EFFICIENT GRADIENT-BASED ALGORITHM FOR TRAIN ING DEEP LEARNING MODELS WITH MANY NONLIN EAR ACTIVATIONS

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ABSTRACT

This research paper presents a novel algorithm for training deep neural networks with many nonlinear layers (e.g., 30). The method is based on backpropagation of an approximated gradient, averaged over the range of a weight update. Unlike the gradient, the average gradient of a loss function is proven within this research to provide more accurate information on the change in loss caused by the associated parameter update of a model. Therefore, it may be utilized to improve learning. In our implementation, the efficiently approximated average gradient is paired with RMSProp and compared to the typical gradient-based approach. For the tested deep model with numerous stacked fully-connected layers featuring nonlinear activations on MNIST and Fashion MNIST, the presented algorithm: (a) generalizes better, at least in a reasonable epoch count, (b) in the case of optimal implementation, learning would require less computation time than the gradient-based RMSProp, with the memory requirement of the Adam optimizer, (c) performs well on a broader range of learning rates, therefore it may bring time and energy savings from reduced hyperparameter searches, (d) improves sample efficiency about three times according to median training losses. On the other hand, for a deep sequential convolutional model trained on the IMDB dataset, sample efficiency is improved by about 55%. However, in the case of the tested shallow model, the method performs approximately the same as the gradient-based RMSProp in terms of both training and test loss. The source code is provided at [...].

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1 INTRODUCTION

1.1 AVERAGE GRADIENT

In this research, we focus on solving deep learning problems by calculating the precise influence 037 of potential updates on the loss for each model parameter separately. Our goal is to obtain more precise information about the influence of each parameter on loss than what the gradient provides. Each potential update of a model parameter influences the locally-optimal direction of other model 040 parameters during the same weight update, highlighting the complexity of the problem. The average 041 gradient (defined in Appendix A), unlike the gradient, stores the accurate contribution of each model 042 parameter to the loss delta related to a given weight update (Fig. 1; Eq. 14). Therefore, the average 043 gradient can be utilized to efficiently minimize the loss. In this research, we propose a very fast 044 algorithm to approximate the average gradient. We prove its approximation accuracy, validate the proof using our handcrafted metric to compare batch-loss minimization efficiency between methods, and test our method on various domains and models. Our algorithm in its current form primarily 046 targets very deep models with many nonlinear layers. 047

Due to the tendency to increase model depth along with its width (Tan & Le, 2019) and the popularity of certain nonlinear activation functions, our approach may offer insights for future improvements in practical deep learning. Our primary target is to significantly improve sample efficiency, even at the cost of a moderate increase in computation time, which is essential for practical applications in fields like deep reinforcement learning or reinforcement learning from human feedback (Kirk et al., 2023).
These methods have been used in popular chatbots, such as OpenAI's ChatGPT (OpenAI, 2023) and Anthropic's Claude (Kirk et al., 2023).





(a) *Example 1*. The average gradient suggests a different direction for updating a particular parameter.

(b) *Example 2*. If the average gradient decreases in the same direction as the gradient, it additionally provides more information about the loss landscape.

Figure 1: Comparison of Gradient and Average Gradient. The latter accurately reflects the influence of a parameter update on loss (as described by Equation 3, under the assumptions that f represents the visualized loss function, with x and x' denoting the parameter values before and after an update, respectively). The plots refer to a simple case with only one parameter of the model. However, they can also be understood to present the loss contribution of a parameter during a weight update involving multiple model parameters. Appendix F presents visualizations involving two parameters.

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1.2 GRADIENT OPTIMIZATION AND AVERAGING

Gradient optimization dominates deep learning with optimizers like Stochastic Gradient Descent (Liu et al., 2020), RMSProp (Tieleman et al., 2012), Adam (Kingma & Ba, 2014) or Nadam (Dozat, 2016).
The leading algorithms for training do not change frequently over the years. However, our algorithm or its variants may be used along with first-order optimizers.

Gradient averaging is commonly used in machine learning, but in a distinct scenario than in our approach. Momentum is the running average of gradients over subsequent batches (Liu et al., 2020; Kingma & Ba, 2014; Dozat, 2016). It prevents falling into local minimums and may accelerate learning. Similarly, averaging model parameters may improve convergence and learning speed (Ruppert, 1988; Polyak & Juditsky, 1992; Merity et al., 2017; Wei et al., 2023; Sun et al., 2010), though it requires a significant amount of memory. The technique can be described as averaging a function of gradients, as the averaged model parameters over subsequent updates depend on the gradient values.

Accumulating gradients over a batch is inherent in machine learning. In practice, it is equivalent to averaging gradients computed for multiple inputs. However, this approach alone does not take into account the information about a parameter update (Fig. 1), which remains unknown during its computation. Consequently, it does not guarantee the accuracy of computing the influence of the unknown parameter update on the loss. Nevertheless, batching remains fully compatible with our method and is employed in our implementation.

- Our approach is more closely related to some second-order optimization methods (Tan & Lim, 2019) rather than momentum-based or parameter-averaging techniques. This is due to the utilization of information about the curvature of a loss function during each parameter update (Fig. 1). Recently, one of the most popular algorithms for second-order optimization of neural networks is L-BFGS (Berahas et al., 2016). However, the current methods in this field are impractical for training large models due to their computational inefficiency or substantial memory requirements.
- The integrated gradient, closely related to the average gradient, is used in some neural-network
 explainability techniques (Sundararajan et al., 2017; Khorram et al., 2021; Sattarzadeh et al., 2021).
 However, the approximation algorithms for the integral of the gradient used in the literature are very
 inefficient to compute for every parameter update of a model due to the calculation of the Riemann sum (Hughes-Hallett et al., 2021).

¹⁰⁸ 2 Methods

110 2.1 ALGORITHM

112 All of the best and most popular optimizers for training large neural networks rely on the gradient. Consequently, they explicitly ignore how loss function in terms of model parameters behaves in the 113 range between before and after a potential weight update (Fig. 1). The definition of the gradient 114 implies, that it reflects the accurate influence on loss only for learning rates approaching to zero, 115 which does not hold in practice. Consequently, gradient-based optimizers do not calculate the accurate 116 influence on loss of a potential weight update, which may significantly slow down the learning of very 117 deep models with many nonlinear operators, as our experiments show. The average gradient solves 118 the described problem. Our algorithm efficiently approximates the average gradient, providing more 119 reliable information on the update direction that minimizes the loss. The average gradient (contrary 120 to the gradient) is directly proportional to the loss delta (Fig. 1; Equation 14 in Appendix B), hence it 121 accurately describes the influence on loss of a parameter delta. 122

In our algorithm, given a sequential model, the average gradient is approximated and propagated according to the equation proven in Appendix B:

 $\mathcal{A}_{\theta_k}^{\mathcal{VG}} \nabla_{\theta_k} \ell \cong \mathcal{A}_{\theta_k}^{\mathcal{VG}} \frac{\partial \boldsymbol{x}_k}{\partial \theta_k} \cdot \mathcal{A}_{\boldsymbol{x}_k}^{\mathcal{VG}} \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_k} \cdot \ldots \cdot \mathcal{A}_{\boldsymbol{x}_{n-1}}^{\mathcal{VG}} \frac{\partial \boldsymbol{x}_n}{\partial \boldsymbol{x}_{n-1}} \cdot \mathcal{A}_{\boldsymbol{x}_n}^{\mathcal{VG}} \nabla_{\boldsymbol{x}_n} \ell$

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where ℓ is a loss function, θ_k are parameters of a layer no. i and $(x_k, x_{k+1}, \ldots, x_n)$ are inputs and outputs of subsequent layers of a neural network. The notation $\nabla_x f$ refers to the gradient of some function f for an argument x, and $\frac{\partial f}{\partial x}$ denotes the Jacobian. The average operator \mathcal{AVG} of gradients or Jacobians is defined in Appendix A and aligns with intuition. The averages are aggregated with respect to the parameters of a model (θ_k) or the outputs of subsequent layers ($x_k, x_{k+1}, \ldots, x_n$). The average gradients are propagated in the same manner as the gradients in the standard backpropagation algorithm. The computation based on Equation 1 is fast and memory efficient because the procedure is similar to the standard backpropagation of gradients, which is done according to:

$$\nabla_{\theta_k} \ell = \frac{\partial \boldsymbol{x}_k}{\partial \theta_k} \cdot \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_k} \cdot \dots \cdot \frac{\partial \boldsymbol{x}_n}{\partial \boldsymbol{x}_{n-1}} \cdot \nabla_{\boldsymbol{x}_n} \ell$$
(2)

(1)

The version of our algorithm that consists of two iterations (Algorithm 1) first performs the standard 138 backpropagation (Equation 2) through layer outputs x, and model parameters θ along with parameter 139 update of an optimizer (in the experiments it is RMSProp) to new weight values θ' . Then it is assumed 140 that the absolute value of the parameter delta $|\theta - \theta'|$ of the RMSProp optimizer is good enough to 141 retain it. The second backpropagation is performed for eventual negations of update directions only, 142 where, conversely, the average gradient is propagated (Algorithm 2). Importantly, the range on which 143 the gradient is averaged equals $[\theta, \theta']$ (between parameters before and after the estimated potential 144 update; Algorithm 3). The average derivatives of each nonlinear activation are calculated as follows: 145

$$\mathcal{AVG}_{t \in [x,x']} f'(t) = \frac{\int_x^{x'} f'(t) dt}{x' - x} = \frac{f(x') - f(x)}{x' - x}$$
(3)

where f means an activation function (in the experiments it is either ELU or Tanh activation), x means an input scalar assuming forward propagation using the θ weights, and x' means the corresponding input number assuming forward pass for the θ' . The equation is the one-dimensional analogy of the average gradient and the Jacobian, both of which are defined in Appendix A.

In the case of applying an activation function $f : \mathbb{R} \to \mathbb{R}$, or $f : \mathbb{R}^n \to \mathbb{R}^n$, to a layer output $x = \langle x_1, x_2, ..., x_n \rangle$ (assuming parameters θ), which changes to $x' = \langle x'_1, x'_2, ..., x'_n \rangle$ during the forward pass with updated parameters θ' :

$$\mathcal{AVG}_{t \in [\boldsymbol{x}, \boldsymbol{x}']} \frac{\partial \boldsymbol{f}}{\partial t} = \operatorname{diag}(\langle \mathcal{AVG}_{t_1 \in [x_1, x_1']} f'(t_1), \mathcal{AVG}_{t_2 \in [x_2, x_2']} f'(t_2), \dots, \mathcal{AVG}_{t_n \in [x_n, x_n']} f'(t_n) \rangle)$$
(4)

where each term $\mathcal{AVG}_{(\cdot)} f'(\cdot)$ is defined in Equation 3.

161 Let us define a typical layer, denoted as k, which is parameterized by θ_k . This layer could be a convolutional layer, a fully-connected layer, or another operator that is linear over all or most of

its domain. Let us assume that the layer no. k outputs y_k , which is then passed to an activation f_k . Consequently each part of Equation 1 can be approximated as:

$$\begin{aligned} \mathcal{AVG}_{\boldsymbol{x}_{k}} & \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_{k}} = \mathcal{AVG}_{\boldsymbol{x}_{k}} & \frac{\partial f_{k}}{\partial \boldsymbol{x}_{k}} \cong \mathcal{AVG}_{\boldsymbol{x}_{k}} & \frac{\partial \boldsymbol{y}_{k}}{\partial \boldsymbol{x}_{k}} \cdot \mathcal{AVG}_{\boldsymbol{t} \in [\boldsymbol{y}_{k}, \boldsymbol{y}_{k}']} & \frac{\partial f}{\partial t} \\ \mathcal{AVG}_{\boldsymbol{\theta}_{k}} & \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{\theta}_{k}} = \mathcal{AVG}_{\boldsymbol{\theta}_{k}} & \frac{\partial f_{k}}{\partial \boldsymbol{\theta}_{k}} \cong \mathcal{AVG}_{\boldsymbol{\theta}_{k}} & \frac{\partial \boldsymbol{y}_{k}}{\partial \boldsymbol{\theta}_{k}} \cdot \mathcal{AVG}_{\boldsymbol{t} \in [\boldsymbol{y}_{k}, \boldsymbol{y}_{k}']} & \frac{\partial f}{\partial \boldsymbol{t}} \end{aligned} \tag{5}$$

where the approximation, instead of equality, is the consequence of chaining averages of Jacobians, 168 which can be proven analogously to Equation 1 (see Appendix B). The average operator \mathcal{AVG} of Jacobians is defined in Appendix A. $\mathcal{AVG}_{t \in [y_{k+1}, y'_{k+1}]} \frac{\partial f}{\partial t}$ is defined in Equation 4. Generally, the 170 vast majority of applied neural network operators are either nonlinear activations or linear functions 171 in by far most of their domains (e.g., max pooling, convolution, fully connected, or ReLU). In the 172 case of the nonlinear activations, equations no. 3 and 4 are used to compute the average Jacobians. 173 For linear transformations, such as $y_k(x_k)$ and $y_k(\theta_k)$, the average gradients and Jacobians are easy 174 and fast to compute. However, for implementation simplicity and a slight speedup of computations, 175 broader estimates of the average Jacobians from Equation 5 are applied: 176

$$\begin{aligned} &\mathcal{AVG}_{\boldsymbol{x}_{k}} \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_{k}} = \mathcal{AVG}_{\boldsymbol{x}_{k}} \frac{\partial f_{k}}{\partial \boldsymbol{x}_{k}} \cong \frac{\partial \boldsymbol{y}_{k}}{\partial \boldsymbol{x}_{k}} \cdot \mathcal{AVG}_{\boldsymbol{t} \in [\boldsymbol{y}_{k}, \boldsymbol{y}_{k}']} \frac{\partial f}{\partial \boldsymbol{t}} \\ &\mathcal{AVG}_{\boldsymbol{\theta}_{k}} \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{\theta}_{k}} = \mathcal{AVG}_{\boldsymbol{\theta}_{k}} \frac{\partial f_{k}}{\partial \boldsymbol{\theta}_{k}} \cong \frac{\partial \boldsymbol{y}_{k}}{\partial \boldsymbol{\theta}_{k}} \cdot \mathcal{AVG}_{\boldsymbol{t} \in [\boldsymbol{y}_{k}, \boldsymbol{y}_{k}']} \frac{\partial f}{\partial \boldsymbol{t}} \end{aligned} \tag{6}$$

which use the non-averaged Jacobian $\frac{\partial y_k}{\partial x_k}$. Therefore, intuitively, the broad estimation of $\mathcal{AVG}_{x_k} \frac{\partial x_{k+1}}{\partial x_k}$ is approximately between $\frac{\partial x_{k+1}}{\partial x_k}$ and $\mathcal{AVG}_{x_k} \frac{\partial x_{k+1}}{\partial x_k}$, and analogously for $\mathcal{AVG}_{\theta_k} \frac{\partial x_{k+1}}{\partial \theta_k}$.

Algorithm 1 *Simplified algorithm version for 2 iterations.* Back and forward propagation would be called two times in optimal implementation, where memory requirement would be the same as for Adam optimizer. Over the whole paper, we refer to the optimal implementation as the one that minimizes recomputations, avoids costly statistics during training, and is machine-code optimized to the same extent as optimizers from mainstream libraries. The lines marked as redundant within comments in curly brackets are unnecessary for the optimal operation of the below pseudocode.

0	comments in curry brackets are unnecessary for the optimal operation of the below pseudocode.
1	Input: model: Neural Network Model
2	dataset: Training Dataset
	lossFn: Loss Function
	optimizer: Optimizer
	for all $batch \in dataset$ do
	$modelOutput \leftarrow model(batch.x)$ {It is assumed that model's layers' results are kept inside
	$model\}$
	$modelLoss \leftarrow LossFn(modelOutput, batch.y)$
	$modelCopy \leftarrow model \{Copy model\}$
	$modelCopyOutput \leftarrow modelCopy(batch.x)$ {This inference is redundant if $modelCopy$ gets
	also intermediate layers' results copied}
	$modelCopyLoss \leftarrow LossFn(modelCopyOutput, batch.y)$ {This computation is also redun-
	dant, since it is the same as modelLoss}
	Backpropagate(modelCopyLoss) {Compute the gradients using the standard backpropaga-
	tion procedure. Assume that the gradients are stored inside <i>modelCopy</i> }
	optimizer.Step(modelCopy) {Perform weight update on modelCopy (using the gradients
	stored inside modelCopy)}
	modelCopy(batch.x) {Execute inference to store new layer-wise results in modelCopy}
	AveragedBackpropagation(model, modelCopy, modelLoss) {The procedure is de-
	scribed as Algorithm 2. The parameters of the <i>model</i> are modified within}
	end for

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An algorithm version with *n* backpropagation iterations computes (n-1) times the approximated gradient average, each time based on the previous. The intuition behind this is that a better estimate of the averaged derivatives of nonlinear activations is computed after every iteration (Equation 3; backpropagated according to equations no. 4, 6 and 1). Consequently, each time a more precise estimate of the optimal parameter update $(\Delta \theta)^*$ is obtained (where optimality means that the average gradient is accurately estimated and the parameter update is compliant with it). Therefore, once again

proce	dure AveragedBackpropagation(
mode	l: Neural Network Model
mode	lAfterUpdate: model After Candidate Update of
Par	ameters
mode	lLoss: model's Loss)
Ba	ckpropagate(modelLoss,
r	nodel.Layers.Last().Output) {Compute the gradient of modelLoss in terms of the last
lay	er's output. Let us assume that the gradient is assigned to the grad property of the output
vari	able (model.Layers.Last().Output)}
for	$index \leftarrow (Count(model.Layers) - 1) \mathbf{to} \ 0 \mathbf{do}$
θ	$\theta = model.Layers[index].\theta$ {To simplify notation}
i	f IsNonlinear(model.Layers[index]) then {Calculation of either the gradient or its aver
a	ge, which corresponds to the terms in Equation 6}
	${\bf BackpropagateThroughNonlinearLayer} (model. Layers [index]. Output,$
	$model. {\tt Layers} [index]. {\tt Input}, model After Update. {\tt Layers} [index]. {\tt Output},$
	modelAfterUpdate.Layers[index].Input) {Procedure described as Algorithm 3. Le
	us assume that activations are separate layers (like in equations no. 4 and 6)}
e	lse
	Backpropagate(model.Layers[index].Output, model.Layers[index].Input) {The typ
	ical backpropagation procedure. It propagates the gradient through a layer. Let us assume
	that the gradient is assigned to the grad property of the input variable}
	θ .averagedGrad $\leftarrow \theta$.grad {In this case, for a linear layer, the gradient is treated as its
	average (compare equations no. 6 and 5)}
e	nd if
θ	$' = modelAfterUpdate.Layers[index].\theta$ {Notation simplification}
θ	$\leftarrow \theta + \theta' - \theta \cdot sgn(\theta.averagedGrad)$ {Update by the absolute value of <i>optimizer</i> 's
u	pdate from Algorithm 1: $ \theta' - \theta $, but in the direction of the approximated gradient average
	$gn(\theta.averagedGrad)$
s	
s end	for
end end p	l for rocedure
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267 on the *whole dataset*. The first iteration of our method is the gradient-based RMSProp procedure, 268 hence the change in loss for RMSProp $\Delta_{RMSProp}$ is known for both the same model parameters 269 and data as in the case of the loss delta of our method. Therefore, the sum of loss differences after 269 the updates of both approaches can be easily and measurably compared relatively to the sum of loss

270	Algorithm 3 Backpropagation Through Nonlinear Layer. It is assumed that each input number
271	influences a corresponding single output scalar. This is because, in the experiments, the only
272	operators assumed to be nonlinear during backpropagation of the gradient average are certain
273	activation functions: $\mathbb{R} \to \mathbb{R}$).
274	procedure BACKPROPAGATETHROUGHNONLINEARLAYER(LayerOutput: Tensor
275	LayerInput: Tensor
276	LayerOutputAfterUpdate: Tensor
277	LayerInputAfterUpdate: Tensor)
278	$f \leftarrow \text{Layer function}$
279	for all $(outputNum, inputNum, outputNumUpdated, inputNumUpdated) \in Zip($
280	$LayerOutput, LayerInput, LayerOutputAfterUpdate, LayerInputAfterUpdate) {\rm do}$
281	{The commonly used Zip function illustrates iterating through multiple tensors at once}
282	if $ inputNumUpdated - inputNum > \epsilon$ then {Check if the difference in the inputs is
283	higher than a tiny constant ϵ . The condition prevents division by zero. In the experiments
284	$\epsilon \approx 1.19e-7$
285	$\mathcal{AVG}_{x \in [inputNum, inputNumUpdated]} f'(x) = \frac{outputNumUpdated-outputNum}{inputNumUpdated-inputNum}$ {Eq. 3 and
286	$4\}$
287	$inputNum.averagedGrad \leftarrow \mathcal{AVG}_x f'(x) \cdot outputNum.averagedGrad$
288	{Propagate the average gradient backward using the chain rule. Equations 3 and 4 define
289	the term $\mathcal{AVG}_{t \in [y_k, y'_k]} \frac{\partial f}{\partial t}$ in Equation 6, which is part of Equation 1}
200	else
201	$inputNum$.averagedGrad $\leftarrow f'(inputNum) \cdot outputNum$.averagedGrad {In this
291	case $inputNum \approx inputNumUpdated$, thus $\mathcal{AVG}_{x\in[]} f'(x) \approx f'(inputNum)$ for
292	activation functions. The backpropagation towards input complies with the chain rule
293	(equations no. 6 and 1)}
294	end if
295	end for
296	end procedure

deltas of RMSProp:

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$$\mathcal{RD}_{AG,RMSProp} = \frac{\mathcal{AVG}_{b\in B}\,\Delta_{AG} - \mathcal{AVG}_{b\in B}\,\Delta_{RMSProp}}{|\mathcal{AVG}_{b\in B}\,\Delta_{RMSProp}|} \\ = \frac{\sum_{b\in B}\left(\ell_b(\theta'_{AG,b}) - \ell_b(\theta_b)\right)}{|\sum_{b\in B}\left(\ell_b(\theta'_{RMSProp,b}) - \ell_b(\theta_b)\right)|} - sgn(\sum_{b\in B}\left(\ell_b(\theta'_{RMSProp,b}) - \ell_b(\theta_b)\right))$$
(7)

305 $\mathcal{RD}_{AG,RMSProp}$ is the relative difference in avg. loss deltas of RMSProp and the method based 306 on the average gradient (AG). The AVG operator denotes the arithmetic average. B is the set of all 307 batches. $\Delta_{AG,b}$ is the loss delta assuming a batch b after our algorithm's update of model parameters θ_b to new values $\theta'_{AG,b}$. Notation for RMSProp is analogous. ℓ_b is the loss, assuming data of a 308 batch b. sgn is the sign function. \mathcal{RD} would not be as useful when using momentum because the 309 metric compares the aggregated loss of a single batch per parameter update, whereas momentum 310 contributes to a decrease in loss over many batches per a single parameter update. Without the 311 momentum, \mathcal{RD} significantly increases the statistical confidence in comparing training algorithms 312 because, for *the same* model weights, the losses are compared for each weight update. Keeping the 313 same parameter values for each loss delta reduces the variance of \mathcal{RD} , resulting in a decrease in 314 errors when comparing methods. 315

316317 2.2 MODELS AND TRAINING

Our algorithm was tested on three different models with nonlinear ELU (Clevert et al., 2015) and
Tanh activations. Model A has a small number of layers (Table 1), and the second one, Model B, is
much deeper, with 30 nonlinear layers (Table 2; not counting max pooling as nonlinear). The third
model is a convolutional neural network with 46 nonlinear layers (see Appendix H.1 for the training
details of this model). It was assumed that Model A is trained for 15 epochs, while Model B – 500 in
the case of the gradient-based RMSProp training, and 300 for our method. Grid search was used to
find the optimal learning rates for the standard RMSProp training over the course of all 500 epochs,

while our method was optimized only for 200 epochs (out of 300 during testing). The objective of the hyperparameter search was to minimize the loss that is the smallest over a training. The results of the search for optimal learning rates are shown in Table 3. The epoch counts are tailored to ensure that the training achieves minimal or near-minimal test loss values before the final epoch of the gradient-based RMSProp training. The only loss function used in this research is cross-entropy loss, and the batch size is set to 128 in all experiments.

Table 1: Model A

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Tuble 1. h	1000111		Table 2: Model B is desi	gned to test	the perfor-
	Output	Parameter	mance of our algorithm o	n deep neur	al networks
Layers	Shape	Count	to achieve a reasonable tin	ne of many t	rainings for
Convolution 2D (3×3)			statistical significance of the	ne results. I	ne practical-
+ ELU	(8, 26, 26)	80	ity of the architecture is no	5t prioritized	1.
Convolution 2D (3×3)				Output	Parameter
+ ELU	(8, 24, 24)	584	Layers	Shape	Count
Convolution 2D (5×5)			Convolution 2D (3×3)	-	
+ ELU	(10, 10, 10)	0010	+ ELU	(8, 26, 26)	80
stride = 2 padding = 2	(16, 12, 12)	3216	Max Pooling 2D (2×2)	(8, 13, 13)	
Convolution 2D (3×3)	(10, 10, 10)	2220	$\overline{\text{Convolution 2D} (3 \times 3)}$	() / /	
+ ELU	(16, 10, 10)	2320	+ ELU	(16, 11, 11)	1168
Convolution 2D (3×3)			Max Pooling 2D (2×2)	(16, 5, 5)	
+ ELU	(16, 8, 8)	2320	Flatten	400	
Convolution 2D (5×5)			Linear + Tanh	10	4010
+ ELU	(10, 1, 1)	0.11.0	26 ×		
stride = 2 padding = 2	(16, 4, 4)	6416	Linear + Tanh	10	26 ×110
Flatten	256		Linear + Softmax	10	330
Linear + Softmax	10	2570			8228
		17506			00

Models A and B were trained on two popular image datasets: MNIST (LeCun & Cortes, 2010) and Fashion MNIST (Xiao et al., 2017). Both datasets have the same input size $(28 \times 28 \times 1)$, but their image characteristics are *significantly* different. Moreover, since the method does not have any hyperparameters apart from the learning rate, it is less likely to overfit to a specific experimental setup (model, dataset, and learning rate) and show good results on it while experiencing deficient performance on other setups. We further validated the performance of our algorithm on a deep sequential convolutional model using an NLP benchmark, specifically the IMDB dataset. All details are presented in Appendix H.

3 RESULTS

For the shallow model A, all of the training algorithms are approximately equal (Fig. 2a, Fig. 2b). The relative difference in summed loss deltas (Equation 7) revealed that the algorithm based on the average gradient is only marginally better than the standard RMSprop according to $\mathcal{RD} =$ 1.20e-3 ± 2.7e-4 (0.12% faster minimization of loss with 0.027% of SEM error) on MNIST and $\mathcal{RD} = 5.86e-3 \pm 2.79e-3$ on Fashion MNIST in the case of two iterations. For five iterations, $\mathcal{RD} = 6.47e-4 \pm 9.8e-5$ on MNIST and $\mathcal{RD} = 2.37e-3 \pm 4.5e-4$ on Fashion MNIST.

368 The results of Model B are much more interesting. The version of the algorithm with two iterations 369 is about three times faster at minimizing the median of training losses on both datasets (Fig. 2c; 370 Fig. 2d). Moreover, the mean losses tend to be considerably lower than those for standard RMSProp 371 training, even when repeating the experiments using different weight initialization (see Appendix I). 372 Despite the minority of epochs with high oscillations, the method utilizing the average gradient is 373 approximately two to three times faster in minimizing the mean loss, although this is not clearly 374 visible in the plots. Furthermore, for both versions of our algorithm on both datasets, during from 375 49.3% to 70% of epochs, the average training loss was lower with statistical significance (SEM) than for the gradient-based RMSProp. Conversely, our algorithm was worse in that respect during from 376 0.667% to 2.33% of epochs with statistical significance. The average of minimal training losses on 377 MNIST for the five iterations is 0.0393 ± 0.0058 , which is significantly lower than 0.0883 ± 0.0117

Table 3: *Learning rates*. All hyperparameter searches of Model A consist of five trainings for each
learning rate (LR), while in the case of Model B, it is one training, unless stated otherwise. For Model
B, the losses do not directly predict the performance of the methods, because different epoch counts
are used between the methods. The standard error of the mean is used as the confidence range for the
losses, while for the LRs, the maximum distance to the next best LRs on both sides represents the
errors. The LRs used in the experiments are listed in the "Learning Rate" column.

			Learning	The Most Important Hyperparameter Search Results
Dataset	Model	Method	Rate	[Learning Rate: Avg. of Min. Training Loss]
	Model	RMSProp 2 Iterations	8e-4 8e-4	$\begin{array}{c} 6e{-4}: 8.04e{-3}; \ 7e{-4}: 6.48e{-3}; \ 8e{-4}: 5.69e{-3}\\ 9e{-4}: 5.94e{-3}; \ 10e{-4}: 7.70e{-3}; \ 11e{-4}: 7.39e{-3}\\ \hline 8e{-4}: 0.00555; \ 9e{-4}: 0.00692; \ 1e{-3}: 0.00832 \end{array}$
MNIST		5 Iterations	8e-4	8e-4: 0.00514; 9e-4: 0.00580; 1e-3: 0.00678
1011 (10)1				1.5e-4: 0.194; 2e-4: 0.0979; 2.5e-4: 0.0651
				3e-4: 0.0683; 3.5e-4: 0.191; 4e-4: 0.0759
				The best learning rate of the search <i>after</i> the experiments
				$(10 \text{ trainings per LR in } \{1.5e-4, 2e-4, \dots, 5.5e-4\}):$
				$(3.5e-4 \pm 1.5e-4) : (0.0856 \pm 0.0139)$, (matches the
				performance in our experiments in Section 3)
				The loss for a high learning rate (10 trainings):
	Modal	RMSProp	2.5e-4	$1.5\mathrm{e}{-3}:(2.09\pm0.05)$
	B	2 Iterations	7.5e-4	The learning rate is guessed
	D	5 Iterations	7.5e-4	The learning rate is guessed
		DICOD	1 5 0	1e-3: 0.201; 1.25e-3: 0.186; 1.5e-3: 0.183
	Model	RMSProp	1.5e-3	$\frac{1.75e-3:0.189; 2e-3:0.183; 2.25e-3:0.193}{1.000}$
Fachian	А	2 Iterations	$\frac{1.9e-3}{1.5e-3}$	$\frac{1.8e-3:0.180; \mathbf{1.9e-3}:0.179; 2e-3:0.180}{\mathbf{1.5e} \cdot 2 \cdot 0.179; 2e-3:0.180}$
MNIST		5 Iterations	1.5e-5	$\frac{1.36-3:0.178; 1.06-3:0.179; 1.76-3:0.200}{2e-4:0.356: 2.5e-4:0.331: 3e-4:0.285}$
1011 (15) 1				35e-4:0.349:4e-4:0.487:45e-4:0.459
				The best learning rate of the search <i>after</i> the experiments
				(10 trainings per LR in $\{2e-4, 2.5e-4, \dots, 6e-4\}$):
				$(4e_{-4} + 1.5e_{-4}) : (0.318 + 0.016)$ (matches the
				$(40^{\circ} 4 \pm 1.00^{\circ} 4)$. (0.510 ± 0.010) , (matches the
				The loss for a high learning rate (10 trainings):
		DMCDrow	20 1	The loss for a high learning face (10 trainings). $0_{0} = 4 \cdot (0.641 \pm 0.168)$
	Model	2 Iterations	$\frac{3e-4}{9e-4}$	$\frac{9e-4:(0.041\pm0.108)}{6e-4:0.230:0e-4:0.242:1.2e-3:0.255}$
	В	5 Iterations	$\frac{3e-4}{9e-4}$	$\frac{-0.242, 1.26-3, 0.555}{9e-4 \cdot 0.243, 1.26-3, 0.276}$
		5 noradolla	50 4	JC 1.0.210, 1.00 0.0.210

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for the standard RMSProp. Meanwhile, the two iterations are also perform better than the gradientbased RMSProp, but without statistical significance, achieving 0.0747 ± 0.0188 . Even better averages of minimal training losses were obtained on Fashion MNIST, with the five-iteration and two-iteration versions achieving 0.254 ± 0.017 and 0.257 ± 0.014 respectively, compared to 0.314 ± 0.008 by the gradient-based training.

Plots of the test losses of Model B look very similar to the training losses (Appendix C), showing significant improvements in generalization, which correspond to the lower training losses. On MNIST, the average of best accuracies over training for five iterations is equal to (97.87 ± 0.09) %, which is significantly higher than (96.80 ± 0.78) % and (96.75 ± 0.55) % for the two-iteration version and gradient-based algorithm, respectively. On Fashion MNIST, the analogous results are (88.09 ± 0.35) %, (87.54 ± 0.55) % and (86.57 ± 0.29) %, respectively. Appendix D presents the accuracy plots.

For Model B, the \mathcal{RD} metric (Equation 7) provides a very high confidence of superiority of the average gradient for the high learning rates used for the trainings based on our algorithm (Table 3). On MNIST for two and five iterations, it equals 10.41 ± 1.94 and 1.43 ± 0.29 , respectively. On Fashion MNIST, it is 0.58 ± 0.14 and 0.24 ± 0.04 for both variants, respectively.



Figure 2: Training losses. Only mean curves contain confidence ranges (SEM).

Importantly, for Model B, the average-gradient algorithm dominated also for the learning rates that are optimal for the standard RMSProp training. Multiple metrics favored our algorithm with statistical significance, i.e., $\mathcal{RD} \in [0.0611 \pm 0.0004, 1.07 \pm 0.31]$, despite training counts equal to only two or three (for each of the four experiments).

In the case of Model B, our implementation of the two-iteration variant of the algorithm based on the average gradient (Alg. 1) is nearly three times slower per epoch than the training based on the gradient, while the five iterations (Alg. 4 in Appendix G) are almost eight times slower per epoch.
The estimated runtime of optimal implementation is slightly more than two times longer for the two iterations per epoch when compared to the gradient-based RMSProp, and around six to seven times longer for the five iterations.

For the deep convolutional model on the IMDB dataset, sample efficiency of the two iteration variant of our algorithm achieved about 55% gain in sample efficiency compared to the gradient baseline. The analogous result using four iterations falls between 25% and 30%. See Appendix H.2 for details.

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4 CONCLUSIONS

Surprisingly, modifying the gradient on nonlinear activations in very deep models can significantly
 increase sample efficiency for some deep models, which is a direct conclusion of our experiments.
 For the MNIST and Fashion MNIST benchmarks, the algorithm based on the average gradient offers
 significant benefits compared to the standard RMSProp training for the deep model with many stacked
 fully-connected layers and nonlinear activations: (a) About a threefold increase in sample efficiency

486 in terms of median loss, and about two to three times faster mean loss reduction. This is reached 487 by only two iterations, which optimally require a little more than double the time of computation 488 per epoch in comparison with the gradient-based RMSProp training. Meanwhile, our suboptimal 489 implementation of the two-iteration version of the algorithm needs nearly three times more runtime 490 per epoch than the training based on the gradient. Therefore, the presented method is not only more sample-efficient, but it is also faster and saves energy. (b) Outstanding performance on higher 491 learning rates, which may offer significant benefits in terms of both electricity and time spent on 492 hyperparameter searches. (c) Considerably better generalization, at least in a reasonable epoch 493 The increase in sample efficiency and good performance across a wider range of learning count. 494 rates is confirmed by experiments using different weight initialization (see Appendix I). 495

On the other hand, for a deep sequential convolutional model trained on the IMDB dataset, sample
efficiency is improved by about 55% when using only two iterations of our algorithm (Appendix H.2).
This is the only significant benefit of our algorithm in this experiment, as the variant using more
iterations achieved efficiency between that of the vanilla RMSProp and the two-iteration variant.

500 The \mathcal{RD} (Equation 7) confirms the outstanding results of the other measures. The score of \mathcal{RD} = 501 10.41 ± 1.94 , achieved by the two iterations on MNIST, corresponds to the average speed of batch-loss 502 minimization that is $(1141 \pm 194)\%$ of the speed of the gradient-based RMSProp while using the 503 same absolute values of weight updates. In the other cases of deep models, the average speed of batch-loss minimization ranges from $(2.10 \pm 0.18)\%$ to $(243 \pm 29)\%$. Therefore, even a relatively 504 slight speedup in batch-loss minimization (such as 2.1% on the IMDB dataset) can contribute to 505 a significantly higher gain in sample efficiency. Moreover, it is crucial to note that the highest of 506 the mentioned gains occur at learning rates that are three times higher than the optimal rates for 507 gradient-based training. Generally, high learning rate values may enable rapid learning because model 508 parameters are adjusted faster. Nevertheless, the average gradient is also superior in terms of the 509 average speed of batch-loss minimization when using the optimal learning rates for gradient-based 510 training across all tested models, with statistical significance. This validates the proof in Appendix B, 511 as both the metric and the proof focus on the efficiency of batch-loss minimization. On the other hand, 512 refer to Appendix F for the limitations of our algorithm in estimating the locally optimal update.

Surprisingly, the algorithm version with five iterations is worse than the two iterations according to \mathcal{RD} with higher statistical confidence than for other measures. Across all experiments, the variant is computationally inefficient in terms of the resources required to reduce the loss to a certain level.

In the case of the shallow model with nonlinear ELU activations, the method is only marginally better than the standard gradient-based RMSProp training. This behavior is expected due to the scaling properties of the algorithm (Appendix E).

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5 DISCUSSION

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The successful evaluation using different weight initialization techniques on the NLP and computer
 vision benchmarks, using both deep convolutional architecture and the model based on fully-connected
 layers with nonlinear activations, provides insight into significant improvements in sample efficiency,
 at least for some models. Furthermore, the computational cost associated with these improvements is
 modest. These results are especially important in the field of online learning, where sample efficiency
 is crucial.

529 For very deep models without residual connections, gradient-based training tends to be inefficient 530 (Balduzzi et al., 2017), which we demonstrate how to mitigate. In general, the very deep structure 531 of human brains enables the learning of universal and complex patterns. Therefore, accurately mimicking human brain model could potentially lead to satisfactory results. Our algorithm aims to 532 improve learning in scenarios involving neural structures that are very deep, a feature of provably 533 efficient biological brains that distinguishes them from current AI models. Therefore, the method 534 may contribute to the training of large models in the future, where sample efficiency is needed to 535 learn new tasks on the fly, akin to how people or some animals do. 536

However, at present, the potential modifications to the algorithm are even more intriguing. Not only
is it possible to efficiently calculate the average gradient for linear layers using Eq. 5 instead of Eq. 6,
but Eq. 1 can also be utilized to compute the average gradients over a much larger range than that of
a parameter update to capture the global trend of the loss landscape (see Appendix J for future work).

540 **REPRODUCIBILITY STATEMENT** 6

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We put emphasis on providing detailed descriptions of all experiments. The algorithms (Alg. 4 in 543 Appendix G and Alg. 1 in Section 2, with subprocedures labeled as Alg. 2 and Alg. 3) are described 544 in detail in Section 2.1. The models (Tables 1, 2 and 4 in Appendix H), the learning rates (Tables 3 and 5 in Appendix H), and all other important experiment settings are described in Section 2.2 546 and Appendix H.1. The code, along with environment settings, is available under [...]. Appendix B contains one of our most important theoretical results: the proof of Equation 1 and its superiority 547 548 over the gradient in minimizing the batch loss by accurately indicating how each model parameter individually contributes to the change in the batch loss (Equation 14). The proven potential for 549 batch-loss minimization is verified not only by the \mathcal{RD} metric with high statistical significance but 550 also by comparisons of training losses and other metrics (Section 3).

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634 635 636	A DEFINITION OF AVERAGE GRADIENT/JACOBIAN
637 638	Let us define the average gradient of a function $f(x) : \mathbb{R}^n \to \mathbb{R}$ for some row vector $x \in [a, b]$ (the formula is analogous to the one-dimensional case in Equation 3):
639 640 641	$\mathcal{AVG}_{\boldsymbol{x}\in[\boldsymbol{a},\boldsymbol{b}]}\nabla_{\boldsymbol{x}}f = (\boldsymbol{b}-\boldsymbol{a})^{\circ-1} \circ \int_{\boldsymbol{a}}^{\boldsymbol{b}} \nabla_{\boldsymbol{x}}f \ d\boldsymbol{x} = (\boldsymbol{b}-\boldsymbol{a})^{\circ-1} \circ \int_{0}^{1} \nabla_{\boldsymbol{a}+t\cdot(\boldsymbol{b}-\boldsymbol{a})}f \ dt $ (8)
642 643 644 645	where \circ denotes the elementwise operation of either multiplication or inversion $((\cdot)^{\circ-1})$. However, the cases of vector elements where division by zero occurs are handled differently, using the partial derivative $\frac{\partial f}{\partial x_i}$:

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648 If $f(x) : \mathbb{R}^n \to \mathbb{R}^m$, then using to Equation 8:

$$\mathcal{AVG}_{\mathbf{\mathcal{I}}} \frac{\partial \boldsymbol{f}}{\partial \boldsymbol{x}} = \begin{bmatrix} \mathcal{AVG}_{\boldsymbol{x} \in [\boldsymbol{a}, \boldsymbol{b}]} \nabla_{\boldsymbol{x}} f_1 \\ \mathcal{AVG}_{\boldsymbol{x} \in [\boldsymbol{a}, \boldsymbol{b}]} \nabla_{\boldsymbol{x}} f_2 \end{bmatrix} = \begin{bmatrix} (\boldsymbol{b} - \boldsymbol{a})^{\circ - 1} \circ \int_{\boldsymbol{a}}^{\boldsymbol{b}} \nabla_{\boldsymbol{x}} f_1 \ d\boldsymbol{x} \\ (\boldsymbol{b} - \boldsymbol{a})^{\circ - 1} \circ \int_{\boldsymbol{a}}^{\boldsymbol{b}} \nabla_{\boldsymbol{x}} f_2 \ d\boldsymbol{x} \end{bmatrix}$$

Again, the cases of vector elements where division by zero occurs are handled as follows:

$$\forall i: b_i - a_i = 0 \implies \mathcal{AVG}_{x_i \in [a_i, b_i]} \frac{\partial f}{\partial x_i} = \frac{\partial f}{\partial a_i}$$
(11)

B PROOF OF EQUATION 1 AND ITS LOSS-MINIMIZATION POTENTIAL

B.1 DEFINITION AND PROPERTIES OF AVERAGE GRADIENT OF LOSS

Using Equation 2, the average gradient $\mathcal{AVG}_{\theta_k} \nabla_{\theta_k} \ell$ can be defined without the approximation given in Equation 1:

 $\mathcal{AVG}_{\theta_{k}} \nabla_{\theta_{k}} \ell = \mathcal{AVG}_{(\theta_{k}, \boldsymbol{x}_{k}, \boldsymbol{x}_{k+1}, \dots, \boldsymbol{x}_{n})} \left(\frac{\partial \boldsymbol{x}_{k}}{\partial \theta_{k}} \cdot \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_{k}} \cdot \dots \cdot \frac{\partial \boldsymbol{x}_{n}}{\partial \boldsymbol{x}_{n-1}} \cdot \nabla_{\boldsymbol{x}_{n}} \ell \right)$ (12)

where multiple variables are under the average operator $(\theta_k, x_k, x_{k+1}, \dots, x_n)$. There are numerous ways to define how $(x_k, x_{k+1}, \ldots, x_n)$ depend on the weights and biases θ_k , as they all change together during a parameter update. To compute the average (Equation 12), it can be assumed that the parameters of the layer no. k and the outputs of the layers change linearly with respect to each other, as if they move from θ_k to θ'_k and from (x_k, \ldots, x_n) to (x'_k, \ldots, x'_n) after an update of the parameters of all layers. Under this assumption, the calculation is formulated as follows: while computing the average, the integral contains a function $f_{\theta,k}(t) = \theta_k + t \cdot (\theta'_k - \theta_k)$ for the variable under integration $t \in [0, 1]$ (θ_k and θ'_k denote model parameters before and after an update, respectively). Moreover, the integral involves each layer's output: $f_{x,i}(t) = x_i + t \cdot (x'_i - x_i)$. Finally, the average gradient (Equation 12) is equal to:

$$\mathcal{AVG}_{\theta_{k}} \nabla_{\theta_{k}} \ell = \mathcal{AVG}_{f_{\theta,k}} \nabla_{f_{\theta,k}} \ell = \mathcal{AVG}_{t} \left(\frac{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)}{\partial f_{\theta,k}(t)} \cdot \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)} \cdot \dots \cdot \frac{\partial \boldsymbol{f}_{\boldsymbol{x},n}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},n-1}(t)} \cdot \nabla_{\boldsymbol{f}_{\boldsymbol{x},n}(t)} \ell(t) \right) \\
= \int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)}{\partial f_{\theta,k}(t)} \cdot \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)} \cdot \dots \cdot \frac{\partial \boldsymbol{f}_{\boldsymbol{x},n}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},n-1}(t)} \cdot \nabla_{\boldsymbol{f}_{\boldsymbol{x},n}(t)} \ell(t) dt \tag{13}$$

which is more direct and easier to work with.

Importantly, unlike the gradient, the average gradient $(\mathcal{AVG}_{\theta_k} \nabla_{\theta_k} \ell)$ is directly proportional to the loss-change impact of each model parameter separately $l_{\theta',k} - l_{\theta,k}$ (of the shape of θ_k and θ'_k , unlike the scalar ℓ):

$$\mathcal{AVG}_{\theta_{k}} \nabla_{\theta_{k}} \ell = \mathcal{AVG}_{\theta_{k}} \mathcal{O}_{\theta_{k}} (\sum_{j=0}^{n} \sum_{i=0}^{|\theta_{j}|} \ell_{\theta,j,i}) = \mathcal{AVG}_{\theta_{k}} \nabla_{\theta_{k}} (\sum_{i=0}^{|\theta_{k}|} \ell_{\theta,k,i}) = \mathcal{AVG}_{\theta_{k}} (\operatorname{diag}(\frac{\partial \boldsymbol{l}_{\theta,k}}{\partial \theta_{k}})) = \mathcal{AVG}_{\theta_{k}} (\mathcal{O}_{\theta_{k},i}) = \mathcal{O}_{\theta_{k}} \mathcal{O}_{\theta_{k},i} (\mathcal{O}_{\theta_{k},i}) = \mathcal{O}_{\theta_{k}} \mathcal{O}_{\theta_{k}} (\mathcal{O}_{\theta_{k},i}) = \mathcal{O}_{\theta_{k}} (\mathcal$$

$$= \langle \mathcal{AVG}_{\theta_{k,1}} \ell'_{\theta,k,1}, \dots, \mathcal{AVG}_{\theta_{k,n}} \ell'_{\theta,k,n} \rangle = \langle \frac{\int_{\theta_{k,1}}^{\theta_{k,1}} \ell'_{\theta,k,1} \, d\vartheta}{\theta'_{k,1} - \theta_{k,1}}, \dots, \frac{\int_{\theta_{k,n}}^{\theta_{k,n}} \ell'_{\theta,k,n} \, d\vartheta}{\theta'_{k,n} - \theta_{k,n}} \rangle \tag{14}$$

$$=\langle \frac{\ell_{\theta',k,1}-\ell_{\theta,k,1}}{\theta'_{k,1}-\theta_{k,1}},\ldots,\frac{\ell_{\theta',k,n}-\ell_{\theta,k,n}}{\theta'_{k,n}-\theta_{k,n}}\rangle = (\theta'_k-\theta_k)^{\circ-1} \circ (\boldsymbol{l}_{\theta',k}-\boldsymbol{l}_{\theta,k}) \propto \boldsymbol{l}_{\theta',k}-\boldsymbol{l}_{\theta,k}$$

702 where \circ denotes the elementwise operation of either multiplication or inversion $((\cdot)^{\circ-1})$. diag $(\frac{\partial l_{\theta,k}}{\partial \theta_{k}})$ 703 denotes diagonal elements of the Jacobian matrix. $\ell_{\theta,k,i} \in l_{\theta,k}$ represents the scalar loss contribution 704 of a single model parameter $(\theta_{k,i})$, that can be defined as an integral of the gradient: $\ell_{\theta,k,i} =$ 705 $\int_{C_1}^{\theta_{k,i}} \nabla_{\vartheta} \ell_{\theta} \, d\vartheta + C_2, \text{ for any constant scalars } C_1 \text{ and } C_2. \text{ (Note that in this case, } \ell_{\theta,k,i} \neq \ell_{\theta} + C_{\ell}, \text{ for any constant } C_{\ell}, \text{ because the loss } \ell \text{ also depends on other parameters than } \theta_{k,i}.) \text{ Important}$ 706 707 properties: (a) $\ell_{\theta} = C + \sum_{k=0}^{n} \sum_{i=0}^{|\theta_k|} \ell_{\theta,k,i}$ for a constant C that is invariant across updates of the model parameters θ . (b) The elements of l are related to the difference in loss during 708 709 parameter update: $\ell_{\theta'} - \ell_{\theta} = (\sum_{k=0}^{n} \sum_{i=0}^{|\theta'_k|} \ell_{\theta',k,i}) - (\sum_{k=0}^{n} \sum_{i=0}^{|\theta_k|} \ell_{\theta,k,i}).$ (c) The following equation is satisfied: $\nabla_{\theta} \ell = \nabla_{\theta} (\sum_{k=0}^{n} \sum_{i=0}^{|\theta_k|} \ell_{\theta,k,i}).$ The simple one-dimensional visualization of the proportionality from Equation 14 $(\mathcal{AVG}_{\theta_k} \nabla_{\theta_k} \ell \propto \mathbf{l}_{\theta',k} - \mathbf{l}_{\theta,k})$ is shown in Fig. 1. Note that 710 711 712 the property of proportionality does not hold for the gradient updates (which are utilized by Adam 713 (Kingma & Ba, 2014), RMSProp (Tieleman et al., 2012), and SGD (Ketkar, 2017; Liu et al., 2020)). 714 In the gradient case, during the update step of θ weights, θ' is not used in the calculation of itself. 715 Therefore, $l_{\theta'} - l_{\theta}$ cannot be computed yet, and the accurate influence on loss remains unknown, 716 unlike for the average gradient (Equation 14). The cases of scalar parameters $\theta_{k,i} \in \theta_k$ and $\theta'_{k,i} \in \theta'_k$ 717 where division by zero occurs are handled differently: 718

$$\forall i: \theta'_{k,i} - \theta_{k,i} = 0 \implies \mathcal{AVG}_{\vartheta_{k,i} \in [\theta_{k,i}, \theta'_{k,i}]} \frac{\partial \ell}{\partial \vartheta_{k,i}} = \frac{\partial \ell}{\partial \theta_{k,i}}$$
(15)

Assuming the functions $f_{\theta,k}$ and $f_{x,i}$ from Equation 13 are any functions (but differentiable with respect to each other), Equation 14 remains valid. Therefore, the crucial property of direct proportionality to the loss values does not depend on our previous assumptions about θ_k and x_i . The purpose of these assumptions is to provide a simple example, reduce reasoning abstraction, and simplify further proofs in Sections B.2 and B.3.

B.2 PROOF OF OF EQUATION 1 WITHOUT SPECIFYING PRECISION OF APPROXIMATION

For some function f and some constants C_1, C_2, \ldots, C_n :

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$$\int C_1 \cdot C_2 \cdot \ldots \cdot C_n \cdot f(x) \, dx = C_1 \cdot C_2 \cdot \ldots \cdot C_n \cdot \int f(x) \, dx \tag{16}$$

Similarly, let us denote approximately constant functions as $C'_1(x) \cong C_1, C'_2(x) \cong C_2, \dots, C'_n(x) \cong C_n$ for some $x \in [a, b], a \neq b$. The constant that precisely approximates each function C'(x), is its average: $C'_1(x) \cong AVG C'_1(x) = C_1, C'_2(x) \cong AVG C'_2(x) = C_2, \dots, C'_n(x) \cong AVG C'_n(x) = C_n$. Therefore, similarly to Equation 16:

$$\int_{a}^{b} C'_{1}(x) \cdot \ldots \cdot C'_{n}(x) \cdot f(x) \, dx \cong \underset{x \in [a,b]}{\mathcal{AVG}} C'_{1}(x) \cdot \ldots \cdot \underset{x \in [a,b]}{\mathcal{AVG}} C'_{n}(x) \cdot \int_{a}^{b} f(x) \, dx$$

$$\int_{a}^{b} C'_{1}(x) \cdot \ldots \cdot C'_{n}(x) \cdot f(x) \, dx \cong \int_{a}^{b} \frac{C'_{1}(x)}{b-a} dx \cdot \ldots \cdot \int_{a}^{b} \frac{C'_{n}(x)}{b-a} dx \cdot \int_{a}^{b} f(x) \, dx$$
(17)

which is also approximately equal to both sides of Equation 16. In Equation 17, both approximations are equivalent, because $\mathcal{AVGC}'_i(x) = \int_a^b C'_i(x)/(b-a) \, dx$. For functions $\mathbb{R}^n \to \mathbb{R}^m$, equations no. 16 and 17 are analogous. Note that, in the general case, the different approximations of the terms $C'_i(x) \cong C'_i(a)$ and $C'_i(x) \cong C_i(b)$ are worse than the average: $C'_i(x) \cong \mathcal{AVGC}'_i(x) = C_i$ (which is used further in Section B.3).

Rapid changes in the gradient over the range of an update indicate that the update step is too large, leading to instability and reduced training effectiveness due to excessively large steps in the loss landscape. We assume effective learning, where gradients do not change significantly¹ between updates, ensuring the learning rate is appropriately sized. In this case, the gradient $\nabla_{\theta_k} \ell$ does not change significantly² over the range of a weight update $[\theta, \theta']$. However, these assumptions are

 ¹The magnitude of the gradient change need not be specified, as it suffices that it contributes to the approximations with unspecified bounds in Equations 18 and 19. The accuracy of these approximations is proven in Section B.3.

²See footnote 1.

merely intended to build intuition and are not necessary for this proof. We do not yet assume any specific level of precision in how Equation 17 approximates Equation 13:

$$\mathcal{AVG} \nabla_{\theta_k} \ell = \int_0^1 \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)}{\partial f_{\theta,k}(t)} \cdot \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)} \cdot \dots \cdot \frac{\partial \boldsymbol{f}_{\boldsymbol{x},n}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},n-1}(t)} \cdot \nabla_{\boldsymbol{f}_{\boldsymbol{x},n}(t)} \ell(t) dt$$
$$\cong \int_0^1 \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)} dt \cdot \int_0^1 \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)} dt \cdot \dots \cdot \int_0^1 \frac{\partial \boldsymbol{f}_{\boldsymbol{x},n}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},n}(t)} dt \cdot \int_0^1 \nabla_{\boldsymbol{f}_{\boldsymbol{x},n}(t)} \ell(t) dt$$

$$\int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)}{\partial f_{\theta,k}(t)} dt \cdot \int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)} dt \cdot \dots \cdot \int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},n-1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},n-1}(t)} dt \cdot \int_{0}^{1} \nabla_{\boldsymbol{f}_{\boldsymbol{x},n}(t)} \ell(t) dt \quad (18)$$
$$= \mathcal{A}_{\mathcal{V}}\mathcal{G} \frac{\partial \boldsymbol{x}_{k}}{\partial \theta_{k}} \cdot \mathcal{A}_{\boldsymbol{x}_{k}}\mathcal{G} \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_{k}} \cdot \dots \cdot \mathcal{A}_{\boldsymbol{x}_{n-1}}\mathcal{G} \frac{\partial \boldsymbol{x}_{n}}{\partial \boldsymbol{x}_{n-1}} \cdot \mathcal{A}_{\boldsymbol{x}_{n}}\mathcal{V}\mathcal{G} \nabla_{\boldsymbol{x}_{n}}\ell$$

Applying the notation of Equation 12 to Equation 18, we get:

$$\begin{aligned}
\mathcal{A}_{\theta_{k}}^{\mathcal{V}\mathcal{G}} \nabla_{\theta_{k}} \ell &\cong \int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)}{\partial f_{\theta,k}(t)} dt \int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},k+1}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},k}(t)} dt \cdots \int_{0}^{1} \frac{\partial \boldsymbol{f}_{\boldsymbol{x},n}(t)}{\partial \boldsymbol{f}_{\boldsymbol{x},n-1}(t)} dt \int_{0}^{1} \nabla_{\boldsymbol{f}_{\boldsymbol{x},n}(t)} \ell(t) dt \\
&= \int_{\theta_{k}}^{\theta_{k}'} \frac{\partial \boldsymbol{x}_{k}(\vartheta_{k})}{\partial \vartheta_{k}} d\vartheta_{k} \int_{\boldsymbol{x}_{k}'}^{\boldsymbol{x}_{k}'} \frac{\partial \boldsymbol{x}_{k+1}(\boldsymbol{\chi}_{k})}{\partial \boldsymbol{\chi}_{k}} d\boldsymbol{\chi}_{k} \cdots \int_{\boldsymbol{x}_{n-1}}^{\boldsymbol{x}_{n-1}'} \frac{\partial \boldsymbol{x}_{n}(\boldsymbol{\chi}_{n-1})}{\partial \boldsymbol{\chi}_{n-1}} d\boldsymbol{\chi}_{n-1} \int_{\boldsymbol{x}_{n}}^{\boldsymbol{x}_{n}'} \nabla_{\boldsymbol{\chi}_{n}} \ell d\boldsymbol{\chi}_{n} \\
&= \mathcal{A}_{\theta_{k}}^{\mathcal{V}\mathcal{G}} \frac{\partial \boldsymbol{x}_{k}}{\partial \theta_{k}} \cdot \mathcal{A}_{\boldsymbol{x}_{k}}^{\mathcal{V}\mathcal{G}} \frac{\partial \boldsymbol{x}_{k+1}}{\partial \boldsymbol{x}_{k}} \cdots \mathcal{A}_{\boldsymbol{x}_{n-1}'}^{\mathcal{V}\mathcal{G}} \frac{\partial \boldsymbol{x}_{n}}{\partial \boldsymbol{x}_{n-1}} \cdot \mathcal{A}_{\boldsymbol{x}_{n}}^{\mathcal{V}\mathcal{G}} \nabla_{\boldsymbol{x}_{n}} \ell \end{aligned} \tag{19}$$

where $\theta_k, x_k, x_{k+1}, \dots, x_n$ are all linear functions of t (previously denoted as $f_{\theta,k}, f_{\boldsymbol{x},k}, f_{\boldsymbol{x},k+1}, \dots, f_{\boldsymbol{x},n}$. Therefore, the functions $\boldsymbol{x}_k(\theta_k), \boldsymbol{x}_{k+1}(\boldsymbol{x}_k), \dots, \boldsymbol{x}_n(\boldsymbol{x}_{n-1})$ are known. The edge cases of those scalars within $\theta_k, x_k, x_{k+1}, \ldots, x_n$ that do not depend on t are handled analogously to Equation 15, as in these cases the average gradient equals the gradient.

Despite the provided arguments on why the approximation is applied, the precision of the estimation is not specified, although it is crucial. Therefore, the accuracy of the approximation is described in Section B.3. Otherwise, if the precision of the estimation is not important, then Equation 19 ultimately proves Equation 1. \Box

The analogous reasoning can be applied to prove Equation 5.

In the algorithm, it is also assumed that the average gradient of the loss with respect to the output of the last layer, denoted as $(\mathcal{AVG}_{\boldsymbol{x}_n} \nabla_{\boldsymbol{x}_n} \ell)$, is replaced by the gradient $(\nabla_{\boldsymbol{x}_n} \ell)$. Moreover, in our implementation, the gradients replace the average gradients of layers that are approximately linear (using Equation 6 instead of Equation 5), resulting in a broader approximation in Equation 1. However, the presented reasoning still applies, including the proof of approximation accuracy in Section B.3. See Appendix F for comments on the limitations of our implementation of Equation 1.

PROOF OF SUFFICIENT PRECISION OF APPROXIMATION B.3

Referring to the content of the paragraphs just before and after Equation 17, the approximation in Equation 17 is more precise in the case of $C'_i(x) \cong \mathcal{AVG} C'_i(x) = C_i$ than in the case of approximating $C'_i(x) \cong C'_i(a)$. The average Jacobian of each term in Equation 1 can be denoted as $\mathcal{AVG} C'_i(x)$, while the Jacobian of each term in Equation 2 can be denoted as $C'_i(a)$. For the average Jacobian $\mathcal{AVG} C'_i(x)$, a better estimation in Equation 17 is obtained, as stated in the text near the equation. Consequently, applying Equation 17 to approximate Equation 13 results in a higher precision in estimating Equation 1 when averaging each Jacobian term separately, compared to utilizing the Jacobians without averaging. Therefore, a better approximation of the accurate average gradient is obtained compared to using the gradient. \Box The average gradient is proportional to the change in loss after the corresponding parameter update (Equation 14). Therefore, approximating the average gradient more precisely than current gradient-based methods can lead to more efficient minimization of batch loss, for example, by using Eq. 1. Therefore, learning can be enhanced compared to the potential of gradient-based methods.



Figure 3: Test losses of Model B. Only mean curves contain confidence ranges (SEM).

D TEST ACCURACY CURVES OF MODEL B

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Figure 4: Test accuracy of Model B. Only mean curves contain confidence ranges (SEM).

E SCALING IN TERMS OF MODEL DEPTH

The algorithm based on the average gradient aims to reduce errors of the predicted influence on loss of a parameter update. In the case of the gradient-based approach, the errors arise from the impaired prediction of how inputs to subsequent layers influence their outputs (Fig. 1). Let us model the errors as multiplicative, because each time a fraction of output may be influenced by the error (Balduzzi et al., 2017). Therefore, when compared to the gradient-based algorithm as a baseline, the multiplicative errors are reduced after backpropagation through each nonlinear layer (by computing the average Jacobian of the layer). Consequently, the incorporation of the average gradient exponentially reduces the error in terms of a count of nonlinear layers (that are involved in the backpropagation process). This explains the huge performance-improvement gap between the models for the method based on the average gradient, which emerges from the difference in models' depths. However, the gap is also increased due to the linearity of the ELU activation function in most of its domain, where the gradient equals its average. In this case, our algorithm produces results similar to those of gradient-based optimization.

If the errors (of the predicted influence on loss of a parameter update) are enormous, then the learning
is impossible. Therefore, the learning performance tends to decrease after the error reaches a certain
value for a given model, learning rate, and other parameters. From that point onward, our algorithm
more efficiently reduces the batch loss compared to the gradient-based approach by minimizing the
error in the loss-influence prediction. Importantly, the improvements tend to increase with both the
number of nonlinear layers in a model and the learning rate.

F THREE-DIMENSIONAL COMPARISON OF THE GRADIENT AND THE AVERAGE GRADIENT

In our experiments, during a parameter update, in terms of the average reduction of loss for a batch, our algorithm lies between the gradient (red arrows in Figure 5) and the lowest average gradient (black arrows in Figure 5). Our algorithm does not always find a locally optimal solution (the best in the range of a single parameter update) because:

- a The average gradient is approximated (by using Equation 1 instead of Equation 3, Equation 6 as a substitute of Equation 5, and the non-averaged gradient of the loss with respect to the last layer output).
- b The optimal parameter update may be inaccurately estimated before the average gradient for this parameter update is calculated. Moreover, even after many iterations of Algorithm 4 (Appendix G), the update step may not converge to a locally optimal solution (black vectors in Figure 5).
- c After the first iteration of our algorithm, only the negations of the directions of changes in each parameter are possible. Thus, the search for locally optimal updates is bounded by $2^{|\Theta|}$ combinations, where $|\Theta|$ is the count of trainable parameters.

Nevertheless, the *RD* metric (defined in Equation 7) indicates our algorithm minimizes the batch
 loss more efficiently on average compared to the gradient-based approach.



Figure 5: *Three-dimensional comparison of the gradient and the lowest average gradient in a few example scenarios.* The latter accurately reflects the influence on the loss of a parameter update. Furthermore, it accurately shows how each model parameter individually contributes to the change in the batch loss (Equation 14), which is utilized by our algorithm. Each plot illustrates the loss in terms of two example model parameters, assuming a specific magnitude for each parameter update (represented by the radius of each white circle). The arrows point to the loss values after an update based on the gradient and the average gradient. The average gradient is calculated for the update that minimizes it. Therefore, it points to the minimum loss on each white circle, although this minimum is not always achieved by the approximated average gradient computed by our algorithm.

972 G ALGORITHM VERSION WITH PARAMETERIZED NUMBER OF ITERATIONS 973

,	of iterations is equal to the number of backpropagation calls and inferences in optimal implement The memory requirement of the ideal implementation would be higher than that of Adam by on
	additional scalar size per parameter of the model.
	Input: model: Neural Network Model to Train
	dataset: Training Dataset
	lossFn: Loss Function
	optimizer: Optimizer
	<i>iterCount</i> : Number of Backpropagation Iterations
	for all $batch \in dataset do$
	$modelInitial \leftarrow model$
	$modelCopy \leftarrow model$
	initialOutput \leftarrow modelCopy(batch.x) {It is assumed that modelCopy's layers' result
	kept inside modelCopy}
	$initialLoss \leftarrow LossFn(initialOutput, batch.y)$
	Backpropagate(initialLoss) {Compute the gradients using the standard backpropag
	procedure. Assume that the gradients are stored inside <i>model(opy)</i>
	$modelOutnut A fterUndate \leftarrow modelComu(hatch x)$
	$modelOutput A fterUndate \leftarrow Loss En(modelOutput A fterUndate batch y)$
	for $iter - 1$ $iterCount - 1$ do $\{I \text{ oon } (iterCount - 1) \text{ times because one back}\}$
	$ron agation is done \}$
	if $iter \neq 1$ then
	$modelCopu \leftarrow model$
	$modelCopy(batch.x)$ {For each layer, compute its output, and store it inside $modelC$
	$model \leftarrow modelInitial$
	end if
	<i>initialOutput</i> \leftarrow <i>model</i> (<i>batch.x</i>) {This computation is redundant if layer output
	copied from <i>modelInitial</i> }
	$initialLoss \leftarrow LossFn(initialOutput, batch.y)$ {Analogously, this computation is
	redundant}
	AveragedBackpropagation(model, modelCopy, initialLoss) {The procedure
	scribed as Algorithm 2. The parameters of the <i>model</i> are modified within}
	end for

H CONVOLUTIONAL NEURAL NETWORK ON IMDB

1011 1012 H.1 METHODS

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We refer to Model C (Tab. 4) as our very deep convolutional model, which we tested on the IMDB dataset. This model, primarily composed of convolutional layers, is designed to evaluate the performance of our learning algorithm on a deep convolutional neural network without skip connections. Skip connections simplify the learning task by enabling the network to leverage features that can be extracted by shallower networks (Veit et al., 2016). Our primary goal is to assess the algorithm's capabilities, rather than achieving state-of-the-art results.

1019 IMDB preprocessing includes: (a) Equal split for test and training sets. (b) Duplicate removal. 1020 (c) Punctuation removal. (d) Tokenization. (e) Padding to the length of 122 (mean training-exam-1021 ple length), and keeping the final part of each review. (f) Lemmantization. (g) Vectorization using 1022 GloVe embeddings (Pennington et al., 2014). Finally, the input has the shape (1, 122, 50), where 1023 each input word is converted into its corresponding GloVe embedding with a length of 50. Model C 1024 (Tab. 4) utilizes multiple convolutional layers of shape (1×1) , which are used to change the data 1025 shape and, for each "pixel", to extract features from the outputs of different filters. The neighboring 1026 dimensions of each GloVe embedding do not have any special relationship compared to the distinct

1027			
1028		Output	Parameter
1029	Layers	Shape	Count
1030	Convolution 2D (1×50) , Tanh	(50, 122, 1)	2550
1031	Convolution 2D (1×1) , Tanh	(40, 122, 1)	2040
1032	Convolution 2D (1×1) , Tanh	(35, 122, 1)	1435
1033	Convolution 2D (1×1) , Tanh	(30, 122, 1)	1080
1034	Convolution 2D (1×1) , Tanh	(27, 122, 1)	837
1035	Convolution 2D (1×1) , Tanh	(24, 122, 1)	672
1036	Convolution 2D (1×1) , Tanh	(21, 122, 1)	525
1037	Convolution 2D (1×1) , Tanh	(18, 122, 1)	396
1038	Convolution 2D (1×1) , Tanh	(16, 122, 1)	304
1030	Convolution 2D (1×1) , Tanh	(14, 122, 1)	238
1035	Convolution 2D (1×1) , Tanh	(12, 122, 1)	180
1040	Convolution 2D (1×1) , Tanh	(10, 122, 1)	130
1041	Convolution 2D (1×1) , Tanh	(8, 122, 1)	88
1042	Convolution 2D (1×1) , Tanh	(6, 122, 1)	54
1043	Convolution 2D (1×1) , Tanh	(5, 122, 1)	35
1044	Convolution 2D (3×1) with stride = 2, Tan	h $(5, 60, 1)$	80
1045	25 ×Convolution 2D (3×1) , Tanh	(5, 10, 1)	25×80
1046	Flatten	50	
1047	Linear, Tanh	25	1275
1048	Linear, Tanh	13	338
1049	Linear, Tanh	7	98
1050	Linear, Tanh	4	32
1051	Linear, Softmax	2	10
1052			14397

Table 4: Model C.

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ones. Therefore, we used convolutions with a filter-size dimension equal to either one or all featuresin the GloVe embeddings.

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1058 H.2 RESULTS

Both versions of our algorithm were more sample-efficient than the gradient-based RMSProp, as indicated by the training loss (Fig. 6). In the case of gradient-based training, the trade-off between the 1061 mean and median of the training loss is visible in both Figs. 6a and 6b. The tendency for instability in 1062 training with a higher learning rate leads to the occurrence of outliers, also in terms of whole worse 1063 trainings, which lower the mean. However, the median is resistant to these outliers. This can also 1064 be observed in Figures 7b and 8b. To evaluate both the mean and median, considering the trade-off between them, we compared the methods by averaging the median and mean losses. This approach 1066 provides a consistent comparison result across both learning rates of the gradient-based RMSProp 1067 (Fig. 6). Using this evaluation method, the performance of gradient-based RMSProp at epoch 200 1068 is approximately equal to the results of the two iterations of our method at epochs 125 and 130, in 1069 Figs. 6a and 6b, respectively. This translates to a sample efficiency between 53% and 60% higher 1070 in favor of the two iterations of our algorithm compared to the vanilla RMSProp. Surprisingly, the performance of the four iterations falls between the other methods, with the sample efficiency gain 1071 ranging from 25% to 30%. The $\mathcal{RD} = 0.0394 \pm 0.0053$ metric also favors the two-iteration variant, 1072 outperforming the $\mathcal{RD} = 0.0210 \pm 0.0018$ achieved by the four iterations. 1073

1074 The test-accuracy (Fig. 8) and test-loss curves (Fig. 7) should be interpreted in the context that 1075 the objective of the hyperparameter search is dependent solely on the training loss. In addition, 1076 considering the trade-off between the mean and median losses, which occurs between the lower and 1077 higher learning rates of the vanilla RMSProp, slightly better mean generalization in the gradient 1078 training for the low learning rate (Fig. 7a) does not imply generally better test performance. However, 1079 the comparison using the same learning rate (Fig. 7b) indicates that the two-iteration variant achieves 1079 the most stable test-loss performance. 1080Table 5: Learning rates for the IMDB dataset. The "Learning Rate" column presents the final chosen1081learning rates for the experiments. A repeated loss of 0.6931 is equivalent to the lack of training. The1082table includes the results of the final experiments; however, the results are clipped to 150 epochs for1083the variants of our algorithm. The best results and the learning rates chosen for the experiments are1084marked in bold.



rate of 3.641e-4.

(b) Gradient-based RMSProp with the lower learning rate of **6.906e**-4.



Figure 6: *Training loss of Model C*. Only mean curves contain confidence ranges (SEM).

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Due to suboptimal backpropagation of the average gradient through activations in our implementation, it has a bigger computational overhead for models applying activations to large feature maps.
 Therefore, our implementation is computationally slower relatively to the gradient-based training for Model C than in the case of Model B.

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(a) Gradient-based RMSProp with the lower learning rate of **3.641e**-4.

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Figure 8: Test accuracy of Model C. Only mean curves contain confidence ranges (SEM).

1176 I EXPERIMENTS WITH ALTERNATIVE WEIGHT INITIALIZATION FOR MODEL B

1178 We repeated the experiments for Model B on the MNIST and Fashion MNIST datasets due to 1179 suboptimal parameter initialization, which resulted in vanishing gradients at the start of the training. During the repeated experiments, we initialized the weights using the Glorot uniform method (Glorot 1180 & Bengio, 2010), which is specifically designed to initialize layers with nonlinear activations such as 1181 Tanh or Sigmoid. A gradient-magnitude gain of $\frac{5}{3}$ for Tanh activations was used, as recommended 1182 by the PyTorch library. Biases were initialized to zero. The gradient magnitudes were examined to 1183 ensure they fell within a satisfactory range after initialization. Training length was reduced to 125 1184 epochs for the gradient-based RMSProp and 50 epochs for two iterations of our method to test a 2.5x 1185 learning speedup. 1186

1187 As expected, similar magnitudes of learning rates performed well in the trainings using different weight initializations (compare Tables 3 and 6). The losses using our method are significantly lower

Iut	sie o. Resuits for afferent tearning rates.
	The Most Important Results
Dataset Method	[Learning Rate: Avg. of Min. Training Loss]
	8 trainings per each learning rate:
	$1e-4: 0.0945 \pm 0.0062; 1.5e-4: 0.0821 \pm 0.0121;$
	$1.75e-4: 0.0750 \pm 0.0102; 2e-4: 0.0519 \pm 0.0057;$
	$2.25e-4: 0.0862 \pm 0.0171; \ \mathbf{2.5e-4}: 0.0417 \pm 0.0035$
	$2.75e-4: 0.0515 \pm 0.0094; 3e-4: 0.0749 \pm 0.0187;$
RMSProp	$3.5e-4: 0.0536 \pm 0.0095; 4e-4: 0.0604 \pm 0.0188;$
MNILST (125 epochs)	50 trainings: $2.5e-4: 0.0478 \pm 0.0032;$
WIN151	3 trainings per each learning rate:
	$1.54e-4: 0.128 \pm 0.0068; 4.61e-4: 0.0351 \pm 0.0029;$
	8 trainings per each learning rate:
	$6e-4: 0.0320 \pm 0.0013; 8e-4: 0.0256 \pm 0.0019;$
	$9e-4: 0.0245 \pm 0.0005; 1e-3: 0.0255 \pm 0.0021;$
2 Iterations	$1.1e-3: 0.0253 \pm 0.0016; \ 1.2e-3: 0.0246 \pm 0.0005;$
(50 epochs)	$1.4e-3: 0.0255 \pm 0.0002;$
	8 trainings per each learning rate:
	$2.5e-4: 0.329 \pm 0.022; 3e-4: 0.352 \pm 0.024;$
	$3.5e-4: 0.315 \pm 0.025; 4e-4: 0.347 \pm 0.024;$
	$4.5e-4: 0.374 \pm 0.029; 5e-4: 0.396 \pm 0.025;$
	$5.5e-4: 0.310 \pm 0.024; 6e-4: 0.371 \pm 0.025;$
RMSProp	$6.5e-4: 0.368 \pm 0.034; \ 7e-4: 0.407 \pm 0.021;$
Fashion (125 epochs)	50 trainings: $3.5e-4: 0.344 \pm 0.010;$
MNIST	3 trainings per each learning rate:
	$1.33e-4: 0.467 \pm 0.002; 1.75e-4: 0.463 \pm 0.001;$
	8 trainings per each learning rate:
	$6e-4: 0.269 \pm 0.003; 8e-4: 0.255 \pm 0.002;$
	$9e-4: 0.255 \pm 0.002; 1e-3: 0.254 \pm 0.003;$
	$1.1e-3: 0.253 \pm 0.003; \ \mathbf{1.2e-3}: 0.245 \pm 0.001;$
	1.4_{0} $3 \cdot 0.252 \pm 0.003 \cdot 1.5_{0}$ $3 \cdot 0.261 \pm 0.004 \cdot 1.5_{0}$
	$1.4e^{-3}$. 0.252 ± 0.003 , $1.5e^{-3}$. 0.201 ± 0.004 ,
2 Iterations	$1.6e-3: 0.252 \pm 0.003; 1.8e-3: 0.201 \pm 0.004;$ $1.6e-3: 0.253 \pm 0.003; 1.8e-3: 0.273 \pm 0.008;$

Table 6. Results for different learning rates

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after 2.5 times fewer epochs. Therefore, the two iterations of our method increase sample efficiency by more than 2.5 times. Good performance of the method across different parameter-initialization distributions is essential for its practical application as it contributes to robustness. Importantly, our algorithm maintains its performance gain compared to the gradient-based RMSProp in scenarios involving vanishing gradients, as demonstrated in the main experiments.

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J FUTURE WORK

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Interesting directions for further experiments include: (a) Computing the average gradients over a 1232 much larger range than that of a parameter update to capture the global trend of the loss landscape. 1233 (b) More accurate approximation of the average Jacobians using Equation 5 instead of Equation 6. 1234 This would enable computing the average Jacobians of linear operators. Therefore, the algorithm 1235 based on the average gradient may enhance trainings of deep models without nonlinear activations. 1236 Moreover, the usage of Equation 5 may further improve the performance in the case of many 1237 nonlinear activations because of the increased precision in approximating the average gradient. 1238 (c) Incorporation of the momentum into our algorithm. Preferably Nesterov momentum (Dozat, 1239 2016) should be used. If not, the average gradient would also be calculated for the momentum part of the update step. This could often reverse the direction of the momentum for a model parameter, 1240 thereby impairing the effectiveness of the entire momentum procedure. (d) Development of similar 1241 algorithms, but with update steps, that, for a given model parameter, vary in size over the iterations

of the average-gradient computation. By adjusting the step size of each model parameter to the absolute value of the average gradient, the learning process may be enhanced. (e) Tests of the method on large and very deep architectures, that are used in practice and contain many nonlinear layers. (f) More research on how the method scales up (Appendix E), also in relation to the number of neurons in layers of neural networks. (g) Experiments with learning without forgetting (Li & Hoiem, 2017) and online learning. Sample efficiency may be very beneficial there.