10-301/601: Introduction to Machine Learning Lecture 5 – Model **Selection** 

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### **Lecture 5 Polls**

#### 0 done

 $\sim$  50 underway construction of the const

### **Front Matter**

- Announcements:
	- · PA1 released 5/18, due 5,
- · Recommended Readings:
	- · Daumé III, Chapter 2: Lim

Recall: Setting  $k$ 

- $\cdot$  When  $k = 1$ :
	- many, complicated decision boundaries
	- may overfit
- $\cdot$  When  $k = N$ :
	- no decision boundaries; always predicts the most common label in the training data
	- · may underfit
- $\cdot$  k controls the complexity of the hypothesis set  $\Longrightarrow$  k affects how well the learned hypothesis will generalize

Recall: Setting  $k$ 

- Theorem:
	- $\cdot$  If k is some function of N s.t.  $k(N) \rightarrow \infty$  and  $\frac{k(N)}{N}$  $\boldsymbol{N}$  $\rightarrow 0$ as  $N \rightarrow \infty$  ...
	- ... then (under certain assumptions) the true error of a  $kNN$  model  $\rightarrow$  the Bayes error rate
- Practical heuristics:
	- $\cdot k = |\sqrt{N}|$
	- $\cdot k = 3$
- $\cdot$  This is a question of **model selection**: each value of  $k$ corresponds to a different "model"

# Model Selection

- A **model** is a (typically infinite) set of classifiers that a learning algorithm searches through to find the best one (the "hypothesis space")
- **· Model parameters** are the numeric values or structure that are selected by the learning algorithm
- **Hyperparameters** are the tunable aspects of the model that are not selected by the learning algorithm

#### **Example: Decision Trees**

- $\cdot$  Model = set of all possible trees, potentially narrowed down according to the hyperparameters (see below)
- Model parameters = structure of a specific tree e.g., splits, split order, predictions at leaf nodes,
- Hyperparameters = splitting criterion, maxdepth, tie-breaking procedures, etc...

# Model Selection

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### **Example:**  $kNN$

 $\cdot$  Model = set of all possible nearest neighbors classifiers

• Model parameters = none! kNN is a "nonparametric model"

• Hyperparameters =  $k$ 

Model **Selection** with **Test Sets** 

Given  $\mathcal{D} = \mathcal{D}_{train} \cup \mathcal{D}_{test}$ , suppose we have multiple candidate models:

 $\mathcal{H}_1, \mathcal{H}_2, ..., \mathcal{H}_M$ 

• Learn a classifier from each model using only  $\mathcal{D}_{train}$ :  $h_1 \in \mathcal{H}_1, h_2 \in \mathcal{H}_2, \dots, h_M \in \mathcal{H}_M$ 

Evaluate each one using  $D_{test}$  and choose the one with lowest test error:

> $\dot{m} = \argmin_{m \in \{1, ..., M\}} err(h_m, D_{test})$  $m \in \{1,...,M\}$

Model Selection with Test Sets? Given  $\mathcal{D} = \mathcal{D}_{train} \cup \mathcal{D}_{test}$ , suppose we have multiple candidate models:

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 $\cdot$  Is  $err(h_{\hat{m}}, D_{test})$  a good estimate of  $err(h_{\hat{m}})$ ?

Model Selection with **Validation Sets**  Given  $\mathcal{D} = \mathcal{D}_{train} \cup \mathcal{D}_{val} \cup \mathcal{D}_{test}$ , suppose we have multiple candidate models:  $\mathcal{H}_1, \mathcal{H}_2, ..., \mathcal{H}_M$ 

• Learn a classifier from each model using only  $D_{train}$ :  $h_1 \in \mathcal{H}_1, h_2 \in \mathcal{H}_2, \dots, h_M \in \mathcal{H}_M$ 

Evaluate each one using  $\mathcal{D}_{val}$  and choose the one with lowest *validation* error:

 $\dot{m} = \argmin$  $m \in \{1,...,M\}$  $err(h_m, D_{val})$ • Now  $err(h_{\hat{m}}, D_{test})$  is a good estimate of  $err(h_{\hat{m}})$ !

Hyperparameter **Optimization** with **Validation Sets** 

Given  $\mathcal{D} = \mathcal{D}_{train} \cup \mathcal{D}_{val} \cup \mathcal{D}_{test}$ , suppose we have multiple candidate hyperparameter settings:  $\theta_1, \theta_2, ..., \theta_M$ 

- Learn a classifier for each setting using only  $\mathcal{D}_{train}$ :  $h_1, h_2, ..., h_M$
- Evaluate each one using  $\mathcal{D}_{val}$  and choose the one with lowest *validation* error:

 $\dot{m} = \underset{m \in \{1, \ldots, M\}}{\text{argmin}} \text{err}(h_m, D_{val})$  $m \in \{1,...,M\}$ 

• Now  $err(h_{\hat{m}}, D_{test})$  is a good estimate of  $err(h_{\hat{m}})$ !

Setting  $k$ for  $kNN$ with **Validation Sets** 



#### $k$  NN train and validation errors on Fisher Iris data

How should we partition our dataset?



#### $k$  NN train and validation errors on Fisher Iris data



• Use each one as a validation set once:



- Let  $h_{-i}$  be the classifier learned using  $\mathcal{D}_{-i} = \underline{\mathcal{D}} \backslash \underline{\mathcal{D}}_i$  (all folds other than  $\underline{\mathcal{D}}_i$ ) and let  $e_i = err(h_{-i}, D_i)$
- $\cdot$  The K-fold cross validation error is

$$
err_{CV_K} = \frac{1}{K} \sum_{i=1}^{K} e_i
$$

- Given  $D$ , split  $D$  into  $K$  equally sized datasets or folds:  $\mathcal{D}_1, \mathcal{D}_2, \dots, \mathcal{D}_K$
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• Special case when  $K = N$ : Leave-one-out cross-validation

• Choosing between  $M$  candidates requires training  $MK$  times

# **Summary**



# Hyperparameter **Optimization**

Given  $\mathcal{D} = \mathcal{D}_{train} \cup \mathcal{D}_{val} \cup \mathcal{D}_{test}$ , suppose we have multiple candidate hyperparameter settings:

 $\theta_1, \theta_2, ..., \theta_M$ 

- Learn a classifier for each setting using only  $\mathcal{D}_{train}$ :  $h_1, h_2, ..., h_M$
- Evaluate each one using  $\mathcal{D}_{val}$  and choose the one with lowest *validation* error:

 $\hat{m} = \underset{m \in \{1, \ldots, M\}}{\text{argmin}} \text{err}(h_m, \mathcal{D}_{val})$  $m \in \{1,...,M\}$ 

• Now  $err(h_{\hat{m}}^+, D_{test})$  is a good estimate of  $err(h_{\hat{m}}^+)$ !

Pro tip: train your final model using *both* training and validation datasets

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 $\hat{m} = \underset{m \in \{1, \ldots, M\}}{\text{argmin}} \text{err}(h_m, \mathcal{D}_{val})$  $m \in \{1,...,M\}$ 

• Train a new model on  $\mathcal{D}_{train}$  ∪  $\mathcal{D}_{val}$  using  $\theta_{\widehat{m}}$ , $\left\langle h_{\widehat{m}}^{\pm}\right\rangle$ 

• Now  $err(h_{\hat{m}}^+, D_{test})$  is a good estimate of  $err(h_{\hat{m}}^+)$ !

How do we pick hyperparameter settings to try?

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# General **Methods** for Hyperparameter **Optimization**

- · Idea: set the hyperparameter performance metric of the me
- Issue: if we have many hyper take on lots of different value test all possible combinations
- Commonly used methods:
	- Grid search
	- Random search
	- · Bayesian optimization (us to optimize the hyperpara https://arxiv.org/pdf/181
		- Evolutionary algorithms
		- Graduate-student descen

# Grid Search vs. Random Search (Bergstra and Bengio, 2012)

**Grid Layout** 



# In general, which hyperparameter optimization method do you think will perform better?



# **Grid Search**

## Random Search

Start the presentation to see live content. For screen share software, share the entire screen. Get help at pollev.com/app

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# Grid Search vs. Random Search (Bergstra and Bengio, 2012)



Important parameter

Grid and random search of nine trial  $f(x, y) = g(x) + h(y) \approx g(x)$  with Above each square  $g(x)$  is shown in  $h(y)$  is shown in yellow. With grid se in three distinct places. With random distinct values of  $g$ . This failure of gr the exception in high dimensional hy Key Takeaways

- Differences between training, validation and test datasets in the model selection process
- Cross-validation for model selection
- Relationship between training, hyperparameter optimization and model selection
- Grid search vs. random search for hyperparameter optimization