Neural networks and Backpropagation

\$ echo "Data Sciences Institute"



Yesterday

- Overview
- Computation graph view of neural networks
- Linear operation followed by non-linear activation
 ...But what is the linear operation, really?

Today

- A closer look at what's going on in a "neuron"
- Backpropagation: how do we train a neural network?

Neural Network for classification

Vector function with tunable parameters θ

$$\mathbf{f}(\cdot; heta):\mathbb{R}^N o (0,1)^K$$

Sample s in dataset S:

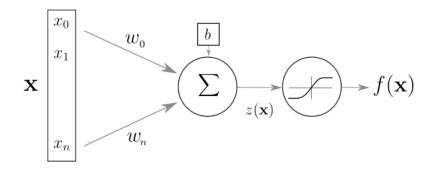
- input: $\mathbf{x}^s \in \mathbb{R}^N$
- ullet expected output: $y^s \in [0,K-1]$

Output is a conditional probability distribution:

$$\mathbf{f}(\mathbf{x}^s; \theta)_c = P(Y = c | X = \mathbf{x}^s)$$



Artificial Neuron



$$z(\mathbf{x}) = \mathbf{w}^T \mathbf{x} + b$$

$$f(\mathbf{x}) = g(\mathbf{w}^T \mathbf{x} + b)$$

- $\mathbf{x}, f(\mathbf{x})$ input and output
- $z(\mathbf{x})$ pre-activation
- \bullet w, b weights and bias
- g activation function

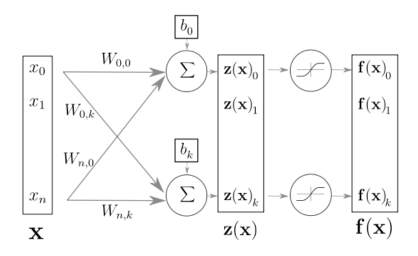




Concrete Example

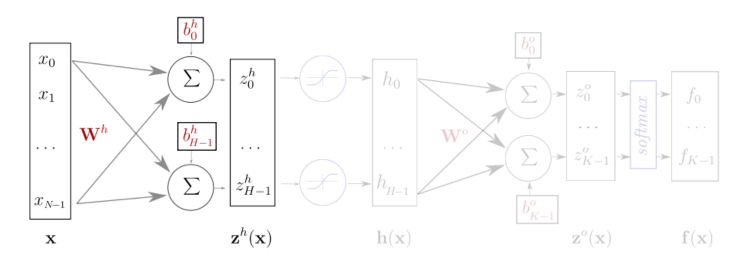
- Say we have two input dimensions x_1 and x_2 and one output dimension f(x) (sometimes, \hat{y} the predicted value of y is used instead of f(x))
- ullet Our weights and biases could be W=[3,-2] and b=1
- Our non-linearity could be ReLU: g(z) = max(0, z)
- ullet Now $z(x)=3x1-2x_2+1$ and $f(x)=max(0,3x_1-2x_2+1)$
- Every neuron in a neural network is a function like this!

Layer of Neurons (Vectorization)



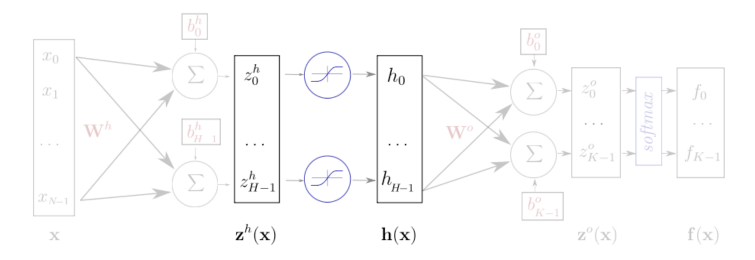
$$f(x) = g(z(x)) = g(Wx + b)$$

• **W**, **b** now matrix and vector

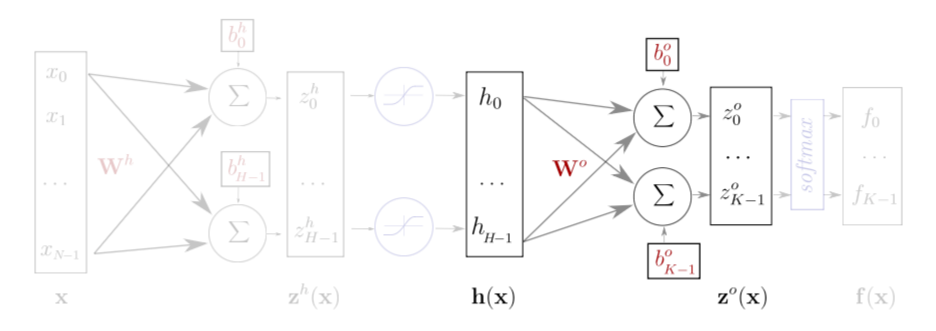


$$ullet \mathbf{z}^h(\mathbf{x}) = \mathbf{W}^h\mathbf{x} + \mathbf{b}^h$$

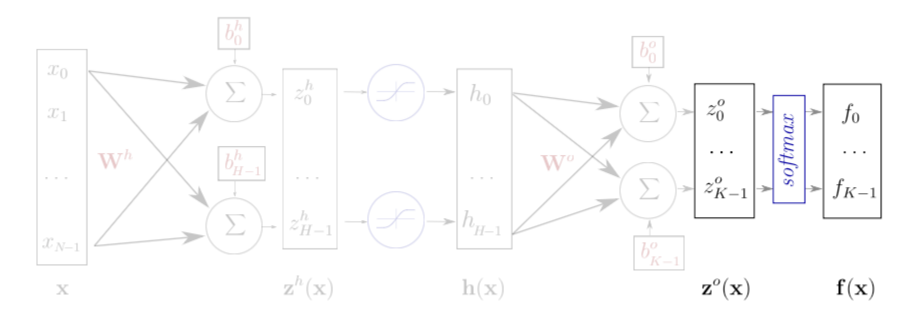




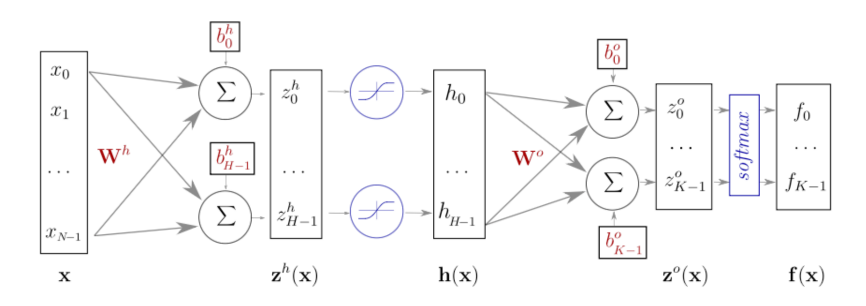
•
$$\mathbf{h}(\mathbf{x}) = g(\mathbf{z}^h(\mathbf{x})) = g(\mathbf{W}^h\mathbf{x})$$



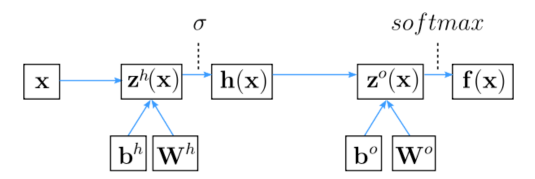
$$ullet \mathbf{z}^o(\mathbf{x}) = \mathbf{W}^o\mathbf{h}(\mathbf{x}) + \mathbf{b}^o$$

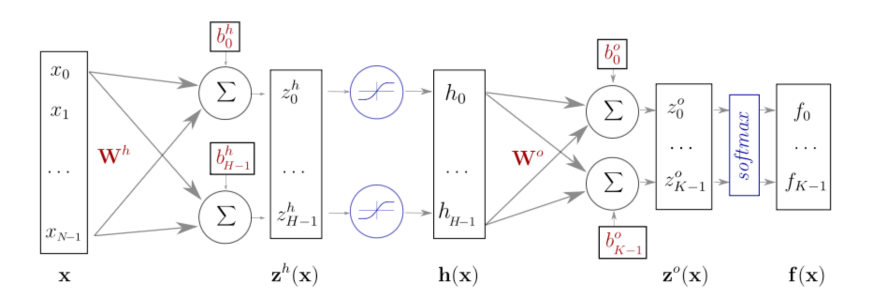


 $ullet \mathbf{f}(\mathbf{x}) = softmax(\mathbf{z}^o) = softmax(\mathbf{W}^o\mathbf{h}(\mathbf{x}) + \mathbf{b}^o)$



Alternate representation

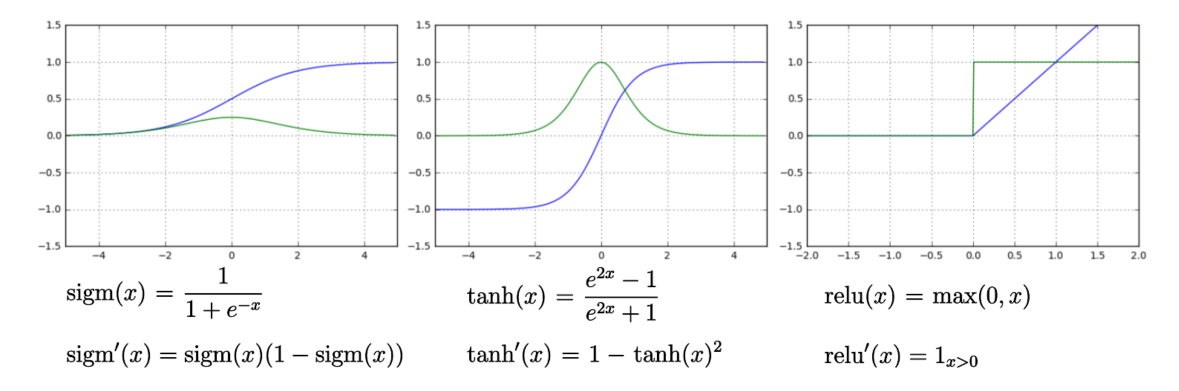




Keras implementation

```
model = Sequential()
model.add(Dense(H, input_dim=N))  # weight matrix dim [N * H]
model.add(Activation("tanh"))
model.add(Dense(K))  # weight matrix dim [H x K]
model.add(Activation("softmax"))
```

Element-wise activation functions



- blue: activation function
- green: derivative



Softmax function

$$softmax(\mathbf{x}) = rac{1}{\sum_{i=1}^n e^{x_i}} \cdot egin{bmatrix} e^{x_2} \ \vdots \ e^{x_n} \end{bmatrix}$$

- vector of values in (0, 1) that add up to 1
- ullet for example, ${f x}=[1,2,3]$ becomes $rac{1}{e^1+e^2+e^3}\cdot [e^1,e^2,e^3]^T=[0.09,0.24,0.67]$
- $p(Y = c|X = \mathbf{x}) = \operatorname{softmax}(\mathbf{z}(\mathbf{x}))_c$
- the pre-activation vector $\mathbf{z}(\mathbf{x})$ is often called "the logits"





Training the network

Find parameters that minimize the **negative log likelihood** (or cross entropy)

The loss function for a given sample $s \in S$:

$$l(\mathbf{f}(\mathbf{x}^s; heta),y^s) = nll(\mathbf{x}^s,y^s; heta) = -\log\mathbf{f}(\mathbf{x}^s; heta)_{y^s}$$

$$y^{s} = 3$$

example
$$y^s=3$$

$$l(\mathbf{f}(\mathbf{x}^s;\theta),y^s)=l\begin{pmatrix} f_0\\ \dots\\ f_3\\ \dots\\ f_{K-1} \end{pmatrix}, \begin{matrix} 0\\ \dots\\ 1\\ \dots\\ 0 \end{pmatrix}=-\log\ f_3$$



Training the network

Find parameters $\theta = (\mathbf{W}^h; \mathbf{b}^h; \mathbf{W}^o; \mathbf{b}^o)$ that minimize the **negative log likelihood** (or cross entropy)

The loss function for a given sample $s \in S$:

$$l(\mathbf{f}(\mathbf{x}^s; \theta), y^s) = nll(\mathbf{x}^s, y^s; \theta) = -\log \mathbf{f}(\mathbf{x}^s; \theta)_{y^s}$$

The cost function is the negative likelihood of the model computed on the full training set (for i.i.d. samples):

$$L_S(heta) = -rac{1}{|S|} \sum_{s \in S} \log \mathbf{f}(\mathbf{x}^s; heta)_{y^s}$$



Training the network

- Now we have a mathematical function representing the network
- And we have a way of measuring how good it is
- How do we find the parameters that minimize the loss?

Gradient Descent

- Let's imagine we only have one parameter θ
- ullet We can compute the derivative of the loss with respect to heta
- The derivative, $\frac{dL}{d\theta}$, tells us the slope of the loss function at a given point
- If $\frac{dL}{d\theta}>0$, increasing heta will increase the loss, and vice versa
- To minimize loss, we adjust θ in the opposite direction of $\frac{dL}{d\theta}$
- This is done using the update rule: $heta= heta_{old}-\eta rac{dL}{d heta}$

Gradient Descent

- We can use gradient descent to play "guess what number I'm thinking of"
- If your guess is too high, you decrease it
- If your guess is too low, you increase it
- The error function is a parabola
- By finding the lowest point on the parabola, you find the best guess

Implementing Gradient Descent

- Start with an initial guess for θ
- Calculate $\frac{dL}{d\theta}$ using the current value of θ
- Update θ using the update rule
- Repeat the process until the change in loss is below a threshold or a set number of iterations is reached
- The choice of learning rate η is crucial: too high, and we may overshoot the minimum; too low, and convergence will be slow

Stochastic Gradient Descent

- Traditional Gradient Descent uses the entire dataset to compute the gradient, which can be computationally expensive
- Stochastic Gradient Descent (SGD) updates the parameters using only a single data point (or a small batch)
- In SGD, for each iteration, a data point (or batch) is randomly selected to compute the gradient
- Since only a subset of data is used, the gradient estimation can be noisy, leading to a less smooth path towards the minimum
- However, SGD is much faster than traditional gradient descent

Stochastic Gradient Descent

Initialize θ randomly

For E epochs perform:

- ullet Randomly select a small batch of samples $(B\subset S)$
 - \circ Compute gradients: $\Delta =
 abla_{ heta} L_B(heta)$
 - \circ Update parameters: $heta \leftarrow heta \eta \Delta$
- ullet Repeat until the epoch is completed (all of S is covered) Stop when reaching criterion:
- nll stops decreasing when computed on validation set



Computing Gradients

Output Weights:
$$\frac{\partial l(\mathbf{f}(\mathbf{x}),y)}{\partial W_{i,j}^o}$$

Output Weights:
$$\frac{\partial l(\mathbf{f}(\mathbf{x}), y)}{\partial W_{i,j}^o}$$
 Hidden Weights: $\frac{\partial l(\mathbf{f}(\mathbf{x}), y)}{\partial W_{i,j}^h}$

Output bias:
$$\frac{\partial l(\mathbf{f}(\mathbf{x}), y)}{\partial b^o}$$

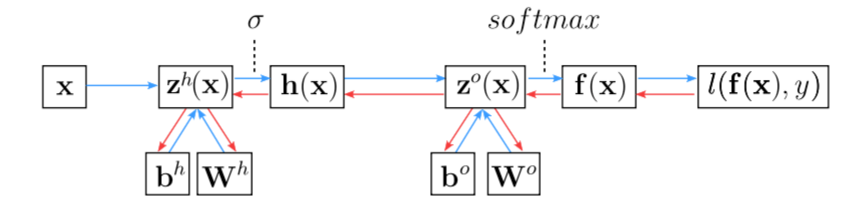
Output bias:
$$\frac{\partial l(\mathbf{f}(\mathbf{x}),y)}{\partial b_i^o}$$
 Hidden bias: $\frac{\partial l(\mathbf{f}(\mathbf{x}),y)}{\partial b_i^h}$

- The network is a composition of differentiable modules
- We can apply the "chain rule"

Chain rule

- Mathematical theorem that lets us compute derivatives when functions are inside other functions
- Remember, our neural network is a composition of functions: f(x) = g(h(x))
- The chain rule tells us how to compute $\frac{df}{dx}$
- $\bullet \ \frac{df}{dx} = \frac{df}{dg} \frac{dg}{dh} \frac{dh}{dx}$
- In English: The derivative of the overall network with respect to its input is the product of derivatives of each function in the network

Backpropagation



- Compute partial derivatives of the loss
- For any given function in the network, we can compute how changing its parameters will affect the loss
- In other words, we can find how much each parameter's value contributes to the loss

Initialization and Learning Tricks

Initialization and normalization

- Input data should be normalized to have approx. same range:
 - standardization or quantile normalization
- Initializing weights:
 - Zero is a saddle point: no gradient, no learning
 - Constant init: all neurons compute the same function
 - \circ Solution: random init, ex: $w \sim \mathcal{N}(0, 0.1)$
 - Better inits: Xavier Glorot and Kaming He & orthogonal
- Biases can (should) be initialized to zero

SGD learning rate

- Very sensitive:
 - \circ Too high \rightarrow early plateau or even divergence
 - Too low → slow convergence
 - $\circ~$ Try a large value first: $\eta=0.1$ or even $\eta=1$
 - Divide by 10 and retry in case of divergence
- Large constant LR prevents final convergence
 - \circ multiply η_t by eta < 1 after each update
 - \circ or monitor validation loss and divide η_t by 2 or 10 when no progress
 - See ReduceLROnPlateau in Keras



Momentum

Accumulate gradients across successive updates:

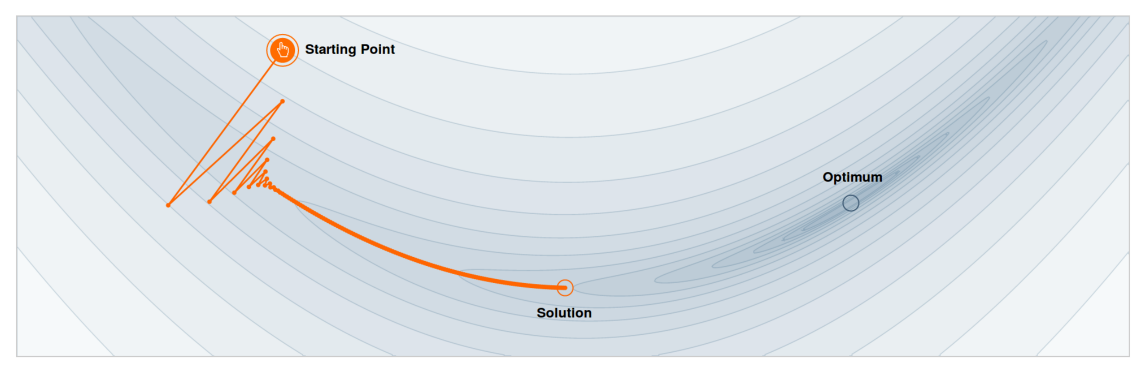
$$m_t = \gamma m_{t-1} + \eta
abla_{ heta} L_{B_t}(heta_{t-1})$$

$$\theta_t = \theta_{t-1} - m_t$$

 γ is typically set to 0.9

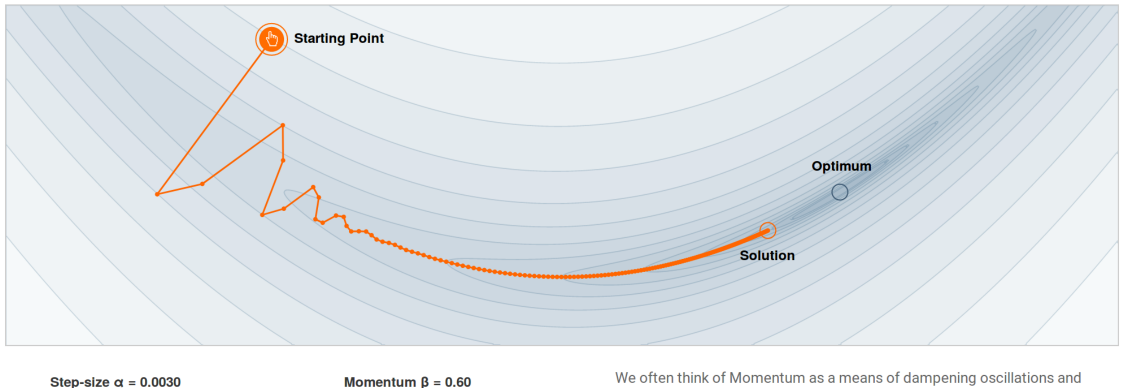
Larger updates in directions where the gradient sign is constant to accelerate in low curvature areas





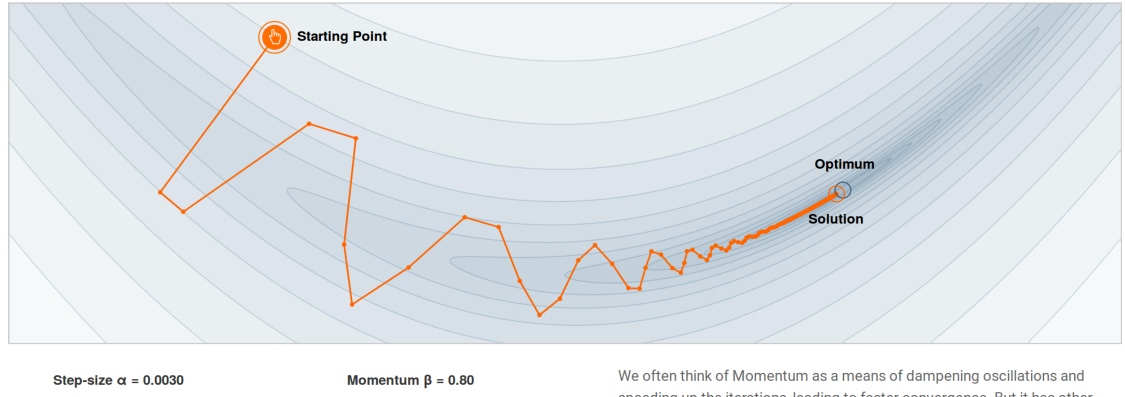


We often think of Momentum as a means of dampening oscillations and speeding up the iterations, leading to faster convergence. But it has other interesting behavior. It allows a larger range of step-sizes to be used, and creates its own oscillations. What is going on?



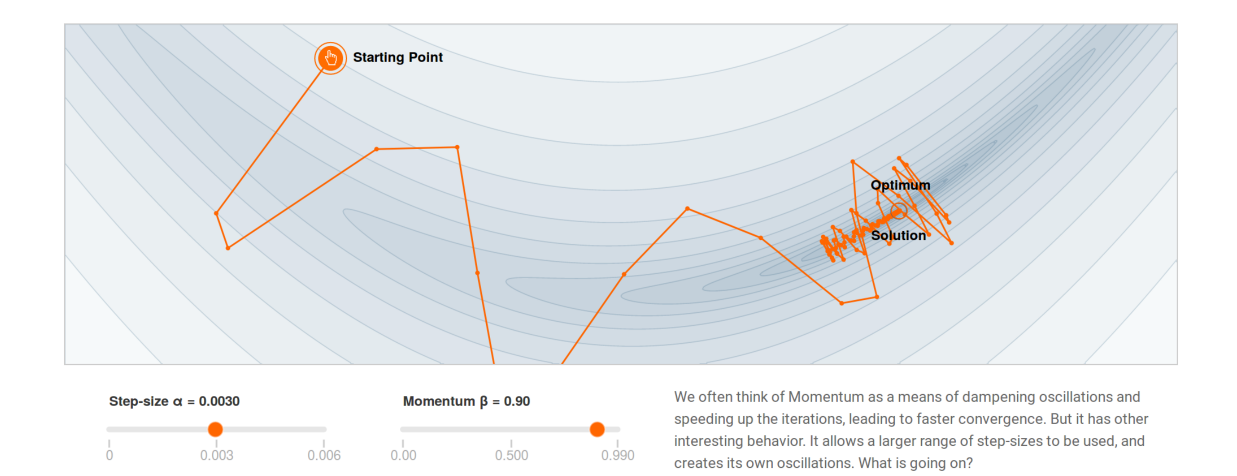


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Alternative optimizers

- SGD (with Nesterov momentum)
 - Simple to implement
 - \circ Very sensitive to initial value of η
 - Need learning rate scheduling
- Adam: adaptive learning rate scale for each param
 - \circ Global η set to 3e-4 often works well enough
 - Good default choice of optimizer (often)
- Many other promising methods:
 - RMSProp, Adagrad, Adadelta, Nadam, ...
 - Often takes some experimentation to find the best one



The Karpathy Constant for Adam





3e-4 is the best learning rate for Adam, hands down.

4:01 AM - 24 Nov 2016

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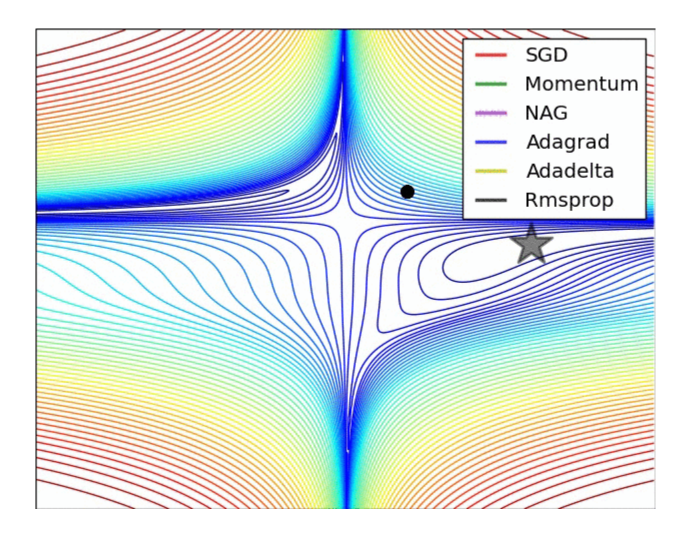








Optimizers around a saddle point





Next: Lab 2!