# **Optimization in Machine Learning**

# **First order methods SGD**



#### **Learning goals**

- $\bullet$  SGD
- Stochasticity
- **•** Convergence
- Batch size



# **STOCHASTIC GRADIENT DESCENT**

NB: We use *g* instead of *f* as objective, bc. *f* is used as model in ML.

 $q: \mathbb{R}^d \to \mathbb{R}$  objective, *g* **average over functions**:

$$
g(\mathbf{x}) = \frac{1}{n} \sum_{i=1}^{n} g_i(\mathbf{x}), \qquad g \text{ and } g_i \text{ smooth}
$$

Stochastic gradient descent (SGD) approximates the gradient

$$
\nabla_{\mathbf{x}} g(\mathbf{x}) = \frac{1}{n} \sum_{i=1}^{n} \nabla_{\mathbf{x}} g_i(\mathbf{x}) \quad := \quad \mathbf{d} \quad \text{by}
$$
\n
$$
\frac{1}{|J|} \sum_{i \in J} \nabla_{\mathbf{x}} g_i(\mathbf{x}) \quad := \quad \hat{\mathbf{d}},
$$

with random subset *J* ⊂ {1, 2, ..., *n*} of gradients called **mini-batch**. This is done e.g. when computing the true gradient is **expensive**.



# **STOCHASTIC GRADIENT DESCENT / 2**

#### **Algorithm** Basic SGD pseudo code

- 1: Initialize  $\mathbf{x}^{[0]}$ ,  $t=0$
- 2: **while** stopping criterion not met **do**
- 3: Randomly shuffle indices and partition into minibatches  $J_1, \ldots, J_K$  of size m
- 4: **for** *k* ∈ {1, ..., *K*} **do**
- 5:  $t \leftarrow t + 1$
- 6: Compute gradient estimate with  $J_k$ :  $\hat{\mathbf{d}}^{[t]} \leftarrow \frac{1}{m} \sum_{i \in J_k} \nabla_{\mathbf{x}} g_i(\mathbf{x}^{[t-1]})$
- 7: Apply update:  $\mathbf{x}^{[t]} \leftarrow \mathbf{x}^{[t-1]} \alpha \cdot \hat{\mathbf{d}}^{[t]}$
- 8: **end for**
- 9: **end while**
	- **Instead of drawing batches randomly we might want to go through the**  $g_i$ sequentially (unless *g<sup>i</sup>* are sorted in any way)
	- Updates are computed faster, but also more stochastic:
		- In the simplest case, batch-size  $m := |J_k|$  is set to  $m = 1$
		- If *n* is a billion, computation of update is a billion times faster
		- **But** (later): Convergence rates suffer from stochasticity!

# **SGD IN ML**

In ML, we perform ERM:

$$
\mathcal{R}(\boldsymbol{\theta}) = \frac{1}{n} \sum_{i=1}^{n} \underbrace{L\left(y^{(i)}, f\left(\mathbf{x}^{(i)} | \boldsymbol{\theta}\right)\right)}_{g_i(\boldsymbol{\theta})}
$$

$$
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$$

 $\bullet$  for a data set

$$
\mathcal{D} = \left( \left( \boldsymbol{x}^{(1)}, y^{(1)} \right), \ldots, \left( \boldsymbol{x}^{(n)}, y^{(n)} \right) \right)
$$

- a loss function  $L\left(y, f(\mathbf{x})\right)$ , e.g., L2 loss  $L\left(y, f(\mathbf{x})\right) = (y f(\mathbf{x}))^2,$
- and a model class  $f$ , e.g., the linear model  $f\left(\mathbf{x}^{(i)}\mid\boldsymbol{\theta}\right)=\boldsymbol{\theta}^{\top}\mathbf{x}$ .

### **SGD IN ML / 2**

For large data sets, computing the exact gradient

$$
\mathbf{d} = \frac{1}{n} \sum_{i=1}^{n} \nabla_{\boldsymbol{\theta}} L\left(y^{(i)}, f\left(\mathbf{x}^{(i)} | \boldsymbol{\theta}\right)\right)
$$

may be expensive or even infeasible to compute and is approximated by

$$
\hat{\mathbf{d}} = \frac{1}{m} \sum_{i \in J} \nabla_{\boldsymbol{\theta}} L\left(y^{(i)}, f\left(\mathbf{x}^{(i)} \mid \boldsymbol{\theta}\right)\right),
$$

for  $J \subset 1, 2, ..., n$  random subset.

**NB:** Often, maximum size of *J* technically limited by memory size.



### **STOCHASTICITY OF SGD**





Minimize  $g(x_1, x_2) = 1.25(x_1 + 6)^2 + (x_2 - 8)^2$ . **Left:** GD. **Right:** SGD. Black line shows average value across multiple runs. (Source: Shalev-Shwartz et al., Understanding Machine Learning, 2014.)

### **STOCHASTICITY OF SGD / 2**

Assume batch size  $m = 1$  (statements also apply for larger batches).

- **(Possibly) suboptimal direction:** Approximate gradient  $\hat{\mathbf{d}} = \nabla_{\mathbf{x}} q_i(\mathbf{x})$  might point in suboptimal (possibly not even a descent!) direction
- **Unbiased estimate:** If *J* drawn i.i.d., approximate gradient **d**ˆ is an unbiased estimate of gradient  $\mathbf{d} = \nabla_{\mathbf{x}} g(\mathbf{x}) = \sum^{n}$ *i*=1  $\nabla_{\mathbf{x}}g_i(\mathbf{x})$ :

$$
\mathbb{E}_{i}\left[\nabla_{\mathbf{x}}g_{i}(\mathbf{x})\right]=\sum_{i=1}^{n}\nabla_{\mathbf{x}}g_{i}(\mathbf{x})\cdot\mathbb{P}(i=i) \n=\sum_{i=1}^{n}\nabla_{\mathbf{x}}g_{i}(\mathbf{x})\cdot\frac{1}{n}=\nabla_{\mathbf{x}}g(\mathbf{x}).
$$

**Conclusion:** SGD might perform single suboptimal moves, but moves in "right direction" **on average**.



**Example:**  $g(\mathbf{x}) = \sum_{i=1}^{5} g_i(\mathbf{x})$ ,  $g_i$  quadratic. Batch size  $m = 1$ .



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In iteration 5, SGD performs a suboptimal move away from the minimum.



X  $\times\overline{\times}$ 

**Blue area**: Each  $-\nabla g_i(\mathbf{x})$  points towards minimum. **Red area** ("confusion area"):  $-\nabla g_i(\mathbf{x})$  might point away from minimum and perform a suboptimal move.

At location **x**, "confusion" is captured by variance of gradients

$$
\frac{1}{n}\sum_{i=1}^n \|\nabla_{\mathbf{x}}g_i(\mathbf{x}) - \nabla_{\mathbf{x}}g(\mathbf{x})\|^2
$$

- If term is 0, next step goes in gradient direction (for each *i*)
- If term is small, next step *likely* goes in gradient direction
- $\bullet$  If term is large, next step likely goes in direction different than gradient

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# **CONVERGENCE OF SGD**

As a consequence, SGD has worse convergence properties than GD.

**But:** Can be controlled via **increasing batches** or **reducing step size**.

#### **The larger the batch size** *m*

- the better the approximation to  $\nabla_{\mathbf{x}} g(\mathbf{x})$
- the lower the variance
- the lower the risk of performing steps in the wrong direction

#### **The smaller the step size**  $α$

- the smaller a step in a potentially wrong direction
- the lower the effect of high variance

As maximum batch size is usually limited by computational resources (memory), choosing the step size is crucial.

X X

# **EFFECT OF BATCH SIZE**



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SGD for a NN with batch size  $\in \{0.5\%, 10\%, 50\%\}$  of the training data. The higher the batch size, the lower the variance.