A Space-Time Approach to Analyzing Stochastic Gradient Descent

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Abstract

We analyze of Stochastic Gradient Descent (SGD) at small learning rates. Unlike prior analyses based on stochastic differential equations, our theory models discrete time and hence non-Gaussian noise. We prove that gradient noise systematically pushes SGD toward flatter minima. We characterize when and why flat minima overfit less than sharp minima. We generalize the Akaike Info. Criterion (AIC) to a smooth estimator of overfitting, hence enabling gradient-based model selection. We show how non-stochastic GD with a modified loss function may emulate SGD. We verify our predictions on convnets for CIFAR-10 and Fashion-MNIST.

1. Introduction

Practitioners benefit from the intuition that SGD approximates noiseless GD (Bottou, 1991). In this paper, we refine that intuition by showing how gradient noise *biases* learning toward certain areas of weight space.

Departing from prior work, we model discrete time and hence non-Gaussian noise. Indeed, we derive corrections to continuous-time, Gaussian-noise approximations such as ordinary and stochastic differential equations (ODE, SDE). For example, we construct a loss landscape on which SGD eternally cycles counterclockwise, a phenomenon impossible with ODEs. Our experiments on image classifiers show that even a single evaluation of our force laws may predict SGD's motion through macroscopic timescales, e.g. long enough to decrease error by 0.5 percentage points.

Our work offers a novel interpretation of SGD as a superposition of concurrent interactions between weights an data, each represented by a diagram analogous to those of Feynman (1949); Penrose (1971). In the conclusion, we discuss this bridge to physics — and its relation to Hessian methods and natural GD — as topics for future research.

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1.1. Example of diagram-based reasoning

Our theory analyzes SGD in terms of combinatorial objects we call *diagrams*. Deferring details, we illustrate how our theory yields non-trivial results via short arguments. First, we list how components of diagrams encode statistics of the loss $l_x(\theta)$ at weight θ and datapoint x:

$$G \triangleq \mathbb{E}_{x} \left[\nabla l_{x}(\theta) \right] \triangleq$$

$$H \triangleq \mathbb{E}_{x} \left[\nabla \nabla l_{x}(\theta) \right] \triangleq$$

$$C \triangleq \mathbb{E}_{x} \left[(\nabla l_{x}(\theta) - G)^{2} \right] \triangleq$$

$$J \triangleq \mathbb{E}_{x} \left[(\nabla \nabla \nabla l_{x}(\theta)) \right] \triangleq$$

$$S \triangleq \mathbb{E}_{x} \left[(\nabla l_{x}(\theta) - G)^{3} \right] \triangleq$$

Table 1. Notation. Throughout, G, H, J denote the 1st, 2nd, and 3rd derivatives of the loss function. We write C, S for the 2nd and 3rd cumulants of the gradient distribution. We differentiate w.r.t. the weight θ and we take expectations w.r.t. the datapoints x. Note: the tensors J, S have three indices. Each $\nabla^d l_x$ corresponds to a node with d thin edges emanating, and fuzzy outlines connect nodes that occur within the same expectation.

We may connect Table 1's diagrams together to obtain *complete diagrams* without loose ends. For example, we may connect two copies of $G = {}^{\downarrow}$ with one copy of $H = {}^{\vee}$ to obtain ${}^{\frown}$.* If we run SGD for T gradient steps with learning rate η starting at θ_0 , then by Taylor expansion we may express the expected test loss at the final weight θ_T in terms of the statistics in Table evaluated at the initialization θ_0 . Diagrams organize the computation of this Taylor series.

Main Idea (Informal). There is a method to assign to any complete diagram a number that depends on η , T. SGD's expected test loss is a sum, over all complete diagrams, of these numbers. We incur only an $o(\eta^d)$ error if we consider only diagrams with at most d edges.

Example 1 (How does non-Gaussian noise affect test loss?). Assume † θ_0 minimizes the test loss and that we run SGD for 1 epoch with batch size 1. The skew S is 0 for Gaussians, and we seek the effect of non-zero S. To compute the leading-order effect of S on test loss, we identify the fewest-edged complete diagrams containing $S = \bigcup_{i=1}^{N} S_i$. In this

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^{*} We color nodes for convenient reference (e.g. to a diagram's "green nodes"). As mere labels, colors lack mathematical meaning.

† for simplicity. Our theory is not limited to this setting.

case, there is one such diagram: Then, working in a basis that diagonalizes ηH , we obtain the leading-order effect of S on test loss (with error $o(\eta^3)$):

$$-\frac{\eta^3}{6} \sum_{\mu\nu\lambda} S_{\mu\nu\lambda} \frac{1 - \exp(-T\eta(H_{\mu\mu} + H_{\nu\nu} + H_{\lambda\lambda}))}{\eta(H_{\mu\mu} + H_{\nu\nu} + H_{\lambda\lambda})} J_{\mu\nu\lambda}$$

Remark 1. The S, the three H's, and the J above respectively correspond to 's group of red nodes, three thin edges, and green node. Each diagram encodes many Taylor terms, and the fact that we may evaluate each diagram as a whole is an advantage of our calculational framework. Intuitively, each diagram gives the net effect of a certain combination of gradients (G), noise (C, S, \cdots) and curvature (H, J, \cdots) . After developing our theory more precisely, we will return to these intuitive interpretations.

2. Background and Notation

2.1. Loss landscape

We henceforth fix a space \mathcal{M} of weights on which a loss function $l: \mathcal{M} \to \mathbb{R}$ is defined. SGD operates on unbiased estimates of l drawn from some fixed probability distribution \mathcal{D} . We thus denote by $(l_n: 0 \le n < N)$ an i.i.d. training sequence of such estimates. We will refer both to n and to l_n as training points. We likewise write l_x for a sample from \mathcal{D} independent from the training sequence. We assume the regularity conditions listed in Appendix FILL IN, e.g. that l, l_x are analytic and that all moments exist.

E.g.: our theory applies to tanh networks with cross entropy loss on bounded data — and with arbitrary weight sharing, skip connections, soft attention, dropout, and weight decay.

2.2. Tensor conventions

Adopting Einstein's convention, we implicitly sum repeated Greek indices: if A_{μ} , B^{μ} are the coefficients of a covector A and a vector B^* , indexed by basis elements μ , then $A_{\mu}B^{\mu} \triangleq \sum_{\mu} A_{\mu} \cdot B^{\mu}$. To expedite dimensional analysis, we regard the learning rate as an inverse metric $\eta^{\mu\nu}$ that converts a gradient covector into a displacement vector (Bonnabel, 2013), and we use η to raise indices: e.g., in $H^{\mu}_{\lambda} \triangleq \eta^{\mu\nu} H_{\nu\lambda}$, η raises one of $H_{\mu\nu}$'s indices. Another example is $C^{\mu}_{\mu} \triangleq \sum_{\mu\nu} \eta^{\mu\nu} \cdot C_{\nu\mu}$. Standard syntactic constraints make manifest which expressions transform naturally.

We say two expressions agree to order η^d when their difference, divided by some homogeneous degree-d polynomial of η , tends to 0 as η shrinks. Their difference is then $\in o(\eta^d)$.

2.3. SGD terminology

SGD decreases the objective l via T steps of discrete-time η -steepest[†] descent on the estimates l_n . We describe SGD in terms of N, T, B, E, M: N counts training points, T counts updates, B counts points per batch, E = TN/B counts epochs, and M = E/B = T/N. Concretely, SGD performs T = NM updates of the form:

$$\theta^{\mu} \longleftrightarrow \theta^{\mu} - \eta^{\mu\nu} \nabla_{\nu} \left(\frac{1}{B} \sum_{n \in \mathcal{B}_{i}} l_{n}(\theta) \right)$$

We write l_t for the loss $\frac{1}{B} \sum_{\mathcal{B}_t} l_n$ on the *t*th batch. no **VANILIA!**

2.4. Diagrams and embeddings

Though a rough, intuitive understanding of concepts such as *diagram* suffices for absorbing this paper's main results, the following definitions may help the reader who wishes to follow our mathematics closely.

Definition 1 (Diagrams). A *diagram* is a finite rooted tree equipped with a partition of its nodes. We draw the tree using thin edges. By convention, we draw each node to the right of its children; the root is thus always rightmost. We draw the partition by connecting the nodes within each part via fuzzy ties. For example, has 2 parts. We insist on using as few fuzzy ties as possible so that, if d counts edges and c counts ties, then d+1-c counts the parts of the partition. There may be multiple ways to draw a single diagram, e.g.

Definition 2 (Spacetime). The *spacetime* associated with an SGD run is the set of pairs (n, t) where the nth datapoint participates in the tth gradient update. Spacetimes thus encode the batch size, training set size, and epoch number.

Definition 3 (Embedding Diagrams into Spacetime). An *embedding* of a diagram D into a spacetime is an assignment of D's non-root nodes to pairs (n, t) such that each node occurs at a time t' strictly after each of its children and such that two nodes occupy the same row n if and only if they inhabit the same part of D's partition.

Definition 4 (A Diagram's Un-resummed Value). The *un-resummed value* of a diagram D is the product of the values of each part p in its partition. The value of a part p with |p| many nodes is the expectation $\mathbb{E}_x\left[(\nabla l_x(\theta))^{|p|}\right]$. The edges of D's tree indicate how to multiply the values of these parts: each edge indicates a contraction. For instance, since the training points are independent:

$$\triangleq \mathbb{E}_{n,n',n''} \left[(\nabla_{\mu} l_n) (\nabla_{\nu} l_n) (\nabla^{\mu} \nabla^{\nu} \nabla_{\lambda} l_{n'}) (\nabla^{\lambda} l_{n''}) \right]$$

^{*} Vectors/covectors are also called column/row vectors.

 $^{^{\}dagger}$ To define "steepness" requires a metric on l 's domain. We regard $\eta^{\mu\nu}$ as an (inverse) metric.

Implicit in the three raised indices are three factors of η . We write value(D) for D's un-resummed value, or D when clear

Definition 5 (An Embedding's Re-summed Value). The *re-summed value* rvalue $_f(D)$ of an embedding f of a diagram D is the same as the un-resummed value of D, save for one change having to do with edges. Consider an edge between two nodes embedded to (n,t) and $(n',t+\Delta t)$. Whereas value(D) has a factor of $\eta^{\mu\nu}$ for this edge, rvalue $_f(D)$ instead has a factor of $((I-\eta H)^{\Delta t-1})^{\mu}_{\lambda}\eta^{\lambda\nu}$. Here, $1 \leq \Delta t$ is the temporal distance between the two nodes' embeddings.

We will often seek *differences*, e.g. between ODE's and SGD's test loss or between a test loss and a train loss. We thus define a compact notation for differences of diagrams:

Definition 6 (Fuzzy Outlines Denote Noise's Net Effect). A diagram drawn with one *fuzzy outline* denotes the difference between the versions with and without fuzzy ties. E.g.:



We define a diagram drawn with more than one fuzzy outline as the fully tied version minus all the versions with fewer fuzzy outlines (these are the Möbius sums of (Rota, 1964)):



Definition 7 (Irreducible Diagrams). A diagram, drawn with fuzzy outlines instead of ties, is *irreducible* when none of its degree-2 non-root nodes participates in fuzzy outlines.

3. Diagram Calculus for SGD

3.1. Recipe for SGD's expected test loss

Our Main Theorem expresses SGD's test loss as a sum over diagram embeddings. Recalling that a diagram with d edges is $O(\eta^d)$, we may read this Theorem as a Taylor series in the learning rate. In practice, we truncate the series to small d, thus focusing on the few-edged diagrams.

Theorem 1 (Test Loss as a Path Integral). For any T: for η sufficiently small, SGD's expected test loss is

$$\sum_{\substack{D \\ irreducible \\ f}} \sum_{\substack{embeddings \\ f}} \frac{1}{\left|\operatorname{Aut}_f(D)\right|} \frac{\operatorname{rvalue}_f(D)}{(-B)^{\left|\operatorname{edges}(D)\right|}}$$

Here, D ranges through irreducible diagrams drawn with fuzzy outlines instead of ties, f ranges through embeddings of D into the SGD's spacetime, and $|\operatorname{Aut}_f(D)|$ counts the graph automorphisms of D that preserve f's assignment of nodes to (n,t) pairs. As a reminder, B is the batch size.

Though the combinatorics of embeddings and graph automorphisms may seem forbidding, our focus on few-edged diagrams will make this counting nearly trivial.

Theorem 2 (Long-Term Behavior at a Local Minimum). When SGD is initialized at a local minimum of test loss, and when $\nabla \nabla l_x$ is bounded below by some positive form that doesn't depend on x, then the dth-order truncation of Theorem 1 converges as T diverges.

Remark 2 (Approximation by integrals). In practice, we approximate sums over embeddings by integrals over times and $(I - \eta H)^t$ by $\exp(-\eta H t)$. This incurs a multiplicative error of $1 + o(\eta)$ that preserves leading order results. So diagrams induce easily evaluated integrals of exponentials.

Remark 3 (Using Un-resummed Values). value(D) is simpler to work with than $\operatorname{rvalue}_f(D)$. Theorem 1 remains true if we replace each $\operatorname{rvalue}_f(D)$ by $\operatorname{value}(D)$, so long as we drop the constraint that D be irreducible and we use diagrams drawn with fuzzy ties instead of fuzzy outlines. However, Theorem 2's convergence guarantee no longer applies, and empirically we find deteriorated predictions.

Remark 4 (Variants). The above gives SGD's expected test loss. What if we seek train instead of test losses? Or net weight displacements instead of losses? Or variances instead of expectations? Theorem 1 and Remark 3 have simple analogues for each of these 2³ possibilities, which we discuss in the appendix.

3.2. Single-Epoch, Singleton-Batch SGD

For SGD with 1 epoch and batch size 1, Theorem 1 then specializes to:

Proposition 1. Single-epoch singleton-batch SGD has expected test loss

$$\sum_{0 \leq d < \infty} \frac{(-1)^d}{d!} \sum_{D} |\operatorname{ords}(D)| \binom{N}{P-1} \frac{d!}{\prod d_p!} \operatorname{value}(D)$$

where D has P parts with sizes d_p . Here, D ranges over d-edged diagrams none of whose parts contains any of its nodes' ancestors, and $|\operatorname{ords}(D)|$ counts the total orderings of D's nodes s.t. children precede parents and parts are contiguous.

So a diagram with d thin edges and c fuzzy ties (hence d+1-c parts), contributes $\Theta\left((\eta T)^d T^{-c}\right)$ to SGD's test loss. Intuitively, ηT measures the physical time of descent and T^{-1} measures the coarseness of time discretization. We thus regard Proposition 1 as a double series in $(\eta T)^d T^{-c}$, where each term isolates the dth order effect of time and the cth order effect of noise. Indeed, c counts fuzzy ties and hence the c=0 terms do not model correlations and hence do not model noise. That is, the c=0 terms give an ODE approximation to SGD. The remaining terms give the corrections due to noise. See Table 2.

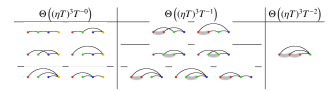


Table 2. **Degree-3 diagrams for** B = M = 1 **SGD's test loss**. The 6 diagrams have (4 + 2) + (2 + 2 + 3) + (1) total orderings relevant to Proposition 1. **Left:** (d, c) = (3, 0). Diagrams for ODE behavior. **Center:** (d, c) = (3, 1). 1st order deviation of SGD away from ODE. **Right:** (d, c) = (3, 2). 2nd order deviation of SGD from ODE with appearance of non-Gaussian statistics.

4. Insights from the Formalism

4.1. SGD descends on a C-smoothed landscape

Integrating rvalue f() over embeddings f, we see:

Corollary 1 (Minima flat w.r.t. C attract SGD). *Initialized* at a test minimum, vanilla SGD's weight moves to order η^2 with a long-time-averaged* expected velocity of

$$v^{\pi} = C_{\mu\nu} \left(F^{-1} \right)_{\rho\lambda}^{\mu\nu} J_{\sigma}^{\rho\lambda} \left(\frac{I - \exp(-T\eta H)}{T\eta H} \eta \right)^{\sigma\pi}$$

per timestep. Here, $F = \eta H \otimes I + I \otimes \eta H$, a 4-valent tensor.

The intuition behind the Corollary is that the diagram contains a subdiagram = CH; by a routine check, this subdiagram is the leading-order loss increase when we convolve the landscape with a C-shaped Gaussian. Since connects the subdiagram to the test measure-

ment via 1 edge, it couples to the linear part of the test loss and hence represents a displacement of weights away from high *CH*. In short, reveals that *SGD descends* on a covariance-smoothed landscape. See Figure 1 (right).

An un-resummed version of this result was first reported by Yaida (2019b); however, for fixed T, the un-resummed result scales with η^3 while Corollary 1 scales with η^2 . The discrepancy occurs, intuitively, because the re-summed analysis accounts for the accumulation of noise from many updates, hence amplifying the contribution of C. Our experiments verify our scaling law.

Unlike Wei & Schwab (2019), we make no assumptions of thermal equilibrium, fast-slow mode separation, or constant covariance. This generality reveals a novel dynamical phenomenon, namely that the velocity field above need not be conservative (see Section 5.4)

4.2. Curvature controls overfitting

Integrating rvalue_f() and rvalue_f() yields:

Corollary 2 (Flat, Sharp Minima Overfit Less). *Initialized* at a test minimum, pure GD's test loss is to order η

$$\frac{1}{2N} C_{\mu\nu} \Big((I - \exp(-\eta T H))^{\otimes 2} \Big)_{\rho\lambda}^{\mu\nu} \Big(H^{-1} \Big)^{\rho\lambda}$$

above the minimum. This vanishes when H does. Likewise, pure GD's generalization gap is to order η :

$$\frac{1}{N} C_{\mu\nu} \left(I - \exp(-\eta T H) \right)_{\lambda}^{\nu} \left(H^{-1} \right)^{\lambda\mu}$$

In contrast to the later-mentioned Takeuchi estimate, this does not diverge as H shrinks.

Corollary 2's generalization gap converges after large T to $C_{\mu\nu}(H^{-1})^{\mu\nu}/N$, also known as Takeuchi's Information Criterion (TIC). In turn, C=H is the Fisher metric in the classical setting of maximum likelihood (ML) estimation (in well-specified models) near the "true" test minimum, so we recover AIC (number of parameters)/N. Unlike AIC, our more general expression is descendably smooth, may be used with MAP or ELBO tasks instead of just ML, and makes no model well-specification assumptions.

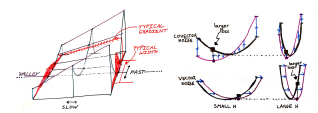


Figure 1. Re-summation reveals novel phenomena. Left: The entropic force mechanism: gradient noise induces a flow toward minima with respect to to the covariance. Though our analysis assumes neither thermal equilibrium nor fast-slow mode separation, we label "fast and slow directions" to ease comparison with Wei & Schwab (2019). Here, red densities denote the spread predicted by a re-summed $C^{\mu\nu}$, and the spatial variation of curvature corresponds to $J_{\mu\nu\lambda}$. Right: Noise structure determines how curvature affects overfitting. Geometrically, for (empirical risk minimization on) a vector-perturbed landscape, small Hessians are favored (top row), while for covector-perturbed landscapes, large Hessians are favored (bottom row). Corollary 2 shows how the implicit regularization of fixed- ηT descent interpolates between the two rows.

4.3. Effects of epochs and of batch size

Corollary 3 (Epoch Number). To order η^2 , one-epoch SGD has $\left(\frac{M-1}{M}\right)\left(\frac{B+1}{B}\right)\left(\frac{N}{2}\right)\left(\nabla_{\mu}C_{\nu}^{\nu}\right)G^{\mu}/2$ less test loss than M-epoch SGD with learning rate η/M .

Analyzing \hookrightarrow , we find that we may cause GD to mimic SGD using any smooth unbiased estimator \hat{C} of C:

^{*} That is, T so large that $C \exp(-\eta KT)$ is negligible. Appendix ?? gives a similar expression for general T.

Corollary 4 (Batch Size). The expected test loss of pure SGD is, to order η^2 , less than that of pure GD by $\frac{M(N-1)}{2} \left(\nabla_{\mu}C_{\nu}^{\nu}\right)G^{\mu}/2$. Moreover, GD on a modified loss $\tilde{l}_n = l_n + \frac{N-1}{4N} \hat{C}_{\nu}^{\nu}(\theta)$ has an expected test loss that agrees with SGD's to second order. We call this method GDC.

4.4. SDE is blind to non-Gaussian noise

Stochastic Differential Equations (SDE: see Liao et al. (2018)) are a popular theoretical approximation to SGD, but SDE and SGD differ in several ways. For instance, the interepoch noise correlations in multi-epoch SGD measurably affect SGD's final test loss (Corollary 3), but SDE assumes uncorrelated gradient updates. Even if we restrict to single-epoch SDE, differences arise due to time discretization and, more interestingly, due to non-gaussian noise.

Corollary 5 (SGD Differs from ODE, SDE). The test loss of single-epoch, singleton-batch SGD deviates from ODE by $\frac{T}{2}$ $C_{\mu\nu}H^{\mu\nu} + o(\eta^2)$. Its leading order deviation from SDE is due to non-Gaussian noise: -(T/6) $+ o(\eta^3) = -(T/6)S_{\mu\nu\lambda}J^{\mu\nu\lambda} + o(\eta^3)$.*

For finite N, this Corollary separates SDE from SGD. Conversely, as $N \to \infty$ with ηN fixed and C scaling with \sqrt{N} , SGD converges to SDE, but generalization and optimization respectively become trivial and computationally intractable.

5. Experiments

We focus on experiments whose rejection of the null hypothesis (and hence support of our theory) is so drastic as to be visually obvious. For example, in Figure 4, (Chaudhari & Soatto, 2018) predicts a velocity of 0 while we predict a velocity of $\eta^2/6$. Throughout, I bars and + marks denote a 95% confidence interval based on the standard error of the mean, in the vertical or vertical-and-horizontal directions, respectively. See Appendix ?? for experimental procedure including architectures and sample size.

5.1. Basic predictions

We test Theorem 3 on smooth convnets on CIFAR-10 and Fashion-MNIST. Our order η^3 predictions agree with experiment up to $\eta T \approx 10^0$ (Figure 2, left). Likewise, Corollary 3 correctly predicts the effect of multi-epoch training (Appendix ??) for $\eta T \approx 10^{-1/2}$. These tests verify that our proofs hide no mistakes of proportionality or sign.

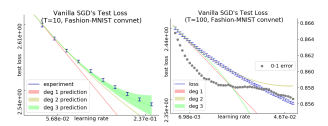


Figure 2. Perturbation models SGD for small ηT . Test loss vs learning rate on a Fashion-MNIST convnet, with un-re-summed predictions. Left: For the instance shown and all 11 other initializations unshown, our degree-3 prediction agrees with experiment through $\eta T \approx 10^0$, which corresponds to a decrease in 0-1 error of $\approx 10^{-3}$. Right: For larger ηT , our predictions can break down. Here, the order-3 prediction holds until the 0-1 error improves by $5 \cdot 10^{-3}$. Beyond this, close agreement with experiment is coincidental.

5.2. Emulating small batches with large ones

By Corollary 4, SGD avoids high-C regions more than GD We artificially correct GD accordingly, yielding an optimizer, GDC, that indeed behaves like SGD on a range of landscapes (Figure 3 (left)). It may be important to emulate SGD's avoidance of high-C regions because we C controls the rate at which each new update increases the generalization gap[†] (Figure 3 (right)).

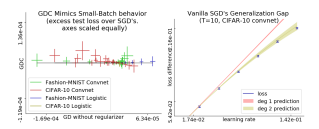


Figure 3. C controls generalization and distinguishes GD from SGD. Left: With equal-scaled axes, this plot shows that GDC matches SGD (small vertical variation) better than GD matches SGD (large horizontal variation) in test loss, for a variety of learning rates ($\approx 10^{-3} - 10^{-1}$) and initializations (zero and several Xavier-Glorot trials) on logistic and architectures for image classification. Here, T = 10. **Right**: CIFAR-10 generalization gaps. For the instance shown and all 11 other initializations unshown, the degree-2 prediction agrees with experiment through $\eta T \approx 5 \cdot 10^{-1}$.

The connection between generalization and covariance was first established by Roberts (2018) in the case T=2 and to order η^2 . In fact, that work conjectures the possibility of emulating GD with SGD. This sub-section extends that work by generalizing to arbitrary T and arbitrary orders η^d , and by concretely defining GDC.

In these experiments, we used a covariance estimator $\hat{C} \propto \nabla l_x (\nabla l_x - \nabla l_y)$ evaluated on two batches x, y that evenly

^{*} This expression differs from the more exact expression of Example 1 because here we use Remark 3's substitution. One may check that the two expressions agree to leading order.

[†]Reminder: for us, generalization gap is test minus train loss.

partition the train set. For typical architectures, we may compute $\nabla \hat{C}$ with the same memory and time as the usual gradient ∇l_t , up to a multiplicative constant.

5.3. Comparison to continuous time

Consider fitting a centered normal $\mathcal{N}(0, \sigma^2)$ to some centered standard normal data. We parameterize the landscape by $h = \log(\sigma^2)$ so that the Fisher information matches the standard dot product (Amari, 1998). The gradient at sample x and weight σ is then $g_x(h) = (1 - x^2 \exp(-h))/2$. Since $x \sim \mathcal{N}(0, 1), g_x(h)$ will be affinely related to a chi-squared, and in particular non-Gaussian. At h = 0, the expected gradient vanishes, and the test loss of vanilla SGD only involves diagrams with no singleton leaves; to third order, it is $\cdot + \frac{T}{2} \longrightarrow + \binom{T}{2} \longrightarrow + \frac{T}{6} \longrightarrow$ In particular, the $\binom{T}{2}$ differs from $T^2/2$ and hence contributes to the timediscretization error of SDE as an approximation for SDE. Moreover, non-Gaussian noise contributes via to that error. Appendix ?? shows that SDE and one-epoch SGD indeed differ. For multi-epoch SGD, the effect of overfitting to finite training data further separates SDE and SGD.

5.4. Nonconservative entropic force

To test Corollary 1's predicted force, we construct a counterintuitive loss landscape wherein, for arbitrarily small learning rates, SGD steadily increases the weight's z component despite 0 test gradient in that direction. Our mechanism differs from that discovered by Chaudhari & Soatto (2018). Specifically, because in this landscape the force is η -perpendicular to the image of ηC , that work predicts an entropic force of 0. This disagreement in predictions is possible because our analysis does not make any assumptions of equilibrium, conservatism, or continuous time.

So, even in a valley of global minima, SGD will move away from minima whose Hessian aligns with the current covariance. However, by the time it moves, the new covariance might differ from the old one, and SGD will be repelled by different Hessians than before. Setting the covariance to lag the Hessian by a phase, we construct a landscape in which this entropic force dominates. This "linear screw" landscape has 3-dimensional $w \in \mathbb{R}^3$ (initialized to 0) and 1-dimensional $x \sim \mathcal{N}(0, 1)$:

$$l_x(w) \triangleq \frac{1}{2}H(z)(w,w) + x \cdot S(z)(w)$$

Here, $H(z)(w, w) = w_x^2 + w_y^2 + (\cos(z)w_x + \sin(z)w_y)^2$ and $S(z)(w) = \cos(z - \pi/4)w_x + \sin(z - \pi/4)w_y$. There is a valley of global minima defined by x = y = 0. If SGD is initialized there, then to leading order in η and for large T, the resummed theory predicts a z-speed of $\eta^2/6$ per timestep. Our

re-summed predictions agree for with experiment for ηT so large that the weight moves about 5 times the landscape's natural length scale of 2π (Figure 4, left).

It is routine to check that, by stitching together copies of this example, we may cause SGD to travel along paths that are closed loops or unbounded curves. We may even add a small linear component so that SGD steadily climbs uphill.

5.5. Sharp and flat minima both overfit less

Prior work has varyingly found that *sharp* minima overfit less (after all, l^2 regularization increases curvature) or that *flat* minima overfit less (after all, flat minima are more robust to small displacements in weight space). Corollary 2 reconciles these competing intuitions by showing how the relationship of generalization and curvature depends on the learning task's noise structure and how the metric η^{-1} mediates this distinction (Figure 1, right).

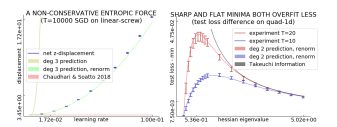


Figure 4. Re-summed predictions excel even for large ηT for SGD near minima. Left: On Linear Screw, the persistent entropic force pushes the weight through a valley of global minima not at a $T^{1/2}$ diffusive rate but at a directional T^1 rate. Since Hessians and covariances are bounded throughout the valley and the effect appears for all sufficiently small η , the effect is not a pathological artifact of well-chosen learning rate or divergent covariance noise. The net displacement of $\approx 10^{1.5}$ well exceeds the z-period of 2π . Right: For Mean Estimation with fixed covariance and a range of Hessians, initialized at the true minimum, the test losses after fixed- ηT optimization are smallest for very small and very large curvatures. This evidences our prediction that both sharp and flat minima overfit less and that TIC's singularity is suppressed.

Because the TIC estimates a smooth hypothesis class's generalization gap, it is tempting to use it as an additive regularization term. However, since the TIC is singular where the Hessian is singular, it gives insensible results for overparameterized models. Indeed, Dixon & Ward (2018) report numerical difficulties requiring an arbitrary cutoff.

Fortunately, by Corollary 2, the implicit regularization of gradient descent both demands and enables a singularity-removing correction to the TIC (Figure 4, right). The resulting *Stabilized TIC* (STIC) uses the metric η^{-1} implicit in gradient descent to threshold flat from sharp minima*. It

^{*} The notion of H's width depends on a choice of metric. Prior work chooses this metric arbitrarily. We show that choosing η^{-1} is

thus offers a principled method for optimizer-aware model selection easily compatible with automatic differentiation systems. By descending on STIC, we may tune smooth hyperparameters such as l_2 coefficients. Experiments on an artificial Mean Estimation problem (task in Appendix ??, plot in Appendix ??) recommend STIC for model selection when H is negligible compared to C/N as in the noisy, small-N regime. Because diagonalization typically takes time cubic in dimension, exact STIC regularization is most useful for small models on noisy and limited data.

6. Related Work

It was Kiefer & Wolfowitz (1952) who, in uniting gradient descent (Cauchy, 1847) with stochastic approximation (Robbins & Monro, 1951), invented SGD. Since the development of back-propagation for efficient differentiation (Werbos, 1974), SGD has been used to train connectionist models including neural networks (Bottou, 1991), in recent years to remarkable success (LeCun et al., 2015).

Several lines of work quantify the overfitting of SGD-trained networks (Neyshabur et al., 2017a). For instance, Bartlett et al. (2017) controls the Rademacher complexity of deep hypothesis classes, leading to generalization bounds that are optimizer-agnostic. However, since SGD-trained networks generalize despite their seeming ability to shatter large sets (Zhang et al., 2017), one infers that generalization arises from the aptness to data of not only architecture but also optimization (Neyshabur et al., 2017b). Others have focused on the implicit regularization of SGD itself, for instance by modeling descent via stochastic differential equations (SDEs) (e.g. Chaudhari & Soatto (2018)). However, per Yaida (2019a), such continuous-time analyses cannot treat covariance correctly, and so they err when interpreting results about SDEs as results about SGD for finite trainsets.

Following Roberts (2018), we avoid continuous-time approximations and Taylor-expand around $\eta=0$. We hence extend that work beyond leading order and beyond 2 time steps, allowing us to compare, for instance, the expected test losses of multi-epoch and one-epoch SGD. We also quantify the overfitting effects of batch size, whence we propose a regularizer that causes large-batch GD to emulate small-batch SGD. In doing so, we establish a precise version of the relationship — between covariance, batch size, and generalization — conjectured by Jastrzebski et al. (2018).

While we make rigorous, architecture-agnostic predictions of learning curves, these predictions become vacuous for large η . Other discrete-time dynamical analyses allow large η by treating deep generalization phenomenologically, whether by fitting to an empirically-determined correlate of Rademacher bounds (Liao et al., 2018), by exhibiting

a natural choice because it leads to a prediction of the gen. gap.

generalization of local minima *flat* with respect to the standard metric (see Hoffer et al. (2017), Keskar et al. (2017), Wang et al. (2018)), or by exhibiting generalization of local minima *sharp* with respect to the standard metric (see Stein (1956), Dinh et al. (2017), Wu et al. (2018)). Our work reconciles those seemingly clashing claims.

Others have perturbatively analyzed descent: Dyer & Gur-Ari (2019) perturb in inverse network width, employing Feynman-'t Hooft diagrams to correct the Gaussian Process approximation for a specific class of deep networks. Meanwhile, (Chaudhari & Soatto, 2018) and Li et al. (2017) perturb in learning rate to second order by approximating noise between updates as Gaussian and uncorrelated. In neglecting correlations and heavy tails, that work neither extends to higher orders not describes SGD's generalization behavior. By contrast, we use Feynman-Penrose diagrams to compute test and train losses to arbitrary order in learning rate. Our method accounts for non-Gaussian and correlated noise and applies to any sufficiently smooth architecture. For example, since our work does not rely on informationgeometric relationships between C and H (Amari, 1998)*, it applies to inexact-likelihood landscapes such as VAEs'.

7. Conclusion

We present a diagram-based method for studying stochastic optimization on short timescales. Theorem 1 justifies long-time predictions of SGD's dynamics near minima. Our theory answers the following questions.

Which Minima Overfit Less? By analyzing \longrightarrow , we find that flat and sharp minima both overfit less than minima of curvature comparable to $(\eta T)^{-1}$. Flat minima are robust to vector-valued noise, sharp minima are robust to covector-valued noise, and medium minima attain the worst of both worlds. We thus reconcile prior intuitions that sharp (Keskar et al., 2017; Wang et al., 2018) or flat (Dinh et al., 2017; Wu et al., 2018) minima overfit worse. These considerations lead us to a smooth generalization of AIC enabling hyperparameter tuning by gradient descent.

Which Minima Does SGD Prefer? Analyzing , we refine Wei & Schwab (2019) to nonconstant, nonisotropic covariance to reveal that SGD descends on a loss landscape smoothed by the *current* covariance C. In particular, SGD moves toward regions flat with respect to C. As C evolves, the smoothing mask and thus the effective landscape evolves. These dynamics are generically nonconservative. In contrast to Chaudhari & Soatto (2018)'s SDE approximation, SGD does not generically converge to a limit cycle.

Can GD Emulate SGD? By analyzing , we prove the

^{*} Disagreement of *C* and *H* is typical in modern learning (Roux et al., 2012; Kunstner et al., 2019).

conjecture of Roberts (2018), that large-batch GD can be made to emulate small-batch SGD. We show how to do this by adding a multiple of an unbiased covariance estimator to the descent objective. This emulation is significant because, while small batch sizes can lead to better generalization (Bottou, 1991), modern infrastructure increasingly rewards large batch sizes (Goyal et al., 2018).

7.1. Consequences

Our analysis of which minima (among a valley of minima) SGD prefers — and our characterization of when SGD overfits less in certain minima — together offer insight into SGD's success in training over-parameterized models.

Our results may also help to analyze fine-tuning procedures such as the meta-learning of MAML (Finn et al., 2017). Indeed, those methods seek models initialized near minima and tunable to new data through a small number of updates, a setting matched to our theory's assumptions.

Since our predictions depend only on loss data near initialization, they break down after the weight moves far from initialization. Our theory thus best applies to small-movement contexts, whether for long times (large ηT) near an isolated minimum or for short times (small ηT) in general.

Yet, even short-time predictions show how curvature and noise — and not just averaged gradients — repel or attract SGD's current weight. For example, we proved that SGD in a valley moves toward regions flat with respect to the current covariance C. Much as meteorologists understand how warm and cold fronts interact despite the intractability of long-term weather forecasting, we quantify the counterintuitive dynamics governing SGD's short-time behavior.* Our results enhance the intuitions of practitioners — e.g. that "SGD descends on the train loss" — by summarizing the effect of noise in closed-form dynamical laws valid in each short-term interval of SGD's trajectory.

7.2. Questions

The diagram method opens the door to exploration of Lagrangian formalisms and curved backgrounds[†]:

Question 1. Does some least-action principle govern SGD; if not, what is an essential obstacle to this characterization?

Lagrange's least-action formalism intimately intertwines with the diagrams of physics. Together, they afford a modular framework for introducing new interactions as new terms or diagram nodes. In fact, we find that some *higher-order* methods — such as the Hessian-based update

 $\theta \iff \theta - (\eta^{-1} + \lambda \nabla \nabla l_t(\theta))^{-1} \nabla l_t(\theta)$ parameterized by small η, λ — admit diagrammatic analysis when we represent the λ term as a second type of diagram node. Though diagrams suffice for computation, it is Lagrangians that most deeply illuminate scaling and conservation laws.

Conjecture 1 (Riemann Curvature Regularizes). *For small* η, SGD's gen. gap decreases as sectional curvature grows.

Though our work so far assumes a flat metric $\eta^{\mu\nu}$, it generalizes to curved weight spaces[‡]. Curvature finds concrete application in the *learning on manifolds* paradigm of Absil et al. (2007); Zhang et al. (2016), notably specialized to Amari (1998)'s *natural gradient descent* and Nickel & Kiela (2017)'s *hyperbolic embeddings*. We are optimistic our formalism may resolve conjectures such as above.

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^{*} Because our analysis holds for any initialization, one may imagine SGD's coarse-grained trajectory as an integral curve of the vector field given by our theory.

[†] Landau and Lifshitz introduce these concepts (1960; 1951).

^{*} One may represent the affine connection as a node, thus giving rise to non-tensorial and hence gauge-dependent diagrams.

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Organization of Appendices

These three appendices respectively serve three functions:

- to explain how to calculate using diagrams;
- to prove our theorems, corollaries, and claims; and
- to specify our experimental methods and results.

A. How to Calculate Expected Test Losses Using Diagrams

Our work introduces a novel technique for calculating the expected learning curves of SGD in terms of statistics of the loss landscape near initialization. Here, we explain this technique. There are **four steps** to computing the expected test loss after a specific number of gradient updates:

- Based on the chosen optimization hyperparameters (namely, batch size, training set size, and number of epochs): draw the spacetime grid that encodes these hyperparameters.
- Based on our desired level of precision, draw all the relevant embeddings of diagrams into the spacetime.
- Evaluate each diagram embedding.
- Sum the embeddings' values to obtain the quantity of interest as a function of the learning rate.

After presenting a small, complete example calculation that follows these four steps, we explain how to perform each of these steps in its own sub-section. We then discuss how diagrams often offer intuition as well as calculational help. Though we focus on the computation of expected test losses, we explain how a small change in the above four steps allows for the computation also of variances (instead of expectations) and of train losses (instead of test losses). We conclude by comparing direct calculation based on our Key Lemma to the diagram method; we point out when and why diagrams streamline computation.

A.1. An example calculation

Let's compute the expected test loss of batchsize-1 SGD on N training points after E epochs. We'll do this calculation to order η^2 , meaning that our answer will be a function of the learning rate η and that its error will shrink faster than quadratically as η becomes small.

First, we identify the relevant spacetime. A spacetime is a set of cells indicating which training points are used in which gradient update. Since our problem has NE many updates, each on a batch of size 1, there will be $NE \times 1$ many cells in the relevant spacetime. We arrange these cells in a

grid whose vertical axis indexes training points and whose horizontal axis indexes training times: FILL IN

Next, we identify the relevant diagram embeddings. The benefit of drawing this diagram.

Then, we evaluate each diagram embedding. The benefit of drawing this diagram.

Finally, we sum the embeddings' values to arrive at an answer. The benefit of drawing this diagram.

- A.2. How to identify the relevant space-time
- A.3. How to identify the relevant diagram embeddings
- A.4. How to evaluate each embedding
- A.5. How to sum the embeddings' values
- A.6. Interpreting diagrams to build intuition
- A.7. How to solve variant problems
- A.8. Do diagrams streamline computation?
- **B.** Assumptions and Proofs
- **B.1.** Setup and assumptions of our theory
- **B.2. Proof of the Key Lemma**
- **B.3. Proof of Theorem 1**
- **B.4.** On Mobius inversion for resummed diagrams
- **B.5. Proof of Theorem 2**
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- C. Exerimental Methods and Results
- C.1. What loss landscapes did we use?
- C.2. Implementing optimizers
- C.3. Evaluating diagrams on a given landscape
- C.4. Software frameworks and hardware
- C.5. Additional figures

C.6. Informal overview of the perturbative approach

Consider running SGD on N training points for T steps with learning rate η , starting at a weight θ_0 . Our method expresses the expectation (over randomly sampled training sets) of quantities such as the final weight (or test or train loss) as a sum of diagrams, where each diagram evaluates to a statistic of the loss landscape at initialization. Diagrams with e edges contribute only $O(\eta^e)$ to the quantities of interest, so for small η we sum only the few-edged diagrams and incur

an $o(\eta^e)$ error term.

The rule for evaluating diagrams is that each degree-d node evaluates to the dth derivative of the test loss l at θ_0 . The edges indicate the order in which those derivatives are multiplied. Most simply, $\cdot = l(\theta_0)$, a 0th derivative. The diagram \cdot evaluates to the dot product ηGG , where $G = \nabla l$ is the gradient of the expected loss l, evaluated at θ_0 . Likewise, $\cdot = \eta^2 GHG$, where $H = \nabla \nabla l$ is the hessian. And $\cdot = \eta^3 GGJG$, where $J = \nabla \nabla \nabla l$ is l's third derivative.

A diagram tells us about the loss landscape but not about

SGD parameters such as T or inter-epoch shuffling. We summarize those parameters as sets of pairs (n, t), one for each participation of the nth datapoint in the tth update. Full-batch GD will have NT many pairs, for instance, while singeleton-batch SGD will have T many pairs.

Each of a diagram's nodes abstractly represents an event at such a pair, and we may *embed* a diagram by assigning to each node a concrete pair (n, t). We will intuitively interpret an embedded edge from (n, t) to (n', t') (from left to right) as depicting information flow from training point n at time t to training point n' at time t'. Thus, we permit only embeddings whose edges have t < t'. The rightmost node represents measurement at test time, so we do not assign a pair (n, t) to it; it does not participate in t < t' constraints.

Theorem (Theorem 3, Informal). Fix SGD parameters, namely N, T, a batch size B, and a deterministic routine to sample each batch from a train set. Sum the diagrams with at most d edges, where a diagram with e edges and c many embeddings is weighted by $c/(-B)^e$. This sum agrees with SGD's expected final test loss to order $o(\eta^d)$.

Example 2. What is SGD's expected test loss to order η^1 ? Two diagrams have ≤ 1 edges: $\cdot = l(\theta_0)$ and $\frown = \eta GG$. For SGD with batchsize 1, \frown has T many embeddings, since its rightmost node must represent the test measurement and its other node can represent any of T many (n, t) pairs. By the Theorem, the answer is $l(\theta_0) - T \cdot \eta GG + o(\eta^1)$.

Example 2 is well-known (e.g. Nesterov (2004)). Indeed, it quantifies the intuition that, in each of T steps, SGD moves the weight by ηG and hence decreases the loss by ηGG . The expression is exact for a noiseless linear landscape, but, because it fails to model how gradients depend on the current weight (curvature) or on the current training point (noise), it is typically an approximation. Diagrams beyond Γ correct the expression by modeling curvature and noise.

Like Example 2, our predictions depend only on loss data near θ_0 and hence break down after the weight moves far from initialization. Our theory thus best applies to small-movement contexts, whether for long times near a minimum or for short times in general. For instance, we analyze SGD overfitting near a minimum (Corollary 2). Invoking

Theorem 1, we analyze how curvature and noise — and not just gradients — repel or attract the evolving weight (Corollary 1).

C.7. Embeddings into spacetime

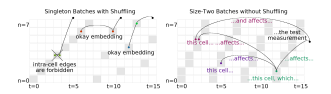


Figure 5. Diagrams in Spacetime Depict SGD's Subprocesses. Two spacetimes with N=8, T=16. Left: Batchsize B=1 with inter-epoch shuffling. Embeddings, legal and illegal, of , and , and . Right: Batchsize B=2 without inter-epoch shuffling. Interpretation of an order η^4 diagram embedding.

To visualize embeddings, we draw (n,t) pairs as shaded cells in an $N \times T$ grid. A diagram embedding is then an assignment of nodes to shaded cells. The t < t' constraint then forbids intra-cell edges (Figure 5 left), and we may interpret each edge as an effect of the past on the future (right). We call an SGD run's set of shaded cells its *spacetime*.

C.8. A first look at curvature and noise

Intuitively, our diagrams' edges depict higher derivatives and hence the test loss l's curvature. However, to study noise and generalization, we need to represent how different datapoints n induce different loss functions l_n . The test loss l is then the expectated value of l_n . We depict correlations and hence noise by a new structure: fuzzy "ties". For example, and are two valid and distinct diagrams. Fuzzy ties determine which derivatives occur within the same expectation, so we have

$$\triangleq \eta^2 \mathbb{E} \left[\nabla l_n \right] \mathbb{E} \left[\nabla \nabla l_n \right] \mathbb{E} \left[\nabla l_n \right] = \eta^2 GHG = \eta^2 \frac{\nabla (GG)}{2} G$$

and, writing C for the covariance of gradients,

$$\triangleq \eta^2 \mathbb{E} \left[\nabla l_n \nabla \nabla l_n \right] \mathbb{E} \left[\nabla l_n \right] = \eta^2 \frac{\nabla \left(GG + C \right)}{2} G$$

The rule is that two nodes connected by a fuzzy tie occur in the same expectation brackets. Since fuzzy ties depict correlations, we demand that each embedding of a diagram sends any two nodes that are connected by a fuzzy tie to pairs (n, t), (n, t') that share a training point index n.

Example 3. When N = T, then singleton-batch SGD permits permits no concretizations of , since the edge constraint t < t' conflicts with the tie constraint n = n' when, as in this case, the permitted (n, t) pairs comprise a bijection between ns and ts.

Example 4. By constrast, when N = T, full-batch GD permits $N\binom{N}{2}$ many concretizations of , since all NT possible (n,t) pairs occur. Those concretizations (n,t),(n,t') have as close analogues the concretizations (n,t),(n',t') in Example 3's setting of the tie-less diagram .

Comparing the two examples above reveals a difference between batchsize-1 and batchsize-N descent for N = T: by the Main Theorem, the latter incurs an additional test loss

$$\frac{c}{(-B)^e}\Big(\longrightarrow - \longrightarrow \Big) = \frac{algebra}{\cdots} = \frac{\eta^2(N-1)}{4}G\nabla C$$

It turns out that \longrightarrow is the only 2-edged diagram whose concretizations in SGD and GD differ. Thus, this test loss difference between SGD and GD is correct to order η^2 .

The above generalizes Roberts (2018)'s T=2 result, proved without diagrams, to arbitrary T. In principle, one could avoid diagrams completely by direct use of our Key Lemma (stated in the Appendix). However, as shown in the Appendix, counting embeddings of diagrams streamlines calculation, yielding shorter and more interpretable arguments compared to direct calculation.

Definition 8 (Un-resummed Value of a Diagram). In the context of a loss landscape and an initial weight θ_0 , a diagram evaluates to the expectation (over all i.i.d. of datapoints to parts) of a product of derivatives, one dth derivative $\nabla^d l(\theta_0)$ for each degree-d node. Each edge denotes a contraction of its two nodes by the inverse metric η . For example,

$$\triangleq \mathbb{E}_{n,n'}\left[(\nabla_{\mu}l_n)(\nabla^{\mu}l_{n'}) \right] (\theta_0)$$

$$\triangleq \mathbb{E}_{n,n',n''}\left[(\nabla_{\mu}l_n)(\nabla_{\nu}l_n)(\nabla^{\mu}\nabla^{\nu}\nabla_{\lambda}l_{n'})(\nabla^{\lambda}l_{n''}) \right] (\theta_0)$$

We write value(D) for a diagram D's value, or D when clear. **Definition 9** (Fuzzy Outlines Denote Noise's Net Effect). We may join any two parts p, \tilde{p} of a diagram D to obtain a new diagram $D_{p\tilde{p}}$. For instance, (\bigcirc)_{red blue} \triangleq \bigcirc . Since fuzzy ties denote correlation and noise, differences such as $D_{p\tilde{p}} - D$ quantify noise's net effect. So, for convenience, we define a diagram with fuzzy *outlines* as the difference between its fuzzy tied and untied versions, e.g.:

$$\triangleq$$
 (\longrightarrow)_{green blue} - \longrightarrow = \longrightarrow -

Definition 10 (Embedding-Sensitive Values). Let $\operatorname{rvalue}_f(D)$ be the expected value of D's corresponding tensor expression, where instead of using η to contract two tensors embedded to times $t, t + \Delta t$, we use $K(\Delta t) = (I - \eta H)^{\Delta t - 1} \eta$. Actually, it will be most convenient to let rvalues represent a *difference* from the noiseless case. For example, to compute rvalue(\frown), we will replace η by $K(\Delta t)$ in \frown) instead of in \frown). This

way, each diagram represents a net effect of noise. For the small diagrams we consider, we obtain rvalues by replacing fuzzy ties by fuzzy outlines; larger diagrams present complications addressed in Appendix ??.

Theorem 3 (Test Loss as a Path Integral). For all T: for η sufficiently small, SGD's expected test loss is

$$\sum_{D} \sum_{embeddings\ f} \frac{1}{\left| \operatorname{Aut}_{f}(D) \right|} \frac{\operatorname{value}(D)}{(-B)^{\left| \operatorname{edges}(D) \right|}}$$

Here, D is a diagram whose root r does not participate in any fuzzy edge, f is an embedding of D into spacetime, and $|\operatorname{Aut}_f(D)|$ counts the graph-automorphisms of D that preserve f's assignment of nodes to cells. If we replace D by $\left(-\sum_{p\in\operatorname{parts}(D)}(D_{rp}-D)/N\right)$, where r is D's root, we obtain the expected generalization gap (test minus train loss).

By Proposition 1, a diagram with d thin edges and f fuzzy ties (hence d+1-c parts), contributes $\Theta\left((\eta T)^d T^{-c}\right)$ to vanilla SGD's test loss.

Intuitively, ηT measures the physical time of descent and T^{-1} measures the coarseness of time discretization. We thus obtain a double series in $(\eta T)^d T^{-c}$; the c=0 terms correspond to a noiseless, discretization-agnostic (hence ODE) approximation to SGD, the the remaining terms model time-discretization and noise. See Table 2.

C.9. Exploiting curvature

Intuitively, the order- η^d truncation of Theorem 3's series depends on simple loss statistics near initialization, so it will fail when ηT is large enough for the weight to drift far from initialization. An especially interesting case where weights do not drift far is the case of SGD dynamics near an isolated minimum. A generic minimum is characterized among critical points by its curvature, so we analyze the case where H is positive. In doing so, we follow prior work that uses lower bounds on the loss landscape's curvature to restrict the hypothesis space to a small basin near a minimum and thus sharpen analyses of optimization and generalization (Bartlett et al., 2005).

We will incorporate the positive-H assumption into our theory via "re-summation" so that our re-summed order- η^d predictions near isolated minima will remain finite for fixed ηT and arbitrary T. More concretely, whenever we compute a diagram, we will also compute the unboundedly many cousins of that diagram that arise by inserting degree-2 nodes onto thin edges. We will sum these diagrams' contributions to Theorem 3's series, arriving at a closed form expression. Theorem 1 establishes the correctness of this approach. Thus, by thoroughly incorporating curvature information, re-summation will help us reason about long-term equilibrium near an isolated minimum and short-term drifts within a valley of minima.

To illustrate the idea, consider this class of topologically related diagrams: , , , , ..., Intuitively, these diagrams all represent the effect of the leftmost node on the rightmost node, with some number of degree-2 nodes mediating. Since degree-2 nodes evaluate to hessians H, we regard these diagrams as versions of modulated by curvature. Each of the above diagrams has some number of embeddings into spacetime. Here (but not in Theorem 1), we will for simplicity consider embeddings into vanilla SGD's spacetime. Moreover, let us consider only embeddings that map the start and end nodes to fixed cells (n_0, t_0) and (n_+, t_+) separated $\Delta t = t_+ - t_0$ timesteps. We will also temporarily relax the constraint on embeddings by allowing each of the middle nodes to occupy any row — and in particular the same row as other nodes.* Then, a routine invocation of the Binomial Theorem shows that these embeddings together contribute the following to Theorem 3's series:

$$-G(I-\eta H)^{\Delta t-1}\eta G$$

For comparison, the analogous embeddings (in this case, there is only one) of the smallest diagram sum to

$$-GnG$$

which matches like the overall sum if we replace η by an "effective learning rate"

$$K(\Delta t) \triangleq (I - \eta H)^{\Delta t - 1} \eta$$

In the proof of Theorem 1, we see that this generalizes: in order to sum over a class of related diagrams' embeddings, we may sum over embeddings of the smallest diagram in that class, then replace each η corresponding to a duration- Δt edge by $K(\Delta t)$.

Example 5. The family , , , , ... includes variants of where we insert new nodes along 's two thin edges. The diagram evaluates to

$$\frac{1}{2}(\nabla_{\mu}C_{\nu\lambda})\eta^{\nu\lambda}\eta^{\mu\rho}G_{\rho}$$

So the overall family evaluates to

$$\frac{1}{2}(\nabla_{\mu}C_{\nu\lambda})K(\Delta t)^{\nu\lambda}K(\Delta t)^{\mu\rho}G_{\rho}$$

^{*} Because we here allow more embeddings than occur in Theorem 3, we are overcounting. It turns out that our use of differences as mentioned in Definition 10 leads to a telescoping cancellation that exactly counters this overcounting. We offer mathematical details in Appendix E.4 and the proof of Theorem 1. For now, we note that Theorem 1 will abstract away the middle nodes altogether, meaning that the problem of overcounting is relevant only to proof details.