



**MSc in Computer Science**  
at University of Milan

Statistical Methods for Machine Learning  
Kernelized Linear Predictor

course held by **Nicoló Cesa-Bianchi**

Email:  
davide.cogni@studenti.unimi.it

Created by:  
**Davide Cogni**  
mat. 09732A

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# 1 introduction

## 1.1 Project Description

The aim of this project is to implement and use different learning algorithms to train linear predictors that solve the binary classification problem.

Specifically for our case with the given dataset, we need to predict the value of the label in column  $y$  based on the numerical features  $x_1$  through  $x_{10}$ .

As mentioned in the description of the project, I used the zero-one loss as a metric to evaluate the performance of the algorithms.

In the following chapter I describe how I analysed and pre-processed the data set to improve the performance of the modules.

In the following chapters I describe how I implemented each algorithm and how I chose the hyperparameters for them.

I have also analysed how varying the hyperparameters affects the validation and training errors, and explained these empirical results with the theoretical background provided in the lectures.

I implemented the following algorithms:

- Perceptron
- Pegasos
- Logistic Pegasos
- Feature expanded Perceptron (with 2nd degree polynomial expansion)
- Feature expanded Pegasos (with 2nd degree polynomial expansion)
- Feature expanded Pegasos with logistic loss (with 2nd degree polynomial expansion)
- Kernel Perceptron
- Kernel Pegasos

## 1.2 Project Structure

The project is divided into the following folders:

- datasets: contains the provided dataset
- models: a set of pre-trained models, created using the train subcommand and serialized with pickle python module
- src: the source code of the project, the entry point is main.py and can be used both to train the models from the dataset and to run them
- report: this folder contains the source for this report

## 1.3 Usage

The entry point to the project is the file **main.py**.

It can be called with the command line argument and provides two subcommands, *train* and *run*.

The first subcommand requires the name of the algorithm to be trained and stores a predictor in the path provided in the output argument, serialised using the Python pickle module. Example:

```
1 $ python src/main.py train perceptron models/perceptron.pkl
```

Results in the following output:

```
1 [Perceptron]
2 trained perceptron: [ 0.55604295  1.97486085 -2.58700382 -1.74490783  1.91766823 -3.898
3 -0.02419966  3.06371997  0.21648717 -0.79054099  1.
4 test error for perceptron: 0.326
```

The *run* subcommand takes a serialized model and then prints its training and test errors

```
1 $ python src/main.py run models/perceptron.pkl
```

With the following output:

```
1 training error for predictor saved at 'models/perceptron.pkl': 0.322625
2 test error for predictor saved at 'models/perceptron.pkl' : 0.326
```

There are other options available that are described using the '-help' option, they will be described in the next sections of this report as they come up.

## 2 Dataset analysis and preprocessing

### 2.1 Dataset description

The provided dataset contains 10000 points with 10 features named from  $x_1$  to  $x_{10}$  and a label column named  $y$ .

All the feature are floating point values and the dataset is well formed (in the sense that there are no missing values).

The label column contains values that are either  $-1$  or  $+1$ .

There isn't any duplicated data in the training set.

I collect the major statistics from each feature in the dataset: *mean*, *standard deviation*, *min* and *max*.

	x1	x2	x3	x4	x5
min	2.44342055e-03	-7.52493399e+00	9.85724553e+01	-7.07893888e+00	-9.99999717e-01
max	9.38422309e+00	8.30237476e+00	1.01260768e+02	-2.92150729e-06	9.99999998e-01
mean	1.59129826e+00	5.15879411e-01	9.98489361e+01	-1.50413876e+00	7.76447773e-02
std	1.32111881	2.05438485	0.71091203	1.13354878	0.70723419

	x6	x7	x8	x9	x10
min	-6.90697075e+00	-7.14075517e+00	-7.15188951e+00	-5.67739307e+01	-1.00000000e+00
max	8.76030588e+00	9.28726632e+00	6.21145227e+00	-5.42088897e+01	1.00000000e+00
mean	5.18228648e-02	9.75207134e-01	6.35194433e-01	5.19260973e-02	-5.54476783e+01
std	0.70471943	2.16212877	2.21259701	1.76955726	0.71004639

It's clear from the main features that the different features have different ranges and follow different probability distributions.

### 2.2 Feature Scaling

Because the dataset is already well formed and there are no missing values the first thing I do is scale the features.

This step should ensure that the values of the features are in a comparable range.

I tried two approaches: normalization and standardisation.

**Standardization** sets each feature to have a mean of 0 and a standard deviation of 1.

This is achieved by subtracting the feature mean from each value and dividing by its standard deviation:

$$x' = \frac{x - \mu}{\sigma}$$

(where  $\mu$  is the mean and  $\sigma$  is the standard deviation).

In the case of **normalization**, the features are rescaled in a fixed range between 0 and 1 in the following way:

$$x' = \frac{x - x_{min}}{x_{max} - x_{min}}$$

It's important to note that we should perform both approaches using a sound procedure, more specifically, we should avoid data leakage: When calculating the minimum and maximum of the feature, we should only consider the training set and scale the test set only according to these values, without deriving any information from it.

In the same way, we should calculate the mean and standard deviation for the standardization process.



Both of these approaches lead to important improvements and can be demonstrated both from an empirical point of view (see table 2.2) and from a theoretical perspective (explained in the chapter describing perceptron). It's also worth noting that I couldn't run the logistic regression without first rescaling the data, due to a float overflow in the exponentiation for the sigmoid function.

	None	Normalization	Standardization
Perceptron	0.507	0.482	0.326
Pegasos	0.294	0.2805	2807

Comparison of test errors between Perceptron and Pegasos with the three different scaling options.

Is possible to choose which scaling method use with the cli option *preprocess* and choose between *none*, *normalize* and *standardize*.

I used the standardization methods by default because is the one that perform better in the majority of the cases.

## 2.3 Outliers removal

Another approach I considered is removing the outliers from the dataset using the Z-score methods. I calculated the score  $Z = (x - \mu)/\sigma$  for each value where  $\mu$  is the mean and  $\sigma$  is the variance of the feature.

I then removed all the points with a Z-score greater or equals than 3 in absolute value.

I found (and removed) 265 outliers (recall that the dataset has size 10000).

I tried training non-kernelized Perceptron and Pegasos over the modified dataset but the results shows that the dataset is already sufficiently cleaned: in fact it affects the performance of the models in a minimum way with no significative changes, and even in some cases it is (even if only slightly) worsening

	with outliers	without outliers
Perceptron	0.326	0.326
Pegasos	0.2865	0.29
Feature expanded Perceptron	0.087	0.087
Feature expanded Pegasos	0.0555	0.0565

Comparison of test errors (using zero one loss) with and without outliers.

Note that when I refere to feature expansion I intend polynomial feature expansion of degree 2.

Is possible to enable the outliers removal using the cli flag *remove-outliers*.

## 2.4 Feature correlation

I sorted all the point according to each axis and plot on the other axis all the other features to spot eventual correlation and I observed that the feature  $x_2$  and  $x_5$  have a linear correlation (with a negative coefficient, see Fig. 1) as the feature 5 and 9 (with positive coefficient, see Fig. 2).

One possibility in this case during the preprocessing of the data is to remove the correlated features and leave only one of them to avoid redundancy of the data.

I don't follow this approach because there is a sensible noise in the correlation and removing some features can lead to also removing this noise that can encode important information for the model.

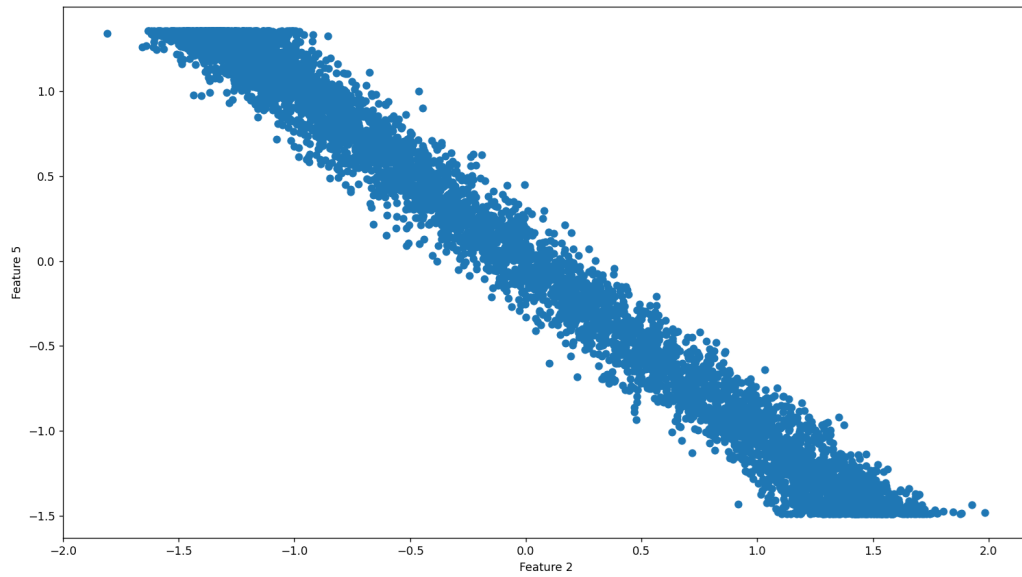


Figure 1: Here there is a clear linear correlation between feature  $x_2$  and  $x_5$ , but very noisy

## 2.5 Feature expansion

To being able to express non-homogeneous linear separators (hyperplane that don't pass through the origin) I add a constant feature of value 1 to each point in the dataset.

Let  $(\mathbf{x})$  be any point in the dataset and  $(\mathbf{w})$  be the linear separator, if we define  $x' = (\mathbf{x}, 1)$  we can define  $w' = (w, c)$ , in that way:

$$w'^T x' = (\mathbf{w}^T \mathbf{x} + c)$$

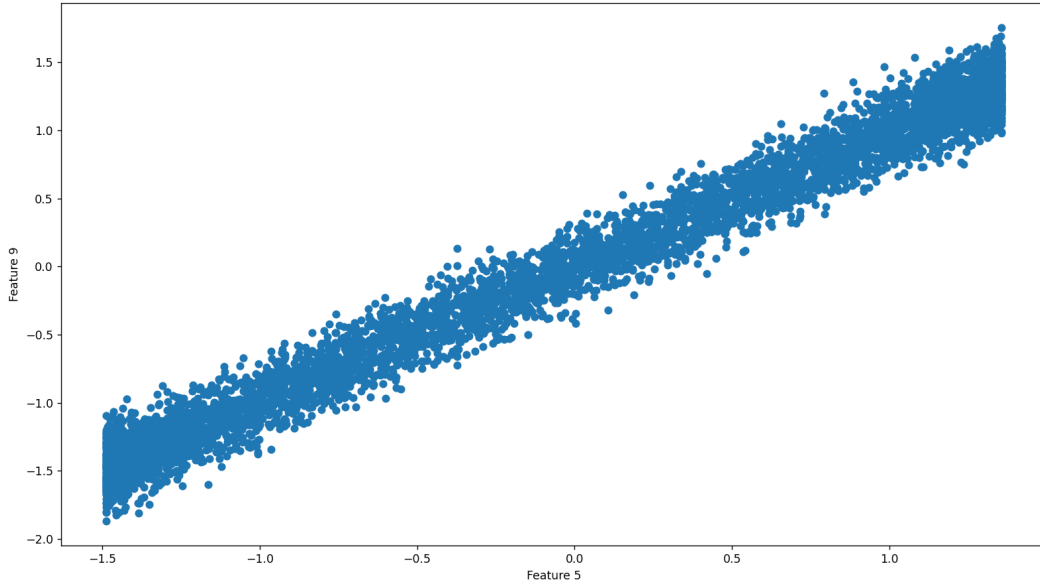


Figure 2: Another correlation is also present between feature  $x_5$  and  $x_9$  (with a negative coefficient)

### 3 Perceptron

The first algorithm that I implemented is the perceptron algorithm.

The perceptron algorithm is used to learn linear classifiers.

Linear classifiers are identified by an hyperplane that separate the input space into two halfspaces, one positive and one negative.

The positive halfspace is called so because the dot product with the normal vector that identify the hyperplane and any point in that halfspace is positive, similarly the negative halfspace has always negative dot products.

One important property of the Perceptron is the convergence (to the ERM) in a finite number of step if the dataset is linearly separable.

This properties is stated by the **Perceptron Convergence Theorem**:

*Let  $(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_m, y_m)$  be a linearly separable training set. Then the Perceptron algorithm returns a linear classifier with zero training error in a finite number of updates*

$$M \leq \left( \min_{\mathbf{u}: \gamma(\mathbf{u}) \geq 1} \|\mathbf{u}\|^2 \right) \left( \max_{t=1, \dots, m} \|\mathbf{x}_t\|^2 \right)$$

where  $\gamma(\mathbf{u})$  is the margin obtained by the linear separator  $\mathbf{u}$

Is possible to show also a bound for non linearly separable cases (That cannot have a 0 training error):

$$M \leq \sum_{t=1}^T h_t(\mathbf{u}) + (\|\mathbf{u}\|X)^2 + \|\mathbf{u}\|X \sqrt{\sum_{t=1}^T h_t(\mathbf{u})} \quad \text{for all } \mathbf{u} \in \mathbb{R}^d$$

This shows a bound on the number of mistakes made by the Perceptron algorithm on any data sequence of arbitrary length  $T$ .

$h_t(\mathbf{u})$  is the hinge loss for the  $t$ -th example, defined as  $h_t(u) = \max\{0, 1 - \mathbf{u}^T \mathbf{x}_t y_t\}$ .

Both the results show a linear dependence with the number of mistakes  $M$  and  $X^2$ , the radius of the smaller sphere that inscribes all the training points.

This also shows why, in our case, both *standardization* and *normalization* are so effective:

We are reducing the radius of this sphere and so having a tighter bound on the number of mistakes with the same number of rounds  $T$ .

### 3.1 Naive version

Here I present the pseudocode for the perceptron algorithm taken from the lecture notes [1].

---

**Algorithm 1:** The Perceptron algorithm

---

**Input:** Training set  $(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_m, y_m)$

```

1  $\mathbf{w} = (0, \dots, 0)$ 
2 while true do
3   for  $i = 1, \dots, m$  do                                     // (epoch)
4     if  $y_i \mathbf{w}^\top \mathbf{x}_i \leq 0$  then
5        $\mathbf{w} \leftarrow \mathbf{w} + y_i \mathbf{x}_i$                              // (update)
6     end
7   if no update in last epoch then
8     break
9 end
Output:  $\mathbf{w}$ 

```

---

My implementation slightly varies from the presented pseudocode: While the above will continue to run until convergence, if the training set is not linearly separable (as in this case) we will never converge and so the algorithm will never terminate.

To avoid this, I use an additional parameter 'max\_epoch' which limits the number of epochs the algorithm can run.

I choose to use a fixed value of 20 for this parameter both for the naive case and over the feature expanded dataset presented in the next section.

Training the perceptron algorithm with the preprocessing methodology described in the previous chapter (For all the algorithms I describe I trained it with the default command line options), I have obtained a training error of 0.322625 and a test error of 0.326.

The linear separator found by the perceptron algorithm has the following features:

(0.55604295, 1.97486085, -2.58700382, -1.74490783, 1.91766823, -3.89876104, -0.02419966, 3.06371997, 0.21648717, -0.79054099, 1) (Recall the features are 11 because we add a constant feature of 1 to the dataset to be able to express non-homogeneous linear hyperplane).

### 3.2 Feature Expansion of 2nd degree

I also trained the perceptron algorithm on a second degree polynomial feature expanded dataset.

In this version of the algorithm it is possible to express hyperplane in a high dimensional feature space, and this can also be interpreted as a polynomial curve (of second degree in this case) in the original space.

For this reason the training and test error of the resulting predictor are significantly better than the previous version.

Specifically I obtained a training error of 0.085375, and a test error of 0.087.

This, of course, came at the cost of significantly increasing the number of features in the datasets and also the computational cost of operations between vectors (such as sums and dot products). As we will see later, we can avoid this cost by using kernels, which allow us to compute the dot product of the feature expansion of two vectors without explicitly computing their expansion. This are the features obtained:

```
(1.87559817e+01 1.13401244e+00 7.68047788e+00 -1.32276315e+01
1.73791816e+01 -5.35283741e+00 1.04738036e+01 6.65533633e+01
3.38354273e+01 5.20890557e+00 -1.40000000e+01 -2.37767269e+00
-1.15700558e+01 3.58887386e+00 -7.70998879e+00 -3.37225323e+00
8.62691037e+00 5.29455094e+00 7.16899813e+01 -2.75843723e+00
-1.44740075e+01 1.87559817e+01 1.58098829e+00 -2.16046998e+01
4.71738784e+00 -9.24568568e-01 2.31787163e+00 -6.88714236e-01
6.36558311e+00 2.24640419e+02 -3.01817078e+01 1.13401244e+00
-1.01581267e+01 -5.54650087e+00 -1.37119017e+01 5.92267047e+00
6.13176127e+00 -1.91940656e+01 1.20197686e+01 -3.78679497e+00
7.68047788e+00 -4.32369339e+00 -2.30745221e-01 -8.05450142e+00
1.43389046e+00 -7.36485907e+01 4.14731220e+00 9.17178493e-01
-1.32276315e+01 9.31783828e-01 -2.90129736e+01 1.10054038e+01
9.29221437e+00 5.16458889e-02 5.24165775e+00 1.73791816e+01
-3.39061351e+00 1.79056771e+01 1.68705706e+01 1.02044524e+01
-7.88104212e+00 -5.35283741e+00 4.04206126e+00 1.28786306e+01
1.01040343e+01 9.69582051e+00 1.04738036e+01 1.03988287e+01
2.78928092e+00 -2.37491314e+01 6.65533633e+01 8.51087821e-01
6.32907160e+00 3.38354273e+01 -2.72097857e+00 5.20890557e+00
-1.40000000e+01)
```

## 4 Support Vector Machine

The Support Vector Machine (SVM) algorithm learn linear classifiers, finding a linear classifier that is the **maximum margin separator hyperplane** and so achieve the maximum margin from all the point in the training set.

Given a linearly separable training set  $S = \{(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_n, y_n)\} \in \mathbb{R}^d \times \{-1, 1\}$  it's possible to find this hyperplane solving the following convex optimization problem with linear constraints.

$$\begin{aligned} \min_{\mathbf{w} \in \mathbb{R}^d} \quad & \frac{1}{2} \|\mathbf{w}\|^2 \\ \text{s.t.} \quad & y_t \mathbf{w}^\top \mathbf{x}_t \geq 1 \text{ for } t = 1, \dots, m \end{aligned}$$

In case the training data is not separable, we try to minimise both how much of each constraint is violated and the margin of the separator.

We express this as another convex problem using the slack variables  $\xi_t$  and a regularisation coefficient  $\lambda$ . If the regularisation coefficient is large, the algorithms will generate a predictor that allows more classification error in the training set.

Conversely, if  $\lambda$  is small, we try to minimise the classification error we have made.

Usually for  $\lambda$  too small, we try to minimise the misclassification, so the training error is small and we are likely to overfit.

Instead, for choices of  $\lambda$  too large, we have a high training error, and if the test error is also high, we will underfit.

In the next subsection 4.1.2, I describe how I choose the hyperparameter of the regularisation coefficient and how the test and training errors vary when I change it more precisely.

$$\begin{aligned} \min_{(\mathbf{w}, \boldsymbol{\xi}) \in \mathbb{R}^{d+m}} \quad & \frac{\lambda}{2} \|\mathbf{w}\|^2 + \frac{1}{m} \sum_{t=1}^m \xi_t \\ \text{s.t.} \quad & y_t \mathbf{w}^\top \mathbf{x}_t \geq 1 - \xi_t & t = 1, \dots, m \\ & \xi_t \geq 0 \text{ for } & t = 1, \dots, m \end{aligned}$$

Now, fix  $\mathbf{w} \in \mathbb{R}^d$ , we can see  $\xi_t = [1 - y_t \mathbf{w}^\top \mathbf{x}_t]_+$  which is the hinge loss  $h_t(\mathbf{w})$ .

The SVM problem can be rewritten as

$$\min_{\mathbf{w} \in \mathbb{R}^d} \quad \frac{\lambda}{2} \|\mathbf{w}\|^2 + \frac{1}{m} \sum_{t=1}^m h_t(\mathbf{w})$$

The optimization problem that describe the Support Vector Machine is optimized using the Pegasos algorithm.

The Pegasos algorithm is a variant of the Stochastic Gradient Descent algorithm, where at each step a point (or a set of points in the mini-batch variant) is sampled randomly from the training set and the current predictor is updated with the negative gradient of the loss of that training example weighted by a learning rate factor  $\eta_t$ .

In case of Pegasos the learning rate factor  $\eta_t$  is choose at each step as  $\frac{1}{\lambda t}$ .

I implement the Pegasos algorithm using the standard variant with the *hinge* loss, and with the *logistic* loss (Described in the Logistic regression subsection 4.2).

Both this functions are convex upper bounds of the zero-one loss, and the  $\lambda$  regularization parameter allow to have a  $\lambda$ -strongly convex function to minimize with the gradient descent.

## 4.1 Naive

As I previously said I implemented Pegasos using two surrogate losses: hinge loss and logistic loss. Now I describe the implementation with hinge loss, that differs from the logistic one only by the update step.

Recall that the hinge loss is defined as  $l(y, \hat{y}) = \max\{0, 1 - y_t \hat{y}\}$

Given  $Z_t = (X_t, Y_t)$  a random sample from the training set, the update rule for Pegasos is:

$$\mathbf{w}_{t+1} = \mathbf{w}_t - \eta_t \nabla \ell_{Z_t}(\mathbf{w}_t)$$

Let be  $s_t$  the realization for the random variable  $Z_t$

Where  $\ell_{s_t}(\mathbf{w}) = [1 - y_{s_t} \mathbf{w}^T x_{s_t}]_+ + \frac{\lambda}{2} \|\mathbf{w}\|^2$  so

$$\nabla \ell_{s_t}(\mathbf{w}) = -y_{s_t} x_{s_t} \mathbb{I}\{h_{s_t}(\mathbf{w}) > 0\} + \lambda \mathbf{w}$$

Let  $\mathbf{v}_t = y_t x_t \mathbb{I}\{h_t(\mathbf{w}_t) > 0\}$  and choosing  $\eta_t = \frac{1}{\lambda t}$  we have

$$\mathbf{w}_{t+1} = \mathbf{w}_t \left(1 - \frac{1}{t}\right) + \frac{1}{\lambda t} \mathbf{v}_t$$

### 4.1.1 Implementation details

I adapted this version of the algorithm from *Pegasos: Primal Estimated sub-GrAdient SOLver for SVM*[\[2\]](#)

---

#### Algorithm 2: Pegasos Algorithm

---

```

Input:  $S, \lambda, T$ 
1 for  $t = 1, 2, \dots, T$  do
2   Choose  $i_t$  uniformly at random.
3   Set  $\eta_t = \frac{1}{\lambda t}$ 
4   if  $y_{it} \mathbf{w}_t^T x_{i_t} < 1$  then
5     Set  $\mathbf{w}_{t+1} \leftarrow (1 - \eta_t \lambda) \mathbf{w}_t + \eta_t y_{it} x_{i_t}$ 
6   end
7   else
8     Set  $\mathbf{w}_{t+1} \leftarrow (1 - \eta_t \lambda) \mathbf{w}_t$ 
9   end
10 end
11 Output  $\mathbf{w}_{T+1}$ 

```

---

The name of the variables are adapted to be consistent with the pseudo code reported.

Between the code presented in the lecture and this one presented in the paper there are some differences:

- The gradient descent update is written in a slightly different way using a conditional statement instead of the classical indicator function, I choose to remain consistent also with this stylistic choice.

- Instead of return the average of all the weight vector calculated at each step, the paper returns only the last one.  
The authors indicate that they note an improvement in performance by returning the last vector instead of the average.  
I accept also this variation.
- The Pegasos pseudocode also describes an optional projection step to clamp the magnitude of the linear predictor, but I don't incorporate it.

Two other approaches that the author suggest but I don't experiment are:

- The author also provides a **mini-batch** version of the Pegasos algorithm.  
The chosen batch is a set of  $k$  random samples from the training set, and the linear predictor is updated with the average loss gradient of all incorrectly predicted samples in the batch.  
The batch size  $k$  can be chosen in advance or treated as another hyperparameter.
- **sampling without replacement**: a random permutation of the training set is chosen and the updates are performed in order on the new sequence of data. In this way, a training point is sampled only once in an epoch. After each epoch, we can choose whether to resume sampling the data sequentially according to the same permutation, or to create a new permutation and sample according to that new order.  
Although the authors report that this approach gives better results than uniform sampling as I did, I haven't experimented with this variant of the algorithm.

#### 4.1.2 Hyperparameter tuning

Unlike the perceptron, the Pegasos algorithm had a hyperparameter to choose from: The regularisation coefficient  $\lambda$ .

To choose the best hyperparameter for this algorithm (and the others that I will implement), I choose the grid search method. I divide the training set into 2 different subsets:  $S_{train}$  and  $S_{dev}$ .

The validation set ( $S_{dev}$ ) is used as a surrogate test set to get an estimate of the risk.

I choose a finite subset of the possible hyperparameter values ( $\Theta_0 \subseteq \Theta$ ) and for each of them I create a predictor  $h_\theta$  which is trained with the chosen hyperparameter on  $S_{train}$ .

The risk is then estimated using the validation error on each predictor, and the one with the lower risk is chosen.

The implementation of the grid search in my codebase is in the function 'grid\_search', I used python know arguments to pass a dictionary where the key is the name of the hyperparameter (the same as used as an argument in the training algorithm) and the value is an iterator containing the subset of possible values ( $\Theta_0$ ).

In this case, I choose the set  $\Theta_0 = \{0.0001, 0.001, 0.01, 0.1, 1, 10, 100\}$ .

$\lambda$	$S_{val}$	$S_{train}$
0.0001	0.3725	0.3521666
0.001	0.2925	0.281333
0.01	0.273	0.27
<b>0.1</b>	<b>0.267</b>	<b>0.2661666</b>
1	0.273	0.269333
10	0.279	0.2775
100	0.2805	0.276333

The values  $S_{val}$  and  $S_{train}$  while  $\lambda$  changes.



Looking at the table we can deduce for which choice of  $\lambda$  we underfit and overfit and in our case the sweet spot is the value for  $\lambda = 0.1$ .

The test error for the best predictor is 0.2935.

We can also see that for smaller value of lambda we have an high test error and model is overfitting while for large value are likely to underfit.

The weight for the Pegasos algorithm are the following:

0.2080856 -0.0066276 0.07067274 -0.26477777 0.24795273 -0.06704699  
0.28821758 0.65130349 0.16619769 -0.06650096 -0.0724

## 4.2 Logistic regression

Logistic regression aim to learn the function  $\eta(\mathbf{x}) = \mathbb{P}(Y = +1 \mid \mathbf{X} = \mathbf{x})$ .

The implementation of logistic regression differs from the standard Pegasos because it uses the logistic loss as surrogate loss and not hinge.

The logistic loss is defines ad follows:

$$\ell(y, \hat{y}) = \log_2(1 + e^{-y\hat{y}}).$$

So the gradient descent update becomes:

$$\mathbf{w}_{t+1} = \mathbf{w}_t + \eta_t \sigma(-y_t \mathbf{w}^T \mathbf{x}_t) y_t \mathbf{x}_t$$

I choose the regularization parameter as described in the previous subsection and obtained the following results.

$\lambda$	$S_{val}$	$S_{train}$
0.1	0.278	0.2718333
<b>1</b>	<b>0.2735</b>	<b>0.274333</b>
10	0.28325	0.27816
100	0.2805	0.278166

The values  $S_{val}$  and  $S_{train}$  while  $\lambda$  changes for the logistic loss.

I have choosen the hyperparameter  $\lambda = 1$  and obtained a test error of: 0.292.

## 4.3 Feature Expansion

I have also trained the two previous algorithms over the polynomial feature expansion of second degree of the whole dataset.

Like in the case of perceptron, since we are able to express a separator in an high dimension we obtain better results than the naive version (at the cost of an increasing number of features in the dataset).

### 4.3.1 Pegasos

This are the following validation and development errors training the Pegasos algorithm the same set of values choosen for the hyperparameter  $\lambda$  as in the naive version.

$\lambda$	$S_{val}$	$S_{train}$
<b>0.001</b>	<b>0.045</b>	<b>0.039833</b>
0.01	0.06	0.050166
0.1	0.0955	0.08833
1	0.1725	0.1525
10	0.219	0.211
100	0.256	0.246
1000	0.2635	0.25233

The test error obtained training the predictor on the whole training set with the regularization coefficient of 0.001 is 0.053 with a training error of 0.044125.

#### 4.3.2 Logistic Regression

$\lambda$	$S_{val}$	$S_{train}$
<b>0.1</b>	<b>0.11</b>	<b>0.1035</b>
1	0.1675	0.15
10	0.197	0.188
100	0.2595	0.2485
1000	0.2505	0.2385

The test error obtained for the feature expanded logistic regression (with  $\lambda = 0.001$ ) is 0.1125 with a training error of 0.1055.

Note that for both versions of logistic regression, not all of the  $\lambda$  parameters used for the hinge loss SVM in the grid search are used in this case.

This is because for values that are too small, I get an overflow error when exponentiating in the sigmoid function.

These values, for which the gradient update calculation is not feasible, are simply discarded.

## 5 Kernel

In the previous section I have showed the benefit of training the algorithms in an high-dimensional space.

This came with the cost of increasing sensibly the number of the feature even in case of a simple polynomial feature expansion of degree 2.

We can have the performance improvement obtained with feature expansion without increasing the dimensionality of the dataset using the kernel trick.

Instead of computing the feature expansion map on evry point and then computing the dot product with the expanded linear predictor we can use a kernel  $K$  that calculate the dot product of two expanded vector without explicitly calculate the expansion.

Let the feature expansion be  $\phi(x)$ , a kernel  $K$  for the feature expansion is defined as

$$K(x, x') = \phi(x)^T \phi(x') \quad \text{for all } x, x' \in R$$

The two kind of kernel used in the project are the **Polynomial kernel** and the **Gaussian kernel**.

The *polynomial kernel* of degree  $n$  is defined as

$$K_n(x, x') = (1 + x^T x')^n$$

The *gaussian kernel* of parameter  $\gamma$  is defined as

$$K_\gamma(x, x') = \exp(-\frac{1}{2\gamma} \|x - x'\|^2)$$

Note that both the kernels accept a parameter (a degree in the polynomial case and gamma for gaussian one) this becomes another hyperparameter of the learning algorithm and it's choosen with the same hyperparameter tuning procedure described in the previous chapters.

Each kernel induce a linear space defined as a set of linear combination of functions  $K(x, \cdot)$ .

$$\mathcal{H}_K \equiv \left\{ \sum_{i=1}^N \alpha_i K(x_i, \cdot) : x_1, \dots, x_N \in \mathcal{X}, \alpha_1, \dots, \alpha_N \in \mathbb{R}, N \in \mathbb{N} \right\}$$

This spaces are called Reproducible Kernel Hilbert Space (RHKS).

A linear predictor trained in these linear spaces induced by kernels can overfit; in the case of polynomial kernels, we can choose a degree that is too high and obtain a linear separator that is also rapresentable by a polynomial curve of degree  $n$  in the original space, which is more likely to separate the training data and obtain a small training error.

For the Gaussian, we should know that the parameter  $\gamma$  is the width of a Gaussian centered on the training point.

If  $\gamma$  is too small, the training error will be small and we will overfit because the Gaussians of different training points almost never overlap and the algorithm behaves like K nearest neighbours with  $K =$ .

### 5.1 Implementation details

I used two classes with the '`__call__`' python magic method to implement the kernels functions.

Both takes two parameter  $X$  and  $X2$ .

They can take two vectors and the kernel behave as mathematicly defined or the parameter can be 2 matrices of size  $m \times d$  and  $n \times d$ .

In the second case a returned matrix  $\mathbf{K}$  ( $m \times n$ ) is defined as  $\mathbf{K}_{i,j} = K(X_i, X2_j)$ .

This is particularly useful when calculating the kernel over the whole dataset, to avoid Python for loops and delegate the calculations to the optimised implementation of the numpy primitives.

Following the implementation of the call method for the two Kernel classes.  
Implementation of the polynomial kernel over  $X$  and  $X2$ .

```

1 def __call__(self, X: np.ndarray, X2: np.ndarray):
2     if X2.ndim == 1:
3         return np.power(np.dot(X, X2) + 1, self.degree)
4     elif X2.ndim == 2:
5         return np.power(np.dot(X, X2.T) + 1, self.degree)

```

Implementation of the gaussian kernel over  $X$  and  $X2$  with the parameter gamma  $\gamma$ .

```

1 def __call__(self, X: np.ndarray, X2: np.ndarray):
2     if X2.ndim == 1:
3         dist = np.linalg.norm(X - X2, 2, axis=1)
4     elif X2.ndim == 2:
5         dist = np.linalg.norm(X[:, np.newaxis, :] - X2[np.newaxis, :, :], axis=2)
6     return np.exp(-dist / self.gamma)

```

## 5.2 Kernelized Perceptron

The first algorithm I implement using the kernel trick is the perceptron.  
Following I provide a pseudocode taken from the lecture notes about this algorithm.

---

### Algorithm 3: Kernel Perceptron

---

```

1  $S \leftarrow \emptyset$ 
2 for  $t = 1, 2, \dots$  do
3     Get next example  $(\mathbf{x}_t, y_t)$ 
4     Compute  $\hat{y}_t = \text{sgn}(\sum_{s \in S} y_s K(\mathbf{x}_s, \mathbf{x}_t))$ 
5     if  $\hat{y}_t \neq y_t$  then
6          $S \leftarrow S \cup \{t\}$ 
7     end
8 end

```

---

Looking at the implementation, I have implemented the kernel perceptron algorithm with an array  $\alpha$  of integer weights for each point of the training set instead of using a set  $S$ , this may seem different from the pseudocode presented, but we can interpret  $\alpha$  as representing a multi-set and have an equivalent algorithm.

This choice, taken from the pseudo code of the kernelized perceptron in the Pegasos paper [2], is adopted because we can express the summation more simply and efficiently thanks to the numpy primitive such as 'np.dot'.

With the kernel version we have a new hyperparameter: The type of kernel used with its parameter (the degree for polynomials and the  $\gamma$  for Gaussians).

We choose the best hyperparameter, as described in the previous chapters.

Among the possible values, we choose 1, 2, 3, 4 for the possible degrees of the polynomial kernel, and 0.01, 0.1, 1 and 10 for the possible values of  $\gamma$ .

We can make some considerations about the data in the table 8 based on the previous theoretical section of this chapter: As we already explained for smaller values of  $\gamma$  in the Gaussian kernel the training error is 0 and the algorithm overfits with this parameters.

Polynomial Kernel $n$	Gaussian Kernel $\gamma$	$S_{val}$	$S_{train}$
1		0.339	0.3251666
2		0.0685	0.06633
<b>3</b>		<b>0.0585</b>	<b>0.048</b>
4		0.06	0.031666
	0.01	0.2075	0.0
	0.1	0.1805	0.0
	1	0.0685	0.0015
	10	0.072	0.01366

It's also possible to know in which cases for the polynomial kernel we are overfitting, observe that in the case of  $n = 4$  we have a value  $S_{val}$  significantly greater than  $S_{train}$ .

In contrast, for  $n < 3$  we also have both a high  $S_{val}$  and a high  $S_{train}$ , indicating underfitting.

The polynomial kernel with  $n = 3$  is the best one even if the  $S_{val}$  higher than  $S_{train}$  indicates a slight overfitting.

After choosing this hyperparameter we obtain a training error of 0.034125 and a test error of 0.0495.

### 5.3 Kernelized Pegasos

---

#### Algorithm 4: Pegasos Algorithm

---

```

1 I implement the kernelized version of the Pegasos algorithm following the pseudocode from the
  paper, reported here: Input:  $S, \lambda, T$ 
2 Initialize: Set  $\alpha_1 = 0$ 
3 for  $t = 1, 2, \dots, T$  do
4   Choose  $i_t \in 0, \dots, |S|$  uniformly at random.
5   For all  $j \neq i_t$ , set  $\alpha_{t+1}[j] = \alpha_t[j]$ 
6   if  $y_{i_t} \frac{1}{\lambda t} \sum_j \alpha_t[j] y_{i_t} K(\mathbf{x}_{i_t}, \mathbf{x}_j) < 1$  then
7     Set  $\alpha_{t+1}[i_t] = \alpha_t[i_t] + 1$ 
8   end
9   else
10    Set  $\alpha_{t+1}[i_t] = \alpha_t[i_t]$ 
11  end
12 end
13 Output  $\alpha_{T+1}$ 

```

---

As in the previous algorithm, I fixed a number of rounds  $T = 100000$  and tuned the algorithm over the hyperparameters  $\lambda$  (the regularization coefficient) and a kernel that can be either polynomial of degree 1, 2, 3 or 4, or gaussian with a  $\gamma$  parameter of one between 0.01, 0.1, 1, 10.

The best hyperparameters obtained by the grid search procedure are a regularization coefficient  $\lambda = 0.1$  and a polynomial kernel of degree 3.

Note that the best performing kernel is the same both for *kernelized perceptron* and for *Pegasos*.

We obtain a training error on the whole data set of 0.035875 and a test error of 0.043.

We can see from the test error that this is the best performing method implemented.

This is also because we are using a 3rd degree polynomial kernel and we are training a linear predictor in a higher dimensional space than the one I obtained with the 2nd degree polynomial expansion.

In the table 13 we can also see for which values of  $\lambda$  and kernel degree the predictor overfits, fixing the other hyperparameter.

Fixing the regularization coefficient  $\lambda = 0.1$  we can experimentally confirm that for high degrees of the kernel (4) the predictor overfits and has a high validation error and for smaller values (1 or 2) it

$\lambda$	Polynomial Kernel n	Gaussian Kernel $\gamma$	$S_{val}$	$S_{train}$
0.001	1		0.275	0.2738333333333333
0.001	2		0.0605	0.055
0.001	3		0.053	0.045
0.001	4		0.057	0.026166666666666668
0.001		0.01	0.1675	0.0
0.001		0.1	0.1465	0.0
0.001		1	0.114	0.06633333333333333
0.001		10	0.157	0.13683333333333333
0.01	1		0.2755	0.26916666666666667
0.01	2		0.055	0.04833333333333333
0.01	3		0.049	0.037166666666666667
0.01	4		0.051	0.025
0.01		0.01	0.167	0.0
0.01		0.1	0.15	0.0
0.01		1	0.2275	0.21266666666666667
0.01		10	0.2495	0.246
0.1	1		0.2645	0.26316666666666666
0.1	2		0.084	0.07616666666666666
<b>0.1</b>	<b>3</b>		<b>0.044</b>	<b>0.043</b>
0.1	4		0.05	0.023166666666666665
0.1		0.01	0.166	0.0
0.1		0.1	0.1465	0.0
0.1		1	0.2385	0.2255
0.1		10	0.2545	0.257
1	1		0.2725	0.26683333333333333
1	2		0.144	0.131
1	3		0.0825	0.072
1	4		0.0535	0.029
1		0.01	0.167	0.0
1		0.1	0.1445	0.0
1		1	0.2415	0.2245
1		10	0.26	0.25883333333333336
10	1		0.285	0.27966666666666667
10	2		0.193	0.17766666666666667
10	3		0.143	0.13383333333333333
10	4		0.0705	0.05433333333333333
10		0.01	0.1675	0.0
10		0.1	0.1445	0.0
10		1	0.2365	0.225
10		10	0.274	0.265
100	1		0.2805	0.27616666666666667
100	2		0.2015	0.19333333333333333
100	3		0.2185	0.21433333333333332
100	4		0.111	0.10033333333333333
100		0.01	0.169	0.0
100		0.1	0.144	0.0
100		1	0.2405	0.223
100		10	0.2845	0.2813333333333333

underfits.

Conversely, by fixing the chosen kernel and varying  $\lambda$ , we can see that  $S_{val}$  is slightly higher for values of the regularization coefficient  $< 0.1$  and we are probably overfitting, while for higher values the validation error is much higher and this can signal underfitting.

## 5.4 A note on the complexity of the algorithms

The computational cost of the kernelized versions of the algorithms differs significantly from the naive (or feature-extended) ones.

In the case of the perceptron, the number of samples considered is the number of epochs times the size of the training set (in non-separable cases like the one considered).

While for Pegasos the number of rounds chosen by the user is fixed to 100000 in our case.

The computational complexity for both is linear in the number of samples  $T$  considered ( $O(T)$ ).

For the kernelized versions, instead, for each sample considered, we should compute a linear combination of the learned *alphas* and the kernel between the sample and every other point in the training set.

This means that the computational complexity increases from  $O(T)$  to  $O(T * m)$ , where  $m$  is the size of the training set.

The variant of the kernelized algorithm described in the Pegasos pseudocode is more effective for a smaller number of rounds  $T \leq m$  (since in the worst case it will be  $O(T^2)$ ), but to get better results I trained the algorithm with a larger value for  $T$ .

## 6 Conclusion

I started this report by describing how to use the project and then moved on the preprocessing the data: which methods are used and based on which observation.

Then I implemented the Perceptron, Pegasos and logistic regression algorithms, also with second degree feature expansion and kernelized variants (the latter only for Perceptron and Pegasos).

I have used the lecture notes to introduce the theoretical background for understanding how these algorithms work at the beginning of each chapter, then explained the experimental data based on that.

I have discussed the implementation details and how I choose the hyperparameters for each algorithm.

I have also identified for which hyperparameters the algorithm overfits and underfits, explained using the theoretical prospective.

The best performing algorithm in all variants is the Pegasos algorithm.

We also observed that the feature-extended version performs better than the naive one, but the best results come from implementing the algorithm in combination with kernels.

More specifically, the third degree polynomial kernel is the best performing kernel for both Perceptron and Pegasos, intuitively because the algorithm is able to express a higher degree polynomial curve in the original space.

The feature expansion is of second degree, so it has less expressive power than the chosen kernel.

But increasing the degree of the polynomial curve doesn't necessarily mean finding a better predictor: we also see that for  $n = 4$  the curve overfits the data, giving a low training error but a high test error.

I also observed Gaussian kernels are not effective as the polynomial one because they tend to overfit more easily.



## References

- [1] Cesa-Bianchi Nicoló *Statistical Methods for Machine Learning*. [https://cesa-bianchi.di.unimi.it/MSA/index\\_23-24.html](https://cesa-bianchi.di.unimi.it/MSA/index_23-24.html)
- [2] Shai Shalev-Shwartz, Yoram Singer, Nathan Srebro, Andrew Cotter *Pegasos: Primal Estimated sub-GrAdient Solver for SVM*. <https://home.ttic.edu/~nati/Publications/PegasosMPB.pdf>