

» Polyak Step Size¹

- * *Polyak choice of step size:* $\alpha = \frac{f(x) - f^*}{\nabla f(x)^T \nabla f(x)}$
with $\nabla f(x)^T \nabla f(x) = \sum_{i=1}^n \frac{\partial f}{\partial x_i}(x)^2$ and $f^* = \min_x f(x)$
- * Where does this come from? Recall $f(x + \delta) \approx f(x) + \nabla f(x)^T \delta$.
Choosing $\delta = -\alpha \nabla f(x)$ then

$$f(x - \alpha \nabla f(x)) \approx f(x) - \alpha \nabla f(x)^T \nabla f(x)$$

- * Choosing $\alpha = \frac{f(x) - f^*}{\nabla f(x)^T \nabla f(x)}$ then

$$f(x - \alpha \nabla f(x)) \approx f(x) - \frac{(f(x) - f^*)}{\nabla f(x)^T \nabla f(x)} \nabla f(x)^T \nabla f(x) = f^*$$

- * Scaling step size with $\frac{1}{\nabla f(x)^T \nabla f(x)}$ makes lots of sense \rightarrow when gradient is large the step size is small, when the gradient is small the step size is large.
- * Don't really need to know f^* , can estimate it or just using $f^* = 0$ can work well in ML problems.
- * Much lower computational burden than line search ...

¹Originally proposed in Polyak's book, for more recent work see e.g. <https://arxiv.org/pdf/1905.00313.pdf>

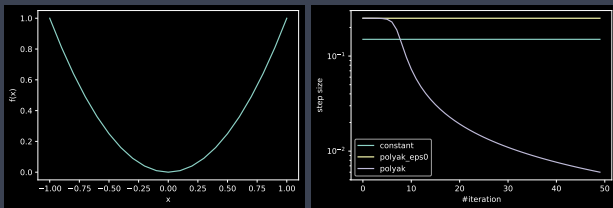
» Polyak Step Size

- * Polyak choice of step size: $\alpha = \frac{f(x) - f^*}{\nabla f(x)^T \nabla f(x)}$
- * When close to minimum then expect $\nabla f(x)$ to be close to zero, and so $\nabla f(x)^T \nabla f(x) \approx 0$. To avoid division by zero, in practice tend to modify Polyak step size to

$$\alpha = \frac{f(x) - f^*}{\nabla f(x)^T \nabla f(x) + \epsilon}$$

where ϵ is a small number e.g. 0.001

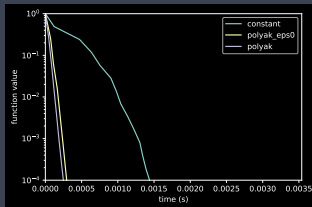
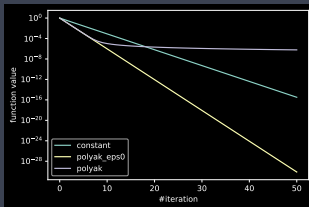
- * Example: $f = x^2$, starting point $x = 1$, $f^* = 0$, $\epsilon = 0.0001$ and $\epsilon = 0$



- * With $\epsilon = 0$ Polyak step size is constant, with $\epsilon = 0.0001$ Polyak step decreases as get close to minimum. Why?

» Polyak Step Size

- * Example: $f = x^2$, starting point $x = 1$, $f^* = 0$, $\epsilon = 0.0001$ and $\epsilon = 0$



- * $df/dx = 2x \rightarrow$ Polyak choice is $\alpha = \frac{f(x) - f^*}{\nabla f(x)^T \nabla f(x) + \epsilon} = \frac{x^2 - 0}{(2x)^2 + \epsilon} = \frac{x^2}{4x^2 + \epsilon}$

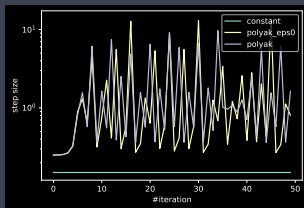
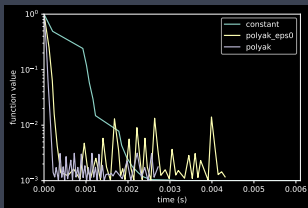
When $\epsilon = 0$, $\alpha = \frac{x^2}{4x^2} = 0.25 \rightarrow$ constant

When $\epsilon = 0.0001$ then when $x^2 \approx 0$ close to minimum $\alpha \approx \frac{x^2}{\epsilon} \approx 0$.

- * Using non-zero ϵ affects behaviour close to minimum, but effect is small if ϵ is small (remember in ML we usually don't need to get v close to min, just close to the noise floor).

» Polyak Step Size

- * Suppose we didn't know f^* exactly but used a lower value. NB: Shouldn't use an f^* larger than true min coz then $f(x) - f^* < 0$ and we might end up with $\alpha < 0$.
- * Example: $f = x^2 + 0.001$, min is now $f(0) = 0.001$ when $x = 0$. Starting point $x = 1$, Polyak $f^* = 0$, $\epsilon = 0.0001$ and $\epsilon = 0$

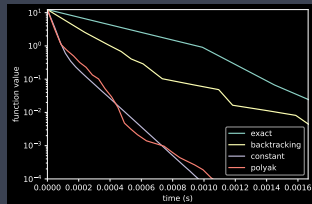
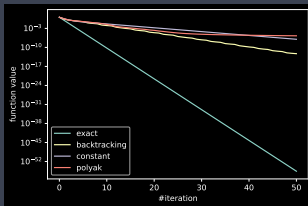


- * Polyak converges quickly, but then step size becomes large and “noisy”
- * When $\epsilon = 0$, $\alpha = \frac{x^2 + 0.001}{4x^2}$. When $x^2 \approx 0$ then $\alpha = \frac{0.001}{4x^2}$ so badly behaved
When $\epsilon = 0.0001$ then when $x^2 \approx 0$ then $\alpha \approx \frac{x^2 + 0.001}{\epsilon} \approx \frac{0.001}{0.0001} = 10$, so too large
- * Might choose to place upper limit on step size?

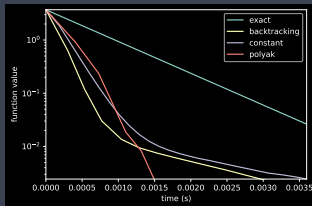
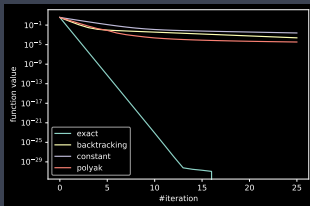
» Examples

Polyak $\epsilon = 0.0001$, $f^* = 0$

* Quadratic:



* Quadratic loss:

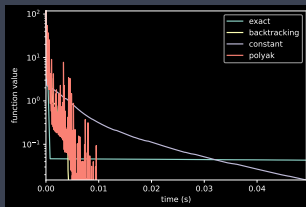
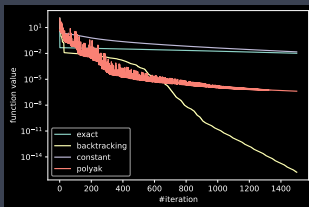


* Since added computational burden of Polyak is low, its competitive with line search wrt convergence vs wall-clock time

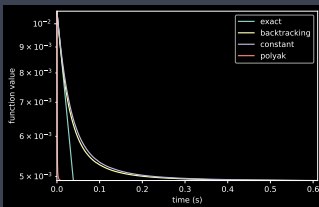
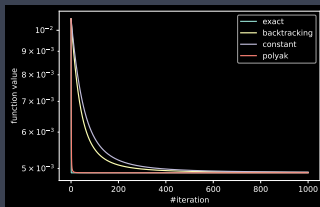
» Examples

Polyak $\epsilon = 0.0001$, $f^* = 0$

- * Rosenbrock function:



- * Toy neural net loss:

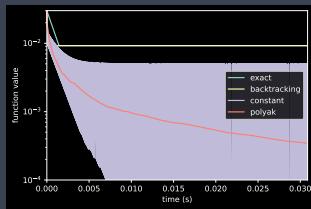
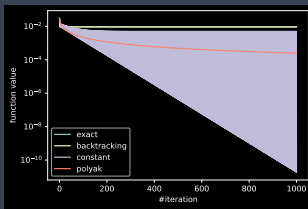


- * Again, Polyak competitive with line search wrt convergence vs wall-clock time
- * Hasn't been much work looking at Polyak for ML

» Examples

Polyak $\epsilon = 0.0001$, $f^* = 0$

- * Non-smooth function $f(x) = |x_1| + x_2^2$



- * Note: Polyak hasn't received much attention for ML \rightarrow but ML usually uses SGD, will come on to this shortly

» Adagrad²

- * Another idea for adapting step size:

$$x_0 = x_0; \alpha_0 = \alpha_0; \text{sum} = \epsilon; t = 0$$

for k in range(num_iters):

$$x_{t+1} = x_t - \alpha_t \frac{df}{dx}(x_t)$$

$$\text{sum} = \text{sum} + \frac{df}{dx}(x_t)^2$$

$$\alpha_{t+1} = \text{alpha}_0 / \text{sum}$$

$$t = t + 1$$

i.e. step size at iteration t is:

$$\alpha_t = \frac{\alpha_0}{\sqrt{\frac{df}{dx}(x_0)^2 + \frac{df}{dx}(x_1)^2 + \dots + \frac{df}{dx}(x_{t-1})^2 + \epsilon}} = \frac{\alpha_0}{\sqrt{\sum_{i=1}^{t-1} \frac{df}{dx}(x_i)^2 + \epsilon}}$$

- * Similar to Polyak, but now divide by *sum* of gradients rather than just by last gradient. Still have to manually select α_0 .
- * Idea: (i) step size should get smaller over time (why?) and (ii) should be larger/smaller depending on past gradients.

²<https://www.jmlr.org/papers/volume12/duchi11a/duchi11a.pdf>, see Section 1.3.1 in particular. Quite mathematical ...

» Adagrad

- * In general, \mathbf{x} is a vector and $\nabla f(\mathbf{x}_t) = [\frac{\partial f}{\partial x_1}(\mathbf{x}), \frac{\partial f}{\partial x_2}(\mathbf{x}), \dots, \frac{\partial f}{\partial x_n}(\mathbf{x})]$ is a vector. So far we've used a single step size:

$$\mathbf{x}_{t+1} = \mathbf{x}_t - \alpha \left[\frac{\partial f}{\partial x_1}(\mathbf{x}), \frac{\partial f}{\partial x_2}(\mathbf{x}), \dots, \frac{\partial f}{\partial x_n}(\mathbf{x}) \right]$$

- * But Adagrad uses a vector of step sizes $[\alpha_1, \alpha_2, \dots, \alpha_n]$ and:

$$\mathbf{x}_{t+1} = \mathbf{x}_t - \left[\alpha_1 \frac{\partial f}{\partial x_1}(\mathbf{x}), \alpha_2 \frac{\partial f}{\partial x_2}(\mathbf{x}), \dots, \alpha_n \frac{\partial f}{\partial x_n}(\mathbf{x}) \right]$$

with

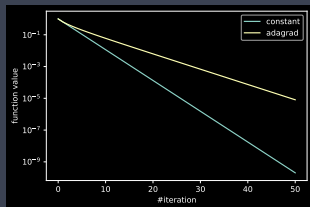
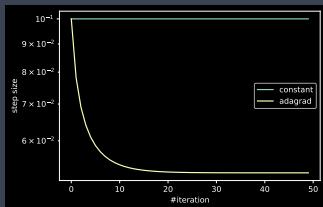
$$\alpha_1 = \frac{\alpha_0}{\sqrt{\sum_{i=1}^{t-1} \left(\frac{\partial f}{\partial x_1}(\mathbf{x}_i) \right)^2 + \epsilon}}$$

$$\alpha_2 = \frac{\alpha_0}{\sqrt{\sum_{i=1}^{t-1} \left(\frac{\partial f}{\partial x_2}(\mathbf{x}_i) \right)^2 + \epsilon}}$$

etc

» Examples

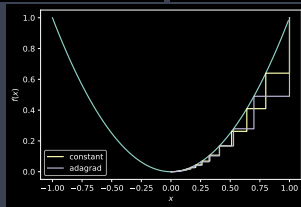
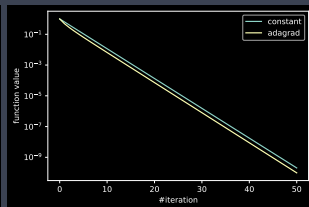
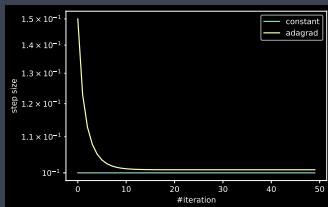
- * Example: $f = x^2$, starting point $x = 1$, Adagrad $\alpha_0 = 0.2$ (selected so that initial step size is 0.1, see below), $\epsilon = 1.0e - 5$, constant $\alpha = 0.1$



- * Adagrad and constant step size initially the same. Adagrad decreases step size, quickly at first (when df/dx is larger) then much more slowly as close to minimum df/dx is small

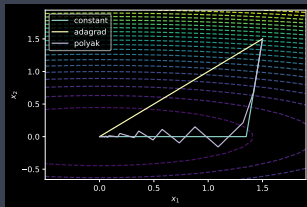
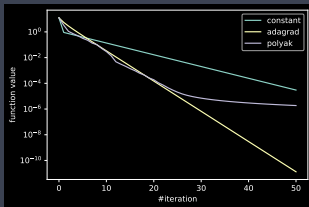
» Examples

- * Adjust α_0 for Adagrad so that later step size matches that constant step size strategy ...
- * Example: $f = x^2$, starting point $x = 1$, Adagrad $\alpha_0 = 0.3$ (larger now), $\epsilon = 1.0e - 5$, constant $\alpha = 0.1$

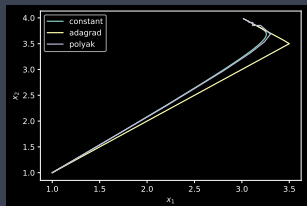
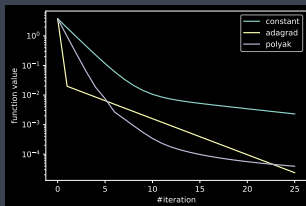


» Examples

- * Quadratic: Adagrad $\alpha_0 = 0.5$, $\epsilon = 1.0e - 5$ (note larger value of α_0), constant $\alpha = 0.1$



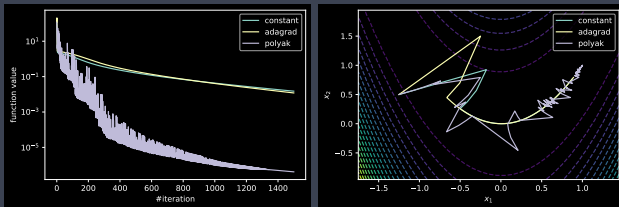
- * Quadratic loss: Adagrad $\alpha_0 = 2.5$, $\epsilon = 1.0e - 5$ (note huge value of α_0), constant $\alpha = 0.5$



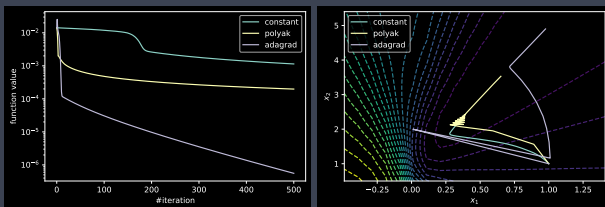
- * Important to tune Adagrad initial step size α_0 , but with tuning can get fast convergence.

» Examples

- * Rosenbrock function: Adagrad $\alpha_0 = 1$, $\epsilon = 1.0e - 5$, constant $\alpha = 0.002$



- * Toy neural net loss: Adagrad $\alpha_0 = 2.5$, $\epsilon = 1.0e - 5$, constant $\alpha = 0.5$

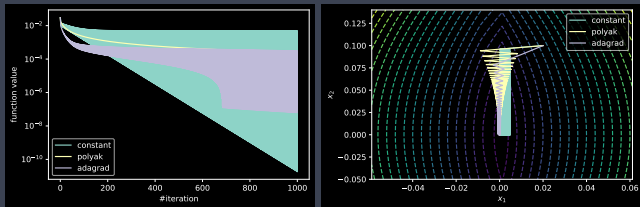


- * Even with tuning α_0 Adagrad isn't faster than constant step size strategy on Rosenbrock function. But fast convergence for neural net loss function.

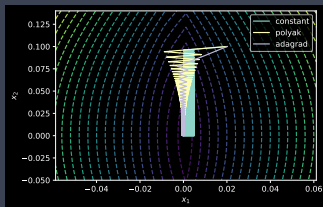
» Examples

Adagrad $\alpha_0 = 0.01, \epsilon = 1.0e - 5$, constant $\alpha = 0.005$

- * Non-smooth function $f(x) = |x_1| + x_2^2$



- * Recall that kink in cost function tends to cause “chattering” when use constant step size close to minimum, by decreasing the step size Adagrad reduces this.



» RMSprop³

- * Adagrad can decrease the step size too aggressively (that's why we needed to make α_0 large), making convergence slow. RMSprop is a tweak to try to address that.
- * Adagrad:

$$\alpha_t = \frac{\alpha_0}{\sqrt{\frac{df}{dx}(x_0)^2 + \frac{df}{dx}(x_1)^2 + \dots + \frac{df}{dx}(x_{t-1})^2 + \epsilon}}$$

- * RMSprop: Gradually forget past values:

$$\alpha_t = \frac{\alpha_0}{\sqrt{(1 - \beta)\beta^t \frac{df}{dx}(x_0)^2 + (1 - \beta)\beta^{t-1} \frac{df}{dx}(x_1)^2 + \dots + (1 - \beta)\frac{df}{dx}(x_{t-1})^2 + \epsilon}}$$

with $0 < \beta \leq 1$. Implement using:

$x_0 = x0$; $\alpha_0 = \alpha0$; $sum = 0$; $t = 0$

for k in range(num_iters):

$$x_{t+1} = x_t - \alpha_t \frac{df}{dx}(x_t)$$

$$sum = \beta sum + (1 - \beta) \frac{df}{dx}(x_t)^2$$

$$\alpha_{t+1} = \alpha0 / (\sqrt{sum} + \epsilon)$$

$$t = t + 1$$

- * Have to manually select α_0 and β .

» RMSprop

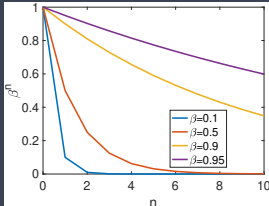
$$* \alpha_t = \frac{\alpha_0}{\sqrt{(1-\beta)(\beta^t \frac{df}{dx}(x_0)^2 + \beta^{t-1} \frac{df}{dx}(x_1)^2 + \dots + \frac{df}{dx}(x_{t-1})^2) + \epsilon}}$$

Implement denominator using:

$$sum = \beta sum + (1 - \beta) \frac{df}{dx}(x_t)^2$$

and then use $\sqrt{sum} + \epsilon$

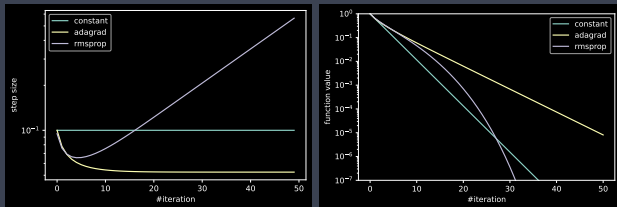
* How to choose β ?



* Typically choose $\beta = 0.9$ or larger. For small β RMSprop reverts to Adagrad.

» Examples

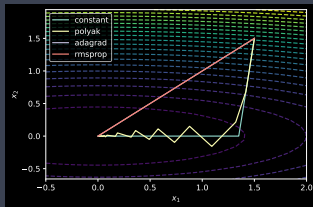
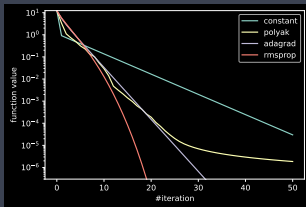
- * Example: $f = x^2$, starting point $x = 1$, RMSprop $\alpha_0 = 0.06$, $\beta = 0.9$, Adagrad $\alpha_0 = 0.2$, $\epsilon = 1.0e - 5$, constant $\alpha = 0.1$



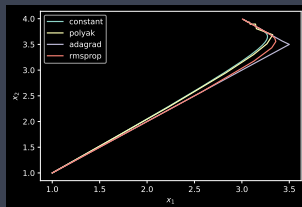
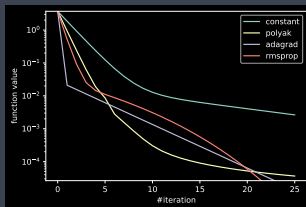
- * Adagrad, RMSprop and constant step size initially the same. RMSprop eventually increases step size - why?

» Examples

- * Quadratic: RMSprop $\alpha_0 = 0.15$, $\beta = 0.9$, $\epsilon = 1.0e - 5$, Adagrad $\alpha_0 = 0.5$, constant $\alpha = 0.1$



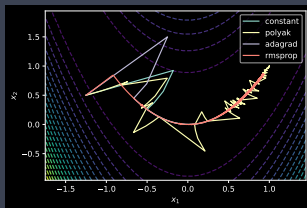
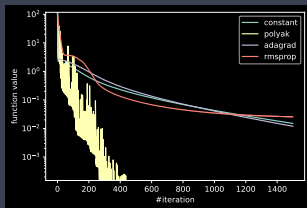
- * Quadratic loss: RMSprop $\alpha_0 = 0.5$, $\beta = 0.9$, $\epsilon = 1.0e - 5$, Adagrad $\alpha_0 = 2.5$, constant $\alpha = 0.5$



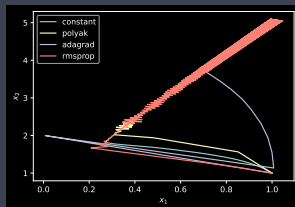
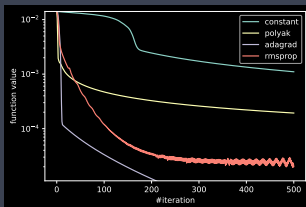
- * Adagrad, RMSprop, Polyak all pretty similar. Note need to tune α_0 for Adagrad, RMSprop.

» Examples

- * Rosenbrock function: RMSprop $\alpha_0 = 0.01$, $\beta = 0.9$, $\epsilon = 1.0e - 5$, Adagrad $\alpha_0 = 1$, constant $\alpha = 0.002$



- * Toy neural net loss: RMSprop $\alpha_0 = 0.05$, $\beta = 0.9$, $\epsilon = 1.0e - 5$, Adagrad $\alpha_0 = 1$, constant $\alpha = 0.75$

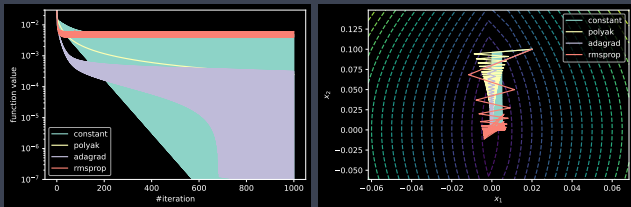


- * Even with tuning α_0 RMSprop/Adagrad are not faster than constant step size strategy on Rosenbrock function. But fast convergence for neural net loss function.

» Examples

RMSprop $\alpha_0 = 0.01$, $\beta = 0.9$, $\epsilon = 1.0e - 5$, Adagrad $\alpha_0 = 0.01$, constant $\alpha = 0.005$

- * Non-smooth function $f(x) = |x_1| + x_2^2$



- * RMSprop less effective than Adagrad at reducing chattering close to minimum of non-smooth function (doesn't reduce step size enough).

» Per co-ordinate step size

- * Adagrad/RMSprop use a vector of step sizes $[\alpha_1, \alpha_2, \dots, \alpha_n]$ and:

$$\mathbf{x}_{t+1} = \mathbf{x}_t - [\alpha_1 \frac{\partial f}{\partial x_1}(\mathbf{x}), \alpha_2 \frac{\partial f}{\partial x_2}(\mathbf{x}), \dots, \alpha_n \frac{\partial f}{\partial x_n}(\mathbf{x})]$$

- * Other classes of approaches that also use a vector α include *Newton* methods and *quasi-Newton* methods.
- * Newton methods use the second-derivative (the derivative of the derivative) to calculate α . For $\nabla f(\mathbf{x}_t) = [\frac{\partial f}{\partial x_1}(\mathbf{x}), \frac{\partial f}{\partial x_2}(\mathbf{x}), \dots, \frac{\partial f}{\partial x_n}(\mathbf{x})]$ then second derivative is a matrix (a 2D array):

$$\begin{pmatrix} \frac{\partial}{\partial x_1} \frac{\partial f}{\partial x_1}(\mathbf{x}) & \frac{\partial}{\partial x_2} \frac{\partial f}{\partial x_1}(\mathbf{x}) & \dots & \frac{\partial}{\partial x_n} \frac{\partial f}{\partial x_1}(\mathbf{x}) \\ \frac{\partial}{\partial x_1} \frac{\partial f}{\partial x_2}(\mathbf{x}) & \frac{\partial}{\partial x_2} \frac{\partial f}{\partial x_2}(\mathbf{x}) & \dots & \frac{\partial}{\partial x_n} \frac{\partial f}{\partial x_2}(\mathbf{x}) \\ \vdots & \vdots & \ddots & \vdots \end{pmatrix}$$

- * Expensive to calculate. Quasi-Newton methods (e.g. *BFGS*) try to approx second derivative more efficiently, but are still computationally costly \rightarrow only suitable for “small” data. Newton/quasi-Newton methods can accelerate convergence vs #iterations but can be slower wrt wall-clock time
- * Can perform poorly on non-convex problems (may not converge at all e.g. see Fig 1 in <https://arxiv.org/pdf/1606.01885v1.pdf>, <https://epubs.siam.org/doi/10.1137/S1052623401383455>)

» Summary

- * Polyak choice of step size works pretty well in examples here, but not well studied for ML problems
- * Adagrad is known to suffer from slow convergence due to decreasing step size too quickly. RMSprop tries to patch up Adagrad. Sometimes helps, sometimes doesn't in examples here
- * Adagrad and RMSprop are implemented in common ML libraries, but Polyak isn't ... hmm?
- * In ML its normal to use SGD, we'll come to this later