# **Lecture 3: Model Selection**

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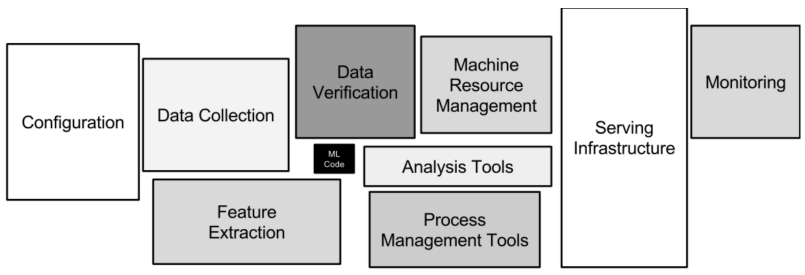
# **Evaluation**

- To know whether we can *trust* our method or system, we need to evaluate it.
- If you cannot measure it, you cannot improve it.
- Model selection: choose between different models in a data-driven way.
- Convince others that your work is meaningful
  - Peers, leadership, clients, yourself(!)
- Keep evaluating relentlessly, adapt to changes

# **Designing Machine Learning systems**

- Just running your favourite algorithm is usually not a great way to start
- Consider the problem at large
  - Do you want to understand phenomena or do black box modelling?
  - How to define and measure success? Are there costs involved?
  - Do you have the right data? How can you make it better?
- Build prototypes early-on to evaluate the above.

- Analyze your model's mistakes
  - Should you collect more, or additional data?
  - Should the task be reformulated?
  - Often a higher payoff than endless finetuning
- Technical debt: creation-maintenance trade-off
  - Very complex machine learning systems are hard/impossible to put into practice
  - See 'Machine Learning: The High Interest Credit Card of Technical Debt'



Only a small fraction of real-world ML systems is composed of the ML code

#### Real world evaluations

- Evaluate predictions, but also how outcomes improve *because of them*
- Feedback loops: predictions are fed into the inputs, e.g. as new data, invalidating models
- The signal your model found may just be an artifact of your biased data
  - When possible, try to *interpret* what your model has learned
  - See 'Why Should I Trust You?' by Marco Ribeiro et al.



(a) Husky classified as wolf

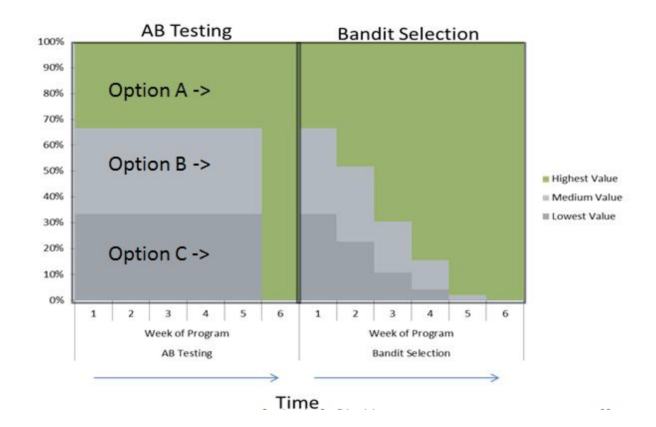


(b) Explanation

- Adversarial situations (e.g. spam filtering) can subvert your predictions
- Do A/B testing (or bandit testing) to evaluate algorithms in the wild

# A/B and bandit testing

- Test a single innovation (or choose between two models)
- Have most users use the old system, divert small group to new system
- Evaluate and compare performance
- Bandit testing: smaller time intervals, direct more users to currently winning system



# Performance estimation techniques

- We do not have access to future observations
- Evaluate models as if they are predicting the future
- Set aside data for objective evaluation
  - How?

# The holdout (simple train-test split)

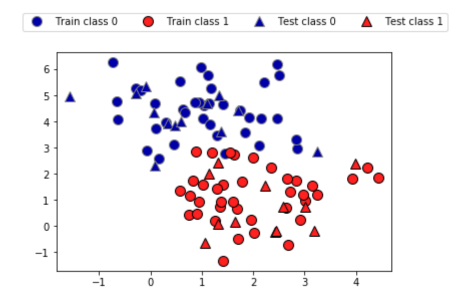
We've already seen the most basic form of evaluation:

- *Randomly* split data (and corresponding labels) into training and test set (75%-25%)
- Train (fit) a model on the training data
- Score a model on the test data (comparing predicted and true labels)
  - We are interested in how well the model generalizes to new (test) data

#### In scikit-learn: train\_test\_split

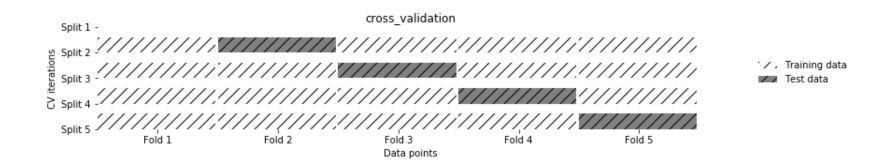
```
X, y = make_blobs(centers=2, random_state=0)
# split data and labels into a training and a test set
X_train, X_test, y_train, y_test = train_test_split(X, y, random_state=0)
# Fit a model to the training set
model = LogisticRegression().fit(X_train, y_train)
# Evaluate on the test data
test_score = model.score(X_test, y_test)
```

Test set score: 0.92



# **Cross-validation**

- What if one random split yields different models (and scores) than another?
- Reduce bias by testing on every point exactly once
- *k-fold cross-validation* (CV): split (randomized) data into *k* equal-sized parts, called *folds* 
  - First, fold 1 is the test set, and folds 2-5 comprise the training set
  - Then, fold 2 is the test set, folds 1,3,4,5 comprise the training set
  - Compute *k* evaluation scores, aggregate afterwards (e.g. take the mean)



#### In scikit-learn:

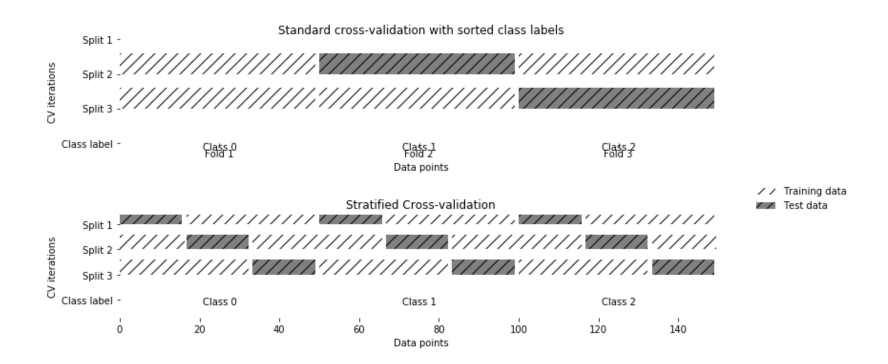
- cross\_val\_score function with learner, data, labels, number of folds
- Returns list of all scores. Models are built internally, but not returned
- Defaults: 3-fold CV, accuracy (classification) or  $\mathbb{R}^2$  (regression)
- Note that there can be quite some *variance* in the results
  - Depends on the stability of the model and the amount of training data
  - Typically, the more training data, the more stable the models

#### Benefits and drawbacks of cross-validation

- More robust: every training example will be in a test set exactly once
  - Model is evaluated on all samples, needs to do well on all
  - With a train-test split, we can be
    - 'lucky': all easy examples in test set
    - o 'unlucky: all hard examples in test set
- Shows how *sensitive* the model is to the exact training set
- Better estimation of true performance
  - 10-fold CV uses 90% of all data for training (vs 75% for holdout)
  - The higher *k*, the more accurate the estimation
- Disadvantage: computational cost, roughly *k* times slower than holdout
- Unstable models (e.g. deep learning) may not converge for every fold

### Stratified K-Fold cross-validation

- If the data is *unbalanced*, some classes have many fewer samples
- Likely that some classes are not present in the test set
- Stratification: *proportions* between classes are conserved in each fold
  - Order examples per class
  - Separate the samples of each class in *k* sets (strata)
  - Combine corresponding strate into folds



#### In scikit-learn:

- Uses stratified cross-validation by default for classification
- Normal cross-validation for regression
- Both are non-randomized (samples are not shuffled beforehand)
  - ordered data (e.g. time series) should never be randomized
- You can build folds manually with KFold or StratifiedKFold
  - randomizable (shuffle parameter)

```
kfold = KFold(n_splits=5, shuffle=False) # Not stratified
cross_val_score(logreg, iris.data, iris.target, cv=kfold
skfold = StratifiedKFold(n_splits=5, shuffle=True)
cross_val_score(logreg, iris.data, iris.target, cv=skfold)

Cross-validation scores KFold(n_splits=5):
[1.     0.933     0.433     0.967     0.433]
Cross-validation scores StratifiedKFold(n_splits=5, shuffle=True):
[1.     1.     0.867     0.967     1.     ]
```

### Can you explain this result?

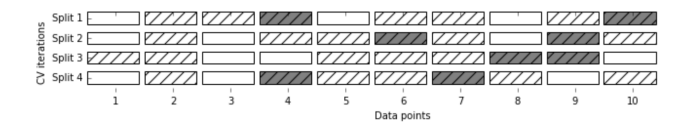
#### Leave-One-Out cross-validation

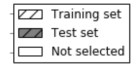
- *k* fold cross-validation with *k* equal to the number of samples
- Completely unbiased (in terms of data splits), but computationally expensive
- But: generalizes *less* well towards unseen data
  - The training sets are correlated (overlap heavily)
  - Overfits on the data used for (the entire) evaluation
  - A different sample of the data can yield different results
- Recommended only for small datasets

```
loo = LeaveOneOut()
scores = cross_val_score(logreg, iris.data, iris.target, cv=loo)
Number of cv iterations: 150
Mean accuracy: 0.95
```

# Shuffle-Split cross-validation

- Samples a number of samples (train\_size) randomly as the training set, and a disjoint number of samples (test\_size) as the test set
- Repeat this procedure n\_iter times, obtaining n\_iter scores
- Handy when using very large datasets
- Example with train\_size = 5, test\_size = 2, n\_iter = 4





#### In scikit-learn:

- ShuffleSplit and StratifiedShuffleSplit (recommended for classification)
- train\_size and test\_size can be absolute numbers or a percentage of the total dataset

```
shuffle_split = StratifiedShuffleSplit(test_size=.5, train_size=.5, n_sp
lits=10)
scores = cross_val_score(logreg, iris.data, iris.target, cv=shuffle_split)

Cross-validation scores:
[0.907 0.96 0.92 0.96 0.973 0.96 0.973 0.96 0.92 0.96 ]
```

Note: this is related to *bootstrapping*:

- Sample *n* (total number of samples) data points, with replacement, as training set (the bootstrap)
- Use the unsampled (out-of-bootstrap) samples as the test set
- Repeat n\_iter times, obtaining n\_iter scores
- Not supported in scikit-learn, use Shuffle-Split instead.
  - With train\_size=0.66, test\_size=0.34
  - You can prove that bootstraps include 66% of all data points on average

# Repeated cross-validation

- Cross-validation is still biased in that the initial split can be made in many ways
- Repeated, or n-times-k-fold cross-validation:
  - Shuffle data randomly, do k-fold cross-validation
  - Repeat n times, yields n times k scores
- Unbiased, very robust, but n times more expensive

# Cross-validation with groups

- Sometimes the data contains inherent groups:
  - Blood analysis results on specific patients
  - Facial expressions of specific people
- With normal cross-validation, data from the same persion may end up in the training *and* test set
- We want to measure how well the model generalizes to *other* people
- We want to make sure that data points fom one person are in *either* the training or test set
  - This is called *grouping* or *blocking*
  - Leave-one-subject-out cross-validation: create test set for each user individually

#### In scikit-learn

- Add an array with group membership to cross\_val\_scores
- Use GroupKFold with the number of groups as CV procedure

```
groups = [0, 0, 0, 1, 1, 1, 2, 2, 2, 3, 3, 3]
scores = cross_val_score(logreg, X, y, groups, cv=GroupKFold(n_splits=4)))

cross_val_score(logreg, X, y, groups, cv=GroupKFold(n_splits=4))
Cross-validation scores:
[0.667 0.667 1. 0.667]
```

# Choosing a performance estimation procedure

No strict rules, only guidelines:

- Always use stratification for classification
- Use holdout for very large datasets (e.g. >1.000.000 examples)
  - Or when learners don't always converge (e.g. deep learning)
- Choose *k* depending on dataset size and resources
  - Use leave-one-out for small datasets (e.g. <500 examples)
  - Use cross-validation otherwise
    - Most popular (and theoretically sound): 10-fold CV
    - Literature suggests 5x2-fold CV is better
- Use grouping or leave-one-subject-out for grouped data

# **Evaluation Metrics and scoring**

Keep the end-goal in mind

## **Evaluation vs Optimization**

- Each algorithm optimizes a given objective function (on the training data)
  - E.g. remember L2 loss in Ridge regression

$$\mathcal{L}_{ridge} = \sum_{i} (y_i - \sum_{j} x_{i,j} w_j)^2 + \alpha \sum_{i} w_i^2$$

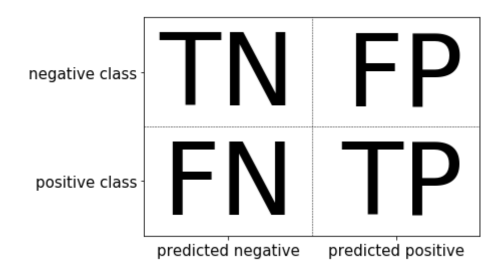
- The choice of function is limited by what can be efficiently optimized
  - E.g. gradient descent requires a differentiable loss function
- We *evaluate* the resulting model with a score that makes sense in the real world
  - E.g. percentage of correct predictions (on a test set)
- We also tune the algorithm's hyperparameters to maximize that score

# **Binary classification**

- We have a positive and a negative class
- 2 different kind of errors:
  - False Positive (type I error): model predicts positive while the true label is negative
  - False Negative (type II error): model predicts negative while the true label is positive
- They are not always equally important
  - Which side do you want to err on for a medical test?

#### **Confusion matrices**

- We can represent all predictions (correct and incorrect) in a confusion matrix
  - n by n array (n is the number of classes)
  - Rows correspond to true classes, columns to predicted classes
  - Each entry counts how often a sample that belongs to the class corresponding to the row was classified as the class corresponding to the column.
  - For binary classification, we label these true negative (TN), true positive (TP), false negative (FN), false positive (FP)



#### **Predictive accuracy**

• Accuracy is one of the measures we can compute based on the confusion matrix:

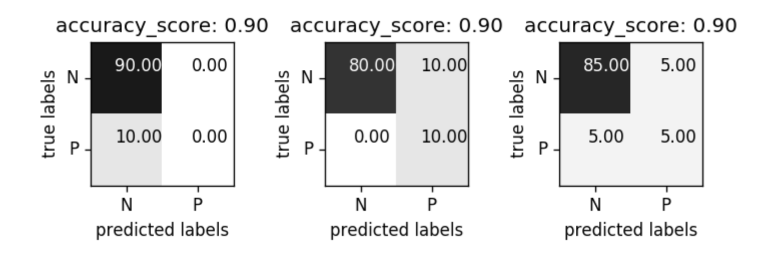
$$Accuracy = \frac{TP + TN}{TP + TN + FP + FN}$$

- In sklearn: use confusion\_matrix and accuracy\_score from sklearn.metrics.
- Accuracy is also the default evaluation measure for classification

```
confusion_matrix(y_test, y_pred):
  [[49  4]
  [ 5 85]]
accuracy_score(y_test, y_pred): 0.9370629370629371
model.score(X_test, y_test): 0.9370629370629371
```

#### The problem with accuracy: imbalanced datasets

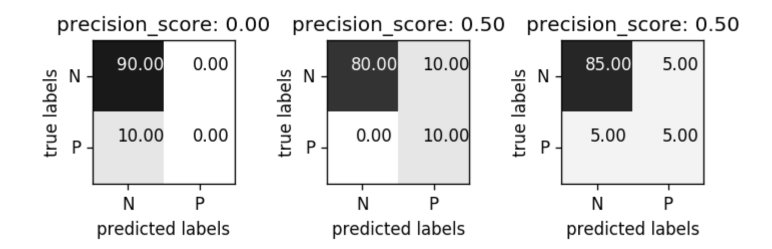
- The type of error plays an even larger role if the dataset is imbalanced
  - One class is much more frequent than the other, e.g. credit fraud
  - Is a 99.99% accuracy good enough?
- Are these three models really equally good?



#### **Precision** is used when the goal is to limit FPs

- Clinical trails: you only want to test drugs that really work
- Search engines: you want to avoid bad search results

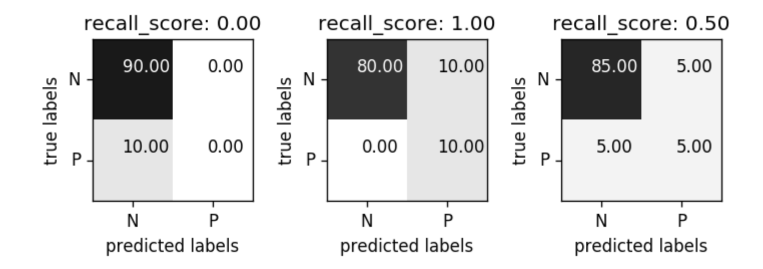
$$Precision = \frac{TP}{TP + FP}$$



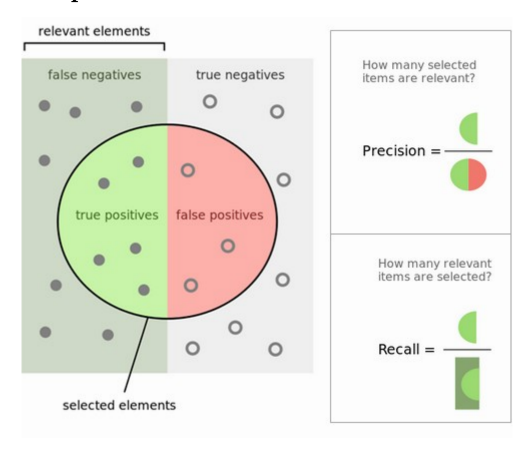
#### **Recall** is used when the goal is to limit FNs

- Cancer diagnosis: you don't want to miss a serious disease
- Search engines: You don't want to omit important hits
- Also know as sensitivity, hit rate, true positive rate (TPR)

$$Recall = \frac{TP}{TP + FN}$$

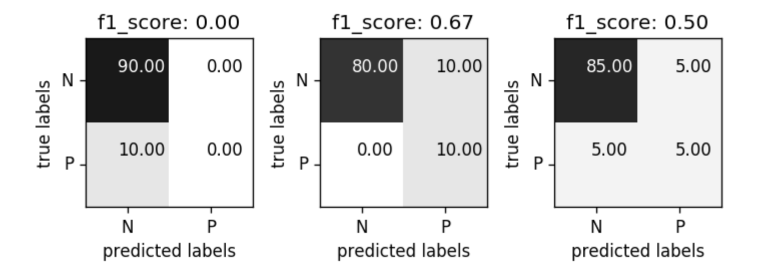


### Comparison



**F1-score** or F1-measure trades off precision and recall:

$$F1 = 2 \cdot \frac{\text{precision} \cdot \text{recall}}{\text{precision} + \text{recall}}$$



### Classification measure Zoo

		True condition			
	Total population	Condition positive	Condition negative	$\frac{\sum Condition\ positive}{\sum Total\ population}$	Accuracy (ACC) = $\frac{\Sigma \text{ True positive} + \Sigma \text{ True negative}}{\Sigma \text{ Total population}}$
Predicted condition	Predicted condition positive	True positive, Power	False positive, Type I error	Positive predictive value (PPV), Precision $= \frac{\Sigma \text{ True positive}}{\Sigma \text{ Predicted condition positive}}$	False discovery rate (FDR) = $\Sigma$ False positive $\Sigma$ Predicted condition positive
	Predicted condition negative	False negative, Type II error	True negative	False omission rate (FOR) = $\Sigma$ False negative $\Sigma$ Predicted condition negative	Negative predictive value (NPV) = $\frac{\Sigma \text{ True negative}}{\Sigma \text{ Predicted condition negative}}$
		True positive rate (TPR), Recall, Sensitivity, probability of detection = $\frac{\Sigma}{\Sigma}$ True positive	False positive rate (FPR), Fall-out, probability of false alarm = $\frac{\Sigma}{\Sigma}$ Condition negative	Positive likelihood ratio (LR+) = $\frac{TPR}{FPR}$	Diagnostic odds F <sub>1</sub> score =
		False negative rate (FNR), Miss rate $= \frac{\Sigma \text{ False negative}}{\Sigma \text{ Condition positive}}$	$\begin{aligned} \text{Specificity (SPC), Selectivity, True negative rate} \\ (\text{TNR}) &= \frac{\Sigma \text{ True negative}}{\Sigma \text{ Condition negative}} \end{aligned}$	Negative likelihood ratio (LR-) = $\frac{FNR}{TNR}$	ratio (DOR) = $\frac{LR+}{LR-}$ $\frac{2}{\frac{1}{Recall} + \frac{1}{Precision}}$

<a href="https://en.wikipedia.org/wiki/Precision">https://en.wikipedia.org/wiki/Precision</a> and recall <a href="https://en.wikipedia.org/wiki/Precision">(https://en.wikipedia.org/wiki/Precision</a> and recall)"

#### Averaging scores per class

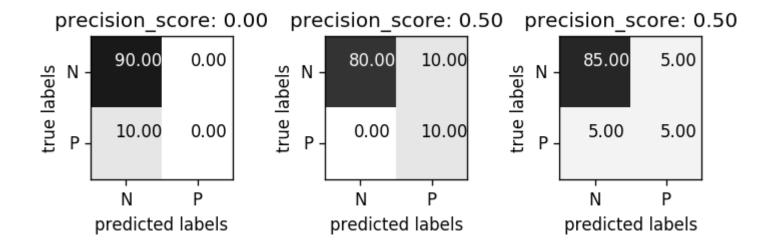
- Study the scores *by class* (in scikit-learn: classification\_report)
  - One class viewed as positive, other(s) als negative
  - Support: number of samples in each class
  - Last line: weighted average over the classes (weighted by number of samples in each class)
- Averaging for scoring measure R across C classes (also for multiclass):
  - micro: count total number of TP, FP, TN, FN
  - macro

$$\frac{1}{C} \sum_{c \in C} R(y_c, \hat{y_c})$$

• weighted ( $w_c$ : ratio of examples of class c)

$$\sum_{c \in C} w_c R(y_c, \hat{y_c})$$

### Example

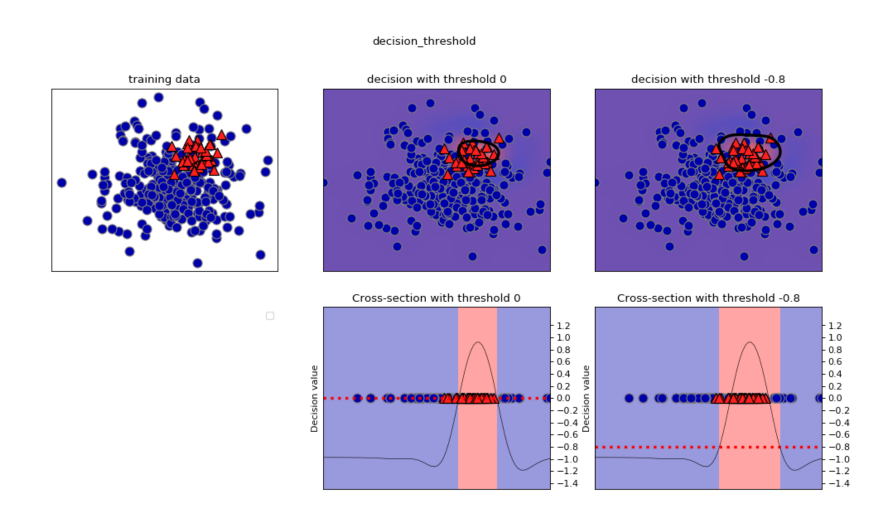


Matrix 1				
	precision	recall	f1-score	support
_				
0	0.90	1.00	0.95	90
1	0.00	0.00	0.00	10
			0.00	100
accuracy	0.45	0.50	0.90	100
macro avg	0.45	0.50	0.47	100
weighted avg	0.81	0.90	0.85	100
Matrix 2				
	precision	recall	f1-score	support
	-			
0	1.00	0.89	0.94	90
1	0.50	1.00	0.67	10
accuracy			0.90	100
macro avg	0.75	0.94	0.80	100
weighted avg	0.95	0.90	0.91	100
Matrix 3				
	precision	recall	f1-score	support
0	0.94	0.94	0.94	90
1	0.50	0.50	0.50	10
			0.00	100
accuracy			0.90	100
macro avg	0.72	0.72	0.72	100
weighted avg	0.90	0.90	0.90	100

# Taking uncertainty into account

- Remember that many classifiers actually return a probability per class
  - We can retrieve it with decision\_function and predict proba
- For binary classification, we threshold at 0 for decision\_function and 0.5 for predict\_proba by default
- However, depending on the evaluation measure, you may want to threshold differently to fit your goals
  - For instance, when a FP is much worse than a FN
  - This is called *threshold calibration*

- Imagine that we want to avoid misclassifying a positive (red) point
- Points within decision boundary (black line) are classified positive
- Lowering the decision treshold (bottom figure): fewer FN, more FP



- Studying the classification report, we see that lowering the threshold yields:
  - higher recall for class 1 (we risk more FPs in exchange for more TP)
  - lower precision for class 1
- We can often trade off precision for recall

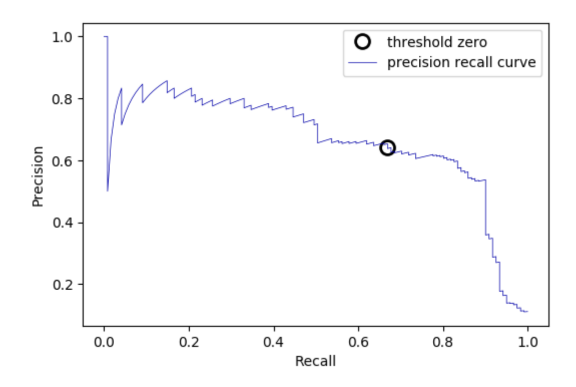
Threshold 0				
	precision	recall	f1-score	support
0	0.91	0.96	0.93	96
1	0.67	0.47	0.55	17
accuracy			0.88	113
macro avg	0.79	0.71	0.74	113
weighted avg	0.87	0.88	0.88	113
Threshold -0.	8			
	precision	recall	f1-score	support
0	0.98	0.92	0.95	96
1	0.65	0.88	0.75	17
accuracy			0.91	113
macro avg	0.81	0.90	0.85	113
weighted avg	0.93	0.91	0.92	113

## **Precision-Recall curves**

- The best threshold depends on your application, should be driven by real-world goals.
- You can have arbitrary high recall, but you often want reasonable precision, too.
- It is not clear beforehand where the optimale trade-off (or *operating point*) will be, so it is useful to look at all possible thresholds
- Plotting precision against recall for all thresholds yields a **precision-recall curve**

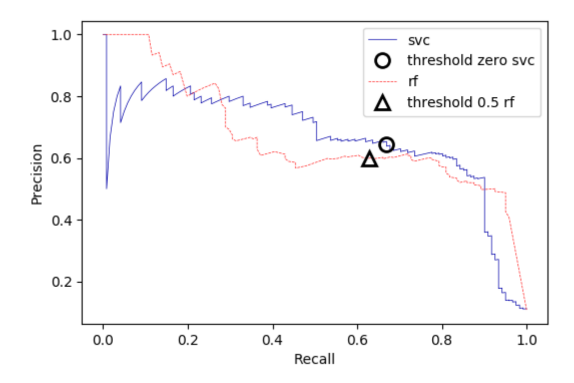
- In scikit-learn, this is included in the sklearn.metrics module
- Returns all precision and recall values for all thresholds
  - Vary threshold from lowest to highest decision function score in the predictions
  - Or from highest to lowest class probability

- The default tradeoff (chosen by the predict method) is shown as *threshold zero*.
  - Higher threshold, more precision (move left)
  - Lower threshold, more recall (move right)
- The closer the curve stays to the upper-right corner, the better
  - High precision and high recall
- Here, it is possible to still get a precision of 0.5 with high recall



### **Model selection**

- Different classifiers work best in different parts of the curve (at different operating points)
- RandomForest (in red) performs better at the extremes, SVM better in center
- The area under the precision-recall curve (AUPRC) is often used as a general evaluation measure



Note that the F1-measure completely misses these subtleties

```
f1_score of random forest: 0.610
f1_score of svc: 0.656
```

- The area under the precision-recall curve is returned by the average\_precision\_score measure
  - It's actually a close approximation of the actual area
- This is a good automatic measure, but also hides the subtleties

Average precision of random forest: 0.660 Average precision of svc: 0.666

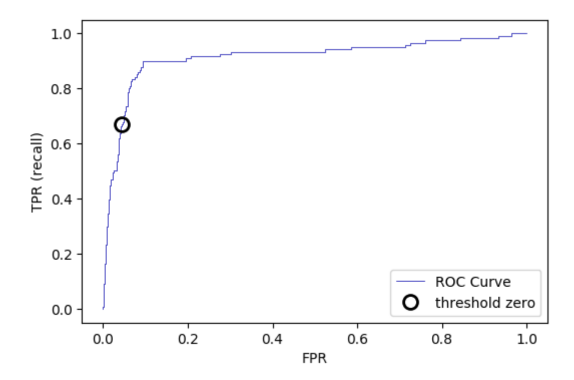
## Receiver Operating Characteristics (ROC) and AUC

- There is another trade-off between recall (true positive rate, TPR) and the false positive rate (FPR).
- The 2D space created by TPR and FPR is called the Receiver Operating Characteristics (ROC) space
- A model will be at one point in this ROC space

$$TPR = \frac{TP}{TP + FN}$$

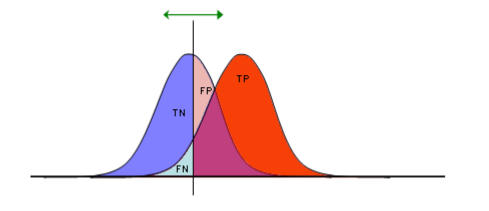
$$FPR = \frac{FP}{FP + TN}$$

- Varying the decision threshold yields the ROC curve
- It can be computed with the roc\_curve function
  - Lower threshold, more recall/TPR, move right
  - High threshold, fewer FPs, move left
- Ideal is close to the top left: high recall, low FPR
- Inspect the curve to find the preferred calibration
  - Here, we can get much higher recall with slightly worse FPR

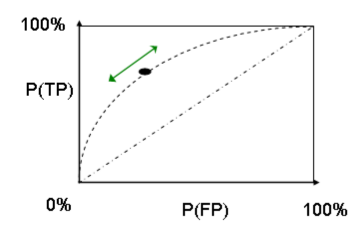


### Visualization

- The blue probability density shows the probability p(x) that the model predicts blue if a data point has a certain predicted probability x to be blue. Same for red.
- In a random classifier the probability densities completely overlap.
- All points with a predicted probability higher than the threshold are predicted positive, others negative
- As we increase the threshold, we'll get fewer FPs, more FNs. We move from right to left along the ROC curve.

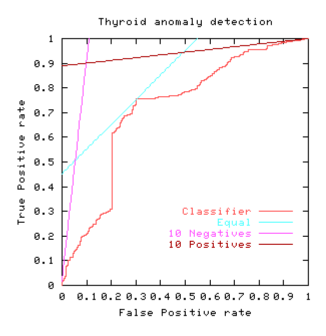


TP	FP
FN	TN
1	1



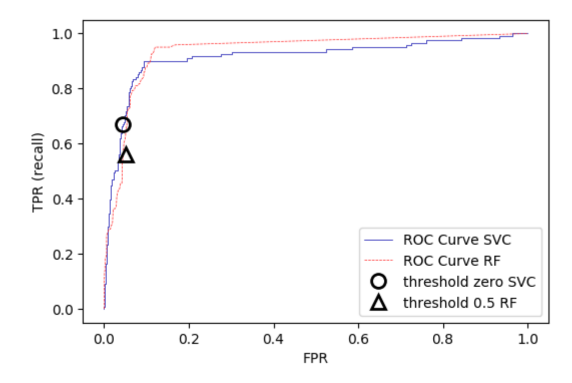
### **ROC** Isometrics

- Different *costs* can be involved for FP and FN
- This yields different *isometrics* (lines of equal cost) in ROC space
- The optimal threshold is the point on the ROC curve where the cost in minimal
  - If a FP and FN are weighted equally, cost lines follow the diagonal (blue line)
  - If a FP is 10 times worse than a FN: pink line
  - IF a FN is 10 times worse than a FP: red line



### **Model selection**

- Again, we can compare multiple models by looking at the ROC curves
- We can calibrate the threshold depending on whether we need high recall or low FPR
- We can select between algorithms (or hyperparameters) depending on the involved costs.



#### Area under the ROC curve

- A good summary measure is the area under the ROC curve (AUROC or AUC)
- Compute using the roc\_auc\_score
  - Don't use auc (uses less accurate trapezoidal rule)

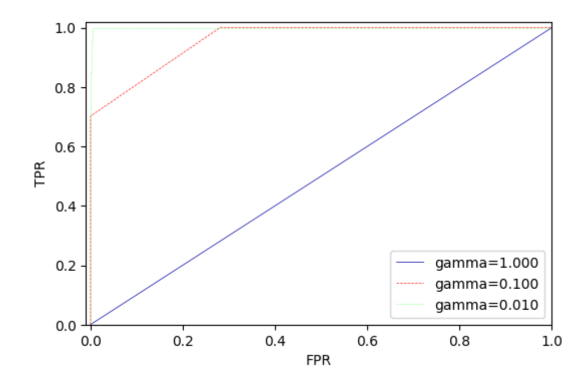
```
rf_auc = roc_auc_score(y_test, rf.predict_proba(X_test)[:, 1])
svc_auc = roc_auc_score(y_test, svc.decision_function(X_test))
AUC for Random Forest: 0.937
AUC for SVC: 0.916
```

### **Imbalanced classes**

- AUC is popular because it is insensitive to class imbalance
  - Random guessing always yields TPR=FPR
  - All points are on the diagonal line, hence an AUC of 0.5
  - Hint: use the visualization of TPR,FPR to see this

- Example: unbalanced digits
  - 3 models, ACC is the same, AUC not
  - If we optimize for ACC, our model could be just random guessing

```
gamma = 1.000 accuracy = 0.90 AUC = 0.5000
gamma = 0.100 accuracy = 0.90 AUC = 0.9582
gamma = 0.010 accuracy = 0.90 AUC = 0.9995
```



## Take home message

- AUC is highly recommended, especially on imbalanced data
- Remember to calibrate the threshold to your needs

## Multi-class classification

- Multiclass metrics are derived from binary metrics, averaged over all classes
- Let's consider the full (10-class) handwritten digit recognition data

### Confusion matrix

```
Accuracy: 0.953
Confusion matrix:

[[37  0  0  0  0  0  0  0  0  0  0]

[ 0  39  0  0  0  0  0  0  0  0  0]

[ 0  0  41  3  0  0  0  0  0  0]

[ 0  0  1  43  0  0  0  0  0  0]

[ 0  0  0  0  38  0  0  0  0  0]

[ 0  1  0  0  0  47  0  0  0  0]

[ 0  1  0  0  0  0  52  0  0  0]

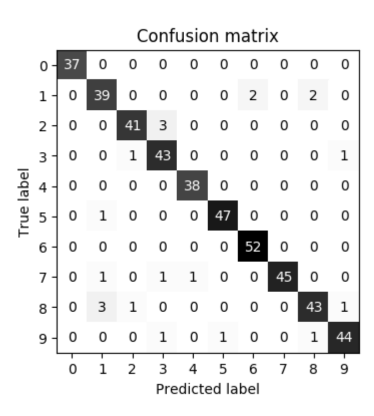
[ 0  1  0  1  1  0  0  45  0  0]

[ 0  3  1  0  0  0  0  0  43  1]

[ 0  0  0  1  0  1  0  0  1  44]
```

## Visualized as a heatmap

• Which digits are easy to predict? Which ones are confused?



## Precision, recall, F1-score now yield 10 per-class scores

	precision	recall	f1-score	support
0	1.00	1.00	1.00	37
1	0.89	0.91	0.90	43
2	0.95	0.93	0.94	44
3	0.90	0.96	0.92	45
4	0.97	1.00	0.99	38
5	0.98	0.98	0.98	48
6	0.96	1.00	0.98	52
7	1.00	0.94	0.97	48
8	0.93	0.90	0.91	48
9	0.96	0.94	0.95	47
accuracy			0.95	450
macro avg	0.95	0.95	0.95	450
weighted avg	0.95	0.95	0.95	450

## Different ways to compute average

- macro-averaging: computes unweighted per-class scores:  $\frac{\sum_{i=0}^{n} score_{i}}{n}$ 
  - Use when you care about each class equally much
- weighted averaging: scores are weighted by the relative size of the classes (support):  $\frac{\sum_{i=0}^{n} score_{i}weight_{i}}{n}$ 
  - Use when data is imbalanced
- micro-averaging: computes total number of FP, FN, TP over all classes, then computes scores using these counts:  $recall = \frac{\sum_{i=0}^{n} TP_i}{\sum_{i=0}^{n} TP_i + \sum_{i=0}^{n} FN_i}$ 
  - Use when you care about each sample equally much

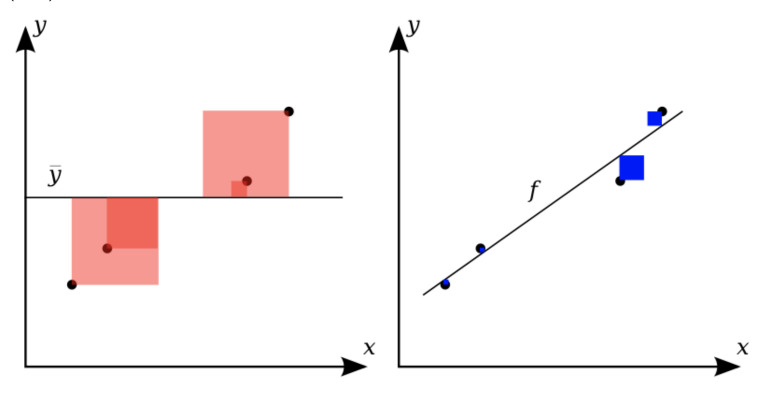
Micro average f1 score: 0.953 Weighted average f1 score: 0.953 Macro average f1 score: 0.954

# **Regression metrics**

Most commonly used are

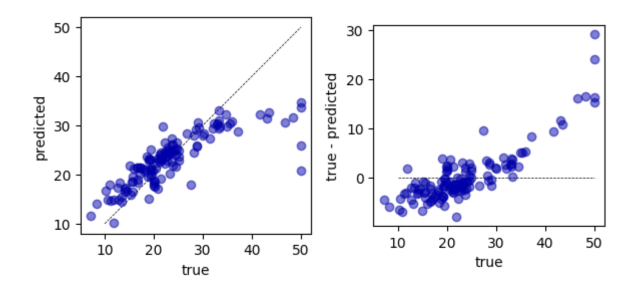
- (root) mean squared error:  $\frac{\sum_{i}(y_{pred_i} y_{actual_i})^2}{n}$
- mean absolute error:  $\frac{\sum_{i} |y_{pred_i} y_{actual_i}|}{n}$ 
  - Less sensitive to outliers and large errors
- R squared (r2):  $1 \frac{\sum_{i} (y_{pred_i} y_{actual_i})^2}{\sum_{i} (y_{mean} y_{actual_i})^2}$ 
  - Ratio of variation explained by the model / total variation
  - Between 0 and 1, but *negative* if the model is worse than just predicting the mean
  - Easier to interpret (higher is better).

• R squared: 1 - ratio of  $\sum_i (y_{pred_i} - y_{actual_i})^2$  (blue) and  $\sum_i (y_{mean} - y_{actual_i})^2$  (red)



## Visualizing errors

- Prediction plot (left): predicted vs actual target values
- Residual plot (right): residuals vs actual target values
  - Over- and underpredictions can be given different costs



# Using evaluation metrics in model selection

- You typically want to use AUC or other relevant measures in cross\_val\_score and GridSearchCV instead of the default accuracy.
- scikit-learn makes this easy through the **scoring** argument
  - But, you need to need to look the <u>mapping between the scorer</u> and the metric (<a href="http://scikit-learn.org/stable/modules/model evaluation.html#model-evaluation">http://scikit-learn.org/stable/modules/model evaluation.html#model-evaluation</a>)

Scoring	Function	Comment
Classification		
'accuracy'	metrics.accuracy_score	
'average_precision'	metrics.average_precision_score	
'f1'	metrics.f1_score	for binary targets
'f1_micro'	metrics.f1_score	micro-averaged
'f1_macro'	metrics.f1_score	macro-averaged
'f1_weighted'	metrics.f1_score	weighted average
'f1_samples'	metrics.f1_score	by multilabel sample
'neg_log_loss'	metrics.log_loss	requires predict_proba support
'precision' etc.	metrics.precision_score	suffixes apply as with 'f1'
'recall' etc.	metrics.recall_score	suffixes apply as with 'f1'
'roc_auc'	metrics.roc_auc_score	
Clustering		
'adjusted_rand_score'	metrics.adjusted_rand_score	
Regression		
'neg_mean_absolute_error'	metrics.mean_absolute_error	
'neg_mean_squared_error'	metrics.mean_squared_error	
'neg_median_absolute_error'	metrics.median_absolute_error	
'r2'	metrics.r2_score	

### Or simply look up like this:

#### Available scorers:

['accuracy', 'adjusted\_mutual\_info\_score', 'adjusted\_rand\_score', 'average \_precision', 'balanced\_accuracy', 'brier\_score\_loss', 'completeness\_scor e', 'explained\_variance', 'f1', 'f1\_macro', 'f1\_micro', 'f1\_samples', 'f1\_weighted', 'fowlkes\_mallows\_score', 'homogeneity\_score', 'jaccard', 'jaccard\_rd\_macro', 'jaccard\_micro', 'jaccard\_samples', 'jaccard\_weighted', 'max\_er ror', 'mutual\_info\_score', 'neg\_log\_loss', 'neg\_mean\_absolute\_error', 'neg\_mean\_squared\_error', 'neg\_mean\_squared\_log\_error', 'neg\_median\_absolute\_error', 'normalized\_mutual\_info\_score', 'precision', 'precision\_macro', 'precision\_micro', 'precision\_samples', 'precision\_weighted', 'r2', 'recall', 'recall\_macro', 'recall\_micro', 'recall\_samples', 'recall\_weighted', 'roc\_auc', 'v measure score']

### Cross-validation with AUC

## Grid Search with accuracy and AUC

- With accuracy, gamma=0.0001 is selected
- With AUC, gamma=0.01 is selected
  - Actually has better accuracy on the test set

```
Grid-Search with accuracy
Best parameters: {'gamma': 0.0001}
Best cross-validation score (accuracy)): 0.970
Test set AUC: 0.992
Test set accuracy: 0.973

Grid-Search with AUC
Best parameters: {'gamma': 0.01}
Best cross-validation score (AUC): 0.997
Test set AUC: 1.000
Test set accuracy: 1.000
```

# Final thoughts

- There exist techniques to correct label imbalance
  - Undersample the majority class, or oversample the minority class
  - SMOTE (Synthetic Minority Oversampling Technique) adds articifial *training* points by interpolating existing minority class points
    - Think twice before creating 'artificial' training data
- Cost-sensitive classification (not in sklearn)
  - Cost matrix: a confusion matrix with a costs associated to every possible type of error
  - Some algorithms allow optimizing on these costs instead of their usual loss function
  - Meta-cost: builds ensemble of models by relabeling training sets to match a given cost matrix
    - Black-box: can make any algorithm cost sensitive (but slower and less accurate)

- There are many more metrics to choose from
  - Cohen's Kappa: accuracy, taking into account the possibility of predicting the right class by chance
    - 1: perfect prediction, 0: random prediction, negative: worse than random
    - With  $p_0$  = accuracy, and  $p_e$  = accuracy of random classifier:

$$\kappa = \frac{p_o - p_e}{1 - p_e}$$

- Balanced accuracy: accuracy where each sample is weighted according to the inverse prevalence of its true class
  - o Identical to macro-averaged recall
- Matthews correlation coefficient: another measure that can be used on imbalanced data
  - 1: perfect prediction, 0: random prediction, -1: inverse prediction

$$MCC = \frac{tp \times tn - fp \times fn}{\sqrt{(tp + fp)(tp + fn)(tn + fp)(tn + fn)}}$$

# **Bias-Variance decomposition**

- When we repeat evaluation procedures multiple times, we can distinguish two sources of errors:
  - Bias: systematic error (independent of the training sample). The classifier always gets certain points wrong
  - Variance: error due to variability of the model with respect to the training sample. The classifier predicts some points accurately on some training sets, but inaccurately on others.
- There is also an intrinsic (noise) error, but there's nothing we can do against that.
- Bias is associated with underfitting, and variance with overfitting
- Bias-variance trade-off: you can often exchange variance for bias through regularization (and vice versa)
  - The challenge is to find the right trade-off (minimizing total error)
- Useful to understand how to tune or adapt learning algorithm

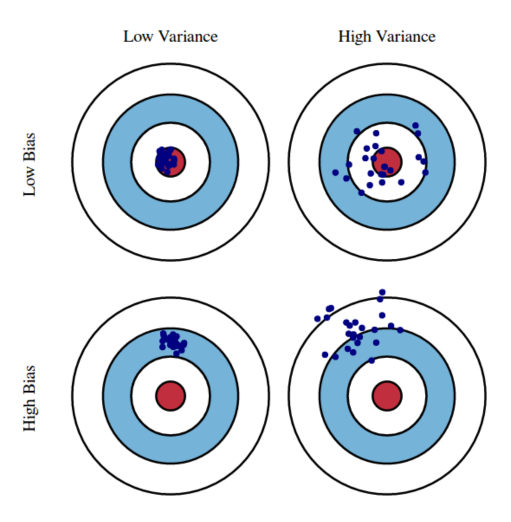
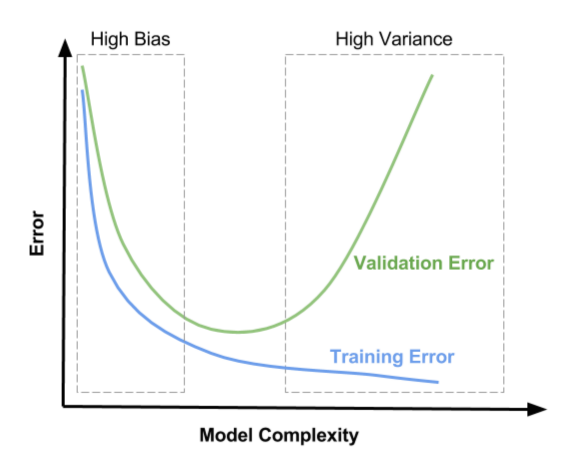


Fig. 1 Graphical illustration of bias and variance.

- Sadly, this is not yet supported by scikit-learn
- How to measure bias and variance (for regression):
  - Take 100 or more bootstraps (or shuffle-splits)
  - For each data point x:
    - $\circ bias(x)^2 = (x_{true} mean(x_{predicted}))^2$
    - $\circ \ variance(x) = var(x_{predicted})$
  - Total bias:  $\sum_{x} bias(x)^2 * w_x$ , with  $w_x$  the ratio of x occurring in the test set
  - Total variance:  $\sum_{x} variance(x) * w_x$

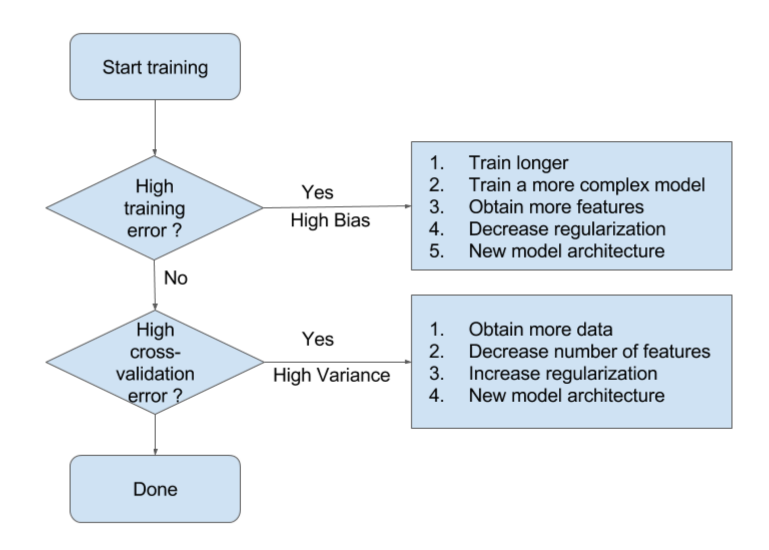
- General procedure for (binary) classification:
  - Take 100 or more bootstraps (or shuffle-splits)
  - Bias for any point x = misclassification ratio
    - If misclassified 50% of the time: bias(x) = 0.5
  - Variance for any point x is  $(1 (P(class_1)^2 + P(class_2)^2))/2$ 
    - $\circ$   $P(class_i)$  is ratio of class i predictions
    - When each class predicted half of the time:  $variance(x) = (1 (0.5^2 + 0.5^2))/2 = 0.25$
  - Total bias:  $\sum_{x} bias(x)^2 * w_x$ , with  $w_x$  the ratio of x occurring in the test data
  - Total variance:  $\sum_{x} variance(x) * w_x$

# Bias-variance and overfitting



- High bias means that you are likely underfitting
  - Do less regularization
  - Use a more flexible/complex model (another algorithm)
  - Use a bias-reduction technique (e.g. boosting, see later)
- High variance means that you are likely overfitting
  - Use more regularization
  - Get more data
  - Use a simpler model (another algorithm)
  - Use a variance-reduction techniques (e.g. bagging, see later)

## Bias-Variance Flowchart (Andrew Ng, Coursera)



# Hyperparameter tuning

Now that we know how to evaluate models, we can improve them by tuning their hyperparameters

We can basically use any optimization technique to optimize hyperparameters:

- Grid search
- Random search

More advanced techniques:

- Local search
- Racing algorithms
- Model-based optimization (see later)
- Multi-armed bandits
- Genetic algorithms

## **Grid Search**

- For each hyperparameter, create a list of interesting/possible values
  - E.g. For kNN: k in [1,3,5,7,9,11,33,55,77,99]
- Evaluate all possible combination of hyperparameter values
  - E.g. using cross-validation
- Select the hyperparameter values yielding the best results
- A naive approach would be to just loop over all combinations

```
for gamma in [0.001, 0.01, 0.1, 1, 10, 100]:
    for C in [0.001, 0.01, 0.1, 1, 10, 100]:
        # for each combination, train and evaluate an SVC
        svm = SVC(gamma=gamma, C=C);
        svm.fit(X_train, y_train);
        score = svm.score(X_test, y_test)
        # if we got a better score, store the score and parameters
        if score > best_score:
            best_score = score
            best_parameters = {'C': C, 'gamma': gamma}

Size of training set: 112     size of test set: 38
Best score: 0.97
Best parameters: {'C': 100, 'gamma': 0.001}
```

### Overfitting the parameters and the validation set

- Simply taking the best performing model yields optimistic results
- We've already used the test data to evaluate each hyperparameter setting!
- Hence, we don't have an independent test set to evaluate these hyperparameter settings
  - Information 'leaks' from test set into the final model
- Solution: Set aside part of the training data to evaluate the hyperparameter settings
  - Select best hyperparameters on validation set
  - Rebuild the model on the training+validation set
  - Evaluate optimal model on the test set

training set	validation set	test set
Model fitting	Hyperparameter optimization	Evaluation

```
# split data into train+validation set and test set
X_trainval, X_test, y_trainval, y_test = train_test_split(
    iris.data, iris.target, random_state=0)
# split train+validation set into training and validation set
X_train, X_valid, y_train, y_valid = train_test_split(
    X_trainval, y_trainval, random_state=1)

Size of training set: 84    size of validation set: 28    size of test set: 38

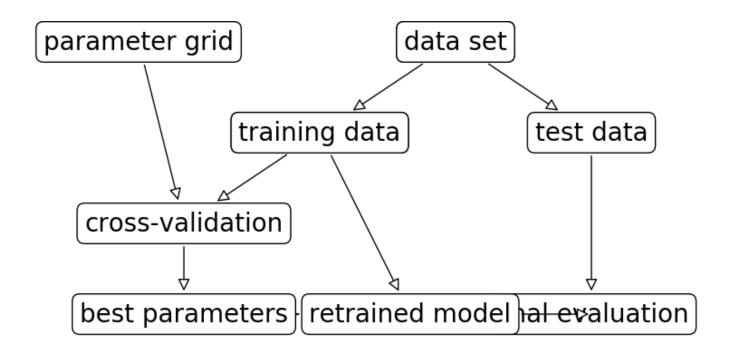
Best score on validation set: 0.96
Best parameters: {'C': 10, 'gamma': 0.001}
Test set score with best parameters: 0.92
```

#### Grid-search with cross-validation

- Same problem as before: the way that we split the data into training, validation, and test set may have a large influence on estimated performance
- We need to use cross-validation again, instead of a single split
- Expensive. Often, 3 or 5-fold CV is enough

```
for gamma in [0.001, 0.01, 0.1, 1, 10, 100]:
              for C in [0.001, 0.01, 0.1, 1, 10, 100]:
                  # train an SVC
                  svm = SVC(gamma=gamma, C=C)
                  # perform cross-validation
                  scores = cross val score(svm, X trainval, y trainval, cv=5)
                  # compute mean cross-validation accuracy
                  score = np.mean(scores)
                  # if we got a better score, store the score and parameters
                  if score > best score:
                      best score = score
                      best parameters = {'C': C, 'qamma': qamma}
Out[51]: SVC(C=100, cache size=200, class weight=None, coef0=0.0,
              decision function shape='ovr', degree=3, gamma=0.01, kernel='rbf',
              max iter=-1, probability=False, random state=None, shrinking=True,
              tol=0.001, verbose=False)
```

# Overall process



#### Grid search in scikit-learn

- Create a parameter grid as a dictionary
  - Keys are parameter names
  - Values are lists of hyperparameter values

- GridSearchCV: like a classifier that uses CV to automatically optimize its hyperparameters internally
  - Input: (untrained) model, parameter grid, CV procedure
  - Output: optimized model on given training data
  - Should only have access to training data

```
grid search = GridSearchCV(SVC(), param grid, cv=5)
          grid search.fit(X train, y train)
          GridSearchCV(cv=5, error score='raise-deprecating',
Out[561:
                       estimator=SVC(C=1.0, cache size=200, class weight=None, coef
          0 = 0.0,
                                      decision function shape='ovr', degree=3,
                                      gamma='auto deprecated', kernel='rbf', max ite
          r=-1,
                                      probability=False, random state=None, shrinkin
          g=True,
                                      tol=0.001, verbose=False),
                       iid='warn', n jobs=None,
                       param grid={'C': [0.001, 0.01, 0.1, 1, 10, 100],
                                    'gamma': [0.001, 0.01, 0.1, 1, 10, 100]},
                       pre dispatch='2*n jobs', refit=True, return train score=Fals
          e,
                       scoring=None, verbose=0)
```

The optimized test score and hyperparameters can easily be retrieved:

```
grid search.score(X test, y test)
Test set score: 0.97
grid search.best params
grid search best score
Best parameters: {'C': 100, 'gamma': 0.01}
Best cross-validation score: 0.97
grid search.best estimator
Best estimator:
SVC(C=100, cache size=200, class weight=None, coef0=0.0,
    decision function shape='ovr', degree=3, gamma=0.01, kernel='rbf',
   max iter=-1, probability=False, random state=None, shrinking=True,
    tol=0.001, verbose=False)
```

# Visualizing hyperparameter impact

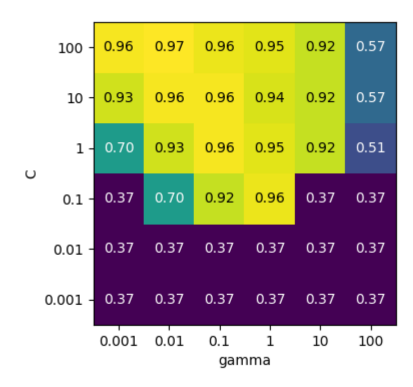
We can retrieve and visualize the cross-validation resulst to better understand the impact of hyperparameters

results = pd.DataFrame(grid\_search.cv\_results\_)

	mean_fit_time	std_fit_time	mean_score_time	std_score_time	•••	split4_test_score	mean_test_score	std_test_score	rank_test_score
0	1.06e-03	5.41e-04	4.11e-04	1.21e-04		0.38	0.37	0.01	22
1	5.87e-04	5.28e-05	2.51e-04	7.88e-06		0.38	0.37	0.01	22
2	6.44e-04	8.58e-05	3.07e-04	1.02e-04		0.38	0.37	0.01	22
3	8.14e-04	1.50e-04	3.85e-04	6.89e-05		0.38	0.37	0.01	22
4	9.86e-04	2.02e-04	4.03e-04	1.13e-04		0.38	0.37	0.01	22

 $5 \text{ rows} \times 15 \text{ columns}$ 

## Visualize as a heatmap



When hyperparameters depend on other parameters, we can use lists of dictionaries to define the hyperparameter space

#### **Nested cross-validation**

- Note that we are still using a single split to create the outer test set
- We can also use cross-validation here
- Nested cross-validation:
  - Outer loop: split data in training and test sets
  - Inner loop: run grid search, splitting the training data into train and validation sets
- Result is a just a list of scores
  - There will be multiple optimized models and hyperparameter settings (not returned)
- To apply on future data, we need to train GridSearchCV on all data again

Cross-validation scores: [0.967 1. 0.9 0.967 1. ]
Mean cross-validation score: 0.96666666666668

### Parallelizing cross-validation and grid-search

- On a practical note, it is easy to parallellize CV and grid search
- cross\_val\_score and GridSearchCV have a n\_jobs parameter defining the number of cores it can use.
  - set it to n\_jobs=-1 to use all available cores.

## Random Search

- Grid Search has a few downsides:
  - Optimizing many hyperparameters creates a combinatorial explosion
  - You have to predefine a grid, hence you may jump over optimal values
- Random Search:
  - Picks n\_iter random parameter values
  - Scales better, you control the number of iterations
  - Often works better in practice, too
    - o not all hyperparameters interact strongly
    - o you don't need to explore all combinations

- Executing random search in scikit-learn:
  - RandomizedSearchCV works like GridSearchCV
  - Has n\_iter parameter for the number of iterations
  - Search grid can use distributions instead of fixed lists

```
RandomizedSearchCV(cv='warn', error score='raise-deprecating',
Out[641:
                             estimator=SVC(C=1.0, cache size=200, class weight=Non
          e,
                                            coef0=0.0, decision function shape='ov
          r',
                                            degree=3, gamma='auto deprecated',
                                            kernel='rbf', max iter=-1, probability=F
          alse,
                                            random state=None, shrinking=True, tol=
          0.001,
                                            verbose=False),
                             iid='warn', n iter=20, n jobs=None,
                             param distributions={'C': <scipy.stats. distn infrastr</pre>
          ucture.rv frozen object at 0x11d861668>,
                                                   'gamma': <scipy.stats. distn infr
          astructure.rv frozen object at 0x11d98a860>},
                             pre dispatch='2*n jobs', random state=None, refit=Tru
          e,
                             return train score=False, scoring=None, verbose=0)
```

# **Summary**

- k-fold Cross-validation
  - Choose k depending on how much data you have
    - Larger k is slower, but allows more training data
    - o 10-fold, 5-fold, 5x2-fold most popular
  - Always use stratification for (imbalanced) classification
  - Train-test split and Shuffle-split: useful for large datasets
  - Use grouping when you want to generalize over groups
- Model selection
  - Don't aggregate over test scores: those have seen the test data
  - Use validation sets to choose algorithms/hyperparameters first
- Optimization
  - Grid Search: exhaustive but simple
  - Random Search: scales better
  - We'll see more advanced techniques later

# **Summary**

- Real-world data is often imbalanced
- False positives may be much worse than false negatives (or vise-versa)
- Binary classification
  - Select metrics that can distinguish different types of errors (precision, recall, f1-score, AUC,...)
  - Calibrate decision thresholds to the task at hand
  - Precision-Recall and ROC curves: choose the best threshold or take area under the curve
- Multiclass classification
  - Macro/Micro/weighted average of per-class scores (one-vs-all)
- Regression
  - (Root) mean squared/absolute error from 0..Inf
  - R2 easier to interpret
- All measures can be used in cross-validation or grid/random search
- Cost-sensitive classification: optimize for any cost matrix or cost function