Handmade KNN

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THEORY

The aim of the *K Nearest Neighbours* algorithm is to predict a target variable y^{test} (categorical for *classification* problem or continuous for *regression problem*) in a *test* data comparing its associated known features $x^{test} = \{x_1^{test}, \dots, x_p^{test}\}$ with the features of a *train* data $x^{train} = \{x_1^{train}, \dots, x_p^{train}\}$ and using the known y^{train} values as a reference for the y^{test} prediction.

Firstly, let's see the main steps of this simple machine learning algorithm.

- 1. Define the inputs $x_{[NxP]}^{train}$, $y_{[Nx1]}^{train}$, $x_{[MxP]}^{test}$ and K.
- 2. For each M observation in the test data $x_{[MxP]}^{test}$, compute the mathematical distance (Euclidean in this paper) to each N observation in the train data $x_{[NxP]}^{train}$.
- 3. For each M observation in the test data $x_{[MxP]}^{test}$, select the K nearest (lower mathematical distance) observations in the train data $x_{[NxP]}^{train}$.
- 4. For each M observation in the test data $x_{[MxP]}^{test}$,
 - 4.1 In regression problem, compute the prediction $\hat{y}^{test} = mean(y_1^{train}, \dots, y_K^{train})$.
 - 4.2 In classification problem, compute the prediction $\hat{y}^{test} = mode(y_1^{train}, \dots, y_K^{train})$.

CODE

The data used is the popular dataset *iris*. Let's predict the categorical variable *Species* (then it is a *classification* problem with 3 categories) with the explanatory features: *Petal.Length* and *Petal.Width*.

Note: It is important to normalize (every observation minus the feature minimum value divided by the feature range) the features so that their units do not affect the mathematical distance. For example, if you compute the mathematical Euclidean distance from a person A = (1.80m, 70kg) and another B = (1.9m, 80kg) the result is

$$distance(A, B) = \sqrt{(1.8 - 1.9)^2 + (70 - 80)^2} = 10$$

But if you change the units to A = (180cm, 70000q) and B = (190cm, 80000q) the result is

$$distance(A, B) = \sqrt{(180 - 190)^2 + (70000 - 80000)^2} = 10000$$

Let's define the inputs $x_{[NxP]}^{train}$ and $y_{[Nx1]}^{train}$

```
duplicates <- duplicated(iris[,c("Petal.Length","Petal.Width")])
X_train <- iris[!duplicates,c("Petal.Length","Petal.Width")]
Y_train <- iris[!duplicates,"Species"]
X_train[,1] <- (X_train[,1]-min(X_train[,1]))/(max(X_train[,1])-min(X_train[,1]))
X_train[,2] <- (X_train[,2]-min(X_train[,2]))/(max(X_train[,2])-min(X_train[,2]))</pre>
```

Let's define $x_{[MxP]}^{test}$ by creating artificial data with all combinations of both features from 0 to 1 by 0.02, it will be interesting for observing the KNN decision boundaries in the following plots. Remember that the range of a normalized feature goes from 0 to 1.

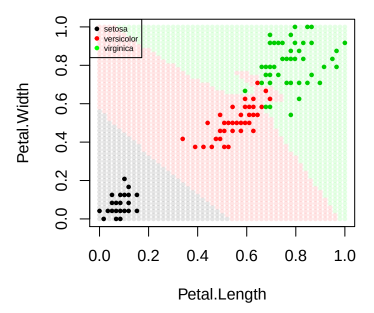
```
X_test <- expand.grid(list(Petal.Length=seq(0,1,0.02),Petal.Width=seq(0,1,0.02)))</pre>
```

Let's create the KNN algorithm function.

```
KNN <- function(X_train, X_test, Y_train, K){</pre>
  # Define function for mathematical distance between observations
  euclidean <- function(a,b){</pre>
    return(sqrt(sum((a-b)^2)))
  # Iterate test observations
  Y test <- c()
  for(i in 1:nrow(X_test)){
    # Iterate train observations and compute all distances to the test observation i
    distances <- c()
    for(j in 1:nrow(X_train)){
      distances <- c(distances, euclidean(X_train[j,1:2], X_test[i,]))</pre>
    }
    # Select K neighbours and compute the prediction for test observation i
    Y_neighbours <- Y_train[order(distances)][1:K]</pre>
    Y_test[i] <- names(which.max(table(Y_neighbours)))</pre>
  return(factor(Y_test))
}
```

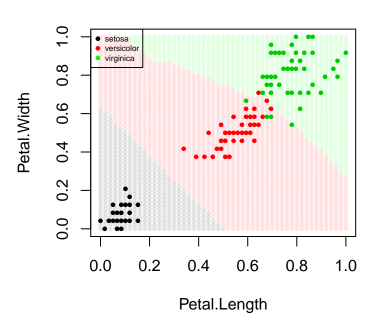
Now, let's apply the KNN function with K=1 and plot the result.

1NN



Now, let's apply the KNN function with K = 11 and plot the result.

11NN



In conclusion, we can observe that the decision boundary is more flexible or complex in 1NN than in 11NN, and it means that the bias of predictions is lower but the variance is higher. The idea is that averaging more neighbours gives a generalized prediction at the expense of bias increase, which can be a positive thing in test data for avoiding over-fitting.