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

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5/23/23

Lecture 5 Polls

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Front Matter

- Announcements:
 - PA1 released 5/18, due 5/25 at 11:59 PM
- Recommended Readings:
 - Daumé III, Chapter 2: Limits of Learning

Recall: Setting k

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

Recall: Setting k

- Theorem:
 - If k is some function of N s.t. $k(N) \to \infty$ and $\frac{k(N)}{N} \to 0$ as $N \to \infty$...
 - ... then (under certain assumptions) the true error of a kNN model \rightarrow the Bayes error rate
- Practical heuristics:

•
$$k = \lfloor \sqrt{N} \rfloor$$

•
$$k = 3$$

• 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

- 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

 Model = set of all possible nearest neighbors classifiers

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

• Hyperparameters = k, distance metric, how to scale features, the -breaking

Model Selection with Test Sets

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

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

• 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 \mathcal{D}_{test} and choose the one with lowest test error:

$$\widehat{m} = \underset{m \in \{1, \dots, M\}}{\operatorname{argmin}} \operatorname{err}(h_m, \mathcal{D}_{test})$$

Model Selection with Test Sets?

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• Evaluate each one using \mathcal{D}_{test} and choose the one with lowest test error:

$$\widehat{m} = \underset{m \in \{1, \dots, M\}}{\operatorname{argmin}} \operatorname{err}(h_m, \mathcal{D}_{test})$$

• Is $err(h_{\widehat{m}}, \mathcal{D}_{test})$ a good estimate of $err(h_{\widehat{m}})$?

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:

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

• 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 \mathcal{D}_{val} and choose the one with lowest *validation* error:

$$\widehat{m} = \underset{\{h_{1}, h_{2}, \dots, h_{M}\}}{\operatorname{argmin}} \operatorname{err}(h_{m}, \mathcal{D}_{val})$$

• Now $err(h_{\widehat{m}}, \mathcal{D}_{test})$ is a good estimate of $err(h_{\widehat{m}})!$

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:

$$\theta_1, \theta_2, \dots, \theta_M$$

• Learn a classifier for each setting using only \mathcal{D}_{train} :

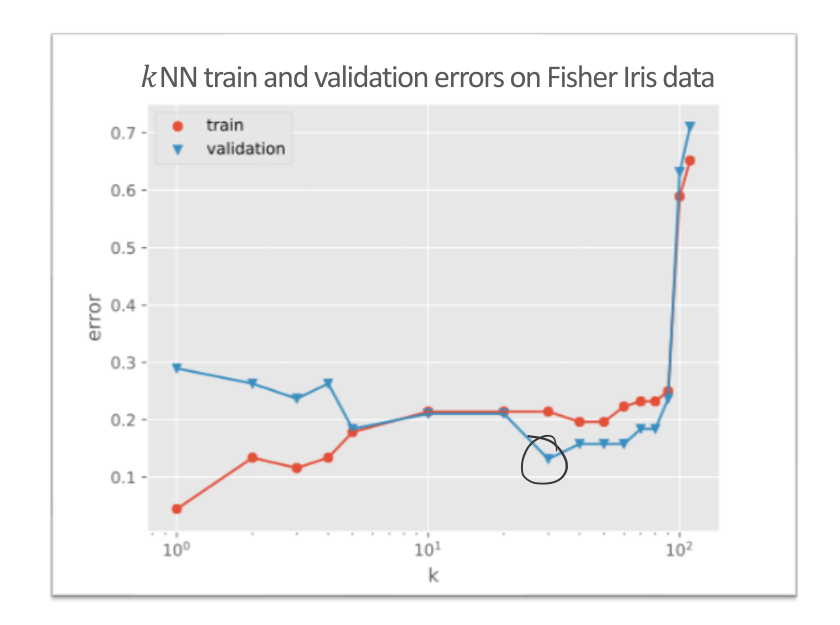
$$h_1, h_2, ..., h_M$$

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

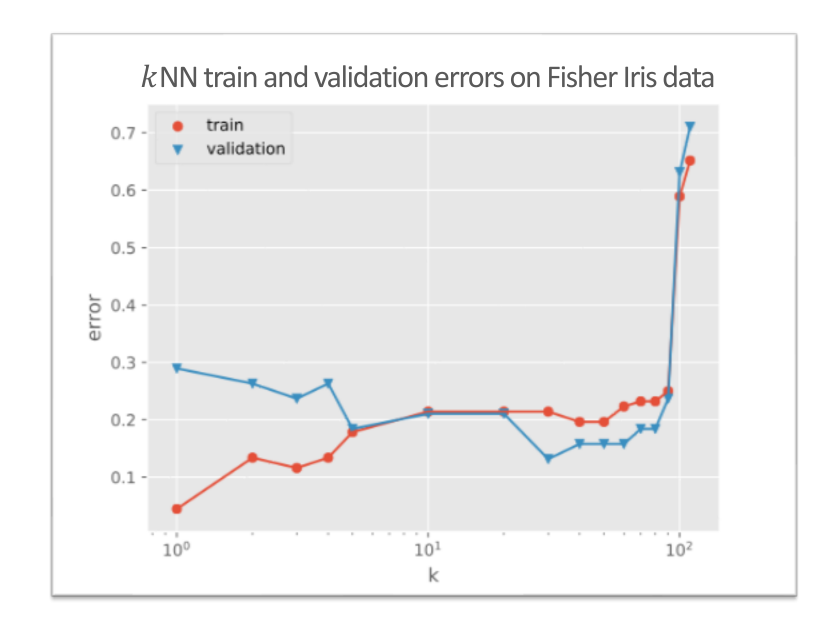
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Setting k for k NN with Validation Sets

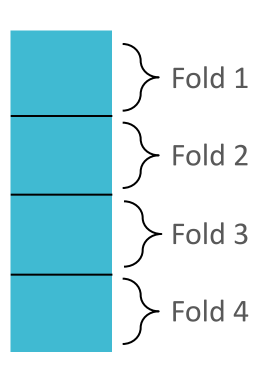


How should we partition our dataset?



• Given \mathcal{D} , split \mathcal{D} into K equally sized datasets or folds:

$$\mathcal{D}_1, \mathcal{D}_2, \dots, \mathcal{D}_K$$

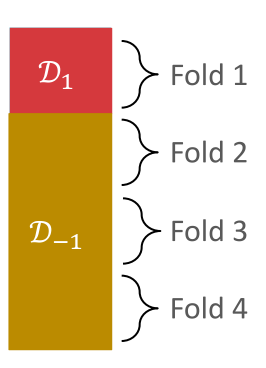


- Let h_{-i} be the classifier learned using $\mathcal{D}_{-i} = \underline{\mathcal{D}} \setminus \underline{\mathcal{D}}_i$ (all folds other than \mathcal{D}_i) and let $e_i = err(h_{-i}, \mathcal{D}_i)$
- The *K*-fold cross validation error is

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

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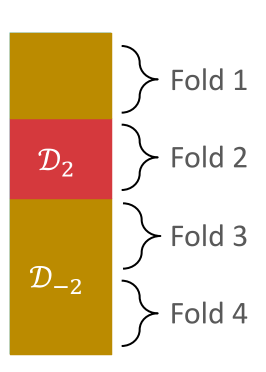


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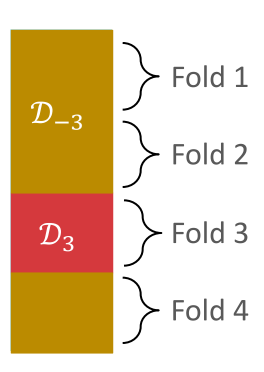


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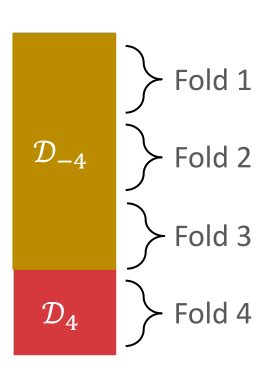


- Fold 1 Fold 2 Let h_{-i} be the classifier learned using $\mathcal{D}_{-i} = \mathcal{D} \setminus \mathcal{D}_i$ (all folds other than \mathcal{D}_i) and let $e_i = err(h_{-i}, \mathcal{D}_i)$
 - The *K*-fold cross validation error is

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

• Given \mathcal{D} , split \mathcal{D} into K equally sized datasets or folds:

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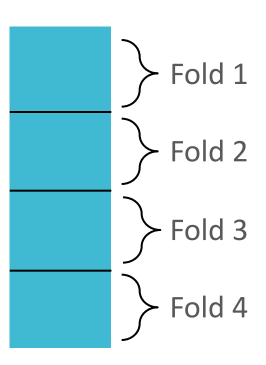
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- The *K*-fold cross validation error is

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

• Given \mathcal{D} , split \mathcal{D} into K equally sized datasets or folds:

$$\mathcal{D}_1, \mathcal{D}_2, \dots, \mathcal{D}_K$$

Use each one as a validation set once:



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

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

- Special case when K = N: Leave-one-out cross-validation
- Choosing between M candidates requires training MK times

Summary

	Input	Output
Training	training datasethyperparameters	 best model parameters
Hyperparameter Optimization	training datasetvalidation dataset	best hyperparameters
Cross-Validation	training datasetvalidation dataset	 cross-validation error
Testing	test datasetclassifier	• test error

Hyperparameter Optimization

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

$$\theta_1, \theta_2, \dots, \theta_M$$

• 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:

$$\widehat{m} = \underset{m \in \{1, \dots, M\}}{\operatorname{argmin}} \operatorname{err}(h_m, \mathcal{D}_{val})$$

• Now $err(h_{\widehat{m}}^+, \mathcal{D}_{test})$ is a good estimate of $err(h_{\widehat{m}}^+)$!

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

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

$$\theta_1, \theta_2, \dots, \theta_M$$

• 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:

$$\widehat{m} = \underset{m \in \{1, \dots, M\}}{\operatorname{argmin}} \operatorname{err}(h_m, \mathcal{D}_{val})$$

- Train a new model on $\mathcal{D}_{train} \cup \mathcal{D}_{val}$ using $\theta_{\widehat{m}}$, $h_{\widehat{m}}^+$
- Now $err(h_{\widehat{m}}^+, \mathcal{D}_{test})$ is a good estimate of $err(h_{\widehat{m}}^+)$!

How do we pick hyperparameter settings to try?

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

$$\theta_1, \theta_2, \dots, \theta_M$$

• 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:

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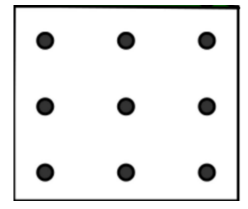
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General Methods for Hyperparameter Optimization

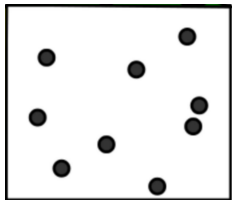
- Idea: set the hyperparameters to optimize some performance metric of the model
- Issue: if we have many hyperparameters that can all take on lots of different values, we might not be able to test all possible combinations
- Commonly used methods:
 - Grid search
 - Random search
- Bayesian optimization (used by Google DeepMind to optimize the hyperparameters of AlphaGo: https://arxiv.org/pdf/1812.06855v1.pdf)
 - Evolutionary algorithms
- Graduate-student descent

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

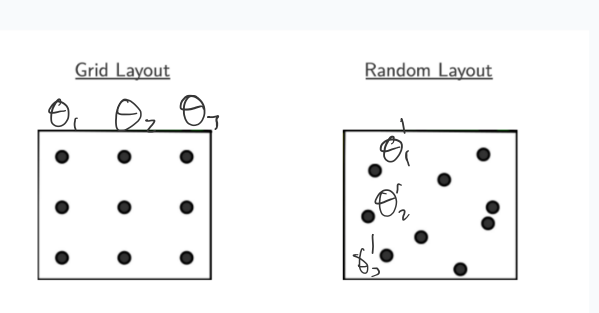
Grid Layout



Random Layout



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

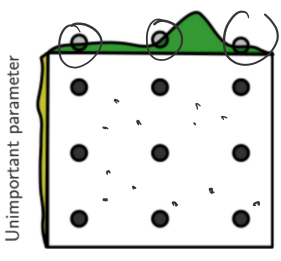


Grid Search

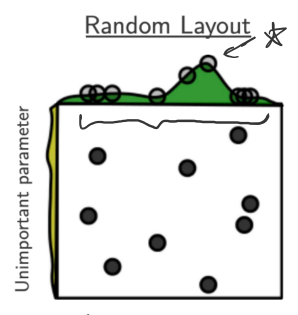
Random Search

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

Grid Layout



Important parameter



Important parameter

Grid and random search of nine trials for optimizing a function $f(x,y) = g(x) + h(y) \approx g(x)$ with low effective dimensionality. Above each square g(x) is shown in green, and left of each square h(y) is shown in yellow. With grid search, nine trials only test g(x) in three distinct places. With random search, all nine trials explore distinct values of g. This failure of grid search is the rule rather than the exception in high dimensional hyper-parameter optimization.

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