

# Transformation pipelines

A transformation pipeline is a sequence (sometimes called a *pipeline*) of independent transformations  $T_1, T_2, \dots, T_t$

$$\tilde{\mathbf{x}}_{(1)} = T_1(\mathbf{x})$$

$$\tilde{\mathbf{x}}_{(2)} = T_2(\tilde{\mathbf{x}}_{(1)})$$

$$\vdots$$

$$\tilde{\mathbf{x}}_{(l+1)} = T_{(l+1)}(\tilde{\mathbf{x}}_{(l)})$$

The final transformed  $\tilde{\mathbf{x}}$  can be implemented as a function  $T$  that is the composition of each transformation function

$$\tilde{\mathbf{x}} = T(\mathbf{x}) = T_t( T_{t-1}(\dots T_1(\mathbf{x}) \dots ) )$$

## Pipelines in sklearn

sklearn has created a generic architecture called Pipeline to simplify this for you.

This module is **not meant to be a tutorial**, only to highlight a very convenient tool in sklearn.

[Here \(https://scikit-learn.org/stable/modules/compose.html\)](https://scikit-learn.org/stable/modules/compose.html) is a link to sklearn's manual.

Let's illustrate with a two-step sequence of transformations on numeric features

- Missing data imputation
  - If feature  $j$  is missing for example  $i$  ( $\mathbf{x}_j^{(i)}$  not defined)
  - Replace it with the median (over all  $m$  examples) of  $\mathbf{x}_j$
- Add a second order polynomial feature to each example

Here is some illustrative code

```
# Illustrative code: this cell is NOT executable from sklearn
import pipeline from sklearn.impute import
SimpleImputer from sklearn.preprocessing import PolynomialFeatures # Create transformers
polynomial_features = PolynomialFeatures(2) imputed_features = SimpleImputer(strategy='median') # Put
the transformers into a Pipeline num_pipe = pipeline.Pipeline([ ("imputer", imputed_features),
("polynomial_features", polynomial_features), ])
```

Each individual transformation has a `fit` and `transform` method, just like a model (e.g., `LinearRegression`)

- `fit` on the transformation `SimpleImputer(strategy='median')` computes the median value of each feature
  - Computes  $\Theta_{\text{transform}}$ , the parameters of this transformation
- `transform` applies the transformation, after it has been fit
  - creates  $\tilde{\mathbf{x}}$  from  $\mathbf{x}$

The Pipeline `num_pipe` defines a *sequence* of transformations

- Each element of the Pipeline is a tuple
  - name of the transformation (for your convenience in referencing it)
  - the transformer

The Pipeline *also* has methods `fit` and `transform`

- `fit` on a Pipeline applies `fit` to each component transformation in turn
- `transform` on a Pipeline applies `transform` to each component transformation in turn

The Pipeline is thus a composition of the individual transformations

$$\tilde{\mathbf{x}} = T(\mathbf{x}) = T_t( T_{t-1}(\dots T_1(\mathbf{x}) \dots) )$$

## **Model objects as part of a pipeline**

Observe that transformers and models both respond to the `fit` and `transform` methods.

This means that you can include a model as the final element of a Pipeline !



```
# Illustrative code: this cell is NOT executable from sklearn import datasets, linear_model
linear_regression = linear_model.LinearRegression() full_pipe = pipeline.Pipeline([("imputer", imputed_features),
("polynomial_features", polynomial_features), ("model", linear_regression)])
```

You can fit the transformations *and* the model in one step

- `full_pipe.fit(X, y)`

In addition to being a concise and convenient notation, pipelines with models as final elements

- Can be used in Cross Validation to avoid "cheating"
- The cross validation code passes in *only* the folds that are the training examples

## Example: Pipeline with model as final element

Let's apply a Pipeline to solve an example encountered in a previous module

- Linear Regression model
- With a second order feature added

Let's get the data again:

```
In [4]: # I will give you the data via a function (so I can easily alter the data in sub  
sequent examples)  
v1, a1 = 1, .005  
lin = recipe_helper.Recipe_Helper(v = v1, a = a1)  
  
X_lin, y_lin = lin.gen_data(num=50)  
  
v2, a2 = v1, a1*2  
curv = recipe_helper.Recipe_Helper(v = v2, a = a2)  
X_curve, y_curve = curv.gen_data(num=50)  
  
X_train, X_test, y_train, y_test = rh.split(X_curve,y_curve)
```

And here's the model, implemented via a Pipeline in `sklearn`

```
In [5]: from sklearn.preprocessing import PolynomialFeatures
import sklearn.preprocessing as pre_proc

# Create the Pipeline
from sklearn.pipeline import make_pipeline
poly_model = make_pipeline(pre_proc.PolynomialFeatures(2),
                           linear_model.LinearRegression())

# Fit the model, running the data transformation first
_= poly_model.fit(X_train.reshape(-1,1), y_train)
```

The `PolynomialFeatures(2)` transformer

- Replace each  $\mathbf{x}$
- With 3 features:  $\mathbf{x}^0, \mathbf{x}^1, \mathbf{x}^2$

That is: it creates an "intercept" dummy ( $\mathbf{x}^0 == 1$ ) plus first and second order features.

A pipeline successively applies transformations, with the result of transformation  $(i - 1)$  fed as input to transformation  $i$ .

Let's look "inside" the pipeline at the stages, and apply them manually.



```
In [6]: # Examine the "stages" of the pipeline
print("Input shape: {shp}".format(shp=X_test.reshape(-1,1).shape) )

# First stage: Create First and Second Order polynomial features
(label_0, model_0) = poly_model.steps[0]
transf_0 = model_0.transform(X_test.reshape(-1,1))
print("{lab:s} returns shape: {shp}".format(lab=label_0, shp=transf_0.shape) )

# Second stage: Linear Regression
(label_1, model_1) = poly_model.steps[1]
transf_1 = model_1.predict( transf_0 )
print("{lab:s} returns shape: {shp}".format(lab=label_1, shp=transf_1.shape) )
```

```
Input shape: (10, 1)
polynomialfeatures returns shape: (10, 3)
linearregression returns shape: (10, 1)
```

```
In [7]: # Prediction based on test set
        y_pred = poly_model.predict(X_test.reshape(-1,1))

        # In and out of sample scores
        print("Score (train): ", poly_model.score(X_train.reshape(-1,1), y_train))
        print("Score (test): ", poly_model.score(X_test.reshape(-1,1), y_test))

Score (train): 1.0
Score (test): 1.0
```

```
In [8]: # Plot the results
        # Create a figure
        fig, axs = plt.subplots(1,2, figsize=(12,6))

        _ = axs[0].scatter(X_train,y_train)

        xfit = np.linspace( X_train[:,0].min(), X_train[:,0].max()).reshape(-1,1)
        yfit = poly_model.predict(xfit)
        _ = axs[0].plot(xfit, yfit);
        _ = axs[0].set_title("Train")

        _ = axs[1].scatter(X_test, y_test, color="blue")
        _ = axs[1].scatter(X_test, y_pred, color="red")

        _ = axs[1].plot(xfit, yfit)
        _ = axs[1].set_title("Test")

        print("R-squared score (test): {:.2f}".format(r2_score(y_test, y_pred)) )

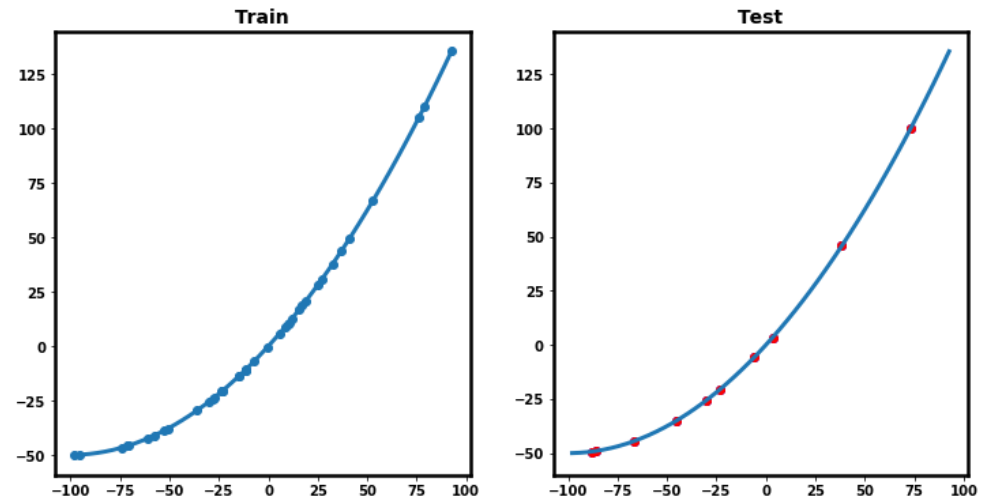
        # Hide the figure for now, will show it in the next slide
        plt.close(fig)
```

R-squared score (test): 1.00

In [9]:

```
fig
```

Out[9]:



Here's ([external/PythonDataScienceHandbook/notebooks/05.04-Feature-Engineering.ipynb#Feature-Pipelines](https://external/PythonDataScienceHandbook/notebooks/05.04-Feature-Engineering.ipynb#Feature-Pipelines)) a slightly longer pipeline from VanderPlas.

- `Imputer` to deal with missing values
- `PolynomialFeatures(degree=2)`
- `LinearRegression()`

## Nested Pipelines

A Pipeline responds to the same methods ( `fit` , `transform` ) as its elements.

This means that a Pipeline can also be used as a *nested* element of an outer Pipeline.

This is very convenient: we will illustrate this in our detailed example for Classification.

## Advanced Pipelines

An issue with Pipelines is that each transformation is applied to *all* the features in an example.

But some transformations need to be applied to *selected* features

- Can't apply numeric transformations to non-numeric data, and vice-versa
- May want to apply a particular transformation to only a subset of features

sklearn facilitates this by

- Allowing you to create "filters" that restrict features to a subset
- Applying one pipeline per subset
- Creating a union of transformed features at the end



## FeatureUnion

sklearn allows you to create complex Pipelines with the FeatureUnion Pipeline

- "glue together" the features of separate Pipelines. For example
  - Create one pipeline to be applied only to numeric features
  - Create one pipeline to be applied only to non-numeric features

Here is some illustrative code:

```
num_pipeline = Pipeline([("select_numeric", DataFrameSelector(["Age", "SibSp", "Parch", "Fare"])),
                          ("imputer", SimpleImputer(strategy="median")), ])
cat_pipeline = Pipeline([("select_cat", DataFrameSelector(["Pclass", "Sex", "Embarked"])),
                          ("imputer", MostFrequentImputer()),
                          ("cat_encoder", OneHotEncoder(sparse=False)), ])
from sklearn.pipeline import FeatureUnion
preprocess_pipeline = FeatureUnion(transformer_list=[("num_pipeline", num_pipeline), ("cat_pipeline", cat_pipeline), ])
```

The first element in each pipeline ( `DataFrameSelector` )

- restricts each example to a selected subset of features

So the numeric and categorical pipelines above transform disjoint groups of features.

The `FeatureUnion` concatenates (along the horizontal axis) the result of the separate transformations.

# ColumnTransformer

There is an *experimental* Pipeline object in `sklearn` called Column Transformers, that is a simplification of the `FeatureUnion` paradigm

- No need to filter the features with `DataFrameSelector`
  - just provide a list of feature names
- No need to use `'FeatureUnion'`

This object *only works* for collections of examples in which we can access features *by name*

- Panda DataFrames !

Here is some illustrative code

```
cat_features = ["Sex", "Pclass"] cat_transformers= Pipeline(steps=[ ('imputer', MostFrequentImputer()),  
('sex_encoder', SexToInt()) ]) cat_pipeline = ColumnTransformer( transformers=[ ("categorical",  
cat_transformers, cat_features) ] )
```

# Creating your own transformations

In addition to supplying a number of built-in transformers, `sklearn` let's use build your own.

- Create an object that is subclass of the types used by built-in transformer
- Provide your own implementation of `fit` and `transform`

Here's a transformation for imputation of missing values using the most frequent value as the substitute.

- Similar in function to the built-in `SimpleImputer(strategy="most_frequent")`

```
In [10]: # Inspired from stackoverflow.com/questions/25239958
from sklearn.base import BaseEstimator, TransformerMixin

class MostFrequentImputer(BaseEstimator, TransformerMixin):
    def fit(self, X, y=None):
        self.most_frequent_ = pd.Series([X[c].value_counts().index[0] for c in
X],
                                     index=X.columns)

        return self

    def transform(self, X, y=None):
        return X.fillna(self.most_frequent_)
```

## Example

We will use Pipelines in our Classification task.



```
In [11]: print("Done")
```

Done

