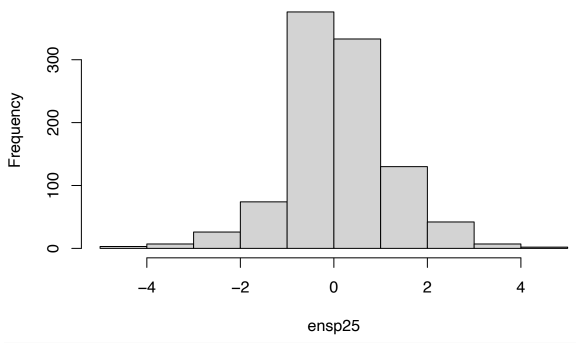
## Histogram of ensp25



```
rm(list=ls())
set.seed(123)
ensp50 = c()
for (j in 1:1000){
  n0bs = 100
  nPreds = 50
  #generate random values from N(0,1)
  dat = rnorm(nObs * (1 + nPreds))
  #create a matrix of the values
  dataset = matrix(data = dat, nrow = nObs, ncol = nPreds + 1)
  dframe = as.data.frame(dataset)
  strs = c("y")
  for (k in 1:nPreds){
    strs = c(strs, paste("x", toString(k), sep=""))
  colnames(dframe) = c(strs)
  #linear model
  model = lm(y ~ ., data=dframe)
  vals = summary(model)$coefficients
  sig_vals = vals[ ,4]
  \#count\ number\ of\ predictors\ with\ val < 0.05
  counter1 = 0
  for (i in 1:length(sig_vals)){
    if (sig_vals[i] <= 0.05){</pre>
      counter1 = counter1 + 1
    }
  }
```

```
#now we check for the significant predictors (those with a val < 0.25)
  keep = c(1)
  for (i in 1:length(sig_vals)){
   if (sig_vals[i] <= 0.25){</pre>
      keep = append(keep, i)
  }
  new_matrix = dataset[ , keep]
  new_dframe = as.data.frame(new_matrix)
  new_names = c("ynew")
  for (k in 1:(length(keep)-1)){
   new_names = c(new_names, paste("x", toString(k), sep=""))
  colnames(new_dframe) = c(new_names)
  #new regression:
  new_model = lm(ynew ~ ., data=new_dframe)
  new_vals = summary(new_model)$coefficients
  new_sig_vals = new_vals[ ,4]
  #count number of predictors with val < 0.05
  counter2 = 0
  for (i in 1:length(new_sig_vals)){
   if (new_sig_vals[i] <= 0.05){</pre>
      counter2 = counter2 + 1
   }
 }
  ensp50 = c(ensp50, (counter2 - counter1))
## Warning in summary.lm(new_model): essentially perfect fit: summary may be
## unreliable
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```