KernelCourse2020_Exercise01

February 6, 2020

1 Exercise 01

Kernel Methods in Machine Learning (CS-E4830)

Release date: 21 of January, 2020

Submission date: 06 of February, 2020 @4PM (no late submission allowed)

Tasks:

- 1. Section ??
- 2. Section ?? (1 Point)
- 3. Section ?? (2 Point)
- 4. Section ?? (2 Point)
- 5. Section ??
- 6. Section ?? (2 Point)
- 7. Section ??(1 Point)

Optional Tasks:

4. Section ?? (no points)

Version: 1.2

Version history: - 1.0: Initial version - 1.1: Fix typo: Parzen Window Clf. dual-variables (alpha) had wrong sign in formula. Issue on MyCourses - 1.2: Fix typo: Gaussian kernel doc-string, put brackets around the denominator within exponential function.

Please add your student number and email address to the notebook into the corresponding cell.

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1.1 Import required python packages

All tasks in this exercise can be solved by using only function and packages imported below. Please do not use any other imports.

```
[7]: import numpy as np
import matplotlib.pyplot as plt
import matplotlib.colors as colors
plt.set_cmap("RdBu_r")

from sklearn.base import BaseEstimator, ClassifierMixin
from sklearn.datasets import make_blobs, make_moons
from sklearn.model_selection import KFold, ParameterGrid, BaseCrossValidator,

→train_test_split
from sklearn.base import clone
```

<Figure size 432x288 with 0 Axes>

1.2 1. Kernel matrix calculation

Your task is to implement two functions to calculate Linear- and Gaussian-kernel matrices for two sets of feature vectors $\mathbf{X}_A \in \mathbb{R}^{n_A \times d}, \mathbf{X}_B \in \mathbb{R}^{n_B \times d}$, where d is the dimension of the feature vectors, and n_a and n_B are the number of examples in set A respectively B.

1.2.1 A. Linear kernel (1 Point)

Task:

 $\hookrightarrow None$, than Y = X

if X_B is None:

11 11 11

Implement missing code parts of the function calculation the linear kernel matrix given two features.

```
X_B = X_A
K_AB = np.dot(X_A, X_B.T)
```

:param X_A : array-like, shape= $(n_samples_A, d)$, feature-matrix of set A: param X_B : array-like, shape= $(n_samples_B, d)$, feature-matrix of set B or A

:return: array-like, shape=(n_samples_A, n_samples_B), kernel matrix

```
# print("shape of K_AB = ", K_AB.shape)
# raise NotImplementedError()

return K_AB
```

```
[3]: X_A = np.array([[1], [2], [3], [4]])
     X_B = np.array([[0], [2], [1]])
     # Test size
     np.testing.assert_equal(linear_kernel(X_A, X_B).shape, (4, 3))
     np.testing.assert_equal(linear_kernel(X_A).shape, (4, 4))
     # Test values
     np.testing.assert_equal(linear_kernel(X_A)[0, 0], 1)
     np.testing.assert_equal(linear_kernel(X_A)[1, 1], 4)
     np.testing.assert_equal(linear_kernel(X_A)[0, 2], 3)
     np.testing.assert_equal(linear_kernel(X_A)[2, 0], 3)
     np.testing.assert_equal(linear_kernel(X_A, X_B)[0, 0], 0)
     np.testing.assert_equal(linear_kernel(X_A, X_B)[0, 1], 2)
     # Test performance
     import time
     time_tolerance = 1 # in seconds
     size = 3000
     t0 = time.time()
     a = linear_kernel(np.random.rand(size), np.random.rand(size))
     t1 = time.time()
     total = t1 - t0
     print('Time to run with size {} = {:.4f} seconds'.format(size, total))
     assert total < time_tolerance, 'Time limit exceeded'</pre>
```

Time to run with size 3000 = 0.0011 seconds

1.2.2 B. Gaussian kernel (1 Point)

Task:

```
return np.exp(num/denom)
     def gaussian_kernel(X_A, X_B=None, sigma=None):
         Calculate the Gaussian kernel matrix, so that
              k_i j = exp(-1/x_i - x_j / 2 / (2 * sigma^2))
         :param X_A: array-like, shape=(n_samples_A, n_features), feature-matrix of (n_samples_A, n_features)
         :param X B: array-like, shape=(n \text{ samples } B, n \text{ features}), feature-matrix of \Box
      \rightarrowset B or None, than X_B = X_A
         :param sigma: scalar, bandwidth parameter
         :return: array-like, shape=(n_samples_A, n_samples_B), kernel matrix
         if X_B is None:
             X^B = X^A
         n_A = X_A.shape[0]
         n_B = X_B.shape[0]
         if sigma is None:
              sigma = np.sqrt(X_A.shape[1] / 2.0)
         X_A_{norm} = np.matrix(np.linalg.norm(X_A, axis = 1)).T
           print("X A norm shape = ", X A norm.shape)
         X_B_norm = np.matrix(np.linalg.norm(X_B, axis = 1))
           print("X_B norm shape = ", X_B_norm.shape)
         X_A_B_{dot} = np.dot(X_A, X_B.T)
           print("X_A_B_dot.shape = ", X_A_B_dot.shape)
           print(np.square(X\_A\_norm).shape, np.square(X\_B\_norm).shape, (2*X\_A\_B\_dot).
      \hookrightarrowshape)
         K_AB = np.square(X_A_norm) + np.square(X_B_norm) - 2*X_A_B_dot
         K_AB = np.exp(-K_AB/(2*np.square(sigma)))
           print("returning K_AB, shape = ", K_AB.shape)
         return K AB
[5]: X_A = np.array([[1], [2], [3], [4]])
     X_B = np.array([[0], [2], [1]])
     # Test size
     np.testing.assert_equal(gaussian_kernel(X_A, X_B).shape, (4, 3))
     np.testing.assert_equal(gaussian_kernel(X_A).shape, (4, 4))
     # Test values
```

```
np.testing.assert_equal(gaussian_kernel(X_A, sigma=1)[0, 0], 1)
np.testing.assert_equal(gaussian_kernel(X_A, sigma=1)[1, 1], 1)
np.testing.assert_equal(gaussian_kernel(X_A, sigma=1)[0, 2], np.exp(-2))
np.testing.assert_equal(gaussian_kernel(X_A, sigma=1)[2, 0], np.exp(-2))
np.testing.assert_equal(gaussian_kernel(X_A, X_B, sigma=1)[0, 0], np.exp(-0.5))
np.testing.assert_equal(gaussian_kernel(X_A, X_B, sigma=1)[0, 1], np.exp(-0.5))
# Test performance
import time
time tolerance = 1 # in seconds
size = 3000
features = 10
t0 = time.time()
a = gaussian kernel(np.random.rand(size, features), np.random.rand(size,

→features))
t1 = time.time()
total = t1 - t0
print('Time to run with size {} = {:.4f} seconds'.format(size, total))
assert total < time_tolerance, 'Time limit exceeded'</pre>
```

Time to run with size 3000 = 0.6673 seconds

1.3 2. Parzen Window Classifier implementation (2 Points)

The Parzen Window Classifier prediction model can be written as:

$$h(\mathbf{x}) = \operatorname{sign}(g(\mathbf{x})) = \operatorname{sign}\left(\sum_{i=1}^{n} \alpha_i \kappa(\mathbf{x}_i, \mathbf{x}) + b\right),$$

with: - $g: \mathbb{R}^d \to \mathbb{R}$ being the **decision function** - $b \in \mathbb{R}$ being the **bias term** defined as:

$$b = \frac{1}{2n_{-}^{2}} \sum_{i,j \in I^{-}} \kappa(\mathbf{x}_{i}, \mathbf{x}_{j}) - \frac{1}{2n_{+}^{2}} \sum_{i,j \in I^{+}} \kappa(\mathbf{x}_{i}, \mathbf{x}_{j}),$$

• α_i 's $\in \mathbb{R}$ being the **dual variables** for all training examples \mathbf{x}_i defined as:

$$\alpha_i = \begin{cases} \frac{1}{n_+} & \text{if } y_i = +1\\ -\frac{1}{n_-} & \text{if } y_i = -1 \end{cases}.$$

We denote the number of positive / negative training examples as n_+ / n_- , and $n = n_+ + n_-$, and I^+ / I^- are the indices of the positive / negative training examples.

Below you find the class-template for the Parzen Window Classifier. It's functionality is split into three parts:

1.3.1 1. Intialization of Classifier Object using init()

A Parzen Window Classifier instance can be created using its constructor and the kernel to be used can be specified, e.g.:

```
est = ParzenWindowClassifier(kernel="gaussian").
```

1.3.2 2. Model Training using fit()

This function takes as input the features of the training examples \mathbf{X}_{train} and their corresponding labels $\mathbf{y}_{train} \in \{-1, 1\}^{n_{train}}$ and estimates the α_i 's and b. The necessary kernel values between the training examples, i.e. $\kappa(\mathbf{x}_i, \mathbf{x}_j)$ are calculated during the fitting process.

```
est.fit(X_train, y_train)
```

1.3.3 3. Prediction for new Examples using predict() and decision_function()

When the model parameters, i.e. b and α_i s, are fitted, than we can make predictions for a new example \mathbf{x} using the function $h(\mathbf{x})$.

```
y_test_pred = est.predict(X_test)
```


Task: Implement the missing code parts of <code>fit()</code>, <code>decision_function()</br>

Hint: The NumPy function <code>np.sum</code> can be used to sum over the elements of a new code sum over the elements of a new code>np.sum</code></br>

```
[8]: class ParzenWindowClassifier(BaseEstimator, ClassifierMixin):
         def __init__(self, kernel="gaussian", sigma=None):
              Parzen Window Classifier
              :param kernel: string, specifying which kernel to use. Can be
      → 'qaussian' or 'linear'.
              :param sigma: scalar, gaussian kernel parameter, can be None if the_{\!\scriptscriptstyle \sqcup}
      \hookrightarrow linear kernel is used.
              11 11 11
              # Parzen Window Classifier model parameter
              self.b = None # bias term
              self.alphas = None # dual variables
              # Training data needed for the prediction phase
              self.X_train = None
              # Set up kernel function
              self.kernel = kernel
              self.sigma = sigma
         def fit(self, X_train, y_train):
```

```
Fit a Parzen Window Classifier using training data
       :param X_train: array-like, shape=(n_samples, n_features), __
\hookrightarrow feature-matrix
        :param y_{train}: array-like, shape=(n_{samples}) or (n_{samples}, 1), label_{\sqcup}
\hookrightarrow vector
       # Calculate the specified kernel
       self.X_train = X_train
       KX_train = self._get_kernel(self.X_train)
       # Get indices of positive and negative examples: I n, I p
       I_n = np.where(y_train==-1)[0]
       I_p = np.where(y_train==1)[0]
       # Count the number of postitive and negative examples: n_n, n_p
       n_n = I_n.size
       n_p = I_p.size
       # Calcualte the bias term: self.b
       if self.kernel == "gaussian":
           kernel_sum_neg = np.sum(gaussian_kernel(X_train[I_n]))
           kernel_sum_pos = np.sum(gaussian_kernel(X_train[I_p]))
       elif self.kernel == "linear":
           kernel_sum_neg = np.sum(linear_kernel(X_train[I_n]))
           kernel_sum_pos = np.sum(linear_kernel(X_train[I_p]))
       self.b = 1/(2*n n**2)*kernel sum neg - 1/(2*n p**2)*kernel sum pos
       # TODO : try it using the already calculated kernel in self._get_kernel
       # Calculate alpha_i's: self.alpha
       self.alphas = np.zeros((n_n + n_p, 1))
       for i in range(self.alphas.size):
            self.alphas[i] = 1/n_p if y_train[i] == 1 else -1/n_n
   def decision_function(self, X):
       Calculate decision function:
            g(x) = sum_i a_i k(x_i, x) + b
       :param X: array-like, shape=(n_samples_test, n_features),_
\hookrightarrow feature-matrix of new data.
       :return: array-like, shape=(n_samples_test,), decision function value<sub>□</sub>
\rightarrow q(x) for all new data points
```

```
if self.alphas is None or self.b is None or self.X_train is None:
           raise RuntimeError("Call fit-function first.")
       # Calculate the specified kernel between the training and test examples
       KX_test_train = self._get_kernel(X, self.X_train)
       # Calculate the value of the decision function for each test example
       g X = np.zeros((X.shape[0],1))
       for i in range(X.shape[0]):
           for j in range(self.X_train.shape[0]):
               g_X[i] += self.alphas[j]*KX_test_train[i,j]
           g_X[i] += self.b
       return g_X.flatten() # output a one-dimensional vector
   def predict(self, X):
       Predict labels using Parzen Window Classifier:
           h(x) = sign(g(x)), with g(x) being the decision function
       :param X: array-like, shape=(n_samples_test, n_features),_
\rightarrow feature-matrix of new data.
       :return: array-like, shape=(n_samples_test,), predicted labels \{-1, 1\}_{\sqcup}
\hookrightarrow for all new data points
       if self.alphas is None or self.b is None or self.X train is None:
           raise RuntimeError("Call fit-function first.")
       # Calculate prediction h(x) = sign(g(x))
       pred = self.decision_function(X)
       h_X = np.where(pred>0, 1, -1)
       return h X
   def _get_kernel(self, X, Y=None):
       Calcualte kernel matrix using specified kernel-function and parameters.
       :param X: array-like, shape=(n samples_A, n_features), feature-matrix

⊥
\hookrightarrow of set A
       :param\ Y:\ array-like,\ shape=(n\_samples\_B,\ n\_features),\ feature-matrix_{\sqcup}
\rightarrow of set B or None, than Y = X
       :return: array-like, shape=(n_samples_A, n_samples_B), kernel matrix
       if self.kernel == "gaussian":
           return gaussian_kernel(X, Y, self.sigma)
```

```
elif self.kernel == "linear":
    return linear_kernel(X, Y)
else:
    raise ValueError("Invalid kernel chosen.")
```

```
[9]: X, y = make_blobs(n_samples=50, centers=[[1, 1], [1.5, 1.5]], cluster_std=[0.1,__
     \hookrightarrow 0.1],
                        random_state=80)
     y[y==0] = -1
     test_est = ParzenWindowClassifier(kernel="linear")
     test_est.fit(X, y)
     # Test correct shape of self.b and self.alphas
     assert(np.isscalar(test_est.b))
     np.testing.assert_equal(test_est.alphas.shape, (50, 1))
     # Test correct shape of the predictions
     X_{\text{test}} = \text{np.array}([[1, 1], [1.5, 1.5], [1, 0], [2, 3]])
     # Validate predictions
     np.testing.assert_equal(test_est.decision_function(X_test).shape, (4,))
     np.testing.assert_equal(test_est.predict(X_test).shape, (4,))
     np.testing.assert_equal(test_est.predict(X_test), np.array([-1, 1, -1, 1]))
     # Validate score
     np.testing.assert_equal(test_est.score(X, y), 1.0)
```

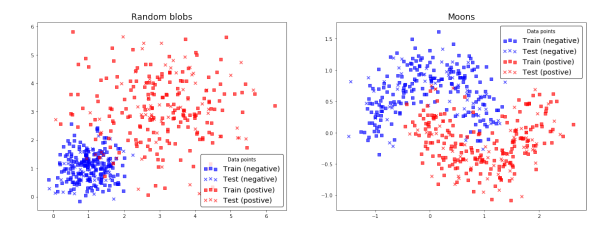
1.4 3. Application of the Parzen Window Classifier

It is time to apply your Parzen Window Classifier to some datasets. For that you are given two synthetic datasets \mathbf{X} . Each is splitted into a training and test subset: \mathbf{X}_{train} (75%) and \mathbf{X}_{test} (25%).

```
X_blobs_train, X_blobs_test, y_blobs_train, y_blobs_test = train_test_split(
    X_blobs, y_blobs, random_state=319)
X_moons_train, X_moons_test, y_moons_train, y_moons_test = train_test_split(
    X_moons, y_moons, random_state=747)
```

We can plot the datasets to get an impression what we are dealing with:

```
[11]: # Plot datasets
     fig, axrr = plt.subplots(1, 2, figsize=(20, 7))
     # Blobs
     for l_str, l_num, col in [("negative", -1, "blue"), ("postive", 1, "red")]:
         axrr[0].scatter(
             X_blobs_train[y_blobs_train==l_num, 0],__
      →X_blobs_train[y_blobs_train==l_num, 1],
             c=col, alpha=0.65, label="Train (%s)" % 1_str, marker="s")
         axrr[0].scatter(
             X_blobs_test[y_blobs_test==l_num, 0], X_blobs_test[y_blobs_test==l_num, u]
      \hookrightarrow1],
             c=col, alpha=0.65, label="Test (%s)" % l_str, marker="x")
     # Blobs
     for l_str, l_num, col in [("negative", -1, "blue"), ("postive", 1, "red")]:
         axrr[1].scatter(
             X_moons_train[y_moons_train==l_num, 0],__
      →X_moons_train[y_moons_train==l_num, 1],
             c=col, alpha=0.65, label="Train (%s)" % l_str, marker="s")
         axrr[1].scatter(
             X moons_test[y_moons_test==l_num, 0], X moons_test[y_moons_test==l_num, u]
      →1],
             c=col, alpha=0.65, label="Test (%s)" % l_str, marker="x")
     axrr[0].set_title("Random blobs", fontsize="xx-large")
     →edgecolor="k")
     axrr[1].set_title("Moons", fontsize="xx-large")
     axrr[1].legend(title="Data points", fontsize="x-large", scatterpoints=3, __
      →edgecolor="k")
     plt.show()
```



1.4.1 A. Implement the hyper parameter optimization (1 point)

Train (fit) your Parzen Window Classifier with Gaussian kernel on the training examples, i.e. X_blobs_train and X_moons_train . To find the optimal Gaussian bandwidth parameter σ we search a grid of different parameter values and score each one using CV.

we search a grid of different parameter values and score each one using CV.

The function gets in a set of training examples and a grid of different parameters, e.g. σ 's, calculates an average validation set score for all of them using cross-validation. Subsequently, a model using all the training data ist used, to train a model with the best set of parameters.

Hints:

- Make use of the Python tutorial (fetch on JupyterHub), if you want to see the pseudo-code of the parameter search.
- Read the documentation of the sklearn KFold function.
- You can calculate the performance score using est.score(...). Read also: https://scikit-learn.org/stable/modules/generated/sklearn.base.ClassifierMixin.html#sklearn.base.ClassifierMixin.score

Task: Implement the missing code parts of the <code>hyper_parameter_search_using_cv</code

```
[15]: def hyper_parameter_search_using_cv(estimator, X, y, param_grid, n_cv_folds=5, random_state=None):

"""

Function calculating the estimator score for a grid of hyper parameters.

:param estimator: object, subclass of RegressorMixin or ClassifierMixin and_□

→BaseEstimator

:param X: array-like, shape=(n_samples, n_features), feature-matrix used_□

→for training

:param y: array-like, shape=(n_samples,) or (n_samples, 1), label vector_□

→used for training

:param param_grid: dictionary,

keys: different parameter names
```

```
values: grid-values for each parameter
    :param n cv folds: scalar, a KFold cross-validation is performed where the \Box
→number of splits is equal the scalar.
    : param\ random\_state:\ scalar,\ RandomState\ instance\ or\ None,\ optional, \\ \sqcup
\hookrightarrow default=None
        If int, random state is the seed used by the random number generator;
        If RandomState instance, random_state is the random number generator;
        If None, the random number generator is the RandomState instance used
        by `np.random`.
    :return: tuple = (
            best estimator,
            param grid as list and corresponding evaluation scores,
            score of best parameter
            best parameter
    11 11 11
    # Get an iterator over all parameters
   param_grid_iter = ParameterGrid(param_grid)
    # Create cross-validation object
   cv = KFold(n_splits=n_cv_folds, random_state=random_state)
   # Store the valdidation set performance scores for all folds and parameters
   perf_scores = np.zeros((cv.get_n_splits(), len(param_grid_iter)))
     print("perf_scores:", perf_scores)
   for fold, (train_set, val_set) in enumerate(cv.split(X, y)):
        # Separate training and validation set from X and y,
        # i.e. X_train, X_val, y_train and y_val
          print("fold: ", fold)
        X_train = X[train_set]
        y_train = y[train_set]
        X val = X[val set]
        y_val = y[val_set]
        for idx, param in enumerate(param_grid_iter):
#
              print("idx,param ", idx, param)
            # Clone the estimator object to get an un-initialized object
            est = clone(estimator)
            # Set model parameters
            est.set_params(**param)
            # Fit the model using training set
            est.fit(X_train, y_train)
```

```
print("est predict: ",est.predict(X_train))
                  \# Calculate the perf. score on validation set for current fold and
       \rightarrow parameter index
                  perf_scores[fold][idx] = est.score(X_val, y_val)
                    print("perf_score: ", perf_scores[fold][idx])
          # Find best performing hyper-parameter
          print("perf_scores: ", perf_scores)
          avg_perf_scores = np.mean(perf_scores, axis=0)
          # Average the perf. scores for each parameter across each fold
          print("avg perf scores: ", np.mean(perf_scores, axis=0))
          idx_best = np.argmax(avg_perf_scores)
          best_perf_score = avg_perf_scores[idx_best]
          best_param = param_grid_iter[idx_best]
          # Fit model using all data with the best parameters
          est = clone(estimator)
          est.set params(**best param)
          est.fit(X, y)
            print("returning: ")
           print((est, {"params": list(param grid iter), "scores": avg perf scores},
      #
                    best_perf_score, best_param))
          return (est, {"params": list(param_grid_iter), "scores": avg_perf_scores},
                  best_perf_score, best_param)
[17]: X, y = make_blobs(n_samples=100, centers=[[1, 1], [1.5, 1.5]], cluster_std=[0.
      -2, 0.75],
                        random_state=100)
      v[v==0] = -1
      _param_grid = {"sigma": [0.05, 0.1, 0.5, 1.]}
      _est, _param_scores, _best_score, _best_param = hyper_parameter_search_using_cv(
          ParzenWindowClassifier(kernel="gaussian"), X, y, _param_grid)
      np.testing.assert_equal(_param_scores["scores"].shape,__
      →(len(_param_grid["sigma"]),))
      np.testing.assert_allclose(_param_scores["scores"], np.array([0.83, 0.91, 0.89,__
      →0.83]))
      np.testing.assert_equal(_best_param, {"sigma": 0.1})
      np.testing.assert_allclose(_best_score, 0.91)
      assert(isinstance( est, ParzenWindowClassifier))
```

```
perf_scores: [[0.5 0.5 0.85 0.85]
 [0.45 0.45 0.9 0.9]
 [0.4 0.4 0.95 0.8]
 [0.65 0.65 0.8 0.75]
 [0.5 0.5 0.9 0.85]]
avg perf scores: [0.5 0.5 0.88 0.83]
        AssertionError
                                                  Traceback (most recent call_
 →last)
        <ipython-input-17-6316946dc64e> in <module>
          9 np.testing.assert_equal(_param_scores["scores"].shape,_
 →(len(_param_grid["sigma"]),))
    ---> 10 np.testing.assert_allclose( param scores["scores"], np.array([0.83,__
 \rightarrow0.91, 0.89, 0.83]))
         11 np.testing.assert equal(best param, {"sigma": 0.1})
         12 np.testing.assert_allclose(_best_score, 0.91)
        /opt/conda/lib/python3.7/site-packages/numpy/testing/_private/utils.py_
 →in assert_allclose(actual, desired, rtol, atol, equal_nan, err_msg, verbose)
                header = 'Not equal to tolerance rtol=%g, atol=%g' % (rtol, atol)
                {\tt assert\_array\_compare(compare, actual, desired,\_} \\

→err_msg=str(err_msg),
    -> 1515
                                     verbose=verbose, header=header,⊔
 →equal_nan=equal_nan)
       1516
       1517
        opt/conda/lib/python3.7/site-packages/numpy/testing/_private/utils.py∪
 →in assert_array_compare(comparison, x, y, err_msg, verbose, header, precision, u
 →equal_nan, equal_inf)
        839
                                            verbose=verbose, header=header,
        840
                                            names=('x', 'y'),
 →precision=precision)
    --> 841
                       raise AssertionError(msg)
        842
                except ValueError:
        843
                    import traceback
```

AssertionError:

```
Not equal to tolerance rtol=1e-07, atol=0

Mismatch: 75%

Max absolute difference: 0.41

Max relative difference: 0.45054945

x: array([0.5, 0.5, 0.88, 0.83])

y: array([0.83, 0.91, 0.89, 0.83])
```

1.4.2 B. Plot validation score for different hyper parameters

With the hyper parameter optimization function at hand, we find the best σ parameter for the two synthetic datasets and also inspect the average validation error for different values of σ .

First let us the parameter grid for the Gaussian kernel bandwidth parameter σ :

```
[18]: param_grid = {"sigma": [0.01, 0.05, 0.1, 0.5, 1, 5, 10, 50]}
```

Now we run the hyper_parameter_search_using_cv function for X_blobs_train and X_blobs_test :

```
est_blobs, scores_blobs, best_score_blobs, best_param_blobs =_u

hyper_parameter_search_using_cv(

ParzenWindowClassifier(kernel="gaussian"), X_blobs_train, y_blobs_train,_u

param_grid,

random_state=737)

print("[Blobs] Best average validation score", best_score_blobs)

print("[Blobs] Best parameter", best_param_blobs)

est_moons, scores_moons, best_score_moons, best_param_moons =_u

hyper_parameter_search_using_cv(

ParzenWindowClassifier(kernel="gaussian"), X_moons_train, y_moons_train,_u

param_grid,

random_state=747)

print("[Moons] Best average validation score", best_score_moons)

print("[Moons] Best parameter", best_param_moons)
```

/opt/conda/lib/python3.7/site-packages/sklearn/model_selection/_split.py:296: FutureWarning: Setting a random_state has no effect since shuffle is False. This will raise an error in 0.24. You should leave random_state to its default (None), or set shuffle=True.

FutureWarning

```
perf_scores: [[0.49333333 0.49333333 0.89333333 0.97333333
0.69333333
0.49333333 0.49333333]
```

```
[0.49333333 0.49333333 0.49333333 0.90666667 0.96
                                                              0.62666667
       0.49333333 0.49333333]
      [0.49333333 0.49333333 0.49333333 0.90666667 0.97333333 0.58666667
       0.49333333 0.49333333]]
     avg perf scores: [0.49333333 0.49333333 0.89866667 0.97066667
     0.61333333
      0.49333333 0.49333333]
     [Blobs] Best average validation score 0.9706666666666667
     [Blobs] Best parameter {'sigma': 1}
     perf_scores: [[0.41333333 0.6 0.90666667 0.89333333 0.84
                                                                            0.8
       0.74666667 0.41333333]
      Γ0.52
                  0.69333333  0.94666667  0.85333333  0.78666667  0.76
       0.81333333 0.52
      [0.53333333 0.81333333 0.94666667 0.89333333 0.8
                                                              0.76
       0.76
                  0.45333333]
      [0.41333333 0.69333333 0.88 0.93333333 0.82666667 0.8
                  0.413333333
       0.8
      [0.58666667 0.88
                             0.93333333 0.93333333 0.92
                                                              0.92
       0.86666667 0.573333333]]
     avg perf scores: [0.49333333 0.736 0.92266667 0.90133333 0.83466667 0.808
      0.79733333 0.47466667]
     [Moons] Best average validation score 0.9226666666666666
     [Moons] Best parameter {'sigma': 0.1}
     Plot validation score for the different parameter values.
     <b>Hint:</b> Both curves should have a single maxima, if you did not change any random seeds,
[20]: fig, axrr = plt.subplots(1, 2, figsize=(14, 4), sharex="row")
      axrr[0].plot(param_grid["sigma"], scores_blobs["scores"], '*-')
      axrr[0].set xscale("log")
      axrr[0].grid(True, which="both", ls="-", alpha=0.5)
      axrr[0].set title("Random blobs: Validation accuracy")
      axrr[0].set_xlabel("Gaussian bandwidth parameter")
      axrr[0].set_ylabel("Mean accuracy (perf. score)")
      axrr[1].plot(param_grid["sigma"], scores_moons["scores"], '*-')
      axrr[1].set_xscale("log")
      axrr[1].grid(True, which="both", ls="-", alpha=0.5)
      axrr[1].set_title("Moons: Validation accuracy")
      axrr[1].set_xlabel("Gaussian bandwidth parameter")
      axrr[1].set_ylabel("Mean accuracy (perf. score)")
```

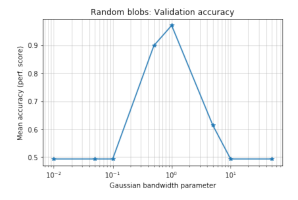
[0.45333333 0.45333333 0.45333333 0.89333333 0.97333333 0.53333333

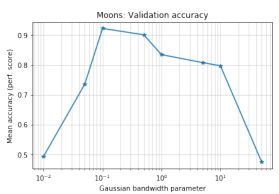
[0.53333333 0.53333333 0.53333333 0.89333333 0.97333333 0.62666667

0.45333333 0.453333333]

0.53333333 0.533333333]

plt.show()





The hyper-parameter optimization was done only using the training data. We now can apply the best model (with the optimal σ parameter) to the training data. We should see a similar performance as for the validation set:

```
[21]: print("[Blobs] Score on test set", est_blobs.score(X_blobs_test, y_blobs_test))
print("[Moons] Score on test set", est_moons.score(X_moons_test, y_moons_test))
```

```
[Blobs] Score on test set 0.936
[Moons] Score on test set 0.872
```

1.5 4. Model visualization: Non-linear vs. Linear (no points)

```
<b>Bonus task:</b>
```

Here we inspect the decision function and how it separates the two classes in the feature space. For that we will evaluate $g(\mathbf{x})$ for \mathbf{x} 's on a regular grid.

- 1. Set up interval to plot decision function, e.g. min. and max. value of our synthetic datasets.
- 2. Create grid-points \mathbf{x} covering the intervall.
- 3. Evaluate $z = g(\mathbf{x})$ for all points.
- 4. Plot the z's for different points and color them according to their sign.

```
[59]: def get_gridpoints(X):

"""

:param X: array-like, shape=(n_samples, d), datset feature matrix

:return: array-like, shape=(n_grid_samples, d), feature vectors on a

→regular grid in the feature space

"""

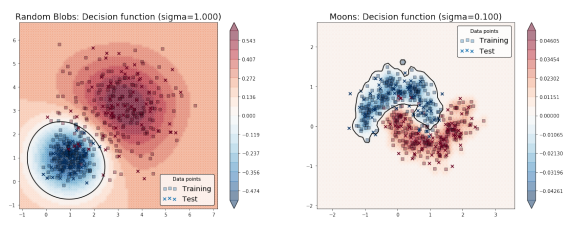
x_min, x_max = X[:, 0].min() - 1, X[:, 0].max() + 1

y_min, y_max = X[:, 1].min() - 1, X[:, 1].max() + 1
```

1.5.1 Decision Function for Gaussian Kernel (Model of previous tasks)

```
[61]: fig, axrr = plt.subplots(1, 2, figsize=(20, 7))
      # plot contours, levels, ...
      pcm = axrr[0].pcolormesh(XX_blobs_grid, YY_blobs_grid, Z_blobs_grid, alpha=0.5,_
      ⇔shading="gouraud",
                         norm=get_color_normalizer(Z_blobs_grid))
      fig.colorbar(pcm, ax=axrr[0], extend="both", orientation="vertical")
      axrr[0].contour(XX_blobs_grid, YY_blobs_grid, Z_blobs_grid, colors=['k'],_
      →linestyles=['-'], levels=[0])
      # plot points
      axrr[0].scatter(X_blobs_train[:, 0], X_blobs_train[:, 1], c=y_blobs_train,__
      →marker="s", edgecolors="k", alpha=0.35)
      axrr[0].scatter(X_blobs_test[:, 0], X_blobs_test[:, 1], c=y_blobs_test,__
      →marker="x", edgecolors="k")
      # plot labels, titles, ...
      axrr[0].legend(["Training", "Test"], title="Data points", fontsize="x-large", u
      ⇒scatterpoints=3, edgecolor="k")
      axrr[0].set_title("Random Blobs: Decision function (sigma=%.3f)" %__
       ⇔best_param_blobs["sigma"],
                       fontsize="xx-large")
```

```
pcm = axrr[1].pcolormesh(XX_moons_grid, YY_moons_grid, Z_moons_grid, alpha=0.5,_
⇒shading="gouraud",
                  norm=get_color_normalizer(Z_moons_grid))
fig.colorbar(pcm, ax=axrr[1], extend="both", orientation="vertical")
axrr[1].scatter(X_moons_train[:, 0], X_moons_train[:, 1], c=y_moons_train,__
→marker="s", edgecolors="k", alpha=0.35)
axrr[1].scatter(X_moons_test[:, 0], X_moons_test[:, 1], c=y_moons_test,__
 axrr[1].legend(["Training", "Test"], title="Data points", fontsize="x-large",
⇔scatterpoints=3, edgecolor="k")
axrr[1].set_title("Moons: Decision function (sigma=%.3f)" %__
→best_param_moons["sigma"],
                fontsize="xx-large")
axrr[1].contour(XX_moons_grid, YY_moons_grid, Z_moons_grid, colors=['k'],_
→linestyles=['-'], levels=[0])
plt.show()
```



1.5.2 Decision Function for Linear Kernel

We have seen that the Gaussian kernel leads to a non-linear decision boundary, i.e. $g(\mathbf{x}) = 0$ (black line in plots). Now we take a look on the linear Parzen Window Classifier, buy using a linear kernel.

```
[62]: est_blobs = ParzenWindowClassifier(kernel="linear")
    est_blobs.fit(X_blobs_train, y_blobs_train)

est_moons = ParzenWindowClassifier(kernel="linear")
    est_moons.fit(X_moons_train, y_moons_train)
```

[63]:

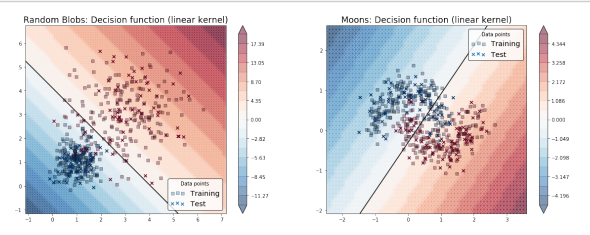
```
[64]: fig, axrr = plt.subplots(1, 2, figsize=(20, 7))
     # plot contours, levels, ...
     pcm = axrr[0].pcolormesh(XX_blobs_grid, YY_blobs_grid, Z_blobs_grid, alpha=0.5,_
      ⇔shading="gouraud",
                       norm=get color normalizer(Z blobs grid))
     fig.colorbar(pcm, ax=axrr[0], extend="both", orientation="vertical")
     axrr[0].contour(XX_blobs_grid, YY_blobs_grid, Z_blobs_grid, colors=['k'],__
      →linestyles=['-'], levels=[0])
     # plot points
     axrr[0].scatter(X_blobs_train[:, 0], X_blobs_train[:, 1], c=y_blobs_train,__
      →marker="s", edgecolors="k", alpha=0.35)
     axrr[0].scatter(X blobs test[:, 0], X blobs test[:, 1], c=y blobs test,
      # plot labels, titles, ...
     axrr[0].legend(["Training", "Test"], title="Data points", fontsize="x-large", u

→scatterpoints=3, edgecolor="k")
     axrr[0].set_title("Random Blobs: Decision function (linear kernel)", ___
      pcm = axrr[1].pcolormesh(XX_moons_grid, YY_moons_grid, Z_moons_grid, alpha=0.5,_u
      ⇔shading="gouraud",
                       norm=get color normalizer(Z moons grid))
     fig.colorbar(pcm, ax=axrr[1], extend="both", orientation="vertical")
     axrr[1].scatter(X_moons_train[:, 0], X_moons_train[:, 1], c=y_moons_train,__
      →marker="s", edgecolors="k", alpha=0.35)
     axrr[1].scatter(X_moons_test[:, 0], X_moons_test[:, 1], c=y_moons_test,__

    dedgecolors="k", marker="x")

     axrr[1].legend(["Training", "Test"], title="Data points", fontsize="x-large", u
      axrr[1].set_title("Moons: Decision function (linear kernel)", ___
      axrr[1].contour(XX_moons_grid, YY_moons_grid, Z_moons_grid, colors=['k'],_
      →linestyles=['-'], levels=[0])
```

plt.show()



[]: