### Gradient Descent and Loss Functions

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(Slides credit to David Rosenberg, He He, et al.)

NYU

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### Lecture Slides

- For those of you who want to take notes on your tablets.
- Otherwise, slides will be shared on the course website after the lecture.



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### Homework 1

- Homework 1 will be released soon. You have until Oct 1 noon (12pm) to finish.
- Submit PDF and code to Gradescope.
- Course website: https://nyu-cs2565.github.io/2024-fall/

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Review: ERM

# Our Machine Learning Setup

#### Prediction Function

A prediction function gets input x and produces an output  $\hat{y} = f(x)$ .

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# Our Machine Learning Setup

#### Prediction Function

A **prediction function** gets input x and produces an output  $\hat{y} = f(x)$ .

#### Loss Function

A loss function  $\ell(\hat{y}, y)$  evaluates an action in the context of the outcome y.

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# Risk and the Bayes Prediction Function

#### Definition

The **risk** of a prediction function  $f: \mathcal{X} \to \mathcal{Y}$  is

$$R(f) = \mathbb{E}\ell(f(x), y).$$

In words, it's the **expected loss** of f on a new example (x,y) drawn randomly from  $P_{\mathfrak{X}\times \mathfrak{Y}}$ .

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#### **Definition**

A Bayes prediction function  $f^*$  is a function that achieves the *minimal risk* among all possible functions:

$$f^* \in \operatorname*{arg\,min}_f R(f)$$
,

• The risk of a Bayes prediction function is called the **Bayes risk**.

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# The Empirical Risk

Let  $\mathfrak{D}_n = ((x_1, y_1), \dots, (x_n, y_n))$  be drawn i.i.d. from  $\mathfrak{P}_{\mathfrak{X} \times \mathfrak{Y}}$ .

#### **Definition**

The **empirical risk** of f with respect to  $\mathfrak{D}_n$  is

$$\hat{R}_n(f) = \frac{1}{n} \sum_{i=1}^n \ell(f(x_i), y_i).$$

- The unconstrained empirical risk minimizer can overfit.
  - i.e. if we minimize  $\hat{R}_n(f)$  over all functions, we overfit.

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## Constrained Empirical Risk Minimization

#### Definition

A hypothesis space  $\mathcal{F}$  is a set of functions mapping  $\mathcal{X} \to \mathcal{Y}$ .

• This is the collection of prediction functions we are choosing from.

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- ullet An empirical risk minimizer (ERM) in  ${\mathcal F}$  is

$$\hat{f}_n \in \underset{f \in \mathcal{F}}{\operatorname{arg\,min}} \frac{1}{n} \sum_{i=1}^n \ell(f(x_i), y_i).$$

- From now on "ERM" always means "constrained ERM".
- So we should always specify the hypothesis space when we're doing ERM.

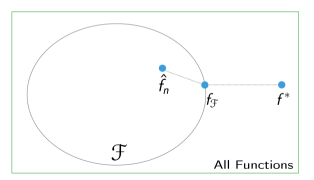
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## Error Decomposition Review

• Excess risk decomposition for function  $\tilde{f}_n$  returned by an optimization algorithm in practice:

Excess 
$$\operatorname{Risk}(\tilde{f}_n) = R(\tilde{f}_n) - R(f^*)$$

$$= \underbrace{R(\tilde{f}_n) - R(\hat{f}_n)}_{\text{optimization error}} + \underbrace{R(\hat{f}_n) - R(f_{\mathcal{F}})}_{\text{estimation error}} + \underbrace{R(f_{\mathcal{F}}) - R(f^*)}_{\text{approximation error}}$$



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- Choose a hypothesis space  $\mathcal{F}$ .
- Use an optimization method to find an empirical risk minimizer  $\hat{f}_n \in \mathcal{F}$ :

$$\hat{f}_n = \arg\min_{f \in \mathcal{F}} \frac{1}{n} \sum_{i=1}^n \ell(f(x_i), y_i).$$

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- Or find a  $\tilde{f}_n$  that comes close to  $\hat{f}_n$
- The machine learning scientist's job:
  - Choose  $\mathcal{F}$  that balances approximation and estimation error.
  - As we get more training data, we can use a bigger  $\mathcal{F}$ .

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## Setup

• Loss:  $\ell(\hat{y}, y) = (y - \hat{y})^2$ 

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- Loss:  $\ell(\hat{y}, y) = (y \hat{y})^2$
- Hypothesis space:  $\mathcal{F} = \{f : \mathbb{R}^d \to \mathbb{R} \mid f(x) = w^T x, w \in \mathbb{R}^d \}$
- Given a data set  $\mathcal{D}_n = \{(x_1, y_1), \dots, (x_n, y_n)\},\$ 
  - Our goal is to find the ERM  $\hat{f} \in \mathcal{F}$ .

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#### Objective Function: Empirical Risk

We want to find the function in  $\mathcal{F}$ , parametrized by  $w \in \mathbb{R}^d$ , that minimizes the empirical risk:

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• How do we solve this optimization problem?

$$\min_{w \in \mathbb{R}^d} \hat{R}_n(w)$$

• (For OLS there's a closed form solution, but in general there isn't.)

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## Unconstrained Optimization

### Setting

We assume that the objective function  $f : \mathbb{R}^d \to \mathbb{R}$  is differentiable.

We want to find

$$x^* = \arg\min_{x \in \mathsf{R}^d} f(x)$$

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### The Gradient

- Let  $f: \mathbb{R}^d \to \mathbb{R}$  be differentiable at  $x_0 \in \mathbb{R}^d$ .
- The gradient of f at the point  $x_0$ , denoted  $\nabla_x f(x_0)$ , is the direction in which f(x) increases fastest, if we start from  $x_0$ .
- The **gradient** of f is the partial derivatives of all dimensions:  $\nabla f(x) = [\partial f/\partial x_1(x), ..., \partial f/\partial x_d(x)].$

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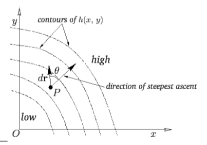


Figure A.111 from Newtonian Dynamics, by Richard Fitzpatrick.

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#### **Gradient Descent**

- Initialize  $x \leftarrow 0$ .
- Repeat:

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• until the stopping criterion is satisfied.

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- until the stopping criterion is satisfied.
- The "step size"  $\eta$  is not the amount by which we update x!
- "Step size" is also referred to as "learning rate" in neural networks literature.

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## Gradient Descent Path

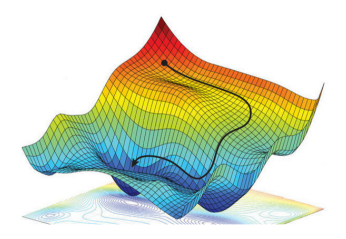


Image credit: Amini et al. Spatial Uncertainty Sampling for End-to-End Control. 2018.

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# Gradient Descent: Step Size

A fixed step size will work, eventually, as long as it's small enough

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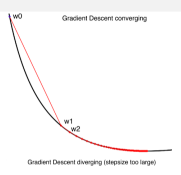
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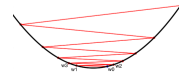
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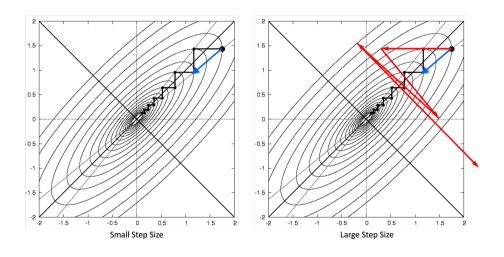
- If  $\eta$  is too large, the optimization process might diverge
- In practice, it often makes sense to try several fixed step sizes
- Intuition on when to take big steps and when to take small steps?





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# 2D Divergence example



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# Notes on Convergence

• Gradient descent with an appropriate step size converges to stationary point (derivative = 0) for differentiable functions.

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## Notes on Convergence

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- Stationary points can be (local) minima, (local) maxima, saddle points, etc.
- Gradient descent can converge to global minimum for convex functions.

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## Convergence Theorem for Fixed Step Size

#### Theorem

Suppose  $f: \mathbb{R}^d \to \mathbb{R}$  is convex and differentiable, and  $\nabla f$  is **Lipschitz continuous** with constant L > 0 (L-smooth), i.e.

$$\|\nabla f(x) - \nabla f(x')\| \le L\|x - x'\|$$

for any  $x, x' \in \mathbb{R}^d$ . Then gradient descent with fixed step size  $\eta \leq 1/L$  converges. In particular,

$$f(x^{(k)}) - f(x^*) \leqslant \frac{\|x^{(0)} - x^*\|^2}{2\eta k}.$$

This says that gradient descent is guaranteed to converge and that it converges with rate O(1/k).

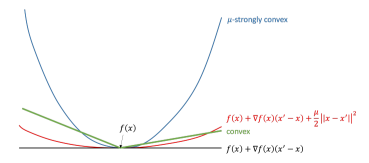
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## Strongly Convex Functions

#### Definition

A function f is  $\mu$ -strongly convex if

$$f(x') \ge f(x) + \nabla f(x) \cdot (x' - x) + \frac{\mu}{2} ||x - x'||^2$$



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# Convergence Theorem for Strongly Convex Functions

#### Theorem

If f is L-smooth and  $\mu$ -strongly convex, and step size  $0 < \eta \leqslant \frac{1}{L}$ , then gradient descent converges with the following inequality:

$$||x^{(k)} - x^*||^2 \le (1 - \eta \mu)^k ||x^{(0)} - x^*||^2$$

This means we can get linear convergence, but it depends on  $\mu$ . If the estimate of  $\mu$  is bad then the rate is not great.

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## Gradient Descent: When to Stop?

- Wait until  $\|\nabla f(x)\|_2 \le \varepsilon$ , for some  $\varepsilon$  of your choosing.
  - (Recall  $\nabla f(x) = 0$  at a local minimum.)

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## Gradient Descent: When to Stop?

- Wait until  $\|\nabla f(x)\|_2 \le \varepsilon$ , for some  $\varepsilon$  of your choosing.
  - (Recall  $\nabla f(x) = 0$  at a local minimum.)
- Early stopping:
  - evalute loss on validation data (unseen held out data) after each iteration;
  - stop when the loss does not improve (or gets worse).

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## Quick recap: Gradient Descent for ERM

- We have a hypothesis space of functions  $\mathfrak{F} = \{f_w : \mathfrak{X} \to \mathfrak{Y} \mid w \in \mathbb{R}^d\}$ 
  - Parameterized by  $w \in \mathbb{R}^d$ .

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# Quick recap: Gradient Descent for ERM

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- Finding an empirical risk minimizer entails finding a w that minimizes

$$\hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^n \ell(f_w(x_i), y_i)$$

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$$\hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^n \ell(f_w(x_i), y_i)$$

- Suppose  $\ell(f_w(x_i), y_i)$  is differentiable as a function of w.
- Then we can do gradient descent on  $\hat{R}_n(w)$

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## Gradient Descent: Scalability

• At every iteration, we compute the gradient at the current w:

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- How does this scale with *n*?
- We have to iterate over all n training points to take a single step. [O(n)]
- Can we make progress without looking at all the data before updating w?

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# Stochastic Gradient Descent

# "Noisy" Gradient Descent

- Instead of using the gradient, we use a noisy estimate of the gradient.
- Turns out this can work just fine!

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## "Noisy" Gradient Descent

- Instead of using the gradient, we use a noisy estimate of the gradient.
- Turns out this can work just fine!
- Intuition:
  - Gradient descent is an iterative procedure anyway.
  - At every step, we have a chance to recover from previous missteps.

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### Minibatch Gradient

• The full gradient is

$$\nabla \hat{R}_n(w) = \frac{1}{n} \sum_{i=1}^n \nabla_w \ell(f_w(x_i), y_i)$$

• It's an average over the **full batch** of data  $\mathcal{D}_n = \{(x_1, y_1), \dots, (x_n, y_n)\}.$ 

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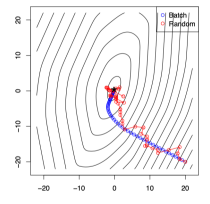
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$$\nabla \hat{R}_{N}(w) = \frac{1}{N} \sum_{i=1}^{N} \nabla_{w} \ell(f_{w}(x_{m_{i}}), y_{m_{i}})$$

### Batch vs Stochastic Methods



(Slide adapted from Ryan Tibshirani)

Rule of thumb for stochastic methods:

- Stochastic methods work well far from the optimum
- But struggle close the the optimum

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$$\mathbb{E}\left[\nabla\hat{R}_{N}(w)\right] = \nabla\hat{R}_{n}(w)$$

• The bigger the minibatch, the better the estimate.

$$\operatorname{Var}\left[\nabla \hat{R}_{N}(w)\right] = \operatorname{Var}\left[\frac{1}{N}\sum_{i}\nabla \hat{R}_{i}(w)\right] = \frac{1}{N^{2}}\operatorname{Var}\left[\sum_{i}\nabla \hat{R}_{i}(w)\right] = \frac{1}{N}\operatorname{Var}\left[\nabla \hat{R}_{i}(w)\right]$$

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- Because of vectorization, the computation cost of minibatches is sublinear

- For convergence guarantee, use **diminishing step sizes**, e.g.  $\eta_k = 1/k$
- Theoretically, GD is much faster than SGD in terms of convergence rate and number of steps:
  - much faster to add a digit of accuracy (more details later)

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  - but most of that advantage comes into play once we're already pretty close to the minimum
  - in many ML problems we don't care about optimizing to high accuracy (why?)

# Step Sizes in Minibatch Gradient Descent

### Minibatch Gradient Descent (minibatch size *N*)

- initialize w = 0
- repeat
  - randomly choose N points  $\{(x_i, y_i)\}_{i=1}^N \subset \mathcal{D}_n$

• 
$$w \leftarrow w - \eta \left[ \frac{1}{N} \sum_{i=1}^{N} \nabla_{w} \ell(f_{w}(x_{i}), y_{i}) \right]$$

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- For SGD, fixed step size can work well in practice.
- Typical approach: Fixed step size reduced by constant factor whenever validation performance stops improving (staircase decay).
- Other schedules: inverse time decay (1/t) etc.

# Convergence of SGD Theorem (Optional)

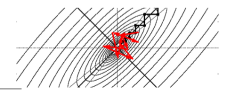
More on why we need a diminishing step size.

#### Theorem

If f is L-smooth and convex, and SGD has bounded variance  $Var(\nabla f(x^{(k)})) \leqslant \sigma^2$  for all k, then SGD with step size  $\eta \leqslant \frac{1}{I}$  satisifies:

$$\min_{k} \mathbb{E}[||\nabla f(x^{(k)}||^{2}] \leqslant \frac{f(x^{(0)}) - f(x^{*})}{\sum_{k} \eta_{k}} + \frac{L\sigma^{2}}{2} \frac{\sum_{k} \eta_{k}^{2}}{\sum_{k} \eta_{k}}$$

The extra term of variance will dominate if the step size does not decrease. <sup>1</sup>



<sup>1</sup> https://www.cs.ubc.ca/~schmidtm/Courses/540-W19/L11.pdf

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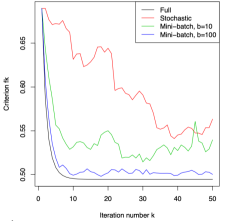
These days terminology isn't used so consistently, so when referring to SGD, always clarify the [mini]batch size.

SGD is much more efficient in time and memory cost and has been quite successful in large-scale ML.

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# Example: Logistic regression with $\ell_2$ regularization

Batch methods converge faster :

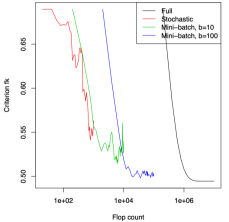


(Example from Ryan Tibshirani)

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# Example: Logistic regression with $\ell_2$ regularization

Stochastic methods are computationally more efficient:

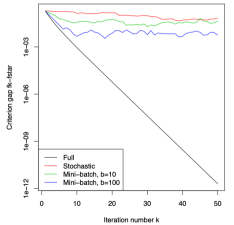


(Example from Ryan Tibshirani)

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## Example: Logistic regression with $\ell_2$ regularization

Batch methods are much faster close to the optimum:



(Example from Ryan Tibshirani)

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Loss Functions: Regression

- Examples:
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- Notation:
  - $\hat{y}$  is the predicted value (the action)
  - y is the actual observed value (the outcome)

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  - what you have to add to your prediction to get the correct answer.
- A loss  $\ell(\hat{y}, y)$  is called **distance-based** if:
  - It only depends on the residual:

$$\ell(\hat{y}, y) = \psi(y - \hat{y})$$
 for some  $\psi: R \to R$ 

2 It is zero when the residual is 0:

$$\psi(0) = 0$$

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• Distance-based losses are translation-invariant. That is,

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- Often you can transform response y so it's translation-invariant (e.g. log transform)

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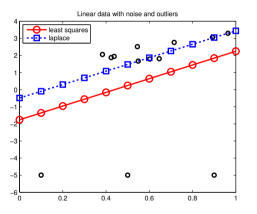
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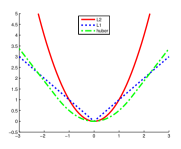
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- Outliers typically have large residuals.
- Square loss much more affected by outliers than absolute loss.

#### Loss Function Robustness

• Robustness refers to how affected a learning algorithm is by outliers.



- Square or  $\ell_2$  Loss:  $\ell(r) = r^2$  (not robust)
- Absolute or Laplace Loss:  $\ell(r) = |r|$  (not differentiable)
  - gives median regression
- **Huber** Loss: Quadratic for  $|r| \le \delta$  and linear for  $|r| > \delta$  (robust and differentiable)
  - Equal values and slopes at  $r = \delta$



# Classification Loss Functions

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How can we optimize the model output?

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- Output space  $\mathcal{Y} = \{-1, 1\}$
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The value f(x) is called the **score** for the input x.

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- In this context, f may be called a score function.
- The magnitude of the score can be interpreted as our confidence of our prediction.

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- The margin is a measure of how correct we are:
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- We want to maximize the margin.
- Most classification losses depend only on the margin (they are margin-based losses).

- If  $\tilde{f}$  is the inference function (1 if f(x) > 0 and -1 otherwise), then
- The **0-1 loss** for  $f: \mathcal{X} \to \{-1, 1\}$ :

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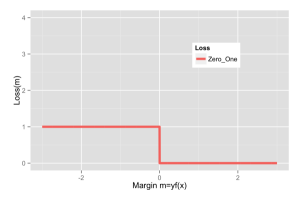
Minimizing empirical 0-1 risk not computationally feasible.

 $\hat{R}_n(f)$  is non-convex, not differentiable, and even discontinuous.

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#### Classification Losses

Zero-One loss:  $\ell_{0-1} = \mathbb{1}[m \leq 0]$ 

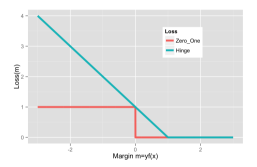


• x-axis is margin:  $m > 0 \iff$  correct classification

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## Hinge Loss

SVM/Hinge loss:  $\ell_{\text{Hinge}} = \max(1 - m, 0)$ 



Hinge is a convex, upper bound on 0-1 loss. Not differentiable at m=1.

We will cover SVM and Hinge loss in more details in future lectures.

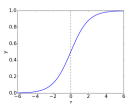
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- Two equivalent types of logistic regression losses, depending on the labels.
- If the label is 0 or 1:
- $\hat{y} = \sigma(z)$ , where  $\sigma$  is the sigmoid function, and  $z = f(x) = w^{\top}x$ .

$$\sigma(z) = \frac{1}{1 + \exp(-z)}$$



CSCI-GA 2565

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CSCI-GA 2565 55 / 60

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Remember the negative sign!

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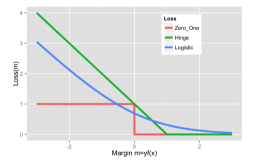
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### Logistic Loss

Logistic/Log loss:  $\ell_{\text{Logistic}} = \log(1 + e^{-m})$ 



Logistic loss is differentiable. Logistic loss always rewards a larger margin (the loss is never 0).

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• Loss 
$$\ell(f(x), y) = (f(x) - y)^2$$
.

- Loss  $\ell(f(x), y) = (f(x) y)^2$ .
- Turns out, can write this in terms of margin m = f(x)y:
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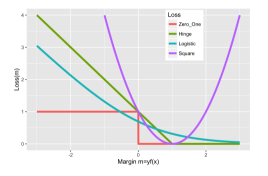
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$$= (1 - m)^{2}$$

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Heavily penalizes outliers (e.g. mislabeled examples).

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