RAProject

Mingzhi Ye

2022/10/10

Get the combined distribution of number of Alpha chains and number of Beta chains in a clonotype

```
IndexnAnBMap=read.csv("IndexnAnBMap.csv",header = F)
IndexnAnBMap=as.matrix(IndexnAnBMap)
```

Get a template dataframe of a clonotype which has one alpha chain and one beta chain

```
sce0 <- readRDS('toyTCRdata.rds')
TemplateClonotype=sce0$contigs[[4]][c(2,1),]</pre>
```

Generate sample pool

```
poolSize=2000
SampleSize=5000
AmiNoAcids=c('A','R','N','D','C','Q','E','G','H','I','L','K','M','F','P','S','T','W','Y','V')
getCDR3 <- function(x){
    cdr3=sample(AmiNoAcids,15, replace=T)
    return(paste(cdr3,collapse = ''))
}

getClonotypeForPool<-function(clonotype_index, CDRAlpha, CDRBeta){
    result=TemplateClonotype
    result$cdr3[1]=CDRAlpha
    result$cdr3[2]=CDRBeta
    return(result)
}

getSamplePool <- function(poolSize){
    Alpha_pool=sapply(1:poolSize,getCDR3,simplify = TRUE)</pre>
```

```
Beta_pool=sapply(1:poolSize,getCDR3,simplify = TRUE)

CDR_Alpha=sample(Alpha_pool,poolSize, replace=T)

CDR_Beta=sample(Beta_pool,poolSize, replace=T)

barcode <- paste('cell', 1:poolSize, replace=T)

samplePool <- lapply(1:poolSize, function(clonotype_index){
    getClonotypeForPool(clonotype_index, CDR_Alpha[clonotype_index], CDR_Beta[clonotype_index])})

samplePool <- SplitDataFrameList(samplePool)
samplePool[,'barcode'] <- barcode
samplePool[,'sample'] <- 'isim'
names(samplePool) <- barcode

# sampleI<-as(sample, 'SplitDFrameList')
return(samplePool)
}
samplePool=getSamplePool(poolSize)</pre>
```

Generate the dataframe recording the abundance for each clonotype in the sample pool

```
clonotypes_in_pool=samplePool[,'cdr3']
concatCDR3<-function(x){
  res=paste(x[1],x[2],sep=' ')
    res=paste(x[1],x[2],sep=' ')
  return(res)
}
clonotypes_in_pool=lapply(clonotypes_in_pool,concatCDR3)
clonotypes_in_pool=unlist(clonotypes_in_pool)
abundance=rep(1/poolSize,poolSize)
realDF<- data.frame(clonotypes_in_pool,abundance)
rownames(realDF) <- NULL
realDF=aggregate(realDF$abundance, by=list(clonotype=realDF$clonotypes_in_pool), FUN=sum)
colnames(realDF) <- c('clonotype','abundance')</pre>
```

Functions for generate a sample

```
getClonotype<-function(clonotype_index, numberA, numberB, errorProb = .01){
    result=DataFrame()
    indexInPool=sample.int(poolSize, 3, replace=FALSE)</pre>
```

```
setAlpha <- hashset()</pre>
setBeta <- hashset()</pre>
if (numberA>=1){
    if(runif(1) < errorProb){</pre>
      result=rbind(result,samplePool[[indexInPool[2]]][1,])
      insert(setAlpha,indexInPool[2])
      result=rbind(result,samplePool[[indexInPool[1]]][1,])
      insert(setAlpha,indexInPool[1])
    numberA=numberA-1
if (numberB>=1){
    if(runif(1) < errorProb){</pre>
      result=rbind(result,samplePool[[indexInPool[3]]][2,])
      insert(setBeta,indexInPool[3])
    }else{
      result=rbind(result,samplePool[[indexInPool[1]]][2,])
      insert(setBeta,indexInPool[1])
    }
    numberB=numberB-1
}
while(numberA>0){
    indexInPool=sample.int(poolSize, 1, replace=TRUE)
    while(setAlpha[indexInPool] == TRUE) {
      indexInPool=sample.int(poolSize, 1, replace=TRUE)
    insert(setAlpha,indexInPool)
    result=rbind(result,samplePool[[indexInPool]][1,])
    numberA=numberA-1
}
while(numberB>0){
    indexInPool=sample.int(poolSize, 1, replace=TRUE)
    while(setBeta[indexInPool] == TRUE) {
      indexInPool=sample.int(poolSize, 1, replace=TRUE)
    insert(setBeta,indexInPool)
    result=rbind(result,samplePool[[indexInPool]][2,])
    numberB=numberB-1
}
return(result)
```

Use the functions to generate a sample with size 50

 $However,\ function\ `clonoStats'\ is\ not\ compatible\ with\ SimpleD-FrameList,\ how\ to\ transform\ SimpleD-FrameList\ to\ CompressedSplitDFrameList$

```
getSample <- function(samplesize){</pre>
     RandomIntegers <- sample(1:(length(IndexnAnBMap)/2), samplesize, replace=T)</pre>
    barcode <- paste('cell', 1:samplesize)</pre>
     sample <- lapply(1:samplesize, function(clonotype_index){</pre>
          getClonotype(clonotype_index, IndexnAnBMap[RandomIntegers[clonotype_index],1], IndexnAnBMap[RandomIntegers[clonotype_index],1], IndexnAnBMap[RandomIntegers[clonotype_index],1]
    })
    sample <- SplitDataFrameList(sample)</pre>
     sample[,'barcode'] <- barcode</pre>
     sample[,'sample'] <- 'sim'</pre>
    names(sample) <- barcode</pre>
     # sample1<-as(sample, 'SplitDFrameList')</pre>
    return(sample)
}
# samplelist=getSample(SampleSize)
# EMpredicted <- clonoStats(samplelist, method = 'EM')</pre>
# UNpredicted <- clonoStats(samplelist, method = 'unique')</pre>
```

Get Variation Distance

```
getDistance_R<-function(clonotype,abundance){
   stimulated<- data.frame(clonotype,abundance)

sum_abundance=sum(stimulated['abundance'])
   stimulated['abundance']=stimulated['abundance']/sum_abundance

merged = merge(x = realDF, y = stimulated, by = "clonotype",all = TRUE)
   merged[is.na(merged)] <- 0
   return (sum(abs(merged['abundance.x']-merged['abundance.y'])))
}</pre>
```

Test

```
timeO=Sys.time()
samplelist=getSample(5000)
time1=Sys.time()
print(time1-time0)
EMpredicted <- clonoStats(samplelist, method = 'EM')
time2=Sys.time()</pre>
```

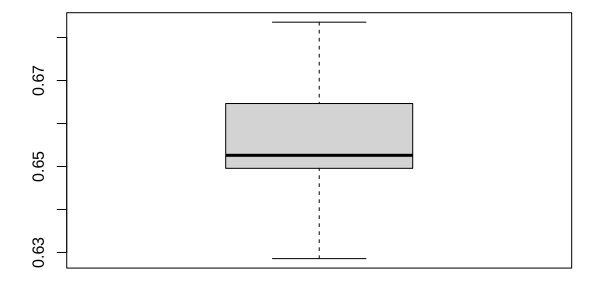
```
print(time2-time1)
distance1=getDistance_R(clonoNames(EMpredicted),clonoAbundance(EMpredicted)[,1])
time3=Sys.time()
print(time3-time2)
UNpredicted <- clonoStats(samplelist, method = 'unique')
time4=Sys.time()
print(time4-time3)
distance2=getDistance_R(clonoNames(UNpredicted),clonoAbundance(UNpredicted)[,1])
time5=Sys.time()
print(time5-time4)</pre>
```

Calculate distances for samples

```
DistanceListFromEM=c()
DistanceListFromUN=c()
for(i in 1:20){
    samplelist=getSample(SampleSize)
    EMpredicted <- clonoStats(samplelist, method = 'EM')
    distance1=getDistance_R(clonoNames(EMpredicted),clonoAbundance(EMpredicted)[,1])
    DistanceListFromEM[i]=distance1
    UNpredicted <- clonoStats(samplelist, method = 'unique')
    distance2=getDistance_R(clonoNames(UNpredicted),clonoAbundance(UNpredicted)[,1])
    DistanceListFromUN[i]=distance2
}</pre>
```

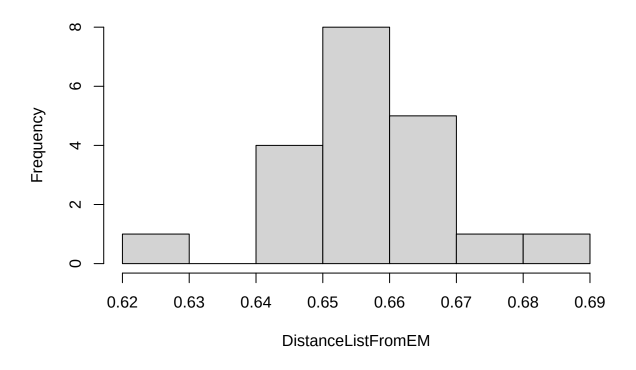
Visualize

```
boxplot(DistanceListFromEM)
```

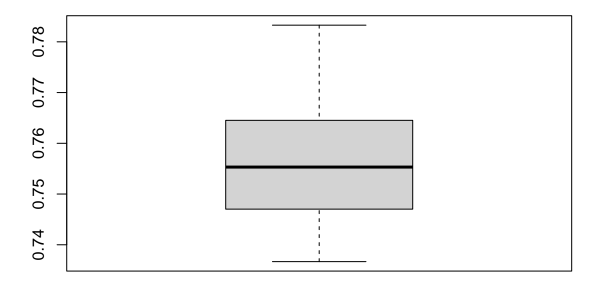


hist(DistanceListFromEM)

Histogram of DistanceListFromEM

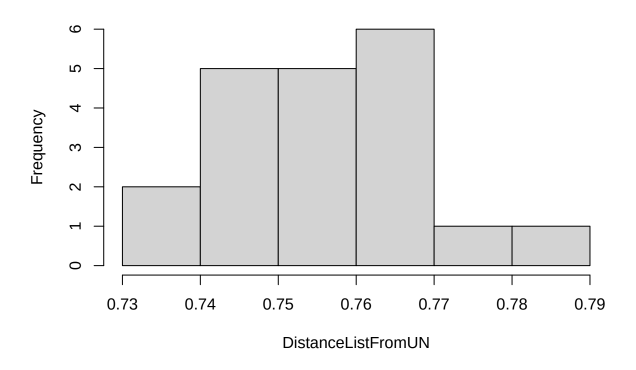


boxplot(DistanceListFromUN)



hist(DistanceListFromUN)

Histogram of DistanceListFromUN



print(median(DistanceListFromEM))

[1] 0.6526006

print(median(DistanceListFromUN))

[1] 0.7553123

print(mean(DistanceListFromEM))

[1] 0.6563997

print(mean(DistanceListFromUN))

[1] 0.7555607