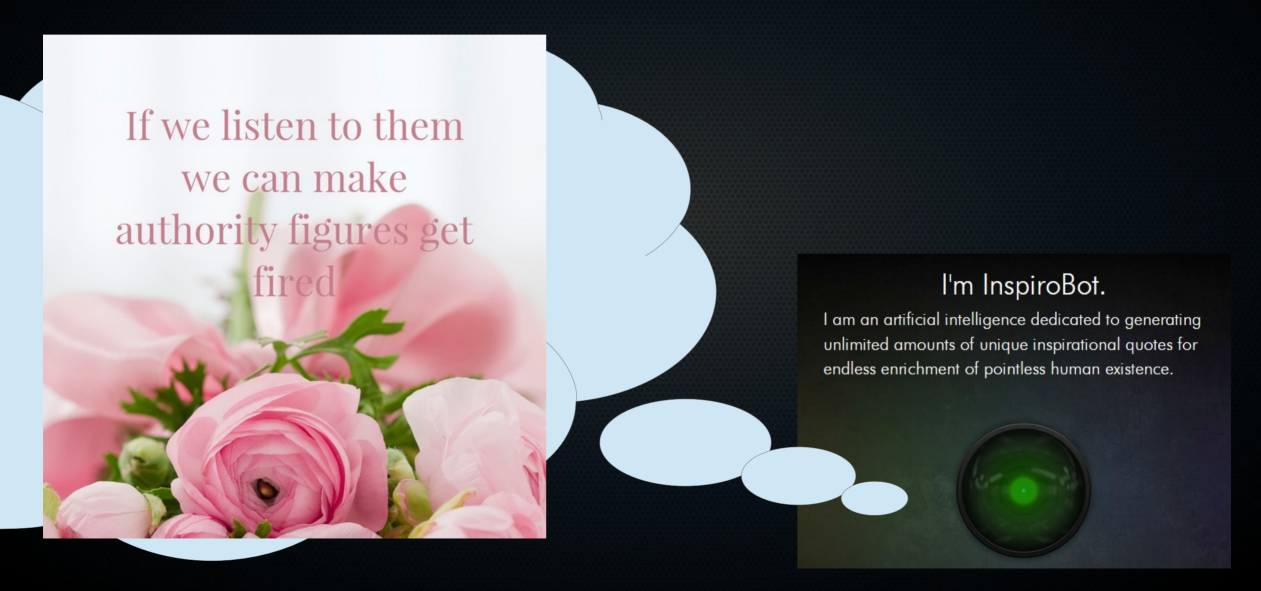
Daily Inspiration



Today

- Recap dim. reduction
- Introduction this week
- Scikit-learn
- Keras

Recap dimensionality reduction

- High dimensional data prevents lots of difficulties: sparseness, proneness to overfitting, etc.
- We can reduce the number of dimensions while keeping most of the information we're interested in.
- Non-linear methods: UMAP and consorts who try to best capture high-dimensional distance properties in low dimensions
- Linear methods: PCA and consorts, where PCA defines linear combinations of old dimensions (i.e. rotations of the data) where the first new PC captures most of the variance, the second the second most, etc.

Introduction this week

So far:

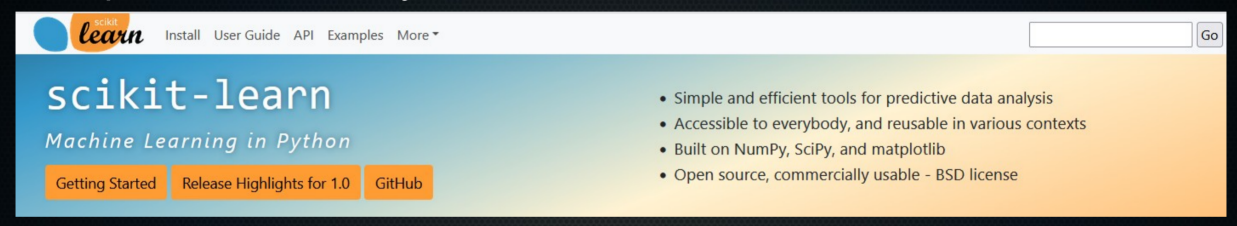
- Supervised learning implemented yourself (linear regression, logistic regression, simple/dense neural network)
- Unsupervised learning implemented yourself (K-means, hierarchical clustering, PCA)
- Intimate knowledge of the fundamentals, of what happens under the hood and why.

Introduction this week

Now:

- Switch to current applied workflows
 - Scikit-learn for supervised and unsupervised ML
 - Keras for easy creation and training of (convolutional) neural networks
- Do a project with these high-level libraries, using the knowledge you've gained

Open-source library



Implements a wealth of ML algorithms

Implements a wealth of ML algorithms and preprocessing/plotting functions:

1.1. Linear Models

1.1.1. Ordinary Least Squares

1.1.2. Ridge regression and

classification

1.1.3. Lasso

1.1.4. Multi-task Lasso

1.1.5. Elastic-Net

1.1.6. Multi-task Elastic-Net

1.1.7. Least Angle Regression

1.1.8. LARS Lasso

1.1.9. Orthogonal Matching Pursuit (OMP)

1.1.10. Bayesian Regression

1.1.11. Logistic regression

1.1.12. Generalized Linear

Regression

1.1.13. Stochastic Gradient Descent

- SGD

1.1.14. Perceptron

1.1.15. Passive Aggressive

Algorithms

1.1.16. Robustness regression:

outliers and modeling errors

1.1.17. Quantile Regression

1.1.18. Polynomial regression: extending linear models with basis functions

1.4. Support Vector Machines

1.4.1. Classification

1.4.2. Regression

1.4.3. Density estimation, novelty

detection

1.4.4. Complexity

1.4.5. Tips on Practical Use

1.4.6. Kernel functions

1.4.7. Mathematical formulation

1.4.8. Implementation details

1.10. Decision Trees

1.10.1. Classification

1.10.2. Regression

1.10.3. Multi-output problems

1.10.4. Complexity

1.10.5. Tips on practical use

1.10.6. Tree algorithms: ID3, C4.5,

C5.0 and CART

1.10.7. Mathematical formulation

1.10.8. Minimal Cost-Complexity Pruning

2.3. Clustering

2.3.1. Overview of clustering methods

2.3.2. K-means

2.3.3. Affinity Propagation

2.3.4. Mean Shift

2.3.5. Spectral clustering

2.3.6. Hierarchical clustering

2.3.7. DBSCAN

2.3.8. OPTICS

2.3.9. BIRCH

2.3.10. Clustering performance evaluation

6.3. Preprocessing data

6.3.1. Standardization, or mean

removal and variance scaling

6.3.2. Non-linear transformation

6.3.3. Normalization

6.3.4. Encoding categorical features

6.3.5. Discretization

6.3.6. Imputation of missing values

6.3.7. Generating polynomial

features

6.3.8. Custom transformers

2.5. Decomposing signals in components (matrix factorization problems)

2.5.1. Principal component analysis (PCA)

2.5.2. Kernel Principal Component

Analysis (kPCA)

2.5.3. Truncated singular value decomposition and latent semantic analysis

2.5.4. Dictionary Learning

2.5.5. Factor Analysis

2.5.6. Independent component analysis (ICA)

2.5.7. Non-negative matrix

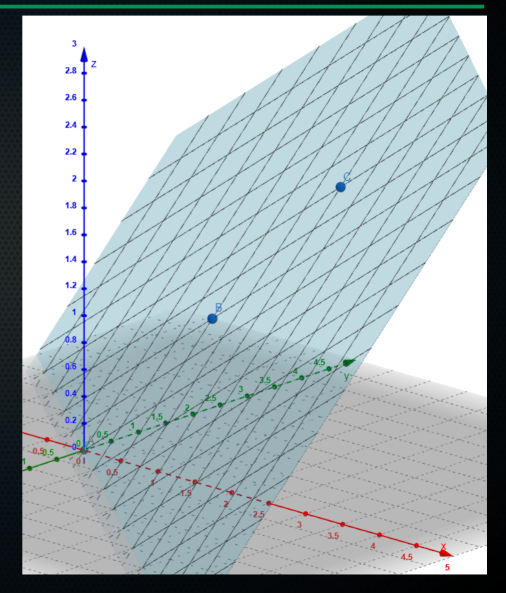
factorization (NMF or NNMF)

2.5.8. Latent Dirichlet Allocation

(LDA)

Similar API for all models:

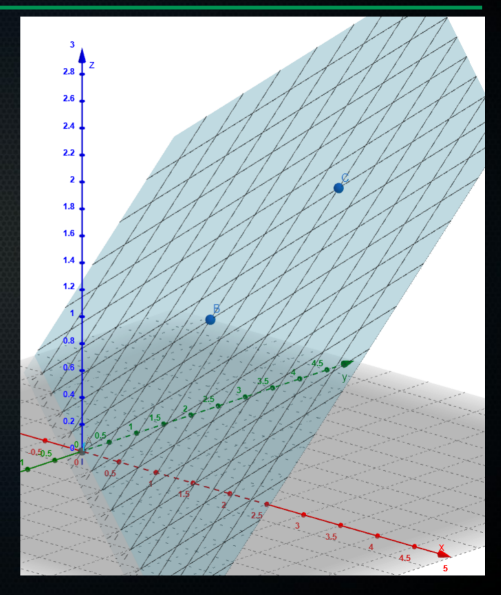
```
>>> from sklearn import linear_model
>>> reg = linear_model.LinearRegression()
>>> reg.fit([[0, 0], [1, 1], [2, 2]], [0, 1, 2])
LinearRegression()
>>> reg.coef_
array([0.5, 0.5])
```



Similar API for all models:

```
>>> from sklearn import linear_model
>>> reg = linear_model.LinearRegression()
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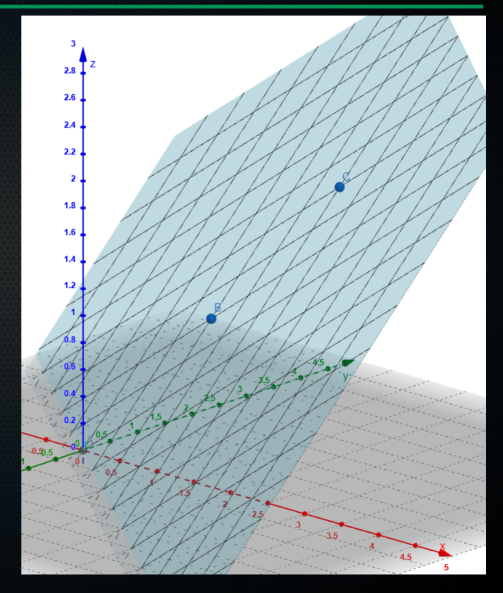
Initiate the model



Similar API for all models:

 Fit the model to data (multivariate linear regression with z predicted from x and y)

$$h_{\theta}(x, y) = [x \quad y] \cdot \theta^{T}$$

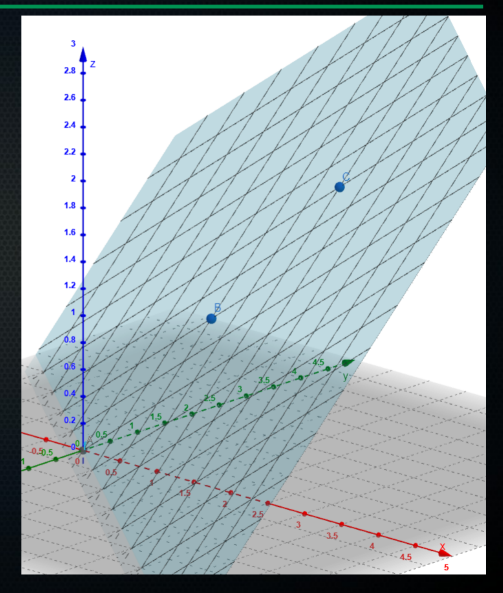


Similar API for all models:

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>>> reg.fit([[0, 0], [1, 1], [2, 2]], [0, 1, 2])
LinearRegression()
>>> reg.coef_
array([0.5, 0.5])
```

Look at fitted coefficients

$$h_{\theta}(x,y) = \begin{bmatrix} x & y \end{bmatrix} \cdot \theta^{T} \theta = \begin{bmatrix} 0.5 \\ 0.5 \end{bmatrix}$$



Similar API for all models:

```
>>> from sklearn import linear_model
>>> reg = linear_model.LinearRegression()
>>> reg.fit([[0, 0], [1, 1], [2, 2]], [0, 1, 2])
LinearRegression()
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array([0.5, 0.5])

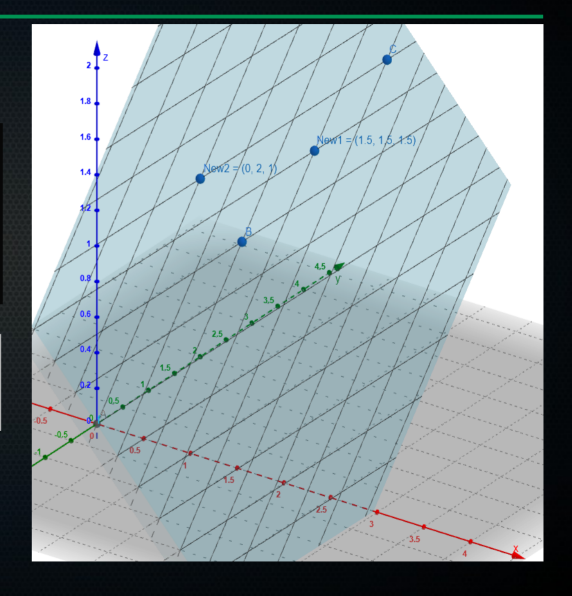
reg.predict([[1.5, 1.5], [0,2]])

array([1.5, 1. ])

θ = 0.5
0.5
```

Predict new data

$$h_{\theta}(x,y) = \begin{bmatrix} 1.5 & 1.5 \\ 0 & 2 \end{bmatrix} \cdot \theta^{T} = \begin{bmatrix} 1.5 \\ 1 \end{bmatrix}$$



Scikit-learn: split data

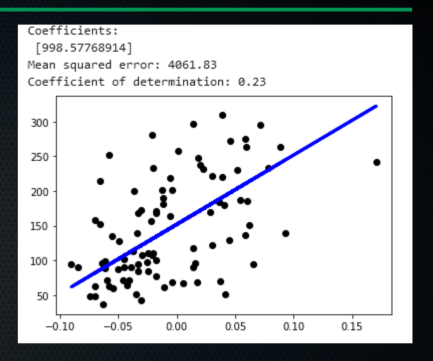
```
import matplotlib.pyplot as plt
import numpy as np
from sklearn import datasets, linear_model
from sklearn.metrics import mean_squared_error, r2_score
from sklearn.model_selection import train_test_split

# Load the diabetes dataset
diabetes_X, diabetes_y = datasets.load_diabetes(return_X_y=True)
```

Data Set Chara	octeristics:				
Number of Instances:	442				
Number of Attributes:	First 10 columns are numeric predictive values				
Target:	Column 11 is a quantitative measure of disease progression one year after baseline				
Attribute Information:	 age age in years sex bmi body mass index bp average blood pressure s1 tc, total serum cholesterol s2 ldl, low-density lipoproteins s3 hdl, high-density lipoproteins s4 tch, total cholesterol / HDL s5 ltg, possibly log of serum triglycerides level s6 glu, blood sugar level 				

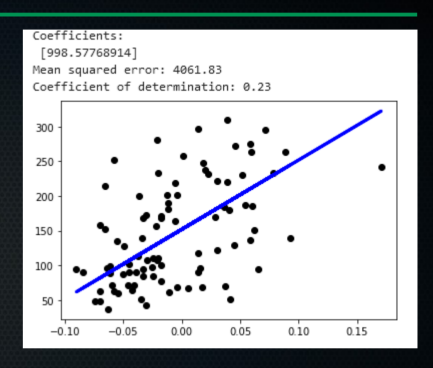
Scikit-learn: split data

```
# Use only one feature
diabetes X = diabetes X[:, np.newaxis, 2]
#split data:
diabetes X train, diabetes X test, diabetes y train, diabetes y test = train test split(
    diabetes_X, diabetes_y, test_size = 0.2, random_state = 42)
# Create linear regression object
regr = linear model.LinearRegression()
# Train the model using the training sets
regr.fit(diabetes X train, diabetes y train)
# Make predictions using the testing set
diabetes y_pred = regr.predict(diabetes X_test)
#MSE and R2 on test set
MSE = mean_squared_error(diabetes_y_test, diabetes_y_pred)
RSQ = r2 score(diabetes y test, diabetes y pred)
```



Scikit-learn: calculate metrics

```
import matplotlib.pyplot as plt
import numpy as np
from sklearn import datasets, linear model
from sklearn.metrics import mean squared error, r2 score
from sklearn.model selection import train test split
# Load the diabetes dataset
diabetes X, diabetes y = datasets.load diabetes(return X y=True)
# Use only one feature
diabetes X = diabetes X[:, np.newaxis, 2]
#split data:
diabetes X train, diabetes X test, diabetes y train, diabetes y test = train test split(
    diabetes X, diabetes y, test size = 0.2, random state = 42)
# Create linear regression object
regr = linear model.LinearRegression()
# Train the model using the training sets
regr.fit(diabetes X train, diabetes y train)
# Make predictions using the testing set
diabetes y pred = regr.predict(diabetes X test)
#MSE and R2 on test set
MSE = mean_squared_error(diabetes_y_test, diabetes_y_pred)
RSQ = r2 score(diabetes y test, diabetes y pred)
```



Scikit-learn: cross-validation

```
from sklearn.model_selection import cross_val_score
#Cross-Validation
cross_val_score_regr = cross_val_score(regr, diabetes_X, diabetes_y, cv = 10)
print('Cross-Validated R^2: \n', cross_val_score_regr)
print('Mean Cross-Validated R^2 %.2f' %np.mean(cross_val_score_regr))

Cross-Validated R^2:
  [0.31895643 0.00210559 0.22467725 0.46430326 0.19557393 0.50309996 0.28038004 0.29884357 0.3088813 0.42764352]
Mean Cross-Validated R^2 0.30
```

- Two lines of code → ten-fold cross-validated linear regression.
- If you want to plot the lines per cross-validation along with train and test data: can use sklearn.cross_validation.Kfold yourself and supply to cross_val score as argument cv.

One-hot encoding:

- Many ML algorithms work on continuous numbers and see 3 > 2.
- If you have categorical variables (color = red | green | blue), you could encode them as 0, 1, 2. But that would make blue ,More coloury than green, which is ,more coloury than red. → wrong!

- One-hot encoding:
 - Many ML algorithms work on continuous numbers and see 3 > 2.
 - If you have categorical variables (color = red | green | blue), you could encode them as 0, 1, 2. But that would make blue ,More coloury' than green, which is ,more coloury' than red. Instead turn into 3 features:

red,	green,	blue
1,	0,	0
0,	1,	0
0,	0,	1

One-hot encoding:

One-hot encoding:

Same .fit() method as estimators

One-hot encoding:

transform() is similar to .predict() for estimators.

One-hot encoding:

Result is:

Sex	Geographic location	Browser
Female	US	Safari
Male	Europe	Safari



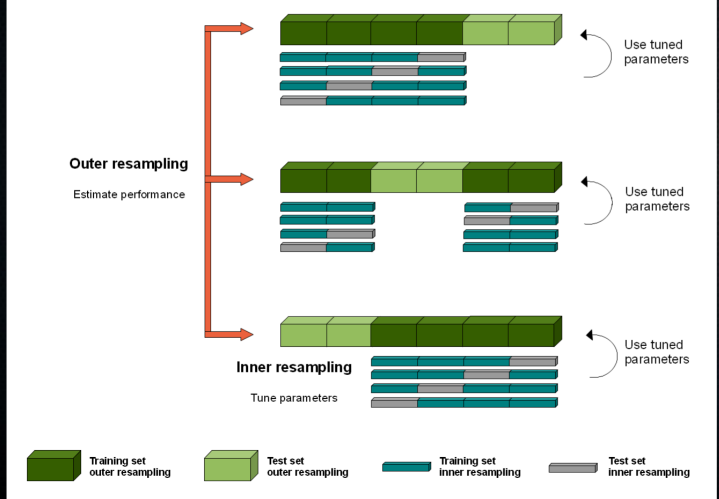
Female		From Europe	From US	Uses Firefox	Uses Safari
1	0	0	1	0	1
0	1	1	0	0	1

Scaling:

```
>>> from sklearn import preprocessing
>>> import numpy as np
>>> X train = np.array([[ 1., -1., 2.],
                  [2., 0., 0.],
...
                       [0., 1., -1.]])
. . .
>>> scaler = preprocessing.StandardScaler().fit(X train)
>>> scaler
StandardScaler()
>>> scaler.mean
array([1. ..., 0. ..., 0.33...])
>>> scaler.scale
array([0.81..., 0.81..., 1.24...])
>>> X scaled = scaler.transform(X train)
>>> X scaled
array([[ 0. ..., -1.22..., 1.33...],
      [ 1.22..., 0. ..., -0.26...],
      [-1.22..., 1.22..., -1.06...]
```

Scikit-learn: hyperparameter optimisation

Gold standard is nested cross-validation



Scikit-learn: hyperparameter optimisation

```
from sklearn.datasets import load_iris
from matplotlib import pyplot as plt
from sklearn.svm import SVC
from sklearn.model_selection import GridSearchCV, cross_val_score, KFold
import numpy as np
```

```
# Load the dataset
iris = load iris()
X iris = iris.data
y iris = iris.target
# Set up possible values of parameters to optimize over
p grid = {"C": [1, 10, 100],
          "gamma": [.01, .1]}
# We will use a Support Vector Classifier with "rbf" kernel
svm = SVC(kernel="rbf")
# Arrays to store scores
nested scores = np.zeros(NUM TRIALS)
# Loop 30 times
NUM TRIALS = 30
for i in range(NUM TRIALS):
    # Choose cross-validation techniques for the inner and outer loops,
    # independently of the dataset.
    # E.g "GroupKFold", "LeaveOneOut", "LeaveOneGroupOut", etc.
    inner cv = KFold(n splits=4, shuffle=True, random state=i)
    outer cv = KFold(n splits=4, shuffle=True, random state=i)
    # Nested CV with parameter optimization
    clf = GridSearchCV(estimator=svm, param_grid=p_grid, cv=inner_cv)
    nested score = cross_val_score(clf, X=X_iris, y=y_iris, cv=outer_cv)
    nested scores[i] = nested score.mean()
print(nested scores)
```

Scikit-learn: hyperparameter optimisation

```
from sklearn.datasets import load_iris
from matplotlib import pyplot as plt
from sklearn.svm import SVC
from sklearn.model_selection import GridSearchCV, cross_val_score, KFold
import numpy as np
```

```
[0.94683499 0.94683499 0.97297297 0.9601707 0.95999289 0.96692745 0.97332859 0.9601707 0.96034851 0.96639403 0.9601707 0.96674964 0.95999289 0.97332859 0.96674964 0.96639403 0.9735064 0.95999289 0.96639403 0.9735064 0.95963727 0.98684211 0.95359175 0.97332859 0.97332859 0.97332859 0.95359175 0.95981508 0.98008535 0.98008535 0.98008535]
```

```
# Load the dataset
iris = load iris()
X iris = iris.data
y iris = iris.target
# Set up possible values of parameters to optimize over
p grid = {"C": [1, 10, 100],
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   outer cv = KFold(n splits=4, shuffle=True, random state=i)
   # Nested CV with parameter optimization
    clf = GridSearchCV(estimator=svm, param_grid=p_grid, cv=inner_cv)
   nested score = cross val score(clf, X=X iris, y=y iris, cv=outer cv)
   nested scores[i] = nested_score.mean()
print(nested scores)
```

Scikit-learn: pipelines

Combine preprocessing steps and classification/regression

```
>>> 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
```

 Correct way is to: scale training data, save means and variances of training data, train classifier, subtract training data means and training data variances from test data, classify. → all done in simple step.

Scikit-learn: pipelines

Combine preprocessing steps and classification/regression

```
>>> from sklearn.pipeline import Pipeline
>>> from sklearn.svm import SVC
>>> from sklearn.decomposition import PCA
>>> estimators = [('reduce_dim', PCA()), ('clf', SVC())]
>>> pipe = Pipeline(estimators)
>>> pipe
Pipeline(steps=[('reduce_dim', PCA()), ('clf', SVC())])
```

Combine a PCA for dimensionality reduction with classification afterwards.

Scikit-learn: pipelines

Easily cross-validate entire pipeline, compare different dim.

reductions:

```
import numpy as np
import matplotlib.pyplot as plt
from sklearn.datasets import load_digits
from sklearn.model_selection import GridSearchCV
from sklearn.pipeline import Pipeline
from sklearn.svm import LinearSVC
from sklearn.decomposition import PCA, NMF
from sklearn.feature_selection import SelectKBest, chi2
```

```
pipe = Pipeline([
    # the reduce dim stage is populated by the param grid
    ('reduce dim', 'passthrough'),
    ('classify', LinearSVC(dual=False, max iter=10000))
1)
N FEATURES OPTIONS = [2, 4, 8]
C \text{ OPTIONS} = [1, 10, 100, 1000]
param grid = [
         'reduce dim': [PCA(iterated power=7), NMF()],
         'reduce dim n components': N FEATURES OPTIONS,
         'classify C': C OPTIONS
         'reduce dim': [SelectKBest(chi2)],
         'reduce dim k': N FEATURES_OPTIONS,
         'classify C': C_OPTIONS
    },
reducer labels = ['PCA', 'NMF', 'KBest(chi2)']
grid = GridSearchCV(pipe, n jobs=1, param grid=param grid)
X, y = load digits(return X y=True)
grid.fit(X, y)
```

Break for practical 1