Exploration of Shared SNPs in Thaps Chr1-unfiltered

June 29, 2017

Rambling exploration of SNP positions shared between two or more of the isolates. Code is included to document it thoroughly, (even if largely uninteresting to anyone else), and I will summarize it as I go.

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1 HISTORY 2

1 History

This was added to SVN 1/26/2014; not sure when it was started, but earliest related emails I see are from 1/21/14.

```
r413 | ruzzo | 2014-01-26 08:22:37 -0800 (Sun, 26 Jan 2014) | 2 lines adding shared-snp analysis.
```

2 Preliminaries

NOTE: Some comments in code and some parts of the text, especially specific numbers and general conclusions, are based on Unqfiltered, Chr1, Medium stringency (i.e., "[[2]]" below) analysis. The broad picture does not appear to change with other choices, but details do, and the text is neither fully parameterized nor fully updated, so proceed with caution.

Load utility R code; do setup:

```
source('../../R/wlr.R') # load util code; path relative this folder or sibling in scripts/larrys
## Running as: ruzzo @ bicycle.cs.washington.edu; SVN Id, I miss you. $Id: wlr.R 2017-06-26 or later $
setup.my.wd('shared-snps') # set working dir; UPDATE if this file moves, or if COPY/PASTE to new file
setup.my.knitr('f-knitr/')
generic.setup('figs-mine/')
```

3 Major Analysis/Performance Parameters.

Choices here control how this file is processed, what data is analyzed, speed, etc. Set them carefully before running "make." Major choices are:

- 1. WHICH SNP TABLES ARE LOADED??? The logical vector load.tb selects the desired combination of SNP tables to load, in the order full.unfiltered, chrl.unfiltered, full.qfiltered, chrl.qfiltered. E.g., load.tb=(T, F, T, F) loads *full* tables for *both* q- and un-qfiltered data. Primary analysis is only performed on one of them, but the others are retained for comparison/debugging.
- 2. WHICH MAIN ANALYSIS??? If multiple tables are loaded, which is used for the main analysis? Parameter pri is a permutation of 1:4, corresponding to load.tb; the first loaded table in that order becomes the analysis focus. The default pri=c(1,2,3,4) looks at un-q-filtered data in preference to q-filtered, and full tables in preference to Chr1 within each group.
 - (Choice of data for the "Table 1" coverage summary in section 5 is independent of this; full genome data is prefered over Chr 1 for both q- and unq-filtered reads; change tset.picker calls near the end of that section to modify this.)
- 3. CLEAR CACHE??? clear.cache=T forces Knitr cache removal at the start of the run; especially important if the previous parameters have changed since the last run.
- 4. HOW MANY BOOTSTRAP REPLICATES??? The variable nboot is a major performance factor; 1000 reps takes several hours. Set to 5 for debug and quick look; 100 or more for final run.
- 5. TRUNCATE TABLES TO Chrs ONLY??? I.e., remove mitochondrial-, plastid-, and BD- contigs.

The following code chunk sets the first four parameters based on where it's run. To prototype/debug on a laptop, faster is better—run on Chr1 with small nboot; when run on the linux servers, I typically do full genomes, more replicates. Just override them if these defaults don't work for you.

```
# for Makefile, params can be command line args, else base on system; see wlr.r for details.
# load.tb order: full.un, chr1.un, full.qfil, chr1.qfil
params <- pick.params(</pre>
 mac = list(load.tb=c(F,T,F,F), pri=1:4, clear.cache=F, nboot= 1, trunc.tables=T), # quick on lap
#linux = list(load.tb=c(F,F,F,T), pri=1:4, clear.cache=F, nboot= 5, trunc.tables=T), # quick qfil on server
 linux = list(load.tb=c(T,F,T,F), pri=1:4, clear.cache=T, nboot=101, trunc.tables=T) # full on server
# Alternatively, edit/uncomment the following to override the above as needed
#params<-pick.params(default=list(load.tb=c(T,T,T,T)),pri=1:4,clear.cache=T,nboot=1000,trunc.tables=T))</pre>
print (params)
# $load.tb
# full.unf chr1.unf full.qf chr1.qf
   FALSE TRUE FALSE TRUE
# $pri
# [1] 1 2 3 4
# $clear.cache
# [1] TRUE
# $nboot
# [1] 5
# $trunc.tables
# [1] FALSE
```

CLEAR CACHE??!! Some code chunks use the knitr cache, but extent of cache consistency checks unknown. If in doubt, delete "cache/" (knitr's) directory to force rebuild. T/F set in params above will/won't force removal (actually, rename):

```
decache (params$clear.cache)
# Rename of 'cache' to 'cache81523' returned TRUE .
```

If still in doubt, also manually remove "00common/mycache/" (mine). Load the main SNP data file(s) based on the parameters set in section 3.

```
# short names to keep the following chunk compact
tb <- params$load.tb
tset <- list(NULL, NULL, NULL, NULL) # tset = 'table set'</pre>
```

```
# see wlr.R for load paths
if(tb[1]) {tset[[1]] <- load.snp.tables(use.chr1.tables = FALSE, data.name='full.tables.01.26.14')}
if(tb[2]) {tset[[2]] <- load.snp.tables(use.chr1.tables = TRUE, data.name='full.tables.01.26.14')}
# Loading ../00common/mycache/snp.tables.chr1.unqfiltered.rda ...Loaded.

if(tb[3]) {tset[[3]] <- load.snp.tables(use.chr1.tables = FALSE, data.name='full.tables.02.25.15')}
if(tb[4]) {tset[[4]] <- load.snp.tables(use.chr1.tables = TRUE, data.name='full.tables.02.25.15')}
# Loading ../00common/mycache/snp.tables.chr1.qfiltered.rda ...Loaded.
# Bandaiding qfiltered tables...</pre>
```

Grrr! I should have excluded non-Chr contigs from full genome runs. Rather than change tons of code below to add mask params, I'm just going to truncate the tables, as follows. (See notes in wlr.r::make.mask for assumptions.)

```
if(params$trunc.tables) {
  for(i in 1:4) {
    if(!is.null(tset[[i]])) {
      first.mito <- match("mitochondria.fasta", tset[[i]][[7]]$Chr)
    if(!is.na(first.mito)) { # will be NA for Chrl tables</pre>
```

```
for(j in 1:7){
    # hmmm... slow; wonder whether head(tset[[i]][[j]], first.mito-1) is faster;
    # ok, simple tests suggest not: system.time(head(data.frame(1:1e7,1:1e7),5e6))
    tset[[i]][[j]] <- tset[[i]][[j]][1:(first.mito-1),]
    }
}
}
else {
    cat('***\n*** DID YOU *REALLY* WANT UNTRUNCATED TABLES???\n***\n')
}

# ***
# *** DID YOU *REALLY* WANT UNTRUNCATED TABLES???
# ***</pre>
```

The tersely-named tset list is sometimes convenient, but give them more descriptive names, too.

```
snp.tables.full.unfiltered <- tset[[1]]; names(tset)[1] <- 'snp.tables.full.unfiltered'
snp.tables.chr1.unfiltered <- tset[[2]]; names(tset)[2] <- 'snp.tables.chr1.unfiltered'
snp.tables.full.qfiltered <- tset[[3]]; names(tset)[3] <- 'snp.tables.full.qfiltered'
snp.tables.chr1.qfiltered <- tset[[4]]; names(tset)[4] <- 'snp.tables.chr1.qfiltered'</pre>
```

Main analysis may just use one of the potentially 4 table sets. Pick it according to the priority specified in section 3, using the shorter name 'snp.tables' for this default choice.

```
snp.tables <- tset.picker(priority=params$pri, table.set=tset)</pre>
```

```
# Sanity check: unlike unqfiltered tables, bug in early code gave qfiltered ones different numbers
# of rows per strain, which breaks much code. Verify this is no longer happening.
check.eq.nrows <- function(tables){</pre>
 if(!is.null(tables)){
   nrow.snp.tables <- unlist(lapply(tables,nrow))</pre>
   print (nrow.snp.tables)
   if(all(nrow.snp.tables == nrow.snp.tables[1])){
     cat('OK, all strains have same number of rows.\n')
     cat('***\n*** Warning: Different strains have different numbers of rows! ***\n***\n')
dummy<-lapply(tset, check.eq.nrows)</pre>
                           1014
                                 1015
            1012
                    1013
                                          3367 1335
# 3042585 3042585 3042585 3042585 3042585 3042585
# OK, all strains have same number of rows.
   1007 1012 1013 1014 1015 3367 1335
# 3042585 3042585 3042585 3042585 3042585 3042585
# OK, all strains have same number of rows.
```

Which tables have we got?:

```
# 'which.snp.tables' return summary of which tables, either as a char string (default), e.g.
# "Chrl-qfiltered", or as vector of 2 strings, e.g. c("full", "unfiltered").
cat('This analysis uses: (', paste(unlist(lapply(tset,which.snp.tables)),collapse=', '), ') SNP tables.\n')
# This analysis uses: ( NULL, Chrl-unfiltered, NULL, Chrl-qfiltered ) SNP tables.
cat('Main shared SNP analysis focuses on', which.snp.tables(snp.tables), '\n')
# Main shared SNP analysis focuses on Chrl-unfiltered
```

A LATEX hack: I want which snp.tables info in doc title/page headers, but it is unknown until now, so the following writes a command definition \whichsnptables into the .aux file, which is read during the next LATEX run, when \begin{document} is processed:

```
\makeatletter
\immediate\write\@auxout{\noexpand\gdef\noexpand\whichsnptables{Chr1-unfiltered}}
\makeatother
```

Subsequent analysis was initially all directed at Chr1. In general, I have *not* updated the discussion to reflect genome-wide analysis.

```
if(exists('snp.tables.chr1.qfiltered') && exists('snp.tables.chr1.unqfiltered')){
  # If have both, where is new unequal to old?
  uneq <- snp.tables.chr1.qfiltered[[1]]$Ref[1:chr1.len] != snp.tables.chr1.unqfiltered[[1]]$Ref[1:chr1.len]
  cat('Sum uneq:', sum(uneq,na.rm=T), '\n')
  cat('Sum NA: ', sum(is.na(uneq)), '\n')
  print(which(is.na(uneq))[1:10])
  seecounts(which(is.na(uneq))[1:4],snp.tables=snp.tables.qfiltered,debug=F)
}</pre>
```

In brief, "snp.tables" will be a list of 7 data frames, one per strain, giving read counts for each nucleotide at each position, SNP calls, etc.:

```
names (snp.tables)
# [1] "1007" "1012" "1013" "1014" "1015" "3367" "1335"
str(snp.tables[[1]])
# 'data.frame': 3042585 obs. of 15 variables:
# $ chr : Factor w/ 66 levels "BD10_65", "BD11_74",..: 39 39 39 39 39 39 39 39 39 39 ...
          : int 1 2 3 4 5 6 7 8 9 10 ...
  $ pos
         : int 0000000000...
  $ snp
 $ Chr : chr "Chr1" "Chr1" "Chr1" "Chr1" ...
 $ Pos : int 1 2 3 4 5 6 7 8 9 10 ...
                 "T" "C" "C" "A" ...
  $ Ref : chr
         : num
                 1 3 4 5 7 7 10 12 13 15 ...
  $ Cov
          : num 0 0 1 0 0 0 0 0 1 0 ...
  Ŝа
 $ q
         : num 0 0 0 0 0 0 0 0 0 0 ...
  $ C
         : num 0 0 0 0 0 0 0 0 0 ...
  $ t
         : num 0 0 0 0 0 0 0 0 0 0 0 ...
: num 0 0 0 0 0 0 0 0 0 0 ...
  $ .match: num 1 3 3 5 7 7 10 12 12 15 ...
 $ exon : logi FALSE FALSE FALSE FALSE FALSE ...
# $ indel : logi FALSE FALSE FALSE FALSE FALSE ...
```

Just for background, also load the desert tables:

```
# from svn+ssh://ceg1.ocean.washington.edu/var/svn/7_strains/trunk/code/snpNB/data
#load('../../data/ungit-data/des.rda')
load('../../data/des.rda')
```

What's the total length of all deserts in each strain? Big deserts (defined as "big.threshold" or longer)?

```
some.desert.stats <- function(big.threshold=0){</pre>
 desert.len <- unlist(lapply(des, function(x) {sum(unlist(lapply(x, function(y) {sum(y[, 'Length'])}))))}))</pre>
 bigdes.len <- unlist(lapply(des, function(x) {sum(unlist(lapply(x, function(y)) {
                                                 sum(y[y[,'Length']>=big.threshold,'Length'])})))))
 rbind(desert.len, desert.pct=round( desert.len / genome.length.constants()$genome.length.trunc * 100),
       bigdes.len, bigdes.pct=round( bigdes.len / genome.length.constants() $genome.length.trunc * 100))
some.desert.stats(big.threshold=50000)
              tp1007 tp1012 tp1013 tp1014 tp1015 thapsIT tp1335
# desert.len 11146526 11332566 5801763 9464213 11251426 6780300 10883723
                                                  36
# desert.pct
                36
                          36
                                 19 30
                                                        22
# bigdes.len 3495805 3936973
                               55365 3627235 3727061
                                                        57119 4046934
             11
                          13
                               0
                                      12
```

I.e., looking at all deserts, about 1/3 of L-clade, 1/5 of H-clade are in deserts, whereas, looking at the largest deserts (> 50k), only about 12% in L-clade (and none in H-clade). Note that the rough stats above include artifactual "deserts" created by gaps in the reference sequence, large genomic deletions, etc. A more careful analysis of this is found in nc-snps.rnw.

4 Refined SNP Calls

4.1 Method

It is appropriate that SNP calls should be conservative, to avoid many false positives, but, when a position is called a SNP in one isolate, we often see a significant number of reads for the same non-reference nucleotide at that position in other isolates, even if they are not called as SNPs. On the other hand, we sometimes see a position called a SNP in two or more isolates, but with *different* pairs of nucleotides, potentially suggesting technical errors. Analysis in this section attempts to refine the SNP calls by looking for issues such as these by looking at all 7 isolates jointly, at each position called a SNP in any of them.

For a given strain, the following function returns a vector of 0:4 to indicate which nonreference nucleotide has the maximum read count at the corresponding position. The values 1..4 indicate that the max count occurred at A, G, C, T, resp. (Ties are resolved arbitrarily (a < g < c < t), which possibly deserves further attention.) The value 0 means all nonreference counts are below threshold, based *either* on absolute count *or* as a fraction of coverage. Default only excludes 0 counts.

```
nref.nuc.new <- function(strain=1, mask=T, thresh.count=0, thresh.rate=0.0){
    # get read count for max nonref nuc
    nref <- apply(snp.tables[[strain]][mask, c('a', 'g', 'c', 't')], 1, max)
    # where does nref count match a (g,c,t, resp) count
    as <- ifelse(nref == snp.tables[[strain]][mask,'a'],1,0)
    gs <- ifelse(nref == snp.tables[[strain]][mask,'g'],2,0)
    cs <- ifelse(nref == snp.tables[[strain]][mask,'c'],3,0)
    ts <- ifelse(nref == snp.tables[[strain]][mask,'t'],4,0)
    # most positions will show 3 zeros and one of 1:4, so max identifies max nonref count;
    # ties broken arbitrarily (a<g<c<t)
    merge <- pmax(as,gs,cs,ts)
    # but if max nonref count is zero or below threshold, return 0
    merge[nref == 0 | nref < thresh.count] <- 0
    merge[nref/snp.tables[[strain]][mask,'Cov'] < thresh.rate] <- 0
    return(merge)
}</pre>
```

Get union and intersection of the sets of called SNPs. ("\$snp" is 0/1.) Also, 5-way (L-clade) and 4-way (L-excluding Gyre).

```
# 4-way union/intersection
u4.snps <- snp.tables[[1]]$snp
i4.snps <- snp.tables[[1]]$snp</pre>
for(i in c(2,5,7)) {
        u4.snps <- pmax(u4.snps, snp.tables[[i]]$snp)
        i4.snps <- pmin(i4.snps, snp.tables[[i]]$snp)</pre>
# 5-way: add gyre
u5.snps <- pmax(u4.snps, snp.tables[[4]]$snp)
i5.snps <- pmin(i4.snps, snp.tables[[4]]$snp)
# 7-way
union.snps <- pmax(u5.snps, snp.tables[[3]]$snp, snp.tables[[6]]$snp)
intersect.snps <- pmin(i5.snps, snp.tables[[3]]$snp, snp.tables[[6]]$snp)</pre>
nu4snps <- sum(u4.snps)</pre>
nu5snps <- sum(u5.snps)
ni4snps <- sum(i4.snps)
ni5snps <- sum(i5.snps)</pre>
nusnps <- sum(union.snps)
nisnps <- sum(intersect.snps)</pre>
c(n4u=nu4snps, n5u=nu5snps, n7u=nusnps, n4i=ni4snps, n5i=ni5snps, n7i=nisnps)
```

```
# n4u n5u n7u n4i n5i n7i
# 18564 18696 47499 14365 7628 1641
```

There are nusnps=47499 positions called as SNPs in one or more strains (but only nisnps=1641 that are shared among all 7). Note that the 4-way union is only modestly larger (1.2923077 times larger) than the 4-way intersection, emphasizing the inherent similarities among these SNP sets. The corresponding 5-way numbers show that Gyre adds relatively little to the 5-way union vs the 4-way union, whereas it removes a fair bit from the 5-way intersection. However, much of that loss is simply because Gyre has fewer called SNPs: only 8331 vs 14365 in the 4-way intersection, and they are highly concordant:

```
sum(snp.tables[[4]]$snp*i4.snps)/sum(snp.tables[[4]]$snp)
# [1] 0.9156164
```

So, a likely source of the Gyre's difference in called SNPs is technical (lower read coverage, higher read error rate) rather than biological.

Inclusion of the 2 H-clade members, however, causes more dramatic changes in both union and intersection numbers. I examine all these relationships in more detail below, but first I examine what I believe to be a significant source of technical error in these comparisons—erroneous SNP calls, especially false negative calls.

It is appropriate that SNP calls should be conservative, to avoid many false positives, but, if a position is called a SNP in one strain, we often see a significant number of reads for the same non-reference nucleotide at that position in other strains, even if they are not called as SNPs. For my purposes below, these will be considered "shared SNPs," based on three different levels of permissiveness. Note that, e.g., $\geq 84\%$ of all positions have zero reads for any non-reference nucleotide, and only a small fraction have 2 or more non-reference reads:

```
nonmatch <- rbind(</pre>
  unlist(lapply(snp.tables, function(x) {sum(x$Cov-x$.match == 0)})),
  unlist(lapply(snp.tables, function(x) {sum(x$Cov-x$.match == 1)})),
  unlist(lapply(snp.tables, function(x){sum(x$Cov-x$.match == 2)})),
  unlist(lapply(snp.tables, function(x) {sum(x$Cov-x$.match == 3)})),
  unlist(lapply(snp.tables, function(x) {sum(x$Cov-x$.match >= 4)})),
 unlist(lapply(snp.tables, function(x){sum((x$Cov-x$.match)[union.snps==0] >= 4)}))
)/nrow(snp.tables[[1]]) *100
rownames (nonmatch) <- c('% ==0','% ==1','% ==2','% ==3','% >=4', '% >=4, nonSNP')
nonmatch
                      1007
                                 1012
                                            1013
                                                        1014
                                                                   1015
                                                                               3367
                                                                                          1335
# % ==0
               92.49473721 88.8682485 84.9571992 86.51028648 90.6336881 86.4695974 85.2300922
# % ==1
                6.37162150 9.3939528 12.1458891 11.81896315 7.9041670 10.9489792 11.6810541
# % ==2
                0.41921590 0.8698853
                                       1.3586145 1.09222914 0.6199991
                                                                         1.1349231
# % ==3
                0.07950476 0.1349182 0.2267480 0.17849953 0.1118128 0.2029196
# % >=4
                0.63492063 0.7329951 1.3115492 0.40002169 0.7303329 1.2435807 0.9002214
# % >=4, nonSNP 0.06267697 0.1274903 0.2943878 0.06037629 0.1217715 0.2716440 0.3085534
```

Build a table of max non-reference nucleotides at each position in the union.snps set. The three criteria are

- [[1]]: any non-zero count at any coverage is considered significant
- [[2]]: (count ≥ 2 and count/coverage ≥ 0.05) is considered significant
- [[3]]: (count > 4 and count/coverage > 0.10) is considered significant

In all three cases, the nonref nucleotide must also be consistent across all strains passing that threshold; see below.

```
non.refs[[1]][,j] <- nref.nuc.new(j, mask=union.snps==1, thresh.count=0, thresh.rate=0.00)
non.refs[[2]][,j] <- nref.nuc.new(j, mask=union.snps==1, thresh.count=2, thresh.rate=0.05)
non.refs[[3]][,j] <- nref.nuc.new(j, mask=union.snps==1, thresh.count=4, thresh.rate=0.10)
}</pre>
```

For comparison, I want to look at unfiltered SAMTools SNP calls. In complete opposition to the measures of consistency imposed above, I'm going to simply force this into the "non.refs" structure constructed above by imagining that any position called a SNP in any strain has its max nonref count on "A", so any given position called a SNP in any strain will automatically be declared "consistent." This will allow the tree-code, etc. given below to work in a uniform way (even though interpretation of the results is different.) Results will be jammed into a 4th component of the "non.refs" list; i.e., we have a 4th criterion:

• [[4]]: all called SNPs at a given position are considered "consistent."

As this case was a late addition to the analysis, the commentary throughout this document has not necessarily been updated to reflect that this case is distinct from the first three.

```
for(j in 1:7){
  non.refs[[4]][,j] <- snp.tables[[j]]$snp[union.snps==1]
}</pre>
```

"non.refs" indicates, among those positions in the union of all called SNPS having any non-reference read count above the thresholds listed above, the non-ref nucleotide having the highest read count in each strain. If, for a given position, the max of this code is the same as the min (among non-zero values), then every strain having over-threshold nonref reads in that position, in fact has most non-reference reads on the *same* nucleotide. These are defined as the "consistent" SNPs.

```
find.consistent <- function(nr) {
  nr.max <- apply(nr,1,max)
  nr.min <- apply(nr,1,function(x) {ifelse(max(x)==0,0,min(x[x>0]))})
  return(nr.min == nr.max)
}
consistent <- lapply(non.refs, find.consistent)</pre>
```

4.2 Save them

```
get.snp.locs.char = function(strain, stringency=2){
           # return char vector of locations of consistent SNPs @ specified stringency & strain
           snps <- filtered.snps$Code$get.snps(strain, stringency)</pre>
           return (names (snps) [snps])
        get.snp.locs.df = function(strain, stringency=2){
           # return data frame (Chr/Pos) of locations of consistent SNPs @ specified stringency & strain
          snplist <- strsplit(filtered.snps$Code$get.snp.locs.char(strain, stringency), ':', fixed=TRUE)</pre>
           # strsplit returns long list of 2-vectors, 1st=chr, 2nd=char position
                                          unlist(lapply(snplist, function(x) {return(x[1])})),
           df <- data.frame(Chr=
                            Pos=as.integer(unlist(lapply(snplist, function(x){return(x[2])}))),
                            stringsAsFactors = FALSE)
           return (df)
       )
# dont't clobber existing .rda, but save if absent. (delete to re-save)
rda.filtered <- paste('../00common/mycache/filtered.snps', which.snp.tables(), 'rda', sep='.')
if(file.exists(rda.filtered)){
 cat('Pre-existing file', rda.filtered, 'unchanged.\n')
} else {
 cat('Saving', rda.filtered, '...')
 save(filtered.snps, file=rda.filtered, compress=TRUE)
 cat('Saved.\n')
# Pre-existing file ../00common/mycache/filtered.snps.Chr1-unfiltered.rda unchanged.
```

Knitr seems to be failing to format the long char string above, which says:

```
cat (filtered.snps$Description)
# Contents of this .rda file:
    * Description: this text
   \star Data -- 5 items defining filtered SNPs, at 4 different stringency levels, as defined
     in shared-snps.rnw:
      * based.on.which.snp.tables: {"Chr1", "full", "trunc"}-{"unfiltered", "qfiltered"},
       depending on which snp tables were used to build this data. ("trunc" = all Chrs.)
      * number.union.snps: the total number of SNPs (SAMtools calls) in the union of SNPs
       across all 7 strains.
      * number.intersection.snps: similar, for the 7-way intersection.
       nusnps/nisnps are easily recalculated from the data below, but their inclusion
       may be convenient, e.g., to quickly see if the .rda represents the full genome
        (nusnps=488848), or the chr 1 subset (nusnps=47499); (redundant with "based.on...";
       numbers above are for unfiltered, perhaps slightly different if qfiltered)
      \star non.ref.nucleotide: 4 arrays, each nusnps x 7, of values 0..4 (0..1 in the 4th
        array). In the 1st 3 arrays, 0 means the given position in the given strain did
       not have nonreference read counts above the corresponding filtering threshold,
       i.e., is NOT a filtered SNP in that strain, whereas 1..4 mean that it did pass
        threshold, for A,C,G,T resp. In the 4th array, this value is just 1/0,
        indicating is/is not a called SNP in that strain.
      * consistent.snps: 4 Bool vectors of length nusnps flagging positions whose nonref
       nucs (wrt to the 4 filtering criteria) are deemed *consistent* across
        all 7 strains. For the 1st 3, this means all nonzero entries of non.ref.nuc
      are equal, i.e., nonref read counts passing threshold are on the SAME nonref
```

```
# nucleotide in all strains having over-threshold counts. Just for comparison
and uniformity of data structures, the 4th is all TRUE, i.e., union of SNPs
# across all strains, without any regard for thresholds or consistency.

# In short, the filtered SNPs according to our medium filtering criteria are
# strains/positions where consistent.snps[[2]]==TRUE and non.ref.nucleotide[[2]]>0.

# Rownames in both non.ref.nucs and consistent define location, e.g. "Chrl:333".

# * Code -- simple routines to extract filtered SNPs in (potentially) convenient formats:

# * get.snps(strain, stringency=2)
# returns nusnps x 1 Bool vector of consistent SNPs @ specified stringency in
# given strain

# * get.snp.locs.char(strain, stringency=2)
# returns n x 1 char vector of locations of consistent SNPs @ specified stringency
# in given strain, e.g. "Chrl:1234", where n == sum(get.snps(...))

# * get.snp.locs.df(strain, stringency=2){
# As above, but returns data frame (char vector Chr, int vector Pos) with the same info.
```

```
str(consistent[[1]])

# Named logi [1:47499] TRUE FALSE TRUE FALSE TRUE TRUE ...
# - attr(*, "names")= chr [1:47499] "Chr1:333" "Chr1:417" "Chr1:435" "Chr1:438" ...
```

```
consistent.count <- unlist(lapply(consistent, sum)); consistent.count
# [1] 36040 46896 47174 47499
inconsistent.count <- consistent.count[4] - consistent.count; inconsistent.count
# [1] 11459 603 325 0
inconsistent.percent <- inconsistent.count/consistent.count[4]*100; inconsistent.percent
# [1] 24.1247184 1.2695004 0.6842249 0.0000000</pre>
```

I.e., of the 47499 positions in which a SNP is called, 36040 are consistent by my loose definition, and 47174 are consistent by my tightest definition. The increase in concordance supports the view that the loose definition is too loose. Perhaps misleadingly, these counts include positions that are "consistent SNPs" in only one strain; more below. (*TODO* I suspect, but have not yet systematically checked, that most of the rest are positions with low coverage and/or very low read counts on the mixture of non-reference nucleotides.)

4.3 Examples: Consistent

Here are a few (nonrandomly selected) prototypical consistent SNPs:

```
esnps <- names(consistent[[2]][consistent[[2]]])</pre>
\texttt{esnps2} < -\texttt{as.integer}(\texttt{unlist}(\texttt{lapply}(\texttt{strsplit}(\texttt{esnps}[\texttt{c}(7,11:13,92)], ':', \texttt{fixed=TRUE}), \texttt{function}(\texttt{x})\{\texttt{x}[2]\})))
seecounts(esnps2, snp.tables=snp.tables)
      chr pos Ref Strain A G C T SNP exon indel nrf rat
#
# 1
     Chr1 567 T
# 2
                       1007 0 0 2 29
                                            O TRUE FALSE
# 3
                       1012 0 0 15 44
                                            1
                                                TRUE FALSE
                       1013 0 0 15 97 0 TRUE FALSE
# 4
# 5
                       1014 0 1 0 33 0 TRUE FALSE
                                             1 TRUE FALSE
                       1015 0
3367 0
# 6
                                 0 9 43
                                 0 16 46
                                             1
                                                 TRUE FALSE
                       1335 0 0 2 116 0 TRUE FALSE
# 8
# 9 Chrl 1053 A
                    1007 39 0 0 5 0 TRUE FALSE
# 10
```

```
# 11
                   1012 55 0 0 12 0 TRUE FALSE
                            1 0 40
# 12
                   1013 17
                                      O TRUE FALSE
                            0 0 0 1
# 13
                   1014 25
                                  5
                                      0
                                         TRUE FALSE
# 14
                   1015 38
                                  20
                                      1
                                         TRUE FALSE
# 15
                   3367 13
                            0 0
                                  9
                                     O TRUE FALSE
# 16
                   1335 71
                           1 0 46
                                     1 TRUE FALSE
# 17 Chrl 1055
# 18
                   1007
                        0
                           41
                              0
                                   2
                                       0
                                         TRUE FALSE
                        1
                           63 0
# 19
                   1012
                                   8
                                      0
                                         TRUE FALSE
# 20
                   1013 1 62 0
                                   8
                                      0
                                         TRUE FALSE
# 21
                   1014 1
                           26 0
                                  8
                                      1
                                         TRUE FALSE
                   1015 0
                                      O TRUE FALSE
# 22
                           44 0
                                  14
                        0
                           27
                               0
                                  0
                                      0
                                         TRUE FALSE
# 23
                   3367
                   1335 0
                           78 0
                                      1 TRUE FALSE
# 24
                                  40
# 25 Chr1 1176 G
# 26
                   1007 2 67 0
                                  0
                                     O FALSE FALSE
                   1012 1
                           68 0
                                   0
                                      O FALSE FALSE
# 27
# 28
                   1013 29
                           73
                               0
                                   0
                                       O FALSE FALSE
# 29
                   1014 1 52
                              0
                                      O FALSE FALSE
                                   0
# 30
                   1015 4 103 0
                                      O FALSE FALSE
                                     1 FALSE FALSE
# 31
                   3367 11 8 0
                                  0
                   1335 1 206 0 0
 32
                                      O FALSE FALSE
# 33 Chr1 8670 A
                   1007 19
                           0 0 7
                                      O TRUE FALSE
# 34
                   1012 36
                           0 0 12
# 35
                                      0 TRUE FALSE
                   1013 44
                            0 0 12
                                      O TRUE FALSE
# 36
                            0
 37
                   1014 10
                               0
                                      0
                                         TRUE FALSE
                                  11
# 38
                   1015 24
                            0
                              0
                                      1
                                         TRUE FALSE
                   3367 18
                            0 0
                                 0
                                     O TRUE FALSE
# 39
                   1335 27 0 0 6 0 TRUE FALSE
```

4.4 Examples: Inconsistent

Here is a brief look at some *in*-consistent positions. E.g., Chr1:2013 shows nontrivial counts on 3 alleles in Wales, as do 2319, 3286, 5002, 5433, whereas 7878 shows a different alternate allele in Italy than in Wales.

```
unc <- names(consistent[[2]][!consistent[[2]]])</pre>
 unc2 <- \textbf{as.integer(unlist(lapply(strsplit(unc[1:10],':',fixed=TRUE),function(x)\{x[2]\})))} 
seecounts(unc2, snp.tables=snp.tables)
           pos Ref Strain A G C T SNP exon indel nrf rat
# 1
    Chr1 2013 T
# 2
                     1007 4
                             0
                                 0 20 0 TRUE FALSE
# 3
                     1012
                          8
                              0
                                  0 34
                                         0
                                            TRUE FALSE
# 4
                     1013
                          9
                              12
                                  0 16
                                         1
                                            TRUE FALSE
# 5
                     1014 1
                              0
                                  0 19
                                         0
                                            TRUE FALSE
                     1015 13
                                   0 24
                                         1 TRUE FALSE
                                  0 36 0 TRUE FALSE
                     3367 10
                              0
                     1335 20
                                  0 68
# 8
                              0
                                         1 TRUE FALSE
# 9
    Chr1 2319
                     1007 0
                             29 22 0
# 10
                                         1 TRUE FALSE
                     1012 0
                             54 26 0
# 11
                                        1 TRUE FALSE
                                         1 TRUE FALSE
# 12
                     1013 19
                              19
                                 18 0
                                         1
# 13
                     1014
                          0
                              25
                                 19
                                     0
                                            TRUE FALSE
# 14
                     1015
                          0
                              54
                                 29
                                     0
                                         1
                                            TRUE FALSE
# 15
                     3367 5
                             0 43 0
                                         O TRUE FALSE
# 16
                     1335 0 132
                                 48 0
                                        1 TRUE FALSE
# 17 Chr1 3286
# 18
                     1007
                           4
                               0
                                  1 17
                                         0
                                            TRUE FALSE
                     1012 9
                                  3 45
# 19
                               0
                                         0
                                            TRUE FALSE
# 20
                     1013 39
                               1
                                 38 12
                                         1
                                            TRUE FALSE
# 21
                     1014 4
                                  6 27
                                         O TRUE FALSE
                               0
# 22
                     1015 11
                                  7 37
                                         O TRUE FALSE
# 23
                     3367 8
                               0
                                 39 10
                                         0
                                            TRUE FALSE
                                 4 75 0 TRUE FALSE
                     1335 15 0
# 24
```

```
# 25 Chr1 5002 T
# 26
                    1007 0 15
                                0 12
                                      O TRUE FALSE
                                       1 TRUE FALSE
0 TRUE FALSE
# 27
                    1012 1
                            23
                                 0 26
                    1013 21
# 28
                             11
                                 0 39
# 29
                    1014 0
                             8
                                 0 12
                                        O TRUE FALSE
# 30
                    1015 0
                            19
                                 0 16
                                      1 TRUE FALSE
# 31
                    3367 0
                             0
                                 0 35
                                       O TRUE FALSE
 32
                    1335 0
                             57
                                 0 60
                                        0 TRUE FALSE
# 33 Chr1 5433
# 34
                    1007 0
                             50
                                 0 3
                                        0 TRUE FALSE
# 35
                    1012 0
                            78
                                 0 5
                                       O TRUE FALSE
                                 0 14
# 36
                    1013 18
                            47
                                        1 TRUE FALSE
                                           TRUE FALSE
# 37
                    1014 9
                             19
                                 0 0
                                        O TRUE FALSE
                         7
                                 0 2
# 38
                    1015
                             63
                                 0 0 0 TRUE FALSE
# 39
                    3367 8 54
# 40
                    1335 6 109
                                0 4 0 TRUE FALSE
# 41 Chr1 7858
# 42
                    1007
                         0
                              0 48 0
                                        0
                                          TRUE FALSE
                             1 61 0
                                        O TRUE FALSE
# 43
                    1012 0
# 44
                    1013 0
                             0 131 10
                                        O TRUE FALSE
# 45
                    1014 0
                             0 34 0
                                        O TRUE FALSE
                                        0
# 46
                    1015 0
                             0
                                74
                                    0
                                          TRUE FALSE
# 47
                    3367 20
                              0
                                8
                                    0
                                        1
                                           TRUE FALSE
                    1335 0
                             0 120 0
                                        O TRUE FALSE
# 48
# 49 Chrl 8914 A
# 50
                    1007 23
                              0
                                0 2
                                        O TRUE FALSE
                    1012 29
                                15
# 51
                              0
                                    0
                                        1
                                           TRUE FALSE
                    1013 25
# 52
                              0
                                 6
                                    0
                                        0
                                          TRUE FALSE
# 53
                    1014 22
                              ()
                                 0 0
                                        O TRUE FALSE
# 54
                    1015 31
                              0
                                5 2
                                        O TRUE FALSE
# 55
                    3367 8
                              0 0 1
                                        0 TRUE FALSE
                    1335 68
                              0
                                 7 0
                                        0
                                          TRUE FALSE
# 57 Chrl 8974
# 58
                    1007 0
                             2
                                6 0
                                        O TRUE FALSE
# 59
                    1012 0
                             2 17 0 0 TRUE FALSE
                    1013 10
                            22
                                        1 TRUE FALSE
# 60
                                   0
                                 4
                                10
# 61
                    1014
                         0
                              1
                                    ()
                                        0
                                           TRUE FALSE
                    1015 0
                              2
                                15
# 62
                                    0
                                        0
                                          TRUE FALSE
# 63
                    3367 2
                             0
                                3 0
                                        O TRUE FALSE
# 64
                    1335 0
                            11 49 0
                                        0 TRUE FALSE
# 65 Chr1 10099
                    1007 17
                             0
                                0 29
                                        O TRUE FALSE
# 66
                                0 36
# 67
                    1012 48
                             0
                                        O TRUE FALSE
                    1013 0
                                        0 TRUE FALSE
# 68
                                 6 68
# 69
                    1014 34
                            0
                                 0 26 0 TRUE FALSE
 70
                    1015 41
                             0
                                 0 38
                                        0 TRUE FALSE
# 71
                    3367 0
                             1
                                 0 14
                                        0
                                           TRUE FALSE
                    1335 55
# 72
                            0
                                 0 68
                                        1 TRUE FALSE
 73 Chrl 15154 A
                                0 0 0 FALSE FALSE
# 74
                    1007 25
                             0
 75
                    1012 56
                             0
                                 0 1
                                        O FALSE FALSE
# 76
                    1013 10
                             0
                                38 10
                                        1 FALSE FALSE
# 77
                    1014 26
                                        O FALSE FALSE
                             0
                                 0 0
# 78
                    1015 37
                              0
                                 0 0
                                       O FALSE FALSE
# 79
                    3367 19
                              0
                                0 13
                                        1 FALSE FALSE
                    1335 70
# 80
                             0 0 3 0 FALSE FALSE
```

4.5 Examples: Homozygous nonref

And at some *homozygous nonreference* positions (defined to be those with nonref fraction > 0.75):

```
hnr <- lapply(snp.tables, function(x) {x$.match/x$Cov < 0.25})  # find them
hnr <- lapply(hnr, function(x) {ifelse(is.na(x), FALSE, x)})  # remove NA
unlist(lapply(hnr, sum))  # count per strain</pre>
# 1007 1012 1013 1014 1015 3367 1335
```

```
# 65 65 6281 23 62 7071 40
```

Hmm, in L-clade, excluding the ref isolate (1335) this tracks time-in culture to some degree; Maybe many of these are in hemizygous regions. Next two chunks lifted from nc-snps to get tables for hemi-deletion.

```
cnv.chronly <- load.cnv.tables('../../data/cnv.txt', chrs.only=TRUE)</pre>
str(cnv.chronly)
# 'data.frame': 1956 obs. of 11 variables:
# $ strain : Factor w/ 7 levels "IT", "tp1007",...: 3 3 3 3 3 3 3 3 3 3 ...
             : Factor w/ 65 levels "BD1_7","BD10_65",...: 38 38 38 38 38 38 38 38 38 38 ...
# $ chr
               : int 10601 112001 215001 358901 536501 554801 673401 781801 806901 853201 ...
: int 13500 116500 221100 370300 538600 559300 685000 787400 811100 855600 ...
  $ start
  $ end
# $ length : int 2900 4500 6100 11400 2100 4500 11600 5600 4200 2400 ...
# $ filtered : logi FALSE FALSE FALSE TRUE FALSE FALSE ...
  $ type : Factor w/ 1 level "CNVnator": 1 1 1 1 1 1 1 1 1 1 ...
  $ cov_ratio: num   0.63738   1.54893   1.65381   0.00204   0.68486   ...
# $ dup_frac : num   0.41188   0.00908   0.01178   0.97997   0.0211   ...
# $ iStart : num 10601 112001 215001 358901 536501 ...
# $ iEnd : num 13500 116500 221100 370300 538600 ...
cnv.chronly[\mathbf{c}(1:4, \mathbf{nrow}(\text{cnv.chronly}) + \mathbf{c}(-1, 0)),]
                                                                           ## first/last few rows
                                end length filtered type cov_ratio dup_frac iStart
       strain chr start
     tp1012 Chr1 10601 13500 2900 FALSE CNVnator 0.63738000 0.41187900 10601
                                                                                                      13500
# 1
                                                                                            112001
     tp1012 Chr1 112001 116500 4500 FALSE CNVnator 1.54893000 0.00907677 tp1012 Chr1 215001 221100 6100 FALSE CNVnator 1.65381000 0.01178470
                                                                                                      116500
       tp1012 Chr1 215001 221100 6100 FALSE CNVnator 1.65381000 0.01178470 215001 tp1012 Chr1 358901 370300 11400 TRUE CNVnator 0.00204431 0.97997300 358901
       tp1012 Chr1 215001 221100
                                                                                                       221100
                                                                                                      370300
# 1955 tp1335 Chr24 259901 278000 18100 FALSE CNVnator 1.41458000 0.38091100 31264334 31282433
# 1956 tp1335 Chr24 286901 289800 2900 FALSE CNVnator 1.74941000 0.74228100 31291334 31294233
```

```
get.cnv.dels <- function(cov.thresh.lo = 0.0,</pre>
                          cov.thresh.hi = 0.8,
                          cnv,
                          snp.tables = NULL,
                          DEBUG = FALSE
  # build list of 7 Bool vectors of genome length, with i-th == T iff
 # * i-th pos is 'NA' in genome seq (if snp.tables are provided), or
 # * in CNVnator call for coverage in half-open [cov.thresh.lo, hi), and
  # * not marked 'filtered' by CNVnator
  cnv.deletions <- vector(mode='list',7)</pre>
                                                          # make list of bool vectors
 if(is.null(snp.tables)){
   # if no tables, assume full
   t.len <- genome.length.constants() $genome.length.trunc
  } else {
    t.len <- nrow(snp.tables[[1]])</pre>
  for(st in 1:7){
   if(is.null(snp.tables)){
     cnv.deletions[[st]] <- logical(t.len)</pre>
                                                                      # all F
    } else {
      cnv.deletions[[st]] <- is.na(snp.tables[[st]]$Pos[1:t.len]) # NA positions in genome</pre>
  }
  strain.names <- c(paste('tp10',c('07',12:15),sep=''),'IT','tp1335')
  names (cnv.deletions) <- strain.names</pre>
  for(i in 1:nrow(cnv)){
   if(!cnv$filtered[i] &&
       cnv$cov_ratio[i] >= cov.thresh.lo &&
       cnv$cov_ratio[i] < cov.thresh.hi)</pre>
      if (DEBUG) {
       print(cnv[i,])
        print(as.character(cnv$strain[i]))
```

Slight discrepancy in H-clade that I should hunt down, but basically OK. (hmm; maybe untrunc tbls.)

```
# the ones we want for the current analysis:
hemi.masks <- get.cnv.dels(0.3, 0.8, cnv.chronly, snp.tables=snp.tables)</pre>
rbind(
 homnr
             = unlist(lapply(hnr,sum)),
            = unlist(lapply(hemi.masks, sum)),
 homnr.unhemi = unlist(lapply(list(1,2,3,4,5,6,7), function(i)(sum(hnr[[i]] & !hemi.masks[[i]]))))
              1007 1012 1013 1014 1015 3367 1335
# homnr
                65
                     65 6281
                                23
                                      62 7071
             11761 16653 24337 11603 19824 49254 15367
# hemi
# homnr.unhemi 65 64 6193 23 52 6962 40
```

```
# based on the thought that hnr in 1335 may reflect errors in the ref seq,
# are they shared with others?
unlist(lapply(hnr, function(x){sum(x & hnr[[7]])}))
                                                                   # hnr shared with 1335
# 1007 1012 1013 1014 1015 3367 1335
  11 15 18 11 14 19 40
# answer: around 300 in each strain, of 558 in NY, genomewide,
# so that seems like a plausibly important factor.
hnr.lclade <- hnr[[1]] | hnr[[2]] | hnr[[4]] | hnr[[5]] | hnr[[7]] # union over L-clade
                                                                    # count all in L-clade
sum(hnr.lclade)
# [1] 152
sum(hnr[[3]] | hnr[[6]])
                                                                    # present in H-clade
# [1] 10348
sum(hnr[[3]] & hnr[[6]])
                                                                    # shared in H-clade
# [1] 3004
# look at a few in L-clade
w.hnr.l <- which (hnr.lclade)
seecounts(w.hnr.l[1:10], snp.tables=snp.tables)
```

#	chr	pos	Ref	Strain	A	G	С	T	SNP	exon	indel	nrf rat	
# 1	Chr1	5397	С										
# 2				1007	0	0	24	27	1	TRUE	FALSE		
# 3				1012	0		34	40	1		FALSE		
				1013	0		12	42			FALSE		
# 4									0				
# 5				1014	1		30	28	1		FALSE		
# 6				1015	0	0	33	35	1	TRUE	FALSE		
# 7				3367	0	0	20	38	1	TRUE	FALSE		
# 8				1335	0	0	29	98	1	TRUE	FALSE		
	Cl 1	20071		1333	O	0	2)	20	_	11(01	1111011		
	CHLI	20071	T			_	_						
# 10				1007	22	0	0	15	1	FALSE	FALSE		
# 11				1012	109	0	0	41	1	FALSE	FALSE		
# 12				1013	28	0	0	33	1	FALSE	FALSE		
# 13				1014	76		0	29		FALSE			
# 14				1015	130			28		FALSE			
# 15				3367	27	0	0	28	0	FALSE	FALSE		
# 16				1335	95	0	0	57	0	FALSE	FALSE		
# 17	Chr1	25350	G										
18				1007	104	31	0	0	1	FALSE	EVICE		
19				1012	171	53				FALSE			
20				1013	209	87	1	0	0	FALSE	FALSE		
21				1014	19	32	0	0	0	FALSE	FALSE		
22				1015	91					FALSE			
23				3367	397					FALSE			
24				1335	80	64	0	0	0	FALSE	F'ALSE		
25	Chr1	26205	T										
26				1007	50	0	0	20	1	FALSE	FALSE		
27				1012		0	0	33		FALSE			
28				1013	224	0	0	69		FALSE			
29				1014	23	0	1	16	1	FALSE	FALSE		
30				1015	88	0	0	33	1	FALSE	FALSE		
31				3367	143	0	0	41	1	FALSE	FALSE		
32				1335	196	0		67		FALSE			
	G1. 1	00040	0	1333	100	0	0	0 /	_	1111011	1111011		
	Chrl	90942	С										
# 34				1007	0	0	15	0	0	FALSE	FALSE		
# 35				1012	0	0	33	0	0	FALSE	FALSE		
# 36				1013	0	0	46	0	0	FALSE	FALSE		
# 37				1014	0		16	0		FALSE			
# 38				1015	0		7			FALSE			
ŧ 39				3367	0	0	56	0	0	FALSE	FALSE		
ŧ 40				1335	0	0	70	0	0	FALSE	FALSE		
# 41	Chr1	149447	T										
42			_	1007	0	0	0	1	0	FALSE	DATER		
43				1012	0					FALSE			
44				1013	0	0	0	1	0	FALSE	FALSE		
45				1014	0	0	0	8	0	FALSE	FALSE		
46					0	0	0	2	0	FALSE	FALSE		
: 47				3367	0	0	0	1	0	EVICE	EVICE		
										FALSE			
48				1335	0	1	0	1	0	FALSE	FALSE		
4 9	Chr1	149457	<NA $>$										
50				1007	<na></na>	<na></na>	<na></na>	<na></na>	0	FALSE	FALSE		
51										FALSE			
52										FALSE			
53				1014	<na></na>	<na></na>	<na></na>	<na></na>	0	FALSE	FALSE		
54				1015	<na></na>	<na></na>	<na></na>	<na></na>	0	FALSE	FALSE		
£ 55										FALSE			
ŧ 56										FALSE			
	Cl- 1	150040	70		/IVI/	/INT/	/INT/	-1VII	U	11100	11100		
	Chrl	156248	А										
58						0				FALSE			
59				1012	7	0	67	0	0	FALSE	FALSE		
				1013	5	0	53	0	0	FALSE FALSE	FALSE		
60													
						0				FALSE			
61				1015	6	0	44	0	0	FALSE	FALSE		
† 61 † 62						0	66	0	0	FALSE	FALSE		
† 61 † 62				3367	9	U	0.0	O					
# 61 # 62 # 63													
# 60 # 61 # 62 # 63 # 64		176517	C	1335	9 62					FALSE			
# 61 # 62 # 63 # 64		176517	С	1335	62	0	31	0	1		FALSE		

```
# 67
                       0
# 68
                           0 4 0 0 TRUE FALSE
                   1013
                                     0 0 TRUE FALSE
0 0 TRUE FALSE
# 69
                   1014
                         0
                             0
                                 6
# 70
                   1015
                         0
                             0
                                 0
                                   0 0 TRUE FALSE
# 71
                   3367 0 0 4
# 72
                   1335 0 0 11 0 0 TRUE FALSE
# 73 Chrl 193761
                   1007
                         0
                             0
                                20
                                    14
                                        1 FALSE FALSE
                             0 19 31 1 FALSE FALSE
# 75
                   1012
                         0
                   1013 0 1 4 6 1 FALSE FALSE
# 76
# 77
                   1014 0 0 9 4 1 FALSE FALSE
                   1015 0 0 28 39 1 FALSE FALSE
# 78
                           0 7 11 0 FALSE FALSE
0 10 43 1 FALSE FALSE
# 79
                         0
                   3367
                       0
# 80
                   1335
# one of those is a little weird:
xx<-snp.tables[[1]][149457,]
for (i in 2:7) {xx <- rbind(xx, snp.tables[[i]][149457,])}</pre>
row.names (xx) <-names (snp.tables)</pre>
# My quess is that Chr/Pos/Ref are left as NA if coverage is zero.
                      Pos Ref Cov a g c t n .match exon indel
          pos snp Chr
      chr
# 1007 Chr1 149457 0 <NA> NA <NA> 0 0 0 0 0 0 0 FALSE FALSE # 1012 Chr1 149457 0 <NA> NA <NA> 0 0 0 0 0 0 0 FALSE FALSE
O FALSE FALSE
                                              1 FALSE FALSE
               O FALSE FALSE
# 3367 Chr1 149457
                                               O FALSE FALSE
```

5 Table 1 stats

Here is a brief summary of per-strain SNP counts, pairwise overlaps, and other conveniently available stats, such as those shown in Table 1 of the paper.

```
snp.counts <- matrix(NA,7,4)</pre>
snp.pctofny <- matrix(NA, 7, 4)</pre>
snp.pctofself <- matrix(NA,7,4)</pre>
snp.inter <- matrix(NA,7,7)</pre>
snp.union <- matrix(NA, 7, 7)</pre>
rownames (snp.counts) <- names (snp.tables)</pre>
rownames (snp.pctofny) <- names (snp.tables)</pre>
rownames (snp.pctofself) <- names (snp.tables)</pre>
rownames (snp.inter) <- names (snp.tables)</pre>
colnames (snp.inter) <- names (snp.tables)</pre>
rownames (snp.union) <- names (snp.tables)</pre>
colnames (snp.union) <- names (snp.tables)</pre>
for(stringency in 1:4){
 cat('\nStringency', stringency, ifelse(stringency==4,'(i.e. raw SAMTools SNP calls)',''),
      ':\n----\n')
  for(i in 1:7){
    f.snps.i <- filtered.snps$Code$get.snps(i, stringency)</pre>
    snp.counts[i,stringency] <- sum(f.snps.i)</pre>
    for(j in i:7){
      f.snps.j <- filtered.snps$Code$get.snps(j, stringency)</pre>
      snp.inter[i,j] <- sum(f.snps.i & f.snps.j)</pre>
      snp.union[i,j] <- sum(f.snps.i | f.snps.j)</pre>
  snp.pctofny [,stringency] <- snp.inter[,7]/snp.counts[7,stringency]</pre>
  snp.pctofself[,stringency] <- snp.inter[,7]/snp.counts[ ,stringency]</pre>
  cat('Union Counts:\n');
                                               print (snp.union)
  cat('Intersect Counts:\n');
                                             print(snp.inter)
```

```
cat('Intersect as percent of union:\n'); print(snp.inter/snp.union*100,digits=3)
# Stringency 1 :
# Union Counts:
  1007 1012 1013 1014 1015 3367 1335
# 1007 17466 18310 30476 18476 18305 30017 18612
# 1012 NA 17836 30594 18774 18574 30130 18883
       NA NA 24999 30148 30643 31962 30648
# 1013
# 1014
        NA
             NA NA 16338 18733 29644 18990
       NA
                NA NA 17814 30160 18852
# 1015
             NA
# 3367 NA NA NA
                     NA NA 24021 30175
# 1335 NA NA
                NA
                      NA NA NA 18039
# Intersect Counts:
      1007 1012 1013 1014 1015 3367 1335
# 1007 17466 16992 11989 15328 16975 11470 16893
# 1012 NA 17836 12241 15400 17076 11727 16992
       NA NA 24999 11189 12170 17058 12390
# 1013
# 1014
        NA
             NA NA 16338 15419 10715 15387
                  NA NA 17814 11675 17001
# 1015
        NA
             NA
# 3367
             NA NA
                       NA NA 24021 11885
        NA
# 1335 NA NA NA NA
                            NA NA 18039
# Intersect as percent of union:
     1007 1012 1013 1014 1015 3367
# 1007 100 92.8 39.3 83.0 92.7 38.2
      NA 100.0 40.0 82.0 91.9 38.9 90.0
# 1012
# 1013 NA NA 100.0 37.1 39.7 53.4 40.4
            NA NA 100.0 82.3 36.1 81.0
# 1014
       NA
# 1015
                 NA NA 100.0 38.7
       NA
            NA
                      NA NA 100.0 39.4
# 3367
       NA
            NA
                 NA
# 1335 NA NA
                NA NA
                          NA NA 100.0
# Stringency 2 :
# Union Counts:
      1007 1012 1013 1014 1015 3367 1335
# 1007 18089 18521 37510 18365 18729 36679 18497
# 1012 NA 18342 37599 18538 18771 36774 18566
           NA 30797 36124 37777 41302 37426
# 1013
        NA
# 1014
       NA
             NA NA 14194 18733 35137 18202
# 1015 NA NA NA NA 18551 36940 18754
# 3367 NA NA NA NA NA 29496 36558
                      NA
# 1335 NA
           NA
                  NA
                           NA NA 17819
# Intersect Counts:
  1007 1012 1013 1014 1015 3367 1335
# 1007 18089 17910 11376 13918 17911 10906 17411
# 1012 NA 18342 11540 13998 18122 11064 17595
        NA
             NA 30797 8867 11571 18991 11190
# 1013
# 1014
        NA
             NA NA 14194 14012 8553 13811
             NA NA NA 18551 11107 17616
# 1015
        NA
# 3367 NA NA NA NA NA 29496 10757
# 1335 NA NA NA NA
                            NA NA 17819
# Intersect as percent of union:
    1007 1012 1013 1014 1015 3367 1335
# 1007 100 96.7 30.3 75.8 95.6 29.7 94.1
# 1012
       NA 100.0 30.7 75.5 96.5 30.1 94.8
       NA NA 100.0 24.5 30.6 46.0 29.9
# 1013
# 1014
            NA
                NA 100.0
                          74.8
                               24.3
       NA
                 NA NA 100.0 30.1 93.9
# 1015
       NA
            NA
                     NA NA 100.0 29.4
# 3367
       NA
           NA
                 NA
# 1335 NA
          NA
                NA NA NA NA 100.0
# Stringency 3 :
# -----
# Union Counts:
# 1007 1012 1013 1014 1015 3367 1335
```

```
# 1007 17107 18179 36938 17408 18347 36105 17958
# 1012 NA 17881 37265 18044 18505 36432 18258
        NA
# 1013
              NA 30364 34100 37453 41272 36889
              NA NA 9791 18257 33046 17322
# 1014
        NA
# 1015
        NA
              NA
                   NA
                       NA 18095 36626 18420
# 3367
       NA
            NA NA NA NA 29135 36011
                       NA NA NA 16984
                 NA
# 1335 NA NA
# Intersect Counts:
       1007 1012 1013 1014 1015 3367 1335
# 1007 17107 16809 10533 9490 16855 10137 16133
# 1012 NA 17881 10980 9628 17471 10584 16607
        NA NA 30364 6055 11006 18227 10459
# 1013
              NA NA 9791 9629 5880 9453
NA NA NA 18095 10604 16659
# 1014
        NA
# 1015
        NA
            NA NA NA NA 29135 10108
# 3367
       NA
# 1335 NA NA NA NA NA NA 16984
# Intersect as percent of union:
     1007 1012 1013 1014 1015 3367 1335
# 1007 100 92.5 28.5 54.5 91.9 28.1 89.8
# 1012 NA 100.0 29.5 53.4 94.4 29.1 91.0
       NA NA 100.0 17.8 29.4 44.2 28.4
# 1013
             NA NA 100.0 52.7 17.8 54.6
NA NA NA 100.0 29.0 90.4
# 1014
        NA
# 1015
        NA
# 3367
                  NA
                      NA NA 100.0 28.1
       NA
             NA
                 NA NA
                            NA NA 100.0
# 1335 NA NA
# Stringency 4 (i.e. raw SAMTools SNP calls) :
# Union Counts:
  1007 1012 1013 1014 1015 3367 1335
# 1007 16530 17707 35005 16864 17989 34289 17382
       NA 17019 35294 17276 18074 34563 17577
        NA NA 25412 30445 35599 39448 34479
# 1013
# 1014
       NA NA NA 8331 17634 29704 16078
# 1015
       NA NA NA NA 17397 34876 17881
       NA NA NA NA NA 24613 33699
# 3367
# 1335
       NA
             NA
                   NA
                       NA
                             NA NA 15582
# Intersect Counts:
      1007 1012 1013 1014 1015 3367 1335
# 1007 16530 15842 6937 7997 15938 6854 14730
# 1012 NA 17019 7137 8074 16342 7069 15024
        NA
# 1013
              NA 25412 3298 7210 10577 6515
              NA NA 8331 8094 3240 7835
# 1014
        NA
       NA NA NA NA 17397 7134 15098
# 1015
# 3367
       NA NA NA NA NA 24613 6496
# 1335 NA NA NA NA NA NA 15582
# Intersect as percent of union:
  1007 1012 1013 1014 1015 3367 1335
# 1007 100 89.5 19.8 47.4 88.6 20.0 84.7
# 1012 NA 100.0 20.2 46.7 90.4 20.5 85.5
       NA NA 100.0 10.8 20.3 26.8 18.9
NA NA NA 100.0 45.9 10.9 48.7
# 1013
                                      18.9
# 1014
# 1015
                  NA NA 100.0 20.5 84.4
             NA
       NA
# 3367
                 NA NA NA 100.0 19.3
       NA NA
# 1335 NA NA
                 NA NA NA NA 100.0
vs.stringency <- cbind(snp.counts, matrix(NA,7,1), round(snp.counts[,1:3]/snp.counts[,4]*100,1))
colnames(vs.stringency) <- c('[[1]]', '[[2]]', '[[3]]', '[[4]]', '----', '[[1]]%', '[[2]]%', '[[3]]%')</pre>
# SNPs vs filtering stringency (raw counts and as % of [[4]]). Medium filter
# adds 10-20% in most cases. Big exception is Gyre, where low coverage,
# high err rate and SAMTools conservatism seemed to seriously undercall:
print (vs.stringency)
      [[1]] [[2]] [[3]] [[4]] ---- [[1]]% [[2]]% [[3]]%
# 1007 17466 18089 17107 16530 NA 105.7 109.4 103.5
# 1012 17836 18342 17881 17019 NA 104.8 107.8 105.1
# 1013 24999 30797 30364 25412 NA 98.4 121.2 119.5
```

```
# 1014 16338 14194 9791 8331 NA 196.1 170.4 117.5
# 1015 17814 18551 18095 17397
                                    NA 102.4 106.6 104.0
# 3367 24021 29496 29135 24613 NA 97.6 119.8 118.4 # 1335 18039 17819 16984 15582 NA 115.8 114.4 109.0
# Intersect NY as % of self (vs stringency):
print(snp.pctofself*100, digits=3)
# [,1] [,2] [,3] [,4]
# 1007 96.7 96.3 94.3 89.1
# 1012 95.3 95.9 92.9 88.3
# 1013 49.6 36.3 34.4 25.6
# 1014 94.2 97.3 96.5 94.0
# 1015 95.4 95.0 92.1 86.8
# 3367 49.5 36.5 34.7 26.4
# 1335 100.0 100.0 100.0 100.0
# Intersect NY as % of NY (vs stringency):
print(snp.pctofny*100, digits=3)
         [,1] [,2] [,3] [,4]
# 1007 93.6 97.7 95.0 94.5
# 1012 94.2 98.7 97.8 96.4
# 1013 68.7 62.8 61.6 41.8
# 1014 85.3 77.5 55.7 50.3
# 1015 94.2 98.9 98.1 96.9
# 3367 65.9 60.4 59.5 41.7
# 1335 100.0 100.0 100.0 100.0
```

Quick look at coverage. Are there any NA?:

Seemingly no. What's average in unq- vs q-filtered:

```
snp.tables.unqfil <- tset.picker(c(1,2), table.set = tset)
snp.tables.qfil <- tset.picker(c(3,4), table.set = tset)
cov.unqfil <- unlist(lapply(snp.tables.unqfil, function(x) {mean(x$Cov)}))
cov.qfil <- unlist(lapply(snp.tables.qfil, function(x) {mean(x$Cov,na.rm=T)}))
cov.both <- rbind(cov.unqfil,cov.qfil,cov.qfil/cov.unqfil)
i <- 1
if(!is.null(snp.tables.unqfil)) {
  rownames(cov.both)[i] <- which.snp.tables(snp.tables.unqfil)
  i <- i+1
}
if(!is.null(snp.tables.qfil)) {
  rownames(cov.both)[i] <- which.snp.tables(snp.tables.qfil)
  i <- i+1
}
if(i==3) {
  rownames(cov.both)[i] <- 'Ratio'
}
cat('Mean Coverage:\n'); cov.both</pre>
```

```
# Mean Coverage:

# 1007 1012 1013 1014 1015 3367 1335

# Chr1-unfiltered 36.2816326 68.2005811 66.6908911 31.2663216 59.4704151 62.3834535 103.9124774

# Chr1-qfiltered 27.5849516 49.2557296 43.2293645 12.4319866 46.9874722 43.4403699 78.7789843

# Ratio 0.7603007 0.7222186 0.6482049 0.3976159 0.7900983 0.6963444 0.7581282
```

5.1 Table 1 Data

Throw together the conveniently-available Table 1 data, in Table 1 row order:

```
# if coverage unavailable, build NA vector
if(!is.null(cov.unqfil)){cov.unqfilv <- cov.unqfil} else {cov.unqfilv <- rep(NA,times=7)}</pre>
if(!is.null(cov.qfil )){cov.qfilv <- cov.qfil } else {cov.qfilv <- rep(NA,times=7)}</pre>
tldata.df <- data.frame(</pre>
 id
           = st.locs(1:7, id=T, loc=F, date=F),
           = st.locs(1:7, id=F, loc=T, date=F),
           = st.locs(1:7, id=F, loc=F, date=T),
 date
  cov.unq = cov.unqfilv,
           = cov.qfilv,
  cov.q
           = snp.counts[,4],
 SNPs.4
           = snp.counts[,2],
 olap.ny.4 = snp.pctofny[,4]*100,
  olap.ny.2 = snp.pctofny[,2]*100
t1row.order < -c(7,1,2,5,3,6,4)
print (t1data.df[t1row.order,],digits=3)
                            loc date cov.unq cov.q SNPs.4 SNPs.2 olap.ny.4 olap.ny.2
                      New York 1958 103.9 78.8 15582 17819 100.0
# 1335 CCMP1335
# 1007 CCMP1007
                      Virginia 1964
                                       36.3 27.6 16530 18089
                                                                      94.5
                                                                                97.7
# 1012 CCMP1012
                  W. Australia 1965
                                       68.2 49.3 17019 18342
                                                                      96.4
                                                                                98.7
# 1015 CCMP1015
                  Puget Sound 1985
                                       59.5 47.0 17397
                                                           18551
                                                                      96.9
                                                                                98.9
                                                    25412
# 1013 CCMP1013
                          Wales 1973
                                             43.2
                                                           30797
                                                                      41.8
                                                                                62.8
                         Italy 2007
                                        62.4 43.4
                                                                                60.4
# 3367 CCMP3367
                                                    24613
                                                           29496
                                                                      41.7
# 1014 CCMP1014 N. Pacific Gyre 1971
                                       31.3 12.4
                                                    8331 14194
                                                                      50.3
```

6 Shared-SNPs P-Value

Text of the main paper quotes a "p-value" for the observed degree of SNP sharing in L-clade (and/or L-clade excluding Gyre) under a null model that these isolates were sampled from a population globally in Hardy-Weinberg equilibrium. Details of this analysis are as follows.

6.1 SNP Concordance

Arbitrarily pick one isolate, say, A, as the "template". Arbitrarily pick a heterozygous (aka "SNP") position in A. Let p_1 , and $q_1 = 1 - p_1$ be the frequencies in the overall population of the two nucleotides observed at that position in A. (Positions having 3 or 4 nucleotide variants segregating in the population are assumed to be negligibly rare.) Under the HWE null model, a second isolate B will also be heterozygous at the same position with probability $2p_1q_1 \le 1/2$. Similarly, this position will be heterozygous in a third isolate C with the same probability, *independently*, and so on for isolates D and E. Overall, (assuming HWE) the probability that a heterozygous position in A is simultaneously heterozygous in the other 4 isolates is at most $1/2^4 = 1/16$. Continuing, suppose we pick a second heterozygous position in A, on a different chromosome with allele frequencies $p_2, q_2 = 1 - p_2$, say. Again assuming HWE, this position will be a SNP in all of B, C, D and E with probability $(2p_2q_2)^4 \le 1/16$, and this is independent of the first position, since segregation on different chromosomes is unlinked. Repeat this at 24 heterozygous positions in A, one per chromosome. Then, the number of five-way concordant positions observed should be dominated by the number observed when sampling from a binomial distribution with parameters n = 24 and p = 1/16, i.e., we expect at most 1/16 = 6.25% of positions to agree, or at most 24/16 = 1.5 five-way concordant positions in total. In sharp contrast,

choosing CCMP 1014 (North Pacific Gyre) as the template, we see many more five-way concordant positions than predicted under these assumptions:

```
gyre.count <- sum(snp.tables[[4]]$snp)
# 'unfil.' => unfiltered for consistency; see below.
unfil.fiveway.count <- sum( snp.tables[[4]]$snp * i4.snps)
unfil.fiveway.percent <- unfil.fiveway.count / gyre.count * 100
unfil.p.value <- pbinom(floor(unfil.fiveway.count/gyre.count*24)-1, 24, 1/16, lower.tail = FALSE)
consistency.comparison <-
    data.frame(
        fiveway.count = unfil.fiveway.count,
        fiveway.percent = unfil.fiveway.percent,
        p.value = unfil.p.value
    )
consistency.comparison

# fiveway.count fiveway.percent p.value
# 1 7628 91.56164 8.700771e-23</pre>
```

Namely, 8331 positions are called as SNPs in CCMP1014, of which 7628 or 91.5616373% are also called as SNPs in *all four* other L-clade isolates. 91.5616373% of 24 is 21.9747929, and the probability of seeing 21 or more "Heads" in 24 flips of a biased coin with $P(\text{Heads}) \leq 1/16$, i.e., our p-value under the HWE null hypothesis, is at most: 8.700771×10^{-23} based on this simple binomial model. This is obviously strong evidence against the null hypothesis. This analysis is potentially overly-simplistic in four respects, addressed below.

- 1. " $2pq \le 1/2$ " is conservative. Neutral theory predicts that most variant nucleotides are rare in the population, so $2pq \ll 1/2$ is to be expected. This should make our quoted p-value very conservative.
- 2. Effect of Erroneous SNP calls. We base our analysis on *predicted* (by SAMTOOLS) heterozygous positions, not absolute-truth, which may affect our conclusions. However,
 - False negatives in A are irrelevant, since we never examine those positions. (This is the motivation for using CCMP1014 as the template; it has the lowest predicted SNP rate, likely due to a high false negative rate in that sequencing run. As noted elsewhere, it had the lowest coverage and lowest sequence quality of the 7 isolates, both of which impare SNP calling.)
 - False negatives in *BCDE* make such positions appear *non*-concordant. For our purpose, this makes our statistic more conservative since it can only deflate a statistic that we argue is nevertheless unexpectedly large.
 - False positive calls in A are conservatively treated, as well: barring simultaneous false-positive calls in all of BCDE, such a position will appear non-concordant, again deflating the statistic. The *false* positive rates in B, C, D and E are unknown, but cannot exceed SAMTOOLS *total* positive rate, which is below 1% in all 7 isolates, suggesting a simultaneous BCDE false positive rate $< 10^{-8}$, which will have a negligible effect.
 - A potentially more serious issue is a true positive in A aligned to false positives in BCD and/or E. (I.e., a position that is polymorphic in the population and heterozygous in A, under the HWE null model is likely to be homozygous for one of the two alleles in one or more of BCDE; false positive SNP calls in all of those isolates would make the site appear concordant, i.e., provide evidence against the null model.) However, (a) my impression is that SAMTOOLS is more prone to false negative calls than to false positive calls (see Section 4), and (b) we would need a high rate of false positives to turn a truely heterozygous but non-concordant A call into a false "concordant" call—I'd expect at most half (especially given point 1 above) of BCDE to be heterozygous, but all would need to be falsely declared heterozygous. Such a high false positive rate on BCDE seems unlikely (see previous bullet), and would likely be counterbalanced by a similarly increased rate of false positives on A, which, as noted, tend to deflate our statistic (previous bullet again).
 - Systematic errors. If there were, say, a sequence-context-dependent bias in the DNA sequencing, mapping
 and/or SNP-calling that tended to suggest (or hide) a SNP at some position, we're going to systematically
 over- (or under-) estimate concordant SNPs across isolates. The discordance of called SNPs between the

L- and H-clades and within the H-clade suggests that this is not a major problem, but it is worth noting as a possibility.

3. Discordant nucleotides at "concordant" SNP positions. A "shared" SNP at a given position might be, say, G/C in one isolate vs T/C in another, reflecting an unexpected tri-allelic position in the population or a technical sequencing error. It is inappropriate to count such a "shared" SNP position as evidence against the null hypothesis, since it isn't clear that it is truely shared. Instead, I will identify such inconsistent positions, based on the "stringency [[2]]" criteria established above, and treat each as non-concordant. I.e., a position will be considered to be a "5-way concordant SNP" if and only if it was called as a SNP by SAMTOOLS (independently) in all 5 L-clade isolates, *and* shows the same dominant non-reference nucleotide in all 5, according to criteria [[2]] above. As it turns out, this correction has a very minor effect on the resulting p-value:

```
# 'unfil.' => Ignoring "consistency"; 'fil.' => Filtering for "consistency";
fil.fiveway.count <- sum((snp.tables[[4]]$snp * i4.snps)[union.snps == 1] & consistent[[2]])</pre>
fil.fiveway.percent <- fil.fiveway.count / gyre.count * 100
fil.p.value <- pbinom(floor(fil.fiveway.count/gyre.count*24)-1, 24, 1/16, lower.tail = FALSE)
# append new stats to previous table for easy comparison
consistency.comparison <-
  rbind (consistency.comparison,
       data.frame(
         fiveway.count = fil.fiveway.count,
         fiveway.percent = fil.fiveway.percent,
         p.value = fil.p.value
 )
rownames(consistency.comparison) <- c('unfiltered', 'consistency.filtered')</pre>
consistency.comparison
                     fiveway.count fiveway.percent
                                                        p.value
                               7628 91.56164 8.700771e-23
# unfiltered
                               7534
                                           90.43332 8.700771e-23
# consistency.filtered
```

In particular, it removes 1.1% of five-way consistent positions (only 94 of 7628 positions), and still shows a highly significant p-value.

4. " $P(E[X]) \neq E[P(X)]$ ". I'm expressing this poorly, but finding the p-value based on the *expected* number of concordant positions is somewhat non-standard. A more typical set-up would use the *actual* value of some statistic, then calculate the probability of observing a value that extreme (or more extreme) under the null model. The fundamental problem is that we have thousands of SNPs, but I don't see an easy way to use more than 24 of them at a time, because potential genetic linkage seemingly destroys statistical independence, which is key to most simple analyses. A somewhat more formal, but still non-standard, approach is the following. Suppose we randomly sample one SNP per chromosome and count the number X of them that are 5-way concordant. What I outlined above calculated the p-value based on E[X], the expected value of X, i.e., P(E[X]). Alternatively, we can calculate E[P(X)], the expected p-value. (They are not the same.) In effect, this averages the p-values that would be seen over many different randomly-sampled sets of 24 SNPs. This is not difficult to calculate. First, the probability that we would observe $0 \le i \le 24$ concordant positions in a sample of 24, given that 90.43% of positions are concordant follows this binomial distribution:

```
x.equals.i.distribution <- dbinom(0:24, 24, fil.fiveway.percent/100)
print(x.equals.i.distribution, digits=3)

# [1] 3.45e-25 7.84e-23 8.52e-21 5.90e-19 2.93e-17 1.11e-15 3.32e-14 8.06e-13 1.62e-11 2.72e-10
# [11] 3.86e-09 4.64e-08 4.75e-07 4.15e-06 3.08e-05 1.94e-04 1.03e-03 4.59e-03 1.69e-02 5.04e-02
# [21] 1.19e-01 2.14e-01 2.76e-01 2.27e-01 8.95e-02</pre>
```

Second, the p-value corresponding to $0 \le i \le 24$ observed concordant positions also follows a different binomial distribution:

```
p.val.of.x.equals.i <- c(1, pbinom(0:23, 24, 1/16, lower.tail = F))
print(p.val.of.x.equals.i, digits=3)

# [1] 1.00e+00 7.88e-01 4.48e-01 1.87e-01 5.95e-02 1.49e-02 3.01e-03 4.99e-04 6.90e-05 8.02e-06
# [11] 7.89e-07 6.60e-08 4.72e-09 2.87e-10 1.49e-11 6.59e-13 2.46e-14 7.66e-16 1.98e-17 4.14e-19
# [21] 6.88e-21 8.70e-23 7.88e-25 4.56e-27 1.26e-29</pre>
```

Finally, the expected (or "average") p-value is just the weighted average of the latter values, weighted by the former:

```
e.of.p.of.x <- sum(x.equals.i.distribution * p.val.of.x.equals.i)
e.of.p.of.x
# [1] 1.398085e-14</pre>
```

This is still highly significant, but weaker than the P(E[X]) analysis, basically because X < E[X] has a fair probability of occurring, and the corresponding p-value P(X) rises rapidly as X declines.

Another way to look at the numbers:

```
pvdf <- data.frame(x.density=x.equals.i.distribution,</pre>
                   x.cdf=cumsum(x.equals.i.distribution),
                   pval.of.x=p.val.of.x.equals.i)
print(pvdf, digits=4)
    x.density
                   x.cdf pval.of.x
# 1 3.454e-25 3.454e-25 1.000e+00
# 2 7.835e-23 7.870e-23 7.875e-01
# 3 8.518e-21 8.596e-21 4.476e-01
    5.904e-19 5.990e-19 1.869e-01
# 5 2.930e-17 2.990e-17 5.950e-02
# 6 1.108e-15 1.138e-15 1.490e-02
# 7 3.317e-14 3.430e-14 3.010e-03
# 8 8.062e-13 8.405e-13 4.994e-04
    1.619e-11 1.704e-11 6.899e-05
# 10 2.722e-10 2.892e-10 8.015e-06
# 11 3.859e-09 4.148e-09 7.887e-07
# 12 4.643e-08 5.058e-08 6.603e-08
# 13 4.755e-07 5.260e-07 4.716e-09
# 14 4.149e-06 4.675e-06 2.875e-10
# 15 3.081e-05 3.549e-05 1.493e-11
# 16 1.942e-04 2.297e-04 6.590e-13
# 17 1.033e-03 1.262e-03 2.456e-14
# 18 4.593e-03 5.855e-03 7.662e-16
# 19 1.689e-02 2.274e-02 1.977e-17
# 20 5.041e-02 7.315e-02 4.143e-19
# 21 1.191e-01 1.923e-01 6.877e-21
# 22 2.145e-01 4.067e-01 8.701e-23
# 23 2.765e-01 6.832e-01 7.884e-25
# 24 2.273e-01 9.105e-01 4.556e-27
# 25 8.951e-02 1.000e+00 1.262e-29
```

E.g., row 9 in that table says that the concordance rate (90%) is so high that a sample of 24 SNPs will almost always have 9 or more five-way concordant positions (probability of fewer is only 1.704e-11), while under the null model, seeing 9 or more is very unlikely (probability at most 6.899e-05). ***AM I OFF-BY-ONE INTERPRETING ROW 9 HERE??***

6.2 Notes

In earlier drafts, an analog of the above analysis was based on the concordance of *refined* SNPs. This now seems to me to be questionable, since the "refined" SNP calling makes SNPs called across L-clade non-independent. OTOH,

the above analysis seems valid: SAMTOOLS was run on each isolate independently, and likewise "criterion [[2]]" is evaluated independently in each strain, and is being used here solely to remove SNP predictions, not to add them. "Systematic errors" as outlined above remain a potential problem, but again discordance with/within H-clade suggests that this is of limited concern.

For completeness, I did a similar analysis including a sample of H-clade comparisons: Gyre vs Italy, NY vs Italy, NY vs Italy+Wales, and of Italy vs Wales. As expected, none of these show a statistically significant p-value, although the $\approx 40\%$ concordance in the 2-way comparisons, while < 1/2 as predicted, is a bit higher than I expected based on "neutral theory implies many rare variants." (I did not bother to include "criterion[[2]] filtering" in these calculations.)

```
# 'gi.twoway' => gyre vs italy 2-way concordance;
# 'ni.twoway' => new york vs italy 2-way concordance;
# not bothering with criterion[[2]] filtering
gi.twoway.count <- sum(snp.tables[[4]]$snp * snp.tables[[6]]$snp)</pre>
gi.twoway.percent <- gi.twoway.count / gyre.count * 100</pre>
gi.p.value <- pbinom(floor(gi.twoway.count/gyre.count*24)-1, 24, 1/2, lower.tail = FALSE)
ny.count <- sum(snp.tables[[7]]$snp)</pre>
ni.twoway.count <- sum(snp.tables[[7]]$snp * snp.tables[[6]]$snp)</pre>
ni.twoway.percent <- ni.twoway.count / ny.count * 100</pre>
ni.p.value <- pbinom(floor(ni.twoway.count/ny.count*24)-1, 24, 1/2, lower.tail = FALSE)
niw.threeway.count <- sum(snp.tables[[7]]$snp * snp.tables[[6]]$snp * snp.tables[[3]]$snp)</pre>
niw.threeway.percent <- niw.threeway.count / ny.count * 100</pre>
niw.p.value <- pbinom(floor(niw.threeway.count/ny.count*24)-1, 24, 1/4, lower.tail = FALSE)
it.count <- sum(snp.tables[[6]]$snp)</pre>
iw.twoway.count <- sum(snp.tables[[6]]$snp * snp.tables[[3]]$snp)</pre>
iw.twoway.percent <- iw.twoway.count / it.count * 100</pre>
iw.p.value <- pbinom(floor(iw.twoway.count/it.count*24)-1, 24, 1/2, lower.tail = FALSE)
consistency.comparison <-
  rbind (consistency.comparison,
       data.frame(
         fiveway.count = c(gi.twoway.count, ni.twoway.count, niw.threeway.count, iw.twoway.count),
         = c(gi.p.value,
                                         ni.p.value,
                                                           niw.p.value,
                                                                                      iw.p.value)
colnames (consistency.comparison) [1:2] <- c('552232way.count', '552232way.percent') # old col names misleading</pre>
rownames(consistency.comparison)[3:6] <- c('gyre.vs.italy', 'new.york.vs.italy',</pre>
                                                                              # new rows
                                         'ny.vs.it.plus.wales', 'it.vs.wales')
consistency.comparison
                      552232way.count 552232way.percent
                                                           p.value
                             7628 91.56164 8.700771e-23
# unfiltered
# consistency.filtered
                                7534
                                              90.43332 8.700771e-23
                                3240
                                              38.89089 9.242052e-01
# gyre.vs.italy
# new.york.vs.italy
                               6496
                                             41.68913 8.462719e-01
                                             24.41278 7.533516e-01
# ny.vs.it.plus.wales
                                3804
                                            42.97323 8.462719e-01
# it.vs.wales
                               10577
```

6.3 P-Value: The Bottom Line

So, what to say in the body of the paper? E[P(X)] is highly significant, and conservative, but complex to explain. P(E[X]) is simpler to explain, but may be criticized as misleading if we aren't very careful in that explanation. I'm slightly leaning towards the last option, but want to sleep on it and draft the key sentence or two before settling.

7 Sharing

The following analysis looks at the sharing patterns among the consistent SNPs. I assume that shared SNPs reflect shared ancestry, and that SNPs accumulate slowly over time. Then, in outline, the story is consistent with what we have seen in other analyses—there seem to be 3 groups: 1013 (Wales) in one, 3367 (Italy) in another, and the other 5 in a third, with some hints as to the order of divergence. A caveat is that in a sexual population, non-shared SNPs do not immediately imply non-shared ancestry; they may merely reflect Hardy-Weinberg capturing a homozygous state

in one isolate vs the other. (Or read errors, etc.) Thus, if we are right that the H-isolates retain sex, then the large number of "private" SNPs in H may be at least partially due to HWE.

Analysis is broken into cases based on how many strains share a particular SNP.

7.1 Code

To categorize SNPs by sharing patterns, first convert the 7-way consistent sharing pattern into a 7-bit binary number, and tabulate based on that:

```
# convert (n x 7) 0-1 matrix to n vector of 0-127
tobin <- function(x) {
  bin <- integer(nrow(x)) # initialized to 0</pre>
  for(i in 1:7){
   bin <- bin*2 + as.integer(x[,i]>0)
  return (bin)
# get full set of patterns
snp.pattern.all <- lapply(non.refs,tobin)</pre>
# prune to just the consistent ones
snp.pattern <- snp.pattern.all</pre>
for(i in 1:3){
 snp.pattern[[i]][!consistent[[i]]] <- NA</pre>
# analogous to built-in ``table'' but simpler. Count entries in an integer
# vector sharing values in a (smallish) range. Result is a 2-column matrix with
# the shared values in col 1 and count of occurrences of that value in col 2.
# Out-of-range values cause subscript error.
mytable <- function(vec, therange=range(vec,na.rm=T)){</pre>
  counts <- matrix(0, nrow=therange[2]-therange[1]+1, ncol=2, dimnames=list(NULL, c('val', 'count')))</pre>
  counts[1:nrow(counts),1] <- therange[1]:therange[2]</pre>
  for(i in 1:length(vec)){
    if(!is.na(vec[i])){
      \verb|counts|| vec[i] - the range[1] + 1, 2| <- counts|| vec[i] - the range[1] + 1, 2| + 1
  return (counts)
pattern.counts < lapply (snp.pattern, function(x) {mytable(x,c(0,127))})
```

To display the results, build a data frame whose i-th row, $0 \le i \le 127$ shows one of the 128 possible sharing patterns, with counts of the numbers of consistent, shared SNPs with that pattern according to criteria c1-c3.

```
bvec <- tobitvec(i-1)</pre>
    mydf[i,'sharedBy']=sum(bvec)
    mydf[i,'tp1007']=flg(bvec[1])
    mydf[i,'tp1012']=flg(bvec[2])
   mydf[i,'tp1013']=flg(bvec[3])
    mydf[i,'tp1014']=flg(bvec[4])
    mydf[i, 'tp1015']=flg(bvec[5])
    mydf[i,'tp3367']=flg(bvec[6])
    mydf[i,'tp1335']=flg(bvec[7])
  for(i in 1:length(listOfTbls)){
    tbl <- listOfTbls[[i]]</pre>
    if(!is.null(tbl)){
     mydf[,9+i] <- tbl[,2] ## count1/2/3/4 are columns 10/11/12/13 in mydf
      #for(j in 1:length(tbl)){
      # k <- as.integer(rownames(tbl)[j]);</pre>
         mydf[k+1,9+i] \leftarrow tbl[j] ## count1/2/3 are columns 10/11/12
      #}
    }
  mydf$pat <-as.octmode(mydf$pat) # display bit pattern in octal</pre>
  return (mydf)
pat.summaries <- pat.summary(pattern.counts)</pre>
```

7.2 Sanity Checks

Some sanity checking: table sums equal to number of consistent positions?

```
all(consistent.count == apply(pat.summaries[,10:13],2,sum))
# [1] TRUE
```

More sanity checking: visually inspect a pattern with small counts, specifically pattern 12, i.e., consistent SNPs shared by only strains 1014 and 1015 (2nd and 3 rows from bottom, binary code $12 = 2^3 + 2^2$). There are only 10 such positions on Chr1. Chr1 2524239 has pattern 12 under criteria c1 and c2 but not c3; Chr1 1088766 has in c2 only. Both look good. Neither position is a *called* SNP except in 1015. However, all but 1 nonreference read agree with the called SNP (the exception being one read in Wales). Both 1014 and 1015 have at least 2 non-reference reads, comprising at least 5% of coverage, and in both strains, those reads are on the same non-reference base, satisfying criterion c2. The other strains have higher coverage and/or lower non-reference counts, so they do not satisfy c2. Position 2524239 also satisfies c1, but not c3, since 2 reads out of 35 is below the 10% threshold. (It is pattern 4 inder c3, i.e., a SNP private to 1015.) Position 1088766 is also pattern 4 under c3 (2 reads out of 56 in 1335 is below both thresholds), and it is not consistent under c1, since the single A read in 1013 is discordant with the other non-reference reads.

```
unlist(lapply(snp.pattern, function(x) {sum(x==12,na.rm=T)}))
# [1] 10 2 5 12

sp1 <- snp.pattern[[1]]==12
sp2 <- snp.pattern[[2]]==12
sp3 <- snp.pattern[[3]]==12
sp4 <- snp.pattern[[4]]==12
c(sum(sp1,na.rm=T), sum(sp2,na.rm=T), sum(sp3,na.rm=T), sum(sp4,na.rm=T))
# [1] 10 2 5 12

r1 <- rownames(non.refs[[1]])[which(sp1)]
r2 <- rownames(non.refs[[2]])[which(sp2)]
r3 <- rownames(non.refs[[3]])[which(sp3)]
r4 <- rownames(non.refs[[4]])[which(sp4)]</pre>
```

```
# [1] "Chr1:1088766" "Chr1:2524239"
c1 <- as.integer(unlist(lapply(strsplit(r1[1:min(20,length(r1))],':',fixed=TRUE),function(x){x[2]})))
\texttt{c2} \leftarrow \textbf{as.integer}(\textbf{unlist}(\textbf{lapply}(\textbf{strsplit}(\texttt{r2}[1:\textbf{min}(20,\textbf{length}(\texttt{r2}))],':',\texttt{fixed=TRUE}),\textbf{function}(\texttt{x})\{\texttt{x}[2]\})))
\texttt{c3} \leftarrow \textbf{as.integer}(\textbf{unlist}(\textbf{lapply}(\textbf{strsplit}(\texttt{r3}[1:\textbf{min}(20,\textbf{length}(\texttt{r3}))],':',\texttt{fixed=TRUE}),\textbf{function}(\texttt{x})\{\texttt{x}[2]\})))
c4 <- as.integer(unlist(lapply(strsplit(r4[1:min(20,length(r4))],':',fixed=TRUE),function(x){x[2]})))
   [1] 198498 914018 1317406 1481838 1501481 1878058 2145849 2388286 2524239 2718093
# [1] 1088766 2524239
# [1] 371484 1210354 1886633 2264683 2898352
c.4
   [1] 518347
                  691730 767408 1049906 1390437 2072951 2254059 2254789 2264683 2823796 2898352
# [12] 2998868
seecounts(c2, snp.tables=snp.tables)
               pos Ref Strain A G C T SNP exon indel nrf rat
     Chrl 1088766 G
# 1
                            1007 0 32 1 0 0 FALSE FALSE
# 3
                           1012 0 39 1 0 0 FALSE FALSE
 4
                            1013 1 74 0 0
                                                O FALSE FALSE
                            1014 0 26
                                        2
#
                                           0
                                                O FALSE FALSE
                            1015 0 38 9 0
                                                1 FALSE FALSE
# 6
                            3367 0 36 1 0 0 FALSE FALSE
# 8
                           1335 0 54 2 0 0 FALSE FALSE
# 9 Chrl 2524239
# 10
                            1007 0 0 37 0
                                               O TRUE FALSE
                           1012 0 0 47 0 0 TRUE FALSE
# 11
                            1013 0 0 62 0 0 TRUE FALSE
# 12
                           1014 0 0 33 2
# 13
                                               O TRUE FALSE
                                                1 TRUE FALSE
# 14
                            1015 0
                                    0 11 15
# 15
                            3367 0
                                    0 41 0
                                                0
                                                    TRUE FALSE
                           1335 0 0 95 0 0 TRUE FALSE
# 16
```

Position 1088766, however, in a good example of the situation that motivated this analysis—one strain has a G/C SNP and 5 of the other 6 strains have nonreference reads consistent with that SNP. Although, excluding 1015, the nonreference read counts are not high enough to justify a SNP call in any strain considered in isolation, the fact that they *consistently* agree with the 1015 SNP suggests that they are real. One alternative hypothesis is that there is some sequence-dependent bias at this locus that favors misreading a G as a C. On the other hand, one could equally well posit a shared SNP, and a locus-dependant bias that *supresses* C reads, explaining the unbalanced readout that we observe. However, it is hard to reconcile either view with the significant strain-specific patterns that we see in the shared SNPs (as seen below). I think a more likely explanation is that (a) there are some number of relatively rare SNPs present in each of the sampled populations, (b) some of these SNPs happened to be present in one or two cells of the roughly 5-10 cells that we believe constituted the founding population of the culture grown for sequencing, and (c) stochastic effects during culture growth and during sequencing may have further perturbed the apparent frequency of each variant, but the bottom line is that the above-threshold presence of consistent non-reference reads is evidence for shared SNPs at the population level (and the proportions of such reads represent estimates of the population-level frequencies of the variants, albeit a noisy estimate at any specific position).

An aside: I was curious to see whether there is any consistent pattern to positions that are called consistent SNPs in all but Italy, so I repeated the above, basically. My summary is that coverage in Italy tends to be below average in these positions, but otherwise they don't stand out. For the record:

```
abit <- snp.pattern[[2]]==125
abit[is.na(abit)]<-F</pre>
```

```
sum(abit)
# [1] 1493
rabit <- rownames(non.refs[[2]])[which(abit)]</pre>
rabits <- rabit[1:20]</pre>
 \texttt{cabit} \leftarrow \textbf{as.integer(unlist(lapply(strsplit(rabits, ':', fixed=TRUE), function(x)\{x[2]\})))} 
cabit
# [1] 1244 1575 6485 7181 7220 7661 8144 8208 8518 8552 8567 8670 8685 14361 15254
# [16] 15280 16103 17587 18904 25546
seecounts(cabit,snp.tables=snp.tables)
      chr
           pos Ref Strain A G C T SNP exon indel nrf rat
# 1
     Chr1 1244 G
                           3 30
                                   Ω
                                      0
                                          O TRUE FALSE
# 2
                                          0
# 3
                     1012 5 54
1013 15 47
                                   0
                                      0
                                             TRUE FALSE
# 4
                                   Ω
                                       Ω
                                          1
                                              TRUE FALSE
# 5
                     1014
                           3 30
                                   0
                                       0
                                          0
                                             TRUE FALSE
# 6
                     1015 21 68
                                   0
                                       0
                                           1
                                              TRUE FALSE
                           0 10
                                      0
                                          0 TRUE FALSE
1 TRUE FALSE
                     3367
                                   0
                     1335 108 108
                                   0
                                      0
# 9
     Chr1 1575
# 10
                     1007 26 11
                                   0
                                       0
                                          O TRUE FALSE
# 11
                     1012 49 27
                                   0 0
                                          O TRUE FALSE
# 12
                     1013 19 28
                                   0
                                      0
                                          0
                                              TRUE FALSE
# 13
                     1014 15 17
                                   0
                                      0
                                          0
                                             TRUE FALSE
# 14
                     1015 46
                               42
                                   0
                                       0
                                              TRUE FALSE
# 15
                     3367
                           0 11
                                   0 0
                                          0
                                             TRUE FALSE
# 16
                     1335 37 99
                                   0
                                      0
                                          0 TRUE FALSE
# 17 Chr1 6485 G
                     1007 26 20
                                   0
                                       0
                                          0
                                             TRUE FALSE
                     1012 33 39
                                   0 0
                                          0
                                             TRUE FALSE
# 20
                     1013
                           54
                               48
                                   0
                                       0
                                          0
                                              TRUE FALSE
# 21
                     1014 13 10
                                       0
                                          0
                                             TRUE FALSE
                     1015 34
                                              TRUE FALSE
                              41
# 23
                     3367
                           0 42
                                   0
                                      0
                                          0 TRUE FALSE
                     1335 71 69
                                   0
                                       0
                                          0 TRUE FALSE
    Chr1 7181 G
                            0 37
                                             TRUE FALSE
                     1012
                           0 66
                                             TRUE FALSE
# 28
                     1013
                               30
                                              TRUE FALSE
                     1014
# 29
                            0 19
                                       0
                                          0
                                             TRUE FALSE
# 30
                     1015
                            0 44
                                  33
                                       0
                                              TRUE FALSE
                           0 33
# 31
                     3367
                                             TRUE FALSE
                           0 94 78
# 32
                     1335
                                       0
                                          0 TRUE FALSE
# 33
     Chr1 7220 C
# 34
                     1007 17
                               0 26
                                           0
                                             TRUE FALSE
# 35
                     1012 45
                               0 31
                                      14
                                              TRUE FALSE
                                          0
# 36
                     1013 112
                                  41
                                      14
                                           0
                                              TRUE FALSE
# 37
                     1014 16
                               1 16
                                              TRUE FALSE
                                          0
# 38
                               0 26
                                              TRUE FALSE
                     1015 66
                               0 24
# 39
                     3367
                                      0
                                          0
                                             TRUE FALSE
                               0 51 25
                                          O TRUE FALSE
# 40
                     1335 68
# 41
     Chrl 7661 T
                     1007
                            0
                               0 10
                                           0
                                             TRUE FALSE
# 42
                                      14
                     1012
                               0
                                              TRUE FALSE
# 43
                           0
                                      24
                                          0
                                  32
# 44
                     1013
                                      23
                                              TRUE FALSE
                            0
                               0
                     1014
                                              TRUE FALSE
# 45
                            0
                               0
                                      11
                                           0
                                   8
                     1015
                                             TRUE FALSE
# 46
                            0
                               0
                                   6 41
                                           0
# 47
                     3367
                            0
                               0
                                  0
                                          0
                                             TRUE FALSE
                                   8 42
# 48
                     1335
                           0
                               0
                                          O TRUE FALSE
# 49 Chrl 8144 G
                     1007 10
                                   0
                                           0
                                             TRUE FALSE
# 50
                              16
                                      0
# 51
                     1012
                                              TRUE FALSE
                           19
                               2.8
                                   0
                                          1
# 52
                     1013 63
                               67
                                   0
                                      0
                                          0
                                              TRUE FALSE
# 53
                     1014
                                   Ω
                                      0
                                          0
                                              TRUE FALSE
# 54
                     1015 18 28
                                   0
                                      0
                                          0
                                             TRUE FALSE
                                      0
# 55
                     3367
                           Ω
                                   0
                                          0
                                             TRUE FALSE
# 56
                     1335 17 58
                                   0 0
                                          1 TRUE FALSE
# 57
     Chr1 8208 G
# 58
                            0 15
                                   0
                                       8
                                           1
                                             TRUE FALSE
# 59
                            0
                              2.8
                                   0 16
                                           0
                                              TRUE FALSE
# 60
                     1013
                            0 24
                                   0 63
                                          0
                                             TRUE FALSE
# 61
                     1014
                            0 15
                                   0
                                      4
                                           0
                                              TRUE FALSE
                                          1
# 62
                     1015
                           0 25
                                   0 13
                                             TRUE FALSE
# 63
                     3367
                           0
                                   0
                                          0
                                             TRUE FALSE
# 64
```

```
# 65 Chr1 8518 T
                      1007 0 0 20 18
                                            1 FALSE FALSE
# 67
                      1012
                            0
                                0 45
                                       30
                                            1 FALSE FALSE
# 68
                             0
                                0
                                    57
                                            1 FALSE FALSE
                      1014
                             0
                                0 10
                                        32
                                            O FALSE FALSE
                      1015
                             0
                                 0
                                            1 FALSE FALSE
                                   41
# 71
                      3367
                             0
                                0 0 11
                                            O FALSE FALSE
                      1335
                             0
                                0 120
                                            1 FALSE FALSE
 73
     Chr1 8552
                      1007
                             3
                                13
                                     0
                                         0
                                            0
                                               TRUE FALSE
                      1012 21 31
                                               TRUE FALSE
                      1013
                            33
                                                TRUE FALSE
 77
                      1014
                                15
                                                TRUE FALSE
 78
                                                TRUE FALSE
                      1015 14
                                22
                                     0
                                         0
                                            0
                      3367 0 28
                                               TRUE FALSE
                                     0
                      1335 27
                                               TRUE FALSE
                                     0
                                         0
     Chr1 8567
# 82
                      1007
                            16
                                18
                                     0
                                         0
                                            1
                                                TRUE FALSE
# 83
                      1012 34
                                35
                                     0
                                         0
                                                TRUE FALSE
                                            1
 84
                      1013
                           66
                                     0
                                                TRUE FALSE
 85
                      1014
                                     0
                                         0
                                            0
                                                TRUE FALSE
# 86
                      1015 17
                                31
                                     0
                                         0
                                                TRUE FALSE
                      3367 29
# 87
                                0
                                     0
                                       0
                                            0
                                               TRUE FALSE
                      1335 59
                                44
                                     0
                                        0
                                               TRUE FALSE
 88
# 89
     Chr1 8670
# 90
                      1007
                            19
                                 0
                                     0
                                            0
                                                TRUE FALSE
                      1012 36
                                     0 12
# 91
                                0
                                            0
                                                TRUE FALSE
                                                TRUE FALSE
# 92
                      1013 44
                                 0
                                     0
                                        12
                                            0
# 93
                      1014 10
                                                TRUE FALSE
                                 0
                                     0
                                            0
# 94
                      1015 24
3367 18
                                               TRUE FALSE
                                0
                                     0
                                       11
                                            1
# 95
                                            0
                                               TRUE FALSE
                                0
                                     0
                                        0
                                               TRUE FALSE
 96
                      1335 27
                                 0
                                     0
                                        6
                                            0
     Chr1 8685
# 97
# 98
                      1007
                                16
                                     0
                                         0
                                            0
                                               TRUE FALSE
                      1012 12
# 99
                                                TRUE FALSE
                                37
                                     0
                                         0
                                            0
# 100
                      1013 18
                                     0
                                         0
                                            1
                                                TRUE FALSE
# 101
                      1014
                                32
                                     0
                                            0
                                                TRUE FALSE
                            5
                      1015 11 35
3367 0 12
                                           1
# 102
                                     0
                                         0
                                               TRUE FALSE
                                       0
# 103
                                     Ω
                                               TRUE FALSE
# 104
                      1335 5 45
                                     0
                                         0
                                            0 TRUE FALSE
# 105 Chr1 14361
                      1007 29
                                 7
                                     0
                                         Ω
                                            O FALSE FALSE
# 106
# 107
                      1012 54
                                     0
                                         0
                                            O FALSE FALSE
                                12
# 108
                      1013 28
                                     0
                                         0
                                            1 FALSE FALSE
# 109
                      1014 22
                                2
                                     1
                                         0
                                            0 FALSE FALSE
                      1015 51
3367 12
                                       0
                                           0 FALSE FALSE
0 FALSE FALSE
 110
                                9
                                    0
# 111
                                1
                                     0
# 112
                      1335 64
                                8
                                     0
                                        0
                                            O FALSE FALSE
# 113 Chr1 15254 T
 114
                      1007 11
                                0
                                    0
                                            1 FALSE FALSE
 115
                      1012 28
                                0
                                    0
                                       53
                                            1 FALSE FALSE
 116
                      1013 39
                                0
                                     0 66
                                            1 FALSE FALSE
 117
                      1014
                                 0
                                     0
                                       14
                                            1 FALSE FALSE
 118
                      1015 18
                                0
                                    0 39
                                            1 FALSE FALSE
 119
                      3367
                            0
                                0
                                    0 89
                                            0 FALSE FALSE
 120
                      1335 15
                                0
                                     0 63
                                            1 FALSE FALSE
 121 Chr1 15280
 122
                             0 14
                                     0 32
                                            1 FALSE FALSE
                      1012
                             0
                                31
                                     0 53
                                            1 FALSE FALSE
 124
                      1013
                             0
                                     0 102
                                            0 FALSE FALSE
 125
                      1014
                             0
                                       40
                                            O FALSE FALSE
                      1015
                            0 22
                                            1 FALSE FALSE
 126
 127
                      3367
                            0
                                0
                                     0 74
                                            O FALSE FALSE
                                26
                                     0 109
                                            1 FALSE FALSE
 128
                      1335
 129 Chrl 16103
                      1007 12
                                 0
                                            1 FALSE FALSE
                      1012
                                            1 FALSE FALSE
 131
                            50
                                         0
                      1013
                            29
                                            1 FALSE FALSE
 132
                      1014 28
 133
                                 0
                                         0
                                            0 FALSE FALSE
                                            1 FALSE FALSE
 134
                      1015 37
                                0 10
                                         0
                      3367 41
                                            0 FALSE FALSE
 135
                                    0
                                         0
                                    12
                                            0 FALSE FALSE
 136
                      1335 56
                                 0
                                         0
# 137 Chr1 17587
                      1007 22
                                            O FALSE FALSE
 138
                      1012 62
                                            O FALSE FALSE
 139
                                     0
 140
                      1013 22
                                12
                                     0
                                            1 FALSE FALSE
# 141
                      1014 22
                                2
                                         0
                                            O FALSE FALSE
                                     0
                      1015 29
# 142
                                     0
                                         0
                                            O FALSE FALSE
                      3367 20
                                            O FALSE FALSE
# 143
                                     0
                                        0
                      1335 82
                               11
                                       0 0 FALSE FALSE
# 144
                                    0
```

```
# 145 Chr1 18904 T
                                    0 34
                                            0 FALSE FALSE
                      1012
                                             O FALSE FALSE
                                     0
                                    0 21
 149
                      1014
                                             0 FALSE FALSE
                       1015
                                     0
 151
                       3367
                                             0 FALSE FALSE
                                27
                                     0 73
                       1335
                                             1 FALSE FALSE
 153 Chr1 25546 A
                       1007
                            31
                                     0 14
                                             1 FALSE FALSE
                       1012
                                             1 FALSE FALSE
 157
                      1014
                                        18
                                             1 FALSE FALSE
# 158
                      1015
                            64
                                        18
                                             1 FALSE FALSE
# 159
                       3367
                                             O FALSE FALSE
# 160
                      1335
                            80
                                     0
                                             0 FALSE FALSE
```

More sanity: there are 83 sites on Chr1 shared by zero strains in the tightest condition. (I.e., SAMTOOLS called it a SNP, but the read counts/proportions fall below our 3rd threshold). Are they due to low coverage? Seemingly yes:

```
zp3 \leftarrow snp.pattern[[3]] == 0
zr3 <- rownames(non.refs[[3]])[which(zp3)]</pre>
zc3 <- as.integer(unlist(lapply(strsplit(zr3[1:min(100,length(zr3))],':',fixed=TRUE),function(x){x[2]})))</pre>
zc3
# [1]
       91284 127986 161271 196862 196864 199166 282391 289344 289363 314132 314661
       438976 447253
                       475823
                               501830 501975
                                               504462 652889 657955 692139 709443
       826899 856950 875379 913014 938651
                                              967184 1036942 1100300 1113225 1181146 1203203
 [34] 1210360 1212223 1224082 1270250 1270251 1348311 1431628 1473437 1516083 1526912 1628300
 [45] 1637082 1686331 1736789 1763837 1782580 1967158 2024930 2075603 2098145 2110716 2194162
# [56] 2242316 2258647 2261176 2325671 2376777 2432898 2441781 2498706 2550796 2554565 2581374
# [67] 2614631 2619528 2659281 2675254 2691279 2703771 2737914 2744068 2802553 2842231 2846930
# [78] 2906880 2931365 2948653 2957936 3014028 3016252
seecounts(zc3[1:5], snp.tables=snp.tables)
            pos Ref Strain A G C T SNP exon indel nrf rat
     chr
    Chr1 91284
                      1007
                             0 0 0 17
                                        0 FALSE FALSE
# 3
                      1012
                             0 0 0 38
                                         O FALSE FALSE
                      1013
                             2 0 0 13
                                         O FALSE FALSE
# 5
                      1014
                             0 0
                                 0 20
                                         O FALSE FALSE
                      1015
                             0 0
                                  0 35
                                         O FALSE FALSE
                                  0 12
                      3367
                             3 0
                                         1 FALSE FALSE
                      1335
                             0 0 0 47
                                         O FALSE FALSE
# 9 Chrl 127986
                            47 0 0 0
                      1007
                                         O TRUE FALSE
# 10
                      1012
                            92 0
                                  0
                                         0
# 11
                                     0
                                            TRUE FALSE
                            19 1
# 12
                      1013
                                            TRUE FALSE
                                 0
                                     0
                                         0
# 13
                      1014 73 0 0 0
                                         0
                                           TRUE FALSE
# 14
                      1015 83 0 0 0
                                        O TRUE FALSE
                      3367 13 3 0
                                            TRUE FALSE
# 15
                                     0
                                         1
                      1335 160 0
                                  0
                                     0
                                         0
                                            TRUE FALSE
# 17 Chr1 161271
                      1007 31 0 0 0
                                         0
                                           TRUE FALSE
# 19
                      1012 47 0 0 0
                                         0 TRUE FALSE
# 20
                      1013
                            18 3
                                  0
                                     0
                                         0
                                            TRUE FALSE
# 21
                      1014
                            30 0
                                  0
                                     0
                                         0
                                            TRUE FALSE
                      1015 59 0 0
# 2.2
                                     0
                                         0
                                            TRUE FALSE
# 23
                      3367
                            8 3 0 0
                                        1
                                            TRUE FALSE
                      1335 102 0 0 0
                                        O TRUE FALSE
# 24
# 25 Chr1 196862
# 26
                      1007
                             0 0 10
                                     0
                                         O FALSE FALSE
# 27
                             0 0 22
                      1012
                                    0
                                         O FALSE FALSE
                             0 0 8 2
# 28
                      1013
                                         0 FALSE FALSE
# 29
                             0 0 14 0
                                         0 FALSE FALSE
                      1014
 30
                      1015
                             0 0 18
                                     0
                                         O FALSE FALSE
# 31
                      3367
                             1 0 4
                                     3
                                         1 FALSE FALSE
                      1335
                            0 0 18 0 0 FALSE FALSE
# 32
```

```
# 33 Chr1 196864 T
                        0 0 0 11 0 FALSE FALSE
# 34
                   1007
                        0 0 1 23
3 0 0 8
# 35
                   1012
                                   O FALSE FALSE
                                  1 FALSE FALSE
# 36
                   1013
# 37
                   1014 0 0 0 12
                                  0 FALSE FALSE
# 38
                   3 0 0 4 1 FALSE FALSE
# 39
                   3367
# 40
                   1335
                        0 0 1 19 0 FALSE FALSE
```

7.3 Main Analysis

Turning to the main analysis, there is a large increase in the number of consistent positions between the loose and medium stringency levels; medium and tight are similar in most respects. The likely interpretation is that the loose criterion is including many "SNPs" induced by read errors, and that either of the tighter criteria are successfully filtering them out. In the interest of simplicity, the narrative below will focus on the shared SNPs at the medium stringency level (the "count2" column in the data frame), although the numbers for all three (sometimes all 4) are displayed. Also note that the prose and some comments in the code were based on the Chr1 analysis, and so may occasionally be off-target for the whole-genome data.

```
# Show a subset of pat.summaries, optionally with totals of count_i in last row, and optionally
# aggregating low-count rows as ``Other''
   sharedBy=c(2,4) selects SNPs shared by 2 or 4 strains,
   subset-as.octmode('35') select those with sharing pattern a subset (optionally proper) of this
    c2.thresh=42 suppresses printout of rows with count2 < 42
   restrict.to=c(0,42,127) restrict to these 3 rows
showgroup <- function(p.summ=pat.summaries, sharedBy=0:7, subset=127, split=NULL, proper.subset=F,
                      total=T, c2.thresh=0, fourteenth=F, restrict.to=NULL) {
  # pick just those bit patterns that are subsets of 'subset'
  pick <- bitwAnd(0:127,bitwNot(subset))==0</pre>
  if (proper.subset) {
    pick[subset+1] <- F
  if(!is.null(split)){ # AND that stradle left/right subtrees
   cosplit <- bitwAnd(subset,bitwNot(split))</pre>
    pick <- pick & bitwAnd(0:127,split)!=0 & bitwAnd(0:127,cosplit)!=0</pre>
  # and have desired shareBy counts
 pick <- pick & (p.summ$sharedBy %in% sharedBy)</pre>
  # and are among the set of interest
  if(!is.null(restrict.to)){
   pick <- pick & (0:127 %in% restrict.to)
  # find rows with low counts
 pick.low <- pick & (p.summ$count2 < c2.thresh)
  # now show them
  show <- p.summ[pick & ! pick.low,]</pre>
  # rename columns just to narrow the printouts
  colnames(show) <- c('Pat','ShrBy','1007', '1012', '1013', '1014', '1015', '3367', '1335',</pre>
                       'count1', 'count2', 'count3', 'count4')
  show[,1] <- format(show[,1]) # convert octal col to char so can override in last row(2)</pre>
  nlow <- sum(pick.low)</pre>
  if(nlow > 0){
   n <- nrow(show)+1
   lows <- apply(p.summ[pick.low, 10:13], 2, sum)</pre>
   show[n,10:13] <- lows
   show[n,1:9] <- ''
    row.names (show) [n] <- 'Other'</pre>
    if(fourteenth){
     # do this: add 14th col just to hold this comment:
      show <- cbind(show,' '='', stringsAsFactors=F)</pre>
      show[n,14] <- paste('(', nlow, 'rows w/ c2 <', c2.thresh, ')')</pre>
   } else {
```

```
## or this (looks a bit funky, but fits across page without line-wrap):
    show[n,1:8] <-c('(', nlow, 'rows', 'w/', 'c2', '<', c2.thresh, ')')
}
if(total){
    n <- nrow(show)+1
    tots <- apply(show[,10:13],2,sum)
    show[n,10:13] <- tots
    show[n,10:13] <- tots
    show[n,1:9] <- ''
    row.names(show)[n] <- 'Total'
    if(ncol(show)==14){show[n,14]<-''}
}
return(show)
}</pre>
```

First, are there any SNPs that are not "consistent SNPs?" Yes, a few in c3. As noted above, they seem to be mainly low-coverage positions.

```
showgroup(pat.summaries,0,total=F) # chr1 totals: 0 0 83

# Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 1 0 0 0 83 0
```

Next, look at completely shared SNPs, those found in all 7 strains.

I.e., of the 46896 consistent positions, 7054 or 15% are shared by all 7 strains.

Next look at singletons, aka private SNPs—SNPs that are called in one strain and no other strain has a significant number of non-ref reads at that position. Presumably these are variants that arose in a given population after it separated from the others.

```
showgroup(pat.summaries,1) # chr1 totals: 9669 18865 19670 23574
       Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 2
                                                            41
#
 .3
                                                      4551
                                                             8813
                                                                    9159
                                                                         10949
       004
                                                       90
                                                            207
                                                                    243
                                                                          385
                                        X
# 9
       010
                                   X
                                                             61
                                                                    7.8
                                                                           113
                                                      4954
                                                             9652
                                                                    9985
# 17
       020
               1
                                                                         11677
                              Χ
       040
                                                       29
                                                                    95
# 33
                                                              61
                                                                           174
# 65
                                                        1.0
                                                               30
                                                                     40
               1
                                                                           141
# Total
                                                      9669 18865 19670 23574
```

The import of shared/private SNPs changes between sexual and asexual populations. Presumably asexuals slowly gain and rarely lose private SNPs; shared ones predate separation of the lineages. In sexual lineages, however, SNPs may be rather freely "gained" or "lost," merely by recombination (converting between homo- and heterozygous in the sample we sequenced). Thus, the low private counts for the 5 L-isolates compared to the large count of het positions overall suggest that (a) they are asexual, and (b) none of them has been isolated from the others for very long (if at all). Conversely, the high counts for Italy and Wales suggest that (a) if asexual, they have been separated from each other and from the rest for a long time, but (b) if sexual, there is little surprise: we have $\approx 160 \text{K}$ SNPs shared between the two (90K just in those two (below), plus 70K shared by all 7), and $\approx 90 \text{K}$ additional positions that are het in one but not the other. These are close to, but not exactly equal to, the 1:2:1 ratios we would naively expect from two samples of a single HWE population. The most parsimonious explanation seems to be that the H-clade is sexual, but perhaps some het positions private to each population separates them.

Aside: counts of "consistent" SNPs minus these singletons yeilds count of shared SNPs:

The slightly higher count of shared positions in the medium case further supports this choice for subsequent analysis.

Next look at consistent SNPs shared between just a pair of isolates.

	harramar	· / / / /	+ a		2 2)	# aba	1 00	+ a .	7611	0.57	10 01	72 602	1	
S	howgrou	.b (bs	at.Sumr	naries	5, 4)	# Cnr	1 CO	unts:	/041	954	9 94	72 692	±	
#		Pat	ShrBy	1007	1012	1013	1014	1015	3367	1335	count1	count2	count3	count4
#	4	003	2						X	X	239	16	28	31
#	6	005	2					X		X	11	18	35	52
#	7	006	2					X	Х		141	13	21	41
#	10	011	2				X			X	6	7	7	14
#	11	012	2				X		Х		179	18	2	5
#	13	014	2				X	Х			10	2	5	12
#	18	021	2			Х				X	243	9	11	9
#	19	022	2			Х			Х		5920	9365	9094	6177
#	21	024	2			X		X			125	12	28	46
#	25	030	2			X	X				222	18	5	9
#	34	041	2		X					X	3	4	13	36
#	35	042	2		X				X		150	1	16	27
#	37	044	2		X			Х			5	13	75	155
#	41	050	2		X		X				7	2	5	7
#	49	060	2		X	X					165	4	26	28
#	66	101	2	Х						X	1	2	13	20
#	67	102	2	X					X		93	6	7	25
#	69	104	2	Х				Х			5	9	34	107
#	73	110	2	Х			X				2	2	4	7
#	81	120	2	X		X					105	9	9	31
#	97	140	2	X	X						9	19	34	85
#	Total										7641	9549	9472	6924

I.e., of the 9549 paired SNPs, 9365 or 98.1% are found between Italy and Wales, with comparatively few shared between any other pairs (only).

SNPs shared among exactly 3 isolates are relatively rare. (The 5 trios containing both Italy and Wales predominate in the loose set, probably because they share many pairs that become triples with the addition of a few read errors.)

	1	/	. 1		. 21	// . 7	7		1 1 2 0	0.0	1	71 100	Λ	
S	howgro	up (pa	at.sumr	narıe	S,3)	# chr	1 CO	unts:	1438	29	4 6	71 103	4	
#		Dat	ShrBu	1007	1012	1013	101/	1015	3367	1335	count 1	count?	count3	count /
#		007	3	1007	1012	1013	TOTA	X		1333 X	9	2	9	10
#		013	3				Х		X	X	17	3	2	2
			-									_		2
#		015	3				X			X	7	6	4	/
#		016	3				Х	X			9	1	2	2
#	20	023	3			X			X	X	327	20	29	17
#	22	025	3			X		X		X	9	4	12	21
#	23	026	3			X		X	X		185	27	31	32
#	26	031	3			X	X			X	20	2	0	0
#	27	032	3			X	X		X		324	18	8	5
#	29	034	3			X	X	X			11	3	1	1
#	36	043	3		X				X	X	21	8	14	6
#	38	045	3		X			X		X	6	26	130	131
#	39	046	3		X			X	X		11	12	34	55
#	42	051	3		X		X			Х	0	1	2	4
#	43	052	3		X		X		X		9	0	2	1
#	45	054	3		X		X	X			1	6	17	18
#	50	061	3		Х	Х				Х	12	2	14	12
#		062	3		X				Х		227	17	37	36
#		064	3		X			Х			9	9	36	60
- 11		- 0 -	0								_	_	0 0	0.0

#	57	070	3		Х	Х	Х				13	1	1	2
#	68	103	3	X					X	Х	11	4	4	8
#	70	105	3	X				Х		Х	4	5	25	63
#	71	106	3	Х				Х	Х		3	9	15	27
#	7.4	111	3	Χ			Χ			X	1	0	1	1
#	7.5	112	3	X			X		X		4	1	0	0
#	77	114	3	X			X	X			1	2	2	8
#	82	121	3	X		X				X	8	0	4	4
#	83	122	3	X		X			X		134	10	10	26
#	85	124	3	Х		X		X			11	7	20	35
#	89	130	3	Х		X	X				7	1	2	1
#	98	141	3	X	X					X	2	5	15	40
#	99	142	3	Х	Χ				X		9	1	9	15
#	101	144	3	X	X			X			6	75	167	355
#	105	150	3	Χ	Χ		Χ				0	0	4	6
#	113	160	3	X	X	X					10	6	8	23
#	Total										1438	294	671	1034

Four-way sharing is more common, but dominated by the coastal (i.e., non-Gyre) L-clade isolates. This is likely a reflection of the strong 5-way sharing among the L-clade, from which the Gyre commonly drops out due to the lower coverage/higher error rate in that sequencing run.

show	group (pa	at.sumr	maries	s,4) ;	# chr	1 coi	ınts:	564	1346	2552	3479		
	, (F.			. ,									
#	Pat	ShrBy	1007	1012	1013					count1		count3	count4
# 16	017	4				X	X	X	X	1	5	1	2
# 24	027	4			X		X	X	X	22	14	28	24
# 28	033	4			X	X		X	X	30	2	4	6
# 30	035	4			X	X	X		X	4	2	1	0
# 31	036	4			X	X	X	X		12	4	1	3
# 40	047	4		X			X	X	X	2	18	42	60
# 44	053	4		X		X		X	X	1	0	2	2
# 46	055	4		X		X	X		X	6	12	29	36
# 47	056	4		X		X	X	X		3	2	2	5
# 52	063	4		X	X			X	X	18	11	24	21
# 54	065	4		X	X		X		X	9	17	41	48
# 55	066	4		X	Х		Х	X		25	37	102	76
# 58	071	4		X	Х	X			X	2	1	2	0
# 59	072	4		Х	X	X		Х		8	6	3	2
# 61	074	4		X	Х	X	Х			2	3	2	7
# 72	107	4	X				X	Х	X	5	5	11	16
# 76	113	4	X			X		Х	X	1	2	0	1
# 78	115	4	X			X	X		X	0	4	3	9
# 79	116	4	X			X	X	Х		0	1	0	5
# 84	123	4	Х		X			Х	X	10	9	8	8
# 86	125	4	X		X		X		X	4	3	13	16
# 87	126	4	X		X		X	Х		12	20	21	43
# 90	131	4	X		X	X			X	0	1	0	2
# 91	132	4	Х		X	X		Х		9	1	2	1
# 93	134	4	Х		X	X	X			2	2	1	3
# 10	0 143	4	X	Х				Х	X	1	3	5	20
# 10:	2 145	4	X	Х			X		X	320	1005	1973	2585
# 10	3 146	4	Х	X			X	Х		7	34	65	140
# 10	6 151	4	X	Х		X			X	1	4	5	14
# 10	7 152	4	X	Х		X		Х		2	1	1	4
# 10	9 154	4	X	Х		X	X			16	53	49	103
# 11	4 161	4	Х	X	X				X	9	5	9	18
# 11	5 162	4	Х	X	X			Х		11	12	21	33
# 11		4	Х	X	X		X			8	47	80	163
# 12	1 170	4	Х	Х	X	X				1	0	1	3
# To	tal									564	1346	2552	3479

Five-way sharing is much more common, and is strongly dominated by the 5 L-clade isolates.

```
showgroup(pat.summaries,5) # chr 1 counts: 3969 5047 4624 6125
# Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
```

8 TREES 35

#	32	037	5			Х	Х	Х	Х	Χ	17	10	8	5
#	48	057	5		X		Х	Х	X	Χ	5	7	9	17
#	56	067	5		X	Х		Х	Х	X	33	78	201	104
#	60	073	5		X	X	X		X	X	3	3	6	3
#	62	075	5		X	Х	X	X		X	7	11	13	11
#	63	076	5		X	X	X	X	X		9	11	17	7
#	8.0	117	5	X			X	X	X	X	0	2	1	7
#	88	127	5	X		X		X	X	X	11	17	30	47
#	92	133	5	X		X	X		X	X	3	2	0	0
#	94	135	5	X		X	X	X		X	2	1	3	7
#	95	136	5	X		X	X	X	X		4	6	3	5
#	104	147	5	X	X			X	X	X	125	307	590	1160
#	108	153	5	X	X		X		X	X	3	3	1	7
#	110	155	5	X	X		X	X		X	3484	3912	2642	3228
#	111	156	5	X	X		X	X	X		8	15	15	43
#	116	163	5	X	X	X			X	X	12	18	23	33
#	118	165	5	X	X	X		X		X	201	453	787	1140
#	119	166	5	X	X	X		X	X		30	157	247	254
#	122	171	5	Х	X	X	X			X	3	6	2	7
#	123	172	5	X	X	X	X		X		2	6	3	5
#	125	174	5	X	X	X	X	X			7	22	23	35
#	Total										3969	5047	4624	6125

Six-way sharing is also common, with the sets *excluding Gyre*, Italy, or Wales having the most mutually-shared SNPs.

S	<pre>showgroup(pat.summaries,6) # chr 1 counts: 4166 4741 5312 4722</pre>													
#		Pat	ShrBy	1007	1012	1013	1014	1015	3367	1335	count1	count2	count3	count4
#	64	077	6		X	X	X	X	X	X	43	48	60	26
#	96	137	6	X		X	X	X	X	X	11	6	12	14
#	112	157	6	X	X		X	X	X	X	1343	1192	839	1343
#	120	167	6	X	X	X		X	X	X	951	1879	3320	1852
#	124	173	6	X	X	X	X		X	X	17	9	7	3
#	126	175	6	X	X	X	X	X		X	1756	1493	997	1416
#	127	176	6	X	X	X	X	X	X		45	114	77	68
#	Total										4166	4741	5312	4722

8 Trees

So, overall, the picture looks like a long shared history (7054 7-way shared positions), followed by a split of the 5 L-isolates from the 2 H-isolates, then a long shared history in the 5 (3912 quintuples), in parallel with a long shared history in H- (9365 pairs), then separate histories in Italy and Wales (>8813 "private" SNPs in each, although again if they are sexual, many of these just reflect HWE), and very limited differentiation among the 5 L-isolates.

Branch lengths of course depend on filtering criteria used (and, of course, full vs Chr1 differ by about a factor of 10), but the tree *topology* appears to be fairly stable. Various versions are drawn below, exactly to explore how robust this story is. I think we should go with "medium stringency" SNP filtering (based on un-qfiltered reads).

NOTE: Much of this analysis make less sense for q-filtered read data, since (a) the point of the SNP filtering was to try to correct for noise in the raw reads, which may (or may not; haven't looked closely, yet) be largely fixed by qfiltering (e.g., "loose" or no SNP filtering may be more appropriate, post-q-filtering, esp. if we had re-run SAMTools to call SNPs based on the q-filtered reads), and (b) tree topology *does* appear to change, in that Gyre's coverage has been so sharply reduced by qfiltering that it clearly stands aside from the others (and that's confirmed by bootstrap), but this also seems to be clearly a technical rather than a biological artifact. SO, code below will run on q-filtered data, but *is not tuned to it*. Likewise, most comments in the prose below were made to describe the un-q-filtered data, and *are misleading and in some cases flatly wrong* for qfiltered data, but it doesn't seem worthwhile to bother with a rewrite...

Trees are coded in newick format, which doesn't seem to tolerate line-breaks; print with line-wrap:.

8 TREES 36

```
# wrap a long char string across multiple lines in printout
cat.hardwrap <- function(str,width=80){
    while(nchar(str)>width){
        cat(substr(str,1,width),'\n')
        str <- substr(str,width+1,nchar(str))
    }
    cat(str,'\n')
}</pre>
```

Trees are built as follows. Code for drawing, especially, is specific to the topology of the medium tree, and placement of some of the figure elements have been hand-optimized for this case; drawings for the other variants will not be as pretty.

```
# set up for tree figs
# the newick parser in ape seems to be confused by commas and parens in
# tip names, and blanks are not allowed, so replace by *, <, >, _, resp.
newick.name <- function(name) {</pre>
 name <- gsub(' ', '_', name, fixed=TRUE)
name <- gsub(',', '*', name, fixed=TRUE)
  name <- gsub('(', '<', name, fixed=TRUE)
name <- gsub(')', '>', name, fixed=TRUE)
  return (name)
# undo above changes
newick.name.undo <- function(name) {
#name <- gsub('_', ' ', name, fixed=TRUE) # unnecessary; ape plot routine handles this one
name <- gsub('*', ',', name, fixed=TRUE)
name <- gsub('<', '(', name, fixed=TRUE)
name <- gsub('>', '(', name, fixed=TRUE)
name <- gsub('>', ')', name, fixed=TRUE)
  return(name)
# make a newick string from tree; see it below
# 'pre' is prefixed to ccmpid; 'nb' optionally included;
# 'alt' can be used instead of pre/ccmp/nb/where for less formal labeling
# 'newstyle'==T => new node label: [nb_]where[(pre-less-id)]
# 'newstyle'==F => old node label: [nb_][pre id]where
newickize <- function(tree,pre='CCMP',nb=TRUE,alt=F,newstyle=TRUE) {</pre>
  if(is.null(tree$where)){
                               ogether newick from subtrees
     \verb|sub1| <- \verb|newickize| (tree\$sub1, pre=pre, nb=nb, alt=alt, newstyle=newstyle)|\\
     sub2 <- newickize(tree$sub2,pre=pre,nb=nb,alt=alt,newstyle=newstyle)</pre>
     new <- paste( '(', sub1, ',', sub2, ')', sep='')</pre>
     if(!is.null(tree$length)){
       # internal node, add length
       return(paste(new, ':', tree$length, sep=''))
    return(paste(gsub(' ', '_', new), ';', sep=''))
       # label is
                     just alt; if alt omitted, default to where
       new <- newick.name(ifelse( is.null(tree$alt), tree$where, tree$alt ))</pre>
     } else {
       if(newstyle){
          # new node
          new <- ifelse( nb && !is.null(tree$nb), paste(tree$nb, '_', sep =''), '' )</pre>
          new <- newick.name(paste(new, tree$where, sep=''))</pre>
          new <- ifelse( is.null(tree$id), new, paste(new, '_(', tree$id, ')', sep='') )</pre>
          new <- newick.name(new)</pre>
       } else {
          # old style node label = [nb_][pre id]where
          new <- ifelse( nb && !is.null(tree$nb), paste(tree$nb, '_', sep =''), '' )</pre>
         new <- ifelse( is.null(tree$id), new, paste(new, pre, tree$id, '_', sep='') )
new <- newick.name(paste(new, tree$where, sep=''))</pre>
     #add length to either
     new <- paste(new, ':', tree$length, sep='')</pre>
  return (new)
\# Make a tree as nested lists, **based on the chr1, count2 topology**, but using any of the counts.
```

```
Internal nodes have subtrees sub1/2 and length
    Root has sub1/2, but no length
    Leaves have where, length, optionally, id, alt, nb. (Omit id for 'outgroup'. Use 'alt' for less formal labeling in cartoon version; it defaults to 'where'. Use 'nb' to add abode annotations for legend.)
\# The single parameter v is any of the 4 count vectors contained in pat.summaries (most conveniently
  indexed in octal). E.g., make.tree(pat.summaries[,'count2']) reproduces the count2 tree.
  (This was previously built by hand-pasting the edge lengths; tree.by.hand is retained in appendix
# for comparison, & its counts are in comments below).
make.tree <- function(v) {</pre>
  pat.count <- function(pat, pat.counts=v){return(pat.counts[1+strtoi(pat,8)])}</pre>
    list(
       sub1 = list(
         sub1 = list(
           sub1 = list(id=3367, length=pat.count('002'), where='Venice, Italy', alt='Venice'), #8813
            sub2 = list(id=1013, length=pat.count('020'), where='Wales, UK'),
            length=pat.count('022')),
                                                                                                                    #9365
         sub2 = list(
            sub1 = list(
              sub1 = list(
                sub1 = list(id=1007, length=pat.count('100'), nb='e', where='Virginia, USA'), #30
sub2 = list(id=1012, length=pat.count('040'), nb='d', where='Perth, W. Australia', alt='Perth'), #61
                 length=pat.count('140')),
               sub2 = list(
                 sub1 = list(id=1015, length=pat.count('004'),nb='c', where='Washington, USA', alt='Puget Sound'), #207
sub2 = list(id=1335, length=pat.count('001'), nb='b', where='New York, USA', alt='NY'), #41
                 length=pat.count('005')),
                                                                                                                     #18
              length=pat.count('145')),
                                                                                                                     #1005
            sub2 = list(id=1014, length=pat.count('010'), nb='a', where='N. Pacific Gyre'),
length=pat.count('155')),
                                                                                                                     #61
                                                                                                                     #3912
          length=pat.count('177')),
                                                                                                                      #7054
       sub2 = list(length=0, where='outgroup')
  return (thetree)
```

Code to plot a tree given newick description. Again, code is somewhat general, but has some specializations tied to the medium-stringency, full-genome, un-qfiltered data.

```
# run following 2 lines after an R upgrade
# update.packages()
# install.packages("ape")
library (ape)
show.tree <- function(newick.str=newick.medium,
                       col.edge ='darkblue', lwd.edge =2,
                       col.elabel='darkblue',
                                                               cex.elabel=0.8, font.elabel=3,
                       col.arrow ='red', lwd.arrow=1.5, cex.arrow =0.9, font.arrow =4, col.clade ='black', lwd.clade=1, cex.clade =1.0, font.clade =3,
                       col.legbox='beige',
                                                               cex.legend=0.8,
                       col.tip ='darkblue',
                                                                                 font.tip =4,
                       plusx=FALSE, pltdebug=FALSE, total.snps=consistent.count[2]){
  ####
  # ADJUST NEWICK & GET LENGTHS, COORDINATES
  newick.str.noout <- sub('outgroup','_',newick.str) # Hide outgroup ('_' prints as blank)</pre>
  the.tree <- read.tree(text=newick.str.noout)
  ## nasty hack: ape's newick parser seems to be confused by commas, () in tip labels, so
  ## newickize replaced them by '*<>'; before plotting, I want to convert them back, and hope
  ## this doesn't break anything else... And if a revised version of ape changes the internal
  ## representation of a tree, this may need to be redone
  the.tree$tip.label <- newick.name.undo(the.tree$tip.label)
  # extract branch lengths as char string of comma-separated numbers via pattern matching hack:
  # lengths always preceded by color
  lengths.ch <- strsplit(paste(newick.str,':'),'[^0-9][^:]*:')[[1]]</pre>
  # then convert to ints, dropping empty string at front
  lengths.int <- scan(what=integer(), quiet=T, sep=',',text=lengths.ch[-1])</pre>
  # then to data frame with named rows; a..q are terminal branches; others are internal.
  \# a..e match legend in plot; f/g = wales/italy. lengths appear in postfix order of
  # newick tree, and ape draws the 1st of them at the bottom of the plot.
  lmed <- data.frame(lengths=lengths.int,</pre>
                     row.names=c('g','f','fg','e','d','de','c','b','bc','bcde','a','abcde','all','out'))
```

```
# extract counts needed for legend:
#leg.counts <- c( 61, 41,207, 61, 30, 1005, 18, 19) #by hand, medium chr1 leg.counts <- lmed[c('a','b','c','d','e','bcde','bc','de'),1]
  discord <- total.snps - sum(lmed$lengths)</pre>
  #tree.labels <- list( ## x,y,text; coords are all picked by eye</pre>
  # 3000, 3.62, paste(lmed['all' ,1], 'shared by 7', sep='\n'), # 7054
# 8900, 5.75, paste(lmed['abcde',1], 'by 5' , sep='\n'), # 3912
  # 0900, 5.75, paste(lmed['abcde',1], 'by 5' , , sep='\n'), # 7054
# 12000, 1.50, paste(lmed['fg' ,1], 'shared by 2', sep='\n'), # 9365
# 21000, 2.00, paste(lmed['f' ,1], 'only\nin Wales'), # 9652
# 21000, 1.00, paste(lmed['g' ,1], 'only\nin Italy'), # 8813
# 11500, 4.50, 'x')
# aptomating y n'
  # automating x-placement, below; retain above for comparison...
  tip <- integer(7) # x coords of tree tips
  tip[1] <-sum(lmed[c('all','fg','g'),1])
  tip[2] <-sum(lmed[c('all','fg','f'),1])
  tip[3] <-sum(lmed[c('all', 'abcde', 'bcde', 'de', 'e'), 1])
 tip[4] <-sum(lmed[c'all','abcde','bcde','de','d'),1])
tip[5] <-sum(lmed[c'all','abcde','bcde','bc','c'),1])
tip[6] <-sum(lmed[c'all','abcde','bcde','bc','b'),1])</pre>
  tip[7] <-sum(lmed[c('all', 'abcde', 'a'), 1])</pre>
  inode <- integer(5) # x coords of (some) internal nodes</pre>
  inode[1] <- 0
                                                                          # root
                                                                          # lca of all
  inode[2] <- lmed['all',1]</pre>
  inode[5] <- sum(lmed[c('all','abcde','bcde'),1]) # lca L-clade, nonGyre
tree.labels <- list( ## x,y,text; y coords partially picked by eye
sum(inode[c(1,2)])/2, 3.62, paste(lmed['all' ,1], 'shared by 7', sep='\n'), # 7054
sum(inode[c(2,4)])/2, 5.75, paste(lmed['abcde',1], 'by 5' , sep='\n'), # 3912
sum(inode[c(2,3)])/2, 1.50, paste(lmed['fg' ,1], 'shared by 2', sep='\n'), # 9365
(inode[3]+tip[2])/2, 2.00, paste(lmed['f' ,1], 'only\nin 1013'), # 9652
(inode[3]+tip[1])/2, 1.00, paste(lmed['g' ,1], 'only\nin 3367'), # 8813
sum(inode[c(4,5)])/2, 4.35, '*')</pre>
 tree.labels <- list( ## x,y,text; y coords partially picked by eye sum(inode[c(1,2)])/2, 3.62, paste(lmed['all' ,1], 'in 7', sep='\n'), # 7054 sum(inode[c(2,4)])/2, 5.75, paste(lmed['abcde',1], 'in 5', sep='\n'), # 3912
     sum(inode[c(2,3)])/2, 1.50, paste(lmed['fg' ,1], 'in 2', sep='\n'), # 9365
    (inode[3]+tip[2])/2, 2.00, paste(lmed['f' (inode[3]+tip[1])/2, 1.00, paste(lmed['g' sum(inode[c(4,5)])/2, 4.35, '*')
                                                                       ,1], 'In 2 , Sep- \n' , " 9652
,1], 'only\nin 1013'), # 9652
                                                                       ,1], 'only\nin 3367'),
  ####
  # BOGUS PLOT
  # a messy bit: need string widths to set xlim; but strwidth needs x-scale so must plot first.
  # M plot completely invisible, overlay 2nd plot via par(new=F...) .
  # PROVISIONALLY set x.lim here at about 30% wider than tree; fine tune it for the real plot
  # based on strwidth(tip labels) below.
  provisional.tree.x.lim <- 1.3 * max(tip) # <== PROVISIONAL plot width</pre>
  plot(0,0, type='n', bty='n', xaxt='n', yaxt='n', xlab='', ylab='', xlim=c(0,provisional.tree.x.lim), ylim=c(0,7))
  tiplabel.x <- integer(7)
  for(i in 1:7){
     # see warning above about internals of the.tree; labels have '_', printed as ' '.
     tiplabel.x[i] <- tip[i]+strwidth(gsub('_',' ',the.tree$tip.label[i],fixed=T), font=font.tip)
  # visually show tip coords & max x to debug placement issues
  plt.debug <- function(tree.x.lim, tip, tiplabel.x, spx=NULL, spy=NULL) {
     if(pltdebug){ # F to hide/T to show debug
       cat('Tip labels:', paste(the.tree$tip.label,sep='',collapse='/'), '\n')
        axis(2) # useful only for placing labels
       for(i in 1:7){
          points(c(tip[i],tiplabel.x[i]),c(i,i)) # debug: do I have right tip coordinates?
       lines(rep(tree.x.lim,2),c(0,7)) # where is right edge?
       if(!is.null(spx)) {
  points(spx,spy) # show spline control points, for tweaking
```

```
plt.debug(provisional.tree.x.lim, tip, tiplabel.x)
label.end.H <- max(tiplabel.x[1:2])</pre>
label.end.L <- max(tiplabel.x[3:7])</pre>
clade.dx <- strwidth ('x') # space between clade marker line and its label
xdel <- 3*clade.dx
                          # space between labeled clade tips and marker line
tree.x.lim <- 1.03*(max(tiplabel.x)+xdel) # <== FINAL plot width
if(pltdebug){cat('Plot width hacking:', provisional.tree.x.lim, tree.x.lim, tree.x.lim/1.03/max(tip), clade.dx)}
par(new=T) # I.e., NOT starting a new plot
####
# REAL PLOT
plot (the.tree,
    x.lim = tree.x.lim,
     y.lim = c(0,7),
     font=font.tip, label.offset=100,
tip.color=col.tip, edge.color=col.edge,
                                                    # bold-italic, nudged slightly right
     edge.width=lwd.edge,
                                                    # 5th is bottleneck edge: 14th is outgroup
    edge.lty=c(1,1,1,1, 1 ,1,1,1,1,1,1,1,1,0)
lines(00+c(0,0),c(3.5,6),col='white',lwd=6)
                                                    # Hide vertical line to outgroup
axis(1, pos=0.25, at=seq(0,25,by=5)*10^round(log10(max(tip)/25)))
if(pltdebug) \{ text(tip[1]+100, 1.0, 'Venice, Italy (3367)', adj=0, font=font.tip) \}
# BOTTLENECK ANNOTATION
# spline/elipse control points (spy/y) & tweaks thereto (dx/y)
dx \leftarrow 0.01 * tree.x.lim
dy <- .04
spx <- c(7400, 7400, 9900, 10500) # by eye, chrl, for comparison
spx <- c(inode[2]+dx, inode[2]+dx, inode[4]-3*dx, inode[4]-dx)
spy <- c(3.8, 3.9, 5.6-dy, 5.6-dy)
plt.debug(tree.x.lim, tip, tiplabel.x, spx, spy)
if(T){
  #elipse version, defined by rect thru 2 middle pts of spx/y
  spf<-function(x){</pre>
    ifelse(x <= spx[2], spy[1],
           ifelse(x >= spx[3], spy[4],
                 spy[2]+(spy[3]-spy[2])*sqrt(pmax(0,1-((x-spx[3])/(spx[3]-spx[2]))^2))))
} else {
  spf <- splinefun(spx, spy, method='hyman')</pre>
serx <- seq(spx[1], spx[length(spx)], length.out=50)</pre>
sery <- spf(serx)
tailx <- spx[1]
taily <- spy[1]
headx <- spx[4]
heady <- spy[4]
arrows(headx, heady, headx+tree.x.lim*1e-3, heady, length=.1, col=col.arrow, lwd=lwd.arrow)
lines(rev(serx), rev(sery), lty=c(5,1),col=col.arrow, lwd=lwd.arrow)
bottle.txt <- "inbreeding\nLoH / LoS"
if(T){
  \text{text} ((headx+tailx)/2+(headx-tailx)*(-.01), (heady+taily)/2+(heady-taily)*(-.10),
       bottle.txt, srt=66, font=font.arrow, cex=cex.arrow, col=col.arrow)
} else {
  # experiment at wrapping text along curved path; not too pretty, but retain for now, maybe revisit
  bottlec <- strsplit (bottle, split=NULL) [[1]]
  for(i in 1:length(bottlec)){
    text(xser[i],yser[i],bottlec[i], srt=65, font=4, cex=.7, col=col.arrow)
####
# CLADE ANNOTATION
clade.L.x <- label.end.L + xdel</pre>
clade.H.x <- label.end.H + xdel
dy <-.33
```

```
lines(rep(clade.L.x,2),c(3-dy,7+dy),lwd=lwd.clade,col=col.clade)
  lines(rep(clade.H.x,2),c(1-dy,2+dy),lwd=lwd.clade,col=col.clade)
  text(clade.L.x+clade.dx,5.0, L-clade', srt=90, font=font.clade, cex=cex.clade, col=col.clade)
  text(clade.H.x+clade.dx,1.5,'H-clade',srt=90,font=font.clade,cex=cex.clade,col=col.clade)
  # parameter plusx controls whether we try to annotate b/c (+) and d/e (x) sharing in tree; I think
  # it looks cluttered, rather than adding clarity, so I vote no, but code is here, in case.
                                                                                                       "Logic,"
  # if any, for my symbol choice is that + overlaid on x looks like the * at the next level; this
  # analogy is more visible if we use pch 3/4/8 rather than Courier or Helvetica chars, but probably
  # should use same in both tree & legend, which will take a modicum of additional work.
  legend.text <- c('a: only in 1014 ',
                     'b: only in 1335
                     'c: only in 1015
                    'd: only in 1012
                    'e: only in 1007
                    '*: shared by bcde',
paste(ifelse(plusx,'+:',' '),'shared by b/c '),
                    paste(ifelse(plusx,'x:',' '),'shared by d/e ')
  legend.text <- c('a: only in 1014 ',</pre>
                    'b: only in 1335 ',
'c: only in 1015 ',
                     'd: only in 1012
                     'e: only in 1007
                     '*: in bcde
                    paste(ifelse(plusx, '+:', ' '), 'in bc
                    paste(ifelse(plusx,'x:',' '),'in de
                     'Discordant SNPs
  legend.text <- paste(legend.text, format(c(leg.counts, discord), width=4), sep=' - ')</pre>
  legend.text <- paste(legend.text,' ') # add a little more right margin in box</pre>
  opar <- par(family='mono',cex=cex.legend)</pre>
  \textbf{legend}(\texttt{'topright'}, \texttt{legend=legend.text}, \texttt{cex=cex.legend}, \texttt{inset=} \textbf{c}(0.05, 0), \texttt{bg=col.legbox}, \texttt{box.col=col.legbox})
  par (opar)
  if(plusx){
    points(tree.labels[[16]],tree.labels[[17]]+.14,pch=8,col=col.elabel)
    points(tree.labels[[16]]+200, tree.labels[[17]]+1, pch=3, col=col.elabel)
    points(tree.labels[[16]]+200,tree.labels[[17]]-1,pch=4,col=col.elabel)
  ####
  # EDGE LENGTHS
  for(i in seq(1,length(tree.labels)-ifelse(plusx, 5, 2), by=3)){
    if(F){ # T for \n in edge labels; F to remove (except "by
      text(tree.labels[[i]], tree.labels[[i+1]], tree.labels[[i+2]])
    } else {
      # points(tree.labels[[i]], tree.labels[[i+1]], pch=3,col='green') # for debugging
      \textbf{text}(\texttt{tree.labels[[i]], tree.labels[[i+1]], sub(' \n([^z])', ^ \n'), tree.labels[[i+2]]),}
           pos=3, offset=.4, font=font.elabel, col=col.elabel,cex=cex.elabel)
caption <- function(stringency, which.tables=which.snp.tables(string.val=F)) {</pre>
 caption.where <- '(UNKNOWN genome subset).'
 if(which.tables[1]=='Chr1') {caption.where <- 'on Chr1.'}
if(which.tables[1]=='full') {caption.where <- 'genome-wide.'}</pre>
  if(which.tables[1] == 'trunc') {caption.where <- 'all Chrs.'}</pre>
  cap.stringency <- c(</pre>
     loose SNP filters.',
    'medium SNP filters.
    'strict SNP filters.',
    'unfiltered SNPs.')
  cap <- paste('Tree based on', which.tables[2], 'reads and', cap.stringency[stringency],</pre>
                    `Lengths\'\' are numbers of shared/private SNPs', caption.where)
  return (cap)
```

Trees based on all four SNP filtering criteria are shown below. Their topologies are exactly the same, although the branch lengths are different. In all four, the length of the branch labeled "*" is probably inflated by lower coverage and higher error rate in 1014, which may mask further legitimate sharing between it and the other L-isolates. The branch

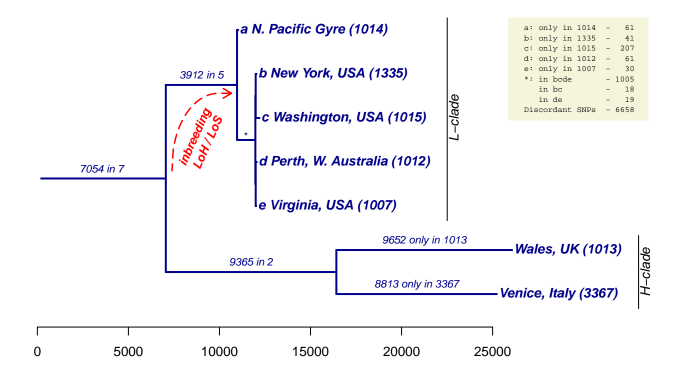


Figure 1: Proposed fig. for paper: Tree based on unfiltered reads and medium SNP filters. "Lengths" are numbers of shared/private SNPs on Chr1.

lengths among the other 4 are too short for their topology to be convincing without a more rigorous analysis (e.g., a bootstrap test), but detail there is irrelevant to the story.

My sense is that the "medium" version is the best for the paper, made here and shown in Fig 1. In theory, this should look exactly like Fig 3, but something is apparently different between Knitr and direct-to-pdf. (Increasing fig.width in Knitr's chunk headers from 8 (as in the pdf call below) to 9 helps somewhat, but probably still best to make the paper fig directly rather than via Knitr.)

```
###
#
# MAKE PDF FOR PAPER
#
if(which.snp.tables() == 'trunc-unfiltered') {
    paperfig.path <- paste('figs-mine/Fig3-paperfig-medium-tree-', which.snp.tables(), '.pdf', sep='')
} else {
    paperfig.path <- paste('figs-mine/paperfig-medium-tree-', which.snp.tables(), '.pdf', sep='')
}
pdf(paperfig.path, width=8, height=5, onefile=TRUE, family='Helvetica', fonts='Courier', pointsize=10)
newick.medium <- newickize(make.tree(pat.summaries[,'count2']))
show.tree(newick.medium, total.snps=consistent.count[2], pltdebug=F)
dev.off()
# pdf
# pdf
# 2</pre>
```

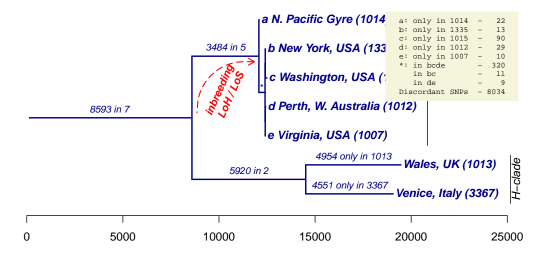


Figure 2: Tree based on unfiltered reads and loose SNP filters. "Lengths" are numbers of shared/private SNPs on Chr1.

```
# fig.paths for knitr chunks below; .h for "hand-made" trees; plain for automatic chrl/full versions
myfigpath <- paste(getwd(), '/figs-knitr/newick-', which.snp.tables(), '-', sep='')
myfigpath.h <- paste(getwd(), '/figs-knitr/newick-', sep='')</pre>
```

Figure 2, i.e., criteria [[1]]:

```
newick.loose <- newickize(make.tree(pat.summaries[,'count1']))
show.tree(newick.loose, total.snps=consistent.count[1])</pre>
```

Figure 3, i.e. [[2]]:

```
# newick.medium <- newickize(tree.by.hand)
# simple.newick.medium <- newickize(tree.by.hand, alt=TRUE)
newick.medium <- newickize(make.tree(pat.summaries[,'count2']))
simple.newick.medium <- newickize(make.tree(pat.summaries[,'count2']), alt=TRUE)
show.tree(newick.medium, total.snps=consistent.count[2])</pre>
```

Figure 4, i.e. [[3]]:

```
newick.strict <- newickize(make.tree(pat.summaries[,'count3']))
show.tree(newick.strict, total.snps=consistent.count[3])</pre>
```

Figure 5, i.e. [[4]]:

```
newick.unfiltered <- newickize(make.tree(pat.summaries[,'count4']))
show.tree(newick.unfiltered, total.snps=consistent.count[4])</pre>
```

Some other versions of the trees are included in the appendix.

Counts for all tree edges in the medium tree:

```
#pat.summaries[c(128,110,102,6,97,19,9,2,5,33,65,17,3),]
tree.edges <- c(128,110,102,6,97,19,9,2,5,33,65,17,3)-1
non.edges <- setdiff(0:127, tree.edges)
sg.edges <- showgroup(restrict.to=tree.edges); sg.edges</pre>
```

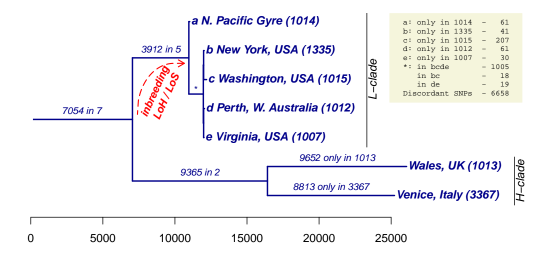


Figure 3: Tree based on unfiltered reads and medium SNP filters. "Lengths" are numbers of shared/private SNPs on Chr1.

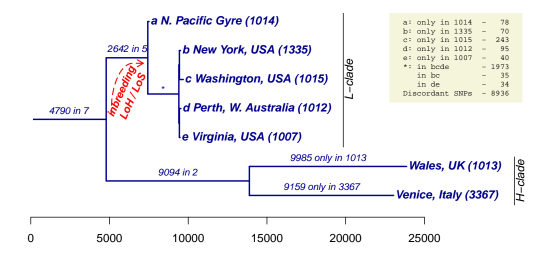


Figure 4: Tree based on unfiltered reads and strict SNP filters. "Lengths" are numbers of shared/private SNPs on Chr1.

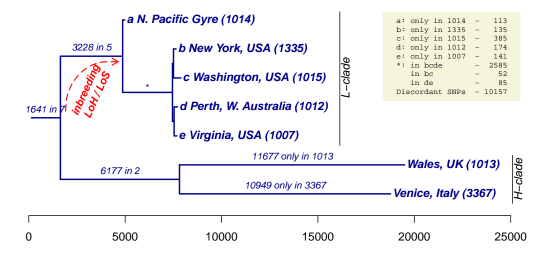


Figure 5: Tree based on unfiltered reads and unfiltered SNPs. "Lengths" are numbers of shared/private SNPs on Chr1.

```
Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 2
         002
         004
                                                                       207
                                                                               243
                                                                                       385
         005
                                                                                       113
         010
                                                                         61
  17
                                                              4954
                                                                       9652
                                                                              9985
                                                                                      6177
         022
                                                              5920
                                                                       9365
                                                                              9094
  33
                                                                29
                                                                        61
                                                                                95
  65
                                                                10
                                                                         30
                                                                                       141
  97
         140
                       Χ
                                                                         19
                                                                                34
                                                                                        85
         145
                       Χ
                                                                              1973
                                                                                      2585
  110
                                                              3484
                                                                              2642
                                                                                      3228
                                                                       3912
  128
         177
                                                              8593
                                                                       7054
                                                                              4790
                                                                                      1641
                                                             28006
                                                                             38238
                                                                                     37342
                                                                     40238
```

Counts for the top 10 discordant patterns, i.e., SNPs whose sharing pattern does not match any of the bifurcations in the tree:

```
tenth <- sort (showgroup (restrict.to=non.edges) [-(length (non.edges) +1), 'count2'], decreasing=T) [10]
sg.non.edges <- showgroup(restrict.to=non.edges, c2.thresh = tenth); sg.non.edges</pre>
         Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3
 56
        067
                                                                       78
                                                                             201
                                                               33
                                                                                     104
        144
                                                                              167
                                                                                     355
 104
         147
                                                                              590
                                                                                    1160
  109
         154
                                                                                     103
 112
         157
                                                             1343
                                                                              839
                                                                              787
 118
         165
                                                        Χ
                                                              201
                                                                      453
                                                                                    1140
 119
         166
                                                                              247
                                                                                     254
                                                        Χ
                                                                                    1852
  120
                                                                     1879
                                                                             3320
        175
 126
                                  Χ
                                                             1756
                                                                     1493
                                                                             997
                                                                                    1416
 127
        176
                                                               45
                                                                      114
                                                                                      68
 Other
               105 rows
                                            53
                                                             3528
                                                                      857
                                                                             1662
                                                                                    2362
                                                             8034
                                                                            8936
# Total
                                                                     6658
                                                                                   10157
```

And percent of discordant SNPs:

In short, the sharing pattern observed at 6658 or 14.2% of the 46896 medium-stringency consistent SNPs positions observed across all 7 isolates are discordant with the medium tree. (The strict tree has slightly more.)

A majority of the discordant SNPs fall into one of three patterns: 6-way sharing excluding Gyre (likely a technical artifact since the low coverage in Gyre reduces our power to detect SNPs there), or 6-way sharing excluding one of the two H-isolates (likely a reflection of sexuality in the H-clade—SNP positions in a population in Hardy-Weinberg equilibrium are fairly likely to be homozygous for the reference allele in a given individual).

```
third.biggest <- sort(showgroup(pat.summaries,6)[-8,'count2'],decreasing=T)[3]
big.three <- showgroup (pat.summaries, 6, c2.thresh = third.biggest); big.three
       Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
                             X X X X X 1343 1192
X Y Y ...
       157 6 X X
167 6 X X
175 6 X X
# 112
                                                                         1343
 120
      167
175
                                                                   3320
                                                                          1852
                                  X X
              6 X
                            X
# 126
                                                     1756
                                                            1493
                                                                   997
                                                                          1416
# Other (
             4 rows w/ c2 < 1192
                                                      116
                                                                           111
# Total
big.three.frac <- sum(big.three[1:3,'count2'])/discordy$count2; big.three.frac
# [1] 0.6854911
```

I.e., 68.5% of discordant SNPs fall into one of these three categories.

Out of curiousity: what is the ratio of full genome to Chr 1 branch lengths. Except for the shortest few, generally $\approx 10x$, as expected given the length of Chr 1:

```
# (vectors derived by editing Newick strings, and in that order)
print(
    c(Italy=86155, Wales=95697, IW=89598, Virg=330, Aust=632, VA=1296,
        Puget=2113, NY=658, PNY=480, four=10059, Gyre=568, five=39517, all=69526) /
    c(Italy=8813, Wales=9652, IW=9365, Virg=30, Aust=61, VA=19,
        Puget=207, NY=41, PNY=18, four=1005, Gyre=61, five=3912, all= 7054),
    digits=3)

# Italy Wales IW Virg Aust VA Puget NY PNY four Gyre five all
# 9.78 9.91 9.57 11.00 10.36 68.21 10.21 16.05 26.67 10.01 9.31 10.10 9.86

round(genome.length.constants()$genome.length.trunc / genome.length.constants()$chr1.length, digits=4)
# [1] 10.2879
```

9 Semi-Automated Tree-Building

Slightly formalizing the process above: Look for the bifurcation of the 7 strains that maximizes the number of shared SNPs *within* each side of the partition while minimizing the number and fraction of SNPs that are shared by subsets that include at least one strain on each side of the partition. The 2/5 split is the winner, with 6418 SNPs in confict with that partition (16% of the 39842 SNPs not shared by all 7; Chr1 data). The runner-up places the Gyre in a group by itself (7079 = 18% in conflict).

```
treepart <- function(p.summ=pat.summaries, root=127, verbose=T, stringency='count2'){
  root.shared <- p.summ[root+1, stringency]
  df<-NULL
  for(i in 1:floor(root/2)){
    if(bitwAnd(i,root)==i && i < root-i){</pre>
```

```
11 <- showgroup (p.summ, subset=i, split=NULL, proper.subset=F, total=T)
    1 <- 11[nrow(11), stringency]</pre>
    r1 <- showgroup (p.summ, subset=root-i, split=NULL, proper.subset=F, total=T)
    r <- r1[nrow(r1), stringency]
    c1 <- showgroup(p.summ, subset=root, split=i, proper.subset=T, total=T)</pre>
    c <- c1[nrow(c1), stringency]</pre>
    df <- rbind(df, data.frame(pat=i,left=l,right=r,both=l+r,cross=c,all=l+r+c,ratio=c/(l+r+c),
                                   best='',stringsAsFactors=F))
df$pat<-as.octmode(df$pat)</pre>
maxl <- which.max(df$left)</pre>
maxr <- which.max(df$right)</pre>
maxb <- which.max(df$both)</pre>
minc <- which.min(df$cross)
minr <- which.min(df$ratio)</pre>
df$best[c(max1, maxr, maxb, minc, minr)] <- '<'</pre>
df$best[maxl] <- paste(df$best[maxl], 'L') # max Left</pre>
df$best[maxr] <- paste(df$best[maxr], 'R') # max Right</pre>
df$best[maxb] <- paste(df$best[maxb], 'B') # max Both (L+R)</pre>
df$best[minc] <- paste(df$best[minc], 'C') # min Cross</pre>
df$best[minr] <- paste(df$best[minr], 'O') # min ratiO (Cross/(Left+Right+Cross)</pre>
if (verbose) {
  same <- all (maxl==c (maxr, maxb, minc, minr))</pre>
  cat('root:',
                      format (as.octmode (root), width=3),
      '; shared:', root.shared,
       '. max 1', format(as.octmode(df$pat[max1]), width=3),
', max r', format(as.octmode(df$pat[maxr]), width=3),
       ', max r',
       ', max both', format (as.octmode (df$pat[maxb]), width=3),
       ', min cross', format(as.octmode(df$pat[minc]), width=3),
       ', min ratio', format(as.octmode(df$pat[minr]), width=3),
        . \nAll the same?:', same,
       '\n')
  cat('\n')
return (df)
```

```
treepart()
# root: 177 ; shared: 7054 . max 1 077 , max r 010 , max both 022 , min cross 022 , min ratio 022 .
# All the same?: FALSE
    pat left right both cross all
     01 41 29077 29118 10724 39842 0.2691632
02 8813 17400 26213 13629 39842 0.3420762
     03 8870 10338 19208 20634 39842 0.5178957
# 3
# 4 04 207 28345 28552 11290 39842 0.2833693
# 5
     05 266 28142 28408 11434 39842 0.2869836
     06 9033 9956 18989 20853 39842 0.5233924
07 9110 9866 18976 20866 39842 0.5237187
# 6
     1.0
           61 32702 32763 7079 39842 0.1776768
# 8
                                                           < R
     11 109 28694 28803 11039 39842 0.2770694
# 10 12 8892 11759 20651 19191 39842 0.4816776
# 11 13 8959 10160 19119 20723 39842 0.5201295
# 12 14 270 28163 28433 11409 39842 0.2863561
# 13 15 342 28006 28348 11494 39842 0.2884895
# 14 16 9115 9849 18964 20878 39842 0.5240199
# 15 17 9213 9781 18994 20848 39842 0.5232669
# 16 20 9652 16099 25751 14091 39842 0.3536720
      21 9702 9470 19172 20670 39842 0.5187993
# 17
# 18 22 27830 5594 33424 6418 39842 0.1610863 < B C O
# 19 23 27916 542 28458 11384 39842 0.2857286
# 20 24 9871 9119 18990 20852 39842 0.5233673
# 21 25 9943 9016 18959 20883 39842 0.5241454
# 22 26 28089 239 28328 11514 39842 0.2889915
# 23 27 28213 175 28388 11454 39842 0.2874856
```

```
# 24 30 9731 10772 20503 19339 39842 0.4853923
               9303 19093 20749 39842 0.5207821
# 2.5
      31
         9790
 26
      32 27945
                1520 29465 10377 39842 0.2604538
      33 28045
                 414 28459 11383 39842 0.2857035
# 28
     34 9955
                9014 18969 20873 39842 0.5238944
 29
     35 10044
                8931 18975 20867 39842 0.5237438
     36 28214
                162 28376 11466 39842 0.2877868
 30
  31
      37
         28375
                 110 28485 11357 39842 0.2850510
 32
     40
            61 28554 28615 11227 39842 0.2817881
 33
     41
           106 28330 28436 11406 39842 0.2862808
 34
     42
          8875 10122 18997 20845 39842 0.5231916
 35
     43
          8944 10017 18961 20881 39842 0.5240952
           281 28125 28406 11436 39842 0.2870338
      44
           370 28005 28375 11467 39842 0.2878119
 37
     45
 38
     46
          9120
               9835 18955 20887 39842 0.5242458
# 39
     47
          9253
               9773 19026 20816 39842 0.5224637
          124 28358 28482 11360 39842 0.2851262
 40
     50
# 41
      51
           177
               28189 28366 11476 39842 0.2880377
     52
          8956 10008 18964 20878 39842 0.5240199
# 42
# 43
          9036
               9926 18962 20880 39842 0.5240701
# 44
     54
           352 27986 28338 11504 39842 0.2887405
 45
      55
           467 27885 28352 11490 39842 0.2883891
# 46
      56
          9212
               9743 18955 20887 39842 0.5242458
# 47
     57
          9386
                9691 19077 20765 39842 0.5211837
                9297 19014 20828 39842 0.5227649
 48
     60
         9717
         9773
                9175 18948 20894 39842 0.5244215
 49
     61
        27913
  50
      62
                 396 28309 11533 39842 0.2894684
 51
      63 28024
                 313 28337 11505 39842 0.2887656
         9958
                9006 18964 20878 39842 0.5240199
 52
     64
     65 10079
                8931 19010 20832 39842 0.5228653
 53
                143 28386 11456 39842 0.2875358
 54
     66 28243
                  93 28624 11218 39842 0.2815622
      67 28531
      70 9799
                9180 18979 20863 39842 0.5236434
 56
 57
      71
          9866
                9087 18953 20889 39842 0.5242960
 58
     72 28037
                 312 28349 11493 39842 0.2884644
                 246 28413 11429 39842 0.2868581
 59
      73 28167
                8912 18966 20876 39842 0.5239697
         10054
      75 10217
                8849 19066 20776 39842 0.5214598
 61
# 62
      76 28399
                  73 28472 11370 39842 0.2853772
     77 28807
                  30 28837 11005 39842 0.2762161
```

Comparing the 5/2 split to the second-place NPG/rest split (below), the former has fewer pattern instances in conflict with the split (6418 vs 7079), as well as somewhat more random distribution of the conflicting patterns (92 vs 62 rows), whereas the 1/6 split has the majority of its conflicts (3912 of 7079, or 55%) concentrated in one pattern—the 5 NE strains. Collectively, these seem to favor the 5/2 split as the correct "history."

```
showgroup(pat.summaries,split=strtoi('022'), subset=127, proper.subset=T, c2.thresh=100)
        Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 104
                                                           125
                                                                   307
                                                                           590
                                                                                 1160
        147
                      Χ
                                           Χ
# 112
        157
                      Χ
                           Х
                                           Χ
                                                 Χ
                                                      Χ
                                                           1343
                                                                  1192
                                                                           839
                                                                                 1343
# 118
        165
                                 Χ
                                           Х
                                                            201
                                                                   453
                                                                           787
                                                                                 1140
                           Χ
                                                      Χ
# 119
        166
                                 Χ
                                           Χ
                                                 X
                                                            30
                                                                   157
                                                                           247
                                                                                  254
 120
        167
                 6
                      Χ
                           Χ
                                 Χ
                                           Χ
                                                 Χ
                                                      Χ
                                                            951
                                                                  1879
                                                                          3320
                                                                                 1852
                                                           1756
                                                                  1493
                                                                                 1416
# 126
        175
                      Χ
                                                                           997
                                                                           77
# 127
        176
                6
                      X
                           X
                                X
                                      Χ
                                           X
                                                            45
                                                                   114
                                                                                   68
# Other
                85 rows
                                         100
                                                           3493
                                                                   823
                                                                          1387
                                                                                 1771
                          w/
# Total
                                                           7944
                                                                  6418
                                                                          8244
                                                                                 9004
showgroup(pat.summaries, split=strtoi('010'), subset=127, proper.subset=T, c2.thresh=100)
        Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 110
        155
                5
                      Χ
                           Χ
                                      Χ
                                           Χ
                                                      Χ
                                                           3484
                                                                  3912
                                                                          2642
                                                                                 3228
  112
        157
                                      Χ
                                           Χ
                                                           1343
                                                                  1192
                                                                           839
                                                                                 1343
                 6
                      Χ
                           Χ
                                                      Х
# 126
        175
                                      Χ
                                           Χ
                                                           1756
                                                                  1493
                                                                           997
                                                                                 1416
                 6
                      Х
                           Χ
                                                                           77
        176
                                Χ
                                           Χ
# 127
                 6
                      X
                           X
                                      Χ
                                                            4.5
                                                                   114
                                                                                   68
                                                           1095
                                                                           368
# Other
                58 rows
                                c2
                                          100
                                                 )
                                                                   368
                                                                                   522
          (
                          w/
                                                          7723
                                                                  7079
# Total
                                                                          4923
                                                                                 6577
```

Below is the full summary of shared SNPs that do *not* directly correspond to tree splits, e.g. deep coalescence, independent coincident mutations, false positives/false negatives in the shared SNP calls, loss of SNPs in hemizygous regions, etc. (Additionally, SAMTools' SNP calls exclude positions it judges to be homozygous, and I think it operates without regard to the reference sequence, so homozygous nonreference positions, while rare except in IT/Wales, often are not called SNPs by SAMTools, but are relevant for this analysis. Provided the position is called a SNP in some other isolate, the consistency filtering we've done above should recover it, but this is still worth keeping in mind when examining the data.)

First, here are SNPs that "coalesce" on the branch from the LCA of bcde, i.e., shared among some nonempty, proper subset of bcde other than bc or de. There are 8 such patterns: any of the 4 choose 3 trios plus any of the 4 pairs having exactly one of bc.

```
sg4 <- showgroup(pat.summaries, subset=strtoi('0145'), split=5, proper.subset = F)</pre>
sg4
        Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
#
 34
        041
            2
                                                          3
                                                                4
                                                                      13
                                                                        7.5
 37
        044
                2
                          Χ
                                                           5
                                                                 13
                                                                               155
                                         X
 38
        045
                                          Χ
                                                    Χ
                                                           6
                                                                 26
                                                                       130
                                                                              131
 66
                                                    Χ
                                                           1
                                                                  2
                                                                        13
# 69
       104
                                         Χ
                                                                        34
                                                                              107
                     Χ
                                                                 5
# 70
       105
                                                                       25
                                                                 5
                                                                       1.5
# 98
        141
               3
                     X
                          X
                                                    Χ
                                                                               40
 101
        144
                     Χ
                          Χ
                                                                 75
                                                                       167
                                                                              355
                                                               1005
                                                                      1973
# 102
        145
                4
                     X
                          Χ
                                         X
                                                         320
                                                                             2585
                                                         352
                                                             1144 2445
# Total
                                                                             3492
sq4n \leftarrow nrow(sq4)
sg4pct <- round(sg4$count2[sg4n-1]/sg4$count2[sg4n]*100,1)</pre>
# [1] 87.8
```

So, of the 1144 SNPs found only in bcde, 87.8% have a sharing pattern consistent with the given tree structure. Similarly, we analyze patterns relative to the root of the L-clade (14 patterns—any nonempty proper subset of bcde together with a):

```
sg5 <- showgroup(pat.summaries, subset=strtoi('0155'), split=8, proper.subset = F)</pre>
sq5
        Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 10
        011
                                     Χ
                                                           6 7 7 14
# 13
        014
                                     Χ
                                          Χ
                                                                          5
                                                                                 12
# 14
        015
                3
                                                     Χ
                                                            7
                                                                   6
                                                                          4
                                                                                 7
                                     Χ
        050
                                     Χ
                                                                   2
# 41
        0.51
                          Χ
                                     Χ
                                                            0
# 42.
                3
                                                                   1
                                                                         17
# 45
        054
                                     Χ
                                                            1
                                                                  6
# 46
        055
                4
                          Χ
                                     Χ
                                                     Χ
                                                            6
                                                                  12
                                                                         29
 73
        110
                2
                     Χ
                                     Χ
                                                                          4
 74
        111
                3
                                     Χ
                                                            1
                                                                   0
                                                                          1
# 77
        114
                3
                     X
                                     Χ
                                          Χ
                                                            1
                                                                   2.
                                                                          2.
 78
        115
                          Χ
                                                                   Ω
# 105
        150
                3
                                     X
                                                            \cap
                                                                                 6
 106
                4
                                     Χ
                                                                   4
                                                                          5
        151
                     Χ
                          Χ
                                                            1
                                                                                14
                                                                       49
                                                                53
# 109
        154
                4
                                     Χ
                                                           16
                                                                                103
        155
# 110
                5
                                                         3484
                                                                3912
                                                                       2.642
                                                                               3228
# Total
                                                         3542
                                                                4013
                                                                       2779
sa5n \leftarrow nrow(sa5)
sg5pct <- round(sg5$count2[sg5n-1]/sg5$count2[sg5n]*100,1)</pre>
```

I.e., of the 4013 SNPs found only in abcde, 97.5% have a sharing pattern consistent with the given tree structure. Finally, how many SNPs have patterns inconsistent with the 5-2 split, i.e., include at least one strain on each side of the 5-2 split, but not shared by all 7?

sa7 <-	showe	roup (r	nat ei	ımmar	ies s	subset	=127	. snl	 t = st =	toi('A'	22!) ni	oner si	ıbset=F)
sg7	STOMO	Jroup (I	vat.Sl	ATTILITAL -	LCD, S	Japsel		, sbr	LC-SCI	.001 (02	, pi	oper.st	woet-r)
#	Do+	ShrDr	1007	1012	1012	1014	1015	3367	1335	count 1	count2	count?	COUN+ A
# # 4	003	Shrby 2	1007	1012	1013	1014	1015	3307 X	1335 X	239	16	28	31
# 7	006	2					X			141	13	21	41
# 8	007	3					X	Χ	Х	9	2	9	10
‡ 11	012	2				Χ		Х		179	18	2	5
12	013	3				X		Х	Χ	17	3	2	2
15	016	3				X	X		V	9	1 5	2	2 2
16 18	017 021	4 2			Х	Χ	Х	Х	X X	243	9	11	9
20	023	3			X			Х	X	327	20	29	17
21	024	2			X		Χ			125	12	28	46
22	025	3			X		X		Χ	9	4	12	21
23	026	3			Χ		Х			185	27	31	32
24	027	4			X	3.7	Х	Χ	Х	22	14	28	24
: 25 : 26	030 031	2			X	X			v	222	18	5	9
20	031	3			X	X		Х	Χ	324	18	8	5
28	033	4			X	X		X	Х	30	2	4	6
‡ 29	034	3			Х	Х	Х			11	3	1	1
ŧ 30	035	4			Х	Х	Х		Х	4	2	1	0
31	036	4			X	X	Х			12	4	1	3
32	037	5 2		V	Х	Χ	Х		Χ	17 150	10	8 1.6	5 27
36	042	3		X				X	Х	21	1 8	16 14	6
: 39	045	3		Х			Х		Λ	11	12	34	55
40	047	4		X			X		X	2	18	42	60
43	052	3		X		X		X		9	0	2	1
44	053	4		Χ		Χ		Х	Χ	1	0	2	2
47	056	4		X		X	X		7.7	3	2	2	5
‡ 48 ‡ 49	057 060	5 2		X	Х	Χ	Х	Х	Χ	5 165	7	9 26	17 28
± 50	061	3		Х	Х				Х	12	2	14	12
51	062	3		X	X			Х		227	17	37	36
52	063	4		Χ	X			X	X	18	11	24	21
53	064	3		X	X		X			9	9	36	60
54	065	4		X	X		X		Х	9	17	41	48
55	066	4		X	X		X		V	25	37	102	76 104
56 57	067 070	5		X	X	X	Χ	X	Χ	33 13	78 1	201	104
: 58	071	4		X	Х	Х			Х	2	1	2	0
59	072	4		X	X	X		Х		8	6	3	2
60	073	5		X	X	X		X	X	3	3	6	3
61	074	4		X	Х	Х	Х			2	3	2	7
62	075	5		X				3.7	Χ	7		13	11
63 64	076 077	5 6		X	X	X	X		Х	9 43	11 48	17 60	7 26
67	102	2	Х		Λ	Λ	Λ	Х	Λ	93	6	7	25
68	103	3	X					X	X	11	4	4	8
71	106	3	X				X			3	9	15	27
72	107	4	X				Х		Х	5	5	11	16
75	112	3	X			X		X	3.7	4	1	0	0
76	113 116	4	X			X	Х	X	Χ	1 0	2	0	1 5
: 80	117	5	X			Х	X		Х	0	2	1	7
81	120	2	X		Х					105	9	9	31
82	121	3	X		X				X	8	0	4	4
83	122	3	X		X			Χ		134	10	10	26
ŧ 84 ŧ 85	123	4	X		X			Χ	X	10	9	8	8
	124 125	3	X		X		X		V	11 4	7	20	35 16
86 87	125	4	X		X		X X		Χ	12	20	13 21	43
88	127	5	X		X		X		Х	11	17	30	47
89	130	3	X		X	X				7	1	2	1
90	131	4	X		X	X			X	0	1	0	2
91	132	4	Χ		Х	Χ		Χ		9	1	2	1

```
# 92
         133
                   5
  93
                                     Χ
                                           Χ
         134
                   4
                         Χ
                                                 Χ
                                                                     2.
                                                                                      1
                                                                                              3
  94
         135
                         Χ
                                     Χ
                                           Χ
                                                 Χ
                                                                     2
                                                                              1
                                                                                      3
  95
         136
                   5
                         Х
                                     Χ
                                           Χ
                                                 Χ
                                                       Χ
                                                                     4
                                                                              6
                                                                                      3
                                                                                              5
 96
         137
                   6
                                           Χ
                                                 Χ
                                                       Χ
                                                                    11
                                                                              6
                                                                                     12
                                                                                             14
                         Χ
  99
         142
                   3
                                                       Χ
                                                                     9
                                                                                      9
                                                                                             15
 100
         143
                                                                             3
                                                                                      5
                   4
                         Χ
                               Χ
                                                       Χ
                                                             Χ
                                                                     1
  103
         146
                   4
                         Χ
                               Χ
                                                 Χ
                                                       Χ
                                                                            34
                                                                                     65
                                                                                            140
 104
         147
                   5
                               Χ
                                                 Χ
                                                       Χ
                                                                   125
                                                                            307
                                                                                    590
                                                                                           1160
# 107
         152
                                           Χ
                                                       Χ
                                                                     2
                         Χ
                               Χ
                                                                             1
                                                                                     1
                                                                                              4
 108
         153
                   5
                               Χ
                                           Χ
                                                       Χ
                                                             Χ
                                                                     3
                                                                             3
                                                                                      1
                   5
                                                                     8
                                                                            15
                                                                                     15
 111
         156
                         Χ
                               Χ
                                           Χ
                                                 Χ
                                                       Χ
                                                                                             43
 112
         157
                         Χ
                                                 Χ
                                                                  1343
                                                                          1192
                                                                                    839
# 113
         160
                   3
                         Χ
                               X
                                                                             6
# 114
         161
                   4
                         Χ
                                     Χ
                                                                                             18
                               Χ
# 115
         162
                   4
                                     Χ
                                                       Χ
                                                                    11
                                                                            12
                                                                                     21
                                                                                             33
                                                                            18
                                                                                     23
  116
         163
                   5
                                                                    12
                                                                                             33
                         Χ
                               Χ
                                     Χ
                                                             Χ
                                                       Χ
 117
         164
                                                                            47
                                                                                     80
                                                                                            163
# 118
                   5
                                                                           453
                                                                                    787
         165
                         Χ
                                     Χ
                                                 Χ
                                                                                           1140
                               X
                                                             Χ
# 119
         166
                                     Χ
                                                 Χ
                                                       Χ
                                                                    30
                                                                           157
                                                                                   247
                                                                                            254
 120
         167
                   6
                               Χ
                                     Χ
                                                       Χ
                                                             Χ
                                                                   951
                                                                          1879
                                                                                   3320
                                                                                           1852
  121
         170
                   4
                         Χ
                               Χ
                                     Χ
                                           Χ
                                                                             0
# 122
         171
                   5
                         Χ
                               Χ
                                     Χ
                                           Χ
                                                                     3
                                                                              6
# 123
                                                                     2.
                   5
                                           Χ
                                                       Χ
                                                                                      3
                                                                                              5
                         X
                               X
                                     X
                                                                              6
                                                                             9
                                                                                     7
 124
         173
                                           Х
                                                       Χ
                                                                    17
                                                                            2.2
 125
         174
                   5
                         Χ
                               Χ
                                     Χ
                                           Χ
                                                 Χ
                                                                                     2.3
                                                                                             3.5
  126
         175
                   6
                         Χ
                               Χ
                                     Χ
                                           Χ
                                                 Χ
                                                                  1756
                                                                          1493
                                                                                    997
                                                                                           1416
# 127
         176
                   6
                         Χ
                               Χ
                                     Χ
                                           Χ
                                                 Χ
                                                       Χ
                                                                    45
                                                                           114
                                                                                             68
                                                                  8593
                                                                          7054
                                                                                   4790
# 128
         177
                   7
                                           Χ
                                                 Χ
                                                       Χ
                                                                                           1641
                               Χ
                                                                         13472
# Total
                                                                                  13034
sa7n \leftarrow nrow(sa7)
        <- round(sg7$count2[sg7n-1]/sg7$count2[sg7n]*100,1)</pre>
sq7pct
# [1] 52.4
```

A more compact version of that table, showing only the larger counts:

```
thresh <- signif(.02 * sg7$count2[sg7n],1)</pre>
thresh
# [1] 300
showgroup(pat.summaries, subset=127, split=strtoi('022'), proper.subset=F, c2.thresh = thresh)
        Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 104
        147
                      Χ
                                                           125
# 112
        157
                 6
                      Χ
                           Х
                                           Χ
                                                 Χ
                                                      X
                                                           1343
                                                                  1192
                                                                           839
                                                                                 1343
# 118
        165
                                 Χ
                                           Χ
                                                           201
                                                                  453
                                                                           787
                                                                                 1140
                      Χ
                           Χ
                                                      Χ
 120
        167
                                           Χ
                                                      Х
                                                            951
                                                                  1879
                                                                          3320
                                                                                 1852
                                           Χ
                                                           1756
                                                                  1493
                                                                          997
 126
        175
                 6
                      Χ
                           Χ
                                 Χ
                                      Χ
                                                      X
                                                                                 1416
                                           Χ
                                                                          4790
  128
                      Χ
                           X
                                 Χ
                                      Χ
                                                           8593
                                                                  7054
                                                                                  1641
                                                                          1711
                87 rows
                                                           3568
 Other
                           w/
                                c2
                                                                  1094
                                                                                 2093
                                                          16537
                                                                13472
                                                                         13034
                                                                                10645
# Total
```

So, of the 13472 SNPs found both in the L- and H-clades, 52.4% have a sharing pattern consistent with the given tree structure, i.e., are found in all 7 isolates. Among the others, three patterns dominate—(i) the 6-way pattern excluding the Gyre is the largest, plausibly explained by 7-way sharing from which the Gyre drops out due to low coverage/high error rate, (ii) the 6-way excluding Italy, and (iii) ditto for Wales. Origin of the later two cases is unclear, but may partly reflect Hardy-Weinberg—some positions that are *population-level* SNPs in those isolates will be homozygous-reference in the CCMP founder cell for IT or Wales. If I take the 7-way shared SNP count (69526) as a surrogate approximating the number of population-level SNPs in either IT or Wales that are shared with the L-clade, then I might expect, based on HWE, roughly half that number to to be lost (become homozygous) in IT, and a similar number in Wales. However, the observed counts of these positions are lower by ≈20K than I might have guessed

from HWE, perhaps suggesting that IT and Wales are distinct populations, each with a pool of many thousand private polymorphisms.

In aggregate:

```
untreelike <-
    sg7$count2[sg7n]-sg7$count2[sg7n-1] +
    sg5$count2[sg5n]-sg5$count2[sg5n-1] +
    sg4$count2[sg4n]-sg4$count2[sg4n-1]
untreelike

# [1] 6658

consistent.count[2]

# [1] 46896

unpct <- round(untreelike/consistent.count[2]*100,1)
unpct

# [1] 14.2</pre>
```

I.e., 6658 or 14.2% of the 46896 consistent SNPs identified (by criterion 2) across all 7 isolates are discordant with the assumed tree.

Overall, based on this data, I take the following to be obvious: (a) separation of the He-isolates from the L-isolates (and from each other??), and (b) near-identity of the L-isolates. Due to the small counts, the exact topology among the L-isolates (esp. bcde) is uncertain, but *any* topology there is consistent with the asexual/clonal/global-expansion hypothesis, so there is little point in examining this subtree more carefuly. Again, we believe the (apparent) slight separation of the Gyre from the other L-isolates is largely driven by technical artifacts (lower coverage/higher error rates) in the sequencing rather than by biological effects. However, the discord between Gyre SNPs and others is the major substantive ambiguity in the offered tree. Nevertheless, in the next section we show by a bootstrap analysis that the offered placement of Gyre with respect to the other 4 L-isolates is strongly supported by the data.

9.1 Bootstrap

How robust is the inferred tree? Italy/Wales seem clearly related to each other but separate from the other 5. Likewise, the 4 coastal L-isolates seem to be closely related, with little data to separate them (and perhaps little sense in trying). So, the key question here is whether the top level bifurcation is 2/5 or NPG/6. Here, we do a simple bootstrap test (on c2 numbers only) to see whether the 2/5 split is consistently the most parsimonious.

```
n2 <- sum(pattern.counts[[2]][,2]); n2
# [1] 46896
```

Conceptually, we sample, with replacement, n2=46896 SNP positions from among the 46896 positions declared consisent SNPs according to criterion c2, and recalculate the statistics examined above to see whether the 2/5 split again minimizes conflicting sharing patterns. This resampling/calculation is repeated nboot times (set near front of file). Since all that matters is the sharing pattern, this procedure is expedited by actually sampling 46896 independent integers in the range 0:127 with probabilities proportional to the pattern counts given in column 2 of pattern.counts[[2]]. The sample is then tabulated in a 128 row table analogous to pattern.summaries, for analysis by showgroups/treepart, as above.

```
boot.sample <- sample(0:127,n2,replace=T,prob=pattern.counts[[2]][,2])
str(boot.sample)

# int [1:46896] 125 1 16 117 127 16 2 127 16 127 ...

boot.count <- mytable(boot.sample,c(0,127))
boot.count[c(1:4,125:128),] # show a few rows</pre>
```

```
val count
# [1,]
      0 0
# [2,]
       1
# [3,]
          8854
# [4,]
       .3
          14
# [5,] 124
           19
# [6,] 125 1408
# [7,] 126
           118
# [8,] 127
boot.counts <- list(NULL, boot.count, NULL) # dummy list with just c2 summaries
cor(pattern.counts[[2]][,2],boot.counts[[2]][,2]) # just curious - how correlated are they?
boot.summaries <- pat.summary(boot.counts)</pre>
showgroup(boot.summaries,c2.thresh=400) #show a few rows
      Pat ShrBy 1007 1012 1013 1014 1015 3367 1335 count1 count2 count3 count4
# 3
                                                 NA 8854 NA NA
# 17
                                                 NA 9650
# 19
      022
                                                 NA 9410
# 102
     145 4 X X
                                                 NA 999
                                                             NA
                                            X
             5 X X
6 X X
# 110
      155
                                   Χ
                                                 NA
                                                      3881
                                                              NA
                                       X X
# 112
      157
                                   Χ
                                                 NA
                                                      1223
                                                              NA
                                           Χ
# 118
      165
                                                      421
                                   Χ
                                                 NA
                                                             NA
# 120
     167
                                                  NA 1866
     175
           6 X
7 X
                    X
                                                 NA 1408
# 126
                               X
                                  X
                                           X
                                                              NA
                                                                    NA
                         X
                                                      7032
# 128
      177
                     X
                          Χ
                               Х
                                        Χ
                                                  NA
          118 rows
                                                      2152
# Other (
                     w/
                          c2
                                  400
                                        )
                                                  NA
                                                              NA
                                                                    NA
                                                 NA 46896 NA NA
# Total
```

Tree partition analysis (and how to pluck out only the best rows based on 3 smallest cross counts and "best" criteria):

```
tp <- treepart(boot.summaries,root=127) ; tp</pre>
# root: 177 ; shared: 7032 . max 1 077 , max r 010 , max both 022 , min cross 022 , min ratio 022 .
# All the same?: FALSE
                                   all
    pat left right both cross
           47 29232 29279 10585 39864 0.2655278
     02 8854 17306 26160 13704 39864 0.3437688
     03 8915 10386 19301 20563 39864 0.5158288
         239 28465 28704 11160 39864 0.2799518
           305 28241 28546 11318 39864 0.2839153
     06 9103 9984 19087 20777 39864 0.5211971
         9184 9878 19062 20802 39864 0.5218242
63 32785 32848 7016 39864 0.1759984
     10
          117 28834 28951 10913 39864 0.2737558
      11
# 10
     12
          8932 11773 20705 19159 39864 0.4806091
         9003 10204 19207 20657 39864 0.5181868
     13
          304 28268 28572 11292 39864 0.2832631
# 12
          380 28097 28477 11387 39864 0.2856462
# 13
     15
         9184 9868 19052 20812 39864 0.5220751
# 14
     16
          9282 9789 19071 20793 39864 0.5215984
# 15
     20 9650 16149 25799 14065 39864 0.3528246
# 16
     21 9709 9556 19265 20599 39864 0.5167319
22 27914 5626 33540 6324 39864 0.1586394 < B C O
# 18
                608 28615 11249 39864 0.2821844
     23 28007
     24 9903 9182 19085 20779 39864 0.5212472
# 2.0
     25 9985 9067 19052 20812 39864 0.5220751
# 21
                263 28469 11395 39864 0.2858469
     26 28206
     27 28339
                190 28529 11335 39864 0.2843418
     30 9733 10813 20546 19318 39864 0.4845976
     31 9803 9378 19181 20683 39864 0.5188391
# 26
      32 28035 1573 29608 10256 39864 0.2572747
     33 28145
                471 28616 11248 39864 0.2821593
      34 9989 9070 19059 20805 39864 0.5218995
# 28
     35 10088 8982 19070 20794 39864 0.5216235
# 30
     36 28334
                178 28512 11352 39864 0.2847682
# 31 37 28505 122 28627 11237 39864 0.2818834
# 32 40 73 28672 28745 11119 39864 0.2789233
# 33 41 122 28435 28557 11307 39864 0.2836394
```

```
# 34 42 8927 10160 19087 20777 39864 0.5211971
# 35 43 8999 10047 19046 20818 39864 0.5222256
# 36 44 329 28221 28550 11314 39864 0.2838150
# 37 45 421 28089 28510 11354 39864 0.2848184
# 38 46 9202 9845 19047 20817 39864 0.5222005
            9340 9772 19112 20752 39864 0.5205699
# 40 50 140 28470 28610 11254 39864 0.2823099
              196 28296 28492 11372 39864 0.2852699
# 42 52 9009 10046 19055 20809 39864 0.5219998
       53 9091 9956 19047 20817 39864 0.5222005
# 44 54 402 28073 28475 11389 39864 0.2856964
# 45 55 514 27965 28479 11385 39864 0.2855960
# 46 56 9292 9749 19041 20823 39864 0.5223510
# 47 57 9461 9688 19149 20715 39864 0.5196418
# 48 60 9726 9366 19092 20772 39864 0.5210716
# 49 61 9787 9246 19033 20831 39864 0.5225517
# 50 62 28007 438 28445 11419 39864 0.2864489
# 51 63 28121 351 28472 11392 39864 0.2857716
# 51 63 28121
                      351 28472 11392 39864 0.2857716
# 52 64 10002 9051 19053 20811 39864 0.5220500
# 53 65 10127 8973 19100 20764 39864 0.5208710
# 54 66 28383 153 28536 11328 39864 0.2841662
                      97 28789 11075 39864 0.2778196
# 55 67 28692
# 56 70 9814 9256 19070 20794 39864 0.5216235
# 57 71 9886 9161 19047 20817 39864 0.5222005
# 58 72 28144 356 28500 11364 39864 0.2850692
# 59 73 28276
                     284 28560 11304 39864 0.2835641
       74 10098 8959 19057 20807 39864 0.5219496
# 60
      75 10264 8893 19157 20707 39864 0.5194411 76 28546 82 28628 11236 39864 0.2818583
# 61
# 62
# 63 77 28971 33 29004 10860 39864 0.2724262
```

```
otp <- order(tp[,'cross'])[1:3]  # 3 smallest 'cross' counts
btp <- which(tp[,'best'] != '')  # 'best' by Left/Right/Both/Cross/ratiO
toptp <- unique(c(otp,btp,18,8))  # above, plus 5/2, 6/1 splits
print(tp[toptp,])  # show the winners

# pat left right both cross all ratio best
# 18 22 27914 5626 33540 6324 39864 0.1586394 < B C O
# 8 10 63 32785 32848 7016 39864 0.1759984 < R
# 26 32 28035 1573 29608 10256 39864 0.2572747
# 63 77 28971 33 29004 10860 39864 0.2724262 < L</pre>
```

Now repeat the above nboot times, and summarize results:

```
nboot <- params$nboot # default from params set in section 2</pre>
nboot \leftarrow ((nboot+2) \%\% 4) * 4 + 1 # summary is cleaner if n mod 4 == 1, so int median/quartiles
cat('***\n*** Doing', nboot, 'bootstrap replicates.\n***\n')
# *** Doing 5 bootstrap replicates.
bcor <- numeric(nboot)</pre>
b52cross <- integer(nboot)
b61cross <- integer(nboot)</pre>
brev <- logical(nboot)</pre>
for(i in 1:nboot){
 boot.sample <- sample(0:127,n2,replace=T,prob=pattern.counts[[2]][,2])
  boot.count <- mytable(boot.sample,c(0,127))</pre>
  boot.counts <- list(NULL, boot.count, NULL) # dummy list with just c2 summaries
  boot.summaries <- pat.summary(boot.counts)</pre>
  tp <- treepart(boot.summaries,root=127, verbose=F)</pre>
  bcor[i] <- cor(pattern.counts[[2]][,2],boot.counts[[2]][,2]) # just curious - how correlated are they?
  b52cross[i] <- tp[18,'cross']</pre>
  b61cross[i] <- tp[ 8,'cross']</pre>
  brev[i] <- (b52cross[i] > b61cross[i])
  if (brev[i]) {
   # show the unexpected ones; probably breaks w/ cache
   otp <- order(tp[,'cross'])[1:3]
   btp <- which(tp[,'best'] != '')</pre>
  toptp <- unique(c(otp,btp,18,8))
```

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```
print(tp[toptp,])
}

# summarize:
corsummary <- t(as.matrix(c(summary(bcor), sd=sd(bcor))))
row.names(corsummary) <- 'bcor'
bdelta <- b6lcross-b52cross
brevp <- 100*brev  # make it percent reversed instead of logical
thesummary <- rbind(summary(b52cross), summary(b61cross), summary(c(bdelta)), summary(brevp))
row.names(thesummary) <- c('b52cross', 'b61cross', 'b61-b52', '% rev')
thesummary <- cbind(thesummary, sd=c(sd(b52cross), sd(b61cross), sd(bdelta), sd(brevp)))</pre>
```

SUMMARY: In 5 bootstrap replicates, we saw 0 samples with the 6/1 split having fewer conflicts than the 5/2 split, and the minimum separation between them was \approx 6.6 sigma, hence highly statistically significant.

```
# 'opt' hacking is trying to force knitr to show more digits of bcor in summary, as Rstudio does, but
# it still fails... Bottom line is the correlation seems to be > .999 in all samples, rounds to 1.0,
# as seen in this run of 1001 samples cut/paste from Rstudio:
      Min. 1st Qu. Median Mean 3rd Qu. Max. sd
              0.9998" " 0.9999" " 0.9999" " 1" " 1" " 0.00003462"
# > max(bcor)
# [11 0.9999915
o.opts <- options(digits=7, width=127)</pre>
format (rbind (corsummary, thesummary), scientific=F, digits=4, drop0trailing=T)
# Min. 1st Qu. Median Mean 3rd Qu. Max. sd
# bcor " 0.99989616" " 0.99991299" " 0.99994527" " 0.99994012" " 0.99996772" " 0.99997845" " 0.
# b52cross "6286" "6310" "6316" "6352" "6389" "6459" "71.
# b61cross "6877" "7000" "7004" "6988.4" "7011" "7050" "65.
                                                                                                        " 65.
# b61cross "08//"
# b61-b52 " 552" " 567"
# % rev " 0" " 0"
                                                         " 567" " 615" " 636.4
" 0" " 0" " 0"
                                                                                         " 734"
                                                                                                         " 83.
                                                                                         ... 0 ...
                                                                                                            0"
options(o.opts)
```

Based on this, it is reasonable to claim that we are confident that the tree topology is as shown in the earlier figures, with the exception of the exact order of the splits with the 4 NE coastal isolates.

10 Notes

This section is a random brain dump of limitations of the current analysis, ideas for improvements, etc. In the main, these may not be worth doing, unless we see significant holes or get pushed by reviewers, etc, but I wanted to catalog before we forget them.

Noise: Various sources of "noise" in the data:

- 1. Read errors, low read depth perhaps fixed by medium/strict thresholding
- 2. Deep coalescence
- 3. Skew because 1335 is the reference. (Julie notes we could partially fix this by remapping based on discovered SNPs, tho that wouldn't fix gross misassembly in 1335, e.g. collapsed or misordered tandem duplicates, or segments missing in 1335 that are present in one or more other strains, etc.; much harder to fix those, let's just hope they are rare...)
- 4. Varying error rates and sequencing depth among the 7. E.g., plausibly the 1000 SNPs shared by 4 but not by Gyre are a result of lower read depth (we missed a SNP that is actually present) and/or higher error rates (causing a position to appear inconsistent in gyre) in the gyre data. I can't think of a way to correct for this effect. It might be possible, perhaps by simulation, to estimate the size of the effect and see whether it could explain ≈1000 SNPs.
- 5. Varying numbers of founder cells in the sequencing cultures. (Again, I made some attempts at modeling this, but nothing very satisfactory yet.)

- 6. Tri-allelic positions where stochastic fluctuation in sequence sampling promotes the rare allele to prominence. (Julie replies: "isn't this the same as more than one founder cell? If they are diploid there should only ever be two alleles, unless there were random and very rare, thus unlikely, trisomy events?" I agree, but it is a concrete example of an effect of multiple founders that might be important. Not sure this is the most important such effect...)
- 7. Gaps/indels alignments are likely to be of lower quality in the vicinity of an indel, so, maybe lower coverage/more SNPs. We ignored them. Does this add any systematic bias? e.g. if one strain had more indels than another, would this confound other analyses? unclear. Julie suggested a paper titled "Barking up the wrong tree-length: yada yada yada gap penalties"; maybe relevant?

Other Items/Potential To Dos:

- 1. any spacial structure to various sub-classes?
- 2. after top level split, should I reanalyze halves of partition in isolation? said another way, I think the tree-building is sensible, but not sure it's optimal.
- 3. if we believe no sex, then I think gain of SNP should be more common than loss of SNP, since the later can only happen by (a) mutation reverting to reference, (b) second mutation matching nonreference, (c) homologous repair (look for blocks of LOH), or (d) false negative e.g. from low read depth. Does tree-building appropriately weight the gain vs loss cases? (Does it even care?)
- 4. should we weight coding and/or nonsynonomous SNPs more heavily? Julie says "you do not want to weight the coding or nonsynonomous/coding SNPs because for time you want the more clock-like neutral mutations." I.e., I got this backwards. Maybe should redo tree based on noncoding SNPs only.
- 5. We could also do an actual parsimony analysis based on 2-state model (homozygous-ref vs not), but I'm not quite sure how to handle this in a mixed sex/nosex case.
- 6. Might be interesting to look at sharing just within (shared?) deserts. Given tree model above and expectation that bottleneck followed split of H- from L-clades, I would expect little or no sharing of L-clade desert SNPs with H-clade; sharing between It/Wales might suggest "desert" is actually a region under strong purifying selection (e.g. a gene); sharing/non-sharing within L-clade deserts might suggest more about evo history of the 5.

11 Appendix: Old Trees, etc.

Tangents, old stuff of historical interest at best, etc..

11.1 HWE Sharing

Tangent: As a function of nonref allele freq, assuming HWE, what is probability that nonref allele will be seen in k strains, $0 \le k \le 4$ (Fig 6).

```
myfigpath.h <- paste(getwd(), '/figs-knitr/', sep='')</pre>
```

```
p <- (0:20)/20
q <- 1-p
r <- 2*p*q+p^2
plot( p, 1*q^0*r^4, type='b',pch='4', ylab="share prob")
points(p, 4*q^2*r^3, type='b',pch='3')
points(p, 6*q^4*r^2, type='b',pch='2')
points(p, 4*q^6*r^1, type='b',pch='1')
points(p, 1*q^8*r^0, type='b',pch='0')</pre>
```

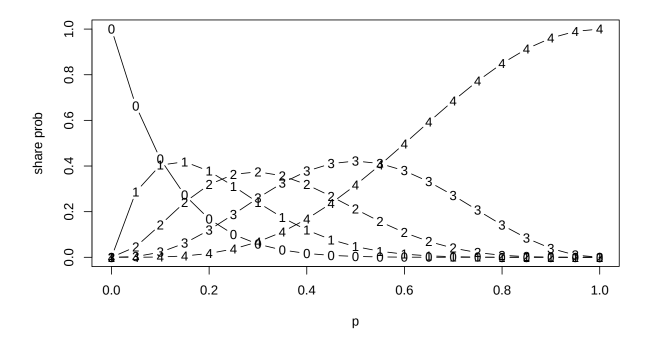


Figure 6: Sharing Probability

11.2 Old Tree Stuff

All based on un-q-filtered reads.

The first pass at the tree analysis was the Chr1 tree, *loose criteria* (c1); it is rendered via http://iubio.bio.indiana.edu/treeapp/treeprint-form.html as Fig 7, and in newick format is:

```
newick.chr1.loose <- '(((tp3367_Italy:4551,tp1013_Wales:4954):5920,(((tp1007_Virginia:10,tp1012_Australia:29):9,
cat.hardwrap(newick.chr1.loose)

# (((tp3367_Italy:4551,tp1013_Wales:4954):5920,(((tp1007_Virginia:10,tp1012_Austra
# lia:29):9,(tp1015_Puget_Sound:90,tp1335_NY:13):11):320,tp1014_Gyre:22):3484):859
# 3,outgroup:0);</pre>
```

Chr 1 tree based on *medium criteria* (c2) has exactly the same topology is, although the branch lengths are different. As noted earlier, the length of the branch labeled "*" is probably inflated by lower coverage and higher error rate in 1014, which may mask further legitimate sharing between it and the other L-isloates. The branch lengths among the other 4 are too short for its topology to be convincing without a more rigorous analysis (e.g., a bootstrap test).

Chr1 tree, medium criteria, in newick format:

```
newick.chr1.med <- '(((tp3367_Italy:8813,tp1013_Wales:9652):9365,(((e_tp1007_Virginia:30,d_tp1012_Australia:61):1
cat.hardwrap(newick.chr1.med)

# (((tp3367_Italy:8813,tp1013_Wales:9652):9365,(((e_tp1007_Virginia:30,d_tp1012_Au
# stralia:61):19,(c_tp1015_Puget_Sound:207,b_tp1335_NY:41):18):1005,a_tp1014_Gyre:
# 61):3912):7054,outgroup:0);</pre>
```

NOTE: In early code, tree was not being recalculated; it was defined by constants in the following code chunk, hand-copied from the analysis above.

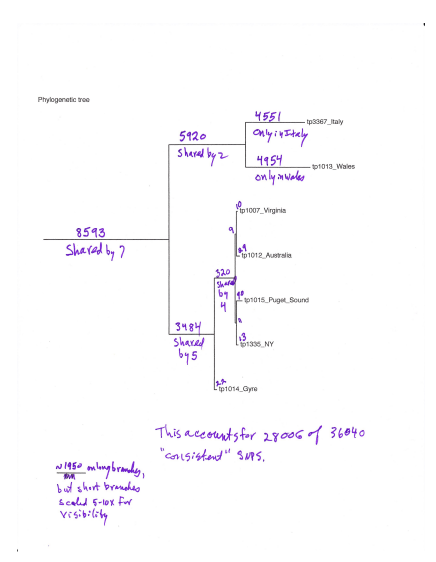


Figure 7: Inferred Tree, based on Chr1, un-q-filtered reads, loose criteria. (Note: to visually resolve the edges among the 5, their lengths were scaled by 5x - 10x in this figure, but not in the newick description shown in the text.)

```
# tree parameters as nested lists
    Internal nodes have subtrees sub1/2 and length
    Root has sub1/2, but no length
    Leaves have where, length, optionally, id, alt, nb. (Omit id for 'outgroup'. Use 'alt' for less formal labeling in cartoon version; it defaults to 'where'. Use 'nb' to add abode annotations for legend.)
# This hand-made version is now subsumed by make.tree; retained for comparison
tree.bv.hand <-
  list(
    sub1 = list(
      sub1 = list(
        sub1 = list(id=3367, length=8813, where='Venice, Italy', alt='Venice'),
        sub2 = list(id=1013, length=9652, where='Wales, UK'),
         length=9365),
             = list(
        sub1 = list(
          sub1 = list(
            sub1 = list(id=1007, length=30, nb='e', where='Virginia, USA'),
             sub2 = list(id=1012, length=61, nb='d', where='Perth, W. Australia', alt='Perth'),
             length=19),
           sub2 = list(
            sub1 = list(id=1015, length=207,nb='c', where='Washington, USA', alt='Puget Sound'),
             sub2 = list(id=1335, length=41, nb='b', where='New York, USA', alt='NY'),
             length=18).
         sub2 = list(id=1014, length=61, nb='a', where='N, Pacific Gyre').
         length=3912),
      length=7054),
    sub2 = list(length=0, where='outgroup')
# historical, format example, and debug help:
oldwick.medium <- '(((CCMP3367_Italy:8813,CCMP1013_Wales:9652):9365,(((e_CCMP1007_Virginia:30,d_CCMP1012_Australia:61):19,(c_CCMP</pre>
# with simpler labeling for cartoon simple.oldwick.medium <- '(((Italy:8813, Wales:9652):9365, (((Virginia:30, Australia:61):19, (Puget:207, NY:41):18):1005, Gyre:61):3912
cat.hardwrap(oldwick.medium)
# (((CCMP3367 Italy:8813,CCMP1013 Wales:9652):9365,(((e CCMP1007 Virginia:30,d CCM
# P1012_Australia:61):19,(c_CCMP1015_Puget_Sound:207,b_CCMP1335_NY:41):18):1005,a_
# CCMP1014_NPG:61):3912):7054,outgroup:0);
cat.hardwrap(simple.oldwick.medium)
# (((Italy:8813, Wales:9652):9365,(((Virginia:30, Australia:61):19,(Puget:207, NY:41)
# :18):1005, Gyre:61):3912):7054, outgroup:0);
```

Two other versions of the tree, for possible use in figs in the main paper.

Figure 8: [** as of 10/4/2015, this fig and next have stray stars on virginia, wales labels; probably due to hacking with commas in newick; not worth fixing unless we resurrect these trees for some purpose, but if so, see use of newick.name.undo in show.tree as probable fix. **]

```
tree.scale <- ifelse(which.snp.tables(string.val=F)[1]=='Chr1', 1, 10)
tree.x.lim <- 3e4 * tree.scale
the.simple.tree <- read.tree(text=simple.newick.medium)
plot(the.simple.tree, x.lim = tree.x.lim)
axis(1)</pre>
```

Figure 9:

```
plot(the.simple.tree, x.lim = tree.x.lim)
axis(1,(0:4)*7000*tree.scale,(0:4)*7000*tree.scale)
```

At some much earlier point, Tony ran the whole-genome version of the then-current code above, and manually entered tree branch lengths/legend for the resuting tree, shown in Fig 10. Code above can now automatically generate such a tree, but retain the following for comparison. The basic story seems clear—same topology and branch lengths scaled by about 10x, which is completely reasonable given that Chr1 is about 10% of the genome. Note that this tree is not being recalculated; it is defined by constants in the following code chunk.

```
fullgenome.newick.medium <- '(((3367_Italy:86155,1013_Wales:95697):89598,(((e_1007_VA:330,d_1012_Australia:632):1296,(c_1015_WA:2cat.hardwrap(fullgenome.newick.medium)

# (((3367_Italy:86155,1013_Wales:95697):89598,(((e_1007_VA:330,d_1012_Australia:63
# 2):1296,(c_1015_WA:2113,b_1335_NY:658):480):10059,a_1014_NPG:568):39517):69526,o
# utgroup:0);</pre>
```

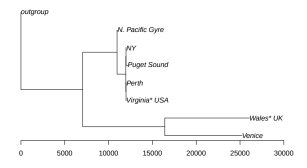


Figure 8: Tree based on unfiltered reads and medium SNP filters. "Lengths" are numbers of shared/private SNPs on Chr1. (no edge labels, nolegend)

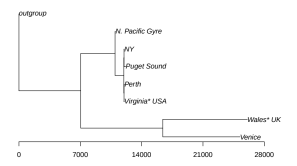


Figure 9: Tree based on unfiltered reads and medium SNP filters. "Lengths" are numbers of shared/private SNPs on Chr1. (no edge labels, no legend, short scale bar)

Figure 10:

```
library(ape)
the.fullgenome.tree <- read.tree(text=fullgenome.newick.medium)
plot(the.fullgenome.tree, x.lim = fullgenome.tree.x.lim)
axis(1) #; axis(2) useful only for placing labels
opar <- par(family='mono',cex=.8)
legend('topright', legend=fullgenome.legend.text)
par(opar)
for(i in seq(1,length(fullgenome.tree.labels)-2,by=3)){
   text(fullgenome.tree.labels[[i]], fullgenome.tree.labels[[i+1]], fullgenome.tree.labels[[i+2]])
}</pre>
```

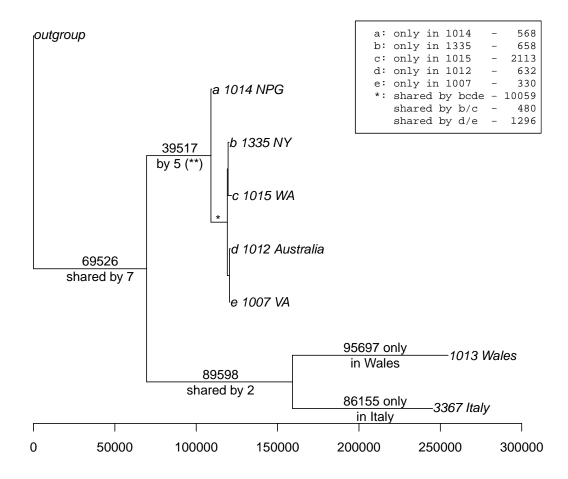


Figure 10: Tree based on unqfiltered reads and medium SNP filters. "Lengths" are numbers of shared/private SNPs genome-wide. (By-hand legacy version)