Pooled Analysis - supplementary

Aimee Taylor and James Watson

Preamble

Load R packages, functions and data.

Data processing

```
# First we remove MS data for which there are no recurrent data
N_episodes_typed = table(MS_pooled$ID[!duplicated(MS_pooled$Episode_Identifier)])
MS_pooled = filter(MS_pooled, ID %in% names(N_episodes_typed[N_episodes_typed>1]))
# recreate pooled summary dataset
MS_pooled_summary = MS_pooled[!duplicated(MS_pooled$Episode_Identifier),]
writeLines(sprintf('Number of individuals with at least two typed episodes analysed: %s',
                   length(unique(MS_pooled$ID))))
## Number of individuals with at least two typed episodes analysed: 212
writeLines(sprintf('Number of episodes in individuals with at least two typed episodes analysed: %s',
                   length(unique(MS_pooled$Episode_Identifier))))
## Number of episodes in individuals with at least two typed episodes analysed: 705
writeLines(sprintf('Number of typed recurrences analysed: %s',
                   length(unique(MS_pooled$Episode_Identifier[MS_pooled$Episode>1]))))
## Number of typed recurrences analysed: 493
First we define the set of microsatellite markers used in this analysis:
MSs_all = c("PV.3.502", "PV.3.27", "PV.ms8",
            "PV.1.501", "PV.ms1", "PV.ms5",
            "PV.ms6", "PV.ms7", "PV.ms16")
```

Number of episodes used to compute frequencies: 212

Run model on easy individuals

Extra computations for VHX: complex individuals

We remove the IDs that can be straightforwardly calculated:

```
ind_calculated = which(MS_pooled_summary$Episode_Identifier %in% Thetas_full_post_TAgnostic$Episode_Identifier %in% Thetas_full_post_TAgnostic$Episode_Ide
```

We use a multinomial-dirichlet model with subjective weight ω . $\omega = 0$ recovers the empirical allele frequencies.

We blow up the pooled analysis into all pairs within individuals:

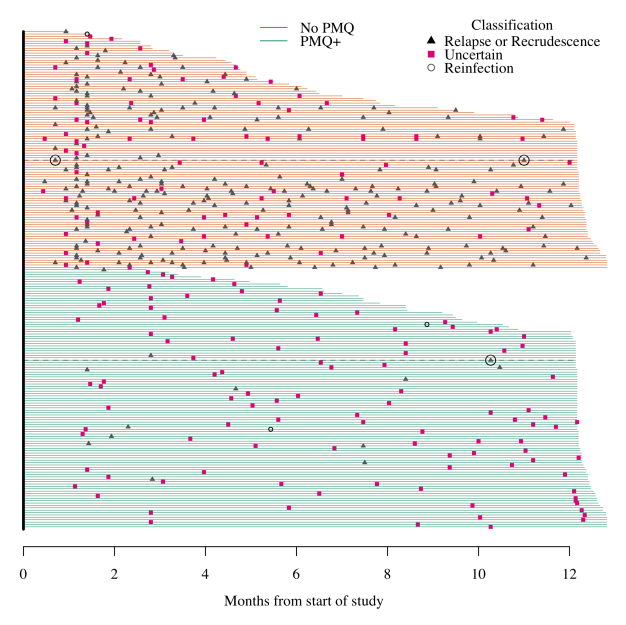
Construct adjacency graphs and compute probabilities of relapse and reinfection.

```
MS_pooled_summary$L_or_C_state = MS_pooled_summary$TotalEpisodes = NA
MS_pooled_summary$L_lower = MS_pooled_summary$L_upper = MS_pooled_summary$L_median = NA
MS_pooled_summary$C_lower = MS_pooled_summary$C_upper = MS_pooled_summary$C_median = NA
MS_pooled_summary$I_lower = MS_pooled_summary$I_upper = MS_pooled_summary$I_median = NA
# Arrange by complexity
# Get single rows per episode (throw away the extra MOI information)
MS inflated summary = MS inflated[!duplicated(MS inflated$Episode Identifier) &
                                  MS_inflated$Episode==2,]
Results_Inflated_TAgnostic$Episode_Identifier = as.character(Results_Inflated_TAgnostic$Episode_Identif
for(i in 1:nrow(MS_inflated_summary)){
 if(!length(which(MS_inflated_summary$Episode_Identifier[i] ==
                  Results_Inflated_TAgnostic$Episode_Identifier))>0){
   MS_inflated_summary = MS_inflated_summary[-i,]
   print('removing')
}
## [1] "removing"
## [1] "removing"
Results_Inflated_TAgnostic$ID_True = NA
Results_Inflated_TAgnostic$First_EpNumber = NA
Results_Inflated_TAgnostic$Second_EpNumber = NA
# The ordering has changed so need to be careful about naming
for(i in 1:nrow(Results_Inflated_TAgnostic)){
 Results_Inflated_TAgnostic$ID_True[i] =
   MS_inflated_summary$ID_True[ind_MS_inflated]
 Results_Inflated_TAgnostic$First_EpNumber[i] =
   MS_inflated_summary$First_EpNumber[ind_MS_inflated]
 Results_Inflated_TAgnostic$Second_EpNumber[i] =
   MS_inflated_summary $Second_EpNumber[ind_MS_inflated]
}
# Iterate through the ones we can calculate in one go
episodes_full_model = unique(Thetas_full_post_TAgnostic$Episode_Identifier)
cols remove = grep('Episode_Identifier', colnames(Thetas_full_post_TAgnostic))
Thetas_full_post_TAgnostic = Thetas_full_post_TAgnostic[, -cols_remove]
for(ep in episodes full model){
 ind1 = (MS_pooled_summary$Episode_Identifier==ep)
 ind2 = rownames(Thetas_full_post_TAgnostic)==ep
 ## Summaries for relapse
 L_cols = grep('L',colnames(Thetas_full_post_TAgnostic))
 MS_pooled_summary$L_upper[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,L_cols]),
                                           probs=upperCI, na.rm = T)
 MS_pooled_summary$L_lower[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,L_cols]),
                                           probs=lowerCI, na.rm = T)
 MS_pooled_summary$L_median[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,L_cols]),
                                            probs=0.5, na.rm = T)
```

```
## Summaries for recrudescence
 C_cols = grep('C',colnames(Thetas_full_post_TAgnostic))
 MS_pooled_summary$C_upper[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,C_cols]),
                                            probs=upperCI, na.rm = T)
 MS_pooled_summary$C_lower[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,C_cols]),
                                            probs=lowerCI, na.rm = T)
 MS_pooled_summary$C_median[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,C_cols]),
                                             probs=0.5, na.rm = T)
 ## Summaries for reinfection
 I_cols = grep('I',colnames(Thetas_full_post_TAgnostic))
 MS_pooled_summary$I_upper[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,I_cols]),
                                            probs=upperCI, na.rm = T)
 MS_pooled_summary$I_lower[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,I_cols]),
                                            probs=lowerCI, na.rm = T)
 MS_pooled_summary$I_median[ind1] = quantile(unlist(Thetas_full_post_TAgnostic[ind2,I_cols]),
                                             probs=0.5, na.rm = T)
 # Just going to classify on relapse versus reinfection
 if(!is.na(MS pooled summary$L upper[ind1])){
   if (MS pooled summary $L upper[ind1] + MS pooled summary $C upper[ind1] < Epsilon lower) {
     MS_pooled_summary$L_or_C_state[ind1] = 'I'
   } else if(MS_pooled_summary$L_lower[ind1]+MS_pooled_summary$C_lower[ind1] > Epsilon_upper){
     MS_pooled_summary$L_or_C_state[ind1] = 'L'
     MS_pooled_summary$L_or_C_state[ind1] = 'Uncertain'
 } else {
   MS_pooled_summary$L_or_C_state[ind1] = NA
# Now iterate through the complex ones
for(i in 1:length(IDs_remaining)){
 id = IDs_remaining[i]
 Doubles_Thetas = filter(Results_Inflated_TAgnostic, ID_True==id)
 for(ep in unique(Doubles Thetas$Second EpNumber)){
    # indices on the MS pooled summary
   ind1 = which(MS pooled summary$ID==id & MS pooled summary$Episode==ep)
    # indices on DOubles thetas: looking for relapse evidence
   ind2 = which(Doubles Thetas$Second EpNumber == ep)
    # index for recrudescence evidence
   ind3 = which(Doubles Thetas$Second EpNumber == ep &
                  Doubles_Thetas$First_EpNumber == (ep-1))
   best_match_relapse = which.max(Doubles_Thetas$L_median[ind2])
   if(length(best_match_relapse)>0){
     MS_pooled_summary$L_lower[ind1] = Doubles_Thetas$L_min[ind2[best_match_relapse]]
     MS_pooled_summary$L_upper[ind1] = Doubles_Thetas$L_max[ind2[best_match_relapse]]
     MS_pooled_summary$L_median[ind1] = Doubles_Thetas$L_median[ind2[best_match_relapse]]
     if(length(ind3)>0){
```

```
MS_pooled_summary$C_lower[ind1] = Doubles_Thetas$C_min[ind3]
        MS_pooled_summary$C_upper[ind1] = Doubles_Thetas$C_max[ind3]
        MS_pooled_summary$C_median[ind1] = Doubles_Thetas$C_median[ind3]
      MS_pooled_summary$I_lower[ind1] = Doubles_Thetas$I_min[ind2[best_match_relapse]]
      MS_pooled_summary$I_upper[ind1] = Doubles_Thetas$I_max[ind2[best_match_relapse]]
      MS_pooled_summary$I_median[ind1] = Doubles_Thetas$I_median[ind2[best_match_relapse]]
    if(!is.na(MS_pooled_summary$C_median[ind1])){
      if(MS_pooled_summary$L_upper[ind1] < MS_pooled_summary$L_lower[ind1]){</pre>
        writeLines(sprintf('Problem with ID %s',id))
        stop()
      }
      if(MS_pooled_summary$L_upper[ind1]+MS_pooled_summary$C_upper[ind1] < Epsilon_lower){</pre>
        MS_pooled_summary$L_or_C_state[ind1] = 'I'
      } else if(MS_pooled_summary$L_lower[ind1]+MS_pooled_summary$C_lower[ind1] > Epsilon_upper){
        MS_pooled_summary$L_or_C_state[ind1] = 'L'
      } else {
        MS_pooled_summary$L_or_C_state[ind1] = 'Uncertain'
      }
    }
 }
}
MS pooled summary$Drug = MS pooled summary$FU = NA
for(id in MS_pooled_summary$ID){
  ind = MS pooled summary$ID==id
  MS_pooled_summary$TotalEpisodes[ind] = max(MS_pooled_summary$Episode[ind])
  MS_pooled_summary$Drug[ind] = as.numeric(
    Combined_Time_Data$arm_num[Combined_Time_Data$patientid==id][1] == 'CHQ/PMQ') + 2
  MS_pooled_summary$FU[ind] = Combined_Time_Data$FU_time[Combined_Time_Data$patientid==id][1]
}
MS_pooled_summary $Plotting_pch_Values =
  as.numeric(mapvalues(MS_pooled_summary$L_or_C_state, from = c('L', 'Uncertain', 'I'), to = c(17,15,1)))
MS_pooled_summary$Plotting_col_Values =
  as.numeric(mapvalues(MS_pooled_summary$L_or_C_state, from = c('L', 'Uncertain', 'I'), to = 1:3))
```

Coatney plot (genetic data informed only)



The Coatney style plot is showing 488 recurrences in 209 individuals Break down the results depending on whether PMQ was received or not:

There are 121 recurrences after PMQ+. The breakdown as % of classification is as follows:

##
I L Uncertain
2 11 88

 $\mbox{\tt \#\#}$ There are 367 recurrences after no PMQ. The breakdown as % of classification is as follows:

##

##

I L Uncertain ## 0 75 24

8.211 sec elapsed