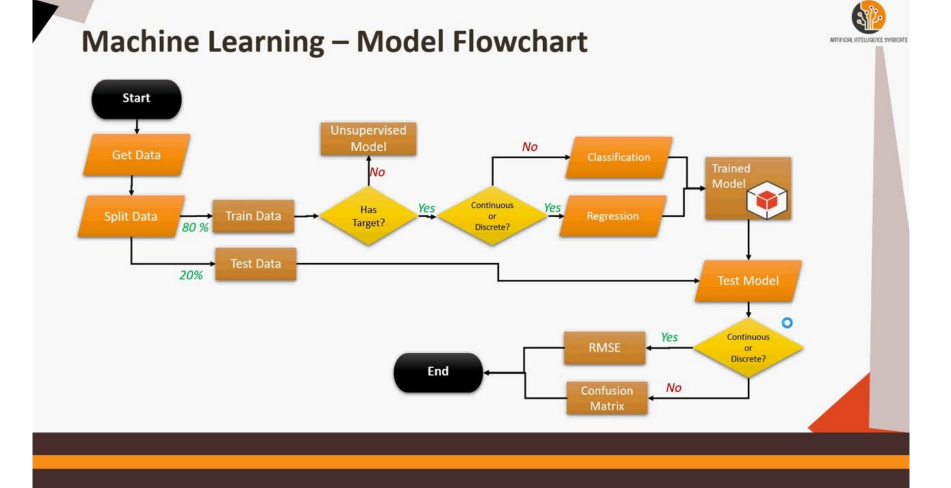
# The Machine Learning Pipeline (workflow)

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## The Machine Learning Pipeline

- The pipeline is composed of all of the computation and analysis required to get from the beginning to the end of a machine learning project
- Gather data
- Clean and preprocess data
- Exploratory data analysis
- Split data into training, validation and testing sets (holdout vs cross-validation)
- Develop/import model
- Train model
- Test model
- Hyperparameter tuning
- Report and visualize results



#### **Gather Data**

- We need data
- Good data is hard to find
  - No cure all method to easily finding and extracting data
  - Be persistent
- Data almost always needs to be cleaned and preprocessed
  - Think about, how can I make this data look like an Excel spreadsheet/Pandas dataframe
- Generally try to read in data with pd.read\_csv()

#### Clean and Preprocess data

- Read data and parse into a pandas dataframe or numpy array
- Take care of NAN values and outliers
  - Imputation or dropping
  - NaN values propagate through calculations
  - Outlier detection (kNN, support vector machine, covariance)
- Use sklearn's StandardScalar() to scale your data to mean of 0 with a standard deviation of 1
- Make necessary data transformations
  - For dates, pd.to\_datetime() should work, but I've had issues with it
  - You can also count number of days or hours from the beginning of the dataset and use this as the date/time variable
  - Units should be in the same system (kg, m, s; lb, ft, s)
  - Encode string-type variables
  - Transformations should be done before using StandardScalar()

#### **Encoding**

Convert
 categorical
 variables
 (strings) to
 numbers

# Fill Missing or NaN Values (sklearn)

#### sklearn.impute.lterativelmputer

Multivariate imputer that estimates each feature from all the others.

```
>>> # explicitly require this experimental feature
>>> from sklearn.experimental import enable_iterative_imputer # noqa
>>> # now you can import normally from sklearn.impute
>>> from sklearn.impute import IterativeImputer
```

#### 6.4.7. Estimators that handle NaN values

Some estimators are designed to handle NaN values without prepr type (cluster, regressor, classifier, transform):

- Estimators that allow NaN values for type regressor:
  - HistGradientBoostingRegressor
- Estimators that allow NaN values for type classifier:
  - HistGradientBoostingClassifier
- Estimators that allow NaN values for type transformer:
  - IterativeImputer
  - KNNImputer
  - MaxAbsScaler
  - MinMaxScaler
  - MissingIndicator
  - PowerTransformer
  - QuantileTransformer
  - RobustScaler
  - SimpleImputer
  - StandardScaler
  - VarianceThreshold

## Fill Missing or NaN Values (Pandas)

```
In [54]: dff.fillna(dff.mean())
Out[54]:
  0.271860 - 0.424972
                       0.567020
   0.276232 -1.087401 -0.673690
  0.113648 -1.478427
                       0.524988
3 -0.140857 0.577046 -1.715002
4 -0.140857 -0.401419 -1.157892
5 -1.344312 -0.401419 -0.293543
6 -0.109050 1.643563 -0.293543
 0.357021 -0.674600 -0.293543
8 -0.968914 -1.294524 0.413738
  0.276662 - 0.472035 - 0.013960
In [55]: dff.fillna(dff.mean()["B":"C"])
Out[55]:
  0.271860 - 0.424972 0.567020
  0.276232 -1.087401 -0.673690
  0.113648 -1.478427
                       0.524988
        NaN 0.577046 -1.715002
        NaN -0.401419 -1.157892
5 -1.344312 -0.401419 -0.293543
 -0.109050
            1.643563 -0.293543
  0.357021 - 0.674600 - 0.293543
8 -0.968914 -1.294524
                       0.413738
9 0.276662 -0.472035 -0.013960
```

#### Fill gaps forward or backward

Using the same filling arguments as reindexing, backward:

```
In [45]: df
Out[45]:
                           three
        one
                   two
        NaN -0.282863 -1.509059
             1.212112 -0.173215
   0.119209 - 1.044236 - 0.861849
f -2.104569 -0.494929
                       1.071804
        NaN -0.706771 -1.039575
In [46]: df.fillna(method="pad")
Out[46]:
                           three
                   two
        one
        NaN -0.282863 -1.509059
a
            1.212112 -0.173215
        NaN
   0.119209 -1.044236 -0.861849
f -2.104569 -0.494929
                       1.071804
h - 2.104569 - 0.706771 - 1.039575
```

#### Filling missing values: fillna

fillna() can "fill in" NA values with non-NA data in a

#### Replace NA with a scalar value

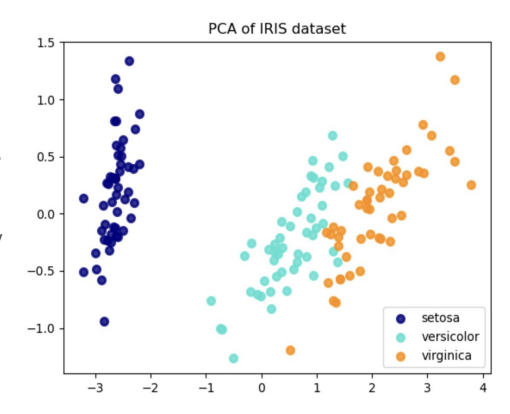
```
In [42]: df2
Out[42]:
                  two
                          three four
        NaN -0.282863 -1.509059
        NaN 1.212112 -0.173215
  0.119209 - 1.044236 - 0.861849
                                       True 20
f -2.104569 -0.494929 1.071804
        NaN -0.706771 -1.039575 bar
In [43]: df2.fillna(0)
Out[431:
                          three four
                  two
  0.000000 -0.282863 -1.509059
   0.000000 1.212112 -0.173215
   0.119209 -1.044236 -0.861849
f -2.104569 -0.494929 1.071804
h 0.000000 -0.706771 -1.039575
In [44]: df2["one"].fillna("missing")
Out[44]:
     missina
     missing
    0.119209
    -2.104569
```

## **Exploratory Data Analysis**

- Understand and communicate what is going on with your data at a general level
- Pandas profiler
  - Basic stats for whole dataframe and individual variables.
  - Scatter plots and correlation heatmap of all variables
  - If you can't install for whatever reason, you can use matplotlib to generate the same plots
- Feature Analysis
  - Unsupervised methods
    - Principal Component Analysis (PCA)
    - Outlier detection
  - Supervised methods
    - Linear Discriminant Analysis (LDA)
    - Random Forest
  - PCA, LDA, and factor analysis all look for linear combinations of variables that best explain the data

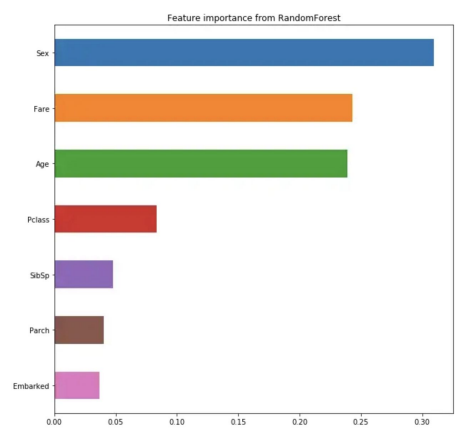
## Principal Component Analysis (PCA)

- "Linear dimensionality reduction using Singular Value Decomposition of the data to project it to a lower dimensional space"
- Separates data into new dimensions based on the combinations of attributes that account for most of the variance in the data (explanatory power)
- LDA is like PCA but it looks for attributes that explain the most variance between classes



```
from sklearn.ensemble import RandomForestClassifier
rf_clf = RandomForestClassifier(n_estimators = 500, max_depth=12)
rf_clf.fit(X_train, y_train)
rf y pred = rf clf.predict(X val)
```

## Random Forest Feature Engineering



#### **Titanic Dataset**

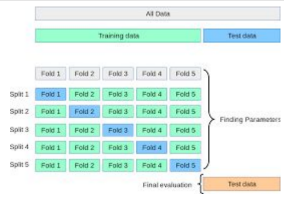
Name	Variable explanation	
pclass	Passenger Class (1 = 1st;2 = 2nd;3 = 3rd)	
Survived	Survival (0 = no, 1 = yes)	
Name	Passenger name	
Sex	Gender of passenger	
Age	Age of passenger	
Sibsp	(number of siblings/spouses aboard)	
Parch	(number of parents/children aboard)	
Ticket	Ticket number	
Fare	Passenger fare (£)	
Cabin	Cabin	
Embarked	Port of Embarkation (C = Cherbourg; Q = Queenstown; S = Southampton)	
Boat	Lifeboat	
Body	Body Identification Number	
Home.dest	Home/Destination	

 $\underline{\text{https://towardsdatascience.com/a-starter-pack-to-exploratory-data-analysis-with-python-pandas-seaborn-and-scikit-learn-a77889485baf}$ 

#### Training and testing splits

- We use training data to fit the model
- We use testing data (unseen during training) to test the model
  - Data leakage happens when the model somehow has access to testing samples during training
- Holdout
  - Only one train-test split
- Cross-Validation (CV)
  - Divide dataset into 6 parts
  - Train on five parts, test on 6th
  - Switch training/testing parts until all pairs are complete

```
>>> from sklearn.svm import SVC
>>> from sklearn.preprocessing import StandardScaler
>>> from sklearn.datasets import make_classification
>>> from sklearn.model selection import train test split
>>> from sklearn.pipeline import Pipeline
>>> X, y = make_classification(random_state=0)
>>> X train, X test, y train, y test = train test split(X, y,
                                                        random state=0)
>>> pipe = Pipeline([('scaler', StandardScaler()), ('svc', SVC())])
>>> # The pipeline can be used as any other estimator
>>> # and avoids leaking the test set into the train set
>>> pipe.fit(X train, y train)
Pipeline(steps=[('scaler', StandardScaler()), ('svc', SVC())])
>>> pipe.score(X test, y test)
0.88
```



#### Model Development

- Choose the appropriate model to solve the problem
- Whole lecture on this later
- Classification or Regression?
- Supervised, unsupervised or reinforcement learning?
  - https://scikit-learn.org/stable/supervised\_learning.html
  - https://scikit-learn.org/stable/unsupervised\_learning.html
- Read the Sklearn documentation for the model you pick to make sure you are using it in the right situation and passing the correct arguments to the model function
- Hyperparameter tuning (more on this later)
  - <a href="https://scikit-learn.org/stable/modules/grid-search.html">https://scikit-learn.org/stable/modules/grid-search.html</a>

# **Model Training**

We use the .fit() method for all training

- We use **fit()** method only on the training data. Why? Because we don't know what our testing data (unseen data) is, hence using the **fit()** method on the test data would not give us a good estimate of how our model is performing.
- We use **transform()** method on train data as well as test data as we need to perform transformation in both cases.

#### In Case Of Transformers

Transformers are for pre-processing the data before modelling.

- fit () This method goes through the training data, calculates the parameters (like mean ( $\mu$ ) and standard deviation ( $\sigma$ ) in StandardScaler class ) and saves them as internal objects.
- transform() The parameters generated using the fit() method are now used and applied to the training data to update them.
- **fit \_Transform()** This method may be more convenient and efficient for modelling and transforming the training data simultaneously.

https://medium.com/nerd-for-tech/difference-fit-transform-and-fit-transform-method-in-scikit-learn-b0a4efcab804

- lin\_reg.fit(X\_train\_diab, y\_train\_diab)
  y\_pred=lin\_reg.predict(X\_test\_diab)
  print ('R2: ',lin\_reg.score(X\_test\_diab,y\_test\_diab))
  print ('RMSE: ',mean squared error(y pred,y test\_diab,squared=False))
- R2: 0.4526066021617382 RMSE: 53.85325698491439
- knn\_class=skn.KNeighborsClassifier(n\_neighbors=i)
  knn\_class.fit(X\_train\_iris, y\_train\_iris)
  y pred=knn class.predict(X test iris)

## Model Testing/Scoring

- We use the .predict() method to generate predictions of our testing data
- If we use .predict(), we then have to generate a score by comparing the predicted data ("y\_pred") with the true data ("y\_test")
- If we use .score(), Sklearn does the prediction and score generation together
- The .score() method uses the arguments (x\_test,y\_test), but if you call one of the specific scoring functions on the right, the arguments are (y\_pred,y\_test)

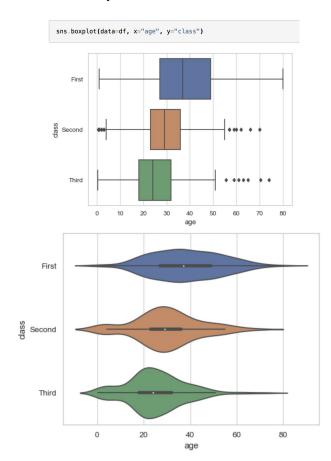
```
lin_reg.fit(X_train_diab, y_train_diab)
y_pred=lin_reg.predict(X_test_diab)
print ('R2: ',lin_reg.score(X_test_diab,y_test_diab))
print ('RMSE: ',mean_squared_error(y_pred,y_test_diab,squared=False))
```

Scoring	Function
Classification	
'accuracy'	metrics.accuracy_score
'balanced_accuracy'	<pre>metrics.balanced_accuracy_score</pre>
'top_k_accuracy'	<pre>metrics.top_k_accuracy_score</pre>
'average_precision'	<pre>metrics.average_precision_score</pre>
'neg_brier_score'	metrics.brier_score_loss
'f1'	metrics.fl_score
'f1_micro'	metrics.fl_score
'f1_macro'	metrics.fl_score
'f1_weighted'	metrics.fl_score
'f1_samples'	metrics.fl_score
'neg_log_loss'	metrics.log_loss
'precision' etc.	metrics.precision_score
'recall' etc.	metrics.recall_score
'jaccard' etc.	metrics.jaccard_score
'roc_auc'	metrics.roc_auc_score
Regression	
'explained_variance'	metrics.explained_variance_score
'max_error'	metrics.max_error
'neg_mean_absolute_error'	metrics.mean_absolute_error
'neg_mean_squared_error'	metrics.mean_squared_error
'neg_root_mean_squared_error'	metrics.mean_squared_error
'neg_mean_squared_log_error'	metrics.mean_squared_log_error
'neg_median_absolute_error'	metrics.median_absolute_error
ʻr2'	metrics.r2_score
'neg_mean_poisson_deviance'	metrics.mean_poisson_deviance
'neg_mean_gamma_deviance'	metrics.mean_gamma_deviance
'neg_mean_absolute_percentage_error'	metrics.mean_absolute_percentage_error

## Report and Visualize Results (Regression)

#### Regression

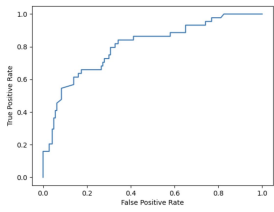
- Scatter plots with line of best fit
- For multiple linear regression, you can use 3D plots (x,y, and z coordinates), color, and scatter point size to add extra dimensions to the plot
- Correlation matrix/heatmap (already done with Pandas profiler)
- Box or violin plots of scores for different models/experiments/runs
  - Test different models under a range of changing variables/parameters (range of scores for each model)
  - Test the range of results for probabilistic models (different result after each run, like with neural nets)

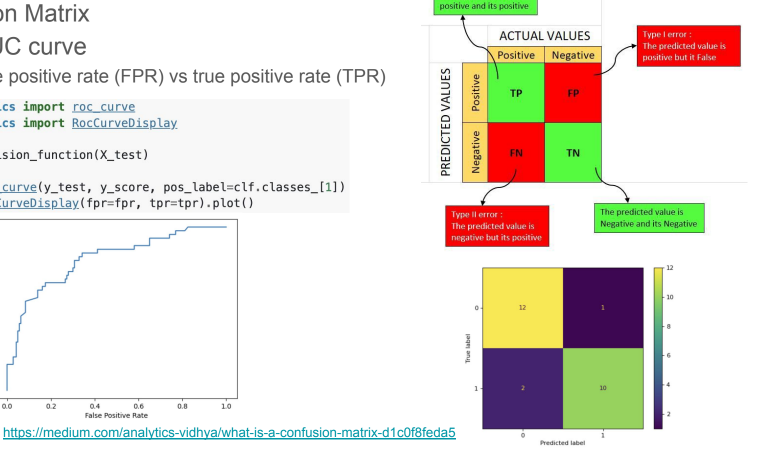


# Report and Visualize Results (Classification)

- Confusion Matrix
- **ROC-AUC** curve
  - False positive rate (FPR) vs true positive rate (TPR)

```
from sklearn.metrics import roc curve
from sklearn.metrics import RocCurveDisplay
y_score = clf.decision_function(X_test)
fpr, tpr, _ = roc curve(y_test, y_score, pos_label=clf.classes_[1])
roc display = RocCurveDisplay(fpr=fpr, tpr=tpr).plot()
```





The predicted value is

Example Walkthrough of Iris dataset confusion matrix and best nearest neighbors searcher

```
import sklearn.metrics as skm
best i=0
equal list=[]
for i in range(1,100):
    knn class=skn.KNeighborsClassifier(n neighbors=i)
    knn class.fit(X train iris, y train iris)
    y pred=knn class.predict(X test iris)
    if i==1:
        best mat=skm.confusion matrix(y pred,y test iris)
        best_score=(best_mat[0,0]+best_mat[1,1]+best_mat[2,2])/30
        best i=1
    else:
        new mat=skm.confusion matrix(y pred,y test iris)
        if (new mat[0,0]+new mat[1,1]+new mat[2,2])/30>best score:
            best mat=skm.confusion matrix(y pred,y test iris)
            best score=(new mat[0,0]+new mat[1,1]+new mat[2,2])/30
            best i=i
        elif new mat[0,0]+new mat[1,1]+new mat[2,2]==best score:
            equal list.append(i)
print ('Classification Performance: ',best score,' NN = ',str(best i))
```

Classification Performance: 1.0 NN = 1

## Help for Hw1 Q2

```
In [16]: from sklearn.datasets import load iris, load diabetes
         iris = load iris()
         diab = load diabetes()
         from sklearn.model selection import train test split
         X train iris, X test iris, y train iris, y test iris = train test split(iris.data, iris.target, test size=0.2)
         X train diab, X test diab, y train diab, y test diab = train test split(diab.data, diab.target, test size=0.2)
In [17]: import sklearn
         import sklearn.linear model
         log reg=sklearn.linear model.LogisticRegression(max iter=200)
         log reg.fit(X train iris, y train iris)
         y pred=log reg.predict(X test iris)
         print ('accuracy: ',log reg.score(X test iris,y test iris))
         accuracy: 0.966666666666667
In [18]: import sklearn.metrics as skm
         mat=skm.confusion matrix(y pred,y test iris)
         disp=skm.ConfusionMatrixDisplay(mat)
         disp.plot()
Out[18]: <sklearn.metrics. plot.confusion matrix.ConfusionMatrixDisplay at 0x7fe43913adf0>
            0 -
```