

On the difficulty of training Recurrent Neural Networks

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Abstract

There are two widely known issues with properly training Recurrent Neural Networks, the *vanishing* and the *exploding* gradient problems detailed in Bengio *et al.* (1994). In this paper we attempt to improve the understanding of the underlying issues by exploring these problems from an analytical, a geometric and a dynamical systems perspective. Our analysis is used to justify a simple yet effective solution. We propose a gradient norm clipping strategy to deal with exploding gradients and a soft constraint for the vanishing gradients problem. We validate empirically our hypothesis and proposed solutions in the experimental section.

1. Introduction

A recurrent neural network (RNN), e.g. Fig. 1, is a neural network model proposed in the 80's (Rumelhart *et al.*, 1986; Elman, 1990; Werbos, 1988) for modeling time series. The structure of the network is similar to that of a standard multilayer perceptron, with the distinction that we allow connections among hidden units associated with a time delay. Through these connections the model can retain information about the past inputs, enabling it to discover temporal correlations between events that are possibly far away from each other in the data (a crucial property for proper learning of time series).

While in principle the recurrent network is a simple and powerful model, in practice, it is unfortunately hard to train properly. Among the main reasons why this model is so unwieldy are the *vanishing gradient*

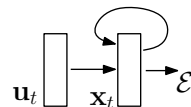


Figure 1. Schematic of a recurrent neural network. The recurrent connections in the hidden layer allow information to persist from one input to another.

and *exploding gradient* problems described in Bengio *et al.* (1994).

1.1. Training recurrent networks

A generic recurrent neural network, with input \mathbf{u}_t and state \mathbf{x}_t for time step t , is given by equation (1). In the theoretical section of this paper we will sometimes make use of the specific parametrization given by equation (11)¹ in order to provide more precise conditions and intuitions about the everyday use-case.

$$\mathbf{x}_t = F(\mathbf{x}_{t-1}, \mathbf{u}_t, \theta) \quad (1)$$

$$\mathbf{x}_t = \mathbf{W}_{rec}\sigma(\mathbf{x}_{t-1}) + \mathbf{W}_{in}\mathbf{u}_t + \mathbf{b} \quad (2)$$

The parameters of the model are given by the recurrent weight matrix \mathbf{W}_{rec} , the biases \mathbf{b} and input weight matrix \mathbf{W}_{in} , collected in θ for the general case. \mathbf{x}_0 is provided by the user, set to zero or learned, and σ is an element-wise function (usually the *tanh* or *sigmoid*). A cost \mathcal{E} measures the performance of the network on some given task and it can be broken apart into individual costs for each step $\mathcal{E} = \sum_{1 \leq t \leq T} \mathcal{E}_t$, where $\mathcal{E}_t = \mathcal{L}(\mathbf{x}_t)$.

One approach that can be used to compute the necessary gradients is Backpropagation Through Time (BPTT), where the recurrent model is represented as

¹ This formulation is equivalent to the more widely known equation $\mathbf{x}_t = \sigma(\mathbf{W}_{rec}\mathbf{x}_{t-1} + \mathbf{W}_{in}\mathbf{u}_t + \mathbf{b})$, and it was chosen for convenience.

a deep multi-layer one (with an unbounded number of layers) and backpropagation is applied on the unrolled model (see Fig. 2).

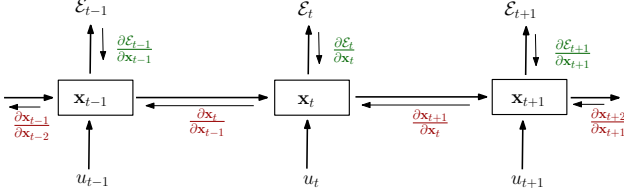


Figure 2. Unrolling recurrent neural networks in time by creating a copy of the model for each time step. We denote by \mathbf{x}_t the hidden state of the network at time t , by \mathbf{u}_t the input of the network at time t and by \mathcal{E}_t the error obtained from the output at time t .

We will diverge from the classical BPTT equations at this point and re-write the gradients (see equations (3), (4) and (5)) in order to better highlight the exploding gradients problem. These equations were obtained by writing the gradients in a sum-of-products form.

$$\frac{\partial \mathcal{E}}{\partial \theta} = \sum_{1 \leq t \leq T} \frac{\partial \mathcal{E}_t}{\partial \theta} \quad (3)$$

$$\frac{\partial \mathcal{E}_t}{\partial \theta} = \sum_{1 \leq k \leq t} \left(\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} \frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k} \frac{\partial^+ \mathbf{x}_k}{\partial \theta} \right) \quad (4)$$

$$\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k} = \prod_{t \geq i > k} \frac{\partial \mathbf{x}_i}{\partial \mathbf{x}_{i-1}} = \prod_{t \geq i > k} \mathbf{W}_{rec}^T \text{diag}(\sigma'(\mathbf{x}_{i-1})) \quad (5)$$

$\frac{\partial^+ \mathbf{x}_k}{\partial \theta}$ refers to the “immediate” partial derivative of the state \mathbf{x}_k with respect to θ , i.e., where \mathbf{x}_{k-1} is taken as a constant with respect to θ . Specifically, considering equation 2, the value of any row i of the matrix $(\frac{\partial^+ \mathbf{x}_k}{\partial \mathbf{W}_{rec}})$ is just $\sigma(\mathbf{x}_{k-1})$. Equation (5) also provides the form of Jacobian matrix $\frac{\partial \mathbf{x}_i}{\partial \mathbf{x}_{i-1}}$ for the specific parametrization given in equation (11), where diag converts a vector into a diagonal matrix, and σ' computes the derivative of σ in an element-wise fashion.

Note that each term $\frac{\partial \mathcal{E}_t}{\partial \theta}$ from equation (3) has the same form and the behaviour of these individual terms determine the behaviour of the sum. Henceforth we will focus on one such generic term, calling it simply the gradient when there is no confusion.

Any gradient component $\frac{\partial \mathcal{E}_t}{\partial \theta}$ is also a sum (see equation (4)), whose terms we refer to as *temporal* contributions or *temporal* components. One can see that each such temporal contribution $\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} \frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k} \frac{\partial^+ \mathbf{x}_k}{\partial \theta}$ measures how θ at step k affects the cost at step $t > k$. The factors

$\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ (equation (5)) transport the error “in time” from step t back to step k . We would further loosely distinguish between *long term* and *short term* contributions, where long term refers to components for which $k \ll t$ and short term to everything else.

2. Exploding and Vanishing Gradients

As introduced in Bengio *et al.* (1994), the *exploding gradients* problem refers to the large increase in the norm of the gradient during training. Such events are caused by the explosion of the long term components, which can grow exponentially more than short term ones. The *vanishing gradients* problem refers to the opposite behaviour, when long term components go exponentially fast to norm 0, making it impossible for the model to learn correlation between temporally distant events.

2.1. The mechanics

To understand this phenomenon we need to look at the form of each temporal component, and in particular at the matrix factors $\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ (see equation (5)) that take the form of a product of $t - k$ Jacobian matrices. *In the same way a product of $t - k$ real numbers can shrink to zero or explode to infinity, so does this product of matrices* (along some direction \mathbf{v}).

In what follows we will try to formalize these intuitions (extending a similar derivation done in Bengio *et al.* (1994) where only a single hidden unit case was considered).

If we consider a linear version of the model (i.e. set σ to the identity function in equation (11)) we can use the *power iteration method* to formally analyze this product of Jacobian matrices and obtain tight conditions for when the gradients explode or vanish (see the supplementary materials for a detailed derivation of these conditions). It is *sufficient* for the largest eigenvalue λ_1 of the recurrent weight matrix to be smaller than 1 for long term components to vanish (as $t \rightarrow \infty$) and *necessary* for it to be larger than 1 for gradients to explode.

We can generalize these results for nonlinear functions σ where the absolute values of $\sigma'(x)$ is bounded (say by a value $\gamma \in \mathcal{R}$) and therefore $\|\text{diag}(\sigma'(\mathbf{x}_k))\| \leq \gamma$.

We first **prove** that it is *sufficient* for $\lambda_1 < \frac{1}{\gamma}$, where λ_1 is the absolute value of the largest eigenvalue of the recurrent weight matrix \mathbf{W}_{rec} , for the *vanishing gradient* problem to occur. Note that we assume the parametrization given by equation (11). The Jacobian matrix $\frac{\partial \mathbf{x}_{k+1}}{\partial \mathbf{x}_k}$ is given by $\mathbf{W}_{rec}^T \text{diag}(\sigma'(\mathbf{x}_k))$. The 2-norm of this Jacobian is bounded by the product of

the norms of the two matrices (see equation (6)). Due to our assumption, this implies that it is smaller than 1.

$$\forall k, \left\| \frac{\partial \mathbf{x}_{k+1}}{\partial \mathbf{x}_k} \right\| \leq \|\mathbf{W}_{rec}^T\| \|\text{diag}(\sigma'(\mathbf{x}_k))\| < \frac{1}{\gamma} \gamma < 1 \quad (6)$$

Let $\eta \in \mathbb{R}$ be such that $\forall k, \left\| \frac{\partial \mathbf{x}_{k+1}}{\partial \mathbf{x}_k} \right\| \leq \eta < 1$. The existence of η is given by equation (6). By induction over i , we can show that

$$\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} \left(\prod_{i=k}^{t-1} \frac{\partial \mathbf{x}_{i+1}}{\partial \mathbf{x}_i} \right) \leq \eta^{t-k} \frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} \quad (7)$$

As $\eta < 1$, it follows that, according to equation (7), long term contributions (for which $t - k$ is large) go to 0 exponentially fast with $t - k$. \square

By inverting this proof we get the *necessary* condition for *exploding gradients*, namely that the largest eigenvalue λ_1 is larger than $\frac{1}{\gamma}$ (otherwise the long term components would vanish instead of exploding). For tanh we have $\gamma = 1$ while for sigmoid we have $\gamma = 1/4$.

2.2. Drawing similarities with Dynamical Systems

We can improve our understanding of the exploding gradients and vanishing gradients problems by employing a dynamical systems perspective, as it was done before in Doya (1993); Bengio *et al.* (1993).

We recommend reading Strogatz (1994) for a formal and detailed treatment of dynamical systems theory. For any parameter assignment θ , depending on the initial state \mathbf{x}_0 , the state \mathbf{x}_t of an autonomous dynamical system converges, under the repeated application of the map F , to one of several possible different attractor states (e.g. point attractors, though other type of attractors exist). The model could also find itself in a chaotic regime, case in which some of the following observations may not hold, but that is not treated in depth here. Attractors describe the asymptotic behaviour of the model. The state space is divided into basins of attraction, one for each attractor. If the model is started in one basin of attraction, the model will converge to the corresponding attractor as t grows.

Dynamical systems theory tells us that as θ changes slowly, the asymptotic behaviour changes smoothly almost everywhere except for certain crucial points where drastic changes occur (the new asymptotic behaviour ceases to be topologically equivalent to the old one). These points are called bifurcation boundaries and are caused by attractors that appear, disappear or change shape.

(Doya, 1993) hypothesizes that such bifurcation crossings could cause the gradients to explode. We would like to extend this observation into a sufficient condition for gradients to explode, and for that reason we will re-use the one-hidden unit model (and plot) from (Doya, 1993) (see Fig. 3).

The x-axis covers the parameter b and the y-axis the asymptotic state \mathbf{x}_∞ . The bold line follows the movement of the final point attractor, \mathbf{x}_∞ , as b changes. At b_1 we have a bifurcation boundary where a new attractor emerges (when b decreases from ∞), while at b_2 we have another that results in the disappearance of one of the two attractors. In the interval (b_1, b_2) we are in a rich regime, where there are two attractors and the change in position of boundary between them, as we change b , is traced out by a dashed line. The vector field (gray dashed arrows) describe the evolution of the state \mathbf{x} if the network is initialized in that region.

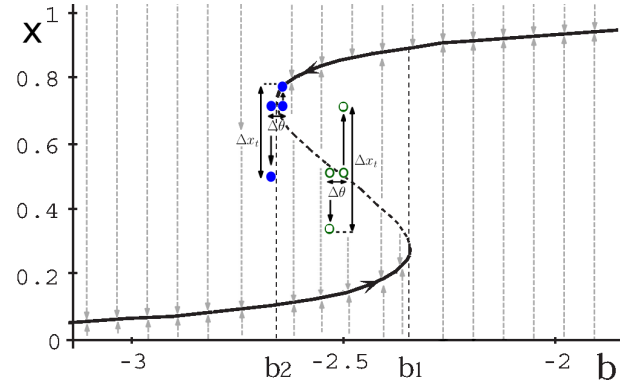


Figure 3. Bifurcation diagram of a single hidden unit RNN (with fixed recurrent weight of 5.0 and adjustable bias b ; example introduced in Doya (1993)). See text.

We show that there are two types of events that could lead to a large change in \mathbf{x}_t , with $t \rightarrow \infty$. One is crossing a boundary between basins of attraction (depicted with a unfilled circles), while the other is crossing a bifurcation boundary (filled circles). For large t , the Δx_t resulting from a change in b will be large even for very small changes in b (as the system is attracted towards different attractors) which leads to a large gradient.

It is however *neither necessary nor sufficient* to cross a bifurcation for the gradients to explode, as bifurcations are global events that could have no effect locally. Learning traces out a path in the parameter-state space. If we are at a bifurcation boundary, but the state of the model is such that it is in the basin of attraction of one attractor (from many possible attractors) that does not change shape or disappear when the bifurcation is crossed, then this bifurcation will not affect learning.

Crossing boundaries between basins of attraction is a *local* event, and it is *sufficient* for the gradients to explode. If we assume that crossing into an emerging attractor or from a disappearing one (due to a bifurcation) qualifies as crossing some boundary between attractors, that we can formulate a *sufficient* condition for gradients to explode which encapsulates the observations made in Doya (1993), extending them to also normal crossing of boundaries between different basins of attractions. Note how in the figure, there are only two values of b with a bifurcation, but a whole range of values for which there can be a boundary crossing.

Another limitation of previous analysis is that they only consider autonomous systems and assume the observations hold for input-driven models. In (Bengio *et al.*, 1994) input is dealt with by assuming it is bounded noise. The downside of this approach is that it limits how one can reason about the input. In practice, the input is supposed to drive the dynamical system, being able to leave the model in some attractor state, or kick it out of the basin of attraction when certain triggering patterns present themselves.

We propose to extend our analysis to input driven models by folding the input into the map. We consider the family of maps F_t , where we apply a different F_t at each step. Intuitively, for the gradients to explode we require the same behaviour as before, where (at least in some direction) the maps F_1, \dots, F_t agree and change direction. Fig. 4 describes this behaviour.

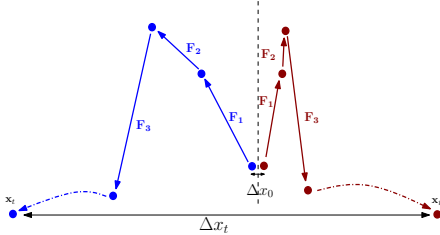


Figure 4. This diagram illustrates how the change in \mathbf{x}_t , $\Delta\mathbf{x}_t$, can be large for a small $\Delta\mathbf{x}_0$. The blue vs red (left vs right) trajectories are generated by the same maps F_1, F_2, \dots for two different initial states.

For the specific parametrization provided by equation (11) we can take the analogy one step further by decomposing the maps F_t into a fixed map \tilde{F} and a time-varying one U_t . $F(\mathbf{x}) = \mathbf{W}_{rec}\sigma(\mathbf{x}) + \mathbf{b}$ corresponds to an input-less recurrent network, while $U_t(\mathbf{x}) = \mathbf{x} + \mathbf{W}_{in}\mathbf{u}_t$ describes the effect of the input. This is depicted in in Fig. 5. Since U_t changes with time, it can not be analyzed using standard dynamical systems tools, but \tilde{F} can. This means that when a boundary between basins of attractions is crossed for \tilde{F} , the state will move towards a different attractor,

which for large t could lead (unless the input maps U_t are opposing this) to a large discrepancy in \mathbf{x}_t . Therefore studying the asymptotic behaviour of \tilde{F} can provide useful information about where such events are likely to happen.

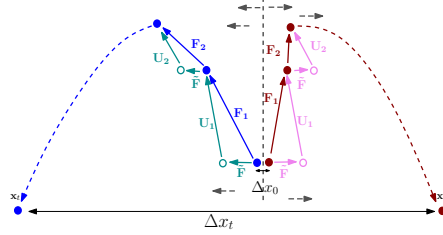


Figure 5. Illustrates how one can break apart the maps F_1, \dots, F_t into a constant map \tilde{F} and the maps U_1, \dots, U_t . The dotted vertical line represents the boundary between basins of attraction, and the straight dashed arrow the direction of the map \tilde{F} on each side of the boundary. This diagram is an extension of Fig. 4.

One interesting observation from the dynamical systems perspective with respect to vanishing gradients is the following. If the factors $\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ go to zero (for $t - k$ large), it means that \mathbf{x}_t does not depend on \mathbf{x}_k (if we change \mathbf{x}_k by some Δ , \mathbf{x}_t stays the same). This translates into the model at \mathbf{x}_t being close to convergence towards some attractor (which it would reach from anywhere in the neighbourhood of \mathbf{x}_k).

2.3. The geometrical interpretation

Let us consider a simple one hidden unit model (equation (8)) where we provide an initial state x_0 and train the model to have a specific target value after 50 steps. Note that for simplicity we assume no input.

$$x_t = w\sigma(x_{t-1}) + b \quad (8)$$

Fig. 6 shows the error surface $\mathcal{E}_{50} = (\sigma(x_{50}) - 0.7)^2$, where $x_0 = .5$ and σ to be the sigmoid function.

We can more easily analyze the behavior of this model by further simplifying it to be linear (σ then being the identity function), with $b = 0$. $x_t = x_0 w^t$ from which it follows that $\frac{\partial x_t}{\partial w} = t x_0 w^{t-1}$ and $\frac{\partial^2 x_t}{\partial w^2} = t(t-1) x_0 w^{t-2}$, implying that when the first derivative explodes, so does the second derivative.

In the general case, when the gradients explode they do so along some directions \mathbf{v} . This says that there exists, in such situations, a vector \mathbf{v} such that $\frac{\partial \mathcal{E}_t}{\partial \theta} \mathbf{v} \geq C \alpha^t$, where $C, \alpha \in \mathbb{R}$ and $\alpha > 1$. For the linear case (σ is the identity function), \mathbf{v} is the eigenvector corresponding to the largest eigenvalue of \mathbf{W}_{rec} . If this bound is tight, we hypothesize that in general *when gradients*

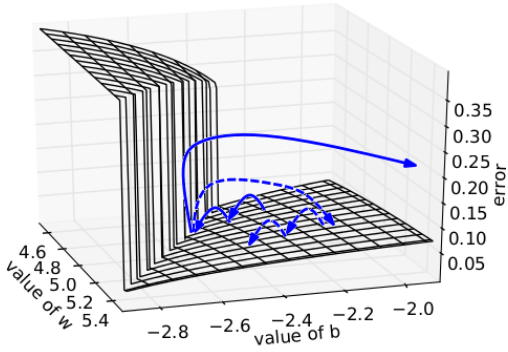


Figure 6. We plot the error surface of a single hidden unit recurrent network, highlighting the existence of high curvature walls. The solid lines depicts standard trajectories that gradient descent might follow. Using dashed arrow the diagram shows what would happen if the gradients is rescaled to a fixed size when its norm is above a threshold.

explode so does the curvature along \mathbf{v} , leading to a wall in the error surface, like the one seen in Fig. 6.

If this holds, then it gives us a simple solution to the exploding gradients problem depicted in Fig. 6.

If both the gradient and the leading eigenvector of the curvature are aligned with the exploding direction \mathbf{v} , it follows that the error surface has a steep wall perpendicular to \mathbf{v} (and consequently to the gradient). This means that when stochastic gradient descent (SGD) reaches the wall and does a gradient descent step, it will be forced to jump across the valley moving perpendicular to the steep walls, possibly leaving the valley and disrupting the learning process.

The dashed arrows in Fig. 6 correspond to *ignoring the norm of this large step*, ensuring that the model stays close to the wall. The key insight is that all the steps taken when the gradient explodes are aligned with \mathbf{v} and ignore other descent direction (i.e. the model moves perpendicular to the wall). At the wall, a small-norm step in the direction of the gradient therefore merely pushes us back inside the smoother low-curvature region besides the wall, whereas a regular gradient step would bring us very far, thus slowing or preventing further training. Instead, with a bounded step, we get back in that smooth region near the wall where SGD is free to explore other descent directions.

The important addition in this scenario to the classical high curvature valley, is that we assume that the valley is wide, as we have a large region around the wall where if we land we can rely on first order methods to move towards the local minima. This is why just clipping the gradient might be sufficient, not requiring the use a second order method. Note that this algo-

rithm should work even when the rate of growth of the gradient is not the same as the one of the curvature (a case for which a second order method would fail as the ratio between the gradient and curvature could still explode).

Our hypothesis could also help to understand the recent success of the Hessian-Free approach compared to other second order methods. There are two key differences between Hessian-Free and most other second-order algorithms. First, it uses the full Hessian matrix and hence can deal with exploding directions that are not necessarily axis-aligned. Second, it computes a new estimate of the Hessian matrix before each update step and can take into account abrupt changes in curvature (such as the ones suggested by our hypothesis) while most other approaches use a smoothness assumption, i.e., averaging 2nd order signals over many steps.

3. Dealing with the exploding and vanishing gradient

3.1. Previous solutions

Using an L1 or L2 penalty on the recurrent weights can help with exploding gradients. Given that the parameters initialized with small values, the spectral radius of \mathbf{W}_{rec} is probably smaller than 1, from which it follows that the gradient can not explode (see necessary condition found in section 2.1). The regularization term can ensure that during training the spectral radius never exceeds 1. This approach limits the model to a simple regime (with a single point attractor at the origin), where any information inserted in the model has to die out exponentially fast in time. In such a regime we can not train a generator network, nor can we exhibit long term memory traces.

Doya (1993) proposes to pre-program the model (to initialize the model in the right regime) or to use *teacher forcing*. The first proposal assumes that if the model exhibits from the beginning the same kind of asymptotic behaviour as the one required by the target, then there is no need to cross a bifurcation boundary. The downside is that one can not always know the required asymptotic behaviour, and, even if such information is known, it is not trivial to initialize a model in this specific regime. We should also note that such initialization does not prevent crossing the boundary between basins of attraction, which, as shown, could happen even though no bifurcation boundary is crossed.

Teacher forcing is a more interesting, yet a not very well understood solution. It can be seen as a way of initializing the model in the right regime and the right

region of space. It has been shown that in practice it can reduce the chance that gradients explode, and even allow training generator models or models that work with unbounded amounts of memory (Pascanu and Jaeger, 2011; Doya and Yoshizawa, 1991). One important downside is that it requires a target to be defined at every time step.

In Hochreiter and Schmidhuber (1997); Graves *et al.* (2009) a solution is proposed for the vanishing gradients problem, where the structure of the model is changed. Specifically it introduces a special set of units called LSTM units which are linear and have a recurrent connection to itself which is fixed to 1. The flow of information into the unit and from the unit is guarded by an input and output gates (their behaviour is learned). There are several variations of this basic structure. This solution does not address explicitly the exploding gradients problem.

Sutskever *et al.* (2011) use the Hessian-Free optimizer in conjunction with *structural damping*, a specific damping strategy of the Hessian. This approach seems to deal very well with the vanishing gradient, though more detailed analysis is still missing. Presumably this method works because in high dimensional spaces there is a high probability for long term components to be orthogonal to short term ones. This would allow the Hessian to rescale these components independently. In practice, one can not guarantee that this property holds. As discussed in section 2.3, this method is able to deal with the exploding gradient as well. Structural damping is an enhancement that forces the change in the state to be small, when the parameter changes by some small value $\Delta\theta$. This asks for the Jacobian matrices $\frac{\partial \mathbf{x}_t}{\partial \theta}$ to have small norm, hence further helping with the exploding gradients problem. The fact that it helps when training recurrent neural models on long sequences suggests that while the curvature might explode at the same time with the gradient, it might not grow at the same rate and hence not be sufficient to deal with the exploding gradient.

Echo State Networks (Lukoševičius and Jaeger, 2009) avoid the exploding and vanishing gradients problem by not learning the recurrent and input weights. They are sampled from hand crafted distributions. Because usually the largest eigenvalue of the recurrent weight is, by construction, smaller than 1, information fed in to the model has to die out exponentially fast. This means that these models can not easily deal with long term dependencies, even though the reason is slightly different from the vanishing gradients problem. An extension to the classical model is represented by leaky integration units (Jaeger *et al.*, 2007), where

$$\mathbf{x}_k = \alpha \mathbf{x}_{k-1} + (1 - \alpha) \sigma(\mathbf{W}_{rec} \mathbf{x}_{k-1} + \mathbf{W}_{in} \mathbf{u}_k + \mathbf{b}).$$

While these units can be used to solve the standard benchmark proposed by Hochreiter and Schmidhuber (1997) for learning long term dependencies (see (Jaeger, 2012)), they are more suitable to deal with low frequency information as they act as a low pass filter. Because most of the weights are randomly sampled, is not clear what size of models one would need to solve complex real world tasks.

We would make a final note about the approach proposed by Tomas Mikolov in his PhD thesis (Mikolov, 2012) (and implicitly used in the state of the art results on language modelling (Mikolov *et al.*, 2011)). It involves clipping the gradient’s temporal components element-wise (clipping an entry when it exceeds in absolute value a fixed threshold). Clipping has been shown to do well in practice and it forms the backbone of our approach.

3.2. Scaling down the gradients

As suggested in section 2.3, one simple mechanism to deal with a sudden increase in the norm of the gradients is to rescale them whenever they go over a threshold (see algorithm 1).

Algorithm 1 Pseudo-code for norm clipping the gradients whenever they explode

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 $\hat{\mathbf{g}} \leftarrow \frac{\partial \mathcal{E}}{\partial \theta}$ 
if  $\|\hat{\mathbf{g}}\| \geq threshold$  then
     $\hat{\mathbf{g}} \leftarrow \frac{threshold}{\|\hat{\mathbf{g}}\|} \hat{\mathbf{g}}$ 
end if
    
```

This algorithm is very similar to the one proposed by Tomas Mikolov and we only diverged from the original proposal in an attempt to provide a better theoretical foundation (ensuring that we always move in a descent direction with respect to the current mini-batch), though in practice both variants behave similarly.

The proposed clipping is simple to implement and computationally efficient, but it does however introduce an additional hyper-parameter, namely the threshold. One good heuristic for setting this threshold is to look at statistics on the average norm over a sufficiently large number of updates. In our experiments we have noticed that for a given task and model size, training is not very sensitive to this hyper-parameter and the algorithm behaves well even for rather small thresholds.

The algorithm can also be thought of as adapting the learning rate based on the norm of the gradient. Compared to other learning rate adaptation strategies, which focus on improving convergence by collecting statistics on the gradient (as for example in

Duchi *et al.* (2011), or Moreira and Fiesler (1995) for an overview), we rely on the *instantaneous* gradient. This means that we can handle very abrupt changes in norm, while the other methods would not be able to do so.

3.3. Vanishing gradient regularization

We opt to address the vanishing gradients problem using a regularization term that represents a preference for parameter values such that back-propagated gradients neither increase or decrease too much in magnitude. Our intuition is that increasing the norm of $\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ means the error at time t is more sensitive to all inputs $\mathbf{u}_t, \dots, \mathbf{u}_k$ ($\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ is a factor in $\frac{\partial \mathcal{E}_t}{\partial \mathbf{u}_k}$). In practice some of these inputs will be irrelevant for the prediction at time t and will behave like noise that the network needs to learn to ignore. The network can not learn to ignore these irrelevant inputs unless there is an error signal. These two issues can not be solved in parallel, and it seems natural to expect that we need to force the network to increase the norm of $\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ at the expense of larger errors (caused by the irrelevant input entries) and *then* wait for it to learn to ignore these irrelevant input entries. This suggest that moving towards increasing the norm of $\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ can not be always done while following a descent direction of the error \mathcal{E} (which is, for e.g., what a second order method would try to do), and therefore we need to enforce it via a regularization term.

The regularizer we propose below prefers solutions for which the error signal preserves norm as it travels back in time:

$$\Omega = \sum_k \Omega_k = \sum_k \left(\frac{\left\| \frac{\partial \mathcal{E}}{\partial \mathbf{x}_{k+1}} \frac{\partial \mathbf{x}_{k+1}}{\partial \mathbf{x}_k} \right\|}{\left\| \frac{\partial \mathcal{E}}{\partial \mathbf{x}_{k+1}} \right\|} - 1 \right)^2 \quad (9)$$

In order to be computationally efficient, we only use the “immediate” partial derivative of Ω with respect to \mathbf{W}_{rec} (we consider that \mathbf{x}_k and $\frac{\partial \mathcal{E}}{\partial \mathbf{x}_{k+1}}$ as being constant with respect to \mathbf{W}_{rec} when computing the derivative of Ω_k), as depicted in equation (10). Note we use the parametrization of equation (11). This can be done efficiently because we get the values of $\frac{\partial \mathcal{E}}{\partial \mathbf{x}_k}$ from BPTT. We use Theano to compute these gradients (Bergstra *et al.*, 2010; Bastien *et al.*, 2012).

$$\begin{aligned} \frac{\partial^+ \Omega}{\partial \mathbf{W}_{rec}} &= \sum_k \frac{\partial^+ \Omega_k}{\partial \mathbf{W}_{rec}} \\ &= \sum_k \frac{\partial^+ \left(\frac{\left\| \frac{\partial \mathcal{E}}{\partial \mathbf{x}_{k+1}} \mathbf{W}_{rec}^T \text{diag}(\sigma'(\mathbf{x}_k)) \right\|}{\left\| \frac{\partial \mathcal{E}}{\partial \mathbf{x}_{k+1}} \right\|} - 1 \right)}{\partial \mathbf{W}_{rec}} \end{aligned} \quad (10)$$

Note that our regularization term only forces the Jacobian matrices $\frac{\partial \mathbf{x}_{k+1}}{\partial \mathbf{x}_k}$ to preserve norm in the relevant

direction of the error $\frac{\partial \mathcal{E}}{\partial \mathbf{x}_{k+1}}$, not for any direction (i.e. we do not enforce that all eigenvalues are close to 1). The second observation is that we are using a soft constraint, therefore we are not ensured the norm of the error signal is preserved. If it happens that these Jacobian matrices are such that the norm explodes (as $t-k$ increases), then this could lead to the exploding gradients problem and we need to deal with it for example as described in section 3.2. This can be seen from the dynamical systems perspective as well: preventing vanishing gradients implies that we are pushing the model such that it is further away from the attractor (such that it does not converge to it, case in which the gradients vanish) and closer to boundaries between basins of attractions, making it more probable for the gradients to explode.

4. Experiments and Results

4.1. Pathological synthetic problems

As done in Martens and Sutskever (2011), we address the pathological problems proposed by Hochreiter and Schmidhuber (1997) that require learning long term correlations. We refer the reader to this original paper for a detailed description of the tasks and to the supplementary materials for the complete description of the experimental setup.

4.1.1. THE TEMPORAL ORDER PROBLEM

We consider the temporal order problem as the prototypical pathological problem, extending our results to the other proposed tasks afterwards. The input is a long stream of discrete symbols. At two points in time (in the beginning and middle of the sequence) a symbol within $\{A, B\}$ is emitted. The task consists in classifying the order (either AA, AB, BA, BB) at the end of the sequence.

Fig. 7 shows the success rate of standard SGD, SGD-C (SGD enhanced with out clipping strategy) and SGD-CR (SGD with the clipping strategy and the regularization term). Note that for sequences longer than 20, the vanishing gradients problem ensures that neither SGD nor SGD-C algorithms can solve the task. The x -axis is on log scale.

This task provides empirical evidence that exploding gradients are linked with tasks that require long memory traces. We know that initially the model operates in the one-attractor regime (i.e. $\lambda_1 < 1$), in which the amount of memory is controlled by λ_1 . More memory means larger spectral radius, and, when this value crosses a certain threshold the model enters rich regimes where gradients are likely to explode. We see in Fig. 7 that as long as the vanishing gradient prob-

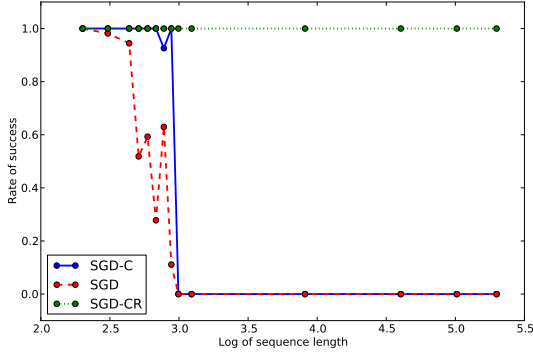


Figure 7. Rate of success for solving the temporal order problem versus log of sequence length. See text.

lem does not become an issue, addressing the exploding gradients problem ensures a better success rate.

When combining clipping as well as the regularization term proposed in section 3.3, we call this algorithm SGD-CR. SGD-CR solved the task with a success rate of 100% for sequences up to 200 steps (the maximal length used in Martens and Sutskever (2011)). Furthermore, we can train a single model to deal with any sequence of length 50 up to 200 (by providing sequences of different lengths for different SGD steps). Interestingly enough, **the trained model can generalize to new sequences that can be twice as long as the ones seen during training.**

4.1.2. OTHER PATHOLOGICAL TASKS

SGD-CR was also able to solve (100% success on the lengths listed below, for all but one task) other pathological problems proposed in Hochreiter and Schmidhuber (1997), namely the *addition* problem, the *multiplication* problem, the *3-bit temporal order problem*, the *random permutation* problem and the *noiseless memorization* problem in two variants (when the pattern needed to be memorized is 5 bits in length and when it contains over 20 bits of information; see Martens and Sutskever (2011)). For the first 4 problems we used a single model for lengths up to 200, while for the *noiseless memorization* we used a different model for each sequence length (50, 100, 150 and 200). The hardest problems for which only one trail out of 8 succeeded was the random permutation problem. In all cases, we observe successful generalization to sequences longer than the training sequences. In most cases, these results outperforms Martens and Sutskever (2011) in terms of success rate, they deal with longer sequences than in Hochreiter and Schmidhuber (1997) and compared to (Jaeger, 2012) they generalize to longer sequences.

Table 1. Results on polyphonic music prediction in negative log likelihood per time step. Lower is better.

DATA SET	DATA FOLD	SGD	SGD+C	SGD+CR
PIANO-MIDI.DE	TRAIN	6.87	6.81	7.01
	TEST	7.56	7.53	7.46
NOTTINGHAM	TRAIN	3.67	3.21	3.24
	TEST	3.80	3.48	3.46
MUSEDATA	TRAIN	8.25	6.54	6.51
	TEST	7.11	7.00	6.99

Table 2. Results on the next character prediction task in entropy (bits/character)

DATA SET	DATA FOLD	SGD	SGD+C	SGD+CR
1 STEP	TRAIN	1.46	1.34	1.36
	TEST	1.50	1.42	1.41
5 STEPS	TRAIN	N/A	3.76	3.70
	TEST	N/A	3.89	3.74

4.2. Natural problems

We address the task of polyphonic music prediction, using the datasets Piano-midi.de, Nottingham and MuseData described in Boulanger-Lewandowski *et al.* (2012) and language modelling at the character level on the Penn Treebank dataset (Mikolov *et al.*, 2012). We also explore a modified version of the task, where we ask the model to predict the 5th character in the future (instead of the next). Our assumption is that to solve this modified task long term correlations are more important than short term ones, and hence our regularization term should be more helpful.

The training and test scores reported in Table 1 are average negative log likelihood per time step. We fixed hyper-parameters across the three runs, except for the regularization factor and clipping cutoff threshold. **SGD-CR provides a statistically significant improvement on the state-of-the-art for RNNs on all the polyphonic music prediction tasks except for MuseData on which we get exactly the same performance as the state-of-the-art (Bengio *et al.*, 2012), which uses a different architecture.** Table 2 contains the results on language modelling (in bits per letter).

These results suggest that clipping the gradients solves an optimization issue and does not act as a regularizer, as both the training and test error improve in general. Results on Penn Treebank reach the state of the art achieved by Mikolov *et al.* (2012), who used a different clipping algorithm similar to ours, thus providing evidence that both behave similarly. The regularized model performs as well as the Hessian-Free trained model.

By employing the proposed regularization term we are able to improve test error even on tasks that are not

dominated by long term contributions.

5. Summary and Conclusions

We provided different perspectives through which one can gain more insight into the *exploding and vanishing gradients* issue. To deal with the exploding gradients problem, we propose a solution that involves clipping the norm of the exploded gradients when it is too large. The algorithm is motivated by the assumption that when gradients explode, the curvature and higher order derivatives explode as well, and we are faced with a specific pattern in the error surface, namely a valley with a single steep wall. In order to deal with the vanishing gradient problem we use a regularization term that forces the error signal not to vanish as it travels back in time. This regularization term forces the Jacobian matrices $\frac{\partial \mathbf{x}_i}{\partial \mathbf{x}_{i-1}}$ to preserve norm only in relevant directions. In practice we show that these solutions improve performance on both the pathological synthetic datasets considered as well as on polyphonic music prediction and language modelling.

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Analytical analysis of the exploding and vanishing gradients problem

$$\mathbf{x}_t = \mathbf{W}_{rec}\sigma(\mathbf{x}_{t-1}) + \mathbf{W}_{in}\mathbf{u}_t + \mathbf{b} \quad (11)$$

Let us consider the term $\mathbf{g}_k^T = \frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} \frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k} \frac{\partial^+ \mathbf{x}_k}{\partial \theta}$ for the linear version of the parametrization in equation (11) (i.e. set σ to the identity function) and assume t goes to infinity and $l = t - k$. We have that:

$$\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k} = (\mathbf{W}_{rec}^T)^l \quad (12)$$

By employing a generic *power iteration method* based proof we can show that, given certain conditions, $\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} (\mathbf{W}_{rec}^T)^l$ grows exponentially.

Proof Let \mathbf{W}_{rec} have the eigenvalues $\lambda_1, \dots, \lambda_n$ with $|\lambda_1| > |\lambda_2| > \dots > |\lambda_n|$ and the corresponding eigenvectors $\mathbf{q}_1, \mathbf{q}_2, \dots, \mathbf{q}_n$ which form a vector basis. We can now write the row vector $\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t}$ into this basis:

$$\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} = \sum_{i=1}^N c_i \mathbf{q}_i^T$$

If j is such that $c_j \neq 0$ and any $j' < j, c_{j'} = 0$, using the fact that $\mathbf{q}_i^T (\mathbf{W}_{rec}^T)^l = \lambda_i^l \mathbf{q}_i^T$ we have that

$$\frac{\partial \mathcal{E}_t}{\partial \mathbf{x}_t} \frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k} = c_j \lambda_j^l \mathbf{q}_j^T + \lambda_j^l \sum_{i=j+1}^n c_i \frac{\lambda_i^l}{\lambda_j^l} \mathbf{q}_i^T \approx c_j \lambda_j^l \mathbf{q}_j^T \quad (13)$$

We used the fact that $|\lambda_i/\lambda_j| < 1$ for $i > j$, which means that $\lim_{l \rightarrow \infty} |\lambda_i/\lambda_j|^l = 0$. If $|\lambda_j| > 1$, it follows that $\frac{\partial \mathbf{x}_t}{\partial \mathbf{x}_k}$ grows exponentially fast with l , and it does so along the direction \mathbf{q}_j . \square

The proof assumes \mathbf{W}_{rec} is diagonalizable for simplicity, though using the Jordan normal form of \mathbf{W}_{rec} one can extend this proof by considering not just the eigenvector of largest eigenvalue but the whole subspace spanned by the eigenvectors sharing the same (largest) eigenvalue.

This result provides a necessary condition for gradients to grow, namely that the spectral radius (the absolute value of the largest eigenvalue) of \mathbf{W}_{rec} must be larger than 1.

If \mathbf{q}_j is not in the null space of $\frac{\partial^+ \mathbf{x}_k}{\partial \theta}$ the entire temporal component grows exponentially with l . This approach extends easily to the entire gradient. If we re-write it in terms of the eigen-decomposition of \mathbf{W} , we get:

$$\frac{\partial \mathcal{E}_t}{\partial \theta} = \sum_{j=1}^n \left(\sum_{i=k}^t c_j \lambda_j^{t-i} \mathbf{q}_j^T \frac{\partial^+ \mathbf{x}_i}{\partial \theta} \right) \quad (14)$$

We can now pick j and k such that $c_j \mathbf{q}_j^T \frac{\partial^+ \mathbf{x}_k}{\partial \theta}$ does not have 0 norm, while maximizing $|\lambda_j|$. If for the chosen j it holds that $|\lambda_j| > 1$ then $\lambda_j^{t-k} c_j \mathbf{q}_j^T \frac{\partial^+ \mathbf{x}_k}{\partial \theta}$ will dominate the sum and because this term grows exponentially fast to infinity with t , the same will happen to the sum.

Experimental setup

Note that all hyper-parameters were selected based on their performance on a validation set using a grid search.

The pathological synthetic tasks

Similar success criteria is used in all of the tasks below (borrowed from Martens and Sutskever (2011)), namely that the model should make no more than 1% error on a batch of 10000 test samples. In all cases, discrete symbols are depicted by a one-hot encoding, and in case of regression a prediction for a given sequence is considered as a success if the error is less than 0.04.

ADDITION PROBLEM

The input consists of a sequence of random numbers, where two random positions (one in the beginning and one in the middle of the sequence) are marked. The model needs to predict the sum of the two random numbers after the entire sequence was seen. For each generated sequence we sample the length T' from $[T, \frac{11}{10}T]$, though for clarity we refer to T as the length of the sequence in the paper. The first position is sampled from $[1, \frac{T'}{10}]$, while the second position is sampled from $[\frac{T'}{10}, \frac{T'}{2}]$. These positions i, j are marked in a different input channel that is 0 everywhere except for the two sampled positions when it is 1. The model needs to predict the sum of the random numbers found at the sampled positions i, j divided by 2.

To address this problem we use a 50 hidden units model, with a tanh activation function. The learning rate is set to .01 and the factor α in front of the regularization term is 0.5. We use clipping with a cut-off threshold of 6 on the norm of the gradients. The weights are initialized from a normal distribution with mean 0 and standard derivation .1.

The model is trained on sequences of varying length T between 50 and 200. We manage to get a success rate of 100% at solving this task, which outperforms the results presented in Martens and Sutskever (2011) (using Hessian Free), where we see a decline in success rate as the length of the sequence gets closer to 200

steps. Hochreiter and Schmidhuber (1997) only considers sequences up to 100 steps. Jaeger (2012) also addresses this task with 100% success rate, though the solution does not seem to generalize well as it relies on very large output weights, which for ESNs are usually a sign of instability. We use a single model to deal with all lengths of sequences (50, 100, 150 200), and the trained model generalizes to new sequences that can be up 400 steps (while the error is still under 1%).

MULTIPLICATION PROBLEM

This task is similar to the problem above, just that the predicted value is the product of the random numbers instead of the sum. We used the same hyper-parameters as for the previous case, and obtained very similar results.

TEMPORAL ORDER PROBLEM

For the temporal order the length of the sequence is fixed to T . We have a fixed set of two symbols $\{A, B\}$ and 4 distractor symbols $\{c, d, e, f\}$. The sequence entries are uniformly sampled from the distractor symbols everywhere except at two random positions, the first position sampled from $[\frac{T}{10}, \frac{2T}{10}]$, while the second from $[\frac{4T}{10}, \frac{5T}{10}]$. The task is to predict the order in which the non-distractor symbols were provided, i.e. either $\{AA, AB, BA, BB\}$.

We use a 50 hidden units model, with a learning rate of .001 and α , the regularization coefficient, set to 2. The cut-off threshold for clipping the norm of the gradient is left to 6. As for the other two task we have a 100% success rate at training a single model to deal with sequences between 50 to 200 steps. This outperforms the previous state of the art because of the success rate, but also the single model generalizes to longer sequences (up to 400 steps).

3-BIT TEMPORAL ORDER PROBLEM

Similar to the previous one, except that we have 3 random positions, first sampled from $[\frac{T}{10}, \frac{2T}{10}]$, second from $[\frac{3T}{10}, \frac{4T}{10}]$ and last from $[\frac{6T}{10}, \frac{7T}{10}]$.

We use similar hyper-parameters as above, but that we increase the hidden layer size to 100 hidden units. As before we outperform the state of the art while training a single model that is able to generalize to new sequence lengths.

RANDOM PERMUTATION PROBLEM

In this case we have a dictionary of 100 symbols. Except the first and last position which have the same value sampled from $\{1, 2\}$ the other entries are ran-

domly picked from $[3, 100]$. The task is to do next symbol prediction, though the only predictable symbol is the last one.

We use a 100 hidden units with a learning rate of .001 and α , the regularization coefficient, set to 1. The cutoff threshold is left to 6. This task turns out to be more difficult to learn, and only 1 out of 8 experiments succeeded. As before we use a single model to deal with multiple values for T (from 50 to 200 units).

NOISELESS MEMORIZATION PROBLEM

For the noiseless memorization we are presented with a binary pattern of length 5, followed by T steps of constant value. After these T steps the model needs to generate the pattern seen initially. We also consider the extension of this problem from Martens and Sutskever (2011), where the pattern has length 10, and the symbol set has cardinality 5 instead of 2.

We manage a 100% success rate on these tasks, though we train a different model for the 5 sequence lengths considered (50, 100, 150, 200).

Natural Tasks

POLYPHONIC MUSIC PREDICTION

We train our model, a sigmoid units RNN, on sequences of 200 steps. The cut-off coefficient threshold is the same in all cases, namely 8 (note that one has to take the mean over the sequence length when computing the gradients).

In case of the Piano-midi.de dataset we use 300 hidden units and an initial learning rate of 1.0 (whir the learning rate halved every time the error over an epoch increased instead of decreasing). For the regularized model we used a initial value for regularization coefficient α of 0.5, where α follows a $1/t$ schedule, i.e. $\alpha_t = \frac{1}{2t}$ (where t measures the number of epochs).

For the Nottingham dataset we used the exact same setup. For MuseData we increased the hidden layer to 400 hidden units. The learning rate was also decreased to 0.5. For the regularized model, the initial value for α was 0.1, and $\alpha_t = \frac{1}{t}$.

We make the observation that for natural tasks it seems useful to use a schedule that decreases the regularization term. We assume that the regularization term forces the model to focus on long term correlations at the expense of short term ones, so it may be useful to have this decaying factor in order to allow the model to make better use of the short term information.

LANGUAGE MODELLING

For the language modelling task we used a 500 sigmoidal hidden units model with no biases (Mikolov *et al.*, 2012). The model is trained over sequences of 200 steps, where the hidden state is carried over from one step to the next one.

We use a cut-off threshold of 45 (though we take the sum of the cost over the sequence length) for all experiments. For next character prediction we have a learning rate of 0.01 when using clipping with no regularization term, 0.05 when we add the regularization term and 0.001 when we do not use clipping. When predicting the 5th character in the future we use a learning rate of 0.05 with the regularization term and 0.1 without it.

The regularization factor α for next character prediction was set to .01 and kept constant, while for the modified task we used an initial value of 0.05 with a $\frac{1}{t}$ schedule.