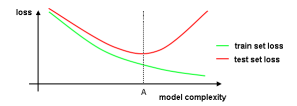
## Ensemble learning, bias-variance decomposition

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### Loss vs. model complexity



#### Comments:

- expected loss on test set is always higher than on train set.
- left to A: model too simple, underfitting, high bias
- right to A: model too complex, overfitting, high variance

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- **5** Sampling ensemble methods

### Bias-variance decomposition

- True relationship  $y = f(x) + \varepsilon$
- This relationship is estimated using random training set  $(X, Y) = \{(x_n, y_n), n = 1, 2...N\}$
- Recovered relationship  $\widehat{f}(x)$ , x-some fixed constant
- Noise  $\varepsilon$  is independent of any  $X,Y,\ \mathbb{E}\varepsilon=0$  and  $Var[\varepsilon]=\sigma^2$

#### Bias-variance decomposition

$$\begin{split} \mathbb{E}_{X,Y,\varepsilon}\{[\widehat{f}(x)-y(x)]^2\} &= \left(\mathbb{E}_{X,Y}\{\widehat{f}(x)\}-f(x)\right)^2 \\ &+ \mathbb{E}_{X,Y}\left\{[\widehat{f}(x)-\mathbb{E}_{X,Y}\widehat{f}(x)]^2\right\} + \sigma^2 \end{split}$$

- Intuition:  $MSE = bias^2 + variance + irreducible error$ 
  - darts intuition

### Proof of bias-variance decomposition

Define for brevity of notation f = f(x),  $\hat{f} = \hat{f}(x)$ ,  $\mathbb{E} = \mathbb{E}_{X,Y,\varepsilon}$ .

$$\mathbb{E}\left(\widehat{f} - f\right)^{2} = \mathbb{E}\left(\widehat{f} - \mathbb{E}\widehat{f} + \mathbb{E}\widehat{f} - f\right)^{2} = \mathbb{E}\left(\widehat{f} - \mathbb{E}\widehat{f}\right)^{2} + \left(\mathbb{E}\widehat{f} - f\right)^{2} + 2\mathbb{E}\left[\left(\widehat{f} - \mathbb{E}\widehat{f}\right)(\mathbb{E}\widehat{f} - f)\right]$$
$$= \mathbb{E}\left(\widehat{f} - \mathbb{E}\widehat{f}\right)^{2} + \left(\mathbb{E}\widehat{f} - f\right)^{2}$$

We used that  $(\mathbb{E}\widehat{f} - f)$  is a constant w.r.t. X, Y and hence  $\mathbb{E}\left[(\widehat{f} - \mathbb{E}\widehat{f})(\mathbb{E}\widehat{f} - f)\right] = (\mathbb{E}\widehat{f} - f)\mathbb{E}(\widehat{f} - \mathbb{E}\widehat{f}) = 0$ .

$$\begin{split} \mathbb{E}\left(\widehat{f} - y\right)^2 &= \mathbb{E}\left(\widehat{f} - f - \varepsilon\right)^2 = \mathbb{E}\left(\widehat{f} - f\right)^2 + \mathbb{E}\varepsilon^2 - 2\mathbb{E}\left[(\widehat{f} - f)\varepsilon\right] \\ &= \mathbb{E}\left(\widehat{f} - \mathbb{E}\widehat{f}\right)^2 + \left(\mathbb{E}\widehat{f} - f\right)^2 + \sigma^2 \end{split}$$

Here  $\mathbb{E}\left[(\widehat{f}-f)\varepsilon\right]=\mathbb{E}\left[(\widehat{f}-f)\right]\mathbb{E}\varepsilon=0$  since  $\varepsilon$  is independent of X,Y.

### Ensemble learning

#### Definition 1

Ensemble learning - using multiple machine learning methods for a given problem and integrating their output to obtain final result.

Synonyms: committee-based learning, multiple classifier systems.

#### **Applications:**

- supervised methods: regression, classification
- unsupervised methods: clustering

#### Ensembles use cases

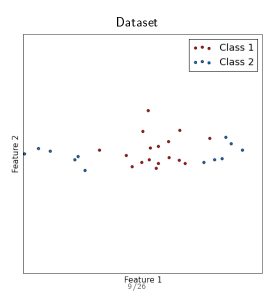
- solving C class classification with many binary classifiers
- underfitting, high model bias
  - existing model hypothesis space is too narrow to explain the true one
  - different oversimplified models have bias in different directions, mutually compensating each other.
- overfitting, high model variance
  - avoid local optima of optimization methods
  - too small dataset to figure out concretely the exact model hypothesis
- when task itself promotes usage of ensembles with features of different nature
  - E.g. computer security:
    - multiple sources of diverse information (password, face detection, fingerprint)
    - different abstraction levels need to be united (current action, behavior pattern during day, week, month)

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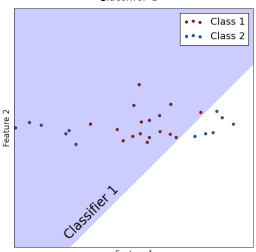
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### Classification: original model space too narrow



## Classification: original model space too narrow

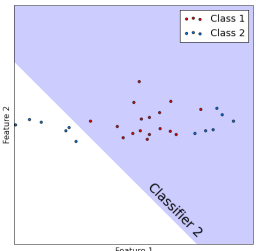
#### Classifier 1



Feature 1

## Classification: original model space too narrow

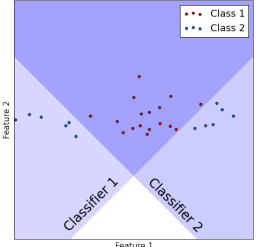




Feature 1

## Classification: original model space too narrow

#### Classifier 1 and classifier 2 combined using AND rule



Feature 1

### Motivation for ensembles

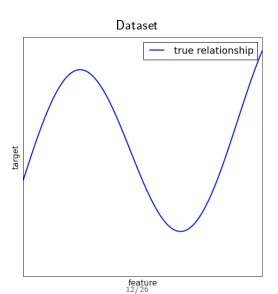
- Consider M classifiers  $f_1(x), ... f_M(x)$ , performing binary classification.
- Let probability of mistake be constant  $p \in (0, \frac{1}{2})$ :  $p(f_m(x) = y) = p \forall m$
- Suppose all models make mistakes or correct guesses independently of each other.
- Let F(x) be majority voting combiner.
- Then  $p(F(x) \neq y) \rightarrow 0$  as  $m \rightarrow \infty$

- 2 Accuracy improvement demos
  - Accuracy improvement for classification
  - Accuracy improvement for regression

Accuracy improvement demos

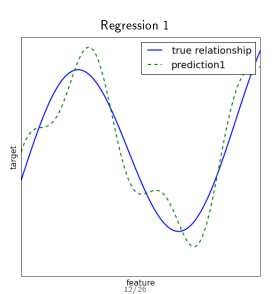
Accuracy improvement for regression

## Regression: high variance



Accuracy improvement demos Accuracy improvement for regression

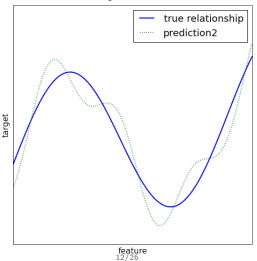
## Regression: high variance



Accuracy improvement demos Accuracy improvement for regression

### Regression: high variance



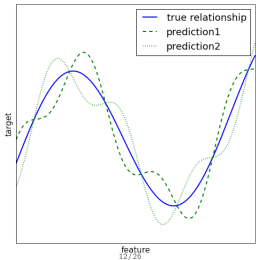


Accuracy improvement demos

Accuracy improvement for regression

Regression: high variance

#### Regression 1 and regression 2 combined using averaging



# Use combination or individual models?

- Consider regression task, performed by individual learners  $f_1(x),...f_K(x)$  and their weighted combination  $F(x) = \sum_{k=1}^K \alpha_k f_k(x)$  with some weights  $\alpha_1,...\alpha_K: \alpha_k \geq 0 \ \forall k, \sum_{k=1}^K \alpha_k = 1$ .
- Take convex loss  $\mathcal{L}(\hat{y} y)$ , such as absolute or square.
- What is better: to use F(x) or take one of  $f_1(x),...f_K(x)$  randomly with probabilities  $\alpha_1,...\alpha_K$ ?

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#### Fixed combiner at class level

### Output of base learner k

Exact class:  $\omega_1$  or  $\omega_2$ .

Combiner predicts  $\omega_1$  if:

- all classifiers predict  $\omega_1$  (AND rule)
- at least one classifier predicts  $\omega_1$  (OR rule)
- at least k classifiers predict  $\omega_1$  (k-out-of-N)
- majority of classifiers predict  $\omega_1$  (majority vote)

Each classifier may be assigned a weight, based on its performance:

- weighted majority vote
- weighted k-out-of-N (based on score sum)

### Fixed combiner - ranking level

#### Output of base learner k

Ranking of classes:

$$\omega_{k_1} \succeq \omega_{k_2} \succeq \ldots \succeq \omega_{k_C}$$

Ranking is equivalent to scoring of each class (with incomparable scoring between classifiers).

#### Definition 2

Let  $B_k(i)$  be the count of classes scored below  $\omega_i$  by classifier k. Borda count B(i) of class  $\omega_i$  is the total number of classes scored below  $\omega_i$  by all classifiers:

$$B(i) = \sum_{k=1}^{K} B_k(i)$$

Combiner predicts  $\omega_i$  where  $i = \arg \max_i B(i)$ 

### Fixed combiner at class probability level

#### Output of base learner k

Vectors of class probabilities:

$$[p^k(\omega_1), p^k(\omega_2), \ldots p^k(\omega_C)]$$

Combiner predicts  $\omega_i$  if  $i = \arg\max_i F(p^1(\omega_i), p^2(\omega_i), \dots p^K(\omega_i))$ 

• F = mean or median.

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### Weighted averaging

Consider regression with K predictor models  $f_k(x)$ , k = 1, 2, ...K. (Alternatively we may consider K discriminant functions in classification)

#### Weighted averaging combiner

$$f(x) = \sum_{k=1}^{K} w_k f_k(x)$$

Naive fitting

$$\widehat{w} = \arg\min_{w} \sum_{i=1}^{N} \mathcal{L}(y_i, \sum_{k=1}^{K} w_k f_k(x_i))$$

will overfit. The mostly overfitted method will get the most weight.

### Linear stacking

- Let training set  $\{(x_i, y_i), i = 1, 2, ...N\}$  be split into M folds.
- Denote fold(i) to be the fold, containing observation i
- Denote  $f_k^{-fold(i)}$  be predictor k trained on all folds, except fold(i).

#### Definition

Linear stacking is weighted averaging combiner, where weights are found using

$$\widehat{w} = \arg\min_{w} \sum_{i=1}^{N} \mathcal{L}(y_i, \sum_{k=1}^{K} w_k f_k^{-fold(i)}(x_i))$$

• For decreased overfitting we may add constraints  $\{w_k \geq 0\}_{k=1}^K$  or regularizer  $\sum_{k=1}^K \left(w_k - \frac{1}{K}\right)^2$ .

### General stacking

#### Definition

Generalized stacking is prediction

$$f(x) = A_{\theta}(f_1(x), f_2(x), \dots f_K(x)),$$

where A is some general form predictor and  $\theta$  is a vector of parameters, estimated by

$$\widehat{\theta} = \arg\min_{\theta} \sum_{i=1}^{N} \mathcal{L}\left(y_i, A_{\theta}\left(f_1^{-fold(i)}(x), f_2^{-fold(i)}(x), \dots f_K^{-fold(i)}(x)\right)\right)$$

- Stacking is the most general approach
- It is a winning strategy in most ML competitions.
- $f_i(x)$  may be:
  - class number (coded using one-hot encoding).
  - vector of class probabilities
  - any initial or generated feature

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### Bagging& random subspaces

### Bagging:

- random selection of samples (with replacement) 12
- efficient for methods with high variance w.r.t. X, Y.

#### Random subspace method:

- random selection of features (without replacement)
- We can apply both methods jointly
- Also we may sample different

<sup>&</sup>lt;sup>1</sup>what is the probability that observation will not belong to bootstrap sample?

<sup>&</sup>lt;sup>2</sup> what is the limit of this probability with  $N \to \infty$ ?

#### Random forests

**Input**: training dataset  $TDS = \{(x_i, y_i), 1 = 1, 2, ...N\}$ ; the number of trees B and the size of feature subsets m. for b = 1, 2, ...B:

- generate random training dataset  $TDS^b$  of size N by sampling  $(x_i, y_i)$  pairs from TDS with replacement.
- build a tree using TDS<sup>b</sup> training dataset with feature selection for each node from random subset of features of size m (generated individually for each node).

**Output**: B trees. Classification is done using majority vote and regression using averaging of B outputs.

#### Comments

- Random forests use random selection on both samples and features
- Step 1) is optional.
- Left out samples may be used for evaluation of model performance.
  - Out-of-bag prediction: assign output to  $x_i$ , i = 1, 2, ...N using majority vote (classification) or averaging (regression) among trees with  $b \in \{b : (x_i, y_i) \notin T^b\}$
  - Out-of-bag quality lower bound for true model quality.<sup>3</sup>
- Less interpretable than individual trees
- +: Parallel implementation
- -: different trees are not targeted to correct mistakes of each other

<sup>&</sup>lt;sup>3</sup>why *lower* bound?

#### Comments

- Extra-Random trees-random sampling of (feature, value) pairs
  - more bias and less variance for each tree
  - faster training of each tree
- RandomForest and ExtraRandomTrees do not overfit with increasing B
- Each tree should have high depth
  - otherwise averaging over oversimplified trees will also give oversimplified model!