



Loss functions and Optimization

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Outline

Loss Functions

Optimization





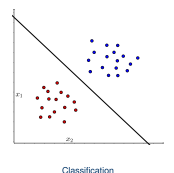
Loss Functions

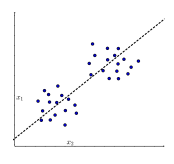




Regression vs. classification

- Regression: Estimate a continuous variable for every input.
- Classification: Estimate a discrete variable for every input.





Regression



Loss function vs. last activation function in a network

The last activation function

- is applied on individual samples
 x_m of the batch
- is present at training and testing
- produces the output, or prediction
- generally produces a vector

The loss function

- combines all M samples and labels
- is only present at training
- produces the loss
- generally produces a scalar



Maximum Likelihood Estimation Reminder

Assume a

- Training set with
 - Observations: $\mathbf{X} = \mathbf{x}_1, \cdots, \mathbf{x}_M$
 - and associated labels $\mathbf{Y} = \mathbf{y}_1, \cdots, \mathbf{y}_M$
- and a model for a conditional probability density function p(y|x)



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Dataset

- Probability to observe \mathbf{y}_m given observation \mathbf{x}_m is $p(\mathbf{y}_m|\mathbf{x}_m)$
- Joint probability is $p(\mathbf{y}_m|\mathbf{x}_m) \cdot p(\mathbf{y}_i|\mathbf{x}_i)$ if they are:
 - Independent
 - and Identically Distributed
- probability to observe **Y** is $\prod_{m=1}^{M} p(\mathbf{y}_m | \mathbf{x}_m)$



Likelihood function

p governed by parameters w

$$\max_{\mathbf{w}} \left\{ \prod_{m=1}^{M} p(\mathbf{y}_{m} | \mathbf{x}_{m}, \mathbf{w}) \right\}$$



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- Maximum not affected by a monotonous transformation
- Maximization to minimization by flipping the sign

• minimize
$$\left\{-\ln\left(\mathit{L}(\mathbf{w})\right)\right\} = \min_{\mathbf{w}} \left\{\sum_{m=1}^{M} -\ln\left(\mathit{p}(\mathbf{y}_{m}|\mathbf{x}_{m},\mathbf{w})\right)\right\}$$



Regression

Assume a univariate Gaussian model:

$$p(y|\mathbf{x},\mathbf{w},eta) = \mathcal{N}(\hat{y}(\mathbf{x},\mathbf{w}), \quad \frac{1}{eta} \quad)$$



Regression

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$$p(y|\mathbf{x}, \mathbf{w}, \beta) = \mathcal{N}(\underbrace{\hat{y}(\mathbf{x}, \mathbf{w})}_{\mu}, \underbrace{\frac{1}{\beta}}_{\sigma})$$



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$$p(y|\mathbf{x}, \mathbf{w}, \beta) = \mathcal{N}(\underbrace{\hat{y}(\mathbf{x}, \mathbf{w})}_{\mu}, \underbrace{\frac{1}{\beta}}_{\sigma})$$
$$= \frac{\sqrt{\beta}}{\sqrt{2\pi}} e^{\beta \frac{-(y_m - \hat{y}(\mathbf{x}_m, \mathbf{w}))^2}{2}}$$



$$L(\mathbf{w}) = \sum_{m=1}^{M} - \ln \left(\frac{\sqrt{\beta}}{\sqrt{2\pi}} e^{\beta \frac{-(y_m - \hat{y}(\mathbf{x}_m, \mathbf{w}))^2}{2}} \right)$$



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This can be generalized to vectors \mathbf{y}_m , $\hat{\mathbf{y}}$:

$$\frac{1}{2} \sum_{m=1}^{M} \|\mathbf{y}_m - \hat{\mathbf{y}}(\mathbf{x}_m, \mathbf{w})\|_2^2$$



Classification using an L-norm

L_2 -loss and L_1 -loss can be applied for classification

- They correspond to variants of minimizing the expected misclassification probability
- They cause slow convergence because they don't penalize heavily misclassified probabilities
- They might be advantageous in situations with extreme label noise



Classification

Assume our network provides us with a probabilistic output p.



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Multi-class generalization: Multinoulli (Categorical, $\mathfrak C$) distribution

• y, which is one-hot encoded

$$\mathfrak{C}(\mathbf{y}|\mathbf{p}) = \begin{cases} \prod_{k=0}^{K} p_k^{y_k} & \text{if } y_k \in \{0,1\} \\ 0 & \text{otherwise} \end{cases}$$



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- We have an unfair coin: $\mathbf{p} = \begin{pmatrix} 0.3 \\ 0.7 \end{pmatrix}$ and observe $\mathbf{y} = \begin{pmatrix} 0 \\ 1 \end{pmatrix}$



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- So the probability to observe $\begin{pmatrix} 0 \\ 1 \end{pmatrix}$ which is tail for this unfair coin is 70%.



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Crossentropy

$$=-\sum_{m=1}^{M}\ln(\hat{y}_k(\mathbf{x}_m,\mathbf{w}))|_{y_{k,m}=1}$$



Relation to the Kullback Leibler Divergence

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We know that our ML estimation for a single sample has the form of cross-entropy:

$$-\sum_{k=0}^{K}\ln\left(\hat{y}_{k}^{y_{k}}\right)=\mathsf{H}(\mathbf{y},\hat{\mathbf{y}})$$

and therefore is equal to minimizing the KL-divergence.



Can we also use cross-entropy for regression?



Can we also use cross-entropy for regression?

- Of course. We just have to make sure $\hat{y}_k \in [0, 1] \forall k$
- This can be achieved using a sigmoid activation function
- y is simply no longer one-hot encoded
- As we've seen before this is equivalent to minimizing KL-divergence





- L2-loss can be used for **regression**
- Cross-entropy-loss can be used for classification



- L₂-loss can be used for regression
- Cross-entropy-loss can be used for classification
- L₂-loss and Cross-entropy-loss can be derived as ML-Estimators from strict probabilistic assumptions
- In absence of more domain knowledge they are your first choices
- They are both intrinsically multi-variate



How does the Perceptron criterion fit into this?

minimize
$$\left\{L(\mathbf{w}) = -\sum_{\mathbf{x}_m \in \mathcal{M}} y_m \cdot (\mathbf{w}^\mathsf{T} \mathbf{x}_m)\right\}$$

• Remember that here $y_m \in -1, 1$ instead of $y_m \in 0, 1$



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- ... and the gradient would vanish almost everywhere

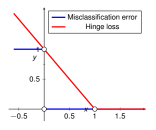


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- What if it was in?
- Than we would just count the number of misclassifications
- ... and the gradient would vanish almost everywhere
- Sounds familiar?
- What did we do about that last time?



Hinge loss



$$L(\mathbf{w}) = \sum_{m=1}^{M} \max(0, 1 - y_m \hat{y}(\mathbf{x}_m, \mathbf{w}))$$

- Classification depends only on the sign
- If the signs match we get a positive value and classify correct
- Hinge loss is a convex approximation to the misclassification loss
- · But what about the gradient?



Suppose we have a convex, differentiable function. Than we have:

$$f(\mathbf{x}) \ge f(\mathbf{x}_0) + \nabla f(\mathbf{x}_0)^T (\mathbf{x} - \mathbf{x}_0) \quad \forall \mathbf{x} \in \mathcal{X}$$



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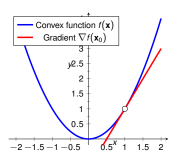
In words: If we follow the gradient from any point of a convex function and check against the function, its value at the same \mathbf{x} will be higher.



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- A vector **g** is a subgradient of a **convex** function f at point $\mathbf{x}_0 \in \mathcal{X}$ if:

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 This is not unique! We get a set of subgradients which we call a subdifferential:

$$\partial f(\mathbf{x}_0) \coloneqq \{\mathbf{g}\}$$



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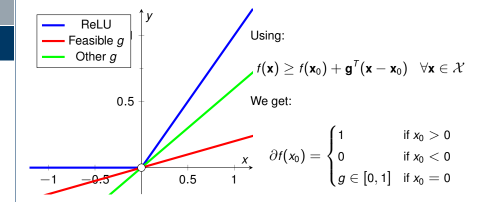
 This is not unique! We get a set of subgradients which we call a subdifferential:

$$\partial f(\mathbf{x}_0) := \{\mathbf{g}\}$$

If f is differentiable at x₀:

$$\partial f(\mathbf{x}_0) = \{\nabla f(\mathbf{x}_0)\}\$$





- We already used this for the ReLU!
- Gradient descent was implicitly generalized to the subgradient algorithm





- Subgradients are a generalization of gradients for convex, non-smooth functions
- The gradient descent algorithm is replaced by the subgradient algorithm for these functions



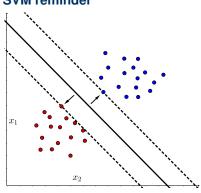
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- This is basically just the solid math why this works
- We use this for the ReLU and Hinge loss so far



SVM reminder



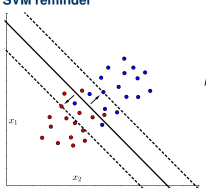
$$min \quad \frac{1}{2} \|\mathbf{w}\|_2^2$$

s.t.
$$\forall m : -(y_m \cdot (\mathbf{w}^T \mathbf{x}_m) - 1) \leq 0$$

$$) \leq 0$$



SVM reminder



$$\min \quad \frac{1}{2} \|\mathbf{w}\|_2^2 + \gamma$$

min
$$\frac{1}{2} \|\mathbf{w}\|_2^2 + \gamma \sum_m \xi_m$$

s.t. $\forall m : -(y_m \cdot (\mathbf{w}^T \mathbf{x}_m) - 1 + \xi_m) \le 0$

$$\forall m: -\xi_m \leq 0$$



• We construct the Lagrangian dual function

$$L(\mathbf{w}) = \frac{1}{2} \|\mathbf{w}\|_{2}^{2} + \gamma \sum_{m=1}^{M} \xi_{m} + \sum_{m=1}^{M} \lambda_{m} (-y_{m} \cdot (\mathbf{w}^{T} \mathbf{x}_{m}) + 1 - \xi_{m}) - \sum_{m=1}^{M} \nu_{m} \xi_{m}$$



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- We construct the Lagrangian dual function
- Remember: $\lambda_m \geq 0$

$$L(\mathbf{w}) = \frac{1}{2} \|\mathbf{w}\|_{2}^{2} + \gamma \sum_{m=1}^{M} \xi_{m} + \sum_{m=1}^{M} \lambda_{m} (-y_{m} \cdot (\mathbf{w}^{T} \mathbf{x}_{m}) + 1 - \xi_{m}) - \sum_{m=1}^{M} \nu_{m} \xi_{m}$$

$$= \frac{1}{2} \|\mathbf{w}\|_{2}^{2} + \sum_{m=1}^{M} (\gamma \xi_{m} - \nu_{m} \xi_{m} - \lambda_{m} \xi_{m}) + \sum_{m=1}^{M} \lambda_{m} (1 - y_{m} \cdot (\mathbf{w}^{T} \mathbf{x}_{m}))$$



- · We construct the Lagrangian dual function
- Remember: $\lambda_m \geq 0$

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- · We construct the Lagrangian dual function
- Remember: $\lambda_m > 0$
- Equivalent "up to an overall multiplicative constant"[1]

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$$\approx \frac{1}{2} \underbrace{\|\mathbf{w}\|_{2}^{2}}_{L2 \text{ regularizer}} + \gamma \sum_{m=1}^{M} \underbrace{\max(0, 1 - y_{m} \cdot (\mathbf{w}^{T} \mathbf{x}_{m}))}_{\text{Hinge loss}}$$



Open points



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Outliers are punished linearly



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• A variant of the hinge loss which penalizes outliers more strongly [4]:

$$L(\mathbf{w}) = \sum_{m=1}^{M} (\max(0, 1 - y_m \hat{y}(\mathbf{x}_m, \mathbf{w})))^2$$



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How to apply SVMs to multi-class problems?



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How to apply SVMs to multi-class problems?

A Hinge loss for multi-class problems [9]:

$$L(\mathbf{w}) = \sum_{m=1}^{M} \sum_{k \neq c}^{K} \max(0, 1 - \hat{y}_{c}(\mathbf{x}_{m}, \mathbf{w}) + \hat{y}_{k}(\mathbf{x}_{m}, \mathbf{w}))$$



Summary

- We have seen we can incorporate an SVM into a neural network
- See [4] for a reference using this
- We've learned before how to deal with the non-smooth objective





Optimization





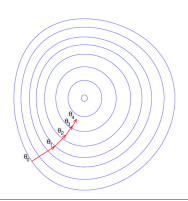
Gradient Descent revisited

Goal: Optimize empirical risk:

$$\mathbb{E}_{\mathbf{x},\mathbf{y}\sim\hat{\rho}_{\mathrm{data}}(\mathbf{x},\mathbf{y})}\big[L(\mathbf{w},\mathbf{x}_m,\mathbf{y}_m)\big] = \frac{1}{M}\sum_{m=1}^{M}L(\mathbf{w},\mathbf{x},\mathbf{y})$$

$$\mathbf{w}^{k+1} = \mathbf{w}^{(k)} - \eta \nabla L(\mathbf{w}^{(k)}, \mathbf{x}, \mathbf{y})$$

- Step size defined by learning rate η
- Gradient with respect to every sample
- Guaranteed to converge to a local minimum





For each iteration...

• Batch Gradient Descent: Use all *M* samples



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 ≪ M random samples

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- Small batches offer regularization effect \Rightarrow need smaller η
- Regains efficiency → the standard case in deep learning



How can this even work?

- Optimization problem is non-convex
- Exponential number of local minima

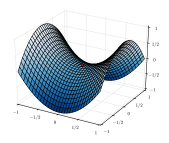


How can this even work?

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Possible Answers (Choromanska et al. 2015, Dauphin et al. 2014)

- High dimensional function
- Local minima exist but very close to global minima
- ... and many of those are equivalent
- → Presumably more critical: saddle points
 - Local minimum might be better than global minima (overfitting!)



Source: https://upload.wikimedia.org/wikipedia/commons/1/1e/Saddle_point.svg



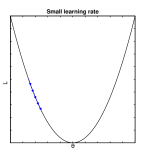
Another possible answer

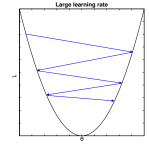
Possible answer (Percy Liang, NIPS 2016)

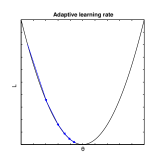
- "overprovisioning"
- Many different ways how a network can approximate the desired relationship
- Only needs to find one
- This has been verified experimentally by learning random labels [10]



SGD – Learning Rate Choice







- η too small: long training time
- η too large: miss optima
- Practice: "learning rate decay": adapt η gradually (e.g.: start with $\eta=0.01$ and divide every x epoch by 10)





By performing line search?



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- Multiple evaluations necessary, while we could take multiple steps
- The direction is extremely noisy anyway
- Still people have presented methods [8]



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- L-BFGS doesn't perform well outside of batch settings
- · A report on this was presented by Google [7]



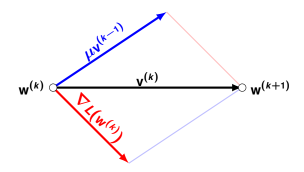
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Parameter update based on current and past gradients:

$$\begin{aligned} \mathbf{v}^{(k)} &= \underbrace{\boldsymbol{\mu}}_{\text{momentum}} \mathbf{v}^{(k-1)} - \eta \, \nabla L(\mathbf{w}^{(k)}) \\ \mathbf{w}^{(k+1)} &= \mathbf{w}^{(k)} + \mathbf{v}^{(k)} \end{aligned}$$



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- commonly: $\mu = \{0.9, 0.95, 0.99\}$ (or adaptive: small \rightarrow large)
- + Overcomes poor Hessian & variance in SGD \rightarrow dampened oscillations
- + Acceleration
- Still learning rate decay needed!



Nesterov Accelerated Gradient (NAG) / Nesterov Momentum

"Look ahead" - compute the gradient in the direction we're going anyway!

$$\mathbf{v}^{(k)} = \mu \mathbf{v}^{(k-1)} - \eta \, \nabla L(\underbrace{\mathbf{w}^{(k)} + \mu \mathbf{v}^{(k-1)}}_{\text{approx. of next parameters}})$$

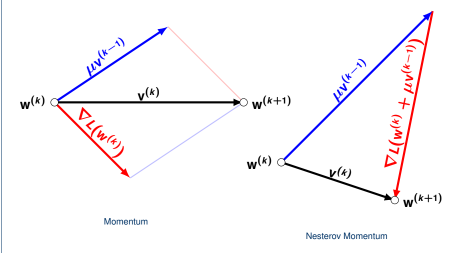
$$\mathbf{w}^{(k+1)} = \mathbf{w}^{(k)} + \mathbf{v}^{(k)}$$

We can rewrite this to use the conventional gradient:

$$\mathbf{v}^{(k)} = \mu \mathbf{v}^{(k-1)} - \eta \, \nabla L(\mathbf{w}^{(k)})$$
$$\mathbf{w}^{(k+1)} = \mathbf{w}^{(k)} - \mu \mathbf{v}^{(k-1)} + (1+\mu)\mathbf{v}^{(k)}$$

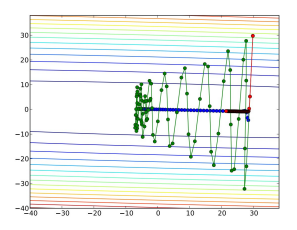


How does this compare to momentum?





Example for an advantage of NAG



GD (red), momentum (green), NAG (blue)



What if our features have different needs?



What if our features have different needs?

- Suppose some features are activated very infrequently
- ... while others are updated very often



What if our features have different needs?

- Suppose some features are activated very infrequently
- ... while others are updated very often
- We'd need individual learning rates for every parameter in the network
- Large (small) learning rates for infrequent (frequent) parameters and parameters with small (large) gradient magnitudes



AdaGrad

$$\mathbf{g}^{(k)} = \nabla L(\mathbf{w}^{(k)})$$

$$\mathbf{r}^{(k)} = \mathbf{r}^{(k-1)} + \mathbf{g}^{(k)} \odot \mathbf{g}^{(k)}$$

$$\mathbf{w}^{(k+1)} = \mathbf{w}^{(k)} - \frac{\eta}{\sqrt{\mathbf{r}^{(k)}} + \epsilon} \odot \mathbf{g}^{(k)}$$

- Adaptive Gradient
- Adaption based on all past squared gradients
- We use ⊙ to emphasize the element-wise multiplication



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- Adaptive Gradient
- Adaption based on all past squared gradients
- We use ⊙ to emphasize the element-wise multiplication
- + Individual learning rates
- Learning rate decreases too aggressively



RMSProp

$$\begin{aligned} \mathbf{g}^{(k)} &= \nabla L(\mathbf{w}^{(k)}) \\ \mathbf{r}^{(k)} &= \rho \mathbf{r}^{(k-1)} + \frac{1 - \rho}{\rho} \mathbf{g}^{(k)} \odot \mathbf{g}^{(k)} \\ \mathbf{w}^{(k+1)} &= \mathbf{w}^{(k)} - \frac{\eta}{\sqrt{\mathbf{r}^{(k)}} + \epsilon} \odot \mathbf{g}^{(k)} \end{aligned}$$

- Hinton suggests $\rho =$ 0.9, $\eta =$ 0.001
- + The aggressive decrease is fixed
- We still have to set the learning rate



Adadelta

$$\mathbf{g}^{(k)} = \nabla L(\mathbf{w}^{(k)})$$

$$\mathbf{r}^{(k)} = \rho \mathbf{r}^{(k-1)} + (1-\rho)\mathbf{g}^{(k)} \odot \mathbf{g}^{(k)}$$

$$\boldsymbol{\Delta}_{x} = -\frac{\sqrt{\mathbf{h}^{(k-1)}} + \epsilon}{\sqrt{\mathbf{r}^{(k)}} + \epsilon} \odot \mathbf{g}^{(k)}$$

$$\mathbf{h}^{(k)} = \rho \mathbf{h}^{(k-1)} + (1-\rho)\boldsymbol{\Delta}_{x} \odot \boldsymbol{\Delta}_{x}$$

$$\mathbf{w}^{(k+1)} = \mathbf{w}^{(k)} + \boldsymbol{\Delta}_{x}$$

- Suggested: $\rho = 0.95$
- + No learning rate



Adam

$$\begin{split} \mathbf{g}^{(k)} &= \nabla L(\mathbf{w}^{(k)}) \\ \mathbf{v}^{(k)} &= \mu \mathbf{v}^{(k-1)} + (1-\mu)\mathbf{g}^{(k)} \\ \mathbf{r}^{(k)} &= \rho \mathbf{r}^{(k-1)} + (1-\rho)\mathbf{g}^{(k)} \odot \mathbf{g}^{(k)} \\ \end{split}$$
 Bias correction:
$$\hat{\mathbf{v}}^{(k)} &= \frac{\mathbf{v}^{(k)}}{1-\mu^k} \quad \hat{\mathbf{r}}^{(k)} = \frac{\mathbf{r}^{(k)}}{1-\rho^k} \\ \mathbf{w}^{(k+1)} &= \mathbf{w}^{(k)} - \eta \frac{\hat{\mathbf{v}}^{(k)} + \epsilon}{\sqrt{\hat{\mathbf{r}}^{(k)}} + \epsilon} \end{split}$$

- Short for Adaptive Moment Estimation
- Suggested: $\mu =$ 0.9, $\rho =$ 0.999, $\eta =$ 0.001
- + Robustness
- · Combination w. NAG exists ("Nadam")



AMSGrad

Adam empirically observed to fail to converge to an optimal/good solution



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- Recent insight by Reddi et al. [5]: Adam (and similar methods) do not guarantee convergence for convex problems (error in original convergence proof)
- AMSGrad [5] "fixes" Adam to ensure non-increasing step size:

$$\hat{\mathbf{v}}^{(k)} = \max(\hat{\mathbf{v}}^{(k-1)}, \mathbf{v}^{(k)})$$



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- Effect has to be shown in larger experiments
- Lesson: Keep your eyes open!



Summary

- SGD + Nesterov momentum + learning rate decay
- + Often converges most reliably
- + Still used in many state-of-the-art papers
- Learning rate decay needs to be adjusted
- Adam
- + Individual learning rates
- + Learning rate very well behaved
- Loss curves harder to interpret
- Not discussed: Distributed gradient descend



Practical recommendations

- Start by using minibatch SGD with momentum
- Mostly keep to the default momentum
- Give Adam a try when you have a feeling for your data
- When in need for individual learning rates use Adam
- Start by using the default parameters for Adam
- Adjust the learning rate first
- Keep your eyes open for unusual behavior (see AMSGrad)

NEXT TIME

ON DEEP LEARNING



Coming Up

- How can we deal with spatial correlation in features?
- Why do we hear so much about convolution in neural networks?
- How can we incorporate invariances into network architectures?



Comprehensive Questions

- What are our standard loss functions for classification and regression?
- What assumptions do our standard loss functions imply?
- What is a subdifferential at a point x₀?
- How can we optimize a non-smooth convex function?
- What if somebody tells you, to use an SVM because it is superior?
- What is Nesterov Momentum?
- Describe Adam.



Further Reading

- Link for details on Maximum Likelihood estimation and the basic loss functions.
- Link [6] for insights about some loss functions
- Link [10] for a troubling insight, that deep networks can learn arbitrary random labels





References





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