Artificial Neural Networks (Gerstner). Exercises for week 6 Regularization and Tricks of the Trade

Exercise 1. Dropout (from exam 2019)

We have a deep network of 2n hidden layers (n > 2) of neurons with sharp threshold functions g(a) = 1 for a > 0 and zero otherwise. After training with dropout, somewhere in hidden layer n, we have a hidden neuron i which receives input from 4 hidden neurons in layer n - 1. All weights onto neuron i are equal to one and the threshold of neuron i is 2.7.

Each of the four hidden neurons j in layer n-1 receives input from the same 2 neurons in layer n-2. The weight vectors and thresholds of the four neurons in layer n-1 are:

- j=1 (1,0) and threshold 0
- j=2 (1,0) and threshold 0.5
- j=3 (1,1) and threshold 1
- j=4 (1,-1)and threshold 1
 - a. Qualitatively sketch the two-dimensional space representing the activity of the 2 neurons in layer n-2 and indicate the region (by shading it with crosses x x x) in which neuron i responds positively.
 - b. Dropout: Remove neurons j = 1 and j = 4 in layer n 1, rescale the weights appropriately, and sketch the input space where neuron i responds positively (by shading it with crosses $x \times x$).
 - c. Dropout: Remove neurons j=2 and j=3 in layer n-1, rescale the weights appropriately, and sketch the input space where neuron i responds positively (by shading it with crosses $x \times x$).
 - d. Your friend Adam claims: 'Dropout might be a useful trick, but nobody understands how it works'. Your friend Berthilde claims 'Dropout is good for generalization and easy to understand'. Comment on your results (think also of the other 4 combinations of dropping out two neurons) and relate your results to the claims of your friends.

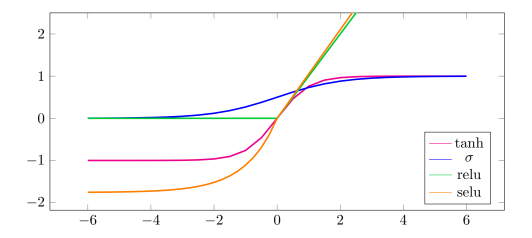
Exercise 2. Cross-validation

Assume K variants of a model, with model k having test error $E_k = E_0 + \epsilon_k$, where ϵ_k has mean $\mathbb{E}\left[\epsilon_k\right] = 0$, auto-variance $\mathbb{E}\left[\epsilon_k^2\right] = v$ and co-variance $\mathbb{E}\left[\epsilon_k\epsilon_n\right] = c$.

- a. What is the expected value of the *test error*, i.e. the expected test error of the model obtained by averaging over all variants?
- b. What is the variance of the average test error? Does the number K of variants play a role?
- c. Consider implementing K-fold cross validation. If every E_k is the error of one of the folds, how do variance v and correlation $\rho = \frac{c}{v}$ change with respect to K? How does the variance of the average test error behave as K varies, assuming that we have large number of sample points?

Exercise 3. Different activation functions

The choice of the non-linearity function g(x) can have a significant impact on learning speed and final performance. Which non-linearity is best, is still an active research question; the favorite non-linearity in the last century was probably the hyperbolic tangent $\tanh(x)$; since 2010, the rectified linear unit $\mathrm{relu}(x) = \max(0,x)$ is highly popular and there is a fair chance that the new favorite will be the scaled exponential linear unit $\mathrm{selu}(x) = \lambda x$ if x > 0 and $\mathrm{selu}(x) = \alpha(\exp(x) - 1)$ otherwise, with $\lambda \approx 1.0507$ and $\alpha \approx 1.75814$. Currently, it seems that the key concepts to discuss the different non-linearities are, first, linearity problem, second the vanishing gradient problem and, third, the bias shift problem.



a. Linearity problem

- (i) Show that a multi-layer neural network with linear activation function g(x) = x is equivalent to a single layer linear network. Hint: the product of two matrices is again a matrix.
- (ii) Assume that in each layer the inputs follow a Normal distribution with mean zero and small variance, i.e. $\sigma^2 \ll 1$. For which of the activation functions $\sigma(x) = 1/(1 + \exp(-x))$, $\tanh(x)$, $\mathrm{relu}(x)$ and $\mathrm{selu}(x)$ is a deep network basically equivalent to a linear network for this input distribution? Hint: Consider the case $\sigma^2 \to 0$ using a Taylor expansion around 0.

b. Vanishing gradient problem

- (i) Assume now the inputs are such that they also fall into the non-linear regimes. For simplicity we assume that in each layer the activations are $a_1 = -10$, $a_2 = -5$, $a_3 = -1$, $a_4 = 1$, $a_5 = 5$, $a_6 = 10$. Without a calculator, determine the fraction of values close to zero of $g(a_i)$ and $g'(a_i)$ for all i and $g = \sigma$, tanh, relu, selu. For example, for tanh none of the values $\tanh(-10)$, $\tanh(-5)$, ..., $\tanh(10)$ is close to zero but 4/6 = 2/3 of the values of $\tanh' = 1 \tanh^2$ are close to zero.
- (ii) The update of a weight w_{ij} is proportional to $g'(a_i) \cdot g(a_j)$. Determine the fraction of $g'(a_i) \cdot g(a_j)$ that are close to zero considering all combinations of a_i and a_j and all activations $g = \sigma$, tanh, relu, selu.
- (iii) The δ 's in backpropagation are in each layer multiplied with g'. Consider backpropagation through 3 layers, i.e. terms like $g'(a_i)g'(a_j)g'(a_k)$. Determine the fraction of such terms that are close to zero for $g = \sigma$, tanh, relu, selu.

c. Bias shift problem

Consider a simple classification task. The data exist in \mathbb{R}^N . Data points from C_0 (with target t=0) are uniformly distributed in each dimension such that $x_i \in [1,2]$ for i=1...N. Data points from C_1 (with target t=1) are uniformly distributed in each dimension such that $x_i \in [3,4]$ for i=1...N. We want to learn to classify points using a logistic sigmoid unit trained with the cross-entropy loss; from last week, this results in the weight update rule

$$\Delta w_i = \eta \cdot (t - y) \cdot x_i$$

where
$$y = \sigma\left(\sum_{i}^{N} w_{i} x_{i}\right)$$
.

Points are presented one at a time (i.e. stochastic gradient descent).

(i) Assume we start with all weights $w_i = 0$ and present the point \mathbf{x}^a from C_0 , update the weights, then present \mathbf{x}^b . Give the drive $a = \sum_i^N w_i x_i^b$ of the output unit in response to \mathbf{x}^b , in terms of η , \mathbf{x}^a and \mathbf{x}^b . Note: we do not yet need to specify which class \mathbf{x}^b belongs to.

- (ii) In general, we can encounter oscillations in stochastic gradient descent if a single training example strongly affects the network output for instance, if it results in the same network output for any possible input.
 - We assume that if a < -5, $y \approx 0$, and if a > 5, $y \approx 1$. Under what conditions will the network output y be the same for all possible inputs \mathbf{x}^b after the first training step? Can we choose a small enough η to prevent this, independent of N? What if we had chosen \mathbf{x}^a from C_1 instead?
- (iii) A common input normalization technique to to remove the mean from the dataset, such that $E[x_i] = 0$ across all dimensions x_i . Assume that each data point has an equal probability of coming from either C_0 and C_1 . What are the new data ranges for C_0 and C_1 after removing the mean? Repeating step (ii), do we get the same result?
- (iv) Consider a deep network where each hidden layer uses one of the following activation functions: tanh, σ , relu, or selu. Given what we've seen above, can you suggest one of the activation functions? Note that one layer's output is another layer's input.
- d. Summarize your results by ranking the different activation functions for each of the problems discussed in this exercise.

	linearity problem	vanishing gradient problem	bias shift problem
tanh			
σ			
relu			
selu			

Exercise 4. Normalization of activations across multiple layers

In class we have seen, that by an appropriate normalization of the input patterns (for each input component: zero mean, unit standard deviation) combined with a Gaussian distribution of input weights $(\langle w_{ij}^{(1)} \rangle = 0 \text{ and } \langle [w_{ij}^{(1)}]^2 \rangle = 1/N)$ we can ensure that the activation variable of neurons in the first layer has mean $\langle a_i^{(1)} \rangle = 0$ and variance $\langle [a_i^{(1)}]^2 \rangle = 1$. In the following we assume that the distribution of activations in layer 1 is standard Gaussian, i.e. $a_i^{(1)} \sim N(0,1)$.

The aim of the exercise is to go by induction from layer n to layer n+1. We start in layer 1.

Assume that neuron j in layer 1 has a rectified linear activation function, i.e., $x_j^{(1)} = [a_j^{(1)}]_+$.

- a. What is the mean $\langle x_j^{(1)} \rangle$?
- b. Assume that the weights in layer 2 are initialized with zero mean and variance $\langle [w_{kj}^{(2)}]^2 \rangle = [c^2]/N_1$ where N_1 is the number of hidden neurons in the first layer.

What is the mean activation $\langle \tilde{a}_k^{(2)} \rangle$ in layer 2? Here $\tilde{a}_k^{(2)} = \sum_{j=1}^{N_1} w_{kj}^{(2)} x_j^{(1)}$. The total activation of neuron k in layer 2 is $a_k^{(2)} = \tilde{a}_k^{(2)} - \theta_k$.

What value should you choose for the threshold θ_k in layer 2, so that $\langle a_k^{(2)} \rangle = 0$ in layer 2?

- c. Assume that you found a threshold so that $\langle a_k^{(2)} \rangle = 0$. Calculate the variance $\langle [a_k^{(2)}]^2 \rangle$ as a function of the constant c.
- d. Choose c such that the variance is one.
- e. Can you now go from layer 2 to layer 3?