1-ProjectDoc

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Goup work 1

File SNPdata * tab delimited * chromosome_position as row names * rows contains 0 for AA, 1 for Aa, 2 for aa (0 or 1 or 2 mutated alleles) * patient_id as columns * in column -> 1:1200 are patients, 1201:2000 are control subjects

qcalculation

- 1. Take in input numerical matrix in SNPdata like format
- 2. Calculate MAF q for each SNP (chromosome position)
- 3. return maf vector

```
qcalculation <- function(SNPdata) {
  calcq <- function(d) {
    N = length(d)
    AA = length(d[d == 0])
    Aa = length(d[d == 1])
    aa = length(d[d == 2])
    q = (aa * 2 + Aa) / (2 * N)
    return(q)
}

out = as.data.frame(apply(SNPdata, 1, calcq)) # apply calcq to rows
  colnames(out)="MAF"
  return(out)
}</pre>
```

HWEtest

- 1. Take in input numerical matrix in SNPdata like format
- 2. Compute a HWE test for each SNP given as input by calculating the χ^2_{obs} from the data
 - $\chi^2_{obs} = \sum_{i \in \{AA, Aa, aa\}} \frac{O_i Np_i}{N}$ where $p_{AA} = p^2, p_{Aa} = 2pq, p_{aa} = q^2$ and O_i are the number of observed AA,Aa,aa from the provided data.
 - By computing the p value using the function pchisq (DO NOT use directly the chisq.test() function)
 - pchisq(chi_squared, 1, lower.tail = FALSE)
- 3. Return the HWE test p-values for each SNP as a vector of numeric values with names corresponding to the SNP IDs with the same order they had in the input matrix

```
HWEtest <- function(SNPdata) {</pre>
 calcp<- function(d) {</pre>
   N = length(d)
   ## Calculate observed values -----
   AA = length(d[d == 0])
   Aa = length(d[d == 1])
   aa = length(d[d == 2])
   0 = c(AA, Aa, aa)
   ## Expected frequencies -----
   q = (aa * 2 + Aa) / (2 * N)
   p = 1 - q
   prob_expec=c(p^2,2*p*q,q^2)
   ## ChiSquared -----
   chi_squared=sum((0-N*prob_expec)^2/(N*prob_expec))
   ## pvalue -----
   pvalue <- pchisq(chi_squared, 1, lower.tail = FALSE)</pre>
   return(pvalue)
 }
 out = as.data.frame(apply(SNPdata, 1, calcp))
 colnames(out)=c("pvalue")
 return(out)
}
```

VARIANTanalysis

- 1. take as input
 - filepath of file with SNPdata like format
 - indCTRL vector indicating which column has data from control subjects
 - MAFth treshold to filter SNPs lower MAF, default 0.01
 - HWEalpha significance level alpha to be used to filter SNP with lower p-values because possibly not in HWE
- 2. read the file and analyse variants
 - filter out the one with MAF < MAFth \lor HWE-p-value < HWEalpha
 - SNPdata = SNPdata[(qcalculation(SNPdata)>MAFth & HWEtest(SNPdata)[2]>HWEalpha)
 ,]
 - calculate χ^2_{obs}
 - use HWEtest, applied to control only (as in suggestion)
 - calculate p-values
 - use HWEtest, applied to control only (as in suggestion)
- 3. Compute q-value for each SNP using Benjamini-Hockberg procedure
 - $qvalue_i = pvalue_i \times \frac{rank_i}{N}$, $i \in 1, N$, $rank_i$ equal to position of variant in list ordered by pvalue
- 4. Return matrix with data AA_ctrl, Aa_ctrl, aa_ctrl, AA_case, Aa_case, aa_case, pval, qval.

```
VARIANTanalysis <-
  function(filepath,
           indCTRL,
           MAFth = 0.01,
           HWEalpha = 0.01) {
  ## Input and parameters setup ----
  SNPdata <- read.table("SNPdata.txt", header = TRUE, sep = "\t")</pre>
  SNPdata = SNPdata[qcalculation(SNPdata[indCTRL,]) > MAFth &
                      HWEtest(SNPdata[indCTRL,]) > HWEalpha, ]
  `%notin%` <- Negate(`%in%`)</pre>
  N = dim(SNPdata)[2]
  indPATI = 1:N
  indPATI = indPATI[indPATI %notin% indCTRL]
  ## Function definitions
  calcallele <- function(d, indCTRL, indPATI) {</pre>
    AA = length(d[d == 0])
    Aa = length(d[d == 1])
    aa = length(d[d == 2])
    0 = c(AA, Aa, aa)
    return(0)
  calcchip<- function(d) {</pre>
    N = length(d)
    0 <- calcallele(d)</pre>
    q = (0[3] * 2 + 0[2]) / (2 * N)
    p = 1 - q
    prob_expec=c(p^2,2*p*q,q^2)
    chi_squared=sum((0-N*prob_expec)^2/(N*prob_expec))
    pvalue <- pchisq(chi_squared, 2, lower.tail = FALSE)</pre>
    return(c(chi_squared, pvalue))
  }
  ## Calculations -----
  0 = as.data.frame(cbind(
                                                        #observed data
        t(apply(SNPdata[, indCTRL], 1, calcallele)),
        t(apply(SNPdata[, indPATI], 1, calcallele))
      ))
  t=as.data.frame(t((apply(SNPdata[indCTRL],1, calcchip))))
  chisquared=t[,1]
                                                        # chisquared
  pvalues = t[,2]
                                                        # pvalues
  ### calculate q
                                                        # qvalues
  r = order(unlist(pvalues), decreasing = FALSE)
  #qvalues = pvalues * N / r
  qvalues = pvalues * r / N # statistical book alpha*rank/total
  ## Final Output
  out = as.data.frame(cbind(0, pvalues, qvalues))
  colnames(out) <-c("AA_ctrl","Aa_ctrl","aa_ctrl","AA_case","Aa_case","aa_case","pval","qval")
```

```
return(as.data.frame(out))
}
```

Tests

```
SNPdata <- read.table("SNPdata.txt", header = TRUE, sep = "\t")
vec_maf <- qcalculation(SNPdata = SNPdata)
vec_HWE_chi_pvalue <- HWEtest(SNPdata)
vec_VarAnal = VARIANTanalysis("SNPdata.txt", 1201:2000, MAFth = 0.05)</pre>
```