CPSC 340 Assignment 3 (Due 2021-10-15 at 11:59pm)

Pedram Amani - 73993008 Henry Xu - 40728164

Important: Submission Format [5 points]

Please make sure to follow the submission instructions posted on the course website. We will deduct marks if the submission format is incorrect, or if you're not using LATEX and your handwriting is at all difficult to read – at least these 5 points, more for egregious issues. Compared to assignment 1, your name and student number are no longer necessary (though it's not a bad idea to include them just in case, especially if you're doing the assignment with a partner).

1 Matrix Notation and Minimizing Quadratics [12 points]

1.1 Converting to Matrix/Vector/Norm Notation [6 points]

Using our standard supervised learning notation (X, y, w) express the following functions in terms of vectors, matrices, and norms (there should be no summations or maximums).

1. $\max_{i \in \{1,2,\ldots,n\}} |w^T x_i - y_i|$. This is "brittle regression".

Answer: $||Xw - y||_{\infty}$

2. $\sum_{i=1}^{n} v_i (w^T x_i - y_i)^2 + \frac{\lambda}{2} \sum_{j=1}^{d} w_j^2$. This is regularized least squares with a weight v_i for each training example: Hint: You can use V to denote a diagonal matrix that has the values v_i along the diagonal. What does $a^T V b$ look like in summation form (for some arbitrary vectors a, b)?

Answer: $(w^T X^T - y^T)V(Xw - y) + \frac{\lambda}{2}w^T w$

3. $\left(\sum_{i=1}^{n}|w^{T}x_{i}-y_{i}|\right)^{2}+\frac{1}{2}\sum_{j=1}^{d}\lambda_{j}|w_{j}|$. This is L1-regularized least squares with a different regularization strength for each dimension: Hint: You can use Λ to denote a diagonal matrix that has the λ_{j} values along the diagonal.

Answer: $||Xw - y||^2 + \frac{1}{2}||\Lambda w||_1$

Note: you can assume that all the v_i and λ_i values are non-negative.

1.2 Minimizing Quadratic Functions as Linear Systems [6 points]

Write finding a minimizer w of the functions below as a system of linear equations (using vector/matrix notation and simplifying as much as possible). Note that all the functions below are convex, so finding a w with $\nabla f(w) = 0$ is sufficient to minimize the functions – but show your work in getting to this point.

1

1. $f(w) = \frac{1}{2} ||w - v||^2$ (projection of v onto real space).

Answer: Using notation $[a_i]$ to mean the column vector $\begin{bmatrix} a_1 \\ a_2 \\ \dots \\ a_n \end{bmatrix}$, we have:

$$\nabla f(w) = 0 = \frac{1}{2} \left[\partial_{w_i} (w_i - v_i)^2 \right] = [w_i - v_i] = w - v \implies w = v$$

2. $f(w) = \frac{1}{2} ||Xw - y||^2 + \frac{1}{2} w^T \Lambda w$ (least squares with weighted regularization).

Answer: Using the same notation as 1.2.1 and taking x_{i} to mean the i^{th} column vector in X:

$$\nabla f(w) = 0 = \frac{1}{2} \left[\partial_{w_i} \sum_j \left(\left(w^T x_j - y_j \right)^2 + \lambda_j w_j^2 \right) \right] = \left[\sum_j \left(w^T x_j - y_j \right) x_{ji} + \lambda_i w_i \right]$$
$$= \left[x_{:i}^T (Xw - y) + \lambda_i w_i \right] = X^T (Xw - y) + \Lambda w \implies \left(X^T X + \Lambda \right) w = X^T y$$

3. $f(w) = \frac{1}{2} \sum_{i=1}^n v_i (w^T x_i - y_i)^2 + \frac{\lambda}{2} \|w - w^{(0)}\|^2$ (weighted least squares shrunk towards non-zero $w^{(0)}$).

Answer: Using the same notation as 1.2.1 and taking x_{i} to mean the i^{th} column vector in X:

$$\nabla f(w) = 0 = \frac{1}{2} \left[\partial_{w_i} \sum_{j} \left(v_j \left(w^T x_j - y_j \right)^2 + \lambda_j \left(w_j - w_j^{(0)} \right)^2 \right) \right] = \left[\sum_{j} v_j \left(w^T x_j - y_j \right) x_{ji} + \lambda_i \left(w_i - w_i^{(0)} \right) \right]$$

$$= \left[x_{:i}^T V(Xw - y) + \lambda_i \left(w_i - w_i^{(0)} \right) \right] = X^T V(Xw - y) + \Lambda \left(w - w^{(0)} \right)$$

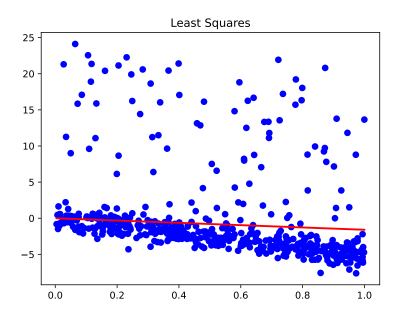
$$\implies \left(X^T V X + \Lambda \right) w = X^T V y + \Lambda w^{(0)}$$

Above we assume that v and $w^{(0)}$ are $d \times 1$ vectors, and Λ is a $d \times d$ diagonal matrix (with positive entries along the diagonal). You can use V as a diagonal matrix containing the v_i values along the diagonal.

Hint: Once you convert to vector/matrix notation, you can use the results from class to quickly compute these quantities term-wise. As a spot check, make sure that the dimensions match for all quantities/operations: to do this, you may need to introduce an identity matrix. For example, $X^TXw + \lambda w$ can be re-written as $(X^TX + \lambda I)w$.

2 Robust Regression and Gradient Descent [41 points]

If you run python main.py -q 2, it will load a one-dimensional regression dataset that has a non-trivial number of 'outlier' data points. These points do not fit the general trend of the rest of the data, and pull the least squares model away from the main downward trend that most data points exhibit:



Note: we are fitting the regression without an intercept here, just for simplicity of the homework question. In reality one would rarely do this. But here it's OK because the "true" line passes through the origin (by design). In Q3.1 we'll address this explicitly.

A coding note: when we're doing math, we always treat y and w as column vectors, i.e. if we're thinking of them as matrices, then shape $n \times 1$ or $d \times 1$, respectively. This is also what you'd usually do when coding things in, say, Matlab. It is not what's usually done in Python machine learning code, though: we usually have y.shape == (n,), i.e. a one-dimensional array. Mathematically, these are the same thing, but if you mix between the two, you can really easily get confusing answers: if you add something of shape (n, 1) to something of shape (n,), then the NumPy broadcasting rules give you something of shape (n, n). This is a very unfortunate consequence of the way the broadcasting rules work. If you stick to either one, you generally don't have to worry about it; we're assuming shape (n,) here. Note that you can go from something of shape (n, 1) to (n,) by, for example, using two_d_array.squeeze(1) (which checks that the axis at index 1, the second one, is length 1 and then removes it). You can go in the other direction with, for instance, one_d_array[:, np.newaxis] (which says "give me the whole first axis, then add another axis of length 1 in the second position").

2.1 Weighted Least Squares in One Dimension [8 points]

One of the most common variations on least squares is weighted least squares. In this formulation, we have a weight v_i for every training example. To fit the model, we minimize the weighted squared error,

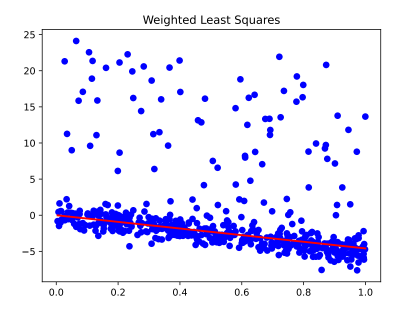
$$f(w) = \frac{1}{2} \sum_{i=1}^{n} v_i (w^T x_i - y_i)^2.$$

In this formulation, the model focuses on making the error small for examples i where v_i is high. Similarly, if v_i is low then the model allows a larger error. Note: these weights v_i (one per training example) are completely different from the model parameters w_j (one per feature), which, confusingly, we sometimes also call "weights." The v_i are sometimes called *sample weights* or *instance weights* to help distinguish them.

Complete the model class, WeightedLeastSquares (inside linear_models.py), to implement this model. (Note that Q1.2.3 asks you to show how a similar formulation can be solved as a linear system.) Apply this model to the data containing outliers, setting v = 1 for the first 400 data points and v = 0.1 for the last 100 data points (which are the outliers). Hand in your code and the updated plot.

Answer: https://numpy.org/doc/stable/reference/generated/numpy.diag.html

```
def fit(self, X, y, v):
    V = np.diag(v)
    self.w = solve(X.T @ V @ X, X.T @ V @ y)
```



2.2 Smooth Approximation to the L1-Norm [8 points]

Unfortunately, we typically do not know the identities of the outliers. In situations where we suspect that there are outliers, but we do not know which examples are outliers, it makes sense to use a loss function that is more robust to outliers. In class, we discussed using the sum of absolute values objective,

$$f(w) = \sum_{i=1}^{n} |w^{T} x_{i} - y_{i}|.$$

This is less sensitive to outliers than least squares, but it is non-differentiable and harder to optimize. Nevertheless, there are various smooth approximations to the absolute value function that are easy to optimize. One possible approximation is to use the log-sum-exp approximation of the max function¹:

$$|r| = \max\{r, -r\} \approx \log(\exp(r) + \exp(-r)).$$

 $^{^1 \}text{Other possibilities}$ are the Huber loss, or $|r| \approx \sqrt{r^2 + \epsilon}$ for some small $\epsilon.$

Using this approximation, we obtain an objective of the form

$$f(w) = \sum_{i=1}^{n} \log \left(\exp(w^{T} x_{i} - y_{i}) + \exp(y_{i} - w^{T} x_{i}) \right).$$

which is smooth but less sensitive to outliers than the squared error. Derive the gradient ∇f of this function with respect to w. You should show your work but you do <u>not</u> have to express the final result in matrix notation.

Answer: Using the same notation as 1.2.1 and taking x_{i} to mean the i^{th} column vector in X:

$$\nabla f(w) = \left[\partial_{w_i} \sum_{j} \log \left(2 \cosh \left(w^T x_j - y_j \right) \right) \right] = \left[\sum_{j} \frac{\partial_{w_i} \left(\cosh \left(w^T x_j - y_j \right) \right)}{\cosh \left(w^T x_j - y_j \right)} \right]$$

$$= \left[\sum_{j} \frac{\partial_{w_i} \left(w^T x_j - y_j \right) \cdot \sinh \left(w^T x_j - y_j \right)}{\cosh \left(w^T x_j - y_j \right)} \right] = \left[\sum_{j} x_{ji} \cdot \tanh \left(w^T x_j - y_j \right) \right]$$

$$= \left[x_{ij}^T \tanh \left(Xw - y \right) \right] = X^T \tanh \left(Xw - y \right)$$

2.3 Gradient Descent: Understanding the Code [5 points]

Recall gradient descent, a derivative-based optimization algorithm that uses gradients to navigate the parameter space until a locally optimal parameter is found. In optimizers.py, you will see our implementation of gradient descent, taking the form of a class named OptimizerGradientDescent. This class has a similar design pattern as PyTorch, a popular differentiable programming and optimization library. One step of gradient descent is defined as

$$w^{t+1} = w^t - \alpha^t \nabla_w f(w^t).$$

Look at the methods named get_learning_rate_and_step() and break_yes(), and answer each of these questions, one sentence per answer:

1. Which variable is equivalent to α^t , the step size at iteration t?

Answer: In get_learning_rate_and_step(), the step size is alpha = self.learning_rate which is not changed between iterations.

2. Which variable is equivalent to $\nabla_w f(w^t)$ the current value of the gradient vector?

Answer: In get_learning_rate_and_step(), g_old is the current gradient vector.

3. Which variable is equivalent to w^t , the current value of the parameters?

Answer: In get_learning_rate_and_step(), w_old is the current value of parameters.

4. What is the method break_yes() doing?

Answer: break_yes() determines whether to end the gradient descent iterations, by checking whether the desired tolerance or the maximum number of iterations have been reached.

2.4 Robust Regression [20 points]

The class LinearModelGradientDescent is the same as LeastSquares, except that it fits the least squares model using a gradient descent method. If you run python main.py -q 2.4 you'll see it produces the same fit as we obtained using the normal equations.

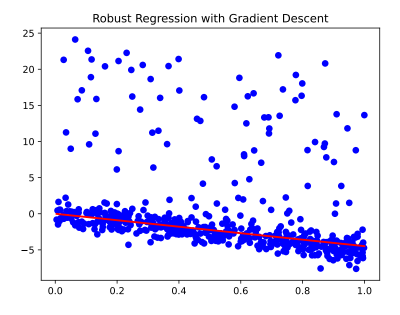
The typical input to a gradient method is a function that, given w, returns f(w) and $\nabla f(w)$. See funObj in LinearModelGradientDescent for an example. Note that the fit function of LinearModelGradientDescent also has a numerical check that the gradient code is approximately correct, since implementing gradients is often error-prone.²

An advantage of gradient-based strategies is that they are able to solve problems that do not have closed-form solutions, such as the formulation from the previous section. The class LinearModelGradientDescent has most of the implementation of a gradient-based strategy for fitting the robust regression model under the log-sum-exp approximation.

2.4.1 Implementing the Objective Function [15 points]

Optimizing robust regression parameters is the matter of implementing a function object and using an optimizer to minimize the function object. The only part missing is the function and gradient calculation inside fun_obj.py. Inside fun_obj.py, complete FunObjRobustRegression to implement the objective function and gradient based on the smooth approximation to the absolute value function (from the previous section). Hand in your code, as well as the plot obtained using this robust regression approach.

```
def evaluate(self, w, X, y):
    f = np.sum(np.log(2 * np.cosh(X @ w - y)))
    g = X.T @ np.tanh(X @ w - y)
    return f, g
```



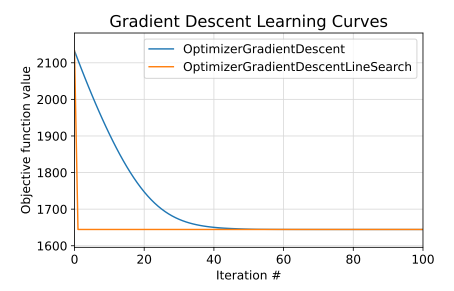
2.4.2 The Learning Curves [5 points]

Using the same dataset as the previous sections, produce the plot of "gradient descent learning curves" to compare the performances of OptimizerGradientDescent and OptimizerGradientDescentLineSearch for robust regression, where **one hundred (100) iterations** of gradient descent are on the x-axis and the **objective function value** corresponding to each iteration is visualized on the y-axis (see gradient descent

 $^{^2}$ Sometimes the numerical gradient checker itself can be wrong. See CPSC 303 for a lot more on numerical differentiation.

lecture). Use the default learning_rate for OptimizerGradientDescent. Submit this plot. According to this plot, which optimizer is more "iteration-efficient"?

Answer: OptimizerGradientDescentLineSearch is much more iteration-efficient, converging in merely 2 iterations.



3 Linear Regression and Nonlinear Bases

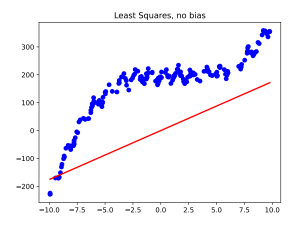
In class we discussed fitting a linear regression model by minimizing the squared error. In this question, you will start with a data set where least squares performs poorly. You will then explore how adding a bias variable and using nonlinear (polynomial) bases can drastically improve the performance. You will also explore how the complexity of a basis affects both the training error and the validation error.

3.1 Adding a Bias Variable [8 points]

If you run python main.py -q 3, it will:

- 1. Load a one-dimensional regression dataset.
- 2. Fit a least-squares linear regression model.
- 3. Report the training error.
- 4. Report the validation error.
- 5. Draw a figure showing the training data and what the linear model looks like.

Unfortunately, this is an awful model of the data. The average squared training error on the data set is over 28000 (as is the validation error), and the figure produced by the demo confirms that the predictions are usually nowhere near the training data:



The y-intercept of this data is clearly not zero (it looks like it's closer to 200), so we should expect to improve performance by adding a bias (a.k.a. intercept) variable, so that our model is

$$y_i = w^T x_i + w_0.$$

instead of

$$y_i = w^T x_i$$
.

In file linear_models.py, complete the class LeastSquaresBias, that has the same input/model/predict format as the LeastSquares class, but that adds a bias variable (also called an intercept) w_0 (also called β in lecture). Hand in your new class, the updated plot, and the updated training/validation error.

Hint: recall that adding a bias w_0 is equivalent to adding a column of ones to the matrix X. Don't forget that you need to do the same transformation in the **predict** function.

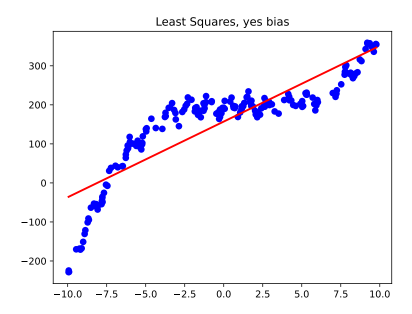
Answer: https://numpy.org/doc/stable/reference/generated/numpy.pad.html

Training error = 3551.3

Validation error = 3393.9

```
class LeastSquaresBias:
    def fit(self, X, y):
        X = np.pad(X, ((0, 0), (0, 1)), 'constant', constant_values=(0, 1))
        self.w = solve(X.T @ X, X.T @ y)

def predict(self, X):
        X = np.pad(X, ((0, 0), (0, 1)), 'constant', constant_values=(0, 1))
        return X @ self.w
```



3.2 Polynomial Basis [10 points]

Adding a bias variable improves the prediction substantially, but the model is still problematic because the target seems to be a *non-linear* function of the input. Complete LeastSquarePoly class, that takes a data vector x (i.e., assuming we only have one feature) and the polynomial order p. The function should perform a least squares fit based on a matrix Z where each of its rows contains the values $(x_i)^j$ for j=0 up to p. E.g., LeastSquaresPoly.fit(x,y) with p=3 should form the matrix

$$Z = \begin{bmatrix} 1 & x_1 & (x_1)^2 & (x_1)^3 \\ 1 & x_2 & (x_2)^2 & (x_2)^3 \\ \vdots & & & \\ 1 & x_n & (x_n)^2 & (x_N)^3 \end{bmatrix},$$

and fit a least squares model based on it. Submit your code, and a plot showing training and validation error curves for the following values of p: 0, 1, 2, 3, 4, 5, 10, 20, 30, 50, 75, 100. Clearly label your axes, and use a logarithmic scale for y by plt.yscale("log") or similar, so that we can still see what's going on if there are a few extremely large errors. Explain the effect of p on the training error and on the validation error.

NOTE: large values of p may cause numerical instability. Your solution may look different from others' even with the same code depending on the OS and other factors. As long as your training and validation error curves behave as expected, you will not be penalized.

Note: you should write the code yourself; don't use a library like sklearn's PolynomialFeatures.

Note: in addition to the error curves, the code also produces a plot of the fits themselves. This is for your information; you don't have to submit it.

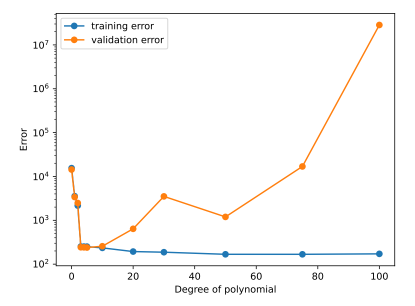
Answer: Model complexity increases with the hyper-parameter p. Therefore, a higher value of p results in a lower training error while the validation error increases due to overfitting.

```
class LeastSquaresPoly:
    def __init__(self, p):
        self.leastSquares = LeastSquares()
        self.p = p

def fit(self, X, y):
        Z = self._poly_basis(X)
        self.w = solve(Z.T @ Z, Z.T @ y)

def predict(self, X):
        return self._poly_basis(X) @ self.w

def __poly_basis(self, X):
        Z = np.empty((len(X), self.p + 1))
        for i in range(0, self.p + 1):
              Z[:, i] = (X ** i).squeeze(1)
        return Z
```



4 Very-Short Answer Questions [24 points]

Answer the following questions (in a sentence or two).

1. Suppose that a training example is global outlier, meaning it is really far from all other data points. How is the cluster assignment of this example by k-means? And how is it set by density-based clustering?

Answer: k-means: The cluster containing the outlier is shifted towards the outlier value after each iteration as the non-outlier values get moved to other clusters. After convergence (depending on initial condition and the number k), there will likely be a single cluster containing only the outlier. density-based: Since the outlier value is far from all other data points, if the threshold distance hyperparameter is chosen appropriately, it will be categorized in its own cluster.

2. Why do we need random restarts for k-means but not for density-based clustering?

Answer: Because the clustering result of k-means is sensitive to initial mean values, running with multiple random initial conditions and taking the mode gives a better clustering (otherwise we might get stuck in a local minimum in the parameter space). On the contrary, density-based clustering gives the same clustering independent of the initial data points.

3. Can hierarchical clustering find non-convex clusters?

Answer: Density-based clustering can find non-convex clusters, so obviously hierarchical density-based clustering can as well.

While k-means can only find convex clusters, hierarchical k-means can yield a non-convex cluster that contains several convex sub-clusters.

4. For model-based outlier detection, list an example method and problem with identifying outliers using this method.

Answer: Say we want to find the outliers in the final CPSC 340 grades. We could model the grade distribution with a normal distribution $\mathcal{N}(\mu, \sigma)$ and consider the grades that are more than 3σ as outliers. This method of outlier detection fails when our model fails to capture the complexity of our data (i.e. a bimodal grade distribution since some students come from a non-computer-science background).

5. For graphical-based outlier detection, list an example method and problem with identifying outliers using this method.

Answer: Say we are interested in identifying unreasonably-priced houses based on some metrics (square-footage, location, etc.). We could plot prices vs. square-footage on a 2D scatter plot and identify the data points that do not follow the trend as outliers. The problem is that we are more-orless limited to looking at a single feature (as higher dimensional plots are very difficult to work with).

6. For supervised outlier detection, list an example method and problem with identifying outliers using this method.

Answer: Say we are interested in identifying credit fraud provided to some metrics about a transaction. We can train a sophisticated model on labeled data (including prior cases of credit fraud) for this purpose. But our model may fail to identify new types of fraud because our training set lacked similar examples.

7. If we want to do linear regression with 1 feature, explain why it would or would not make sense to use gradient descent to compute the least squares solution.

Answer: The cost of solving the normal equation (with 1 feature) exactly is O(n). And the cost of finding the solution numerically with gradient descent is O(nm) where m is the number of iterations.

Since finding the exact solution costs less, it does not make sense to do gradient descent for d=1.

8. Why do we typically add a column of 1 values to X when we do linear regression? Should we do this if we're using decision trees?

Answer: Adding a column of 1s to X effectively adds an extra shift parameter to our model w which is the y-intercept.

Doing so for decision trees makes no sense for several reasons. Most obviously, a column of 1s as a feature does not help a decision tree differentiate between different examples.

9. Why do we need gradient descent for the robust regression problem, as opposed to just using the normal equations? Hint: it is NOT because of the non-differentiability. Recall that we used gradient descent even after smoothing away the non-differentiable part of the loss.

Answer: Because for robust regression (regardless of our choice of a smooth approximation to the absolute value function), $\nabla f = 0$ is not a linear system and thus cannot be solved exactly. Therefore, we resort to a numerical solution using gradient descent.

10. What is the problem with having too small of a learning rate in gradient descent? What is the problem with having too large of a learning rate in gradient descent?

Answer: A small learning rate can lead to increased computation cost since more iterations are needed to get to the objective function's minimum.

A large learning rate can also lead to lower iteration-efficiency as we may miss the minimum and keep oscillating between the left and right sides of the minimum.

11. What is the purpose of the log-sum-exp function and how is this related to gradient descent?

Answer: The log-sum-exp function is a smooth and differentiable approximation to the absolute value function. Since log-sum-exp is differentiable at x = 0, its gradient is defined at x = 0 and does not cause issues in gradient descent (unlike the absolute value function).

12. What type of non-linear transform might be suitable if we had a periodic function?

Answer: We can fit to the $\sin(x)$ of our data. More generally, we can fit to the Fourier transform of our data.