Optimization in the "Big Data" Regime 2: SVRG & Tradeoffs in Large Scale Learning.

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Work on your project milestones

- read/related work summary
- some empirical work

Today:

- Review: optimization of finite sums, (dual) coordinate ascent
- New: SVRG (for sums of loss functions); Tradeoffs in large scale learning How do we optimize in the "big data" regime?

Machine Learning and the Big Data Regime...

goal: find a *d*-dim parameter vector which minimizes the loss on *n* training examples.

- have *n* training examples $(x_1, y_1), \ldots, (x_n, y_n)$
- have parametric a classifier *h*(*x*, *w*), where *w* is a *d* dimensional vector.

$$\min_{w} L(w) \text{ where } L(w) = \sum_{i} \operatorname{loss}(h(x_i, w), y_i)$$

• "Big Data Regime": How do you optimize this when *n* and *d* are large? memory? parallelization?

Can we obtain linear time algorithms to find an ϵ -accurate solution? i.e. find \hat{w} so that

$$L(\hat{w}) - \min_{w} L(w) \leq \epsilon$$

Review: Stochastic Gradient Descent (SGD)



• SGD update rule: at each time t,

sample a point (x_i, y_i) $w \leftarrow w - \eta (w \cdot x_i - y_i) x_i$

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 Problem: even if w = w_{*}, the update changes w.
 Rate: convergence rate is O(1/ε), with decaying η simple algorithm, light on memory, but poor convergence rate • What about more general convex problems? e.g.

$$\min_{w} L(w) \text{ where } L(w) = \sum_{i} \operatorname{loss}(h(x_i, w), y_i)$$

- the basic idea (formalized with duality) is pretty general for convex $\mathrm{loss}(\cdot).$
- works very well in practice.
- memory: SDCA needs O(n + d) memory, while SGD is only O(d).
- What about an algorithm for non-convex problems?
 - SDCA seems heavily tied to the convex case.
 - Is there an algo that is highly accurate in the convex case *and* sensible in the non-convex case?

L smooth and μ -strongly convex case

- Suppose L(w) is μ strongly convex.
- Suppose each loss loss(·) is *L*-smooth
- To get ϵ accuracy:
 - # iterations to get ε-accuracy:

 $\frac{\mathbf{L}}{\mu\epsilon}$

(see related work for precise problem dependent parameters)

• Computation time to get *\epsilon*-accuracy:

$$\frac{L}{\mu\epsilon}a$$

(assuming O(d) cost pre gradient evaluation.)

(another idea) Stochastic Variance Reduced Gradient (SVRG)

1 exact gradient computation: at stage *s*, using \tilde{w}_s , compute:

$$\nabla L(\widetilde{w}_{s}) = \frac{1}{n} \sum_{i=1}^{n} \nabla \operatorname{loss}(h(x_{i}, \widetilde{w}_{s}), y_{i})$$

2 variance reduction + SGD: initialize $w \leftarrow \tilde{w}_s$. for *m* steps,

sample a point (x, y) $w \leftarrow w - \eta \left(\nabla \operatorname{loss}(h(x, w), y) - \nabla \operatorname{loss}(h(x, \widetilde{w}_s), y) + \nabla L(\widetilde{w}_s) \right)$

3 update and repeat: $\widetilde{w}_{s+1} \leftarrow w$.

• unbiased updates: What is the mean of the blue term?

 $\mathbb{E}[\nabla \operatorname{loss}(h(x, \widetilde{w}_s), y) - \nabla L(\widetilde{w}_s)] = ?$

where the expectation is for a random sample (x, y).

- If $\widetilde{w} = w_*$, then no update.
- Memory is O(d).
- No "dual" variables.

Applicable to non-convex optimization.

- set $m = L/\mu$.
- # of gradient computations to get ϵ accuracy:

$$\left(n+\frac{L}{\mu}\right)\log 1/\epsilon$$

- a gradient evaluation is at point (*x*, *y*).
 - SVRG: # of gradient computations to get ϵ accuracy:

$$\left(n+\frac{L}{\mu}\right)\log 1/\epsilon$$

• # of gradient evaluations for batch gradient descent:

$$n rac{ ilde{L}}{\mu} \log 1/\epsilon$$

where L̃ is the smoothness of L(w).
of gradient computations for SGD:

$$\frac{L}{\mu\epsilon}$$

• How many gradient evaluations does it take to find w so that:

$$\|\nabla L(w)\|^2 \le \epsilon^2$$

(i.e. "close" to a stationary point)

- Rates: the number of gradient evaluations, at a point (*x*, *y*), is:
 - GD: O(n/ε)
 - SGD: O(1/ε²)
 - SVRG: $O(n + n^{2/3}/\epsilon)$

Does SVRG work well in practice?

- Many issues sources of "error"
- approximation error: our choice of a hypothesis class
- estimation error: we only have *n* samples
- optimization error: computing exact (or near-exact) minimizers can be costly.
- How do we think about these issues?

The true objective

• hypothesis map $x \in \mathcal{X}$ to $y \in \mathcal{Y}$.

- have *n* training examples $(x_1, y_1), \ldots, (x_n, y_n)$ sampled i.i.d. from \mathcal{D} .
- Training objective: have a set of parametric predictors $\{h(x, w) : w \in W\},\$

$$\min_{w \in \mathcal{W}} \hat{L}_n(w) \text{ where } \hat{L}_n(w) = \frac{1}{n} \sum_{i=1}^n \operatorname{loss}(h(x_i, w), y_i)$$

• True objective: to generalize to \mathcal{D} ,

$$\min_{w \in \mathcal{W}} L(w) \text{ where } L(w) = \mathbb{E}_{(X,Y) \sim \mathcal{D}} \text{loss}(h(X,w),Y)$$

Optimization: Can we obtain linear time algorithms to find an ϵ -accurate solution? i.e. find \hat{h} so that

$$L(\hat{w}) - \min_{w \in \mathcal{W}} L(w) \leq \epsilon$$

• Let h^* is the Bayes optimal hypothesis, over all functions from $\mathcal{X} \to \mathcal{Y}$.

 $h^* \in \operatorname{argmin}_h L(h)$

• Let w* is the best in class hypothesis

 $w^* \in \operatorname{argmin}_{w \in \mathcal{W}} L(w)$

• Let *w_n* be the *empirical risk minimizer*:

 $w_n \in \operatorname{argmin}_{w \in \mathcal{W}} \hat{L}_n(w)$

• Let \tilde{w}_n be what our algorithm returns.

Observe:

$$L(\tilde{w}_n) - L(h^*) = L(w^*) - L(h^*)$$
Approximation error
+ $L(w_n) - L(w^*)$ Estimation error
+ $L(\tilde{w}_n) - L(w_n)$ Optimization error

- Three parts which determine our performance.
- Optimization algorithms with "best" accuracy dependencies on L
 n
 may not be best.
 Forcing one error to decrease much faster may be wasteful.





• Vary the number of examples



• Vary the number of examples



• Optimal combination depends on training time budget.

Measuring a mean:

$$L(\mu) = \mathbb{E}(\mu - y)^2$$

The minima is at $\mu = \mathbb{E}[\mathbf{y}]$.

- With *n* samples, the Bayes optimal estimator is the sample mean: $\hat{\mu}_n = \frac{1}{n} \sum_i y_i$.
- The error is:

$$\mathbb{E}[L(\hat{\mu}_n)] - L(\mathbb{E}[y]) = \frac{\sigma^2}{n}$$

 σ^2 is the variance and the expectation is with respect to the n samples.

• How many samples do we need for ϵ error?

- SGD: Is $O(1/\epsilon)$ reasonable?
- GD: Is log 1/eps needed?
- SDCA/SVRG: These are also log 1/eps but much faster.

- Can generalize as well as the sample minimizer, w_n? (without computing it exactly)
- For a wide class of models (linear regression, logistic regression, etc), we have that the estimation error is:

$$\mathbb{E}[L(w_n)] - L(w^*) = \frac{\sigma_{\text{opt}}^2}{n}$$

where $\sigma_{\rm opt}^2$ is a problem dependent constant.

• What is the computational cost of achieving exactly this rate? say for large n?

SGD:

$$w_{t+1} \leftarrow w_t - \eta_t \nabla \mathrm{loss}(h(x, w_t), y)$$

- An (asymptotically) optimal algo:
 - Have η_t go to 0 (sufficiently slowly)
 - (iterate averaging) Maintain the a running average:

$$\overline{w_n} = \frac{1}{n} \sum_{t \le n} w_t$$

• (Polyak & Juditsky, 1992) for large enough *n* and with one pass of SGD over the dataset:

$$\mathbb{E}[L(\overline{w_n})] - L(w^*) \stackrel{n \to \infty}{=} \frac{\sigma_{\text{opt}}^2}{n}$$

Some slides from "Large-scale machine learning revisited", Leon Bottou 2013.