

Creating train, test, and validation datasets

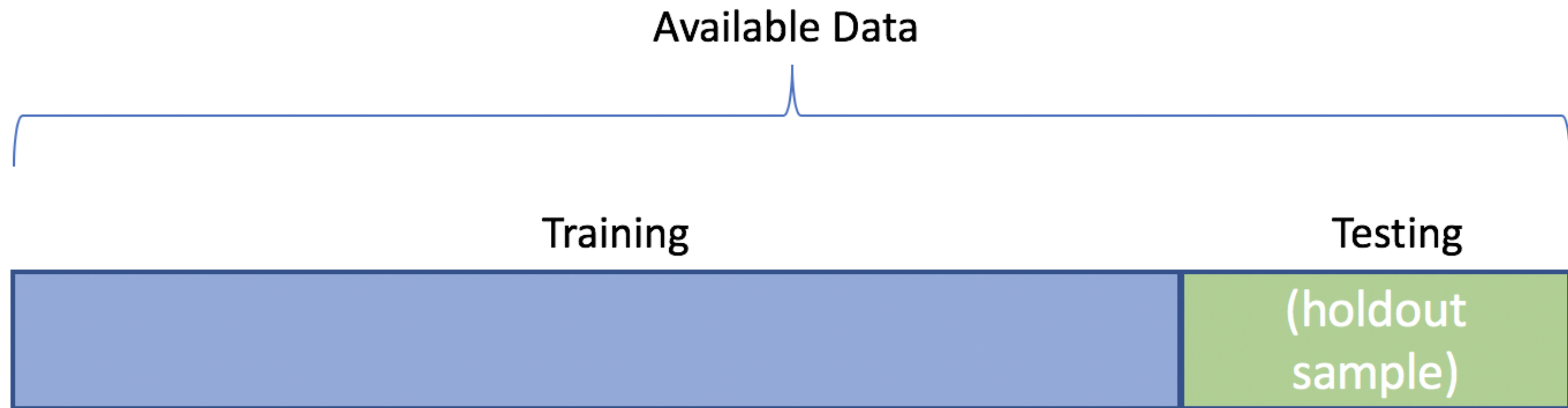
MODEL VALIDATION IN PYTHON



Kasey Jones
Data Scientist

Traditional train/test split

- Seen data (used for training)
- Unseen data (unavailable for training)



Dataset definitions and ratios

Dataset	Definition
Train	The sample of data used when fitting models
Test (holdout sample)	The sample of data used to assess model performance

Ratio Examples

- 80:20
- 90:10 (used when we have little data)
- 70:30 (used when model is computationally expensive)

The X and y datasets

```
import pandas as pd

tic_tac_toe = pd.read_csv("tic-tac-toe.csv")
X = pd.get_dummies(tic_tac_toe.iloc[:,0:9])
y = tic_tac_toe.iloc[:, 9]
```

Python courses covering dummy variables:

- [Supervised Learning](#)
- [Preprocessing for Machine Learning](#)

Creating holdout samples

```
X_train, X_test, y_train, y_test =\
    train_test_split(X, y, test_size=0.2, random_state=1111)
```

Parameters:

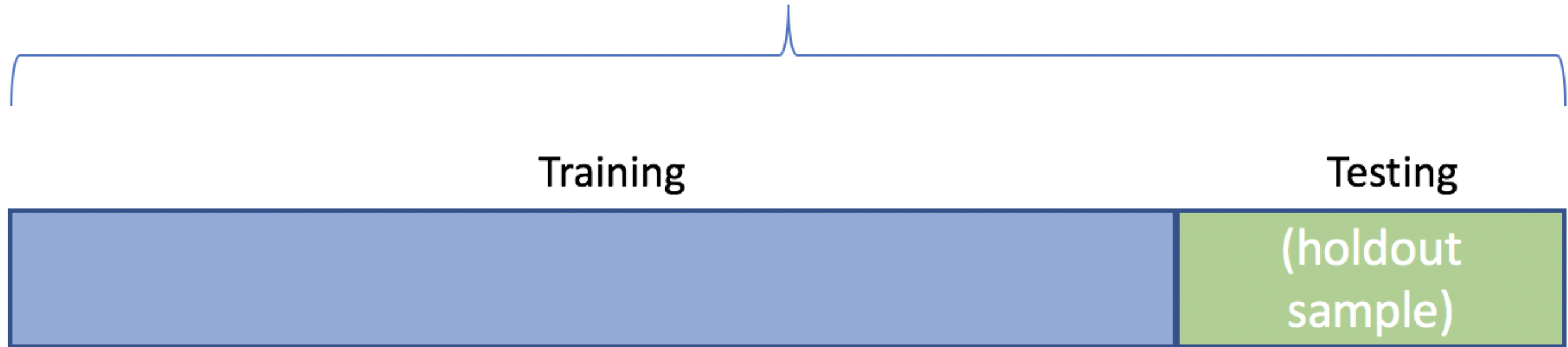
- test_size
- train_size
- random_state

Dataset for preliminary testing?

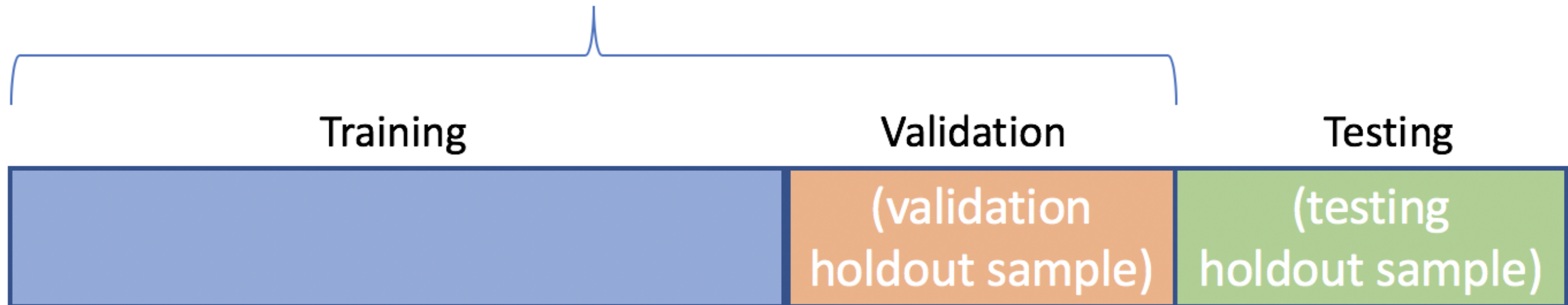
What do we do when testing different model parameters?

- 100 *versus* 1000 trees

Available Data



New Available Data



Train, validation, test continued

```
X_temp, X_test, y_temp, y_test =\
    train_test_split(X, y, test_size=0.2, random_state=1111)
```

```
X_train, X_val, y_train, y_val =\
    train_test_split(X_temp, y_temp, test_size=0.25, random_state=11111)
```


It's holdout time

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Accuracy metrics: regression models

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Regression models

12.2 points

15 gallons of gas

\$1,323,492

6 new puppies

4,320 people

Mean absolute error (MAE)

$$MAE = \frac{\sum_{i=1}^n |y_i - \hat{y}_i|}{n}$$

- Simplest and most intuitive metric
- Treats all points equally
- Not sensitive to outliers

Mean squared error (MSE)

$$MSE = \frac{\sum_{i=1}^n (y_i - \hat{y}_i)^2}{n}$$

- Most widely used regression metric
- Allows outlier errors to contribute more to the overall error
- Random family road trips could lead to large errors in predictions

MAE vs. MSE

- Accuracy metrics are always application specific
- MAE and MSE error terms are in different units and should not be compared

Mean absolute error

```
rfr = RandomForestRegressor(n_estimators=500, random_state=1111)
rfr.fit(X_train, y_train)
test_predictions = rfr.predict(X_test)
sum(abs(y_test - test_predictions))/len(test_predictions)
```

9.99

```
from sklearn.metrics import mean_absolute_error
mean_absolute_error(y_test, test_predictions)
```

9.99

Mean squared error

```
sum(abs(y_test - test_predictions)**2)/len(test_predictions)
```

```
141.4
```

```
from sklearn.metrics import mean_squared_error  
mean_squared_error(y_test, test_predictions)
```

```
141.4
```


Accuracy for a subset of data

```
chocolate_preds = rfr.predict(X_test[X_test[:, 1] == 1])  
mean_absolute_error(y_test[X_test[:, 1] == 1], chocolate_preds)
```

8.79

```
nonchocolate_preds = rfr.predict(X_test[X_test[:, 1] == 0])  
mean_absolute_error(y_test[X_test[:, 1] == 0], nonchocolate_preds)
```

10.99

Let's practice

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Classification metrics

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Classification metrics

- Precision
- Recall (also called sensitivity)
- Accuracy
- Specificity
- F1-Score, and its variations
- ...

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- Precision
- Recall (also called sensitivity)
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Confusion matrix

		Predicted Values	
		0	1
Actual Values	0	23 (TN)	7 (FP)
	1	8 (FN)	62 (TP)

True Positive: Predict/Actual are both 1

True Negative: Predict/Actual are both 0

False Positive: Predicted 1, actual 0

False Negative: Predicted 0, actual 1

```
from sklearn.metrics import confusion_matrix
cm = confusion_matrix(y_test, test_predictions)
print(cm)
```

```
array([[ 23,   7],
       [   8,  62]])
```

```
cm[<true_category_index>, <predicted_category_index>]
cm[1, 0]
```

```
8
```

Accuracy

Predicted Values			
		0	1
Actual Values	0	23	7
	1	8	62

$$\frac{23(\text{TN}) + 62(\text{TP})}{23 + 7 + 8 + 62} = .85$$

Precision

Predicted Values			
		0	1
Actual Values	0	23	7
	1	8	62

$$\frac{62(\text{TP})}{62(\text{TP}) + 7(\text{FP})} = .90$$

Recall

Predicted Values			
		0	1
Actual Values	0	23	7
	1	8	62

$$\frac{62(\text{TP})}{62(\text{TP})+8(\text{FN})} = .885$$

Accuracy, precision, recall

```
from sklearn.metrics import accuracy_score, precision_score, recall_score  
accuracy_score(y_test, test_predictions)
```

```
.85
```

```
precision_score(y_test, test_predictions)
```

```
.8986
```

```
recall_score(y_test, test_predictions)
```

```
.8857
```

Practice time

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The bias-variance tradeoff

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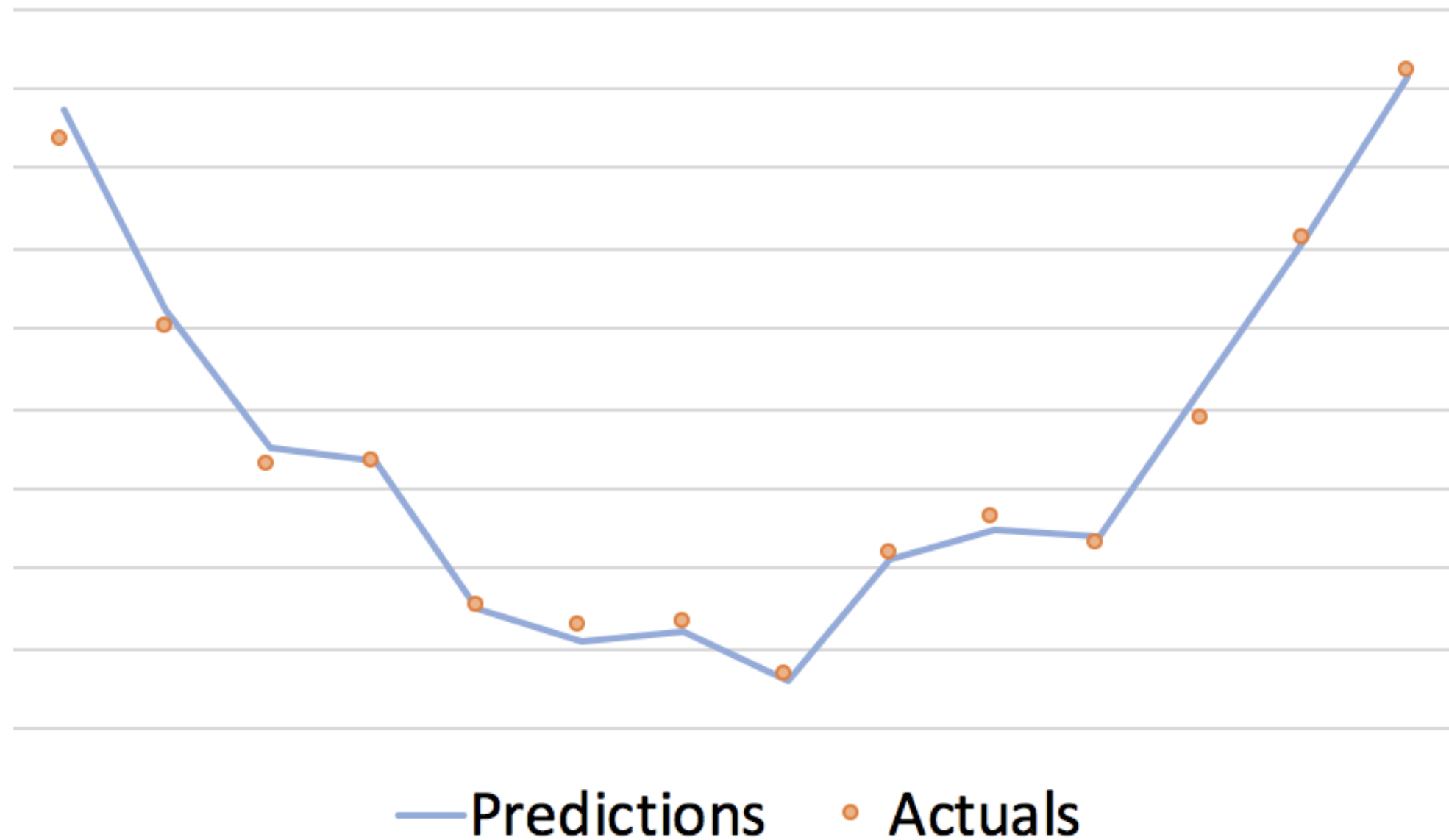


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Variance

- Variance: following the training data too closely
 - Fails to generalize to the test data
 - Low training error but high testing error
 - Occurs when models are overfit and have high complexity

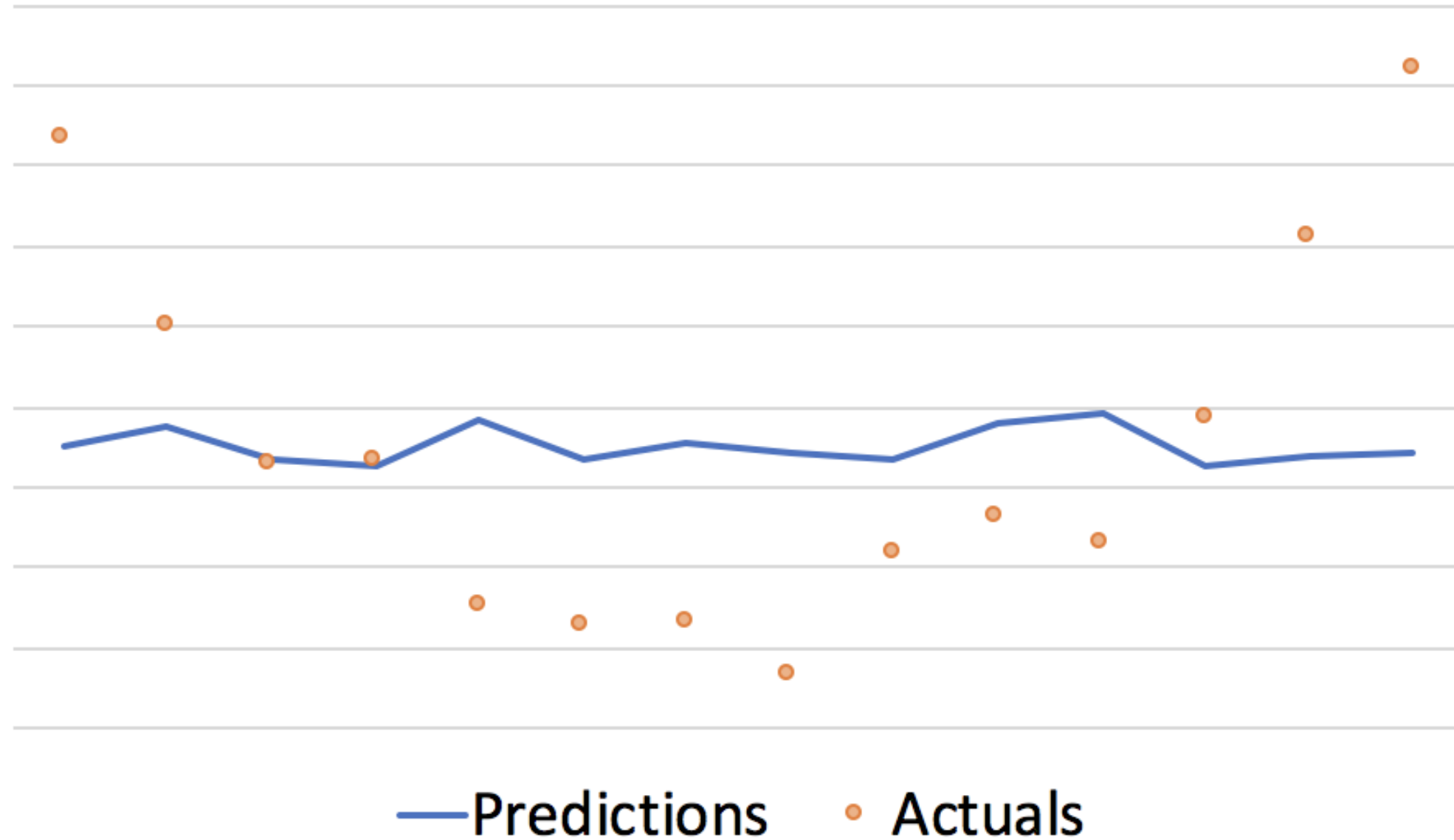
Overfitting models (high variance)



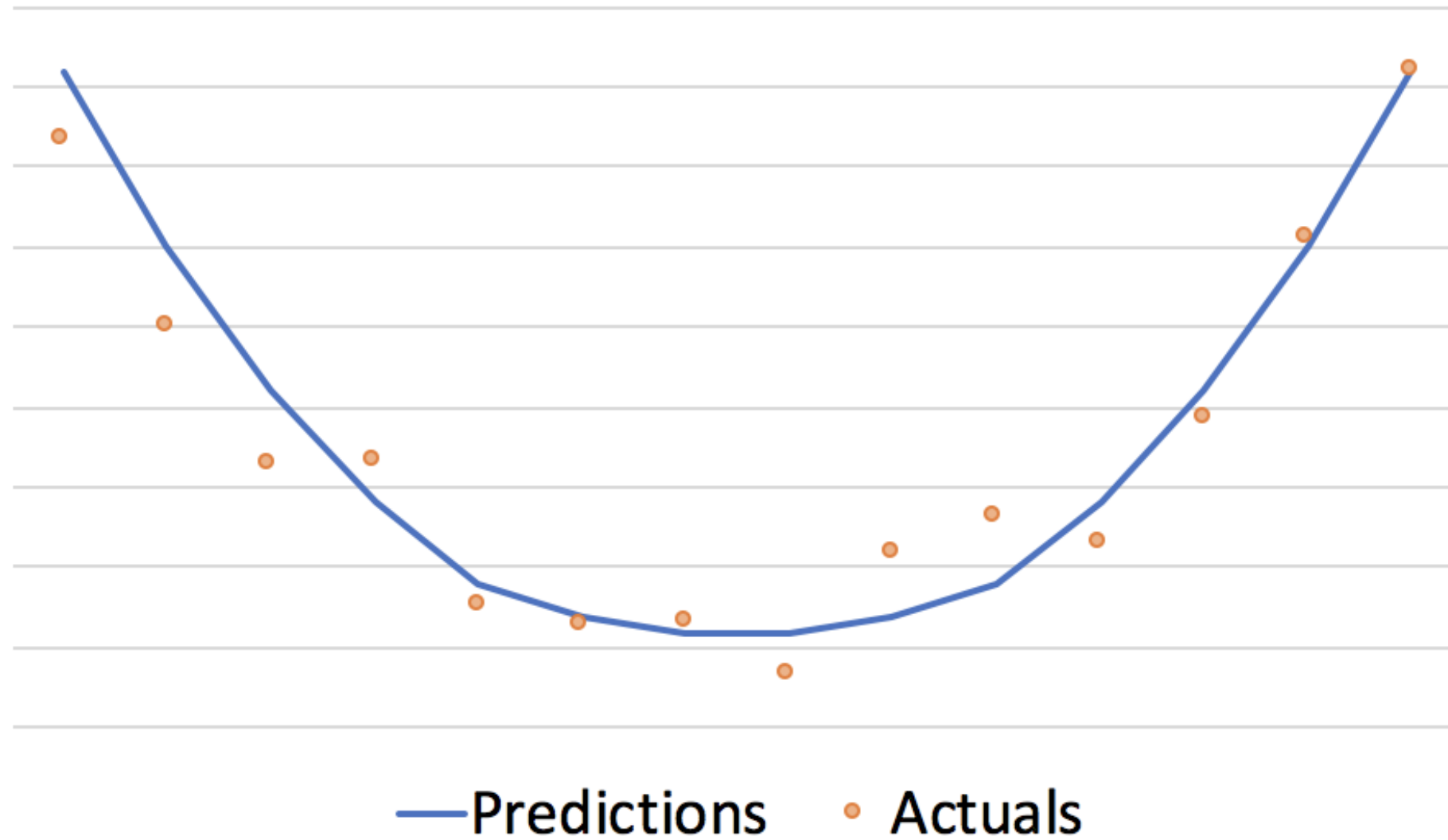
Bias

- Bias: failing to find the relationship between the data and the response
 - High training/testing error
 - Occurs when models are underfit

Underfitting models (high bias)



Optimal performance



- Bias-Variance Tradeoff

Parameters causing over/under fitting

```
rfc = RandomForestClassifier(n_estimators=100, max_depth=4)
rfc.fit(X_train, y_train)

print("Training: {0:.2f}".format(accuracy_score(y_train, train_predictions)))
```

Training: .84

```
print("Testing: {0:.2f}".format(accuracy_score(y_test, test_predictions)))
```

Testing: .77

```
rfc = RandomForestClassifier(n_estimators=100, max_depth=14)
rfc.fit(X_train, y_train)

print("Training: {0:.2f}".format(accuracy_score(y_train, train_predictions)))
```

```
Training: 1.0
```

```
print("Testing: {0:.2f}".format(accuracy_score(y_test, test_predictions)))
```

```
Testing: .83
```

```
rfc = RandomForestClassifier(n_estimators=100, max_depth=10)
rfc.fit(X_train, y_train)

print("Training: {0:.2f}".format(accuracy_score(y_train, train_predictions)))
```

```
Training: .89
```

```
print("Testing: {0:.2f}".format(accuracy_score(y_test, test_predictions)))
```

```
Testing: .86
```

**Remember, only you
can prevent
overfitting!**

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