Indirect Selection

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Path Normalization

Tajima's D

R function for Tajima'D calculation

$$D = \frac{\pi - \theta_W}{\sqrt{Var(\pi - \theta_W)}}$$

```
TajimaD <- function(sfs){</pre>
    #' sfs (site frequency spectrum): number of singletons, doubletons, ..., etc
    n <- length(sfs) + 1 # number of chromosomes</pre>
    ss <- sum(sfs) # number of segregating sites
    a1 \leftarrow sum(1 / seq_len(n-1))
    a2 \leftarrow sum(1 / seq_len(n-1)^2)
    b1 \leftarrow (n + 1) / (3 * (n - 1))
    b2 \leftarrow 2 * (n^2 + n + 3)/(9 * n * (n - 1))
    c1 <- b1 - 1/a1
    c2 \leftarrow b2 - (n + 2)/(a1 * n) + a2 / a1^2
    e1 <- c1 / a1
    e2 < - c2 / (a1^2 + a2)
    Vd \leftarrow e1 * ss + e2 * ss * (ss - 1)
    theta_pi \leftarrow sum(2 * seq_len(n-1) * (n - seq_len(n-1)) * sfs)/(n*(n-1))
    theta_w <- ss / a1
    res <- (theta_pi - theta_w) / sqrt(Vd)
    return(res)
}
```

Site Freq Spectrum

Simulate one SFS

```
df <- data.frame(allele=c(1,2,3,4,5), frq=c(20,3,1,1,1))</pre>
TajimaD(sfs=df$frq)
#install.packages("ggplot2")
library(ggplot2)
fsize=18 #font size
p1 <- ggplot(data=df,aes(x=df$allele, y=df$frq)) +
        geom_bar(stat="identity", position=position_dodge()) +
        theme bw() +
        theme(#axis.text.x=element_blank(), #axis.ticks.x=element_blank(),
          axis.text=element_text(size=fsize),
          axis.title=element_text(size=fsize, face="bold"),
          legend.title = element_text(size=fsize, face="bold"),
          legend.text = element_text(size=fsize)) +
      xlab("# of individuals with derived alleles") +
      ylab("Counts")
p1
```

```
# Site
Freq
Spec-
trum
Simulate
two
SFS
"'r df1
data.frame(allele=c(1,2,3,4,5),
frq=c(20,2,1,1,1),
sel="Sweep")
df2 <-
data.frame(allele=c(1,2,3,4,5),
frq=c(6,1,2,2,4),
sel="Neutral")
df <-
rbind(df1,
df2
TajimaD(sfs=df1frq)TajimaD(sfs =
df2frq
```

```
p2 <-
gg-
plot(df,
aes(x=allele,
y = frq,
fill=sel))
geom_bar(stat="identity",
posi-
tion=position_dodge())
xlab("#
of in-
divid-
uals
with
de-
rived
alle-
les") +
ylab("Counts")
#scale_fill_manual(values=c("#E69F00","#56B4E9",
"#009E73"))
\#scale\_x\_discrete(labels=c("-
log10(mu)","-
log10(nu)","Ne*s"))
theme (legend. position\\
"top",
leg-
end.title
= ele-
ment_blank(),
axis.text=element\_text(size=10),
axis.title=element_text(size=fsize,
face="bold"),
leg-
end.text
= ele-
ment_text(size=fsize))
p2 "'
```

Obtain SFS from the sequencing data

Now we switch from our local computer to HCC

```
ssh ID@crane.unl.edu
cd YOUR Git Repo
git pull
```

```
# request a computing node
srun --qos=short --nodes=1 --licenses=common --ntasks=4 --mem 8G --time 1:30:00 --pty bash
module load R
R
```

```
# Ob-
tain
SFS
from
the se-
quenc-
ing
data
# Ob-
tain
SFS
from
the se-
quenc-
ing
data
r df
read.csv("cache/Mt_derived_alleles.csv")
<-
table(df$da)
TajimaD(sfs=sfs)
```

Calculate Tajima's D for windows (10 kb)

```
df <- read.csv("cache/Mt_derived_alleles.csv")</pre>
names(df)[1:2] <- c("chr", "pos")</pre>
winsize = 10000
df$win <- round(df$pos/winsize,0) + 1</pre>
res <- data.frame()</pre>
sfs0 <- data.frame(Var1=1:19, value=0)</pre>
for(i in 1:58){
  sub <- subset(df, win %in% i)</pre>
  tem <- as.data.frame(table(sub$da))</pre>
  if(nrow(tem) > 0){
    newsfs <- merge(sfs0, tem, by="Var1", all.x=TRUE)</pre>
    newsfs[is.na(newsfs$Freq),]$Freq <- 0</pre>
    out <- data.frame(win=i, tajimad = TajimaD(sfs=newsfs$Freq))</pre>
    res <- rbind(res, out)</pre>
  }
}
```

Visualize the Tajima'D results

Scatter plot

```
pdf("graphs/tajimad_res.pdf", height=5, width=10)
plot(x=res$win, y=res$tajimad, pch=16, col="red", xlab="Physical Position (10-kb)", ylab="Tajima D")
dev.off()
```

Histogram

```
pdf("graphs/hist_tajimad_res.pdf", height=5, width=5)
hist(res$tajimad, xlab="Tajima D", ylab="Frequency")
dev.off()
```

General feature for- $_{\mathrm{mat}}$ (GFF) from EnsemblPlants Maize reference genome change largedata\lab6 folder: bash cd largedata mkdir lab6 cd lab6

We will download and unzip the Mt GFF3 file

```
wget ftp://ftp.ensemblgenomes.org/pub/plants/release-46/gff3/zea_mays/Zea_mays.B73_RefGen_v4.46.chromos
### then unzip it
gunzip Zea_mays.B73_RefGen_v4.46.chromosome.Mt.gff3.gz
```

General feature format (GFF) version 3

```
۷2
  ۷1
                     ٧3
                           ۷4
                                  V5 V6 V7 V8
1 Mt Gramene chromosome
                           1 569630
2 Mt ensembl
                   gene 6391
                                6738
3 Mt
        NCBI
                   mRNA 6391
                                6738
4 Mt
                   exon 6391
        NCBI
                                6738
5 Mt
        NCBI
                    CDS 6391
                                6738
6 Mt ensembl
                   gene 6951
                                8285
    ID=chromosome:Mt;Alias=AY506529.1,NC_007982.1;Is_circular=true
2
    ID=gene:ZeamMp002;biotype=protein_coding;description=orf115-a1;
3
    ID=transcript:ZeamMp002;Parent=gene:ZeamMp002;
4
    Parent=transcript:ZeamMp002;Name=ZeamMp002.exon1;constitutive=1;ensembl_end_phase=0;
5
    ID=CDS:ZeamMp002;Parent=transcript:ZeamMp002;
6
    ID=gene:ZeamMp003;biotype=protein_coding;description=orf444
```

General feature format (GFF) version 3

```
V1 V2 V3 V4 V5 V6 V7 V8

1 Mt Gramene chromosome 1 569630 . . .

2 Mt ensembl gene 6391 6738 . + .

V9

1 ID=chromosome:Mt;Alias=AY506529.1,NC_007982.1;Is_circular=true
2 ID=gene:ZeamMp002;biotype=protein_coding;description=orf115-a1;
```

- 1 **sequence**: The name of the sequence where the feature is located.
- 2 **source**: Keyword identifying the source of the feature, like a program (e.g. RepeatMasker) or an organization (like ensembl).
- 3 feature: The feature type name, like "gene" or "exon".
 - In a well structured GFF file, all the children features always follow their parents in a single block (so all exons of a transcript are put after their parent "transcript" feature line and before any other parent transcript line).
- 4 start: Genomic start of the feature, with a 1-base offset.
 - This is in contrast with other 0-offset half-open sequence formats, like BED.

General feature format (GFF) version 3

```
۷1
٧2
VЗ
۷4
V5 V6
V7 V8
1 Mt
Gramene
chromosome
569630
. 2
Mt
ensembl
gene
6391
6738
. +
. V9
1
ID=chromosome:Mt;Alias=AY506529.1,NC_007982.1;Is_circular=tru
ID=gene:ZeamMp002;biotype=protein_coding;description=orf115-a
```

- 5 end: Genomic end of the feature, with a 1-base offset.
 - This is the same end coordinate as it is in 0-offset half-open sequence formats, like BED.
- 6 score: Numeric value that generally indicates the confidence of the source in the annotated feature.
 - A value of "." (a dot) is used to define a null value.
- 7 strand: Single character that indicates the strand of the feature.
 - it can assume the values of "+" (positive, or 5' -> 3'), "-", (negative, or 3' -> 5'), "." (undetermined).

$\operatorname{Gen-}$ eral feature for- $_{\mathrm{mat}}$ (GFF) version3

```
۷1
٧2
VЗ
V4
V5 V6
V7 V8
1 Mt
Gramene
chromosome
569630
. 2
Mt
ensembl
gene
6391
6738
. V9
1
ID=chromosome:Mt;Alias=AY506529.1,NC_007982.1;Is_circular=tru
ID=gene:ZeamMp002;biotype=protein_coding;description=orf115-a
```

- 8 phase: phase of CDS (means CoDing Sequence) features.
 - The phase indicates where the feature begins with reference to the reading frame.
 - The phase is one of the integers 0, 1, or 2, indicating the number of bases that should be removed from the beginning of this feature to reach the first base of the next codon.
- 9 attributes: All the other information pertaining to this feature.
 - The format, structure and content of this field is the one which varies the most between the three competing file formats.

Use R to process the GFF3 file

```
# install.package("data.table")
library("data.table")

## simply read in wouldn't work
gff <- fread("largedata/lab6/Zea_mays.B73_RefGen_v4.46.chromosome.Mt.gff3", skip="#", header=FALSE, dat

## grep -v means select lines that not matching any of the specified patterns
gff <- fread(cmd='grep -v "#" largedata/lab6/Zea_mays.B73_RefGen_v4.46.chromosome.Mt.gff3', header=FALS.

rename each column

names(gff) <- c("seq", "source", "feature", "start", "end", "score", "strand", "phase", "att")
table(gff$feature)</pre>
```

Use R to process the GFF3 file

Get the start and end positions for each gene

```
gene <- subset(gff, feature %in% "gene")
gene$geneid <- gsub(".*gene:|;biotype.*", "", gene$att)</pre>
```

Calculate Tajima's D for each gene

```
df <- read.csv("cache/Mt_derived_alleles.csv")
names(df)[1:2] <- c("chr", "pos")

res <- data.frame()
sfs0 <- data.frame(Var1=1:19, value=0)
for(i in 1:nrow(gene)){
   sub <- subset(df, chr %in% gene$seq[i] & pos > gene$start[i] & pos < gene$end[i])
   tem <- as.data.frame(table(sub$da))
   if(nrow(tem) > 0){
      newsfs <- merge(sfs0, tem, by="Var1", all.x=TRUE)
      newsfs[is.na(newsfs$Freq),]$Freq <- 0
      out <- data.frame(win=i, gene=gene$geneid[i], tajimad = TajimaD(sfs=newsfs$Freq))
      res <- res[order(res$tajimad),]</pre>
```