Stochastic Gradient Descent

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Gradient Descent for Empirical Risk - Scaling Issues

Gradient Descent for Empirical Risk and Averages

- Suppose we have a hypothesis space of functions $\mathcal{F} = \{f_w : \mathcal{X} \to \mathcal{A} \mid w \in \mathsf{R}^d\}$
 - Parameterized by $w \in \mathbb{R}^d$.
- ERM is to find w minimizing

$$\hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^n \ell(f_w(x_i), y_i)$$

- Suppose $\ell(f_w(x_i), y_i)$ is differentiable as a function of w.
- Then we can do gradient descent on $\hat{R}_n(w)$...

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Gradient Descent: How does it scale with n?

• At every iteration, we compute the gradient at current w:

$$\nabla \hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^n \nabla_w \ell(f_w(x_i), y_i)$$

- We have to touch all n training points to take a single step. [O(n)]
- Will this scale to "big data"?
- Can we make progress without looking at all the data?

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Stochastic Gradient Descent

"Noisy" Gradient Descent

- We know gradient descent works.
- But the gradient may be slow to compute.
- What if we just use an estimate of the gradient?
- Turns out that can work fine.
- Intuition:
 - Gradient descent is an interative procedure anyway.
 - At every step, we have a chance to recover from previous missteps.

• The full gradient is

$$\nabla \hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^n \nabla_w \ell(f_w(x_i), y_i)$$

- It's an average over the **full batch** of data $\mathcal{D}_n = \{(x_1, y_1), \dots, (x_n, y_n)\}.$
- Let's take a random subsample of size *N* (called a **minibatch**):

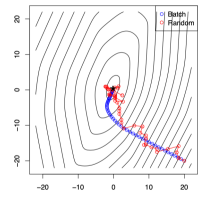
$$(x_{m_1}, y_{m_1}), \ldots, (x_{m_N}, y_{m_N})$$

The minibatch gradient is

$$\nabla \hat{R}_{N}(w) = \frac{1}{N} \sum_{i=1}^{N} \nabla_{w} \ell(f_{w}(x_{m_{i}}), y_{m_{i}})$$

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Batch vs Stochastic Methods



(Slides adapted from Ryan Tibshirani)

Rule-of-thumb for stochastic methods:

- Stochastic methods work well far from the optimum
- But struggles close the the optimum

• What can we say about the minibatch gradient? It's random. What's its expectation?

$$\mathbb{E}\left[\nabla \hat{R}_{N}(w)\right] = \frac{1}{N} \sum_{i=1}^{N} \mathbb{E}\left[\nabla_{w} \ell(f_{w}(x_{m_{i}}), y_{m_{i}})\right]$$

$$= \mathbb{E}\left[\nabla_{w} \ell(f_{w}(x_{m_{1}}), y_{m_{1}})\right]$$

$$= \sum_{i=1}^{n} \mathbb{P}(m_{1} = i) \nabla_{w} \ell(f_{w}(x_{i}), y_{i})$$

$$= \frac{1}{n} \sum_{i=1}^{n} \nabla_{w} \ell(f_{w}(x_{i}), y_{i})$$

$$= \nabla \hat{R}_{n}(w)$$

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Minibatch Gradient Properties

• Minibatch gradient is an unbiased estimator for the [full] batch gradient:

$$\mathbb{E}\left[\nabla\hat{R}_{N}(w)\right] = \nabla\hat{R}_{n}(w)$$

• The bigger the minibatch, the better the estimate.

$$\frac{1}{N}\mathsf{Var}\left[\nabla\hat{R}_1(w)\right] = \mathsf{Var}\left[\nabla\hat{R}_N(w)\right]$$

- Tradeoffs of minibatch size:
 - Bigger $N \implies$ Better estimate of gradient, but slower (more data to touch)
 - Smaller $N \implies$ Worse estimate of gradient, but can be quite fast

Convergence of SGD

- For convergence guarantee, use **diminishing step sizes**, e.g. $\eta_k = 1/k$ (dampens noise in step direction)
- Theoretically, GD is much faster than SGD in terms of convergence rate
 - much faster to add a digit of accuracy on the minimum
 - but most of that benefit happens once you're already pretty close
- However, in many ML problems we don't care about optimizing to high accuracy

Step Sizes in Minibatch Gradient Descent

Minibatch Gradient Descent (minibatch size N)

- initialize w = 0
- repeat
 - randomly choose N points $\{(x_i, y_i)\}_{i=1}^N \subset \mathcal{D}_n$

•
$$w \leftarrow w - \eta \left[\frac{1}{N} \sum_{i=1}^{N} \nabla_{w} \ell(f_{w}(x_{i}), y_{i}) \right]$$

- For SGD, fixed step size can work well in practice.
- Typical approach: Fixed step size reduced by constant factor whenever validation performance stops improving.
- Other tricks: Bottou (2012), "Stochastic gradient descent tricks"

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Summary

- Gradient descent or "full-batch" gradient descent
 - Use full data set of size *n* to determine step direction
- Minibatch gradient descent
 - ullet Use a random subset of size N to determine step direction
- Stochastic gradient descent
 - Minibatch with N=1.
 - Use a single randomly chosen point to determine step direction.

These days terminology isn't used so consistently, so always clarify the [mini]batch size.

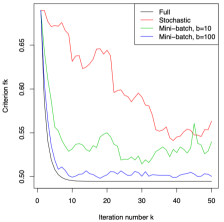
SGD is much more efficient in time and memory cost and has been quite successful in large-scale ML.

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Practical Comparison of GD vs SGD

Logistic regression with ℓ_2 regularization

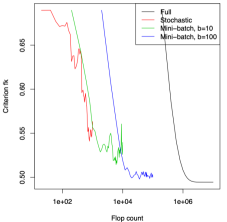
Batch methods converge faster



(Example from Ryan Tibshirani)

Logistic regression with ℓ_2 regularization

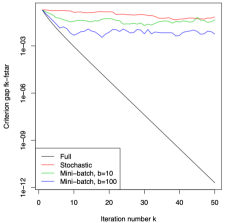
Stochastic methods are computationally more efficient



(Example from Ryan Tibshirani)

Logistic regression with ℓ_2 regularization

Batch methods are much faster close to the optimum



(Example from Ryan Tibshirani)

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