## het

## TMJ

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When you click the **Knit** button a document will be generated that includes both content as well as the output of any embedded R code chunks within the document. You can embed an R code chunk like this:

read the 012 file and combine genotype with individual

```
setwd("~/Google Drive/final_analysis_LRS/pilot_22Apr2016_final/workdir")
genos <- read.delim("out.012", header=FALSE)
indiv <- read.table("out.012.indv", quote="\"", comment.char="")
genos$V1 <- NULL
gi <- cbind(indiv, genos)
row.names(gi) <- gi$V1
gi=gi[,-1]
tgi <- as.data.frame(t(gi))</pre>
```

divide into seperate files and calculate each hybrid tool combination seperately

```
samBC <- tgi[ ,1:3]
samBM <- tgi[ ,4:6]
gatBC <- tgi[ ,7:9]
gatBM <- tgi[ ,10:12]</pre>
```

script used to build loop: samBCt <- samBC[samBC[,1]==0 & samBC[,2]==2 & samBC[,3]!=-1,] dim(samBCt) totLoci <- sum(samBCt[,1]==0) totLoci refAllele <- sum(samBCt[,3]==0) refAllele altAllele <- sum(samBCt[,3]==2) altAllele hetAllele <- sum(samBCt[,3]==1) hetAllele

hetaccuracy <- hetAllele/totLoci hetaccuracy\*100

function for finding hybrid accuracy rate for each file. I.e. if parents are 0 and 2, are hets 1?

```
hfx<-function(file){
filet <- file[file[,1]==0 & file[,2]==2 & file[,3]!=-1,]
dim(filet)
refAllele <- sum(filet[,3]==0)
refAllele
altAllele <- sum(filet[,3]==2)
altAllele
hetAllele <- sum(filet[,3]==1)
hetAllele
totLoci <- sum(filet[,1]==0)
totLoci
hetaccuracy <- hetAllele/totLoci
hetaccuracy*100
}</pre>
```

run the function across the 4 subfiles

```
hetfiles <- list(samBC, samBM, gatBC, gatBM)
resultcorrecthybrid <- lapply(hetfiles, hfx)
#hetfilenames <- list("samBC", "samBM", "gatBC", "gatBM")
#correcthybrid <- as.data.frame(cbind(hetfilenames, resultcorrecthybrid))
#correcthybrid$resultcorrecthybrid <- as.numeric(correcthybrid$resultcorrecthybrid)
```

count number of hets in hybrid

```
script used to build loop: length(which(samBC$Xpress_0183.R_2015_08_24_16_56_10_user_PG1-118-NCSU-2.fastq == 1)) gatBC2 <- gatBC[gatBC[,1]!=-1 & gatBC[,2]!=-1 & gatBC[,3]==1,] dim(gatBC2) totLoci <- sum(gatBC2[,3]==1) totLoci correctcalls <- sum(gatBC2[,1]==0 & gatBC2[,2]==2) correctcalls accuracy <- correctcalls/totLoci accuracy
```

make a file if parents are correct. I.e. if hybrid is a 1 are parents 0 and 2?

```
correctparentfx <- function(file){
file2 <- file[file[,1]!=-1 & file[,2]!=-1 & file[,3]==1,]
dim(file2)
totLoci <- sum(file2[,3]==1)
totLoci
correctcalls <- sum(file2[,1]==0 & file2[,2]==2)
correctcalls
accuracy <- correctcalls/totLoci
accuracy*100
}</pre>
```

make a table with het accuracy rates for each hybrid pair

```
resultcorrectparents <- lapply(hetfiles, correctparentfx)
hetfilenames <- list("samBC", "samBM", "gatBC", "gatBM")
hetaccuracy <- as.data.frame(cbind(hetfilenames, resultcorrecthybrid, resultcorrectparents))
hetaccuracy$resultcorrecthybrid <- as.numeric(hetaccuracy$resultcorrecthybrid)
hetaccuracy$resultcorrectparents <- as.numeric(hetaccuracy$resultcorrectparents)
hetaccuracy$hetfilenames <- as.character(hetaccuracy$hetfilenames)
hetaccuracy</pre>
```

```
hetfilenames resultcorrecthybrid resultcorrectparents
##
## 1
            \mathtt{samBC}
                               82.70440
                                                      91.00346
## 2
            samBM
                               78.68020
                                                      88.06818
## 3
             gatBC
                               97.66355
                                                      91.26638
                                                      93.54839
## 4
            gatBM
                               95.08197
```

```
write.csv(hetaccuracy, file="hetaccuracies.csv")
```

for table with averages

```
samavghyb <- (hetaccuracy[1,2]+hetaccuracy[2,2])/2
gatavghyb <- (hetaccuracy[3,2]+hetaccuracy[4,2])/2
samavgpar <- (hetaccuracy[1,3]+hetaccuracy[2,3])/2
gatavgpar <- (hetaccuracy[3,3]+hetaccuracy[4,3])/2</pre>
```

output table and file with averages

```
tool <- list("samavg", "gatavg")
correcthyb <- list(samavghyb, gatavghyb)
correctpar <- list(samavgpar, gatavgpar)
averages <- as.data.frame(cbind(tool, correcthyb, correctpar))
averages$correcthyb <- as.numeric(averages$correcthyb)
averages$correctpar <- as.numeric(averages$correctpar)
averages$tool <- as.character(averages$tool)
averages</pre>
## tool correcthyb correctpar
## 1 samavg 80.69230 89.53582
## 2 gatavg 96.37276 92.40738
```

write.csv(averages, file="hetaccuraciesaverages.csv")