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The gradient descent acts like follows:

$$x_{k+1} = x_k - \frac{\alpha_k}{n} \sum_{i=1}^n \nabla f_i(x)$$
 (GD)

• Iteration cost is linear in n.

 $f \to \min_{x,y,.}$ 

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 $x_{k+1} = x_k - \frac{\alpha_k}{n} \sum_{i=1}^{n} \nabla f_i(x)$ 

Let's/ switch from the full gradient calculation to its unbiased estimator, when we randomly choose 
$$i_k$$
 index of point at each iteration uniformly:

- $x_{k+1} = x_k \alpha_k \nabla f_i$ ,  $(x_k)$
- With  $p(i_k = i) = \frac{1}{n}$ , the stochastic gradient is an unbiased estimate of the gradient, given by:

 $\mathbb{E}[\nabla f_{i_k}(x)] = \sum_{i=1}^n p(i_k = i) \nabla f_i(x) = \sum_{i=1}^n \frac{1}{n} \nabla f_i(x) = \frac{1}{n} \sum_{i=1}^n \nabla f_i(x) = \nabla f(x)$ 

This indicates that the expected value of the stochastic gradient is equal to the actual gradient of f(x).  $f \to \min_{x,y,z}$  Finite-sum problem



(GD)

(SGD)

Stochastic iterations are n times faster, but how many iterations are needed?

If  $\nabla f$  is Lipschitz continuous then we have:

Assumption	Deterministic Gradient Descent	Stochastic Gradient Descent
PL	$O(\log(1/arepsilon))$	O(1/arepsilon)
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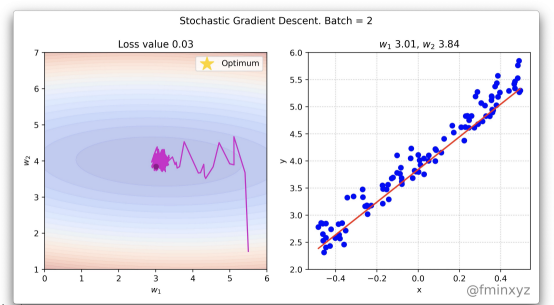
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- Momentum and Quasi-Newton-like methods do not improve rates in stochastic case. Can only improve constant factors (bottleneck is variance, not condition number).



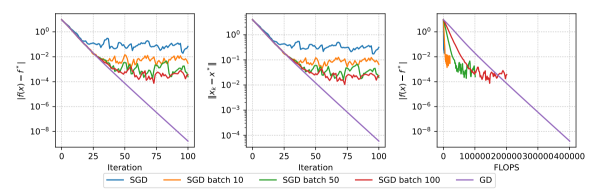
### SGD with constant stepsize does not converge



#### Main problem of SGD

$$f(x) = \frac{\mu}{2} ||x||_2^2 + \frac{1}{m} \sum_{i=1}^m \log(1 + \exp(-y_i \langle a_i, x \rangle)) \to \min_{x \in \mathbb{R}^n}$$

Strongly convex binary logistic regression. m=200, n=10, mu=1.





**Principle:** reducing variance of a sample of X by using a sample from another random variable Y with known expectation:

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#### Application to gradient estimation?

• SVRG: Let  $X = \nabla f_{i_k}(x^{(k-1)})$  and  $Y = \nabla f_{i_k}(\tilde{x})$ , with  $\alpha = 1$  and  $\tilde{x}$  stored.

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- $X Y = \nabla f_i$ ,  $(x^{(k-1)}) \nabla f_i$ ,  $(\tilde{x})$

• Maintain table, containing gradient  $g_i$  of  $f_i$ , i = 1, ..., n

 $f \to \min_{x,y,z}$  Variance reduction methods

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Set all other  $g_i^{(k)} = g_i^{(k-1)}$ ,  $i \neq i_k$ , i.e., these stay the same

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- SAG gradient estimates are no longer unbiased, but they have greatly reduced variance
- Isn't it expensive to average all these gradients? Basically just as efficient as SGD, as long we're clever:

$$x^{(k)} = x^{(k-1)} - \alpha_k \underbrace{\left(\frac{1}{n}g_i^{(k)} - \frac{1}{n}g_i^{(k-1)} + \underbrace{\frac{1}{n}\sum_{i=1}^n g_i^{(k-1)}}_{\text{old table average}}\right)}_{\text{old table average}}$$

new table average

Assume that  $f(x) = \frac{1}{n} \sum_{i=1}^{n} f_i(x)$ , where each  $f_i$  is differentiable, and  $\nabla f_i$  is Lipschitz with constant L.

Denote  $\bar{x}^{(k)} = \frac{1}{k} \sum_{l=0}^{k-1} x^{(l)}$ , the average iterate after k-1 steps.

#### Theorem

SAG, with a fixed step size  $\alpha = \frac{1}{16L}$ , and the initialization

$$g_i^{(0)} = \nabla f_i(x^{(0)}) - \nabla f(x^{(0)}), \quad i = 1, \dots, n$$

satisfies

$$\mathbb{E}[f(\bar{x}^{(k)})] - f^* \le \frac{48n}{k} [f(x^{(0)}) - f^*] + \frac{128L}{k} ||x^{(0)} - x^*||^2$$

where the expectation is taken over random choices of indices.

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- GD:  $\frac{L\|x^{(0)}-x^{\star}\|^2}{2L}$



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- So the first term in SAG bound suffers from a factor of n; authors suggest smarter initialization to make  $f(x^{(0)}) f^*$  small (e.g., they suggest using the result of n SGD steps).

 $f \to \min_{x,y}$ 

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Assume further that each  $f_i$  is strongly convex with parameter  $\mu$ .

Theorem

SAG, with a step size  $\alpha = \frac{1}{16L}$  and the same initialization as before, satisfies

$$\mathbb{E}[f(x^{(k)})] - f^{\star} \le \left(1 - \min\left(\frac{\mu}{16L}, \frac{1}{8n}\right)\right)^{k} \left(\frac{3}{2} \left(f(x^{(0)}) - f^{\star}\right) + \frac{4L}{n} \|x^{(0)} - x^{\star}\|^{2}\right)$$

#### Notes:

• This is linear convergence rate  $\mathcal{O}(\gamma^k)$  for SAG. Compare this to  $\mathcal{O}(\gamma^k)$  for GD, and only  $\mathcal{O}\left(\frac{1}{k}\right)$  for SGD.

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### **SAG** convergence

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- Like GD, we say SAG is adaptive to strong convexity.
- Proofs of these results not easy: 15 pages, computed-aided!



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- For the generalized linear models (this includes LogReg, LLS) you need to store much less memory  $\mathcal{O}(n)$ instead of  $\mathcal{O}(pn)$ .

$$f_i(w) = \varphi(w^T x_i) \leftrightarrow \nabla f_i(w) = \varphi'(w^T x_i) x_i$$



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- When selecting components with a probability proportional to  $L_i$ , the constant L can be reduced from  $\max_i L_i$ to  $\bar{L} = \sum_{i} L_i/N$ :

$$g(x) = \frac{1}{n} \sum_{i=1}^{n} f_i(x)$$

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- To generate with probabilities  $L_i/\sum_i L_j$ , there is an algorithm with complexity  $O(\log N)$ .

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- Two gradient evaluations per inner step.
- Two parameters: length of epochs + step-size  $\gamma$ .
- Linear convergence rate, simple proof.



Very popular adaptive method. Let  $g^{(k)} = \nabla f_{i_k}(x^{(k-1)})$ , and update for  $j = 1, \dots, p$ :

$$v_j^{(k)} = v_j^{k-1} + (g_j^{(k)})^2$$
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- Main weakness is the monotonic accumulation of gradients in the denominator. AdaDelta, Adam, AMSGrad, etc. improve on this, popular in training deep neural networks.
- The constant  $\epsilon$  is typically set to  $10^{-6}$  to ensure that we do not suffer from division by zero or overly large step

sizes.

## RMSProp (Tieleman and Hinton, 2012)

An enhancement of AdaGrad that addresses its aggressive, monotonically decreasing learning rate. Uses a moving average of squared gradients to adjust the learning rate for each weight. Let  $g^{(k)} = \nabla f_{ik}(x^{(k-1)})$  and update rule for  $j = 1, \ldots, p$ :

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- Allows for a more nuanced adjustment of learning rates than AdaGrad, making it suitable for non-stationary problems.
- Commonly used in training neural networks, particularly in recurrent neural networks.

## Adadelta (Zeiler, 2012)

An extension of RMSProp that seeks to reduce its dependence on a manually set global learning rate. Instead of accumulating all past squared gradients, Adadelta limits the window of accumulated past gradients to some fixed size w. Update mechanism does not require learning rate  $\alpha$ :

$$\begin{split} v_j^{(k)} &= \gamma v_j^{(k-1)} + (1-\gamma) (g_j^{(k)})^2 \\ \tilde{g}_j^{(k)} &= \frac{\sqrt{\Delta x_j^{(k-1)} + \epsilon}}{\sqrt{v_j^{(k)} + \epsilon}} g_j^{(k)} \\ x_j^{(k)} &= x_j^{(k-1)} - \tilde{g}_j^{(k)} \\ \Delta x_j^{(k)} &= \rho \Delta x_j^{(k-1)} + (1-\rho) (\tilde{g}_j^{(k)})^2 \end{split}$$

#### Notes:

Adadelta adapts learning rates based on a moving window of gradient updates, rather than accumulating all
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- The method does not require an initial learning rate setting, making it easier to configure.
- Often used in deep learning where parameter scales differ significantly across layers.





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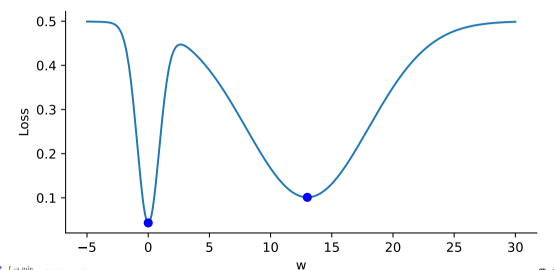
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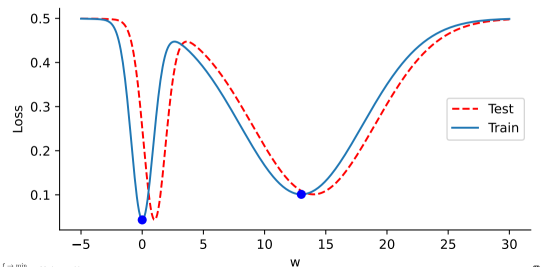
### Wide vs narrow local minima

Узкие и широкие локальные минимумы



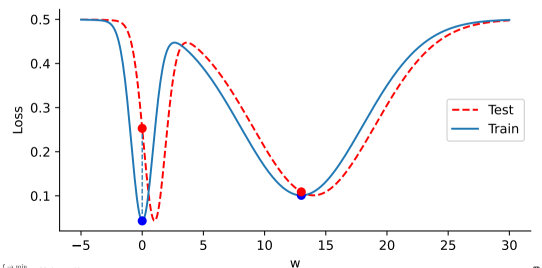
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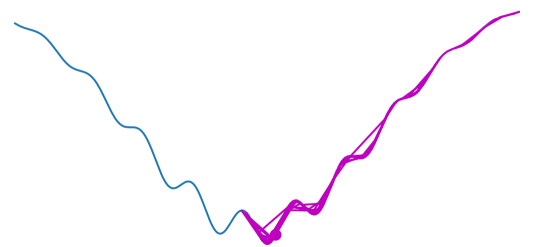
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## Stochasticity allows to escape local minima

Стохастический градиентный спуск выпрыгивает из локальных минимумов



## Local divergence can also be benefitial

Градиентный спуск с большим шагом избегает узкого локального минимума

