CPSC 540: Machine Learning Coordinate Optimization

Mark Schmidt

University of British Columbia

Winter 2018

Last Time: Structured Regularization

• We discussed total variation regularization,

$$\underset{w}{\operatorname{argmin}} f(w) + \sum_{(i,j) \in E} \lambda_{ij} (w_i - w_j)^2,$$

if we want w_i values to be similar across nodes in a graph.

• We discussed structured sparsity,

$$\underset{w}{\operatorname{argmin}} f(w) + \sum_{g \in \mathcal{G}} \lambda_g \|w_g\|,$$

where overlapping groups can enforce patterns of sparsity.

- These regularizers aren't "simple", but several solvers exist.
 - Gradient descent if smooth, inexact proximal gradient for non-smooth.

Transductive Learning

• Our usual supervised learning framework:

$$X = \begin{bmatrix} 0 & 0.7 & 0 & 0.3 & 0 & 0 \\ 0.3 & 0.7 & 0 & 0.6 & 0 & 0.01 \\ 0 & 0 & 0 & 0.8 & 0 & 0 \\ 0.3 & 0.7 & 1.2 & 0 & 0.10 & 0.01 \end{bmatrix}, \quad y = \begin{bmatrix} 1 \\ 1 \\ 0 \\ 1 \end{bmatrix}.$$

In transductive learning, we also have unlabeled examples,

$$\bar{X} = \begin{bmatrix} 0.3 & 0 & 1.2 & 0.3 & 0.10 & 0.01 \\ 0.6 & 0.7 & 0 & 0.3 & 0 & 0.01 \\ 0 & 0.7 & 0 & 0.6 & 0 & 0 \\ 0.3 & 0.7 & 0 & 0 & 0.20 & 0.01 \end{bmatrix},$$

and our goal is to label these particular examples.

Transductive Learning

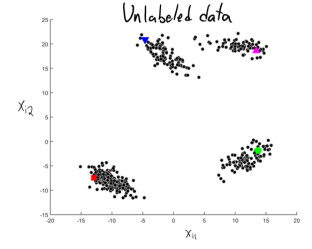
- Transductive learning framework:
 - We have n labeled examples (x^i, y^i) .
 - 2 We have t unabeled examples \bar{x}^i that we want to label.
- This arises a lot:
 - Usually getting unlabeled data is easy but getting labeled data is hard (t >> n).
 - Typically situation: small number labeled examples and huge number of unlabeled.
- Sometimes classifying the data is an intermediate step:
 - Goal is to ulimately use labeled examples to do something else.
 - "I can label a small number of examples, if it helps labeling them all".
- Sometimes it's not possible to obtain labels for any x^i .
 - Predicting gene functions is limited by what we can measure.

Transductive Learning vs. (Semi-)Supervised Learning

- Transductive learning is a special case semi-supervised learning (SSL).
 - Learning with labeled and unlabeled examples (we'll come back to SSL later).
- But transductive SSL has an unusual measure of performance:
 - We don't worry about "test error" (performance on all possible examples).
 - We only care about error for our "test" examples \bar{x}^i .
- Any supervised or semi-supervised method can be used for transduction.
 - Fit model, then apply it to unlabeled examples.
- But in transductive learning, we don't need a model that can predict on new \tilde{x}^i .
 - Some methods don't fit a generic model for mapping from x^i to y^i .

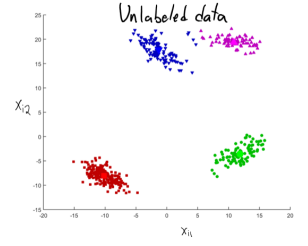
Transductive Learning

- Why should unlabeled data tell us anything about labels?
 - ullet Usually, we assume that similar features o similar labels.



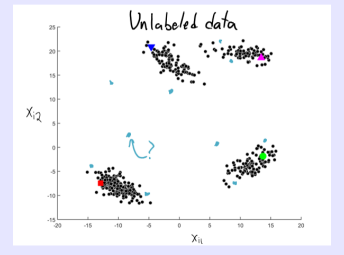
Transductive Learning

- Why should unlabeled data tell us anything about labels?
 - ullet Usually, we assume that similar features o similar labels.



Digression: Transductive vs. Inductive SSL

- In transductive learning we don't need to be able to predict on new examples.
 - In inductive semi-supervised learning goal is to predict well on new examples.



Label Propagation (Graph-Based SSL)

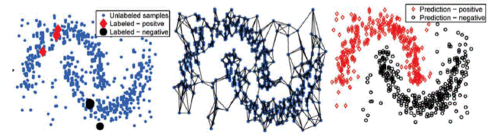
- We could optimize the \bar{y}^i to encourage "similar features \to similar labels".
- Label propagation ("graph-based SSL") method:
 - Define weights w_{ij} saying how similar labeled example i is to unlabled example j.
 - Define weights \bar{w}_{ij} saying how similar unlabeled example i is to unlabeled example j.
 - Find labels \bar{y}^i minimizing a measure of total variation on the label space:

$$\underset{\bar{y} \in \mathbb{R}^t}{\operatorname{argmin}} \sum_{i=1}^n \sum_{j=1}^t w_{ij} (y^i - \bar{y}^j)^2 + \frac{1}{2} \sum_{i=1}^t \sum_{j=1}^t \bar{w}_{ij} (\bar{y}^i - \bar{y}^j)^2.$$

- First term: unlabeled example should get similar labels to "close" labeled examples.
 - "If x^i and \bar{x}^j are similar, then \bar{y}^j should be similar to y^i ."
- Second term: similar unlabeled examples should have similar labels.
 - "Label information 'propagates' through the graph of \bar{y}^i values".

Label Propagation (Graph-Based SSL)

- Label propagation is often surprisingly effective (even with few labeled examples).
- A common choice of the weights (many variations exist):
 - Find the k-nearest neighbours of each example (among labeled and unlabeled).
 - Set w_{ij} and \bar{w}_{ij} to 0 if nodes i and j aren't neighbours.
 - Otherwise, set these to some measure of similarity between features.



Label Propagation for YouTube Tagging and Bioinformatics

- Label propagation doesn't necessarily need features.
 - Consider assigning "tags" to YouTube vidoes (e.g., "cat").



www.youtube.com

- Construct a graph based on sequence of videos that people watch.
 - Give high weight if video 'A' is often followed/preceded by video 'B'.
- Use label propagation to tag all videos.
- Becoming popular in bioinformatics:
 - Label a subset of genes using manual experiments.
 - Find out which genes interact using more manual experiments.
 - Predict function/location/etc. of genes using label propagation.

Label Propagation Variations

- Many variations on label propagation exist:
 - Different ways to choose the graph/weights.
 - Multi-class versions,

$$\underset{\bar{Y} \in \mathbb{R}^{t \times k}}{\operatorname{argmin}} \sum_{i=1}^{n} \sum_{j=1}^{t} w_{ij} \|y^{i} - \bar{y}^{j}\|^{2} + \frac{1}{2} \sum_{i=1}^{t} \sum_{j=1}^{t} \bar{w}_{ij} \|\bar{y}^{i} - \bar{y}^{j}\|^{2}.$$

• Other measures of similarity/distance,

$$\underset{\bar{y} \in \mathbb{R}^t}{\operatorname{argmin}} \sum_{i=1}^n \sum_{j=1}^t f_{ij}(y^i, \bar{y}^j) + \sum_{i=1}^t \sum_{j=1}^t f_{ij}(\bar{y}^i, \bar{y}^j).$$

- Variants where the given labels y^i are also variables (as they might be wrong).
 - Weight gives how much you trust original label.
- ullet Variants where the unlabeled \bar{y}^i are regularized towards a default value.
 - Can reflect that example is really far from any labeled examples.

Outline

Label Propagation

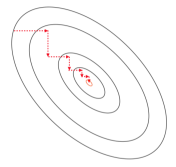
Coordinate Optimization

Beyond Gradient Descent

- For high-dimensional problems we often prefer gradient descent over Newton.
 - Gradient descent requires far more iterations.
 - But iteration cost is only linear in d.
- For very large datasets, even gradient descent iterations can be too slow.
 - If iteration cost is O(nd), we may only be able to do a small number of iterations.
- Two common strategies for yielding even cheaper iterations:
 - Coordinate optimization (today).
 - Stochastic gradient (next time).

Coordinate Optimization

• Each iteration of coordinate optimization only updates on variable:



 \bullet For example, on iteration k we select a variable j_k and set

$$w_{j_k}^{k+1} = w_{j_k}^k - \alpha_k \nabla_{j_k} f(w^k),$$

a gradient descent step on coordinate j_k (other w_i stay the same).

• This variation is called coordinate descent (many variations exist).

Why use Coordinate Descent?

- Theoretically, coordinate descent is a provably bad algorithm:
 - The convergence rate is slower than gradient descent.
 - The iteration cost can be similar to gradient descent.
 - Computing 1 partial derivative may have same cost as computing gradient.
- But it is widely-used in practice:
 - Nothing works better for certain problems.
 - Certain fields think it is the "ultimate" algorithm.
- Renewed theoretical interest began with a paper by Nesterov in 2010:
 - Showed global convergence rate for randomized coordinate selection.
 - Coordinate descent is faster than gradient descent if iterations are d times cheaper.

Problems Suitable for Coordinate Optimization

- For what functions is coordinate descent d times faster than gradient descent?
- The simplest example is separable functions,

$$f(w) = \sum_{j=1}^{d} f_j(w_j),$$

- Here f is the sum of an f_j applied to each w_j , like $f(x) = ||w v||^2 = \sum_{i=1}^d (w_j v_i)^2$.
- Cost of gradient descent vs. coordinate descent:
 - Gradient descent costs O(d) to compute each $f'(w_i^k)$.
 - Coordinate descent costs O(1) to compute the one $f_{i_L}'(w_{i_L}^k)$.
- In fact, for separable functions you should only use coordinate optimization.
 - \bullet The variables w_i have "separate" effects, so can be minimized independently.

Problems Suitable for Coordinate Optimization

• A more interesting example is pairwise-separable functions,

$$f(w) = \sum_{i=1}^{d} \sum_{j=1}^{d} f_{ij}(w_i, w_j),$$

which depend on a function of each pair of variables.

- An example is label propagation.
 - Also includes any quadratic function.
- Cost of gradient descent vs. coordinate descent:
 - Gradient descent costs $O(d^2)$ to compute each f'_{ii} .
 - Coordinate descent costs O(d) to compute d values of f'_{ij} .

Problems Suitable for Coordinate Optimization

• Our label propagation example looked a bit more like this:

$$f(w) = \sum_{j=1}^{d} f_j(w_j) + \sum_{(i,j)\in E} f_{ij}(w_i, w_j),$$

where E is a set of (i, j) pairs ("edges" in a graph).

- Adding a separable function doesn't change costs.
 - We could just combine the f_i with one f_{ij} .
- Restricting (i, j) to E makes gradient descent cheaper:
 - Now costs O(|E|) to compute gradient.
 - Coordinate descent could also cost O(|E|) if degree of j_k is O(|E|).
- Coordinate descent is still d times faster in expectation if you randomly pick j_k .
 - Each f'_{ii} is needed with probability 2/d.
 - So expected cost of O(|E|/d) to compute one partial derivative.

• For the binary label propagation objective,

$$\underset{\bar{y} \in \mathbb{R}^t}{\operatorname{argmin}} \sum_{i=1}^n \sum_{j=1}^t w_{ij} (y^i - \bar{y}^j)^2 + \frac{1}{2} \sum_{i=1}^t \sum_{j=1}^t \bar{w}_{ij} (\bar{y}^i - \bar{y}^j)^2,$$

we can exactly optimize one coordinate given the others.

• Take the gradient with respect to a particular \bar{y}^i ,

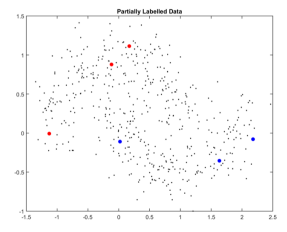
$$\nabla_{\bar{y}^i} f(\bar{y}) = 2 \sum_{j=1}^n w_{ij} (y^j - \bar{y}^i) + 2 \sum_{j=1}^t \bar{w}_{ij} (\bar{y}^i - \bar{y}^j),$$

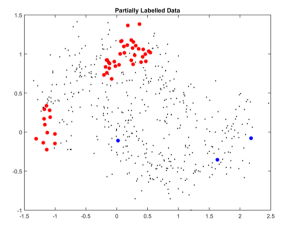
assuming that $\bar{w}_{ij} = \bar{w}_{ii}$ (otherwise, you could replace both by their average).

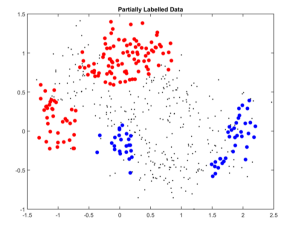
• Solving for \bar{y}^i gives

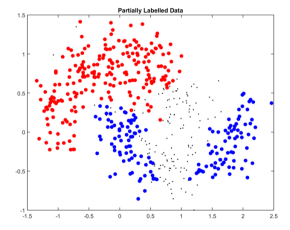
$$\bar{y}^i = \frac{\sum_{j=1}^n w_{ij} y^j + \sum_{j=1}^t \bar{w}_{ij} \bar{y}^j}{\sum_{j=1}^n w_{ij} + \sum_{j=1}^t \bar{w}_{ij}}.$$

• So coordinate optimization takes weighted average of neighbours.

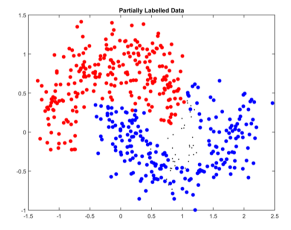




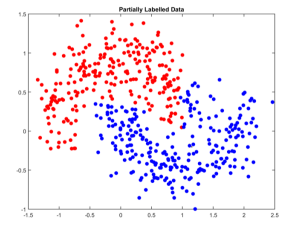




Label Propagation with Coordinate Optimization



Label Propagation with Coordinate Optimization



Analyzing Coordinate Descent

• To analyze coordinate descent, we can write it as

$$w^{k+1} = w^k - \alpha_k e_{j_k} \nabla_{j_k} f(w^k),$$

where "elementary vector" e_i has a zero in every position except j,

$$e_3^T = \begin{bmatrix} 0 & 0 & 1 & 0 & 0 & 0 \end{bmatrix}$$

• We usually assume that each $\nabla_i f$ is L-Lipshitz ("coordinate-wise Lipschitz"),

$$|\nabla_i f(w + \gamma e_i) - \nabla_i f(w)| \le L|\gamma|,$$

which for C^2 functions is equivalent to $|\nabla^2_{ij}f(w)| \leq L$ for all i.

(diagonals of Hessian are bounded)

- This is not a stronger assumption:
 - If the gradient is L-Lipschitz then it's also coordinate-wise L-Lipschitz.

Convergence Rate of Coordinate Optimization

• Coordinate-wise Lipschitz assumption implies a coordinate-wise descent lemma,

$$f(w^{k+1}) \le f(w^k) + \nabla_j f(w^k) (w^{k+1} - w^k)_j + \frac{L}{2} (w^{k+1} - w^k)_j^2,$$

for any w^{k+1} and w^k that only differ in coordinate j.

• Using $\alpha_k=1/L$ (for simplicity), plugging in $(w^{k+1}-w^k)=-(1/L)e_{j_k}\nabla_{j_k}f(w^k)$ gives

$$f(w^{k+1}) \le f(w^k) - \frac{1}{2L} |\nabla_{j_k} f(w^k)|^2,$$

a progress bound based on only updating coordinate j_k .

- If we did optimal update (as in label propagation), this bound would still hold.
 - Optimal update decreases f by at least as much as any other update.

Convergence Rate of Randomized Coordinate Optimization

• Our bound for updating coordinate j_k is

$$f(w^{k+1}) \le f(w^k) - \frac{1}{2L} |\nabla_{j_k} f(w^k)|^2,$$

so progress depends on which j_k that we choose.

• Let's consider expected progress with random selection of j_k ,

$$\begin{split} \mathbb{E}[f(w^{k+1})] &\leq \mathbb{E}\left[f(w^k) - \frac{1}{2L}|\nabla_{j_k}f(w^k)|^2\right] & \text{(expectation wrt } j_k \text{ given } w^k\text{)} \\ &= \mathbb{E}[f(w^k)] - \frac{1}{2L}\mathbb{E}[|\nabla_{j_k}f(w^k)|^2] & \text{(linearity of expectation)} \\ &= f(w^k) - \frac{1}{2L}\sum_{i=1}^d p(j_k = j)|\nabla_j f(w^k)|^2 & \text{(definition of expectation)} \end{split}$$

Convergence Rate of Randomized Coordinate Optimization

• The bound from the previous slide is

$$E[f(w^{k+1})] \le f(w^k) - \frac{1}{2L} \sum_{j=1}^{d} p(j_k = j) |\nabla_j f(w^k)|^2.$$

• Let's choose j_k uniformly in this bound, $p(j_k = j) = 1/d$.

$$\mathbb{E}[f(w^{k+1})] \le f(w^k) - \frac{1}{2L} \sum_{j=1}^d \frac{1}{d} |\nabla_j f(w^k)|^2$$

$$= f(w^k) - \frac{1}{2dL} \sum_{j=1}^d |\nabla_j f(w^k)|^2$$

$$= f(w^k) - \frac{1}{2dL} ||\nabla f(w^k)||^2.$$

Convergence Rate of Randomized Coordinate Optimization

• Our guaranteed progress bound for randomized coordinate optimization,

$$\mathbb{E}[f(w^{k+1}))] \le f(w^k) - \frac{1}{2dL} \|\nabla f(w^k)\|^2.$$

• If we use strongly convexity or PL and recurse carefully (see bonus) we get

$$\mathbb{E}[f(w^k)] - f^* \le \left(1 - \frac{\mu}{dI}\right)^k [f(w^0) - f^*].$$

which means we expect to need $O\left(d\frac{L}{\mu}\log(1/\epsilon)\right)$ iterations.

- Remember that gradient descent needs $O\left(\frac{L}{\mu}\log(1/\epsilon)\right)$ iterations.
- So coordinate optimization needs *d*-times as many iterations?

Randomized Coordinate Optimization vs. Gradient Descent

• If coordinate descent step are d-times cheaper then both algorithms need

$$O\left(\frac{L}{\mu}\log(1/\epsilon)\right),$$

in terms of gradient descent iteration costs.

- So why prefer coordinate optimization?
- The Lipschitz constants L are different.
 - Gradient descent uses L_f and coordinate optimization uses L_c .
 - L_c is maximum gradient changes if you change *one* coordinate.
 - L_f is maximum gradient changes if you change all coordinates.
- Since $L_c \leq L_f$, coordinate optimization is faster.
 - By a factor that could be as large as d.
 - The gain is because coordinate descent allows bigger step-sizes.

Summary

- Transductive learning:
 - Given labeled and unlabeled examples, label the unlabeled examples.
- Label propagation:
 - Transductive learning method minimizing variation in the label space.
- Coordinate optimization: updating one variable at a time.
 - Efficient if updates are *d*-times cheaper than gradient descent.
- Next time: the most important algorithm in machine learning.

Applying Expected Bound Recursively

• Our guaranteed progress bound for randomized coordinate optimization,

$$\mathbb{E}[f(w^{k+1}))] \le f(w^k) - \frac{1}{2dL} \|\nabla f(w^k)\|^2.$$

• If we subtract f^* and use strong-convexity or PL (as before),

$$\mathbb{E}[f(w^{k+1})] - f^* \le \left(1 - \frac{\mu}{dL}\right)[f(w^k) - f^*].$$

• By recursing we get linear convergence rate,

$$\begin{split} \mathbb{E}[\mathbb{E}[f(w^{k+1})]] - f^* &\leq \mathbb{E}\left[\left(1 - \frac{\mu}{dL}\right)[f(w^k) - f^*]\right] \quad \text{(expectation wrt } j_{k-1}) \\ \mathbb{E}[f(w^{k+1})] - f(w^*) &\leq \left(1 - \frac{\mu}{dL}\right)[\mathbb{E}[f(w^k)] - f^*] \quad \text{(iterated expectations)} \\ &\leq \left(1 - \frac{\mu}{dL}\right)^2[f(w^{k-1}) - f^*] \end{split}$$

• You keep alternating between taking an expectation back in time and recursing.