# Deep Learning Book

Chapter 7 Regularization for Deep Learning

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- How to make an algorithm that will perform well not just on the training data, but also on new inputs?
- Many strategies designed to reduce the test error, possibly at the expense of increased training error.
- · These strategies are known collectively as regularization.
- · Many regularization algorithm have been developed.
- Developing more effective regularization strategies is one of the major research efforts in the field.
- In this chapter, we describe regularization in more detail, focusing on regularization strategies for deep models or models that may be used as building blocks to form deep models.

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- There are many regularization strategies.
  - Put extra constrains on a machine learning model. (Adding restrictions on the parameter values.)
  - 2. Add extra terms in the objective function that can be thought of as corresponding to a soft constraint on the parameter values.
- If chosen carefully, these extra constraints and penalties can lead to improved performance on the test set.
- · Sometimes these constraints and penalties are designed to
  - 1. encode specific kinds of prior knowledge.
  - 2. Express a generic preference for a simpler model class in order to promote generalization.
  - make an under-determined problem determined. (Provide more information)
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- An effective regularizer is one that makes a profitable trade, reducing variance significantly while not overly increasing the bias.
- In practice, an overly complex model family does not necessarily include the target function or the true data generating process, or even a close approximation.
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- However, most applications of deep learning algorithms are to domains where the true data generating process is almost certainly outside the model family.
- Deep learning algorithms are typically applied to extremely complicated domains such as images, audio sequences and text, for which the true generation process essentially involves simulating the entire universe.
- To some extent, we are always trying to fit a square peg(the data generating process) into a round hole (our model family)
   『持方枘 (ruì) 而欲内圆凿』.
- What this means is that controlling the complexity of the model is not a simple matter of finding the model of the right size, with the right number of parameters.
- Insteamd, we might find that the best fitting model is a large model that has been regularized appropriately.
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- Regularization has been used for decades prior to the advent of deep learning.
- Linear models allow simple straightforward and effective regularization strategies.
- Most approaches are based on limiting the capacity of models by adding a parameter norm penalty  $\Omega(\theta)$  to the objective function J:

$$J(\boldsymbol{\theta}; X, y) = J(\boldsymbol{\theta}; X, y) + \alpha \Omega(\boldsymbol{\theta})$$

- Setting  $\alpha$  to 0 results in no regularization. Larger values of  $\alpha$  correspond to more regularization.
- · Optimize both J and norm
- $\cdot$  Different  $\Omega$  has different result

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- We penalize **only the weights** of the affine transformation at each layer and leaves the biases unregularized.
- We do not induce too much variance by leaving the biases unregularized.
- Regularizing the bias parameters can introduce a significant amount of under-fitting.
- We therefore use the vector w to indicate all of the weights that should be affected by a norm penalty, while the vector θ denotes all of the parameters, including both w and the unregularized parameters.
- Sometime we use a separate penalty with a different  $\alpha$  coefficient for each layer.
- But it can be expensive to search for the correct value of multiple hyper-parameters, it is still reasonable to use the same weight decay at all layers just to reduce the size of search space.

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# L<sup>2</sup> Parameter Regularization

• The  $L^2$  norm penalty commonly known as weight decay.

$$\Omega(\boldsymbol{\theta}) = \frac{1}{2} \|\mathbf{w}\|_2^2$$

- This regularization strategy drives the weights closer to the origin. (as well as *ridge regression* or *Tikhonov regularization*)
- We can gain some insight into the behavior of weight decay regularization. (assume no bias for simplification)

$$\tilde{J}(w; X, y) = \frac{\alpha}{2} w^{\mathsf{T}} w + J(w; X, y)$$
$$\nabla_{w} \tilde{J}(w; X, y) = \alpha w + \nabla_{w} J(w; X, y)$$

The update

$$w \leftarrow w - \epsilon(\alpha w + \nabla_w J(w; X, y))$$
  
$$w \leftarrow (1 - \epsilon \alpha) w - \epsilon \nabla_w J(w; X, y)$$

- Shrink the weight vector by a constant factor on each step.
- · What happens over the entire course of training?

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- In mathematics, approximation theory is concerned with how functions can best be approximated with simpler functions.
- · local linear approximation and taylor expansion
  - 1. For example, when the independent variable of function  $y = x^3$  changes, which is  $\Delta x$ , the variation of y is

$$\Delta y = (x + \Delta x)^3 - x^3 = 3x^2 \Delta x + 3x(\Delta x)^2 + (\Delta x)^3$$

- 2. When  $\Delta x \rightarrow 0$ , omit last two terms:  $\Delta y = 3x^2 \Delta x$
- 3. In general:

$$\Delta y = f(x_0 + \Delta x) - f(x_0) \approx f'(x_0) \times \Delta x$$

$$\Delta y = f(x) - f(x_0), \ \Delta x = x - x_0$$

$$f(x) - f(x_0) = f'(x_0) \times (x - x_0)$$

$$f(x) = f(x_0) + f'(x_0)(x - x_0)$$

4. In order to improve the precision, we can use second-order approximation, which is the second-order Taylor series expansion.

$$f(x) = f(x_0) + \frac{f'(x_0)}{1!}(x - x_0) + \frac{f''(x_0)}{2!}(x - x_0)^2$$

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$$\Delta y = f(x) - f(x_0), \ \Delta x = x - x_0$$
  

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$$f(x) = f(x_0) + f'(x_0)(x - x_0)$$

 In order to improve the precision, we can use second-order approximation, which is the second-order Taylor series expansion

$$f(x) = f(x_0) + \frac{f'(x_0)}{1!}(x - x_0) + \frac{f''(x_0)}{2!}(x - x_0)^2$$

- In mathematics, approximation theory is concerned with how functions can best be approximated with simpler functions.
- · local linear approximation and taylor expansion
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- Let  $w^* = \arg \min_{w} J(w)$  (unregularized training cost)
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$$\hat{J}(w) = J(w^*) + \frac{1}{2}(w - w^*)^T H(w - w^*)$$

- Where **H** is the Hessian matrix of *J* with respect to **w** evaluated at **w**\*.
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- · We use the variable  $ilde{w}$  to represent the location of the minimum

$$\alpha \tilde{w} + H(\tilde{w} - w^*) = 0$$

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$$\tilde{w} = \frac{Hw^*}{(H + \alpha I)}$$

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- Because **H** is real and symmetric, we can decompose it into a diagonal matrix  $\Lambda$  and an orthonormal basis of eigenvectors, Q, such that  $\mathbf{H} = Q\Lambda Q^{\mathsf{T}}$ .
- Applying the decomposition  $\tilde{\mathbf{w}} = (\mathbf{H} + \alpha \mathbf{I})^{-1}\mathbf{H}\mathbf{w}^*$

$$\tilde{W} = (Q\Lambda Q^{T} + \alpha I)^{-1}Q\Lambda Q^{T}W^{*} 
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- We see that the effect of weight decay is to rescale w\* along the axes defined by the eigenvectors of H.
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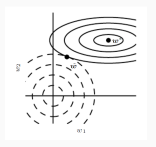
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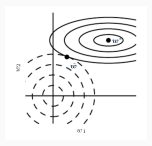
Fig. 1: An illustration of the effect of  $L^2$  (or weight decay) regularization on the value of the optimal  $\mathbf{w}$ 



- The solid ellipses represent contours of equal value of the unregularized objective.
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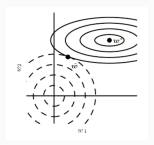
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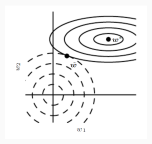
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- How do these effects relate to machine learning in particular?
- We can find out by studying linear regression, the cost function is the sum of squared errors:

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- The new matrix has the addition of  $\alpha$  to the diagonal.
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• L¹ regularization on the model parameter w is defined as:

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- We will now discuss the effect of  $L^1$  regularization on the simple linear regression model, with no bias parameters, that we studied in our analysis of  $L^2$  regularization.
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- Out simple linear model has a quadratic cost function that we can represent via its Taylor series.
- Alternately, we could imaging that this is a truncated Taylor series approximating the cost function of a more sophisticated model.
- · The gradient in this setting is given by

$$\nabla_{w}\widetilde{J}(w) = H(w - w^*)$$

- Because the  $L^1$  penalty does not admit clean algebraic expressions in the case of a full general Hessian, we will also make the further simplifying assumption that the Hessian is a diagonal,  $\mathbf{H} = \operatorname{diag}([H_{1,1}, \ldots, H_{n,n}])$ , where each  $H_{i,i} > 0$ .
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• Our quadratic approximation of the *L*<sup>1</sup> regularized objective function decomposes into a sum over the parameters:

$$\tilde{J}(\boldsymbol{w};\boldsymbol{X},\boldsymbol{y}) = J(\boldsymbol{w}^*;\boldsymbol{X},\boldsymbol{y}) + \sum_{i} \left[ \frac{1}{2} H_{i,i} (\boldsymbol{w}_i - \boldsymbol{w}_i^*)^2 + \alpha |w_i| \right]$$

 The problem of minimizing this approximate cost function has an analytical solution (for each dimension i), with the following form:

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- Consider the situation where w<sub>i</sub>\* > 0 for all i. There are two
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  - 1. The case where  $w_i^* \leq \frac{\alpha}{H_{i,i}}$ . Here the optimal value of  $w_i$  under the regularized objective is simply  $w_i = 0$ . This occurs because the contribution of J(w; X, y) to the regularized objective  $\tilde{J}(w; X, y)$  is overwhelmed—in direction i-by the  $L^1$  regularization which pushes the value of  $w_i$  to zero.
  - 2. The case where  $w_i^* > \frac{\alpha}{H_{i,i}}$ . In this case, the regularization does not move the optimal value of  $w_i$  to zero but instead it just shifts it in that direction by a distance equal to  $\frac{\alpha}{H_{i,i}}$ .
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- In comparison to  $L^2$  regularization,  $L^1$  regularization results in a solution that is more *sparse*.
- Sparsity in this context refers to the fact that some parameters have an optimal value of zero.
- The sparsity property induced by L<sup>1</sup> regularization has been used extensively as a feature selection mechanism.
- Feature selection simplifies a machine learning problem by choosing which subset of the available features should be used.
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# Sparsity? $L^1$ and $L^2$

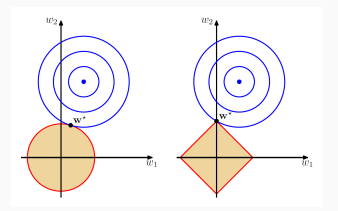


Fig. 2: Plot of the contours of the unregularized error function (blue) along with the constraint region for the quadratic regularizer on the left and the lasso regularizer on the right.

 Consider the cost function regularized by a parameter norm penalty:

$$\tilde{J}(\boldsymbol{\theta}; \mathbf{X}, \mathbf{y}) = J(\boldsymbol{\theta}; \mathbf{X}, \mathbf{y}) + \alpha \Omega(\boldsymbol{\theta})$$

• If we want to constrain  $\Omega(\theta)$  to be less than some constant k, we could construct a generalized Language function

$$\mathcal{L}(\boldsymbol{\theta}, \boldsymbol{\alpha}; X, y) = J(\boldsymbol{\theta}; X, y) + \alpha(\Omega(\boldsymbol{\theta}) - k)$$

The solution to the constrained problem is given by

$$oldsymbol{ heta}^* = rg\min_{oldsymbol{ heta}} \max_{lpha,lpha \geq 0} \mathcal{L}(oldsymbol{ heta},oldsymbol{lpha})$$

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- Solving this problem requires modifying both  $\theta$  and  $\alpha$ .
- Many different procedures are possible–some may use gradient descent, while others may use analytical solutions for where the gradient is zero–but in all procedures  $\alpha$  must increase whenever  $\Omega(\theta) > k$  and decrease whenever  $\Omega(\theta) < k$ .
- · All positive lpha encourage  $\Omega( heta)$  to shrink
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- Many linear models in machine learning, including linear regression and PCA, depend on inverting the matrix  $X^TX$ .
- This is not possible whenever  $X^TX$  is singular.
- This matrix can be singular whenever the data generating distribution truly has no variance in some direction, or when no variance in observed in some direction because there are fewer examples (rows of X) than input features (columns of X).
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- An example is logistic regression applied to a problem where the classes are linearly separable.
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- · In practice, it is limited.
- Create fake data and add it to the training set.
- This approach is easiest for classification.
- A classifier needs to take a complicated, high dimensional input
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$$J = \mathbb{E}_{p(x,y)} \left[ (\hat{y}(x) - y)^2 \right]$$

- The training set with m examples:  $(\mathbf{x}^{(1)}, y^{(1)}), \dots, (\mathbf{x}^{(m)}, y^{(m)})$ .
- We now assume that with each input presentation we also include a random perturbation  $\epsilon_W \mathcal{N}(\epsilon; \mathbf{0}, \eta \mathbf{I})$  of the network weights.
- We denote the perturbed model as  $\hat{y}_{\epsilon_W}(x)$ . The objective function thus becomes:

$$\tilde{J}_{W} = \mathbb{E}_{\rho(\mathbf{x}, y, \epsilon_{W})} \left[ (\hat{y}_{\epsilon_{W}}(\mathbf{x}) - y)^{2} \right] 
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• We study the regression setting, where we wish to train a function  $\tilde{y}(x)$  that maps a set of features x to a scalar using the least-squares cost function between the model predictions  $\tilde{y}(x)$  and the true values y:

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- This form of regularization encourages the parameters to go to regions of parameter space where small perturbations of the weights have a relatively small influence on the output.
- In other words, it pushes the model weights, finding points that are not merely minimal, but minimal surrounded by flat regions (Hochreiter et al. [1995]).

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- · Most datasets have some amount of mistakes in the y labels.
- It can be harmful to maximize  $\log p(y|\mathbf{x})$  when y is a mistake.
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- In the paradigm of semi-supervised learning, both unlabeled examples from P(x) and labeled examples from P(x, y) are used to estimate P(y|x) or predict y from x.
- In the context of deep learning, semi-supervised learning usually refers to learning a representation h = f(x). The goal is to learn a representation so that examples from the same class have similar representations.
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- One can construct models in which a generative model of either (x) or P(x,y) shares parameters with a discriminative model of P(y|x).
- The generative criterion then express a particular form of prior belief about the solution to the supervised learning problem, namely that the structure of P(x) is connected to the structure of P(y|x) in a way that is captured by the shared parameterization.
- By controlling how much of the generative criterion is included in the total criterion, one can find a better trade-off than with a purely generative or purely discriminative training criterion.
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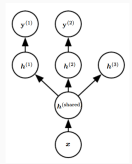
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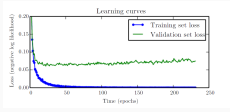
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- · Here is a very common form of multi-task learning.
- Different supervised tasks (predicting  $y^{(i)}$  given x) share the same input x, as well as some intermediate-level representation  $h^{\text{(shared)}}$  capturing a common pool of factors.
- · The model has two kinds of parts:
  - Task-specific parameters (which only benefit from the examples of their task to achieve good generalization). These are the upper layers.
  - Generic parameters, shared across all the tasks (which benefit from the pooled data of all the tasks). These are the lower years.



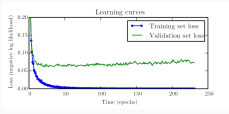
 The factors that explain the variations are shared across two or more tasks.

 When training large models with sufficient representational capacity to overfit the task, we often observe that training error decreases steadily over time, but validation set error begins to rise again.



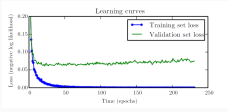
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### Algorithm 1 Early Stopping Algorithm

### Algorithm 2 Early Stopping Algorithm

#### Let *n* be the number of steps between evaluations.

Let *p* be the "patience", the number of times to observe worsening validation set error before giving up.

Let  $\theta_o$  be the initial parameters.

$$\theta \leftarrow \theta_o$$
;  $i \leftarrow 0$ ;  $j \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow v$   
while  $i < p$  do

Update  $\theta$  by running the training algorithm for n steps.

$$i \leftarrow i + n; v' \leftarrow ValidationSetError(\theta)$$

$$J \leftarrow 0; \theta^* \leftarrow \theta; I^* \leftarrow I; V \leftarrow V$$

$$J \leftarrow J + 1$$

nd while

### Algorithm 3 Early Stopping Algorithm

Let n be the number of steps between evaluations.

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#### Algorithm 4 Early Stopping Algorithm

Let n be the number of steps between evaluations.

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Let  $heta_o$  be the initial parameters.

```
m{	heta} \leftarrow m{	heta}_o; i \leftarrow 0; j \leftarrow 0; v \leftarrow \infty; i^* \leftarrow i
while j < p do

Update m{	heta} by running the training algorithm for n
i \leftarrow i + n; v' \leftarrow \text{ValidationSetError}(m{	heta})
if v' < v then
j \leftarrow 0; m{	heta}^* \leftarrow m{	heta}; i^* \leftarrow i; v \leftarrow v'
else
j \leftarrow j + 1
end if
```

### Algorithm 5 Early Stopping Algorithm

Let n be the number of steps between evaluations.

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$$\theta \leftarrow \theta_{o}$$
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while  $j < p$  do  
Update  $\theta$  by running the training algorithm is the product Validation Set Error (4)

if v' < v then

$$j \leftarrow 0$$
;  $\theta^* \leftarrow \theta$ ;  $i^* \leftarrow i$ ;  $v \leftarrow v'$ 

else

$$j \leftarrow j + 1$$

end

nd while

### Algorithm 6 Early Stopping Algorithm

Let n be the number of steps between evaluations.

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$$\theta \leftarrow \theta_0$$
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$$i \leftarrow i + n; v' \leftarrow \text{ValidationSetErr}$$
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else
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#### end while

#### Algorithm 7 Early Stopping Algorithm

Let n be the number of steps between evaluations.

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$$\theta \leftarrow \theta_0$$
;  $i \leftarrow 0$ ;  $j \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow i$  while  $j < p$  do

Update  $\theta$  by running the training algorithm for n steps.

```
i \leftarrow i + n; v' \leftarrow ValidationSetError(\theta)
if v' < v then
j \leftarrow 0; \theta^* \leftarrow \theta; j^* \leftarrow i; v \leftarrow v'
else
j \leftarrow j + 1
end if
```

#### end while

### Algorithm 8 Early Stopping Algorithm

Let n be the number of steps between evaluations.

Let *p* be the "patience", the number of times to observe worsening validation set error before giving up.

Let  $\theta_o$  be the initial parameters.

$$\theta \leftarrow \theta_0$$
;  $i \leftarrow 0$ ;  $j \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow i$  while  $i < p$  do

Update  $\theta$  by running the training algorithm for n steps.

$$i \leftarrow i + n; v' \leftarrow ValidationSetError(\theta)$$
  
if  $v' < v$  then  
 $j \leftarrow 0; \theta^* \leftarrow \theta; i^* \leftarrow i; v \leftarrow v'$   
else  
 $j \leftarrow j + 1$ 

#### end while

### Algorithm 9 Early Stopping Algorithm

Let n be the number of steps between evaluations.

Let *p* be the "patience", the number of times to observe worsening validation set error before giving up.

Let  $\theta_o$  be the initial parameters.

$$\theta \leftarrow \theta_0$$
;  $i \leftarrow 0$ ;  $j \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow i$  while  $j < p$  do

Update  $\theta$  by running the training algorithm for n steps.

$$i \leftarrow i + n; v' \leftarrow ValidationSetError(\theta)$$

if v' < v then

$$j \leftarrow 0$$
;  $\theta^* \leftarrow \theta$ ;  $i^* \leftarrow i$ ;  $V \leftarrow V'$ 

else

$$j \leftarrow j + 1$$

end if

end while

### Algorithm 10 Early Stopping Algorithm

Let n be the number of steps between evaluations.

Let *p* be the "patience", the number of times to observe worsening validation set error before giving up.

Let  $\theta_o$  be the initial parameters.

$$\theta \leftarrow \theta_o$$
;  $i \leftarrow 0$ ;  $j \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow i$  while  $j < p$  do

Update  $\theta$  by running the training algorithm for n steps.

$$i \leftarrow i + n$$
;  $v' \leftarrow ValidationSetError(\theta)$   
if  $v' < v$  then

$$j \leftarrow 0$$
;  $\theta^* \leftarrow \theta$ ;  $i^* \leftarrow i$ ;  $v \leftarrow v'$ 

else

$$j \leftarrow j + 1$$

end if

end while

#### Algorithm 11 Early Stopping Algorithm

Let n be the number of steps between evaluations.

Let *p* be the "patience", the number of times to observe worsening validation set error before giving up.

Let  $\theta_o$  be the initial parameters.

$$\theta \leftarrow \theta_o$$
;  $i \leftarrow 0$ ;  $j \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow i$  while  $j < p$  do

Update  $\theta$  by running the training algorithm for n steps.

$$i \leftarrow i + n; v' \leftarrow \text{ValidationSetError}(\theta)$$
  
if  $v' < v$  then  
 $j \leftarrow 0; \theta^* \leftarrow \theta; i^* \leftarrow i; v \leftarrow v'$   
else

$$j \leftarrow j + 1$$

end if

end while

Best parameters are  $\theta^*$ , best number of training steps is  $i^*$ .

#### Algorithm 12 Early Stopping Algorithm

Let *n* be the number of steps between evaluations. Let p be the "patience", the number of times to observe worsening validation set error before giving up. Let  $\theta_0$  be the initial parameters.  $\theta \leftarrow \theta_0$ ;  $i \leftarrow 0$ ;  $i \leftarrow 0$ ;  $v \leftarrow \infty$ ;  $i^* \leftarrow i$ while i < p do Update  $\theta$  by running the training algorithm for n steps.  $i \leftarrow i + n; v' \leftarrow ValidationSetError(\theta)$ if v' < v then  $i \leftarrow 0$ :  $\theta^* \leftarrow \theta$ :  $i^* \leftarrow i$ :  $v \leftarrow v'$ else  $i \leftarrow i + 1$ end if end while Best parameters are  $\theta^*$ , best number of training steps is  $i^*$ .

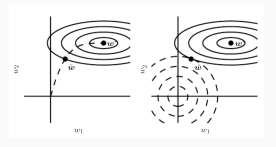
- One way to think of early stopping is as a very efficient hyperparameter selection algorithm.
- In this view, the number of training steps is just another hyperparameter.
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- An additional cost to early stopping is the need to maintain a copy of the best parameters. This cost is generally negligible. (GPU->CPU/MEMORY->HDD).

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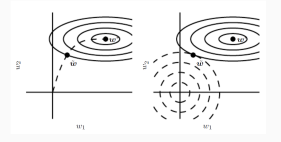
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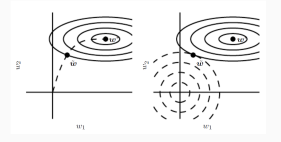
- · How early stopping acts as a regularizer?
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- In order to compare with classical  $L^2$  regularization, we examine a simple setting where the only parameters are linear weights  $(\theta = w)$ .
- We can model the cost function J with a quadratic approximation in the neighborhood of the empirically optimal value of the weights w\*:

$$\hat{J}(\theta) = J(w^*) + \frac{1}{2}(w - w^*)^T H(w - w^*)$$

where H is Hessian matrix of J with respect to w evaluated at  $w^*$ .

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· Under a local Taylor series approximation, the gradient:

$$\nabla_{w} \hat{J}(w) = H(w - w^*)$$

- We are going to study the trajectory followed by the parameter vector during training.
- For simplicity, let us set the initial parameter vector to the origin, that is  $\mathbf{w}^{(0)} = \mathbf{0}$ .
- Let us suppose that we update the parameters via gradient descent:

$$w^{(\tau)} = w^{(\tau-1)} - \epsilon \nabla_{w} J(w^{(\tau-1)})$$

$$= w^{(\tau-1)} - \epsilon H(w^{(\tau-1)} - w^{*})$$

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• Let us now rewrite this expression in the space of the eigenvectors of H, exploiting the eigendecomposition of  $H: H = Q\Lambda Q^T$ , where  $\Lambda$  is a diagonal matrix and Q is an orthonormal basis of eigenvectors.

$$W^{(\tau)} - W^* = (I - \epsilon Q \Lambda Q^{\mathsf{T}})(W^{(\tau-1)} - W^*)$$
  
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• Assuming that  $\mathbf{w}^{(0)} = 0$  and that  $\epsilon$  is chosen to be small enough to guarantee  $|1 - \epsilon \lambda_i| < 1$ , the parameter trajectory during training training after  $\tau$  parameter updates is as follows:

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• In L<sup>2</sup> regularization:

$$\tilde{\mathbf{W}} = \mathbf{Q}(\mathbf{\Lambda} + \alpha \mathbf{I})^{-1} \mathbf{\Lambda} \mathbf{Q}^{\mathsf{T}} \mathbf{W}^* \tag{1}$$

$$Q^{\mathsf{T}}\tilde{\mathbf{w}} = (\Lambda + \alpha \mathbf{I})^{-1}\Lambda Q^{\mathsf{T}}\mathbf{w}^*$$
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$$(I - \epsilon \Lambda)^{\tau} = (\Lambda + \alpha I)^{-1} \alpha$$

- Then *L*<sup>2</sup> regularization and early stopping is equivalent.
- Going even further, by taking logarithms and using the series expansion for  $\log(1+x)$ , if all  $\lambda_i$  are small then:

$$\tau \approx \frac{1}{\epsilon \alpha} \quad ; \quad \alpha \approx \frac{1}{\tau \epsilon}$$
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• That is, under these assumptions, the number of training iterations  $\tau$  plays a role inversely proportional to the  $L^2$  regularization parameter, and the inverse of  $\tau\epsilon$  plays the role of the weight decay coefficient.

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- However, sometimes we may need other ways to express our prior knowledge about suitable values of the model parameters.
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- We have two models performing the same classification task.
- · But with somewhat different input distributions.
- Formally, we have model A with parameters  $\mathbf{w}^{(A)}$  and model B with parameters  $\mathbf{w}^{(B)}$ .
- The two models map the input to different, but related outputs:  $\hat{y}^{(A)} = f(w^{(A)}, x)$  and  $\hat{y}^{(B)} = g(w^{(B)}, x)$ .

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- This kind of approach was proposed by Lasserre et al. [2006], who regularized the parameters of one model, trained as a classifier in a supervised paradigm, to be close to the parameters of another model, trained in an unsupervised paradigm (to capture the distribution of the observed input data).
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- While a parameter norm penalty is one way to regularize parameters to be close to one another, the more popular way is to use constraints: to force sets of parameters to be equal.
- This method of regularization is often referred to as parameter sharing, where we interpret the various models or model components as sharing a unique set of parameters.
- A significant advantage of parameter sharing over regularizing the parameters to be close (via a norm penalty) is that only a subset of the parameters need to be stored in memory.
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- While a parameter norm penalty is one way to regularize parameters to be close to one another, the more popular way is to use constraints: to force sets of parameters to be equal.
- This method of regularization is often referred to as parameter sharing, where we interpret the various models or model components as sharing a unique set of parameters.
- A significant advantage of parameter sharing over regularizing the parameters to be close (via a norm penalty) is that only a subset of the parameters need to be stored in memory.
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- By far the most popular and extensive use of parameter sharing occurs in convolutional neural networks (CNNs) applied to computer vision.
- Natural images have many statistical properties that are invariant to translation.
- CNNs take this property into account by sharing parameters across multiple image locations.
- The same feature (a hidden unit with the same weights) is computed over different locations in the input.
- This means that we can find a object with the same object detector whether the object appears at column i or column i + 1 in the image.
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• A simplified view of this distinction can be illustrated in the context of linear regression:

$$\begin{bmatrix} 18 \\ 5 \\ 15 \\ -9 \\ -3 \\ y \in \mathbb{R}^m \end{bmatrix} = \begin{bmatrix} 4 & 0 & 0 & -2 & 0 & 0 \\ 0 & 0 & -1 & 0 & 3 & 0 \\ 0 & 5 & 0 & 0 & 0 & 0 \\ 1 & 0 & 0 & -1 & 0 & -4 \\ 1 & 0 & 0 & 0 & -5 & 0 \end{bmatrix} \begin{bmatrix} 2 \\ 3 \\ -2 \\ -5 \\ 1 \\ 4 \end{bmatrix}$$

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- Representational regularization is accomplished by the same sorts of mechanisms that we have used in parameter regularization.
- Norm penalty regularization of representation is performed by adding to the loss function J a norm penalty on the representation. This penalty is denoted  $\Omega(h)$ . As before, we denote the regularized loss function by  $\tilde{J}$ :

$$\tilde{J}(\boldsymbol{\theta}; X, y) = J(\boldsymbol{\theta}; X, y) + \alpha \Omega(h)$$

 Just as an L<sup>1</sup> penalty on the parameters induces parameter sparsity, an L<sup>1</sup> penalty on the elements of the representation induces representational sparsity:

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- Others include the penalty derived from a Student-t prior on the representation (Olshausen and Field [2005], Bergstra et al. [2011]) and KL divergence penalties (Larochelle and Bengio [2008]) that are especially useful for representations with elements constrained to lie on the unit interval.
- Lee et al. [2008] and Goodfellow et al. [2009] both provide examples of strategies based on regularizing the average activation across several examples,  $\frac{1}{m} \sum_i \mathbf{h}^{(i)}$ , to be near some target value, such as a vector with 0.01 for each entry.

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$$\underset{h,\|h\|_0 < k}{\operatorname{arg\,min}} \|\mathbf{x} - \mathbf{W}\mathbf{h}\|^2$$

- This problem can be solved efficiently when W is constrained to be orthogonal.
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- The reason that model averaging works is that different models will usually not make all the same errors on the test set.
- · Consider for example a set of k regression models.
- · Suppose that each model makes an error  $\epsilon_i$  on each example, with the errors drawn from a zero-mean multivariate normal distribution with variance  $\mathbb{E}[\epsilon_u^2] = v$  and covariance  $\mathbb{E}[\epsilon_i \epsilon_i] = c$ .
- Then the error made by the average prediction of all the ensemble models is  $\frac{1}{b} \sum_{i} \epsilon_{i}$ .
- The expected squared error of the ensemble predictor is

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- For example, each member of the ensemble could be formed by training a completely different kind of model using a different algorithm or objective function.
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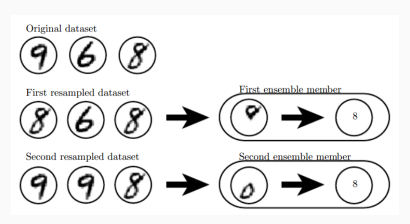


Fig. 3: A cartoon depiction of how bagging works

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- For example, a technique called boosting (Freund et al. [1996]) constructs an ensemble with higher capacity than the individual models.

- Boosting has been applied to build ensembles of neural networks (Schwenk and Bengio [1998]) by incrementally adding neural networks to the ensemble.
- Boosting has also been applied interpreting an individual neural network as an ensemble (Bengio et al. [2006]), incrementally adding hidden units to the neural network.

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- Dropout can be thought of as a method of making bagging practical for ensembles of very many large neural networks.
- Bagging involves training multiple models, and evaluating multiple models on each test example.
- This seems impractical when each model is a large neural network, since training and evaluating such networks is costly in terms of runtime and memory.
- It is common to use ensembles of five to ten neural networks. but more than this rapidly becomes unwieldy.
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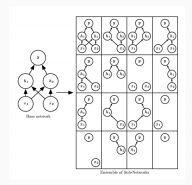
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Specifically, dropout trains the ensemble consisting of all sub-networks that can be formed by removing non-output units from an underlying base network.

- In the most modern neural networks, based on a series of affine transformations and nonlinearities, we can effectively remove a unit from a network by multiplying its output value by zero.
- This procedure requires some slight modification for models such as radial basis function networks, which take the difference between the unit's state and some reference value.



- Here, we present the dropout algorithm in terms of multiplication by zero for simplicity, but it can be trivially modified to work with other operations that remove a unit from the network.
- Recall that to learn with bagging, we define k different models, construct k different datasets by sampling from the training set with replacement, and then train model i on dataset i.
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- Specifically, to train with dropout, we use a minibatch-based learning algorithm that makes small steps, such as stochastic gradient descent.
- Each time we load an example into a minibatch, we randomly sample a different binary mask to apply to all of the input and hidden units in the network.
- The mask for each unit is sampled independently from all of the others.
- The probability of sampling a mask value of one is a hyperparameter fixed before training begins.
- Typically, an input unit is included with probability 0.8 and a hidden unit is included with probability 0.5.
- We then run forward propagation, back-propagation, and the learning update as usual.

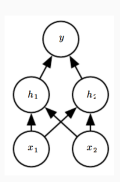
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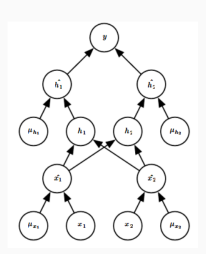
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- More formally, suppose that a mask vector  $\mu$  specifies which units to include, and  $J(\theta, \mu)$  defines the cost of the model defined by parameters  $\theta$  and mask  $\mu$ .
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- To make a prediction, a bagged ensemble must accumulate votes from all of its members. We refer to this process as inference in this context.
- Now, we assume that the model's role is to output a probability distribution.
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- However, there is an even better approach, that allows us to obtain a good approximation to the predictions of the entire ensemble, at the cost of only one forward propagation.
- To do so, we change to using the geometric mean rather than the arithmetic mean of the ensemble member's prediction distributions.
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- The geometric mean of multiple probability distributions is not guaranteed to be a probability distribution.
- To guarantee that the result is a probability distribution, we impose the requirement that none of the sub-models assigns probability 0 to any event, and we renormalize the resulting distribution.
- The unnormalized probability distribution defined directly by the geometric mean is given by

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- Here we use a uniform distribution over  $\mu$  to simplify the presentation, but non-uniform distributions are also possible.
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