

**EE381V: Large Scale Optimization — Spring 2019**

PROBLEM SET FOUR

Constantine Caramanis

Due: Tuesday, February 19, 2019.

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**Computational Problems**

Save your completed code in a file names `hw4.py` or `hw4.ipynb`. Don't use stock optimization code, you should develop the core part of this assignment yourself. All plots should have titles, axis labels, and lines with different colors, markers, and legend labels.

Any question marked by (?) is optional.

1. Consider the problem of robust regression, where some small number of measurements have been potentially completely corrupted. One way to formulate an optimization problem to solve this robust regression is as follows:

$$\begin{aligned} \min_{\beta} : \quad & \|X\beta - \mathbf{y}\|_1 \\ \text{s.t.} : \quad & \beta \in \mathfrak{X}. \end{aligned}$$

The rationale for this formulation stems from the idea that because of the  $\ell^1$ -error, huge errors are not disproportionately penalized, as they would be in the squared error formulation (this is the formulation we have worked with before, including in the previous problem), and therefore the optimal solution is less sensitive to outliers. You will solve this problem using Projected Subgradient Descent, and also Mirror Descent. Let the constraint set be the simplex:

$$\mathfrak{X} = \{\beta : \beta \geq 0, \sum \beta_i = 1\}.$$

- (a) Write down the update for projected gradient descent.

The data for this problem is generated from the 20 newsgroups dataset <sup>1</sup>. First, the documents are represented as a vector of (normalized) word frequencies. Then, Nonnegative Matrix Factorization (NMF) is used to find “topics” which efficiently represent the collection of documents by assuming that each document is a distribution over topics <sup>2</sup>.  $X$  is an  $n \times m$  matrix where  $n$  is the number of unique words considered and  $m$  is the number of topics. Given these topics, we represent a new document as a distribution over topics given by  $\beta$ . Given the (noisy) word frequencies  $y$ , your goal is to recover that distribution over topics  $\beta$ .

- (b) (?) Explore the robustness properties of this formulation. Compare the performance of standard regression (least squares), with the above formulation using the corrupted observations `y_corr.npy`.

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<sup>1</sup>[scikit-learn.org/stable/datasets/twenty\\_newsgroups.html](http://scikit-learn.org/stable/datasets/twenty_newsgroups.html)

<sup>2</sup>See [scikit-learn.org/stable/modules/generated/sklearn.decomposition.NMF.html](http://scikit-learn.org/stable/modules/generated/sklearn.decomposition.NMF.html) and the references therein.

## 2. Mirror Descent

Consider the same problem of robust regression that you had on the previous problem. There, you solved it using SGD. Repeat the problem, but now using Mirror Descent. Specifically:

- Write down the mirror descent update. For this, we will use the mirror map  $\Phi(\beta) = \sum \beta_i \log \beta_i$  that we used in class. Compute the Bregman divergence,  $D_\Phi(\mathbf{x}, \mathbf{y})$  explicitly.
- Using the data in `X.npy` and `y.npy`, and using stepsizes of your choosing, compare the projected subgradient method (from the last problem) with mirror descent. What is  $\beta$ ? Plot the objective above against iterations for both methods (in a single plot).

## 3. Matrix Completion.

In this problem we investigate **low-rank matrix completion**, the problem of finding a low-rank matrix given only a few (randomly sampled entries). While this is (clearly) not possible in general, somewhat remarkably, it is possible once some additional assumptions are made on the problem setup (for example, for a “random” low-rank matrix and random samples). The assumptions required for guarantees on matrix completion are beyond the scope of this class. Our goal is to develop a projected subgradient algorithm to solve such a problem.

Suppose there is a true matrix  $M \in \mathbb{R}^{m \times n}$  that we want to recover, but we are only given elements in the set  $\Omega \subset [m] \times [n]$  (i.e. we know the value of  $m_{ij}$  if  $(i, j) \in \Omega$ ). We want to solve the following constrained optimization problem

$$\begin{aligned} \min_X \quad & \|X\|_* \\ \text{s.t.} \quad & x_{ij} = m_{ij} \text{ for all } (i, j) \in \Omega \end{aligned}$$

where the variable of optimization  $X \in \mathbb{R}^{m \times n}$  is a matrix. Here  $\|\cdot\|_*$  is the “nuclear” norm, equal to the sum of singular values of the matrix. This norm is a convex but not smooth function of  $X$ ; we will implement projected sub gradient descent for this problem.

The sub gradient of the  $\|\cdot\|_*$  function is as follows: for any matrix  $X$ , if its SVD is  $U\Sigma V'$ , then a matrix  $Z \in \partial\|X\|_*$  is in its sub gradient if and only if

$$Z = UV' + W$$

where  $W$  is such that (a) the column and row spaces of  $W$  are perpendicular to the corresponding ones of  $X$ , and (b) the spectral norm  $\|W\|_2 \leq 1$ . Recall that the spectral norm of a matrix is its maximum singular value. Also recall that if  $X$  is rank  $r$ , then the matrices  $U, V$  are of sizes  $m \times r$  and  $n \times r$  respectively, and have orthonormal columns.

- Given a matrix  $X$ , how will you generate an element  $Z \in \partial\|X\|_*$  using a singular value decomposition function in Python (e.g., in `np.linalg`)?
- Given a matrix  $X$ , how will you project it onto the feasible set (i.e. the set of matrices that satisfy the constraints) ?
- Implement projected sub gradient descent with two choices for step sizes:  $\eta_k = \frac{1}{k}$  and  $\eta_k = \frac{1}{\sqrt{k}}$ . You will need to use the files contained in `MatrixCompletion.zip`, which contains two  $100 \times 100$  matrices: a low-rank matrix  $M$ , and the matrix  $O$  that represents the set  $\Omega$  by having entries that are 0 or 1 (in particular,  $O_{ij} = 1$  means  $(i, j) \in \Omega$ ).
- Plot the relative error  $\frac{1}{100^2} \|M - X_k\|_F^2$  between the true matrix and the  $k^{th}$  iterate, as a function of  $k$ , for both step size choices; do so on one plot.

(e) What is the rank of the intermediate iterates? Why is this the case?

#### 4. Stochastic Variance Reduced Gradient Descent (SVRG)

As we discussed in class, decomposable functions of the form

$$\min_{\omega} \left[ F(\omega) = \frac{1}{n} \sum_i^n f_i(\omega) \right],$$

are very common in statistics/ML problems. Here, each  $f_i$  corresponds to a loss for a particular training example. For example, if  $f_i(\omega) = (\omega^\top x_i - y_i)^2$ , then  $F(\omega)$  is a least squares regression problem. The standard gradient descent (GD) update

$$\omega_t = \omega_{t-1} - \eta_t \nabla F(\omega_{t-1})$$

evaluates the full gradient  $\nabla F(\omega) = \frac{1}{n} \sum_i^n \nabla f_i(\omega)$ , which requires evaluating  $n$  derivatives. This can be prohibitively expensive when the number of training examples  $n$  is large. SGD evaluates the gradient of one (or a small subset) of the training examples—drawn randomly from  $1, \dots, n$ —per iteration:

$$\omega_t = \omega_{t-1} - \eta_t \nabla f_i(\omega_{t-1}).$$

In expectation, the updates are equivalent, but SGD has the computational advantage of only evaluating a single gradient  $\nabla f_i(\omega)$ . The disadvantage is that the randomness introduces variance, which slows convergence. This was our motivation in class to introduce the SVRG algorithm.

Given the dataset in `digits.zip`, plot the performance of GD, SGD, and SVRG for logistic regression with  $l_2$  regularization in terms of negative log likelihood on the training data against the number of gradient evaluations for a single training example (GD performs  $n$  such evaluations per iteration and SGD performs 1). Choose the  $l_2$  parameter to optimize performance on the test set. How does the choice of  $m$  affect the performance of SVRG? There should be one plot with a title and three lines with different colors, markers, and legend labels.

- Using any approach, optimize performance of logistic regression on the test set in `news.zip` and compare the performance of your approach to standard SGD. This dataset is the full-dimensional newsgroup dataset (as opposed to the compressed version you worked with previously). The  $X$  matrices are stored in sparse matrix format and can be read using `scipy.io.mmread`. As the dataset is large and high-dimensional, you will have to decide on how best to allocate your computational resources. Try to utilize the sparsity of the data (i.e. don't just convert it to a dense matrix and spend all your time multiplying zeros). You may use any of the techniques covered in class or ideas from outside class (e.g. `momentum`, `variance reduction`, `minibatches`, `adaptive learning rates`, `preprocessing`). Describe your methodology and comment on what you found improved performance and why. Plot the performance (negative log likelihood) of your method against standard SGD in terms of the number of gradient evaluations.

## Written Problems

- (a) Prove that a matrix  $Z$  as described above in the matrix completion problem is indeed a sub gradient to the nuclear norm function at  $X$ . You can use the following fact about the nuclear norm: for any matrix  $M \in \mathbb{R}^{m \times n}$ , let  $s = \min(m, n)$ . Then for any matrices  $A \in \mathbb{R}^{m \times s}$  and  $B \in \mathbb{R}^{n \times s}$  that have orthonormal columns, we have that

$$\|M\|_* \geq \langle M, AB' \rangle$$

- (b) (?) Re-do the computation we did in class, showing that if we use the mirror function

$$\Phi(\mathbf{x}) = \frac{1}{2} \|\mathbf{x}\|_2^2,$$

then the Mirror Descent update for:

$$\begin{aligned} \min_{\mathbf{x}} : & \quad f(\mathbf{x}) \\ \text{s.t.} : & \quad \mathbf{x} \in \mathfrak{X}, \end{aligned}$$

is exactly projected subgradient descent.

- (c) Here you will do some work that helps compute the Mirror Descent update you need for the computational problem above. As we discussed in class, and also as is explained in Section 4.2 of Bubeck's notes, the Mirror Descent update can also be obtained as:

$$x_{t+1} = \arg \min_{x \in \mathcal{X} \cap \mathcal{D}} : \eta \langle g_{x_t}, x \rangle + D_\Phi(x, x_t).$$

Therefore, Mirror Descent is only computationally useful if we can easily solve the problem:

$$\min_{u \in \mathcal{X}} : \langle z, u \rangle + \Phi(u).$$

In this problem, you will show that when  $\mathcal{X}$  is the simplex, i.e.,  $\mathcal{X} = \Delta_n$ , then this problem is indeed easy.

- i. Consider the optimization problem:

$$\begin{aligned} \min : & \quad \langle z, u \rangle + \Phi(u) \\ \text{s.t.} : & \quad \sum_i u_i = 1. \end{aligned}$$

(Note that the constraints  $\{u_i \geq 0\}$  are implicitly included as they are part of  $\text{dom}\Phi$ .) Write the Lagrangian for the problem. The variables will be  $u$  and a single variable  $\lambda$  for the single constraint. Write the KKT conditions for the problem.

- ii. Using the KKT conditions, derive a closed form expression for  $u$  as a function of  $z$ .  
 iii. Now go back to the Mirror Descent update and write explicitly the Mirror Descent update using your work above.
- (d) **Gradient descent and non-convexity**  
 Consider the gradient descent algorithm with fixed step size  $\eta$  for the function  $f(x) = x'Qx$ , where  $Q$  is symmetric but not positive semidefinite. (i.e.,  $Q$  has some negative eigenvalues). Exactly describe the set of initial points from which gradient descent, with *any* positive step size, will diverge. What happens at the other points, if the step size is small enough ?