Neural Network Basics

José R. Dorronsoro
Dpto. de Ingeniería Informática
Escuela Politécnica Superior
Universidad Autónoma de Madrid

Modeling Basics

1 Machine Learning Basics Basic Modeling

Regression Basics
Bias, Variance and Cross Validation
Basic Classification
Logistic Regression

2 Multilayer Perceptrons Rosenblatt's Percep

Classical MLPs
Neural Network Classification
MLP Regularization
Unconstrained Smooth Optimizatior
Computational Costs of MLPs

What Is Machine Learning (ML)?

- Lofty definition: make machines learn!!!
 - Have to make "machines" and "learn" more precise
- The machines of ML: mathematical input—output processes that lend themselves to some form of (numerical) parameterization
- The learning process: adjust the machine's parameters until a goal is reached
- New thing: "goal"?
 - At first sight, get something done
 - · Ultimately, to minimize some error measure
- Summing things up: a ML process tries to find a concrete mathematical/algorithmic input—output parameterized transformation that minimizes an error measure by iteratively adjusting the transformation's parameters

Where Lies ML?

- In the middle of a possibly long process chain
- Before ML starts we must
 - Go from raw to organized data: accessing, gathering, cleaning, formatting, ...
 - Go from organized to (potentially) informative data: extracting basic and derived features
- After ML finishes and start applying a model, we must perform
 - Outcome evaluation: how good/actionable is
 - Outcome exploitation: collect, organize, act
 - Individual model maintenance: monitor performance, tune hyper-parameters
 - Modeling life cycle maintenance: discard old models, introduce new ones and communicate our work/results

Supervised/Unsupervised Models

- ML model types: supervised, unsupervised
- Supervised models:
 - Targets y^p are known and the model tries to predict or estimate them
 - These known targets guide, or supervise, model building
 - Main emphasis here
- Unsupervised models:
 - There are no predetermined or supervising outputs
 - But nevertheless the model is supposed to learn relations or find structure in the data
 - Often as a first step towards a supervised model
- Semisupervised models lie in between: some patterns have targets, some don't

Regression and Classification

- Problems (usually) to be solved by ML models: regression, classification
- In both patterns come in pairs (x, y)
 - x: inputs, predictors, features, independent variables
 - y: target, response, dependent variable; numerical in regression, class labels in classification
- **Regression**: the desired output y is **regressed** into the inputs x to derive a model $\hat{y} = f(x)$
 - We want $y \simeq \hat{y}$ so having $y \hat{y}$ "small" is the natural goal
- Classification: inputs are derived from several classes C_1, \ldots, C_K , to which labels ℓ_k are assigned
 - The model now assigns a label $\ell(x)$ to an input x
 - If x is derived from C_k we want to have $\ell(x) = \ell_k$
 - Here having $\ell(x) \ell_k$ "small" may not make sense

The Boston Housing Regression Problem

- This is a first, widely used "toy" problem
- We want to estimate the median of house values over an area from some information about it which we believe relevant
- Features x: several real estate—related variables of Boston areas
 - · CRIM: per capita crime rate by town
 - RM: average number of rooms per dwelling
 - NOX: nitric oxides concentration (parts per 10 million)
 - AGE: proportion of owner-occupied units built prior to 1940
 - LSTAT: % lower status of the population
 - . .
- Target y: MEDV, median value of owner-occupied homes in \$1,000's

The MNIST Classification Problem

- This is a well known benchmark, moderately large classification problem
- We have 28 \times 28 grey scale image rasters with handwritten digits 0 to 9
- We want to build a classifier that assigns each raster to one of the 0 to 9 digits
- Features x: 28 × 28 image matrices
- Target y: labels [0, 1, ..., 9]
- Examples (0-padded to 32 × 32):







Model Parameterization

- Often individual models are selected through (ideally optimal) parameter sets
 - The parameters (weights) $w \in R^M$ select a concrete f in a model family \mathcal{F}
- Parametric models have a fixed functional form f(x) = f(x; W)
 - Simplest example: linear regression, where x has dimension d,
 M = d + 1 and W = (w₀, w)

$$f(x; w_0, w) = w_0 + \sum_{j=1}^d w_j x_j = w_0 + w \cdot x$$

- Semi-parametric models: also use weights but without a predefined functional form
 - MLPs but also RF or GBR
- Non parametric models do not use weights nor follow any broad functional form
 - Nearest Neighbor models

Issues in Model Building

- There are some initial questions when working with models from a given family F:
 - Which assumptions do they make?
 - How do they work on the input features?
 - How we do build them?
- In turn, these questions lead to another two:
 - How do we select the best model from the given family for the problem at hand?
 - How do we control the model building procedure?
- All of them address fundamental issues that require a moderately deep understanding of what is going on under the model's hood
- This understanding is usually framed in mathematical language

Regression Basics

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How to Build Regression Models

- In general we have a sample S = {x^p, y^p}, 1 ≤ p ≤ N, with x^p the features and y^p the targets
- We want to build a model $\hat{y} = f(x)$ so that $\hat{y}^p = f(x^p) \simeq y^p$;
 - I.e., we want to **regress** y to the f(x)
- The concrete f is chosen within a certain family \mathcal{F}
 - Examples here: linear regression, multilayer perceptrons (MLPs)
 - And also: SVMs, Random Forests (RF), Gradient Boosting (GB), nearest neighbor (NN)
- Natural option to ensure $f(x^p) \simeq y^p$: choose f to minimize the sample **Mean Square Error (MSE)**

$$\widehat{\mathbf{e}}(f) = \widehat{\mathbf{e}}_{\mathcal{S}}(f) = \frac{1}{2N} \sum_{p=1}^{N} (y^p - f(x^p))^2$$

This corresponds to the population error

$$e(w_0, w) = \frac{1}{2} E_{x,y}[(y - f(x))^2] = \frac{1}{2} E[(y - f(x))^2 p(x, y) dx dy]$$
$$= \frac{1}{2} \int (y - f(x))^2 p(x, y) dx dy$$

Model Estimation as Error Minimization

- Thus, we may select a model by solving $\hat{f} = \hat{f}_S = \arg\min_{f \in \mathcal{F}} \hat{e}_S(f)$
 - But this may be quite complicated mathematically and we must simplify it
- For a parametric or semiparametric f(x; W) we can write $\widehat{e}_S(f) = \widehat{e}_S(f(\cdot; W)) = \widehat{e}_S(W)$
- The problem to solve becomes

$$\widehat{\textit{W}}^* = \widehat{\textit{W}}^*_{\mathcal{S}} = \arg\min_{\textit{W}} \widehat{\textit{e}}_{\mathcal{S}}(\textit{W}), \; \text{ i.e., } \; \widehat{\textit{e}}_{\mathcal{S}}(\widehat{\textit{W}}^*) \leq \widehat{\textit{e}}_{\mathcal{S}}(\textit{W}) \; \forall \textit{W}$$

· For linear regression (LR) the sample error is

$$\widehat{e}(w_0,w) = \frac{1}{2N} \sum_{\rho} (y^{\rho} - w_0 - w \cdot x^{\rho})^2$$

• Minimizing the sample error $\hat{e}_{S}(W)$ becomes a problem in mathematical **optimization**

Regression Assumptions

- **Key assumption**: x and y are related as $y = \phi(x) + n$ where
 - $\phi(x)$ is the **true** underlying function
 - *n* is **additive noise** with 0 mean and finite variance σ_N^2
- Our sample is just a particular instance of a deeper sample generation process
- Thus x, n are produced by random variables X, N
 - And so is y, given by $Y = \phi(X) + N$
- Moreover, X and N are independent distributions with densities q(x), ν(n)
- Thus, X and Y (or X and N) have a joint density

$$p(x, y) = p(x, \phi(x) + n) = q(x) \nu(n) = q(x) \nu(y - \phi(x))$$

MSE Decomposition

• We can decompose the MSE error of any model f as

$$2\mathsf{mse}(f) = E_{x,y}[(y - f(x))^{2}] = \int (n + \phi(x) - f(x))^{2} q(x) \nu(n) dx dn$$

$$= \int (n^{2} + 2n(\phi(x) - f(x)) + (\phi(x) - f(x))^{2}) q(x) \nu(n) dx dn$$

$$= \int n^{2} \nu(n) dn + \int (\phi(x) - f(x))^{2} q(x) dx +$$

$$2 \left(\int n\nu(n) dn \right) \left(\int (\phi(x) - f(x)) q(x) dx \right)$$

$$= \sigma_{N}^{2} + E_{x}[(\phi(x) - f(x))^{2}]$$

- Thus for any model we have $mse(t) \ge \sigma_N^2$ always
 - Beware of zero regression errors!!
- And we should focus on achieving on $f \simeq \phi$ (which we don't know!!)

The Best Regression Model

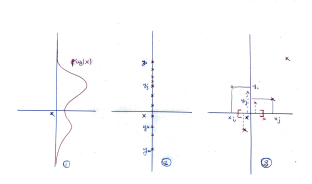
• It is easy to see that the best f is simply $f(x) = E_y[y|x]$, for

$$E_{y}[y|x] = E_{n}[\phi(x) + n] = \int (\phi(x) + n) \nu(n) dn = \phi(x)$$

- Have we finished? In theory yes; in practice, not at all!!!
 - We do not know ν and, thus, cannot compute the required integral
 - If we would have several M values y^j for any x, we could try $\widehat{\phi}(x) = \frac{1}{M} \sum_{i=1}^{M} y^j$
 - But this doesn't happen either
- Now we have two options:
 - Try to stretch the E[y|x] approach, for instance, by k-NN Regression
 - Forget about it and get back to get models f for which we hope $f \simeq \phi$

From The Best Regressor to k-NN

 The left hand side shows the ideal situation but the right one is what we should expect in practice



k-NN Regression

- We can try to use the y^p values of several x^p close to a new x
- We fix k neighbors x^{p_1}, \ldots, x^{p_k} of x and estimate $\hat{y} = \hat{y}(x)$ as

$$\widehat{y}(x) = \frac{1}{k} \sum_{j=1}^{k} y^{p_j}$$

- $\hat{y}(x) = \hat{Y}_k^{NN}(x)$ is the *k*-Nearest Neighbor (NN) regressor
 - This can be refined to weighted versions, such as

$$\widehat{y}(x) = \frac{1}{C_k(x)} \sum_{j=1}^k \frac{1}{\|x^{p_j} - x\|^2} y^{p_j}$$

with
$$C_k(x) = \sum_{j=1}^k \frac{1}{\|x^{p_j} - x\|^2}$$
 a normalizing constant

- But we need that close predictors give predictions that are also close, and that there are enough of them close by
 - This is very unlikely

The Curse of Dimensionality

- Even for low dimensions and large samples, the sample space is essentially empty
- Assume we have 1,000 d-dimensional x patterns whose features have values between 1 and 10
 - In dimension d = 1 there are 100 patterns per unit length
 - But when d = 3 we have just 1 pattern per volume unit
 - And if d = 6 we have just 1 pattern per 1,000 units of volume
 - And in dimension 10 (not a big one nowadays) we have just ... !!!
- Thus, for most problems, there never will be enough close points
- As a consequence, to get k observations we may go too far away from x and the average will not be meaningful
- Therefore, unless we deal with violently non–linear problems, a simple linear model may be better than k–NN regression for moderate dimensions

Linear Models

• Assuming $x \in R^d$, recall the basic linear model is

$$f(x) = w_0 + \sum_{1}^{d} w_i x_i = w_0 + w \cdot x$$

- w_0 complicates notation; to drop it we center x and y so that $E[x_i] = E[y] = 0$; then $w_0 = 0$
- Then we are left with the simpler **homogeneous** model $f(x) = w \cdot x$
- In practice we will always normalize x, for instance to have 0 mean and 1 standard deviation (std) on each feature
 - But not y if we may help it (or it is easy to reverse this)
- But: how do we find w?

1-dimensional Linear Regression (LR)

- Assume that features X and target Y are centered, i.e., have 0 means
- For 1-dimensional patterns x the LR model then becomes f(x) = w x
- And the error is then the function

$$\widehat{e}(w) = \frac{1}{2N} \sum_{p=1}^{N} (w \, x^p - y^p)^2 = \frac{1}{2N} \sum_{p} (w^2 \, (x^p)^2 - 2x^p \, y^p \, w + (y^p)^2$$

$$= \frac{1}{2} \left(\frac{1}{N} \sum_{p} (x^p)^2 \right) w^2 - \left(\frac{1}{N} \sum_{p} x^p \, y^p \right) w + \frac{1}{2N} \sum_{p} (y^p)^2$$

- Thus, $\widehat{e}(w) = aw^2 + bw + c$ with a > 0 and it has obviously a minimum w^*
- To find it we just solve $\hat{e}'(w) = 0$

Solving $\hat{e}'(w) = 0$

• We first compute $\hat{e}'(w)$, for which we have

$$\widehat{e}'(w) = w \left(\frac{1}{N} \sum_{\rho} (x^{\rho})^2\right) - \frac{1}{N} \sum_{\rho} x^{\rho} y^{\rho}$$

• The optimal w^* solves $\hat{e}'(w) = 0$ and is given by

$$w^* = \frac{\frac{1}{N} \sum_{\rho} x^{\rho} y^{\rho}}{\frac{1}{N} \sum_{\rho} (x^{\rho})^2} = \frac{\frac{1}{N} X^t Y}{\frac{1}{N} X^t X} = \frac{\operatorname{covar}(x, y)}{\operatorname{var}(x)}$$

where X and Y denote the $N \times d$ data matrix (vector here) and the $N \times 1$ target vector

General Linear Regression

- Assume again that X and Y are centered
- The LR model becomes now $f(x) = \sum_{i=1}^{d} w_i x_i = w \cdot x$
- If Y is the N × 1 target vector and we organize the sample S in a N × d data matrix X, the sample mse is given by

$$\widehat{e}(w) = \frac{1}{2N} \sum_{p} (w \cdot x^{p} - y^{p})^{2} = \frac{1}{2N} (Xw - Y)^{t} (Xw - Y)$$
$$= \frac{1}{2N} (w^{t} X^{t} Xw - 2w^{t} X^{t} Y + Y^{t} Y)$$

- We have now a **quadratic form** $w^t A w + w^t b + c$, with A a $d \times d$ matrix, b a d vector and c a scalar
- Now we have to solve $\nabla \widehat{e}(w) = 0$, i.e., $\frac{\partial \widehat{e}}{\partial w_i}(w) = 0, 1 \le i \le d$
- It is easy to see that

$$\nabla \widehat{e}(w) = \frac{1}{N} X^t X w - \frac{1}{N} X^t Y = \widehat{R} w - \widehat{b}$$

Solving the Linear Equations

• The optimal \widehat{w}^* must verify $\nabla \widehat{e}(\widehat{w}) = \widehat{R} \ \widehat{w} - \widehat{b} = 0$, where

$$\widehat{R} = \frac{1}{N} X^t X, \ \widehat{b} = \frac{1}{N} X^t Y$$

Over the original, non-centered data matrix we have

$$\widehat{R} = \frac{1}{N}(X - \overline{X})^t(X - \overline{X});$$

i.e., \widehat{R} is the sample covariance matrix

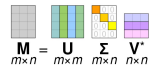
• If \widehat{R} is **invertible**, we just solve the linear system \widehat{R} $\widehat{w} = \widehat{b}$ to get

$$\widehat{w}^* = \widehat{R}^{-1}\widehat{b} = (X^tX)^{-1}X^tY = \operatorname{covar}(X)^{-1}\operatorname{covar}(X,Y)$$

- We won't but we should write \widehat{w}_{S}^{*} , as it is **sample-dependent**
- \widehat{R} is oftent inverted through the **Singular Value Decomposition** (SVD) of the data matrix X



- The **Singular Value Decomposition** (SVD) of a (complex) $m \times n$ matrix M, $m \ge n$, is $M = UDV^*$ where
 - U and V are orthonormal (unitary) matrices with dimensions m × n, n × n respectively
 - D is an $m \times n$ matrix, diagonal on its first n rows and with zeros in its last m n ones
- Graphically, the SVD decomposition of M is



Taken from Wikipedia

- The columns of U and V are called the left and right singular vectors; the diagonal elements in D are the singular values
- We may interpret this image as M performing first a rotation with V*, then feature dilations with Σ (our D) and, finally, another rotation with U
 - Actually both are a roto-reflections

SVD II

- It is easy to see that we can also write $M = \widetilde{U}\widetilde{D}V^*$, with \widetilde{U} having the first n columns of U and \widetilde{D} the top diagonal block of D
 - Assuming M to be real and with a slight abuse of language, we will
 write M = UDV^t
- In our case we can write X = UDV^t, with dimensions N × d, d × d and d × d respectively
- Then $X^tX = VDU^tUDV^t = VD^2V^t$ and $(X^tX)^{-1} = VD^{-2}V^t$ • D^{-2} is a diagonal matrix with $(D^{-2})_{ii} = 1/D_{ii}^2 = 1/\delta_i^2$
- The columns of *U* and *V* are unique up to a sign change
 - So different packages may give seemingly different U and V

Gradient Descent MSE Minimization

- Computing the covariance matrix has a O(N × d²) cost and invert it has a O(d³) cost
 - For big data problems it may not possible to solve analytically the normal equation $\nabla \hat{e}(w) = 0$
- The simplest numerical alternative is gradient descent:
 - Starting from some random w^0 , we iteratively compute

$$w^{k+1} = w^k - \rho_k \nabla \widehat{e}_B(w^k) = w^k - \frac{\rho}{n_B} \left(X_B^t X_B w^k - X_B^t Y \right)$$

over a **mini-batch** X_B with n_B samples and batch error \hat{e}_B

- Component wise: $w_i^{k+1} = w_i^k \rho_k \frac{\partial \hat{e}_B}{\partial w_i}(w^k)$
- ρ_k is the learning rate
- If $w^k \to w^*$, then $\nabla \widehat{e}(w^*) = 0$
 - Since our problems have obviously minima, this should be enough

By the Way ...

- We have just built our first neural network: the linear regression architecture can be seen as a multilayer perceptron (MLP) with no hidden layers
- Thinking of this as an MLP seems far-fetched, as we are missing the hidden layers and the much more complicated gradient computation
- But the layer architecture is there and we also have learning as iterative error minimization, i.e., training
- In fact, we will see that MLPs perform linear regression on the last hidden layer outputs
- And the process from the input to the last hidden layer produces an enhanced feature representation upon which the regression error should be better

Measuring Model Fit

- First option: **Root Square Error** $RSE = \sqrt{\frac{1}{N} \sum (y^p \widehat{y}^p)^2}$
- OK, but how to compare this? We must always have a base model to benchmark our results
- Simplest model: a constant $\hat{y} = w_0$, which reduces to predict the mean $\overline{y} = \frac{1}{N} \sum_{1}^{N} y^p$, with square error

$$\frac{1}{N}\sum (y^p - \overline{y})^2 = \operatorname{Var}(y)$$

We can compare our model against this base model by computing

$$\frac{\sum (y^{p} - \hat{y}^{p})^{2}}{\sum (y^{p} - \overline{y})^{2}} = \frac{\frac{1}{N} \sum (y^{p} - \hat{y}^{p})^{2}}{\frac{1}{N} \sum (y^{p} - \overline{y})^{2}} = \frac{RSE^{2}}{Var(y)}$$

• The widely used R^2 coefficient is simply $R^2 = 1 - \frac{RSE^2}{Var(y)}$

Regularization

- Our regression solution $\widehat{w}^* = (X^t X)^{-1} X^t Y$ won't work if $X^t X$ is not invertible
 - · For instance, when some features are highly correlated
 - We can at most say that X^tX is **positive semidefinite**
- We could fix this working instead with $X^tX + \alpha I$ for some $\alpha > 0$
 - This is then always positive definite and we can thus invert it
- To make this practical, note that $\widehat{\mathbf{w}}^* = \left(\mathbf{X}^t \mathbf{X} + \alpha \mathbf{I}\right)^{-1} \mathbf{X}^t \mathbf{Y}$ minimizes

$$e_R(w) = \frac{1}{2N} \sum_p (y^p - w \cdot x_p^p)^2 + \frac{\alpha}{2} ||w||^2,$$

- This is the Ridge Regression problem
 - Our first example of regularization, a key technique in Machine Learning
 - All ML models must be regularized in some way
- Important issue: how to find the right choice for α ?

Takeaways on Linear Regression

- We introduced supervised models
- We have reviewed the essentials of the linear regression model (always the first thing to try)
- We have considered model estimation as a problem on error minimization
- We have seen how to build linear models analytically and numerically
- 5 We have seen how to measure model fit
- 6 We have introduced regularization

Bias, Variance and Cross Validation

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Sample Bias and Variance

- Remember: all the models f we will build are sample dependent, i.e., f = f_S
- With several **independent** samples S_1, \ldots, S_M , it is natural to use as our best final model the average of their associated $\widehat{f}_{S_m}(x)$ models, i.e.,

$$\frac{1}{M}\sum_{1}^{M}\widehat{f}_{S_m}(x)\simeq E_{S}[\widehat{f}_{S}(x)]=\widehat{f}_{N}(x)$$

- Here we assume x fixed and take the (theoretical) expectation $E_S[\widehat{f}_S(x)]$ over all samples S of size N
- $\widehat{f}_N(x) = E_S[\widehat{f}_S(x)]$ is our ideal **best model** and $\phi(x) \widehat{f}_N(x)$ is its **bias**
- The **variance** of the $\hat{f}_S(x)$ estimates is then

$$V_N(x) = E_S \left[(\widehat{f}_S(x) - \widehat{f}_N(x))^2 \right]$$

Bias Versus Variance

- Recall that our goal is to estimate ϕ in our regression model $y = \phi(x) + n$
- Ideally we would like to have a model such that

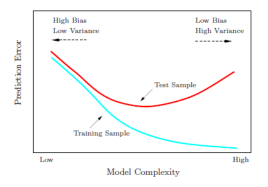
$$\widehat{f}_N(x) - \phi(x) \simeq 0,$$

i.e., a model with small bias

- This should be achievable if we have rich, highly flexible models
- Or with essentially no regularization (or both!)
- But we would also like to have a robust model building procedure which results in a small variance V_N(x) ≈ 0
 - This should be achievable if models are simple with few parameters
 - Or with more severe regularization if not
- Obviously both goals are contradictory to a large extent

The Bias-Variance Tradeoff

 There is thus a tradeoff between bias (low for complex models) and variance (low for simple models)



Taken from Hastie et al., p. 38

Bias vs Variance in k-NN Regression

• Recall that in k-NN regression we choose the k nearest neighbors x^{p_1}, \ldots, x^{p_k} of x and estimate $\hat{y} = \hat{y}(x)$ as

$$\widehat{y}(x) = \frac{1}{k} \sum_{j=1}^{k} y^{\rho_j}$$

- Here the parameter that controls the tradeoff is just k
- If k = N, the sample size, the N-NN estimator is just the mean:
 Y_N^{NN} = ȳ, with very small variance but large bias (it's an obviously bad model!!)
- If k = 1, the 1-NN estimator will have smaller bias but a large variance: changing the sample is very likely to change the sample point nearest to x

Bias vs Variance in Ridge Regression

Recall that the Ridge Regression loss is

$$e_R(w) = \frac{1}{2N} \sum_{p} (y^p - w \cdot x_p^p)^2 + \frac{\alpha}{2} ||w||^2,$$

- Here the parameter that controls the tradeoff is α
- If $\alpha \gg 1$, any non zero w implies a large regularization penalty
 - It is thus likely that $w \simeq 0$ and the Ridge model reduces again to the mean \overline{y} , with small variance but possibly a large bias
- But if $\alpha \simeq 0$, w can wander on the entire \mathbf{R}^d
 - The bias will be then smaller, but the weights w_S and $w_{S'}$ from different samples may be very different, particularly if the covariance is nearly singular

Evaluating Expected Performance

- Recall that we build a model over a train sample but apply it on a test sample
- Thus, before we start applying a model, we should have a reasonably accurate idea of its performance in practice
- I.e., we want to estimate the model's generalization performance
- Estimating the generalization performance only over the sample S used for training results in misleading error values
- The preceding suggests to try to have M independent subsamples S_m and then
 - To compute $\widehat{f}_M(x) = \frac{1}{M} \sum_m \widehat{f}_{S_m}(x) \simeq \widehat{f}_N(x)$
 - To get the error estimate $\hat{e} = \frac{1}{N} \sum_{\rho} (y^{\rho} \hat{f}_{M}(x^{\rho}))^{2}$ over a new, **unseen** sample $S' = \{(x^{\rho}, y^{\rho})\}$
- But since usually we only have a single S, we "simulate" this by Cross Validation (CV) to get our first realistic generalization error estimates

Cross Validation I

Graphically, the general schema of Cross Validation is the following



From Scikit-learn

Cross Validation II

- In more detail, in Cross Validation (CV) and for a global sample S
 we
 - Randomly split $S = S_{tr-val} \cup S_{ts}$
 - Randomly split S_{tr-val} in M subsets S₁,..., S_M
 - Work with *M* **folds**: pairs (S_m, S_m^c) , with

$$S_m^c = S - S_m = \cup_{i \neq m} S_i$$

- Build M different models using the S_m^c as training subsets
- Compute their errors e_m on the folds' validation subsets S_m
- Use these errors' average as a first estimate of the true model performance
- Test it if possible on S_{ts}
- CV can and must be used in any model building procedure
 - Most data science packages have tools to simplify this
- We will also use CV to find **optimal model hyper–parameters** such as α in Ridge Regression

Grid Hyper-parameter Selection

- Consider for Ridge regression a hyperparameter range $\alpha \in [a, A]$, with 0 < a < A
 - In general $a \simeq 0$, $A \gg 1$
 - $\alpha = a$: essentially no penalty, small bias and high variance
 - $\alpha = \Lambda$: large penalty, small variance but high bias
- Select an L+1 point **grid** $\{\alpha_0 = a < \alpha_1 < \ldots < \alpha_L = A\}$
 - The α_i can be equi-spaced, log equi-spaced, random, ...
- At each α_{ℓ}
 - Train M models on the S_m^c using the hyperparameter α_ℓ
 - Average their test errors e_m on the S_m to get the error $e(\alpha_\ell)$ at α_ℓ
- And choose the final (hopefully) optimal hyperparameter α^* as

$$\alpha^* = \operatorname{arg\;min}_{0 \leq \ell \leq L} e(\alpha_\ell)$$

• α^* gives the model with the best expected generalization among all possible α choices

Takeaways on Bias, Variance and CV

- We have stressed that any model estimation is sample-dependent and that this has to be controlled
- We have introduced the bias and variance as the two key components of any model error
- 3 We have discussed bias-variance trade-off
- We have introduced Cross Validation here as a tool to estimate a model's generalization performance
- We have also introduced Cross Validation as a tool to estimate a model's hyper-parameters

The Classification Model

1 Machine Learning Basics

Basic Modeling
Regression Basics
Bias, Variance and Cross Validation

Basic Classification

2 Multilayer Perceptrons

Classical MLPs
Neural Network Classification
MLP Regularization
Unconstrained Smooth Optimization
Computational Costs of MLPs

Regression vs Classification

- Recall that in regression we have numerical continuous targets y
 and want our predictions ŷ to be as close to y as possible
- But in classification we have a finite number of labelled targets for which "selection by closeness" doesn't make sense
- Natural alternative: select the most probable label given the pattern x we have just received
 - The concrete labels used for targets do not matter much anymore
 - Model learning should thus be "target" agnostic
 - And good probability estimates should be quite useful
- Let's analyze this in an example

A First Problem: Pima Indian Diabetes

- We want to diagnose whether a person may have diabetes from some clinical measures
- Features x: clinical measures
 - numPregnant
 - bloodPress
 - massIndex
 - age ...
- Target y: 0 (no diabetes), 1 (diabetes)
- Clear goal but perhaps too radical
- Better: try to estimate the probability P(1|x) of having diabetes depending on the features x we measure

Classification Setup

- We have random patterns ω from M classes, $C_1, \ldots C_M$
- Over each pattern we "measure" d features $x = x(\omega) \in \mathbb{R}^d$
 - x inherits the randomness in ω and becomes a random variable
- A ω has a **prior probability** π_m of belonging to C_m
- Inside each class C_m there is a **conditional class density** f(x|m) that "controls" the appearance of a given x
- The π_m and f(x|m) determine the **posterior probability** P(m|x) that x comes from class C_m
- **Intuition:** we should assign x to the class with the largest P(m|x), that is, work with the classifier

$$\delta(x) = \arg\max_{m} P(m|x)$$

Computing Posterior Probabilities I

- Bayes rule: $P(B|A) = \frac{P(A \cap B)}{P(A)}$
- This requires to work with probabilities, not densities, but $P(\lbrace x \rbrace) = P(m \cap \lbrace x \rbrace) = 0$ and

$$P(m|x) = \frac{P(m \cap \{x\})}{P(\{x\})} = \frac{0}{0} = \dots???$$

But we can use the approximation

$$P(m|x) \simeq P(m|B_{r}(x)) = \frac{P(C_{m} \cap B_{r}(x))}{P(B_{r}(x))} = \frac{P(B_{r}(x)|m)P(C_{m})}{P(B_{r}(x))}$$

$$= \frac{\pi_{m}P(B_{r}(x)|m)}{P(B_{r}(x))} = \pi_{m}\frac{\int_{B_{r}(x)}f(y|m)dy}{\int_{B_{r}(x)}f(z)dz}$$

where we assume that features x are measured independently from classes m

Computing Posterior Probabilities II

• Remember the Fundamental Theorem of Calculus: if $F(x) = \int_a^x f(y)dy$,

$$\lim_{\epsilon \to 0} \frac{1}{2\epsilon} \int_{x_0 - \epsilon}^{x_0 + \epsilon} f(y) dy = \frac{dF}{dx}(x_0) = f(x_0)$$

In d dimensions it becomes

$$g(w) = \lim_{r \to 0} \frac{1}{|B_r(w)|} \int_{B_r(w)} g(z) dz$$

Putting everything toghether, we arrive

$$\begin{array}{rcl}
P(m|x) & = & \lim_{r \to 0} P(m|B_r(x)) = \pi_m \lim_{r \to 0} \frac{\int_{B_r(x)} f(y|m) dy}{\int_{B_r(x)} f(z) dz} \\
& = & \pi_m \lim_{r \to 0} \frac{\frac{1}{|B_r(x)|} \int_{B_r(x)} f(y|m) dy}{\frac{1}{|B_r(x)|} \int_{B_r(x)} f(z) dz} = \frac{\pi_m f(x|m)}{f(x)}
\end{array}$$

The Obviously Optimal Classifier

• This suggests that a natural choice is the **classifier** δ_B

$$\delta_B(x)$$
 = arg max_m $P(m|x)$ = arg max_m $\frac{\pi_m f(x|m)}{f(x)}$
 = arg max_m $\pi_m f(x|m)$

- With some extra work we can show that this **Bayes Classifier** δ_B defines an optimal solution (in some precise sense) of the classification problem
- But ... this doesn't look too practical, for we do not know either π_m or (much harder) f(x|m)

Approximating the Bayes Classifier

- To define δ_B we need to know the prior probabilities π_m and the prior densities f(x|m)
- A reasonable choice for π_m is $\widehat{\pi}_m = \frac{N_m}{N}$, where N_m is the number of patterns of C_m in the sample
- But effective multidimensional density estimates are rather difficult, because of the curse of dimensionality
 - Densities generalize histograms
 - Good histograms need accurate counts of elements nearby
 - But in high dimensions there won't be nearby elements!!
- Options:
 - Restrict possible density models: logistic regression
 - Assume no model and apply a Nearest Neighbor (NN) strategy

The *k*–NN Classifier

- Very simple: at any x consider the subset N_k(x) of its k closest sample points and
 - Let $n_m(x)$ the number of elements of class m in $N_k(x)$
 - Notice that $0 \le n_m(x) \le k$
 - Define $\delta_{kNN}(x) = \arg\max_{m} n_{m}(x)$
- That is, $\delta_{kNN}(x)$ assigns x to the class that has more patterns in $N_k(x)$
- We can partially justify this definition from a Bayesian point of view
- Assume that $B_r(x)$ is the smallest ball that contains $N_k(x)$ and consider the approximations
 - $P(B_r(x)|m) \simeq \frac{n_m(x)}{N_m}$
 - Similarly, $P(B_r(x)) \simeq \frac{k}{N}$
 - And $\pi_m \simeq \frac{N_m}{N}$

A k-NN Example

- Assume N = 200, $N_+ = 150$, $N_{\circ} = 50$
- Set k = 5, fix x and assume $n_{+}(x) = 3$, $n_{\circ}(x) = 2$
- Then $\pi_+ = \frac{3}{4}$, and for our hypothetical $B_r(x)$ we have
 - $P(B_r(x)) = \frac{5}{200} = \frac{1}{40}$
 - $P(B_r(x)|+) = \frac{3}{150} = \frac{1}{5}$
- And, therefore

$$P(+|x) \simeq P(+|B_r(x)) = \frac{\pi_+ P(B_r(x)|+)}{P(B_r(x))} = \frac{\frac{3}{4}\frac{1}{50}}{\frac{1}{40}}$$

$$= \frac{3}{5}$$

k-NN and the Bayes Classifier

We then have

$$P(m|x) \simeq P(m|B_r(x)) = \frac{\pi_m P(B_r(x)|m)}{P(B_r(x))}$$
$$\simeq \frac{N_m}{N} \frac{n_m(x)}{N_m} \frac{1}{\frac{k}{N}} = \frac{n_m(x)}{k}$$

• Therefore δ_{kNN} might be close to δ_B , for

$$\delta_{kNN}(x) = \operatorname{arg max}_{m} n_{m}(x) = \operatorname{arg max}_{m} \frac{n_{m}(x)}{k}$$

$$\simeq \operatorname{arg max}_{m} P(m|x) = \delta_{B}(x)$$

Some *k*–NN Issues

- Q1: How do we choose k? Using CV, of course
- There are no closed form solution and we have to deal again with the bias-variance tradeoff
 - Small variance with large k: if k = N, k-NN classification returns the majority class
 - Small bias with small k: if k = 1 a point very close to x should be in the same class
 - But also large variance: the nearest point to x in another sample may well belong to a different class
- Q2: Is k-NN always meaningful?
 - Well, only if classes are concentrated and far from each other
 - In other words, when the classification problem is easy!!

Measuring Classifier Accuracy

- Consider a two class problem with labels y = 0, 1
- We will call patterns with label 1 positive and those with label 0 negative
 - Usually the positive patterns are the interesting ones: sick people, defaulted loans, . . .
- Let $\hat{y} = \hat{y}(x)$ the label predicted at x; we say that x is a
 - True Positive (TP) if $y = \hat{y} = 1$
 - True Negative (TN) if $y = \hat{y} = 0$
 - False Positive (FP) if y = 0 but $\hat{y} = 1$
 - False Negative (FN) if y = 1 but $\hat{y} = 0$
- These values are combined in many performance measures

Classifier Metrics

- The classifier **accuracy** is $acc = \frac{TP + TN}{N}$
 - The first thing to measure but ...
- It may not be too significant when the number N₀ of negatives is >>> N₁, the number of positives; then
 - The classifier $\delta(x) = 0$ will have a high accuracy $N_0/N \simeq 1$
 - But it will also be useless!!
- First variant: Precision, Recall
 - **Recall**: TP/(TP + FN), i.e., the fraction of positives detected
 - **Precision**: TP/(TP + FP), i.e., the fraction of true alarms issued
- Recall measures how many positive cases we recover, i.e., how effective is our method
- Precision measures the effort we need for that, i.e., its efficiency
- Ideally: high recall, high precision (i.e., effective and efficient!!)
- Another way of presenting the performance is through the confusion matrix

The Confusion Matrix

Standard layout

	P' (Predicted)	N' (Predicted)
P	True Positive	False Negative
(Actual)		
N	False Positive	True Negative
(Actual)		

- Other layouts:
 - Positives (with label 1) at bottom (as done in confusion_matrix of sklearn)
 - Predicted values in rows, real values in columns
- Accuracy and the confusion matrix can be extended to multi-class problems
 - But not other concepts such as recall or precision

Logistic Regression (LogR)

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Linear Regression for Classification?

- k–NN Classifier is simple but also crude; have to look elsewhere
- Building a regression model with targets given by some coding of class labels usually doesn't make sense
- However, for a binary 0–1 response, it can be shown that the $w_0 + w \cdot x$ obtained using linear regression is in fact an estimate of P(1|x)
 - We may thus fix a threshold δ_0 and decide 0 if $w_0 + w\dot{x} < \delta_0$ and 1 otherwise
 - However, we may end up with probability estimates less than 0 o bigger than 1!!!
- Better idea: try to transform the linear output $w_0 + w \cdot x \in (-\infty, \infty)$ into a probability $P(1, x) \in (0, 1)$
 - Logistic regression does this through the logistic or sigmoid function

Logistic Regression (LogR)

We assume

$$P(1|x) = P(1|x; w_0, w) = \frac{1}{1 + e^{-(w_0 + w \cdot x)}}$$

- Then $0 \le P(1|x) \le 1$ for any x
- We then have

$$P(0|x) = 1 - P(1|x) = \frac{e^{-(w_0 + w \cdot x)}}{1 + e^{-(w_0 + w \cdot x)}} = \frac{1}{1 + e^{w_0 + w \cdot x}}$$

- Notice that if $w_0 + w \cdot x = 0$, P(1|x) = P(0|x) = 0.5
- The ratio $\frac{P(1|x)}{P(0|x)} = e^{w_0 + w \cdot x}$ is called the **odds** of x and its log the **log odds** or **logit**
- Thus, the basic assumption in LogR is that the logit is a linear function w₀ + w · x of x
- We have the model f(x; w); we need a loss function L(w₀, w) to minimize for which we use the sample's likelihood

Estimating w_0^* , w^*

- Assume a single sample x, y and two possible model coefficients w_0, w and w'_0, w'
- Denoting by $p = P(y|x; w_0, w)$ and $p' = P(y|x; w'_0, w')$, it is clear that we should prefer w_0 , w if p > p' and w'_0 , w' if not
 - In other words, we prefer the coefficients that give a higher posterior probability
- For an **independent** sample $S = \{(x^p, y^p)\}$, its joint probability under a posterior model $p = P(y|, x, w_0, w)$ is

$$P(Y|X; w_0, w) = \prod_{p=1}^N P(y^p|x^p; w_0, w)$$

• And, again, given two possible model coefficients w_0 , w and w'_0 , w', we should prefer w_0 , w iff

$$P(Y|X; w_0, w) > P(Y|X; w'_0, w')$$

Sample's Likelihood

• Therefore, we can estimate the optimal w_0^* , w^* as

$$\textit{w}_{0}^{*}, \textit{w}^{*} = \text{arg max}_{\textit{w}_{0},\textit{w}}\textit{P}(\textit{Y}|\textit{X};\textit{w}_{0},\textit{w})$$

• By the independence assumption we have

$$P(Y|X; w_0, w) = \prod_{p=1}^{N} P(y^p|x^p; w_0, w)$$

$$= \left\{ \prod_{y^p=1} P(1|x^p) \right\} \left\{ \prod_{y^p=0} P(0|x^p) \right\}$$

$$= \prod_{p=1}^{N} P(1|x^p)^{y^p} P(0|x^p)^{1-y^p}$$

because

• If
$$y^p = 1$$
, $P(1|x^p) = P(1|x^p)^{y^p} = P(1|x^p)^{y^p}$ $P(0|x^p)^{1-y^p}$, and

• If
$$y^p = 0$$
, $P(0|x^p) = P(0|x^p)^{1-y^p} = P(1|x^p)^{y^p} P(0|x^p)^{1-y^p}$

Max Log-Likelihood Estimation

• The **log–likelihood** of w_0 , w given S is then

$$\ell(w_0, w; S) = \log P(Y|X; w_0, w)$$

$$= \sum_{p} \{y^p \log p(1|x^p) + (1 - y^p) \log p(0|x^p)\}$$

$$= \sum_{p} y^p \log \frac{p(1|x^p)}{p(0|x^p)} + \sum_{p} \log p(0|x^p)$$

$$= \sum_{p} y^p (w_0 + w \cdot x^p) - \sum_{p} \log(1 + e^{w_0 + w \cdot x^p})$$

• We can thus estimate the optimal \widehat{w}_0^* , \widehat{w}^* as

$$\widehat{w}_0^*, \widehat{w}^* = \operatorname{arg\ min}_{w_0, w} - \ell(w_0, w; S)$$

Extra bonus: −ℓ is a convex differentiable function of (w₀, w) and, thus, it is enough to solve ∇ℓ(w₀, w) = 0

Newton-Raphson Solution

- However, $\nabla \ell(w_0, w) = 0$ doesn't admit a closed form solution, but only an iterative, numerical one
- We apply the Newton-Raphson iterative method, here equivalent to the general Newton method for function minimization
- For a general weight vector w, and starting with an initial random w⁰, Newton's iterations are

$$\mathbf{w}^{k+1} = \mathbf{w}^k + (\mathcal{H}_{\ell}(\mathbf{w}^k))^{-1} \nabla \ell(\mathbf{w}^k)$$

- $\mathcal{H}_{\ell}(w^k)$ denotes the **Hessian** of ℓ at w^k , which may or may not be invertible
 - Everything is fine if the w^k are close enough to the optimum w* but far away things may get tricky
- Just as before, we can add a regularization term $\frac{\alpha}{2} ||w||^2$ to avoid invertibility problems
- The iterations in Logistic Regression are again typical of many of the model building methods used in Machine Learning

Recap: Learning in ML

- The general approach to learning is the following:
 - A model f(x; w) is chosen
 - Given a sample $S = \{(x^1, y^1), \dots, (x^N, y^N)\}$, we define a **sample** dependent loss function

$$L(w) = L(w|S) = L(y^1, ..., y^N, f(x^1; w), ..., f(x^N; w))$$

L(w) is often minimized from some w⁰ by iterations

$$w^{k+1} = w^k - \rho_k G(w^k, S)$$

with ρ_k a **learning rate** and G some vectorial function

- When $G(w) = \nabla L(w)$ we have gradient descent
- When $G(w) = \mathcal{H}(w)^{-1} \nabla L(w)$ we obtain **Newton's method**
- When the entire sample S is used at each iteration, we speak of batch learning
- When only single patterns (x^{ρ}, y^{ρ}) or small subsamples are used, we speak of **on–line** or **minibatch learning**

By the Way ...

- We have just built our second neural network: the logistic regression architecture can be seen as a multilayer perceptron (MLP) with no hidden layers and sigmoid outputs
- Again, thinking of this as an MLP seems far-fetched, as we are missing the hidden layers
- Now the gradient computations are more involved
- But we also have learning as iterative loss minimization, i.e., training
- In fact, we will see that classification MLPs for two class problems perform logistic regression on the last hidden layer outputs
- And, again, the process from the input to the last hidden layer produces an enhanced feature representation upon which the regression error should be better

What's New from Regression?

- Some things change from regression, some don't
- We should check feature correlations: although less clearly influential, they will affect most models
 - Scatter plots (x_i, y) are usually less informative
- Important: positive and negative-class feature histograms
- The bias-variance trade-off is subtler in classification
- Accuracy, recall, precision are the usual model quality measures
 - But NN training does not directly minimize them
- We use CV with stratified folds to estimate generalization performance
- We also use CV for hyperparameter estimation, as regularization will also be needed
 - In LogR we should minimize $-\ell(w_0, w; S) + \frac{\alpha}{2} ||w||^2$

How to Handle Posterior Probabilities

- If possible, we don't want labels as model outputs but posterior probabilities
- Most models give them as pairs

$$(\widehat{P}(0|x), \widehat{P}(1|x)) = (\widehat{P}(0|x), 1 - \widehat{P}(0|x))$$

- In principle we would decide 1 if $\hat{P}(1|x) > 0.5$ and viceversa, but this may be too crude
- It may be advisable to set a decision threshold 0 < θ < 1 and decide 1 if P(1|x) > 1 θ and 0 if P(1|x) < θ
- For **imbalanced** problems where $\pi_0 \gg \pi_1$ (usually the interesting ones) we would have $\widehat{P}(1|x) \simeq 0$ for most x
 - In this case we may choose another $\theta \ll 0.5$ and suggest 1 if $\widehat{P}(1|x) > \theta$

Takeaways on Classification

- We have introduced the classification problem as one of computing posterior probabilities
- We have found the optimal Bayes classifier and approximated it by k-NN
- We have introduced Logistic Regression and the numerical minimization of its (minus) log-likelihood
- We have introduced several measures of classifier performance
- 5 We have reviewed some practical issues of classification

Rosenblatt's Perceptrons

Machine Learning Basics

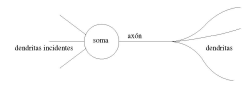
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Basic Neural Models

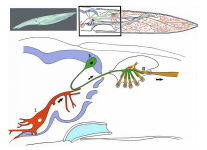
Basic model: Ramón y Cajal's neuron (1900)



- Basic behavior: the neuron either fires or stays at rest depending basically on its inputs
- This translates into an electrical nerve impulse or action potential
- The brain has about 10¹¹ neurons
 - Each one has about 7.000 connections
 - These connections are often recurrent

Hodgkin-Huxley

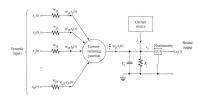
- They developed (circa 1935) the first electro-physiological model to describe the generation and propagation of action potentials in neurons
- They used the giant squid axon for this



From Wikipedia's Squid Giant Synapse

Electronic Neuron

Electronic version: McCulloch–Pitts (1940)



• Since $I = \frac{V}{R} = wV$, with w the conductance, the McCulloch–Pitts neuron output is

$$H\left(\sum_{j=1}^d w_j V_j + I\right) = V$$

with the Heaviside function H ensuring a 0-1 output

 The perceptron is a mathematical formulation of the McC–P neuron where the conductances are to be learned

Rosenblatt's Perceptron

• Given a sample $S = \{(x^p, y^p)\}$ with $y^p = \pm 1$, Rosenblatt's **Perceptrons** (PCPs) are linear machines $w \cdot x$ such that

$$w \cdot x^{\rho} > 0$$
 if $y^{\rho} = 1$
 $w \cdot x^{\rho} < 0$ if $y^{\rho} = -1$

- This can be seen as a homogeneous classification problem
- In more compact form we want for all p

$$y^p \quad w \cdot x^p > 0$$

- If such a separating w exists, it can be computed in many ways
- But Rosenblatt's goal was to **learn** w, i.e., to arrive at a separating w by repeatedly examining the (x^p, y^p) and adjusting w if necessary

Rosenblatt's Delta Rule

• The **Delta Rule** algorithm

```
Start with w=0 While stopping condition not met:

Get a new pattern (x^{\rho},y^{\rho})=\left(x^{\rho(t)},y^{\rho(t)}\right)
if y^{\rho}w\cdot x^{\rho}\leq 0 then:
w=w+y^{\rho}x^{\rho}
```

 Writing w for the new vector and w' for the previous one, notice that

$$y^p \ w \cdot x^p = y^p \ w' \cdot x^p + \|x^p\|^2 > y^p \ w' \cdot x^p$$

Novikov's Theorem

- It states that if the sample S is linearly separable, the Delta rule yields a separating w in a finite number of steps
 - I.e., it learns!!
- More precisely, if for some separating hyperplane w with $\|w\|=1$ we have $\min_p y^p w \cdot x^p = \gamma > 0$, then a separating w' can be found after at most $T=R^2/\gamma^2$ wrong classifications, where $R=\max_p \|x^p\|$
- Notice that γ is the **margin** of the *w* hyperplane. Thus, if

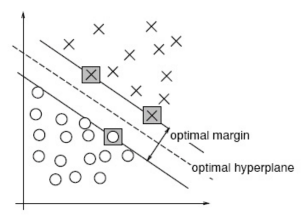
$$\gamma^* = \max_{\|\boldsymbol{w}\|=1} m(\boldsymbol{w}, \boldsymbol{S}) = \max_{\|\boldsymbol{w}\|=1} \min_{\boldsymbol{p}} y^{\boldsymbol{p}} \boldsymbol{w} \cdot \boldsymbol{x}^{\boldsymbol{p}}$$

is the **maximum margin**, we will need at most $R^2/(\gamma^*)^2$ iterations to train a Perceptron

- The maximum margin hyperplane gives the best (smallest) Novikov bound
- And problems with smaller margins require more iterations

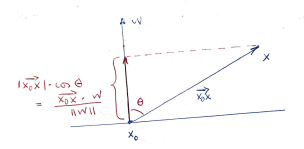
Margins and Generalization

Intuitively, when the margin is large, the problem is easier



Distance to a Line

• The margin of w is just the smallest distance between sample points and the hyperplane defined by w



- This extends to the multidimensional case: if ||w|| = 1 and b = 0, the distance of x to the hyperplane defined by w is $|w \cdot x| = yw \cdot x$
- Thus the Delta Rule corrects weight vectors with negative margins

Novikov's Proof Sketch

- Assume x^t is the t-th pattern such that $y^t w^{t-1} \cdot x^t \le 0$
- Then $||w^t||^2 = ||y^t x^t + w^{t-1}||^2 \le ||x^t||^2 + ||w^{t-1}||^2$ and, thus,

$$||w^t||^2 \le ||x^t||^2 + \ldots + ||x^1||^2 + ||w^0||^2 \le tR^2$$

if $w^0 = 0$ and $||x^p|| \le R$ for all p

 Also, if w' is a 1-norm separating hyperplane i.e., min_p y^pw' · x^p = γ > 0 for all p, we have

$$\mathbf{w}^t \cdot \mathbf{w}' = (\mathbf{y}^t \mathbf{x}^t + \mathbf{w}^{t-1}) \cdot \mathbf{w}' \ge \gamma + \mathbf{w}^{t-1} \cdot \mathbf{w}' \ge \ldots \ge t\gamma + \mathbf{w}^0 \cdot \mathbf{w}'$$

Putting both together we have for all t for which x^t is erroneous

$$t\gamma \le \mathbf{w}^t \cdot \mathbf{w}' \le \|\mathbf{w}^t\| \le \sqrt{t}\mathbf{R}$$

which implies $t \leq R^2/\gamma^2$

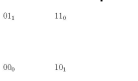
• Thus, the Delta Rule will stop after at most R^2/γ^2 iterations

Problems with Perceptrons

- Rosenblatt's Perceptron caused a big stir on the feasibility of Conexionist Artificial Intelligence but soon its first winter came
- Observation 1: A PCP can learn (obviously) only linearly separable problems; for instance, it can learn the AND and OR predicates



But a PCP cannot learn the XOR predicate



Dichotomies

- Observation 2: for PCPs to be useful, linearly separable problems must be frequent
- Q: how frequent are they?
- A **dichotomy** for $S = \{x^1, x^2, \dots, x^N\}$ is any separation of S in two classes
- It corresponds to any asignment of the ± 1 values to the x^p
- The total number of dichotomies is 2^N
- Thus, the total number of 2-class problems over S is 2^N

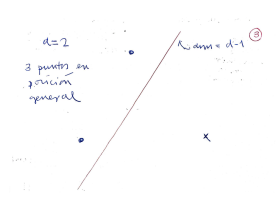
Cover's Theorem

- We have to compare the total number of dichotomies with the number of dichotomies that are linearly separable
- The points in a sample S are said to be in **general position** if there are not d + 1 patterns in a d 1-dimensional hyperplane
- Theorem If S is in general position, the number L (N, d) of linearly separable dichotomies is

$$L(N,d) = \left\{ \begin{array}{cc} 2^N & \text{si } N \leq d+1 \\ 2\sum_{i=0}^d \binom{N-1}{i} & \text{si } N \geq d+1 \end{array} \right\}$$

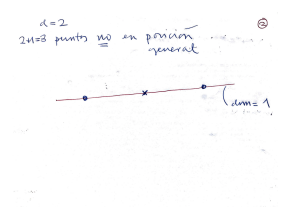
Points in General Position

 Consider d = 2, 3 = d + 1 points and a 1 = d - 1-dimensional hyperplane



Points Not in General Position

• Consider now d = 2 and 3 = d + 1 points **not** on a 1 = d - 1-dimensional hyperplane (i.e., a line)



Thus ...

Notice that for d fixed,

$$\frac{L(N,d)}{2^N} \to 0$$

as $N \to \infty$

- In practice we can expect N ≫ d and the fraction of separable dichotomies will be very small
- Thus, linear PCPs will not be useful
- But a one hidden layer PCP can solve XOR and, in fact, separate any convex region from the rest of the space
- And a two hidden layer PCP can separate any polyhedral region and hence solve any classification problem
- The same is essentially true for regression problems
- But: no algorithm is known to learn these 2-hidden layer PCPs

Why Not Try to Minimize Some Loss?

- Rosenblatt's Perceptrons are natural online machines:
 - Patterns appear sequentially and each may change the current weights
- We can also define a perceptron loss over a given sample as

$$L(w|S) = -\sum_{\{p: y^p w \cdot x^p < 0\}} y^p w \cdot x^p$$

- The local error would thus be $-y^{\rho}w \cdot x^{\rho}$ when $y^{\rho}w \cdot x^{\rho} < 0$ and 0 otherwise
- Since $\nabla_w(-y^\rho \ w \cdot x^\rho) = -y^\rho \ x^\rho$, the Delta Rule can be seen as a kind of local gradient descent on the loss L(w|S):

$$w^{t} = w^{t-1} + y^{p(t)}x^{p(t)} = w^{t-1} - \nabla_{w}\left(-y^{p(t)}x^{p(t)} \cdot w\right)$$

• However L(w|S) is not differentiable and it is not clear what to make of $\nabla_w L(w|S)$...

What Can We Do?

- First option: work with linear models but somehow ensure that d >> N
 - Transform the original d-dimensional features x in new D-dimensional ones such that D >> d
 - We overcome then that, in practice, we usually have $N\gg d$
 - This is the goal of kernel-based Support Vector Machines
- Second option: build trainable PCPs replacing the Heaviside function
 - Work with differentiable PCP transfer function
 - Transform PCP training into a differentiable optimization problem
 - This is what we will do next
- Before doing so, note that Rosenblatt's PCPs can be seen as the first Machine Learning model
- And modern ML started around 1985 along the PCP's ideas

Classical MLPs

1 Machine Learning Basics

Basic Modeling
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Basic Classification
Logistic Regression

2 Multilayer Perceptrons

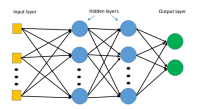
Rosenblatt's Perceptrons

Classical MLPs

Neural Network Classification MLP Regularization Unconstrained Smooth Optimization Computational Costs of MLPs

MLP Architecture

- General layout:
 - An input layer (input)
 - One or several hidden layers
 - One output layer
- Feedforward connections only



• Example: TensorFlow Playground

MLP Connections

- No feedback or lateral conections
- Fully connected layers
- Linear weight connections between units and (usually) non linear activations inside each unit
 - Combined effect of sucesive layers: potentially highly non-linear transformation
- General processing: layered and feedforward
- In practice (1990s), one hidden layer and only sometimes two
- Later (around 2010): Deep Networks with "many" (from 3 to 10) layers
- Now: almost anything!

Unit Activation and Output

• The **activations** a_i^h of a unit in layer h receives the **outputs** o_j^{h-1} from processing in the previous layer

$$a_i^h = \sum_{i=1}^{n_{h-1}} w_{ij}^h o_j^{h-1} + b_i^h,$$

In matrix/vector form:

$$a^h = W^h o^{h-1} + b^h$$

- Output o_i^h of a unit: non linear processing $o_i^h = \varphi(a_i^h)$ of its activation
- In vector form:

$$o^h = \varphi(a^h),$$

where φ is applied over each unit

Activation Functions

- Choices for f:
 - Heaviside (in the very first Rosenblatt's Perceptrons): φ(a) = 0 if a ≤ 0, φ(a) = 1 if a > 0
 - Identity/linear: $\varphi(a) = a$
 - Sigmoid:

$$\varphi(a) = \sigma(a) = \frac{1}{1 + e^{-a}}$$

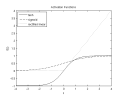
• Hyperbolic tangent:

$$\varphi(a) = \tanh(a) = \frac{e^a - e^{-a}}{e^a + e^{-a}}$$

• Rectified Linear Units (ReLUs): $\varphi(a) = r(a) = \max(0, a)$

Sigmoid, Hyperbolic Tangent, ReLUs

- Sigmoid and tanh: smooth version of Heaviside step function
- For ReLUs r'(x) is either 0 or 1 (hoping x = 0 never happens!!)
 - Many gradient elements will go to 0 and many units will have constant activations no matter their input



From Stanford's UFLDL Tutorial

- Usual activation choices:
 - For hidden units: hyperbolic tangent and, currently, ReLU
 - Linear outputs for regression, sigmoid/soft max outputs for classification

The Forward Pass

- We need first to compute an MLP's outputs through the forward pass
- At each layer h we compute two variable sets
 - The linear **activations** $a^h = W^h o^{h-1} + b^h$ with o^{h-1} the previous layer's output and W^h , b^h the **weights** and **biases** connecting layer h-1 with layer h
 - The nonlinear **outputs** $o^h = \varphi(a^h)$,
- In unit–wise terms: $a_i^h = \sum w_{ii}^h o_i^{h-1} + b_i^h$, $o_i^h = \varphi(a_i^h)$
- Concatenating these operations we arrive at the outputs $\hat{y} = f(x; \mathcal{W})$ of the network, with $\mathcal{W} = (W^1, b^1, \dots, W^H, b^H)$ the network's weight set

The Simplest MLP I

- The Single Hidden Layer (SHL) MLP
 - *D* inputs (determined by the problem at hand)
 - One hidden layer with H units (number to be chosen) and tanh (or what you prefer) activation
 - One linear or sigmoid output (according to the problem at hand)
- Input-hidden processing: denoting inputs by x and the hidden unit ouput as o^H,

$$o_h^H = anh \left(b_h^H + \sum_{j=1}^D W_{hj}^H x_j
ight)$$

• In matrix/vector form: $o^H = \tanh (W^H x + b^H)$

The Simplest MLP II

• Hidden–output processing: since targets are 1–dimensional, we have for the outputs \hat{y}

$$\widehat{y} = \sum_{h=0}^{H} w_h^O o_h^H + b^O,$$

- In vector form: $\hat{y} = w^O \cdot o^H + b^O$
- That is, a linear model on the last hidden layer outputs o^H
- Global process:

$$\widehat{y} = f(x; w^O, W^H, b^O, b^H) = b^O + \sum_h w_h^O \tanh \left(b_h^H + \sum_j W_{hj}^H x_j \right)$$

Or in matrix/vector form

$$\widehat{y} = f(x; w^O, W^H, b^O, b^H) = b^O + w^O \cdot tanh (b^H + W^H x)$$

Or in Python y_pred = np.tanh(x.dot(w_H.T)+ b_H).dot(w_O.T)+ b_O

MLP Error Function

MSE is the standard error function for regression MLPs

$$e(\mathcal{W}) = \frac{1}{2} E_{x,y} \left[(y - f(x; \mathcal{W}))^2 \right] = E_{x,y} \left[e^{\ell}(x, y; \mathcal{W}) \right]$$
$$= \int e^{\ell}(x, y; \mathcal{W}) p(x, y) dx dy$$

with $e^{\ell}(x, y; \mathcal{W})$ denotes the **local error**

$$e^{\ell}(x, y; \mathcal{W}) = \frac{1}{2}(y - \widehat{y})^2 = \frac{1}{2}(y - f(x; \mathcal{W}))^2$$

Minimizing the MSE

 The general idea would be to obtain the optimal W* as a solution of ∇e(W) = 0, where

$$\begin{array}{lcl} \nabla e(\mathcal{W}) & = & E_{x,y} \left[\nabla_{\mathcal{W}} e^{\ell}(x,y;\mathcal{W}) \right] \\ & = & E_{x,y} \left[\nabla_{\mathcal{W}} f(x;\mathcal{W}) (f(x;\mathcal{W}) - y) \right] \end{array}$$

for we have

$$\nabla_{\mathcal{W}} e^{\ell}(x, y; \mathcal{W}) = -(y - f(x; \mathcal{W})) \nabla_{\mathcal{W}} f(x; \mathcal{W})$$
$$= \nabla_{\mathcal{W}} f(x; \mathcal{W}) (f(x; \mathcal{W}) - y)$$

- This suggests to get ∇_W f(x; W) but it is much better to focus on ∇e^ℓ, and
 - We will compute first ∇e^{ℓ} (and then $\nabla e = E[\nabla e^{\ell}]$)
 - We will exploit it to build MLPs through optimization methods

The Forward Pass (again)

- Recall that we compute an MLP's outputs through the forward pass
- At each layer h we compute (and **store**) first the linear **activations** $a^h = w^h o^{h-1} + b^h$ with o^{h-1} the previous layer's output and W^h , b^h the **weights** and **biases** connecting layer h-1 with layer h
- We the compute (and store) the nonlinear **outputs** $o^h = \varphi(a^h)$,
- In unit—wise terms:

$$a_i^h = \sum w_{ij}^h o_j^{h-1} + b_i^h, \quad o_i^h = \varphi(a_i^h)$$

And we also have

$$\frac{\partial a_i^h}{\partial w_{ii}^h} = o_i^{h-1}, \ \frac{\partial o_i^h}{\partial a_i^h} = \varphi'(a_i^h)$$

Computing the Gradient

- The key tool is the chain rule that we will apply backwards from the output layer
- If w_{ij}^h is the general weight connecting unit j in layer h-1 to unit i in layer h, we have

$$\frac{\partial e^{\ell}}{\partial w_{ij}^{h}} = \frac{\partial e^{\ell}}{\partial a_{i}^{h}} \frac{\partial a_{i}^{h}}{\partial w_{ij}^{h}} = \frac{\partial e^{\ell}}{\partial a_{i}^{h}} o_{j}^{h-1} = \delta_{i}^{h} o_{j}^{h-1}$$

- That is, we have an easy to compute term $\frac{\partial a_i^n}{\partial w_{ij}^n}$, and a more complicated but crucial one $\frac{\partial e^{\ell}}{\partial a^h}$
- We refer to $\frac{\partial e^{\ell}}{\partial a_i^h} = \delta_i^h$ as the **generalized error** at unit *i* of layer *h*

Gradient at the Output Layer

- Let's assume that we have layers 1, 2, ..., H 1, H, with H the output layer
- For simplicity, consider a regression problem with linear outputs
- In the output H layer we have

$$e^{\ell} = \frac{1}{2}(y - \hat{y})^2$$
 and $\hat{y} = o^H = a^H = \sum_j w_j^H o^{H-1}$

Thus, the partial derivative is straightforward

$$\delta^{H} = \frac{\partial \mathbf{e}^{\ell}}{\partial \mathbf{a}^{H}} = \frac{\partial \mathbf{e}^{\ell}}{\partial \widehat{\mathbf{y}}} = \widehat{\mathbf{y}} - \mathbf{y}$$

i.e., the generalized error δ^H is here the **standard error**

• Therefore, we have for the output layer H

$$\frac{\partial e^{\ell}}{\partial w_i^H} = (\widehat{y} - y) \frac{\partial a^H}{\partial w_i^H} = (\widehat{y} - y) o_j^{H-1}$$

Now it is (quite) easy to extend this to other layers

Backprop for Other Layers

• Let w_{ij}^h be the weight connecting unit j in layer h-1 with unit i in layer h; recall that then

$$\frac{\partial \boldsymbol{e}^{\ell}}{\partial \boldsymbol{w}_{ij}^{h}} = \frac{\partial \boldsymbol{e}^{\ell}}{\partial \boldsymbol{a}_{i}^{h}} \frac{\partial \boldsymbol{a}_{i}^{h}}{\partial \boldsymbol{w}_{ij}^{h}} = \frac{\partial \boldsymbol{e}^{\ell}}{\partial \boldsymbol{a}_{i}^{h}} \boldsymbol{o}_{j}^{h-1}$$

• And if we have already computed the generalized errors $\delta_k^{h+1} = \frac{\partial e^{\ell}}{\partial a^{h+1}}$ for layer h+1, we have

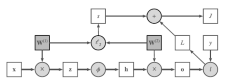
$$\frac{\partial e^{\ell}}{\partial a_{i}^{h}} = \sum_{k} \frac{\partial e^{\ell}}{\partial a_{k}^{h+1}} \frac{\partial a_{k}^{h+1}}{\partial a_{i}^{h}} = \sum_{k} \delta_{k}^{h+1} \frac{\partial a_{k}^{h+1}}{\partial a_{i}^{h}} \\
= \sum_{k} \delta_{k}^{h+1} \frac{\partial a_{k}^{h+1}}{\partial o_{i}^{h}} \frac{\partial o_{i}^{h}}{\partial a_{i}^{h}} = \left[\sum_{k} \delta_{k}^{h+1} w_{ki}^{h+1} \right] \varphi'(a_{i}^{h})$$

Backprop and Deep Networks

- The previous formulae work for any number of hidden layers
 - We can work with deep MLPs
- Moreover, the chain rule can be automated and modern NN packages compute gradients automatically for very general NN architectures
 - Basic tool: automatic symbolic differentiation on computational graphs
 - This is one of the key reasons of the great success of deep neural nets
- Some examples: convolutional nets, residual nets, recurrent NNs, . . .

Automating Backprop

- To deal with such general networks requires automating backprop
- This is done upon the network's computational graph



From Zhang et al., Dive into Deep Learning

- It corresponds to a SHL with a loss J = L + s
- The circles contain differentiable operators
- The shadowed squares contain the variables against which differentiation takes place

Neural Network Classification

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Logistic Regression (again)

- Recall that our Logistic Regression model gave us estimate of P(0|x; w₀, w), P(1|x; w_o, w)
- We then wrote the likelihood of a given sample as

$$P(Y|X; w_0, w) = \prod_{p=1}^{N} P(y^p|x^p; w_0, w)$$

• And then derived the optimum parameters w_0^* , w^* as

$$w_0^*, w^* = \arg \max_{w_0, w} P(Y|X; w_0, w)$$

 Notice that we can see the output of a logistic regression model as that of an NN with no hidden layers and sigmoid outputs

Neural 2-class Classification

 We can follow the same path for a 2 class classification problem changing that output for the one

$$f(x; \mathcal{W}) = w_0 + w \cdot \Phi(x, \mathcal{W}^H)$$

with w_0 , w the weights from the last hidden layer (LHL) to the output and $\Phi(x, \mathcal{W}^H)$ the map from inputs to the LHL outputs

• And now derive the optimum parameters \mathcal{W}^* as

$$\mathcal{W}^* = \operatorname{arg\,min}_{\mathcal{W}} - \sum_{p=1}^{N} \log P(y^p | x^p; \mathcal{W})$$

$$= \underset{w_0, w, \mathcal{W}^H}{\operatorname{arg\,min}} - \sum_{p} y^p \left(w_0 + w \cdot \Phi(x^p, \mathcal{W}^H) \right) + \sum_{p} \log(1 + e^{w_0 + w \cdot \Phi(x^p, \mathcal{W}^H)})$$

- Forward and backpropagation are also straightforward here
- We change things a little bit for K class problems

MLPs for K-class Classification

- We consider an input layer and a number of hidden layers
- Targets are the 1-hot encodings of class labels, so we use K outputs
- We want the MLP's k-th output to estimate the posterior P(k|x)
- The natural output layer activation is thus the softmax function

$$\sigma_j(o) = F_j(x; \mathcal{W}) = \frac{e^{w_0^j + w^j \cdot o}}{\sum_{1}^K e^{w_0^k + w^k \cdot o}}$$

- Here o are the last hidden layer outputs and w^{j} , w_{0}^{j} are the weight and bias from the last hidden layer to the network's j output
- Moreover $\sum_{i} F_{j}(x; \mathcal{W}) = 1$ and we assume $P(j|x) \simeq F_{j}(x; \mathcal{W})$
- For two classes this becomes

$$\sigma_1(o) = \frac{e^{w_0^1 + w^1 \cdot o}}{e^{w_0^0 + w^0 \cdot o} + e^{w_0^1 + w^1 \cdot o}} = \frac{1}{1 + e^{w_0^0 - w_1^1 + (w^0 - w^1) \cdot o}} = \frac{1}{1 + e^{\omega_0 + \omega \cdot o}}$$

We thus get the sigmoid activation of Logistic Regression

Multiclass Log-Loss NN Classification

- We change slightly the preceding for a K class classification problem
- We define an MLP $F(x; W) \in \mathbf{R}^K$ with softmax outputs
 - Recall that $0 \le F_k(x; \mathcal{W}) \le 1$ and $\sum_k F_k(x; \mathcal{W}) = 1$
 - And also we assume that $P(k|x) \simeq \hat{F}_k(x; \mathcal{W})$
- We label now the multiclass targets via the 1–hot encoding, with the target of class k being the vector $e_k = (0, ..., 1, ..., 0)$
- Then, if x ∈ C_k, its label is y = (y₁,..., y_K)^t is e_k, with y_k = 1, and

$$P(k|x) = P(k|x)^{1} = P(k|x)^{y_{k}} = \prod_{c=1}^{K} P(c|x)^{y_{c}} \simeq \prod_{c=1}^{K} F_{c}(x; \mathcal{W})^{y_{c}}$$

The Sample Likelihood

- We have a sample $S = \{x^p, y^p\}$ with (x^p, y^p) in class c(p) and, hence, $y^p = e_{c(p)}$
- We also consider a posterior model $P(c|x, w) = F_c(x, \mathcal{W})$, the probability of getting S = (X, Y) is

$$P(Y|X; \mathcal{W}) = \prod_{1}^{N} P(c(p)|x^{p}; \mathcal{W}) = \prod_{1}^{N} \prod_{c=1}^{K} P(c|x^{p}; \mathcal{W})^{y_{c}^{p}}$$

$$\simeq \prod_{1}^{N} \prod_{c=1}^{K} F_{c}(x; \mathcal{W})^{y_{c}^{p}}$$

• Thus, we estimate W by maximizing the sample's **likelihood**

$$\mathcal{P}(Y|X;\mathcal{W}) = \prod_{p=1}^{N} \prod_{c=1}^{K} F_{c}(x^{p};\mathcal{W})^{y_{c}^{p}}$$

The Cross Entropy Loss

As before, we will work with the sample's log-likelihood, i.e.

$$\ell(\mathcal{W}; S) = \log \mathcal{P}(Y|X; \mathcal{W}) = \sum_{p=1}^{N} \sum_{c=1}^{K} \log \left(F_c(x^p; \mathcal{W})^{y_c^p} \right)$$
$$= \sum_{p=1}^{N} \sum_{c=1}^{K} y_c^p \log F_c(x^p; \mathcal{W})$$

• The **cross–entropy** loss is now simply the negative log–likelihood $-\ell(W; S)$, i.e.,

$$L(\mathcal{W}) = -\ell(\mathcal{W}; S) = -\sum_{p=1}^{N} \sum_{c=1}^{K} y_c^p \log F_c(x^p; \mathcal{W})$$

It is now straightforward to carry this into a NN setting

Takeaways on Clasical MLPs

- 1 They have a layered structure with outputs computed in a forward pass using differentiable activations
- 2 Usual activations: ReLUs, sigmoid, tanh, linear
- MSE is the usual regression cost; we use a variant of cross entropy in classification
- The error function gradients are computed by backpropagation of generalized errors
- 6 Backprop is basically a very simple procedure than can be largely automated
- 6 Regression and classification problems fit very well in this setting

MLP Regularization

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MLPs and Universal Approximation

- In regression we assume $y = \phi(x) + n$ and our goal is to get $f \simeq \phi$
- We say that $\mathcal{F} = \{f(x; \mathcal{W})\}$ is a **Universal Approximation** Family over a domain \mathcal{R} if

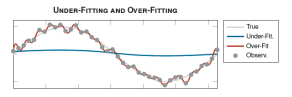
For any $\epsilon > 0$ and any reasonable ϕ , we can find an $f(x; \mathcal{W}_{\phi,\epsilon})$ s.t.

$$e(\mathcal{W}_{\phi,\epsilon}) = \int (\phi(x) - f(x; \mathcal{W}_{\phi,\epsilon}))^2 p(x) dx \le \epsilon$$

- Notice that Universal Approximation is just what we need in regression
- In fact a Single Hidden Layer MLP with enough hidden units is an effective universal approximator
- But it also implies the risk of overfit whose control is crucial

Overfitting in MLPs

- Since MLPs are a UAF, they can also approximate the noise in the sample
 - Given S = {(x^ρ, y^ρ)} if we allow enough hidden units in a SHL MLP we can arrive to a w* s.t. y^ρ = f(x^ρ; w*)
 - We get thus a sample error $\hat{e}(w^*) = 0$ but possibly with a very high generalization error
- I.e., MLPs may have very small bias but possibly large variance



(Ph.D. Thesis of Carlos Alaíz)

Regularization vs Overfitting

- Why is there overfitting?
 - Because we may end up having too many weights with respect to sample size
 - Because we allow these weights to explore the entire weight space
- We can avoid this wandering if we limit w's growth, for which we add a **regularization** term g(||w||) to e(w) with g(r) increasing
- Working with $e_R(w) = e(w) + g(\|w\|)$ we have to **balance** the minimization of e(w) and that of $g(\|w\|)$
- This balanced learning results in better generalization

L₂ Regularization

 The simplest regularization procedure adds a quadratic penalty to the square error e

$$e_R(w) = e(w) + \frac{\lambda}{2} ||w||^2,$$

with λ the **weight decay** factor

- Also known as Tikhonov's regularization (or Ridge Regression for linear models)
- The desired effect is to constrain the evolution of w:
 - In fact, the regularized loss is essentially the Lagrangian of the constrained problem

$$\min_{w} e(w)$$
 subject to $||w||^2 \le \rho$, $\rho > 0$

• The gradient becomes $\nabla e_R(w) = \nabla e(w) + \lambda w$ and gradient descent becomes

$$\mathbf{w}^{k+1} = \mathbf{w}^k - \rho_k(\nabla_{\mathbf{w}}\mathbf{e}(\mathbf{w}^k) + \lambda\mathbf{w}^k)$$

How to choose λ

- Again, the correct choice of λ is crucial
- A small $\lambda \ll$ 1 results in a small regularization effect and overfitting risk appears
- A large \(\lambda \rightarrow 1 \) causes learning to forget about \(e(w) \) and the model will be essentially constant and will underfit
- Usually λ is chosen by using CV by exploring a discrete set of values λ_i,
- The same can essentially done for any other hyper–parameter:
 - Number of hidden layers and of hidden units?
 - Learning rate??
 - Minibatch size???
- But this may become unmanageable if we want to work with many (more than three?) hyperparameters

MLP Ensembles

- Recall that for MLPs e(w) does not have a single minimum
- Moreover, the final MLP depends on the random initial w^0
- And mini-batch training adds extra randomness to the final model
- Therefore, you may never get the same MLP twice!!!
- But we can turn this to our advantage by
 - Starting from K independent initial weights and get K optimal weight sets w_k*
 - Giving as the output the average $f_e(x) = \frac{1}{K} \sum_{1}^{K} f(x; w_k^*)$
- We expect outputs of the form $\hat{y}_k^p = y^p + \epsilon_k^p$ with the errors ϵ_k^p being independent
- Hence $\frac{1}{K}\sum_k \epsilon_k^{p} \simeq 0$ and $\frac{1}{K}\sum_k \widehat{y}_k^{p} \simeq y^{p}$

Takeaways on MLP Regularization

- MLPs have a high risk of overfitting
- Thus, they must be regularized to avoid overfitting
 - The regularization hyperparameter is chosen through cross validation
- MLP training has two random components: the initial point and minibatch selection
 - Thus, each training will converge to a different optimum
- MLP ensembles can take advantage of this
 - They imply extra costs but ensembles are embarrassingly parallelizable

Unconstrained Smooth Optimization

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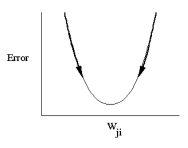
Computational Costs of MLPs

Back to Optimization

- General optimization theory is a key tool in Machine Learning (ML)
- There are two optimization set ups in ML
 - Unconstrained optimization, slightly simpler and the one used for MLPs
 - Constrained optimization, wider and more complex
- In ML we have also to consider the optimization of differentiable and also non differentiable error functions
- MLP optimization: unconstrained and differentiable
- And also batch, i.e., over the entire sample, mini-batch over subsamples or on line, pattern by pattern

Gradient Descent

- We "stretch" the weight set W into a vector w
- We recall that $-\nabla e(w)$ is the maximum descent direction
- First idea: to build a (hopefully convergent) sequence w^k iterating (small) steps along $-\nabla e(w^k)$



Gradient Descent II

In more detail, we start from a random w⁰ and compute

$$w^{k+1} = w^k - \rho_k \nabla_w e(w^k)$$

- ρ_k is the **learning rate** (LR)
- With a small ρ_k we ensure e(w^{k+1}) < e(w^k) (although with possibly a very small descent)
- We can get a better iteration w^{k+1} = w^k − ρ_k*∇_we(w^k) using a ρ_k* given by

$$\rho_k^* = \arg\min_{\rho} e(\mathbf{w}^k - \rho \nabla_{\mathbf{w}} e(\mathbf{w}^k));$$

this is known as line minimization

 These GD methods are called first order methods in part because they only use ∇e

One Dimensional Newton's Method

- Assume a quadratic function q(w) = aw² + bw + c, with a > 0, and a minimum at w*
- We can reach w^* from any w with a step Δw such that

$$0 = q'(w + \Delta w) = 2a(w + \Delta w) + b$$

• We have thus $\Delta w = \frac{-b-2aw}{2a}$, that is,

$$w^* = w - \frac{2aw + b}{2a} = w - \frac{1}{q''(w)}q'(w)$$

 This leads to Newton's method: minimize a general f iteratively using steps

$$w^{k+1} = w^k - \rho_k \frac{1}{f''(w^k)} f'(w^k)$$

with ρ_k a suitable learning rate

One Dimensional Newton's Method II

- Notice how the $\frac{1}{f''(w_k)}$ acts as a self adjusting learning rate
- When the parabola is very sharp (i.e., $a \gg 1$), we will be close to the minimum and the descent can overstep it
- But then $f''(w_k)$ will be big and, hence, $\frac{1}{f''(w_k)}$ moderates the gradient step
- Conversely, when the parabola is very wide (i.e., $a \simeq f''(w_k) \ll 1$), the minimum will be far away and the gradient step small descent will
- But then $f''(w_k)$ will be rather small and, hence, $\frac{1}{f''(w_k)}$ accelerates the gradient step
- We can extend this to several dimensions

Multidimensional Newton's Method

 For a d dimensional w, the Taylor expansion of e at an optimum w* is

$$e(w) \approx e(w^*) + \frac{1}{2}(w - w^*)^t \cdot \mathcal{H}(w^*) \cdot (w - w^*)$$

- $\mathcal{H}(w^*)$ is the **Hessian** of *e* at w^* and $\nabla e(w^*) = 0$
- It follows that $\nabla e(w) \approx \mathcal{H}(w^*) \cdot (w w^*)$ and, therefore,

$$\mathbf{w}^* \approx \mathbf{w} - \mathcal{H}(\mathbf{w}^*)^{-1} \nabla_{\mathbf{w}} \mathbf{e}(\mathbf{w})$$

This suggest to derive the w^k by

$$\mathbf{w}^{k+1} = \mathbf{w}^k - \rho_k \mathcal{H}(\mathbf{w}^k)^{-1} \nabla_{\mathbf{w}} \mathbf{e}(\mathbf{w}^k)$$

which is known as **Newton's Method** (NM)

Variants of Newton's Method

- Theoretically NM converges very fast near w*, but
 - Far from w* convergence is not guaranteed
 - Moreover $\mathcal{H}(w^k)$ may not be invertible
 - Besides, computing $\mathcal{H}(w^k)$ is cumbersome and costly
- The Gauss-Newton (GN) approximation

$$\mathcal{H}(\mathbf{w}) \simeq \mathbf{E}[\nabla \mathbf{e}(\mathbf{w}) \nabla \mathbf{e}(\mathbf{w})^{\tau}],$$

holds for any quadratic cost and simplifies the third problem

- The Levenberg-Marquardt (LM) method deals with the other two problems combining
 - Gradient descent "away" from w*
 - Gauss–Newton "near" w*

although "away" and "near" have to be properly addressed

One-dimensional GN Approximation

• Assume $e(w) = \frac{1}{2} \int (f(x; w) - y)^2 p(x, y) dxdy$; then

$$e'(w) = \int (f(x; w) - y) \frac{\partial f}{\partial w}(x, y) p(x, y) dx dy;$$

$$e''(w) = \int \left(\frac{\partial f}{\partial w}\right)^2 p(x, y) dx dy$$

$$+ \int (f(x; w) - y) \frac{\partial^2 f}{\partial w^2}(x, y) p(x, y) dx dy$$

• Near a minimum w^* we may expect $f(x; w) \simeq y$ and, therefore,

$$e''(w) \simeq \int \left(\frac{\partial f}{\partial w}\right)^2 p(x,y) dxdy > 0$$

 Thus, for square errors, we can use first derivates to approximate e''(w)

General GN Approximation I

In the general case we have

$$\nabla e(w) = E\left[\nabla f(x; w) \left(f(x; w) - y\right)\right]$$

And, therefore,

$$\nabla^2 e(w) = E\left[\nabla^2 f(x; w) \left(f(x; w) - y\right)\right] + E\left[\nabla f(x; w) \nabla f(x; w)^{\tau}\right]$$

- The second term is easy to compute once we have ∇f
- If $w \approx w^*$, $f(x; w) \approx y$; therefore $f(x; w) y \approx 0$,
 - We can ignore the first, more complex, term

General GN Approximation II

• We arrive at $\nabla^2 e(w) \simeq E\left[\nabla f(x;w)\nabla f(x;w)^{\tau}\right]$ or, equivalently,

$$\mathcal{H}_{(i,j)(p,q)}(w) = \left(\frac{\partial^2 e}{\partial w_{ij}\partial w_{pq}}(w)\right) \simeq \left(E\left[\frac{\partial f}{\partial w_{pq}}\frac{\partial f}{\partial w_{ij}}\right]\right)_{(i,j)(p,q)}$$

- $\mathcal{J} = E\left[\nabla f(x; w) \nabla f(x; w)^{\tau}\right]$ is **Fisher's information matrix**
 - J is semidefinite positive
 - And thus not necessarily invertible
- Often only its diagonal is considered and we have

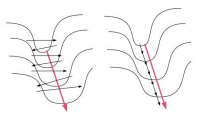
$$\mathcal{H}_{(i,j)(i,j)}(w) \simeq E\left[\left(\frac{\partial f}{\partial w_{ij}}\right)^2\right]$$

Advanced Optimization

- There are many more proposals in unconstrained optimization
- The Conjugate Gradient (CG) and Quasi-Newton (QN) methods are important in classical MLP training
- The basic idea in CG is to replace gradient descent directions g_k = -∇e(w_k) with new conjugate directions that try to keep somehow the previous "good directions"
- The basic idea in QN is to iterate as in NM but with simple approximations A_k to $\mathcal{H}^{-1}(w^k)$ that converge to $\mathcal{H}^{-1}(w^*)$
- When training "small" NNs the Limited-memory Broyden-Fletcher-Goldfarb-Shanno (L-BFGS) QN variant is often used

Accelerating Gradient Descent

- A possibility on the error surface of a NN is to have many long, narrow ravines
 - Gradients bounce in the narrow section, but may be very small along the wider one



Momentum

- Momentum pushes them forward in the wider section (but we have to cope with a new parameter)
- Momentum tries to maintain descent's inertia with a term $\Delta^k = w^k w^{k-1}$, i.e.,

$$\mathbf{w}^{k+1} = \mathbf{w}^k - \rho_k \nabla_{\mathbf{w}} \mathbf{e}(\mathbf{w}^k) + \mu_k \Delta^k$$

- More or less, what a heavy ball would do
- The goal is to keep w^k advancing in "plateaux", i.e., small gradient zones
- Momentum can be seen as a crude approximation of a CG step
- Nice explanation at Why Momentum Really Works

Nesterov's Accelerated Gradient

- Let's rewrite momentum in two steps
 - 1 Define $\Delta^{k+1} = -\rho_k \nabla_w e(w^k) + \mu_k \Delta^k$ and
 - 2 Apply

$$\boldsymbol{w}^{k+1} = \boldsymbol{w}^k + \boldsymbol{\Delta}^{k+1} = \boldsymbol{w}^k - \rho_k \nabla_{\boldsymbol{w}} \boldsymbol{e}(\boldsymbol{w}^k) + \mu_k \boldsymbol{\Delta}^k$$

Nesterov's Accelerated Gradient is a variant of this

$$\widetilde{\Delta}^{k+1} = -\rho_k \nabla_w e(w^k + \mu_k \widetilde{\Delta}^k) + \mu_k \widetilde{\Delta}^k;
 w^{k+1} = w^k + \widetilde{\Delta}^{k+1}$$

 In convex optimization it improves GD and is often highly effective in Deep Network training

Adam

- Adam is currently the most widely used gradient-descent method for deep NN training
- At each step t Adam uses a new random mini-batch to
 - Update exponentially smoothed averages m_t of the gradient g_t and v_t of the the squared gradient g_t² = g_t ⊙ g_t as

$$m_t = \beta_1 m_{t-1} + (1 - \beta_1) g_t, \ v_t = \beta_2 v_{t-1} + (1 - \beta_2) g_t^2;$$

Since we have

$$E[m_t] \simeq (1 - \beta_1^t) E[g_t], \ E[v_t] \simeq (1 - \beta_2^t) E[g_t^2],$$

we compute bias corrections \widehat{m}_t , \widehat{v}_t as

$$\widehat{m}_t = \frac{1}{1 - \beta_1^t} m_t, \ \widehat{v}_t = \frac{1}{1 - \beta_2^t} v_t;$$

• Update weights as $W_t = W_{t-1} - \alpha \frac{\widehat{m}_t}{\sqrt{\widehat{v}_{t+\epsilon}}}$

Understanding Adam

- Default values $\alpha = 0.001, \beta_1 = 0.9, \beta_2 = 0.999$, and $\epsilon = 10^{-8}$ usually work fine
- What is going on in Adam?
- At each step we work with estimates of the average local gradient:

$$\widehat{m}_t \simeq E[\nabla_W f]; \ \widehat{v}_t \simeq \sigma(g_t) \simeq E[(\nabla_W f)^2]$$

Thus, we can first see the Adam iterations

$$W_t = W_{t-1} - \alpha \frac{\widehat{m}_t}{\sqrt{\widehat{v}_t} + \epsilon}$$

as a kind of "normalized" gradient descent

In more detail, we expect

$$g_t^2 = E[(\nabla_W f)^2] \simeq E[\nabla_{W^2}^2 f] = \mathcal{H},$$

we can see Adam as a "dampened" variant of diagonal Gauss-Newton's steps

When to Stop Training

- Typically the $e(W^k)$ error diminishes towards an asymptotic minimum
 - If many units are used, we may arrive to 0 error, which usually implies overfitting
- First solution: to use a separate validation subset V and stop training when the error in V, i.e., the validation error starts growing
 - This early stopping is available in the main NN packages
 - But what do we do for small samples?
- Second solution: get a good regularization
 - Now training stops because of reasons such as computational cost, but not because of overfitting risk
- We must combine both, to save on training time and because overfitting is a serious MLP risk

Takeaways on MLP Optimization

- Gradient descent is the simplest method but possibly also the slowest
- Momentum can be used to speed it up
- Newton's method is much faster but may be very costly and difficult to apply in full form
- All the previous methods require the (usually tricky) selection of a learning rate
- Second order methods such as Conjugate Gradient and Quasi-Newton avoid learning rates and are more efficient but costlier
- Limited Broyden–Fletcher–Goldfarb–Shanno (L-BFGS) is currently the option of choice for "small" MLPs

Big Data

2 Multilayer Perceptrons

Neural Network Classification

Computational Costs of MLPs

MLPs and Big Data?

- Many Vs in Big Data: Volume, Velocity, Variety, Veracity, Value, ...
- Velocity: information flows in data streams that require fast processing and feed back
 - MLPs are rather fast
 - Less than linear models but much more so than SVMs
- Volume is probably the greatest attractive of Big Data
 - Huge samples and/or very large pattern dimension
 - Large impact in model training

Volume in MLPs

- Parallelism is the first answer to Big Volume
- On a standalone machine it may be
 - Passive: let the SO distribute work among several CPU cores, or use GPUs with low level parallelized libraries such as Linpack or BLAS for linear algebra
 - Active: explicitly exploit a problem's parallelism programming an algorithm in, say, OpenMP
- It is easy to passively parallelize the training of several MLPs
 - But memory costs may multiply
- Training an isolated MLP is not CPU parallelizable:
 - There is a sequential layer dependence in Backprop
 - Each training iteration depends on the previous one
 - But matrix-vector multiplications can exploit GPU parallelism

Handling Huge Sample Training

- Two consequences of a large N are
 - Sample doesn't fit in memory and we have to split it somehow
 - Training gets "lost", for in the global gradient $\nabla e = E[\nabla e^{\ell}]$ we average many local gradients that may cancel each other out
- The fist problem has been always present in fields such as analog signal filtering
- Solution for both: adaptive or on line, i.e., pattern by pattern, weight updates
 - It also allows a relatively simple setting for a theoretical analysis
 - But not used today; instead, medium-to-large NNs are trained using mini-batches

MLP Complexity

- MLPs are fast to apply but costly to train
- How can we estimate this cost?
- MLP traning cost is determined by
 - Its architecture, that determines the number of weights to fit and that is also dependent on the input dimension D
 - The full or mini-batch sample size N, that determines the cost of the averages to be computed
 - The training method, with more or less iterations that, in turn, are more or less costly

Forward Pass Complexity

 The number of weights in a single hidden layer (SHL) MLP with D inputs, one output and one hidden layer with H units is

$$(D+1)\times H+(H+1)\simeq HD$$

- Each extra hidden layer pair with (H_1, H_2) units adds $(H_1 + 1) \times H_2 \simeq H_1 H_2$ weights
- The cost per pattern of the feedforward step between layers
 h 1, h is basically that of H_{h-1} × H_h floating point operations
 (FPOs)
- Thus, for a general MLP the cost in floating point operations of a forward pass is ≃ N × (∑_h H_h × H_{h-1})
 - Very fast on GPUs as it theoretically becomes $O(N \times \sum_h H_h)$

The Cost of Computing ∇e^{ℓ}

- Computing a local gradient ∇e^ℓ in a SHL MLP with square error and one output essentially requires to compute
 - *H* components for the hidden to output connections, with a O(1) cost each, for $\frac{\partial e^{\ell}}{\partial w^{\mathcal{Q}}} = (\widehat{y} y)o_h$
 - *DH* components for the input to hidden connections with essentially an O(1) cost each, for $\frac{\partial e^{\ell}}{\partial w_{h}^{H}} = \left(\sum_{i=1}^{L} \delta_{i} w_{ih}^{O}\right) \sigma'(a_{h}) x_{j}$
- In these we are omitting the cost of the partials with respect the bias, but they are much smaller
- Thus, the overall cost of computing ∇e^{ℓ} on a SHL MLP is essentially O(DH)
- More layers with H_h units add a cost $O(H_{h-1} \times H_h)$
 - $O(\sum_h H_h)$ on GPUs

The Cost of Computing ∇e

- For a mini-batch of size N_b, the cost of the mini-batch gradient ∇e of a SHL MLP is O(N_b × D × H)
- And extra layers add a cost $O(N_b \times H_{h-1} \times H_h)$
- This is of the same order of magnitude than the cost of the forward pass
- This also dominates the $O(H_{h-1} \times H_h)$ cost of updating the (H_{h-1}, H_h) weights in gradient descent
- And all these costs have to be multiplied by the number of training epochs

Training Complexity

- The important term in the overall training cost is nEps × cost of ∇e, with nEps the number of epochs
 - One epoch = one pass on the entire sample
- Thus, the globally dominant term in a SHL MLP is

nEps × cost of
$$\nabla e = O(\text{nEps} \times N \times D \times H)$$

with N here the entire sample size

And for each hidden layer we add an extra cost

$$O(nEps \times N \times H_{h-1} \times H_h)$$

- Thus, training many layered, large MLPs can be very costly
 - But linear in sample size N

Takeaways on MLP's Cost

- MLP complexity is determined by its architecture {H_h}, training procedure and sample size N
- The forward and backward MLP passes have basically the same complexity
- Their cost per pattern and layer is $H_{h-1} \times H_h$
- For gradient descent these costs are multiplied by the number nEps of epochs
- First order methods essentially do not add extra complexity
- Second order methods add extra per iteration costs but should require less iterations
- Single MLP training is not CPU parallelizable
- But GPUs can greatly improve MLP processing costs