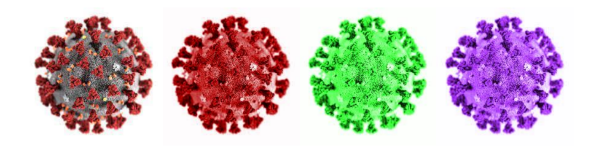
HW3: Regression

In this final assignment, we will try to predict a continuous label using our dataset.

Good Luck!



Instructions

Submission

- Submit by Wednesday, 25.01.2023, 23:59. No late submissions will be accepted.
- Submissions in pairs only in the webcourse.

Your code

- Should be clearly and <u>briefly</u> documented.
- Variables/classes/functions should have meaningful names.
- May be partially reviewed and graded.

Final report

- Should be written in a word processor (Office Word, Google docs, etc.).
 - Should not contain the code itself.
 - Do not submit jupyter notebooks as PDFs.
- o Can be in Hebrew, English, or both.
- You are primarily assessed based on your written report.
- Answer the questions in this instruction file according to their numbering.
- Add concise explanations, figures (outputs of your code), tables, etc.
- You are evaluated for your answers but also for clarity, readability, and aesthetics.
- Tables should include feature names and suitable titles.
- **Plots** should have suitable titles, axis labels, legends, and grid lines (when applicable).

Submit a zip file containing (please use hyphens, not underscores):

- The report PDF file with all your answers, named id1-id2.pdf.
- Your code (choose the relevant options for you):
 - Working with jupyter: a notebook with your code, named id1-id2.ipynb.
 - Working with a "traditional" IDE: one clear main script, named id1-id2.py, and any additional files required for running the main script.
- A completed *LinearRegressor*.py file with your implementation.

Preliminary: Updated Data Loading

Task: Follow the procedure below.

a. Start by **loading** the **new** raw data in HW3_data.csv.

The dataset is almost identical to the one from the previous assignments, but we <u>deleted</u> the binary targets and revealed the continuous contamination level target variable.

b. Make sure the data is **partitioned** correctly to train and test, according to the instructions in the previous assignments.

The train-test partitions **must** be **identical** to the ones you used in HW1 and HW2.

c. Apply the **preprocessing** procedure from the previous assignments (including the normalization steps) on both the training and test sets (but be careful to only use the training set when computing statistics for normalization etc.).

Note: in Lecture 08 we explain why some preprocessing steps (e.g., normalization), should be applied to the validation folds according to statistics computed on the train fold. Here, for simplicity <u>only</u>, you compute these statistics according to all the training samples (before splitting it to train and validation folds).

d. Important: do not use the test set before Section 6.

Throughout this assignment, we mainly focus on regressing the <u>new</u> continuous contamination_level variable.

Section 1: Linear regression implementation

Before we start using sklearn's modules, we want to thoroughly understand the optimization problem we use to solve linear regression – the least squares problem.

We will now implement a non-homogeneous linear regressor with the stochastic gradient descent (SGD) algorithm. The MSE loss definition for such a regressor is:

$$L(\underline{\boldsymbol{w}},b) = \frac{1}{m} \sum_{i=1}^{m} (\underline{\boldsymbol{w}}^{\top} \underline{\boldsymbol{x}}_{i} + b - y_{i})^{2} = \frac{1}{m} \|\boldsymbol{X}\underline{\boldsymbol{w}} + \underline{\boldsymbol{1}}_{m} \cdot b - \underline{\boldsymbol{y}}\|_{2}^{2},$$

where $\underline{\mathbf{1}}_m \in \mathbb{R}^m$ is a vector of ones and $b \in \mathbb{R}$ is a scalar. Using $\underline{\mathbf{1}}_m \cdot b$ allows the same b to be used for all rows (corresponding to all samples).

The gradient with respect to $\underline{w} \in \mathbb{R}^d$ was presented in Tutorial 09:

$$\mathbb{R}^d \ni \nabla_{\underline{w}} L(\underline{w}, b) = \frac{1}{m} 2X^{\top} (X\underline{w} + \underline{1}_m \cdot b - \underline{y})$$

(Q1) In your report, derive (פַתחו) the analytical partial derivative $\frac{\partial}{\partial b}L(\underline{w},b) \in \mathbb{R}$.

To implement our regressor we will inherit from sklearn's BaseEstimator class (like in HW2) for compatibility with scikit-learn API. We will also inherit from RegressorMixin.

This is the only section where we will perform validation without using cross validation.

- For this section only, split your training set into a (new) training subset (80%) and a validation subset (20%).
- Copy the given LinearRegressor module into your notebook / project.
- Complete the following methods
 - o LinearRegressor.loss method so that it computes the objective MSE loss $L(\underline{w},b)$ on a given dataset. Avoid using for loops.
 - o LinearRegressor.gradient method so as to compute the analytic gradients $\nabla_{\underline{w}} L(\underline{w}, b)$ and $\frac{\partial}{\partial b} L$. Avoid using for loops.

Tip: When possible, prefer vector operations (e.g., np.sum, np.linalg.norm).

- o LinearRegressor.fit_with_logs method in the module so as to compute the gradients of the current <u>batch</u> and update the parameters accordingly.
- o LinearRegressor.predict method so as to compute the model prediction.

Like in HW2, you will now verify the correctness of your implementation for the loss and its gradient by plotting the residuals $\|\underbrace{\nabla_{\underline{w}} L(\underline{w}, b)}_{\text{analytic}} - \underbrace{u_{\delta_w}(\underline{w}, b)}_{\text{numeric}}\|_2$ and $\|\underbrace{\frac{\partial}{\partial b} L(\underline{w}, b)}_{\text{analytic}} - \underbrace{u_{\delta_b}(\underline{w}, b)}_{\text{numeric}}\|_2$ as a function of δ_w , δ_b (respectively; over many repeats).

<u>Task</u>: Copy the functions from the given $verify_gradients.py$ into your notebook / project.

Read and understand these functions but do not edit them.

(Q2) Using your preprocessed (and normalized) dataset and contamination_level as our target, generate a plot that compares the numerical gradients to the analytical ones.
Do this by running the following command:

Important: X train should hold the features of your <u>normalized</u> training subset.

y train should hold the contamination level subset training labels.

Attach the plots to your report. No need to discuss these plots, but <u>make sure</u> that they make sense.

<u>Task</u>: Copy the function given in $test_lr.py$ into your notebook / project. Read and understand the function but do not edit it.

(Q3) We now want to evaluate the effects of different learning rates on the learning procedure. Run the following command that plots a graph of the training and validation losses as a function of the iteration number for different learning rates.

Important: x_{val} should hold the features of your (preprocessed) validation subset. y_{val} should hold the contamination_level subset validation labels.

Note: If your model did not converge with any learning rate, you are allowed to alter the lr list variable (but explain this in your report).

This part should also help you verify your implementation (loss should decay).

Attach the plots to your report, briefly discuss the results and <u>justify</u> the demonstrated behaviors. Which learning rate is "optimal" (briefly explain)?

For the best learning rate, does it make sense to increase the number of gradient steps (instead of the default 1500 steps)? Explain.

Now that we have experienced solving and tuning the least squares problem, we are ready for the rest of the assignment... and save the world from Contamination!

Section 2: Evaluation and Baseline

Our general goal is to minimize the generalization MSE, that is $\mathbb{E}_{(x,y)\sim D}[(h(x)-y)^2]$.

As we have learned, in practice, we instead minimize the empirical error $\frac{1}{m}\sum_{i=1}^{m}(h(x_i)-y_i)^2$ on a training set, and tune hyperparameters using a validation set.

In the rest of this assignment, we use the k-fold cross-validation method for better estimating the generalization error, thus improving the tuning procedure. We will use k = 5 folds.

Similar to HW2, we use sklearn to perform <u>cross-validation</u> on the (whole) training set to evaluate the performance of models. As explained earlier, the metric we use for regression is the MSE (using cross validate, set scoring='neg mean squared error').

Simplest baseline

We now train a simple <u>DummyRegressor</u> that always predicts the average contamination_level of the <u>training</u> set. We will use this regressor throughout the assignment as a baseline to which we will compare the performance of our learned regressors.

(Q4) Create a <u>DummyRegressor</u>. Evaluate its performance using cross-validation. In your report, fill in the cross-validated errors of the regressor.

Model	Section	Train MSE	Valid MSE	
		Cross validated		
Dummy	2			

<u>Task</u>: Retrain the dummy regressor on the entire training set (= all its samples) and save it for future use (<u>Sec 6</u>).

Basic hyperparameter tuning

A quick reminder of the tuning process for hyperparameters:

The repeated tuning process (for a single parameter) should include:

- Determining the tested values of the tuned hyperparameter (see <u>numpy.logspace</u>).
 You need to <u>choose</u> suitable values by yourself that will help you optimize the validation error.
- ii. For each value, evaluating a suitable regressor using cross validation (k = 5).
- iii. Plotting (cross-validated) train and validation errors as a function of the hyperparameter.Consider using <u>semilogx</u> or <u>loglog</u> plots.

Also plot a constant line with the validation error of the dummy regressor.

- iv. Reporting the value that yields the optimal validation error and its respective error.
- (Q5) Create an instance of your <u>custom LinearRegressor</u> and evaluate its performance using cross-validation, remember to tune the LR!

 Report the appropriate plots and values detailed above in "the repeated tuning process".

 Fill in the cross-validated errors of the regressor yielded by the optimal hyperparameter.

Model	Section	Train MSE	Valid MSE	
		Cross validated		
Dummy	2	filled	filled	
Linear	2			

(Q6) Had we chosen <u>not</u> to normalize features beforehand, would the training performance of these two models have changed (assume there are no numerical errors)? Explain.

<u>Task</u>: Using the best performing hyperparameter, **retrain** the regressor on the <u>entire</u> training set (= with all its samples) and save it for future use (<u>Sec 6</u>).

Section 3: Linear regression with Lasso

In regularized linear regression, we need to tune the regularization strength (λ in our notation, alpha in sklearn). As mentioned earlier, k-fold cross-validation is performed with the **entire** training set (= all its samples) for the remainder of the experiments.

We will now learn to predict contamination_level from the sklearn...Lasso regressor. Make sure your models are non-homogeneous (fit intercept=True).

- (Q7) Tune the regularization strength of the regressor. Follow the repeated tuning process described earlier. Remember to attach the required plot and specify the optimal strength with its validation error.
 - Remember that plot should have suitable titles, axis labels, grid lines, etc.
- (Q8) Fill in the cross-validated errors of the regressor yielded by the optimal hyperparameter.

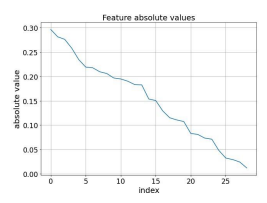
Model	Section	Train MSE Valid MS		
		Cross validated		
Dummy	2	filled	filled	
Linear	2	filled	filled	
Lasso Linear	3			

<u>Task</u>: Using the best performing hyperparameter, **retrain** the regressor on the <u>entire</u> training set (= with all its samples) and save it for future use (<u>Sec 6</u>).

(Q9) Specify the 5 features having the 5 largest coefficients (in absolute value) in the resulting regressor, from the largest to the smallest (among these 5).

We now wish to visualize the resulting coefficients.

- (Q10) Sort and plot the absolute values of the learned coefficients. The x-axis should be the index of the parameter from largest to smallest.
 - The result should roughly look like the illustration to the right (the values are made up).



- (Q11) Why is the magnitude of the coefficients interesting? Explain briefly.
- (Q12) Had we chosen <u>not</u> to normalize features beforehand, would the training performance of the Lasso model have changed (assume there are no numerical errors)? Explain.

Section 4: Polynomial fitting (visualization)

As a detour to better understand polynomial fitting for regression, we now focus on regressing the contamination level, using only PCR 01 and PCR 05.

<u>Task</u>: Create a subset of the training set with these 2 features and contamination_level.

- <u>Task</u>: Visualize the data using plot3d, i.e., plot contamination_level as a function of the PCR_01 and PCR_05 features. (for now, don't use the predictions argument.)
- (Q13) Attach the 3-d plot to your report. What can we understand from this visualization? How should this affect our choice of model to regress contamination level?
- (Q14) We will once again create a baseline, this time our baseline will consist solely of linear Lasso regression (using the two features only), remember again to make sure your models are non-homogeneous (fit_intercept=True).

 Tune for regularization strength as before. Remember to attach required plots, optimal strengths, and optimal validation errors (on this new target).
- <u>Task</u>: Using the best performing hyperparameter, **retrain** the Lasso regressor on the <u>entire</u> training set (= with all its samples).
- (Q15) We will now visualize the model you just trained. Use plot3d to plot all the training samples and their true labels (as before). Also pass your model's predictions in a list to the predictions keyword to compare the true labels to the predicted values. Attach the plot to your report.

We will now try to improve our linear Lasso-regression model by adding quadratic features.

<u>Task</u>: Create a <u>single</u> model that: (1) transforms the original features into <u>2nd</u>-degree polynomial features, (2) normalizes the transformed features, and then (3) trains the linear Lasso regressor on these normalized polynomial features. To do so, we will use a <u>Pipeline</u> like we did in HW2. The following snippet creates such a pipeline:

Complete the first two steps in the pipeline above to make it apply a 2nd-degree PolynomialFeatures transformation, followed by a normalization of your choosing (tip: you are allowed to simply apply a MinMaxScaler for all the transformed features).

(Q16) Why is re-normalization important <u>after</u> applying a polynomial mapping?

Hint: Your answer should revolve around the degrees of the different monomials and focus on mathematical/"optimizational" considerations rather than on statistical ones like in Lecture 08.

(Q17) Tune the regularization strength of a Lasso regressor with the polynomial mapping using cross validation (k = 5 as always). (Remember: on Page 12 in HW2 you saw how to set hyperparameters inside a pipeline. Use <u>cross_validate</u>.)

Remember to attach required plots, optimal strengths, and optimal validation errors.

<u>Task</u>: Using the best performing hyperparameter, **retrain** the Lasso regressor on the <u>entire</u> training set (= with all its samples).

- (Q18) We will now visualize the polynomial model you just trained. Use the new plot3d function to plot all the training samples and their true labels (as before). Pass your model predictions in a list to the predictions keyword to compare the true labels to the predicted values. Attach the plot to your report.
- (Q19) Discuss the results from this section. Compare the two models' capacities to fit the data at hand (use the visualizations, MSE, etc.). This should take 3-6 sentences

Section 5: RandomForest fitting of the CovidScore

Usually, we don't know beforehand which feature mapping is the right one. Furthermore, we might only need to apply each specific feature mapping to different features. For this task, we want to use both our original (pre-processed) features, as well as additional feature mappings such as the RBF feature mapping or the polynomial feature mapping (for some appropriate features). We will then use SciKit's RandomForestRegressor model and the **combination** of both original and transformed features. To approximate the RBF feature mapping (think: why do we need to approximate it?) you can use Scikit's RBFSampler. To apply a feature mapping only to specific features, you can use Scikit's ColumnTransformer. Make sure to build a proper Scikit's Pipeline (as in previous sections), which applies the appropriate normalization, selective feature mapping, and then applies the model (notice: here the normalization should take place before the feature mapping for numerical reasons).

Task: Create copies of your train and test sets (with all the features).

For the RBFSampler: choose (by trial-and-error) appropriate values for the 'gamma' parameter and use the default n_components=100. Also, set the random_state like vou set the seed at the beginning of each assignment.

Note that you can generate multiple mappings (RBF, polynomial etc.) via the ColumnTransformer. Keep in mind that some pre-processing stages might be crucial prior to applying the mapping (think why and at what stage is the normalization sensible).

(Q20) List the features you chose to transform and the transformations that you used, and why.

<u>Tip:</u> we advise transforming only a small subset of the features while <u>keeping</u> the remaining ones (i.e., instead of dropping them).

The Random Forest model is an ensemble of Decision Trees model. Read the following blog post on this model.

We denote the class of all possible (Random Forest models) hypotheses by \mathcal{H}_{RF} .

- (Q21) Explain how (and why) we can expect the training and validation errors (each) of such random forests to change when using the RBF mapping (compared to just using the raw data).
- (Q22) Explain the major difference between a random forest, and the previous polynomial regressors we have used (for the same features).
- (Q23) Tune the 'n_estimators' and 'min_samples_leaf' parameters of the RF model while using RBF mapping using cross-validation with a grid search (see Q8 in Major HW2). Use a similar pipeline to the one created earlier in this section. Remember to attach required heatmaps, optimal hyper-parameters, and optimal train and validation errors.
- **(Q24)** Fill in the train and validation errors of the regressor yielded by the best performing hyperparameter. Remember to compute the errors using <u>cross-validation</u>.

Model	Section	Train MSE Valid M		
		Cross validated		
:	:	filled	filled	
RF Regressor	5			

Task: Train the RF regressor on the entire training set and save it for future use (Sec 6).

Section 6: Testing your models

Important: do not continue to this section until you have finished all previous sections.

Finally, we can let the **test set** come out and play.

At the end of the previous sections, you retrained the tuned models on the <u>entire</u> training set. You will now evaluate the <u>test</u> errors (MSE) for the models, as well as the Dummy baseline.

Remember: do not cross-validate here. Train on the entire training set (= using all its samples) and evaluate on the test set.

(Q25) Complete the entire table

Model	Section	Train MSE	Valid MSE	Test MSE
		Cross validated		Retrained
Dummy	2	filled	filled	
Linear	2	filled	filled	
Lasso Linear	3	filled	filled	
Random Forest	5			

Which model performed best on the test set?

Briefly discuss the results in the table (from an overfitting and underfitting perspective, or any other insightful perspective).