#### Gradient Descent

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## Most AI/ML optimizations must be **numerically** optimized

- One common algorithm is gradient descent
  - Primary algorithm for deep learning
  - Works in very high dimensions

- Other optimization algorithms
  - Expectation Maximization (alternating optimization)
  - Sampling-based optimization (MCMC/Gibb)
  - Greedy optimization (e.g., decision trees)

#### Gradient descent is like taking steps down the steepest descent into a valley



https://www.hackerearth.com/blog/developers/3-types-gradient-descent-algorithms-small-large-data-sets/

Vanilla gradient descent (GD) has simple form

- Objective (Loss) function denoted by  $\mathcal{L}(\theta; \mathcal{D})$ : arg min  $\mathcal{L}(\theta; \mathcal{D})$
- 1. Start at random parameter, e.g.,  $\theta^0 \sim \mathcal{N}(0, 1)$
- 2. Iteratively update parameter via <u>negative</u> <u>gradient</u> of loss function ( $\eta_t$  is step size or

$$\theta^{t+1} = \theta^t - \eta_t \nabla_\theta \mathcal{L}(\theta^t)$$
  
•  $\eta_t$  is learning rate (or step size)

Stochastic gradient descent (SGD) merely uses one sample in the gradient calculation

- The loss function can usually be split into a summation of losses  $\ell(\theta; x_i)$  for each sample  $x_i$ :  $\mathcal{L}(\theta; \mathcal{D}) = \frac{1}{n} \sum_{i=1}^{n} \ell(\theta; x_i)$
- SGD approximates the full gradient by the gradient of a single sample
  - $\nabla_{\theta} \mathcal{L}(\theta^{t}; \mathcal{D}) \approx \nabla_{\theta} \ell(\theta^{t}; x_{i})$
  - Theoretically,  $\mathbb{E}_i[\nabla_{\theta} \ell(\theta^t; x_i)] = \nabla_{\theta} \mathcal{L}(\theta^t; \mathcal{D})$
- ► Loop through all  $x_i \in \mathcal{D}$  $\theta^{t+1} = \theta^t - \eta_t \nabla_\theta \ell(\theta^t; x_i)$
- One pass through dataset
  - GD: 1 large update with O(n) cost
  - SGD: n smaller updates with O(1) cost each

## <u>Stochastic gradient descent (SGD)</u> merely uses one sample in the gradient calculation



Full gradient is average over single sample gradients. This is why it is "stochastic".

https://golden.com/wiki/Stochastic\_gradient\_descent\_(SGD)

# Mini-batch SGD (or just SGD) uses a small batch of samples in the gradient calculation

- Mini-batch SGD approximates the full gradient by the gradient of a batch of samples
  - Sample mini-batch

$$\theta^{t+1} = \theta^t - \eta_t \sum_{k=1}^b \frac{1}{b} \nabla_\theta \ell(\theta^t; x_k)$$

- One pass through dataset
  - ► GD: 1 large update
  - ► SGD: *n* smaller updates
  - Mini-batch SGD:  $\frac{n}{b}$  medium-size updates

#### Gradient descent demo for simplified logistic regression

Learning rate / step size is critical for convergence and correctness of algorithm

- If learning rate is too high, the algorithm could diverge.
  - Diverge means to get farther away from the solution.
- If learning rate too low, the algorithm could take a very long time to converge.



https://www.jeremyjordan.me/nn-learning-rate/

Adaptive learning rates may help (but not always)

- Decreasing step size, η<sub>t</sub> = 1/t
   Intuition: Approaches 0 but can cover an infinite distance since lim<sub>a→∞</sub> Σ<sup>a</sup><sub>t=1</sub> 1/t = ∞
- ADAM Adaptive Moment Estimation
- See <u>https://pytorch.org/docs/stable/optim.html</u> for more options

Parameter initialization can be important if non-convex or step size incorrect

- If convex function, initial parameter θ<sup>0</sup> will not affect final optimization result θ = argmin L(θ).
   Yay! (Assuming appropriate step size.)
- If non-convex, starting position <u>WILL</u> affect final converged θ.
  - Sad day. (But sometimes it's not too bad in practice.)

