	Test Loss	0.683	0.649	0.650
	$\Delta { m Loss}$	0.669	0.606	0.545
	PGD	65.92	65.24	72.16

Results. The data for fig. 6 are given in table 6.