cw1 solution

November 4, 2019

1 COMP3222/6246 Machine Learning Technologies (2019/20)

2 Solution of Coursework 1

Consider the diabetes data set from the scikit-learn python package. The structure of the data set is as follows: Each data point consists of 10 baseline variables (i.e., feature vector x): age, sex, body mass index, average blood pressure, and six blood serum measurements were obtained for each diabetes patient, as well as the response of interest (i.e., output variable y), a quantitative measure of disease progression one year after baseline.

Your task is to identify a good machine learning model to predict the value of y from a given feature vector x. To do so, you can use the above mentioned data set to train your model.

1. What is the python code to load the diabetes data set?

```
[1]: from sklearn.datasets import load_diabetes

diabetes = load_diabetes()
```

2. Uniformly randomly split the data set to training and test data with 80% for training and 20% for testing. Perform linear regression on the training data. What are the model parameters of the solution? What is the RMSE for this model on the training set? Apply this to the test data as well. What is the RMSE there? Repeat the uniformly random split 10 times and write down in a table each time the model parameters, RMSE for training, and RMSE for test data. How much do these values differ?

```
[2]: import math
    from sklearn.preprocessing import StandardScaler
    from sklearn.model_selection import train_test_split
    from sklearn.linear_model import LinearRegression
    from sklearn.metrics import mean_squared_error

for _ in range(10):

    X_train, X_test, Y_train, Y_test = train_test_split(diabetes.data, diabetes.
    →target, test_size=0.2)
    scaler = StandardScaler()
    scaled_X_train = scaler.fit_transform(X_train)
```

```
scaled_X_test = scaler.transform(X_test)
    print()
    lin_reg = LinearRegression()
    lin_reg.fit(scaled_X_train, Y_train)
    print("Hyper-parameters:", lin_reg.intercept_, lin_reg.coef_)
    Y_pred_train = lin_reg.predict(scaled_X_train)
    Y_pred_test = lin_reg.predict(scaled_X_test)
    rmse_train = math.sqrt(mean_squared_error(Y_train, Y_pred_train))
    print("Train RMS:", rmse_train)
    rmse_test = math.sqrt(mean_squared_error(Y_test, Y_pred_test))
    print("Test RMS:", rmse_test)
    print("Difference:", rmse_test - rmse_train)
Hyper-parameters: 152.96033994334277 [ -1.55111077 -12.38941212 23.96334328
15.31390911 -53.10647288
  31.08515011 16.69503909 17.51975235 40.28659377 4.22500358]
Train RMS: 53.172674411690195
Test RMS: 55.69413143782653
Difference: 2.521457026136332
Hyper-parameters: 153.08498583569406 [ -1.87296717 -11.63023155 23.86474236
17.97113429 -19.43800014
  11.86136633 -3.67856736
                            2.65175182 31.93779435 1.14699542]
Train RMS: 51.998929699886794
Test RMS: 59.81949940353726
Difference: 7.820569703650463
Hyper-parameters: 150.29461756373937 [ 1.75481875 -9.65302234 25.3221383
15.80631014 -46.97095782
 27.07218318 13.06060803 15.74934376 37.32435741 2.41030309]
Train RMS: 53.35327512769647
Test RMS: 54.81187957024428
Difference: 1.4586044425478093
Hyper-parameters: 150.1473087818697 [ -0.73803334 -10.66047112 25.40910272
14.0657154 -26.14839611
  15.42677628
               2.3016077
                           10.26105114 28.07057468 4.65446826]
Train RMS: 53.658922558588166
Test RMS: 53.78704965595728
Difference: 0.12812709736911643
Hyper-parameters: 152.87252124645892 [ -1.38363629 -14.22085676 25.74210451
14.87667123 -36.80855366
```

20.12755611 7.42408276 16.39588752 34.50785795 3.3669411]

Train RMS: 52.6029975429194
Test RMS: 58.2852199315305
Difference: 5.682222388611102

Hyper-parameters: 149.8498583569405 [0.15411928 -12.35541429 25.96934585

16.67542104 -39.80109416

25.06465294 4.73935293 6.97522446 34.63874708 2.65463571]

Train RMS: 52.707575896299986 Test RMS: 56.90228760209985 Difference: 4.194711705799861

Hyper-parameters: 155.671388101983 [-2.51094235 -12.51012137 22.92633494

17.47348112 -34.0381815

19.23181967 3.57212537 12.04348132 32.70882597 3.92871841]

Train RMS: 53.11380918225309 Test RMS: 55.96882413027572 Difference: 2.855014948022628

Hyper-parameters: 151.1359773371105 [0.47321325 -12.50591655 23.60392496

15.80771906 -44.27532643

25.99596704 8.46207037 12.12093258 38.19397723 2.78487767]

Train RMS: 53.50982005989561 Test RMS: 53.5598404188519

Difference: 0.050020358956288646

Hyper-parameters: 152.26912181303115 [-0.36832897 -10.54354831 25.31052907

14.71818179 -16.75337884

7.94160914 -5.75342392 4.76468363 26.16567287 3.95109799]

Train RMS: 54.252374747572695 Test RMS: 50.96174831380378 Difference: -3.2906264337689137

Hyper-parameters: 153.59773371104816 [0.27031703 -11.51020193 24.14969283

15.70248577 -26.27630864

14.79536014 1.49538952 5.33553083 30.40939787 5.22679259]

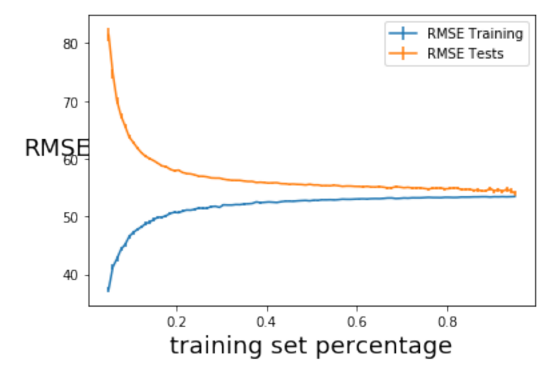
Train RMS: 53.3241281689678 Test RMS: 55.07227828018356 Difference: 1.7481501112157574

3. Investigate whether linear regression is an appropriate model for this data set. Does linear regression fit the data well? Explain your answer in 1-2 paragraphs (i.e., what can you say from the plot?). Hint: gradually increase the size of training data set, and plot the RMSE for both training and test data.

[4]: from sklearn.model_selection import train_test_split from sklearn.linear_model import LinearRegression from sklearn.metrics import mean_squared_error

```
import numpy as np
import math
repetition = 1000
percentages = np.linspace(0.05, 0.95, 100)
mean_rmse_trains = []
mean_rmse_tests = []
ci_rmse_trains = []
ci_rmse_tests = []
for percentage in percentages:
 rmse_trains = []
 rmse_tests = []
 for _ in range(repetition):
   X_train, X_test, Y_train, Y_test = train_test_split(diabetes.data, diabetes.
→target, test_size=1.0-percentage)
   lin_reg = LinearRegression()
   lin_reg.fit(X_train, Y_train)
   Y_pred_train = lin_reg.predict(X_train)
   Y_pred_test = lin_reg.predict(X_test)
   rmse_train = math.sqrt(mean_squared_error(Y_train, Y_pred_train))
   rmse_test = math.sqrt(mean_squared_error(Y_test, Y_pred_test))
   rmse_trains.append(rmse_train)
   rmse_tests.append(rmse_test)
 mean_rmse_train = np.mean(rmse_trains)
 mean_rmse_test = np.mean(rmse_tests)
 ci_rmse_train = 1.96 * np.std(rmse_trains) / math.sqrt(repetition)
 ci_rmse_test = 1.96 * np.std(rmse_tests) / math.sqrt(repetition)
 mean_rmse_trains.append(mean_rmse_train)
 mean_rmse_tests.append(mean_rmse_test)
 ci_rmse_trains.append(ci_rmse_train)
 ci_rmse_tests.append(ci_rmse_test)
import matplotlib.pyplot as plt
plt.errorbar(percentages, mean rmse_trains, yerr=ci_rmse_trains, label="RMSE_"
→Training")
plt.errorbar(percentages, mean_rmse_tests, yerr=ci_rmse_tests, label="RMSE_u
→Tests")
plt.xlabel("training set percentage", fontsize=18)
plt.ylabel("RMSE", fontsize=18, rotation=0)
```

```
plt.legend()
plt.show()
```



In general, linear regression fits the data well which was demonstrated by (1) a low difference between RMSE on training set and RMSE on testing set, and (2) given an increasing training datapoints, the RMSE on testing set decreases gracefully.

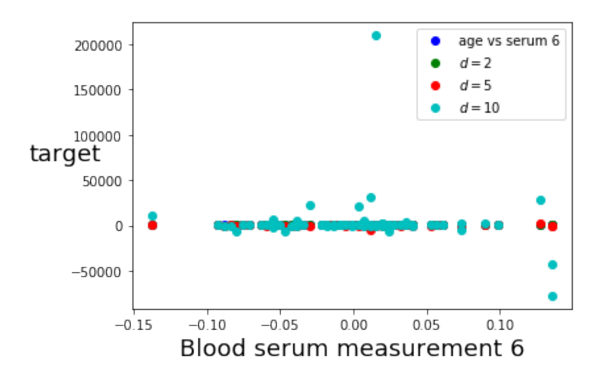
4. Use polynomial regression with degrees d=2.5 and 10 (Still with 80% of data for training and 20% for testing). Plot their predictions in the same plot. Explain what you can see within 1-2 paragraphs.

```
[5]: from sklearn.preprocessing import StandardScaler
from sklearn.preprocessing import PolynomialFeatures

def create_poly(d, x, y):
    poly_features = PolynomialFeatures(degree=d, include_bias=False)
    poly_train_data = poly_features.fit_transform(x)
    lin_reg = LinearRegression()
    lin_reg.fit(poly_train_data, y)
    return lin_reg

def predict_poly(p, d, test):
    poly_features = PolynomialFeatures(degree=d, include_bias=False)
    poly_test_data = poly_features.fit_transform(test)
    return p.predict(poly_test_data)
```

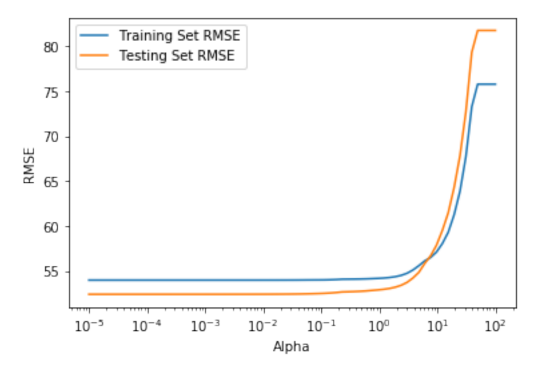
```
def compute_rmse(predict, actual):
        return math.sqrt(mean_squared_error(actual, predict))
     X_train, X_test, Y_train, Y_test = train_test_split(diabetes.data, diabetes.
     →target, test_size=0.2)
     scaler = StandardScaler()
     scaled_X_train = scaler.fit_transform(X_train)
     scaled_X_test = scaler.transform(X_test)
     pol_reg2 = create_poly(2, scaled_X_train, Y_train)
     pol_reg5 = create_poly(5, scaled_X_train, Y_train)
     pol_reg10 = create_poly(10, scaled_X_train, Y_train)
     Y_train_pred2 = predict_poly(pol_reg2, 2, scaled_X_train)
     Y_train_pred5 = predict_poly(pol_reg5, 5, scaled_X_train)
     Y_train_pred10 = predict_poly(pol_reg10, 10, scaled_X_train)
     Y_test_pred2 = predict_poly(pol_reg2, 2, scaled_X_test)
     Y_test_pred5 = predict_poly(pol_reg5, 5, scaled_X_test)
     Y_test_pred10 = predict_poly(pol_reg10, 10, scaled_X_test)
     print("Degree 2: Train RMSE =", compute_rmse(Y_train_pred2, Y_train), ", Test_
     →RMSE =", compute_rmse(Y_test_pred2, Y_test))
     print("Degree 5: Train RMSE =", compute_rmse(Y_train_pred5, Y_train), ", Test_
     →RMSE =", compute_rmse(Y_test_pred5, Y_test))
     print("Degree 10: Train RMSE =", compute_rmse(Y_train_pred10, Y_train), ", Test_
      →RMSE =", compute_rmse(Y_test_pred10, Y_test))
    Degree 2: Train RMSE = 48.06974292662504 , Test RMSE = 59.3764535933219
    Degree 5: Train RMSE = 3.978302254405894e-12 , Test RMSE = 627.8633144860745
    Degree 10: Train RMSE = 1.0729043995804305e-09 , Test RMSE = 24857.38687815741
[6]: # select feature's index and plot
     feat=9
     plt.plot(X_test[:, feat], Y_test, "bo", label="age vs serum 6")
     plt.plot(X_test[:, feat], Y_test_pred2, "go", label="$d=2$")
     plt.plot(X_test[:, feat], Y_test_pred5, "ro", label="$d=5$")
     plt.plot(X_test[:, feat], Y_test_pred10, "co", label="$d=10$")
     plt.xlabel("Blood serum measurement 6", fontsize=18)
     plt.ylabel("target", fontsize=18, rotation=0)
     plt.legend()
     plt.show()
```



In general, polynomial regression performs worse than linear regression since it overfits on the training set. The higher degree it is, the higher RMSE on test set and the lower RMSE on train set it yields.

5. Assuming that we still use 80% of the diabetes data set for training and 20% for testing, evaluate and explain the performance of Lasso with different alpha values within 1-2 paragraphs.

```
for i, alpha_lasso in enumerate(alpha_grid):
  # fit a lasso regressor
  lasso_reg = Lasso(alpha=alpha_lasso)
  lasso_reg.fit(scaled_X_train, Y_train)
  # compute RMSE
  Y_pred_train = lasso_reg.predict(scaled_X_train)
  Y_pred_test = lasso_reg.predict(scaled_X_test)
  train_rmse_grid[i] = math.sqrt(mean_squared_error(Y_train, Y_pred_train))
  test_rmse_grid[i] = math.sqrt(mean_squared_error(Y_test, Y_pred_test))
# plotting
plt.semilogx(alpha_grid, train_rmse_grid, label="Training Set RMSE")
plt.semilogx(alpha grid, test rmse grid, label="Testing Set RMSE")
plt.xlabel("Alpha")
plt.ylabel("RMSE")
plt.legend()
plt.show()
```



Performance of Lasso regression (L1 regularisation) heavily depends on the alpha value. If the alpha is too high, the penalty applies to minimisation problem will be large and a lot of coefficients will be eliminated; thus resulting in a large RMSE where the Lasso regression is unable to capture the trend of the data. However, if the alpha is too low, the penalty will be small and thus no coefficients are eleminated; thus the Lasso might be overfitting.

6. What would be the optimal value of alpha for Lasso on this data set? What are the resulted model parameters? Hint: use cross-validation and grid search to find the optimal alpha.

```
[8]: from sklearn.linear_model import LassoCV
    from sklearn.preprocessing import StandardScaler
    X_train, X_test, Y_train, Y_test = train_test_split(diabetes.data, diabetes.
     →target, test_size=0.2)
    scaler = StandardScaler()
    scaled_X_train = scaler.fit_transform(X_train)
    scaled X test = scaler.transform(X test)
    reg = LassoCV().fit(scaled_X_train, Y_train)
    Y_pred_train = reg.predict(scaled_X_train)
    Y_pred_test = reg.predict(scaled_X_test)
    print("Optimal alpha =", reg.alpha_)
    print("Train RMSE =", math.sqrt(mean_squared_error(Y_train, Y_pred_train)))
    print("Test RMSE =", math.sqrt(mean_squared_error(Y_test, Y_pred_test)))
    print("Coefficients =", reg.coef_)
    Optimal alpha = 2.240262675937059
    Train RMSE = 53.53053398986135
    Test RMSE = 56.15859317975826
    -6.84367938 24.16419391 16.14403784 -1.90915214
                  -10.06163911
                              0.
                                            20.80783338 1.461063087
      -0.
    C:\Local\anaconda3\envs\MLTech\lib\site-
    packages\sklearn\model selection\ split.py:1978: FutureWarning: The default
    value of cv will change from 3 to 5 in version 0.22. Specify it explicitly to
    silence this warning.
      warnings.warn(CV_WARNING, FutureWarning)
```

7. Implement a decision tree as a regression model for this data set. Repeat this 10 times. Visualise the best tree up to depth 5 and report the RMSE on both training and test data. What is the reason that we might get different trees and results? Please explain within 1-2 paragraphs.

```
[25]: from sklearn.tree import DecisionTreeRegressor
    from sklearn.tree import export_graphviz
    from IPython.display import Image
    import numpy as np

# make sure you installed graphviz (exclamation mark is for shell commands)
!apt install graphviz

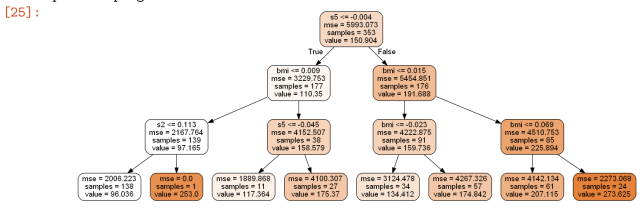
def get_best_decision_tree(max_depth=5, repetition=10):
    # repeat 10 times
    rmse_train = np.zeros(repetition)
    rmse_test = np.zeros(repetition)
    trees = []
```

```
for i in range(repetition):
    # partition dataset
    X_train, X_test, Y_train, Y_test = train_test_split(diabetes.data, diabetes.
 →target, train_size=0.8, test_size=0.2)
    # fit a decision tree regressor
    tree_reg = DecisionTreeRegressor(max_depth=max_depth)
    tree_reg.fit(X_train, Y_train)
    trees.append(tree_reg)
    # compute RMSE
    Y_pred_train = tree_reg.predict(X_train)
    Y_pred_test = tree_reg.predict(X_test)
    rmse_train[i] = math.sqrt(mean_squared_error(Y_train, Y_pred_train))
    rmse_test[i] = math.sqrt(mean_squared_error(Y_test, Y_pred_test))
  # find the "best" decision tree
  best = np.argmin(rmse_test)
  return [best, rmse_train[best], rmse_test[best], trees[best]]
print(get_best_decision_tree(1, 10))
print(get_best_decision_tree(2, 10))
print(get_best_decision_tree(3, 10))
print(get_best_decision_tree(4, 10))
print(get_best_decision_tree(5, 10))
best, _, _, best_tree = get_best_decision_tree(3, 10)
# show it off
export_graphviz(best_tree,
                out_file="my_spiky_tree.dot",
                feature names=diabetes.feature names,
                rounded=True,
                filled=True)
# convert dot file to png file.
!dot -Tpng my_spiky_tree.dot -o my_spiky_tree.png
# display the tree
Image(filename="my_spiky_tree.png")
[1, 65.17590124886883, 63.766902966198344,
```

```
DecisionTreeRegressor(criterion='mse', max_depth=1, max_features=None, max_leaf_nodes=None, min_impurity_decrease=0.0, min_impurity_split=None, min_samples_leaf=1,
```

```
min_samples_split=2, min_weight_fraction_leaf=0.0,
                      presort=False, random_state=None, splitter='best')]
[0, 59.18848398967014, 55.30872342377798, DecisionTreeRegressor(criterion='mse',
max_depth=2, max_features=None,
                      max leaf nodes=None, min impurity decrease=0.0,
                      min impurity split=None, min samples leaf=1,
                      min samples split=2, min weight fraction leaf=0.0,
                      presort=False, random state=None, splitter='best')]
[5, 55.11699124460894, 54.121341512367195,
DecisionTreeRegressor(criterion='mse', max_depth=3, max_features=None,
                      max_leaf_nodes=None, min_impurity_decrease=0.0,
                      min_impurity_split=None, min_samples_leaf=1,
                      min_samples_split=2, min_weight_fraction_leaf=0.0,
                      presort=False, random_state=None, splitter='best')]
[9, 49.15294883269271, 60.03903894424763, DecisionTreeRegressor(criterion='mse',
max_depth=4, max_features=None,
                      max_leaf_nodes=None, min_impurity_decrease=0.0,
                      min_impurity_split=None, min_samples_leaf=1,
                      min_samples_split=2, min_weight_fraction_leaf=0.0,
                      presort=False, random state=None, splitter='best')]
[2, 44.76058829733882, 60.548360116892134,
DecisionTreeRegressor(criterion='mse', max depth=5, max features=None,
                      max_leaf_nodes=None, min_impurity_decrease=0.0,
                      min impurity split=None, min samples leaf=1,
                      min_samples_split=2, min_weight_fraction_leaf=0.0,
                      presort=False, random_state=None, splitter='best')]
```

'apt' is not recognized as an internal or external command, operable program or batch file.



If the random seed which is used to permute features at each split is not fixed, then we will get a different tree at each time the tree is trained.