Handmade Feedforward Neural Network

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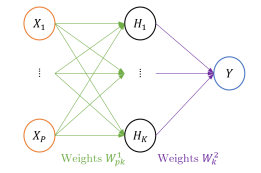
THEORY

The aim of the feedforward neural network is to predict the variable y (categorical for classification problem or continuous for regression problem) knowing the explanatory variables $x = \{x_1, \ldots, x_p\}$.

Firstly, you need to understand the *model structure* which consists of multiple layers of nodes or neurons fully connected with respective weights to the next layer in the network. This model is called feedforward because information flows only in one direction, from the input layer to the output layer.

Note: This paper covers the feedforward neural network with P neurons in the input layer, one hidden layer with K neurons, and one neuron in the output layer.

Input Layer Hidden Layer Output Layer



Activity of neuron $H_{ik} \rightarrow Z_{ik}^2 = X_{i1}W_{1k}^1 + \dots + X_{ip}W_{Pk}^1$ Activation of neuron $H_{ik} \rightarrow A_{ik}^2 = f(Z_{ik}^2)$ Activity of neuron $Y_i \rightarrow Z_i^3 = A_{i1}^2W_1^2 + \dots + A_{iK}^2W_k^2$ Activation of neuron $Y_i \rightarrow A_i^3 = f(Z_i^3)$ with $1 \le k \le K, 1 \le p \le P, 1 \le i \le N$

• Input layer

Corresponds to the input data $X_{[NxP]}$ with N observations and P explanatory variables. Then, the input layer has P nodes.

$$X_{[NxP]} = \begin{bmatrix} x_{11} & \cdots & x_{1P} \\ \vdots & \ddots & \vdots \\ x_{N1} & \cdots & x_{NP} \end{bmatrix}$$

• Hidden layer

The number of hidden layers ranges from one to many and the number of neurons is a tuning parameter with no optimal value for all cases. Using a lot of hidden layers gives name to the concept of *deep learning*.

• Output layer

Corresponds to the output data $Y_{[Nx1]}$ to predict. For predicting a continuous variable, it has only 1 neuron but for a categorical variable, it has as nodes as the number of classes in the variable.

$$Y_{[Nx1]} = \begin{bmatrix} y_1 \\ \vdots \\ y_N \end{bmatrix}$$

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Weights

The weights connect each neuron in the neural network and they are the parameters to optimize. When the *learning* or *training* process is finished, the weights are constant values that connect layers by multiplying them with the outputs of previous layers.

This is the weights matrix that connects the input layer with the hidden layer 1.

$$W_{[PxK]}^{1} = \begin{bmatrix} W_{11}^{1} & \cdots & W_{1K}^{1} \\ \vdots & \ddots & \vdots \\ W_{P1}^{1} & \cdots & W_{PK}^{1} \end{bmatrix}$$

This is the weights matrix that connects the hidden layer 1 with the output layer.

$$W_{[Kx1]}^2 = \begin{bmatrix} W_1^2 \\ \vdots \\ W_K^2 \end{bmatrix}$$

• Neuron activity

Except in the input layer, each node is a neuron with an activity and it is calculated as the linear combination of the outputs in the previous layer (neurons activation) and the weights that connect the previous layer with the actual node.

This is the formula for the activity in the layer 2 or hidden layer 1.

$$Z_{[NxK]}^2 = X_{[NxP]}W_{[PxK]}^1$$

This is the formula for the activity in the layer 3 or the output layer.

$$Z_{[Nx1]}^3 = A_{[NxK]}^2 W_{[Kx1]}^2$$

• Neuron activation

Except in the input layer, each node is a neuron that uses a nonlinear activation function and it means that in every node, its activity value is applied to an activation function. In short, it is the same idea that the *link function* of the *logistic regression* (explained in the paper *Handmade_Logistic_Regression.pdf*) but applied to every neuron in hidden and output layers.

This paper will use the *sigmoid activation function*.

$$f(z) = \frac{1}{1 + e^{-z}} = sigmoid(z)$$

This is the formula for the activation in the layer 2 or hidden layer 1.

$$A_{[NxK]}^2 = f\bigg(Z_{[NxK]}^2\bigg)$$

This is the formula for the activation in the layer 3 or the output layer.

$$\hat{Y}_{[Nx1]} = A_{[Nx1]}^3 = f\left(Z_{[Nx1]}^3\right)$$

Secondly, you need to understand the *learning* or *training* process. Basically, it consists of optimizing the weights that connect layers using the *cost function* which compares the prediction $\hat{Y}_{[Nx1]}$ with the expected output $Y_{[Nx1]}$.

The *learning* or *training* process is an optimization algorithm that repeats a two-phase cycle for each weights combination: *forward propagation* to make the prediction, and *backpropagation* to update the weights in the optimal direction. After repeating this process for a sufficiently large number of training iterations, the network will usually converge to a weights combination with a small prediction error.

Note: Although it seems more complex, it is the same process than in the paper $Handmade_Logistic_Regression.pdf$ or $Handmade_Linear_Regression.pdf$ where in every iteration of the weights optimization we firstly predicted the output and secondly updated the weights using the J gradient.

• Cost function

The prediction of the network is compared with the expected output using an *error function* or *loss function* or *cost function*. This paper will use as *cost function* the *Sum Square Error* divided by 2 (just to simplify the gradient calculation in *backpropagation*).

$$J = \frac{1}{2} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$

• Forward propagation

When an input data is presented to the network, it is propagated forward through the network, layer by layer, until it reaches the output layer and then the final prediction. In the figure of page 1, you can see all the steps for predicting a new observation from X_i until $\hat{Y}_i = A_i^3$.

• Backpropagation

Once the prediction is done through forward propagation, it is time to optimize the weights in order to reduce the value of the error predictions. To optimize or adjust weights properly in backpropagation, it is common to use the gradient descent algorithm which calculates the derivative of the cost function J with respect to the weights and updates the weights in the opposite direction of the gradient. Let's see the mathematical formula for the gradient descendent used in the optimization code.

Derivative of J by W_k^2 :

$$\frac{\partial J}{\partial W_k^2} = \sum_{i=1}^N \frac{2}{2} (y_i - \hat{y_i}) \frac{\partial (y_i - \hat{y_i})}{\partial W_k^2}$$

where

$$\frac{\partial (y_i - \hat{y_i})}{\partial W_k^2} = -\frac{\partial \hat{y_i}}{\partial W_k^2} = -\frac{\partial sigmoid(Z_i^3)}{\partial Z_i^3} \frac{\partial Z_i^3}{\partial W_k^2} = -\frac{e^{-Z_i^3}}{(1 + e^{-Z_i^3})^2} A_{ik}^2$$

then

$$\frac{\partial J}{\partial W_k^2} = \sum_{i=1}^N -(y_i - \hat{y_i}) \frac{e^{-Z_i^3}}{(1 + e^{-Z_i^3})^2} A_{ik}^2$$

Derivative of J by W_{pk}^1 :

$$\frac{\partial J}{\partial W_{pk}^1} = \sum_{i=1}^N \frac{2}{2} (y_i - \hat{y}_i) \frac{\partial (y_i - \hat{y}_i)}{\partial W_{pk}^1}$$

where

$$\frac{\partial (y_i - \hat{y_i})}{\partial W^1_{nk}} = -\frac{\partial \hat{y_i}}{\partial W^1_{nk}} = -\frac{\partial sigmoid(Z^3_i)}{\partial Z^3_i} \frac{\partial Z^3_i}{\partial W^1_{nk}} = -\frac{e^{-Z^3_i}}{(1 + e^{-Z^3_i})^2} \frac{\partial Z^3_i}{\partial A^2_{ik}} \frac{\partial A^2_{ik}}{\partial Z^2_{ik}} \frac{\partial Z^2_{ik}}{\partial W^1_{nk}} = -\frac{e^{-Z^3_i}}{(1 + e^{-Z^3_i})^2} W^2_k \frac{e^{-Z^2_{ik}}}{(1 + e^{-Z^2_{ik}})^2} x_i p^{-2}_i \frac{\partial Z^2_{ik}}{\partial Z^2_{ik}} \frac{\partial Z^$$

then

$$\frac{\partial J}{\partial W_{pk}^1} = \sum_{i=1}^N -(y_i - \hat{y_i}) \frac{e^{-Z_i^3}}{(1 + e^{-Z_i^3})^2} W_k^2 \frac{e^{-Z_{ik}^2}}{(1 + e^{-Z_{ik}^2})^2} x_i p$$

Finally, the gradient of J is used to update the weights in each iteration of the training process

$$W_{[PxK]}^{1new} = W_{[PxK]}^1 - \eta \frac{\partial J}{\partial W_{[PxK]}^1}$$

$$W_{[Kx1]}^{2new} = W_{[Kx1]}^2 - \eta \frac{\partial J}{\partial W_{[Kx1]}^2}$$

where η is the learning rate parameter.

CODE

The data used is the popular dataset *iris*. Let's predict the variable *Petal.Length* with the explanatory features: Sepal.Length and Sepal.Width. It is necessary to scale (divide by maximum value) the target continuous variable because we are using the sigmoid activation function which makes sense to predict $0 \le Y \le 1$.

```
x <- as.matrix(iris[,c("Sepal.Length","Sepal.Width")])
y <- as.matrix(iris[,c("Petal.Length")]/max(iris[,c("Petal.Length")]))
colnames(x) <- c("x1","x2")
colnames(y) <- c("y")</pre>
```

Set the neural network structure or number of neurons in each layer.

```
inputLayerSize <- ncol(x)
outputLayerSize <- ncol(y)
hiddenLayerSize <- 3</pre>
```

Set initial values for the weights randomly.

```
set.seed(1)
w1 <- matrix(rnorm(inputLayerSize*hiddenLayerSize),nrow=inputLayerSize,ncol=hiddenLayerSize)
w2 <- matrix(rnorm(hiddenLayerSize*outputLayerSize),nrow=hiddenLayerSize,ncol=outputLayerSize)</pre>
```

Start the weights optimization (training or learning) with gradient descent.

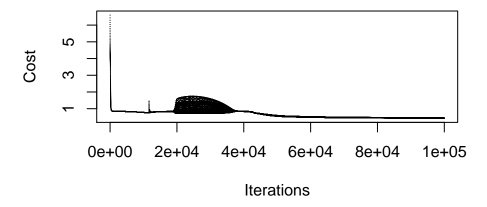
```
sigmoid <- function(z){
   1/(1+exp(-z))
}
derivativeSigmoid <- function(z){
   exp(-z)/((1+exp(-z))^2)
}
costs <- c()
rounds <- 100000
learning <- 0.01
for(i in 1:rounds){
   # Forward propagation: make prediction
   z2 <- x*,**w1  # Activity of layer 2
   a2 <- sigmoid(z2)  # Activation of layer 3
   yHat <- sigmoid(z3)  # Activation of layer 3 or final prediction</pre>
```

```
# Forward propagation: compute the actual cost
  costs <- c(costs, 0.5*sum((y-yHat)^2))
  # Backpropagation: calculate the J gradient by weights
   dJdW1 \leftarrow t(x)%*%(((2/2*(y-yHat)*(-1)*derivativeSigmoid(z3))%*%t(w2))*derivativeSigmoid(z2)) 
  dJdW2 \leftarrow t(a2)%*%(2/2*(y-yHat)*(-1)*derivativeSigmoid(z3))
  # Backpropagation: update the weights
  w1 <- w1-learning*dJdW1
  w2 <- w2-learning*dJdW2
Let's see the optimal estimated weights.
w1
```

```
##
             [,1]
                         [,2]
                                    [,3]
## x1 -0.3253937 -0.2997151 3.689311
## x2 -0.1795942 -0.3568836 -6.270115
w2
##
## [1,]
        -7.058921
## [2,] -11.894069
## [3,]
          2.243619
Let's plot the cost reduction in every iteration.
```

```
plot(costs, xlab="Iterations", ylab="Cost", main="Cost by iterations", cex=0.01)
```

Cost by iterations



Now, compare our model with the implemented function in R package neuralnet for feedforward neural network.

```
library(neuralnet)
data <- data.frame(x,y)</pre>
model <- neuralnet(y~x1+x2,data=data,hidden=hiddenLayerSize,learningrate=learning,</pre>
                    stepmax=rounds,exclude=c(1,4,7,10))
```

These are the optimal weights for the implemented model in R package *neuralnet*. The first row is NA because it corresponds to the *bias* or *intercept* weights which have not been included in our code.

model\$weights

```
## [[1]]
## [[1]][[1]]
##
               [,1]
                             [,2]
                                            [,3]
## [1,]
                 NA
                               NA
                                              NA
## [2,] 5.54799074
                     2.493967519 -0.2449272473
   [3,] 5.40596611 -4.259057181 -0.4992955809
##
## [[1]][[2]]
##
                  [,1]
## [1,]
                    NA
  [2,]
##
         0.4357913590
## [3,]
         0.6078436368
## [4,] -5.8861480345
```

These are the results for the first 10 observations of *iris* dataset.

```
head(data.frame(My_Predition=c(yHat),R_Prediction=model$net.result[[1]],True_Values=c(y)),10)
```

```
##
     My_Predition R_Prediction True_Values
## 1
     0.1814784963 0.2170673912 0.2028985507
## 2
     0.2349977071 0.2858761459 0.2028985507
## 3 0.1459365959 0.1603302129 0.1884057971
     0.1387709589 0.1476275976 0.2173913043
     0.1727102145 0.1952928411 0.2028985507
## 5
## 6
     0.2109175329 0.2453896849 0.2463768116
     0.1378970706 0.1346039599 0.2028985507
## 8 0.1733320933 0.2069756672 0.2173913043
     0.1270273827 0.1221395209 0.2028985507
## 10 0.1974963474 0.2469913656 0.2173913043
```