# **Ensembles**

Following our Recipe for Machine Learning, we may try out several models before deciding on the final one.

Is a single "best" model really best? Is there an alternative?

By combining models with independent errors, we may be able to construct a combined model whose accuracy is better than the best individual model.

The combined models are called an *Ensemble*.

The individual models

- May be of different types:
  - Decision Tree, Logistic Regression, KNN
- May be of the *same* type. Models differ by
  - different parameters/hyper-parameters:
    - Decision Trees of different depths or different features
    - Regression with polynomial features of different degrees
  - training datasets -subsets of full dataset

When the individual models are of the same type

- Each individual model is trained on a *different* subset of the training examples
- This enables the individual models to produce different results
- Makes them more robust to outliers

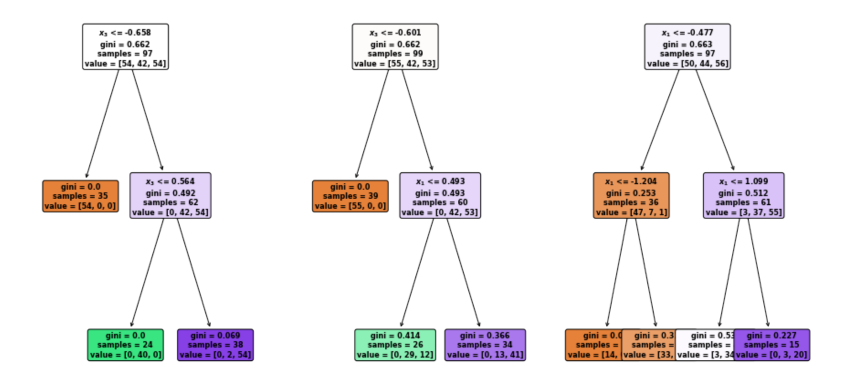
We will shortly explain how the subsets are chosen.

Here is an Ensemble of individual models of the same type: Decision Trees

- classification among 3 classes
- trained on different subsets of the training dataset
  - details to follow: Bagging, Boosting
- ullet we have limited the features used to  ${f x}_1,{f x}_3$  only to make the diagrams smaller

In [7]: | fig\_ens

#### Out[7]:



The individual models are usually quite simple and restricted. • They are weak learners: accuracy only marginally better than chance • But combine to create a strong learner.

If the prediction of an ensemble of M binary classifiers is based on a "majority vote"

- ullet The prediction is incorrect only if  $m' \geq \lceil M/2 
  ceil$  classifiers are incorrect
- The probability of a particular set of  $m^\prime$  models of equal accuracy A all being incorrect is  $(1-A)^{m^\prime}$
- There are

$$\binom{M}{m'}$$

combinations of m' models

ullet So the probability of a correct ensemble prediction when  $m^\prime$  classifiers are incorrect is

$$1-\left(rac{M}{m'}
ight)*(1-A)^{m'}$$

which tends to 1 as M ( and hence,  $m' \geq \lceil M/2 \rceil$  ) increases.

- since (1 A) < 1
- when raised to a power (m') the second term goes to 0

The power of Ensembles comes via the size of M.

Ensembling is independent of the types of the individual models

- A meta-model that can combine many different types of individual models
- Under the assumption of **independent** errors
- Often applied in competitions

## **Ensemble prediction**

Each individual model comes up with a prediction for the target  $\hat{\mathbf{y}}^{(i)}$  of example i, given features  $\mathbf{x}^{(i)}$ .

Let  $p_{(t),c}^{(\mathbf{i})}$ 

- ullet Denote the probability predicted by the  $t^{th}$  individual classifier
- That target  $\mathbf{y^{(i)}}$  is in category  $c \in C$
- Given features  $\mathbf{x}^{(i)}$

The class predicted by the ensemble is the one with highest average (across individual models) probability

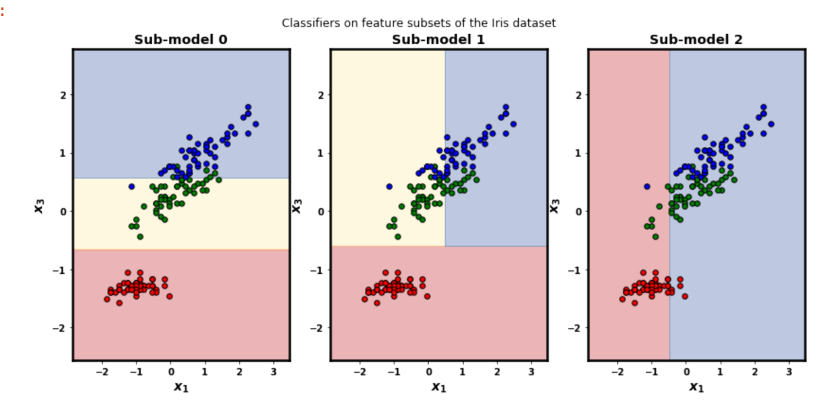
$$\hat{\mathbf{y}}^{(\mathbf{i})} = rgmax \sum_{t=1}^{M} p_{(t),c}^{(\mathbf{i})}$$

Returning to the Ensemble of Decision Trees example, we can plot the decision boundary of each individual model

• 3 classes: red, green, blue
• the boundaries of each model differ
• because they have been trained on different subsets of the full training dataset

In [8]: fig\_submodels

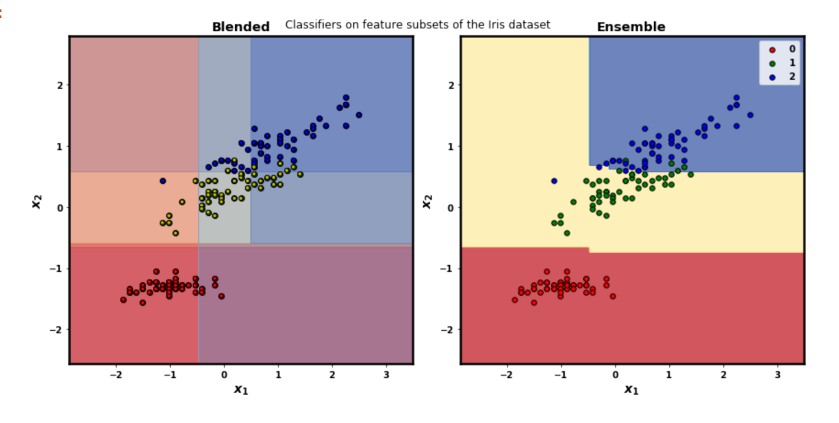
### Out[8]:





In [9]: fig\_sum

### Out[9]:



- The left plot is the super-position
- The right plot is the final boundary of the ensemble

You can see that the combination of the weak learners does a pretty good job!

# Bagging, Bootstrapping

One way to construct multiple weak learners of the same type of model

• Is to train each individual model on a *restricted* set of training examples

Because each individual model is trained on different examples, the predictions made by each are hopefully somewhat independent.

Given the full set of training examples

$$\langle \mathbf{X}, \mathbf{y} \rangle = [\mathbf{x^{(i)}}, \mathbf{y^{(i)}} | 1 \leq i \leq m]$$

we construct a restricted set of examples

$$\langle \mathbf{X}_{(t)}, \mathbf{y}_{(t)} 
angle$$

on which to train the  $t^{th}$  individual model

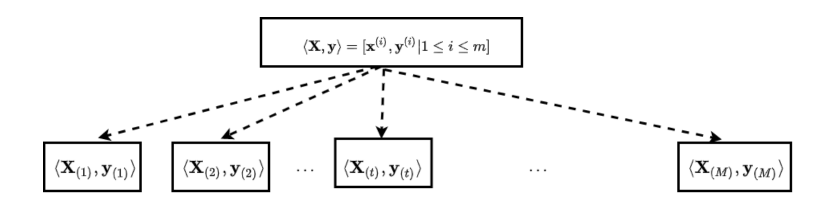
The restricted set is constructed by

- Selecting m examples at random from  $\langle \mathbf{X}, \mathbf{y} 
  angle$
- With replacement
- ullet So it is possible for an example i' to appear more than once in  $\langle \mathbf{X}_{(t)}, \mathbf{y}_{(t)} 
  angle$

This process is called bootstrapping (or bagging) and results in

- $egin{aligned} ullet \left\langle \mathbf{X}_{(t)}, \mathbf{y}_{(t)} 
  ight
  angle \ &= [\mathbf{x}^{(i')}, \mathbf{y}^{(i')} | i' \in \{i_1, \dots, i_m\}] \end{aligned}$
- ullet Where  $i_1,\ldots,i_m$  are the indices of the m chosen examples

### **Bagging**



If each of the m examples in  $\langle \mathbf{X}, \mathbf{y} 
angle$  is chosen with equal probability  $\frac{1}{m}$ 

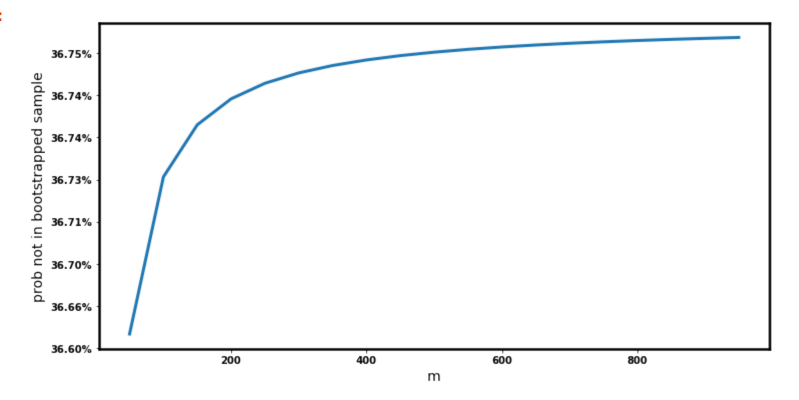
ullet The probability of a particular example i **not** being in  $\langle \mathbf{X}_{(t)}, \mathbf{y}_{(t)} 
angle$  is

$$(1-\frac{1}{m})^m$$

Let's plot this probability as a function of the training dataset size m

In [11]: fig

### Out[11]:



Thus about 63% of the examples in the bootstrapped set are duplicates.

Why is this a potential advantage?

- the model may perform better (in-sample) on duplicated examples
- the model can't overfit to any example that is not in its training set.

### The process of

- Bootstrapping restricted training examples
- Training individual models on the bootstrapped examples
- Aggregating model predictions into a single prediction

is called bagging and each individual training set is called a bag

Bagging has a nice side-effect

- About 37% of the full set of examples are not present in a given bag
- Called out of bag

The out of bag examples thus can be used to test out of sample prediction!

• a built-in validation dataset

## **Random Forests**

A Random Forest

- Is a collection of Decision Trees
- Of restricted power (weak learners)
- Created by Bagging

The learners are made weak by

- Training on a bootstrapped subset
- By limiting the depth of the Decision Tree
- By limiting the choice of feature on which to split a node
  - To a random subset of all features

The result is that the individual models (Decision Trees) are relatively independent.

# **Boosting**

There is another approach to creating ensembles of weak learners.

The method is called boosting

- Rather than create weak learners independently, i.e., a set
- ullet Boosting creates a sequence of weak learners:  $M_{(0)}, M_{(1)}, \ldots, M_{(M)}$
- ullet Where the  $(t+1)^{th}$  individual model in the sequence
- ullet Focuses on correctly predicting those examples incorrectly predicted by the  $t^{th}$  individual model

### **Notation**

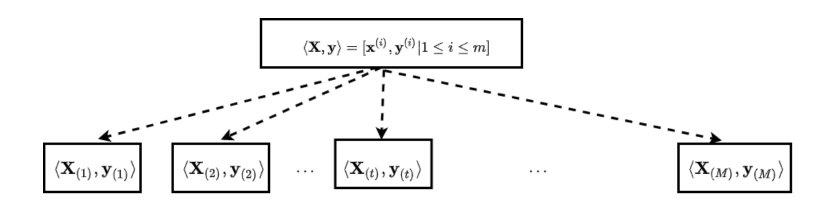
We will be dealing with many sequences. We use subscripts in parentheses to index elements of a sequence.

$$M_{(0)}, M_{(1)}, \ldots, M_{(M)}$$

### Recall:

- when bootstrapping/bagging
- each individual training dataset is drawn simultaneously from the full training dataset

### **Bagging**



#### In contrast

- boosting creates the individual training datasets sequentially
- ullet subset  $\langle \mathbf{X}_{(t+1)}, \mathbf{y}_{(t+1)} 
  angle$  for model  $M_{(t+1)}$
- ullet is chosen to compensate for the errors of **all prior** models  $\{M_{(t')} \mid t' < t\}$

### **Boosting**

$$\langle \mathbf{X}, \mathbf{y} \rangle$$
 ---  $\langle \mathbf{X}_{(1)}, \mathbf{y}_{(1)} \rangle$  ---  $\langle \mathbf{X}_{(2)}, \mathbf{y}_{(2)} \rangle$  ->  $\langle \mathbf{X}_{(t)}, \mathbf{y}_{(t)} \rangle$  ->  $\langle \mathbf{X}_{(t)}, \mathbf{y}_{(t)} \rangle$ 

How do we get an individual model to focus on some particular examples?

- By assigning each example a weight
- Increasing the probability that more heavily weighted examples are included in the training examples for the model
  - examples with poor predictions by earlier models are over-weighted in the subsequent model

Let  $\sup_{(t)}^{(\mathbf{i})}$  denote the weight assigned to example i in the training set for the  $t^{th}$  individual model

The "say" is adjusted from the  $t^{th}$  model to the  $(t+1)^{th}$  individual model

If example i is incorrectly predicted in model t:  $\operatorname{say}_{(t+1)}^{(\mathbf{i})} > \operatorname{say}_{(t)}^{(\mathbf{i})}$ 

If example i is correctly predicted in model t:  $\sup_{(t+1)}^{(\mathbf{i})} < \sup_{(t)}^{(\mathbf{i})}$ 

When bootstrapping, rather than drawing examples with equal probability

- ullet Draw examples for model (t+1) in proportion to it's  $\sup_{(t+1)}^{(\mathbf{i})}$
- So examples that were "problematic" in model t are over-represented in training model (t+1)

Boosting creates a collection of "specialists" (focus on hard to predict examples) Bagging creates a collection of "generalists", each a little better than random

### **AdaBoost**

AdaBoost is a particular model that uses boosting

- The individual models are Decision Trees
  - Usually depth 1; "stumps"
- There is an "importance" associated with each individual model
- Models with higher weight have a greater impact on ensemble prediction

Let

### $importance_{(t)}$

denote the weight of the  $t^{th}$  individual model in the sequence.

- ullet importance $_{(t)}$  is determined by the Performance Metric (e.g., Accuracy) of individual model t
- The class predicted by the ensemble is the one with highest *importance-weighted* average (across individual models) probability

$$\hat{\mathbf{y}^{(\mathbf{i})}} = rgmax_{c} \sum_{t=1}^{M} (p_{(t),c}^{(\mathbf{i})} * \mathrm{importance}_{(t)})$$

Thus, models that are more successful have greater weight.

# **Example: Boosting for a Regression task**

Boosting is often associate with Classification tasks

• for example: the individual models are Decision Trees

Here we show how it may be used for Regression.

Our goal is to solve for the optimal parameters  $\Theta^*$  for the Linear Regression

$$\mathbf{y} = \Theta \cdot \mathbf{x} + \epsilon$$

That is,  $\Theta^*$  minimizes the MSE

$$\Theta^* = \operatorname*{argmin}_{\Theta} rac{1}{m} \sum_i \left( \mathbf{y^{(i)}} - \Theta \cdot \mathbf{x^{(i)}} 
ight)^2 
ight)$$

The Boosting method

ullet creates a sequence of approximations of the parameters  $\Theta$ 

$$\Theta_{(0)},\Theta_{(1)},\dots$$

that approach the optimal  $\Theta^*$ .

We will create a sequence of models  $M_{(0)}, M_{(1)}, \dots, M_{(M)}$ 

Each model's functional form is a Linear

 $\mathsf{Model}\,t$ 

$$egin{array}{lll} \mathbf{e}_{(t)} &=& \Theta_{(t)} \cdot \mathbf{x} + \epsilon_{(t)} \ &=& \hat{\mathbf{e}}_{(t)} + \epsilon_{(t)} \end{array}$$

where

- $\Theta_{(t)}$  are the parameters of the model
- $\mathbf{e}_{(t)}$  is the target (to be defined)
- $\hat{\mathbf{e}}_{(t)}$  is the predicted value
- ullet  $\epsilon_{(t)}$  is the prediction error

## For the first model Model $M_{(0)}$

• target is **y** 

$$\mathbf{e}_{(0)} = \mathbf{y}$$

ullet we ignore the features  ${f x}$  and predict the average

$$\hat{\mathbf{e}}_{(0)} = \bar{\mathbf{y}}$$

ullet the intercept parameters is  $ar{f y}$  , all other parameters are 0

$$\Theta_{(0)} = (ar{\mathbf{y}}, 0, \dots, 0)$$

For subsequent models t+1

- the target  $\mathbf{e}_{(t+1)}$
- ullet is the *error* of the previous model  $M_{(t)}$

$$\mathbf{e}_{(t+1)} = \epsilon_{(t)}$$

## That is

- $\bullet \ \ {\rm the \, goal \, of \, model} \, M_{(t+1)}$
- is to reduce the error of the previous model

#	$\mathbf{e}_{(t)}$	$\hat{\mathbf{e}}_{(t)}$	$\epsilon_{(t)}$
0	$\mathbf{y}$	$\Theta_{(0)}\cdot {f x}$	$\epsilon_{(0)}$
1	$\epsilon_{(0)}$	$\Theta_{(1)}\cdot \mathbf{x}$	$\epsilon_{(1)}$
•			
M	$\epsilon_{(M-1)}$	$\Theta_{(M)}\cdot {f x}$	$\epsilon_{(M)}$

### Unlike in the Linear Regression Model

• we will obtain improve prediction

$$\hat{\mathbf{y}}_{(t)} = \hat{\mathbf{e}}_{(t)}$$

• by modifying the previous prediction

$$\hat{\mathbf{e}}_{(t-1)}$$

directly

- lacktriangledown rather than by modifying parameters  $\Theta_{(t-1)}$
- lacktriangledown we do not solve for  $\Theta_{(t)}$  using Linear Regression

This method, called *Gradient Boosting* will be described shortly.

### **Aside**

Had we solved for improved  $\Theta_{(t)}$  using Linear Regression

- ullet the iterative process would stop after t=1
- ullet error  $\epsilon_{(1)}$  would be uncorrelated with  ${f x}$ 
  - property of Linear Regression

The predictions of the individual models in the ensemble consisting of the first t models  $M_{(0)},\ldots,M_{(t)}$ 

- is combined into the ensemble prediction  $\hat{\mathbf{y}}_{(t)}$
- as the weighted sum of the predictions of the individual models in the ensemble

$$egin{array}{lll} \hat{\mathbf{y}}_{(t)} &=& \sum_{t'=0}^t lpha_{(t)} * \hat{\mathbf{e}}_{(t)} \ &=& \mathbf{x} \cdot \sum_{t'=0}^t lpha_{(t)} * \Theta_{(t')} \end{array}$$

The boosting solution just derives the coefficients  $\Theta^*$  of a direct Linear Regression model

$$\hat{\mathbf{y}} = \Theta^* \cdot \mathbf{x}$$

iteratively, as a sum

$$\Theta^* = \sum_{t'=0}^t lpha_{(t)} * \Theta_{(t')}$$

### We can show this another way

- ullet the final model's error  $\epsilon_{(M)}$
- is the error of a direct linear regression

Since

$$\hat{\mathbf{e}}_{(t)} = \mathbf{e}_{(t)} - \epsilon_{(t)}$$

we can write the ensemble prediction of all M models as

## Convergence to $\Theta^*$

The  $\Theta_{(t)}$  constructed for model  $M_{(t)}$ 

• is an approximation of the optimal  $\Theta^*_{(t)}$  (i.e., the one that Linear Regression solves for) for the model's equation

$$\mathbf{e}_{(t)} = \Theta_{(t)} \cdot \mathbf{x} + \epsilon_{(t)}$$

• for all but the final model (assuming we continue until convergence)

We can readily see the non-optimality in our choice of  $\Theta_{(0)}$ .

We will show (in the section on Gradient Boosting)

• how to construct these approximations

When optimality is achieved

$$\Theta_{(t)}=\Theta_{(t)}^*$$

- the error  $\epsilon_{(t)}$  becomes uncorrelated with  ${f x}$ 
  - mathematical property of Linear Regression
- ullet so that model  $M_{(t')}$  for t'>t will **not be able to** further reduce the error, i.e.

$$\Theta_{(t')} = 0$$

$$\epsilon_{(t')} = \epsilon_{(t)}$$

Thus, there is no point continuing the Boosting process beyond this point.

## **Example: Boosting for a Classification task**

Although we won't construct an example for Classification, there are some important points to consider.

Each model is created from scratch

- ullet Model  $M_{(t+1)}$  does **not** extend model  $M_{(t)}$
- For example, if the models are Decision Trees
  - lacktriangledown the tree for  $M_{(t+1)}$  is not an expansion of the tree for  $M_{(t)}$

Although the models are created independently

their training datasets are constructed sequentially

# **Gradient Boosting**

Gradient Boosting is a "more mathematical" (less operational) approach to boosting.

We associate a Loss Function

$$\mathcal{L}_{(t)}$$

- ullet with the ensemble prediction  $\hat{f y}_{(t)}$
- ullet of the first t models:  $M_{(0)}, M_{(1)}, \ldots, M_{(t)}$

Model  $M_{(t+1)}$  is constructed so as to reduce the ensemble loss

$$\mathcal{L}_{(t+1)} \leq \mathcal{L}_{(t)}$$

In fact, our Regression example above used Gradient Boosting!

Define the Loss for the ensemble consisting of the first t models as the MSE error

$$\mathcal{L}_{(t)} = rac{1}{m} \sum_{i=1}^m {(\mathbf{y^{(i)}} - \hat{\mathbf{y}}_{(t)}^{(i)})^2}$$

where

$$\hat{\mathbf{y}}_{(t)}^{(\mathbf{i})}$$

is the ensemble prediction for training example i.

## Compute the gradient

- of the Loss  $\mathcal{L}_{(t)}$  with respect to ensemble prediction  $\hat{\mathbf{y}}_{(t)}$  $\nabla_{(t)} = \frac{\partial \mathcal{L}_{(t)}}{\partial \hat{\mathbf{y}}_{(t)}}$

$$abla_{(t)} = rac{\partial \mathcal{L}_{(t)}}{\partial \hat{\mathbf{y}}_{(t)}}$$

### By definition of the Gradient

- ullet we can reduce the loss  $\mathcal{L}_{(t+1)}$  of the next model
- ullet by creating ensemble prediction  $\hat{f y}_{(t+1)}$  as

$$\hat{\mathbf{y}}_{(t+1)} = \hat{\mathbf{y}}_{(t)} - lpha * 
abla_{(t)}$$

#### The Gradient

- is in the direction of an increase of Loss
- so we adjust in the *negative* direction of the gradient
- in proportion to  $\alpha$  (the *learning rate*)

The iterative process of reducing the Loss by moving in the direction of the gradient

- is called Gradient Descent
- and is the basis for the optimization used in Deep Learning

Referring back to our Regression example:

We can compute the gradient for the MSE Loss we have chosen:

$$egin{array}{lll} rac{\partial \mathcal{L}_{(t)}}{\partial \hat{\mathbf{y}}_{(t)}} &=& rac{\partial rac{1}{m} \sum_{i=1}^m \left( \mathbf{y^{(i)}} - \hat{\mathbf{y}}_{(t)} 
ight)^{(i)} 
ight)^2}{\partial \hat{\mathbf{y}}_t} \ &=& rac{2}{m} (\mathbf{y^{(i)}} - \hat{\mathbf{y}}_{(t)}^{(i)}) * -1 & ext{chain rule} \ &=& -rac{2}{m} \epsilon_{(t)} & ext{definition of } \epsilon_{(t)} \end{array}$$

So our ensemble predictor (for the larger ensemble of t+1 models)

$$\hat{\mathbf{y}}_{(t+1)}$$

should increase our previous (smaller ensemble of t models) ensemble prediction

$$\hat{\mathbf{y}}_{(t)}$$

proportional to the error

$$\epsilon_{(t)}$$

in order to reduce the Loss

$$\epsilon_{(t)}$$
  $\mathcal{L}_{(t+1)} \leq \mathcal{L}_{(t)}$ 

## That is

- ullet model t
- ullet under-estimates  ${f y}$  by  $\epsilon_{(t)}$ 
  - since

$$\mathbf{y} = \hat{\mathbf{y}} + \epsilon$$

The larger ensemble adds one more term to the summation expression for the ensemble predictor

$$\hat{\mathbf{y}}_{(t+1)} = \hat{\mathbf{y}}_{(t)} + \hat{\mathbf{e}}_{(t+1)}$$

Note that the target  $\mathbf{e}_{(t+1)}$  for model t+1 is

$$\mathbf{e}_{(t+1)} = \epsilon_{(t)}$$

SO

$$\hat{\mathbf{e}}_{(t+1)} = \hat{\epsilon}_{\,(t)}$$

That is

- ullet the term added to  $\hat{f y}_{(t)}$  to obtain  $\hat{f y}_{(t+1)}$
- is approximately equal to the Gradient of Loss
- as required by Gradient Descent

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Done