# **Cyclic Coordinate Descent for Logistic Regression with Lasso regularization**

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#### 1 Methodology

#### 1.1 Selection and generation of datasets

The implemented Cyclic Coordinate Descent for Logsitic Regression with Lasso regularization (LogRegCCD) algorithm was tested on both real and synthetic datasets to evaluate its performance given various metrics such as accuracy, precision, recall, F1 score, balanced accuracy and ROC AUC [1, 2, 3] provided by the scikit-learn [4] Python [5] library.

The synthetic datasets have been created with various values of the following parameters: class prior probability p, number of samples n, number of features d, and the covariance matrix g parameter defined as  $S[i,j]=g^{|i-j|}$ . We have examined all unique combinations of the following parameter values:  $p \in \{0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8\}, n \in \{1000, 1500, 2000\}, d \in \{2, 5, 10, 30\}$  and  $g \in \{0.1, 0.3, 0.5, 0.7, 0.9\}$ . For p and g we have examined more parameter values to further investigate certain trends discussed in further sections. The synthetic datasets were generated according to the task description, that is:

- Generate binary class variable (Y=0 or Y=1) from Bernoulli distribution with class prior probability p.
- Generate feature vector X, such that for class Y=0, X follows d-dimensional multivariate normal distribution with mean vector  $\mu_0 = (0, ..., 0)$  and covariance matrix S where  $S[i, j] = g^{|i-j|}$ .
- For class Y=1, X follows d-dimensional multivariate normal distribution with mean vector  $\mu_1 = (1, \frac{1}{2}, \frac{1}{3}, \dots, \frac{1}{d})$  and the same covariance matrix S.
- Generate n observations using the above steps.

All of the real datasets were taken from the OpenML [6] repository. The datasets were selected based on the criteria of having the amount of features of at least 50% of the amount of samples, having preferably numerical features, low amount of missing values and a multi-class response variable. The datasets that have met these criteria are as follows:

- **ArrhythmiaDataset** A binarized version of the Cardiac Arrhythmia Database, where the aim is to determine the type of arrhythmia from the ECG recordings. This dataset contains 279 features, and 452 samples [7].
- **SpeechTreatmentDataset** The dataset containes phonation samples from patients with voice disorders. The aim of the dataset is to assess whether the voice rehabilitation treatment lead to phonations considered 'acceptable' or 'unacceptable'. The dataset contains 309 features, and 126 samples [8].
- **SemeionDataset** A binarized version of the Semeion Handwritten Digit Dataset, where the aim is to determine the digit from the handwritten samples. This dataset contains 256 features, and 319 samples (only instances 1 and 0 are considered) [9].
- **DBWorldSubjectsDataset** The dataset author collected 64 e-mails from DBWorld newsletter and used them to train different algorithms in order to classify between 'announces of conferences' and 'everything else'. The dataset contains 230 features, and 64 samples [10].

#### 1.2 Details about algorithm implementation and applied optimizations

Logistic Regression is a machine learning method capable of binary classification. It predicts the probability of an outcome by computing the linear combination of input features and weights (Formula 1), then passing it through the sigmoid function presented in Formula 2.

$$z = w_0 + w_1 x_1 + w_2 x_2 + \dots + w_n x_n = \mathbf{w}^T \mathbf{x} + b$$
 (1)

Where x denotes input feature vector, while  $x_1, ..., x_n$  are the elements of that vector, w denotes model weights vector and b is the bias term.

$$\sigma(z) = \frac{1}{1 + e^{-z}} \tag{2}$$

The output of the sigmoid function in range [0,1] denotes the probability that given feature vector x belongs to the positive class. What follows the prediction rule is based on the output of the sigmoid function, if it's larger than a set threshold, such as 0.5, we assign the sample to class 1, otherwise assign to class 0.

To fit the model to the training data one needs to minimize the loss function, in this case Binary Cross-Entropy defined in Formula 3.

$$\mathcal{L} = -\frac{1}{m} \sum_{i=1}^{m} \left[ y^{(i)} \log \hat{y}^{(i)} + \left( 1 - y^{(i)} \right) \log \left( 1 - \hat{y}^{(i)} \right) \right] \tag{3}$$

Where m denotes the number of training examples,  $y^{(i)}$  is the class label and  $\hat{y}^{(i)}$  is the predicted probability. The weights of the model need to be optimized to find the proper fit, this can be achieved by standard gradient descent algorithm. The weights are updated according to the following formulas (Formula 4, Formula 5):

$$w_j := w_j - \alpha \frac{\partial \mathcal{L}}{\partial w_j} \tag{4}$$

$$b := b - \alpha \frac{\partial \mathcal{L}}{\partial b} \tag{5}$$

Where  $\alpha$  is the learning rate, the higher the value the more aggressive weight updates and  $\frac{\partial \mathcal{L}}{\partial w_j}$  is a gradient with respect to weight  $w_j$ .

One of the methods to prevent overfitting of the model to the training data is Lasso Regulaization. Overfitting describes the situation when the trained model can predict samples from the training set very well but struggles on the test set. The loss function with Lasso regularization is defined in Formula 6.

$$\mathcal{L}_{\text{lasso}} = -\frac{1}{m} \sum_{i=1}^{m} \left[ y^{(i)} \log \hat{y}^{(i)} + \left( 1 - y^{(i)} \right) \log \left( 1 - \hat{y}^{(i)} \right) \right] + \lambda \sum_{j=1}^{n} |w_j|$$
 (6)

Where m is the number of training samples,  $y^{(i)}$  is the class label,  $\hat{y}^{(i)}$  is the predicted probability,  $\lambda$  denotes regularization strength.

In essence during the training process, the model will also minimize the absolute sum of the coefficients in addition to the loss function. This will result in some of the weights being set to zero,

effectively reducing the number of features the model is trained on. This can be useful in situations where the number of features is very large and some of them are irrelevant to the prediction task.

Now, to use the Cyclic Coordinate Descent instead of the standard Gradient Descent one needs to minimize the  $\mathcal{L}_{lasso}$  using a different algorithm for updating model weights. However the authors of the 2010 publication entitled *Regularization Paths for Generalized Linear Models via Coordinate Descent* [11] present a more sophisticated approach with certain optimizations.

The logistic regression with lasso regulaization log-likelihood function is approximated using a quadratic approximation presented in Formula 7. This converts the problem into a penalized weighted least squares. The authors also use a regularization path that starts from largest  $\lambda$  where  $\beta=0$  and decreases  $\lambda$  gradually, using previous solutions as warm starts. Instead of computing gradients from scratch with each iteration, the authors propose to use covariance updates (Formula 8). This in turn allows for a more efficient computation of the gradients. For each feature, the optimization problem simplifies to a minimization problem presented in Formula 9.

$$\ell_Q(\beta_0, \beta) = -\frac{1}{2N} \sum_{i=1}^N w_i (z_i - \beta_0 - x_i^T \beta)^2 + C$$
(7)

$$\sum_{i=1}^{N} x_{ij} r_i = \langle x_j, y \rangle - \sum_{k:\beta_k \neq 0} \langle x_j, x_k \rangle \beta_k$$
 (8)

$$\min_{\beta_j} \left[ \frac{1}{2} \sum_{i=1}^N w_i \left( z_i - \beta_0 - \sum_{k \neq j} x_{ik} \beta_k - x_{ij} \beta_j \right)^2 + \lambda |\beta_j| \right]$$
(9)

The algorithm for the cyclic coordinate descent for a given feature j is as follows:

1. Compute partial residuals (excluding  $\beta_i$ )

$$r_i = z_i - (\beta_0 + \sum_{k \neq j} x_{ik} \beta_k)$$

2. Compute the gradient component  $\rho_i$ 

$$\rho_j = \sum_{i=1}^N w_i x_{ij} r_i$$

3. Apply soft-thresholding for L1 regularization

$$\beta_j = \frac{S(\rho_j, \lambda)}{\sum_{i=1}^N w_i x_{ij}^2}$$

$$S(z,\lambda) = \operatorname{sign}(z) \cdot \max(|z| - \lambda, 0)$$

4. Update  $\beta_0$  that is not regularized

$$\beta_0 = \frac{\sum_{i=1}^{N} w_i (z_i - x_i^T \beta)}{\sum_{i=1}^{N} w_i}$$

From a high level overview the presented algorithm consists of:

- 1. Outer Loop: Decrease  $\lambda$  along a regularization path.
- 2. Middle Loop: Update the quadratic approximation using the current  $(\beta_0, \beta)$ .
- 3. Inner Loop: Perform coordinate descent on the penalized weighted least squares problem.

# 2 Impact of dataset parameters: n.p,d,g on the performance of LogRegCCD algorithm

The performance of the LogRegCCD algorithm was evaluated on synthetic datasets with different values of the parameters: class prior probability p, number of samples n, number of features d, and the covariance matrix g parameter defined as  $S[i,j]=g^{|i-j|}$ . The results are presented in Figure 1. The performance was evaluated using the following metrics: ROC AUC and balanced accuracy. The effects of the synthetic dataset features on the performance of the models are as follows:

- p: As the class prior probability diverges from 0.5 the balanced accuracy decrease. The dataset becomes more imbalanced and the model struggles to predict the minority class. Whereas for ROC AUC the effect of the p parameter does not seem to follow a regular pattern. All models achieve highest ROC AUC for p = 0.5 when the dataset is balanced. For values other than 0.5 ROC AUC drops in non regular way.
- n: As the number of samples increases the balanced accuracy increases. The model has more data to learn from and can generalize better. For ROC AUC a similar trend is true up to 1500 samples, after that the ROC AUC stays at a similar level.
- **d**: Both balanced accuracy and ROC AUC increase when the number of features changes from two to five. However, afterwards the performance of the model decreases. This is likely due to the fact that the model is overfitting to the training data. The model has too many features to learn from and it starts to memorize the training data instead of generalizing from it.
- g: The g parameter affects the formula  $S[i,j] = g^{|i-j|}$ . Here, we see a tilted U shaped curved, as the covarianc increase the balanced accuracy decreases until around 0.7 where it starts to rapidly increase.

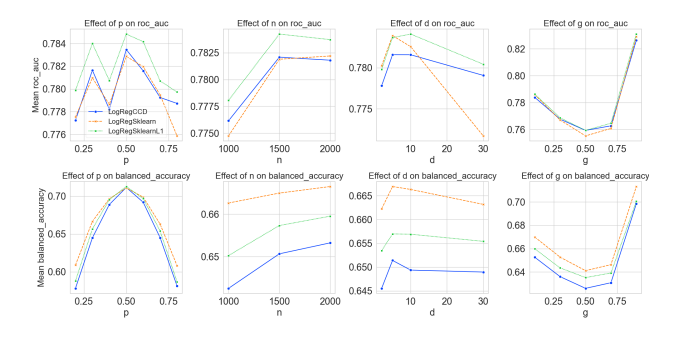


Figure 1: impact of synthetic dataset parameters on the performance of LogRegCCD algorithm

- 3 Benchmark of LogRegCCD with LogisticRegression algorithm
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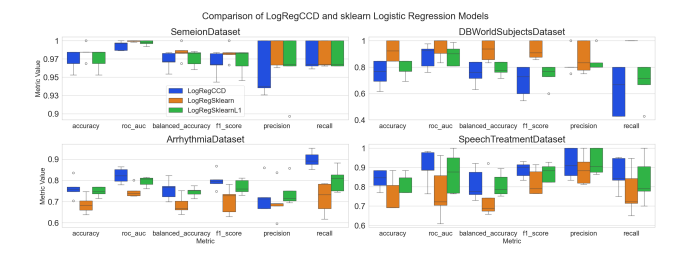


Figure 2: Boxplots of metrics for real datasets

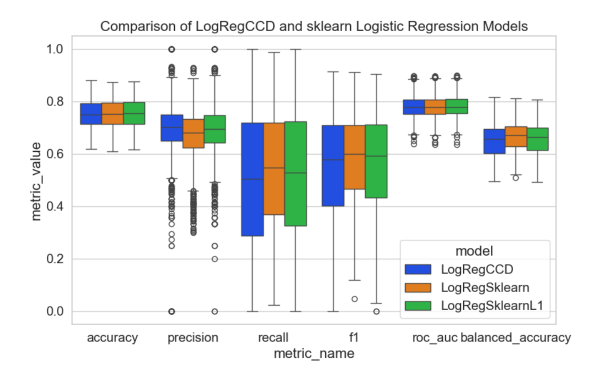
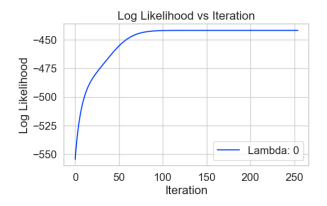


Figure 3: Comparison of LogRegCCD and LogisticRegression on synthetic dataset



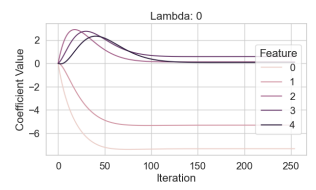


Figure 4: Log likelihood function values depending on iteration for synthetic dataset with  $\lambda=0$ 

Figure 5: Coefficient values depending on iteration for synthetic dataset with  $\lambda=0$ 

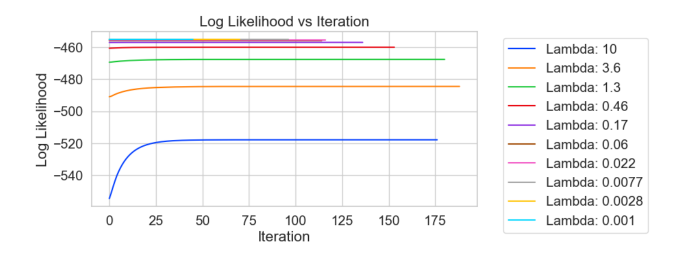


Figure 6: Log likelihood function values depending on iteration for synthetic dataset

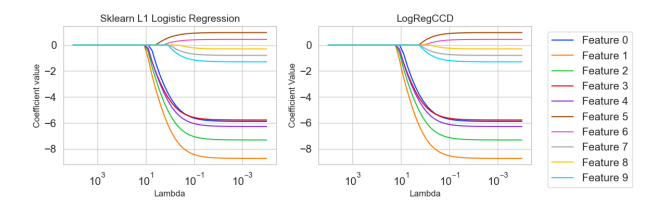


Figure 7: Comparison of LogRegCCD and LogisticRegression on synthetic dataset with redundant features

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