R_Assignment

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2021-03-23

Part I

Data inspection

```
library(tidyverse)
```

Loading the 2 files to be the genotype and pos data frames

```
## -- Attaching packages ----- tidyverse 1.3.0 --
## v ggplot2 3.3.3
                             0.3.4
                    v purrr
## v tibble 3.1.0
                            1.0.5
                   v dplyr
## v tidyr
          1.1.3
                    v stringr 1.4.0
## v readr
         1.4.0
                   v forcats 0.5.1
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                  masks stats::lag()
genotype <- as.data.frame(read.table("fang_et_al_genotypes.txt", sep="\t",header=TRUE))</pre>
pos <- as.data.frame(read.table("snp_position.txt", sep="\t",header=TRUE))</pre>
```

SNP genotypes data

• fang_et_al_genotypes.txt is assigned to be the genotype data frame, which is large dimension data set, so use dim, str, glimpse and etc functions to know the number of rows and columns, column names and their variable types.

```
#str(genotype)
#qlimpse(qenotype)
colnames(genotype)[1:6]
## [1] "Sample_ID" "JG_OTU"
                                 "Group"
                                              "abph1.20"
                                                          "abph1.22"
                                                                       "ae1.3"
genotype[1:6,1:6]
                  JG_OTU Group abph1.20 abph1.22 ae1.3
##
     Sample_ID
## 1
         SL-15 T-aust-1 TRIPS
                                     ?/?
## 2
                                     ?/?
                                               ?/?
         SL-16 T-aust-2 TRIPS
                                                     T/T
## 3
         SL-11 T-brav-1 TRIPS
                                     ?/?
                                               ?/?
                                                     T/T
                                     ?/?
         SL-12 T-brav-2 TRIPS
                                               ?/?
                                                     T/T
## 4
## 5
         SL-18
                  T-cund TRIPS
                                     ?/?
                                               ?/?
                                                     T/T
## 6
          SL-2 T-dact-1 TRIPS
                                     ?/?
                                               ?/?
                                                     T/T
genotype %>%
  group_by(Group) %>%
  count()
## # A tibble: 16 x 2
## # Groups:
                Group [16]
##
      Group
##
      <chr> <int>
##
    1 TRIPS
                22
    2 ZDIPL
##
    3 ZLUXR
                17
##
    4 ZMHUE
                10
   5 ZMMIL
               290
##
   6 ZMMLR
             1256
   7 ZMMMR
               27
##
##
    8 ZMPBA
               900
```

SNP markers information

41

34

75

69

6

7

4

9

- snp_position.txt is assigned to be the pos data frame.
- Using the same function to know the data structure of the pos data frame.
- Replacing the unknown and multiple in Position column to be NA and to know how many numbers of SNP markers, and their maximum and minimum position value in each of chromosome.

```
dim(pos)
```

```
## [1] 983 15
```

9 ZMPIL

10 ZMPJA

11 ZMXCH

12 ZMXCP

13 ZMXIL

14 ZMXNO

15 ZMXNT

16 ZPERR

##

```
sapply(pos, class)[1:6]
##
          SNP_ID
                 cdv_marker_id
                                  Chromosome
                                                  Position
                                                                 alt_pos
##
     "character"
                     "integer"
                                 "character"
                                               "character"
                                                             "character"
## mult_positions
##
     "character"
str(pos)
## 'data.frame':
                  983 obs. of 15 variables:
## $ SNP ID
                   : chr
                              "abph1.20" "abph1.22" "ae1.3" "ae1.4" ...
## $ cdv_marker_id
                              5976 5978 6605 6606 6607 5982 3463 3466 5983 5985 ...
                        : int
                              "2" "2" "5" "5" ...
## $ Chromosome
                        : chr
## $ Position
                        : chr
                              "27403404" "27403892" "167889790" "167889682" ...
                              ...
## $ alt pos
                        : chr
                              ... ... ... ...
##
   $ mult_positions
                        : chr
                              "abph1" "abph1" "ae1" "ae1" ...
##
   $ amplicon
                        : chr
                              "AB042260" "AB042260" "ae1" "ae1" ...
## $ cdv_map_feature.name: chr
                              "abph1" "ae1" "ae1" ...
## $ gene
                        : chr
                              "candidate" "candidate" "candidate" ...
## $ candidate.random
                        : chr
                              8393 8394 8395 8396 8397 8398 8399 8400 8401 8402 ...
## $ Genaissance daa id : int
## $ Sequenom_daa_id
                              10474 10475 10477 10478 10479 10481 10482 10483 10486 10487 ...
                        : int
## $ count_amplicons
                        : int
                              1 0 1 0 0 1 1 0 1 0 ...
##
   $ count_cmf
                        : int 1010010010...
   $ count gene
                        : int 1010011010...
glimpse(pos)
## Rows: 983
## Columns: 15
## $ SNP ID
                        <chr> "abph1.20", "abph1.22", "ae1.3", "ae1.4", "ae1.5"~
## $ cdv_marker_id
                        <int> 5976, 5978, 6605, 6606, 6607, 5982, 3463, 3466, 5~
                        <chr> "2", "2", "5", "5", "5", "1", "3", "3", "4", "4",~
## $ Chromosome
## $ Position
                        <chr> "27403404", "27403892", "167889790", "167889682",~
                        ## $ alt pos
                        ## $ mult_positions
                        <chr> "abph1", "abph1", "ae1", "ae1", "ae1", "ae1", "ba~
## $ amplicon
## $ cdv_map_feature.name <chr> "AB042260", "AB042260", "ae1", "ae1", "ae1", "an1~
## $ gene
                        <chr> "abph1", "abph1", "ae1", "ae1", "ae1", "an1", "ba~
                        <chr> "candidate", "candidate", "candidate", "candidate"
## $ candidate.random
## $ Genaissance_daa_id
                        <int> 8393, 8394, 8395, 8396, 8397, 8398, 8399, 8400, 8~
## $ Sequenom_daa_id
                        <int> 10474, 10475, 10477, 10478, 10479, 10481, 10482, ~
                        <int> 1, 0, 1, 0, 0, 1, 1, 0, 1, 0, 0, 1, 0, 1, 1, 0, 1~
## $ count_amplicons
## $ count_cmf
                        <int> 1, 0, 1, 0, 0, 1, 0, 0, 1, 0, 0, 1, 0, 1, 1, 0, 1~
## $ count gene
                        <int> 1, 0, 1, 0, 0, 1, 1, 0, 1, 0, 0, 1, 0, 1, 1, 0, 1~
colnames(pos)[1:6]
## [1] "SNP_ID"
                      "cdv_marker_id" "Chromosome"
                                                      "Position"
```

"mult_positions"

[5] "alt_pos"

```
pos[1:6,1:6]
##
       SNP_ID cdv_marker_id Chromosome Position alt_pos mult_positions
## 1 abph1.20
                        5976
                                      2 27403404
## 2 abph1.22
                        5978
                                       2 27403892
## 3
                        6605
                                       5 167889790
        ae1.3
## 4
        ae1.4
                        6606
                                       5 167889682
## 5
                        6607
        ae1.5
                                       5 167889821
## 6
        an1.4
                        5982
                                       1 240498509
pos[pos == "unknown"] <- NA</pre>
pos[pos == "multiple"] <- NA</pre>
pos %>%
  group_by(Chromosome) %>%
  summarise(Max=max(Position, na.rm = T), Min=min(Position, na.rm = T), Number=length(Position))
## # A tibble: 11 x 4
##
      Chromosome Max
                             Min
                                          Number
##
      <chr>
                 <chr>
                             <chr>
                                           <int>
##
   1 1
                  "95897171" "10069039"
                                             155
   2 10
                  "96216463" "10432605"
##
                                              53
   3 2
                  "69623323" "10429605"
                                             127
##
                  "95541392" "106631676"
## 4 3
                                             107
                  "78946482" "103665461"
## 5 4
                                              91
                  "945545"
## 6 5
                             "100227859"
                                             122
##
   7 6
                  "98507715" "113705211"
                                              76
                  "43948320" "104898448"
## 8 7
                                              97
                  "83913342" "115257234"
## 98
                                              62
                  "94285743" "104237516"
## 10 9
                                              60
                  11 11
                             11 11
## 11 <NA>
                                              33
```

Data processing

Subset the pos data frame to keep the SNP_ID, Chr, and Pos columns for the merging purpose

• Adjust the variable type of chromosome and Position to be numeric and remove the unknown and multiple which are regarded as NAs out to be posred dataframe.

\$ Position : num 2.74e+07 2.74e+07 1.68e+08 1.68e+08 1.68e+08 ...

Subset the genotype data into maize and teosinte datasets

• Subset the maize and teosinte genotypes by the Group column. Either [which()] or %>% filter() works out.

• Another method.

```
maize <- genotype %>% filter(Group=="ZMMIL" | Group =="ZMMLR" | Group == "ZMMMR")
teosinte <-genotype %>% filter(Group=="ZMPBA" | Group == "ZMPIL" | Group == "ZMPJA")
```

Formatting the maize genotype with SNP information by merging the posred and maize data

• Transform the maize data for merging with posred by SNP_ID column, and descend the Chromosome and Position.

```
maize \leftarrow maize[,c(-2,-3)]
maize[1:6,1:6] ## have a look
##
        Sample_ID abph1.20 abph1.22 ae1.3 ae1.4 ae1.5
## 1210 ZDP_0752a
                        C/G
                                  A/A
                                        T/T
                                               G/G
                                                      C/C
## 1211 ZDP_0793a
                        C/G
                                  A/A T/T
                                               G/G
                                                      C/T
                        C/C
## 1212 ZDP_0612a
                                  A/A T/T
                                               G/G
                                                      C/C
## 1213 ZDP_0602a
                        C/G
                                  A/A G/T
                                               A/G
                                                      C/T
## 1214 ZDP 0581a
                        C/C
                                  A/A
                                        T/T
                                               G/G
                                                      C/T
## 1215 ZDP_0552a
                                       T/T
                        C/G
                                  A/A
                                               G/G
                                                    C/T
maize <- t(maize)</pre>
maize <- cbind(rownames(maize), maize)</pre>
rownames(maize) <- NULL</pre>
colnames(maize) <- maize[1,]</pre>
maize <- maize[-1,]</pre>
maize <- as.data.frame(maize)</pre>
colnames(maize)[1] <- "SNP_ID"</pre>
maizewp <- merge(posred, maize, by = "SNP_ID")</pre>
maizewp <- maizewp %>% arrange(Chromosome, Position)
## maize genotypes with SNP position information
```

Formatting the teosinte genotype with SNP information by merging the posred and teosinte data

• The same methods as with maize for merging data frame.

```
teosinte <- teosinte[,c(-2,-3)]
teosinte <- t(teosinte)
teosinte <- cbind(rownames(teosinte),teosinte)
rownames(teosinte) <- NULL
colnames(teosinte) <- teosinte[1,]</pre>
```

```
teosinte <- teosinte[-1,]
teosinte <- as.data.frame(teosinte)
colnames(teosinte)[1] <- "SNP_ID"
teosintewp <- merge(posred, teosinte, by = "SNP_ID")
teosintewp <- teosintewp %>% arrange(Chromosome, Position)
```

Splitting the maize data into different files by the chromosomes and SNP positions.

• The followings are using loop to separate the maizewp and teosintewp data frames to 10, 10, 10, and 10 files, respectively, by the chromosome and SNP positions in total 40 files. Also, change the missing genotype to be ? or -.

```
chr <- 1:10
for (i in chr) {
   files_inc <- maizewp[maizewp$Chromosome == i,]
   files_inc[files_inc == "?/?"] <- "?"
   if (i < 10) { write.table(files_inc, file = paste("Maize_Chr0",i,"_increase.txt",sep=""),row.names = i
   else {write.table(files_inc, file = paste("Maize_Chr",i,"_increase.txt",sep=""),row.names = FALSE, se
   files_dec <- maizewp[maizewp$Chromosome == i,]
   files_dec[files_dec == "?/?"] <- "-"
   files_dec <- files_dec %>% arrange(desc(Chromosome),desc(Position))
   if (i < 10) { write.table(files_dec, file = paste("Maize_Chr0",i,"_decrease.txt",sep=""),row.names = i
   else {write.table(files_dec, file = paste("Maize_Chr",i,"_decrease.txt",sep=""),row.names = FALSE, see
}</pre>
```

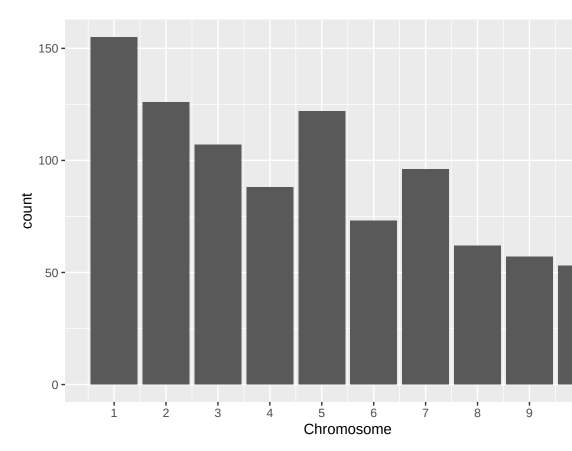
```
chr <- 1:10
for (i in chr) {
  files_inc <- teosintewp[teosintewp$Chromosome == i,]
  files_inc[files_inc == "?/?"] <- "?"
  if (i < 10) { write.table(files_inc, file = paste("Teosinte_Chr0",i,"_increase.txt",sep=""),row.names
  else {write.table(files_inc, file = paste("Teosinte_Chr",i,"_increase.txt",sep=""),row.names = FALSE,
  files_dec <- teosintewp[teosintewp$Chromosome == i,]
  files_dec[files_dec == "?/?"] <- "-"
  files_dec <- files_dec %>% arrange(desc(Chromosome),desc(Position))
  if (i < 10) { write.table(files_dec, file = paste("Teosinte_Chr0",i,"_decrease.txt",sep=""),row.names
  else {write.table(files_dec, file = paste("Teosinte_Chr0",i,"_decrease.txt",sep=""),row.names = FALSE,
}</pre>
```

Splitting the teosinte data into different files by chromosomes and SNP positions.

Part II

Plotting

```
pos %>%
  dplyr::select(SNP_ID, Chromosome, Position) %>%
  drop_na() %>%
  mutate(Chromosome=as.double(Chromosome)) %>%
  ggplot()+
  geom_bar(aes(x=Chromosome))+
  scale_x_continuous(breaks = 1:10)
```



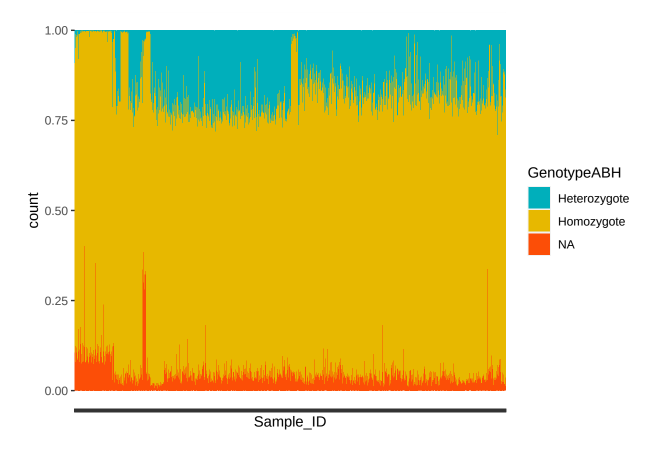
SNPs per chromosome

Missing data and amount of heterozygosity

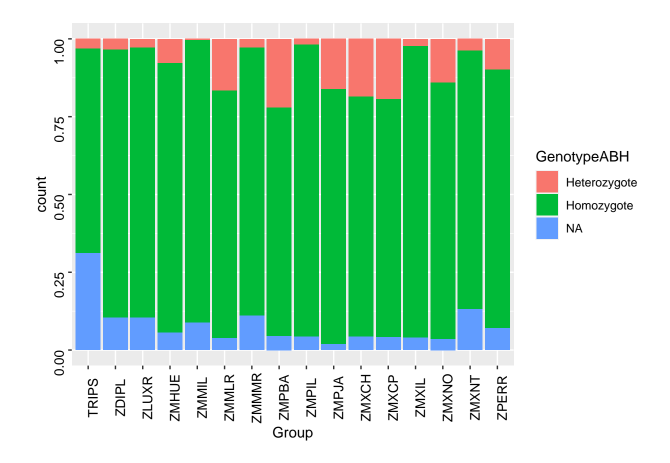
• This chunk is the first version code for substituting the genotypes.

```
## create a function to detect the SNP genotypes
ABH <- function(x) {
   if ( x == "A/A" | x == "C/C" | x == "G/G" | x == "T/T") {
      return("A|B")
   }
   else if (x == "?/?") {
      return("NA")
   }
   else {return("H")}
}
ABH_V <- Vectorize(ABH) ## make the function be a vectorized function</pre>
```

```
genotype3 <- genotype2 %>%
  pivot_longer(3:last_col(), names_to = "SNP", values_to = "Genotype") %>%
 mutate( GenotypeABH = ABH_V(Genotype))
color_Palette = c("#00AFBB", "#E7B800", "#FC4E07")
genotype2 <- genotype[,-2]</pre>
genotype2[1:6,1:6]
##
     Sample_ID Group abph1.20 abph1.22 ae1.3 ae1.4
## 1
         SL-15 TRIPS
                          ?/?
                                   ?/?
                                         T/T
## 2
         SL-16 TRIPS
                          ?/?
                                   ?/?
                                         T/T
                                               ?/?
## 3
         SL-11 TRIPS
                          ?/?
                                   ?/?
                                        T/T
                                               G/G
## 4
         SL-12 TRIPS
                          ?/?
                                   ?/?
                                         T/T
                                               G/G
## 5
         SL-18 TRIPS
                          ?/?
                                   ?/?
                                         T/T
                                               G/G
## 6
         SL-2 TRIPS
                          ?/?
                                   ?/?
                                         T/T
                                               G/G
genotype3 <- genotype2 %>%
  pivot_longer(3:last_col(), names_to = "SNP", values_to = "Genotype") %>%
  mutate( GenotypeABH = ifelse(Genotype%in% c("C/C", "G/G", "A/A", "T/T"), "Homozygote", ifelse(Genotype
ggplot(genotype3)+
  geom_bar(aes(x=Sample_ID, fill=GenotypeABH), position = "fill", width=1)+
  scale_x_discrete(labels=NULL)+
  scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07")) ## change the color palette
```



```
ggplot(genotype3)+
  geom_bar(aes(x=Group, fill=GenotypeABH), position = "fill")+
  theme(axis.text = element_text( angle =90, color="black", size=10, face=1))
```



```
sample_size = posred %>% group_by(Chromosome) %>% summarize(num=n())
library(viridis)
```

The distribution of SNP maker postions on each of chromosomes

Loading required package: viridisLite

```
posred %>%
  left_join(sample_size) %>%
  mutate(myaxis = paste0(Chromosome, "\n", "n=", num)) %>%
  ggplot( aes(x=myaxis, y=Position, fill=as.character(Chromosome)))+
  geom_violin(width=1.4) +
  geom_boxplot(width=0.1, color="grey", alpha=0.2) +
  scale_fill_viridis(discrete = TRUE) +
    theme(
        legend.position="none",
        plot.title = element_text(size=11)
        ) +
        ggtitle("The distribution of SNP postion on each chromosomes") +
        xlab("Chromosome")
```

Joining, by = "Chromosome"

The distribution of SNP postion on each chromosomes

