# Run dmc

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First I'll set up the working environment.

and load the necessary files

```
vcf<-parse.vcf("p4.upd.vcf") #this is the smaller dataset

vcf$SNP<-paste(vcf$`#CHROM`,vcf$POS,sep=".")
scaffs<-levels(as.factor(vcf[,1]))
scaffs[1:22]<-lgs
scaff.starts<-tapply(vcf$POS,vcf$`#CHROM`,max)
scaff.starts<-data.frame(rbind(cbind(names(scaff.starts)),stringsAsFactors = F)
locus.info<-c(colnames(vcf[1:9]),"SNP")</pre>
```

Now I'll go through the steps outline in https://github.com/kristinmlee/dmc/blob/master/dmc\_example.md

#### Calculate neutral F matrix

For this first step (calculating the neutral variance/covariance matrix, F), I'll use linkage groups that have no or few shared outlier loci. I need to (1) Calculate allele frequencies for each population, (2) specify a vector of sample sizes for each population, and (3) specify a string for filename for output. This step is baseline for all parameter sensitivity analyses.

```
calc.allFreqs<-function(vcf,pop.list, pop.labs){</pre>
  allFreqs<-do.call(rbind,lapply(pop.list, function(pop){</pre>
    this.vcf<-cbind(vcf[,1:9],vcf$SNP,vcf[,grep(pop,colnames(vcf))])
    afs<-do.call(rbind,apply(this.vcf,1,calc.afs.vcf))
    return(afs$RefFreq)
  }))
  colnames(allFreqs)<-vcf$SNP</pre>
  rownames(allFreqs)<-pop.labs</pre>
  return(allFreqs)
}
#calculate selected allele frequencies
LG4<-calc.allFreqs(vcf[vcf$\*CHROM\` == "LG4" & vcf$POS < 120000,],pop.list,pop.labs)
saveRDS(LG4, "dmc/selectedRegionAlleleFreqs_p4LG4.RDS")
saveRDS(vcf[vcf$`#CHROM`=="LG4","POS"],"dmc/selectedRegionPositions_p4LG4.RDS")
#calculate neutral allele frequencies
allFreqs<-calc.allFreqs(vcf[vcf$`#CHROM` %in%
                               c("LG3","LG5","LG6","LG7","LG8","LG9","LG11","LG15","LG16","LG17","LG19",
                   pop.list,pop.labs)
saveRDS(allFreqs, "dmc/neutralAlleleFreqs_p4LG4.RDS")
allFreqs<-readRDS("dmc/neutralAlleleFreqs p4LG4.RDS")
```

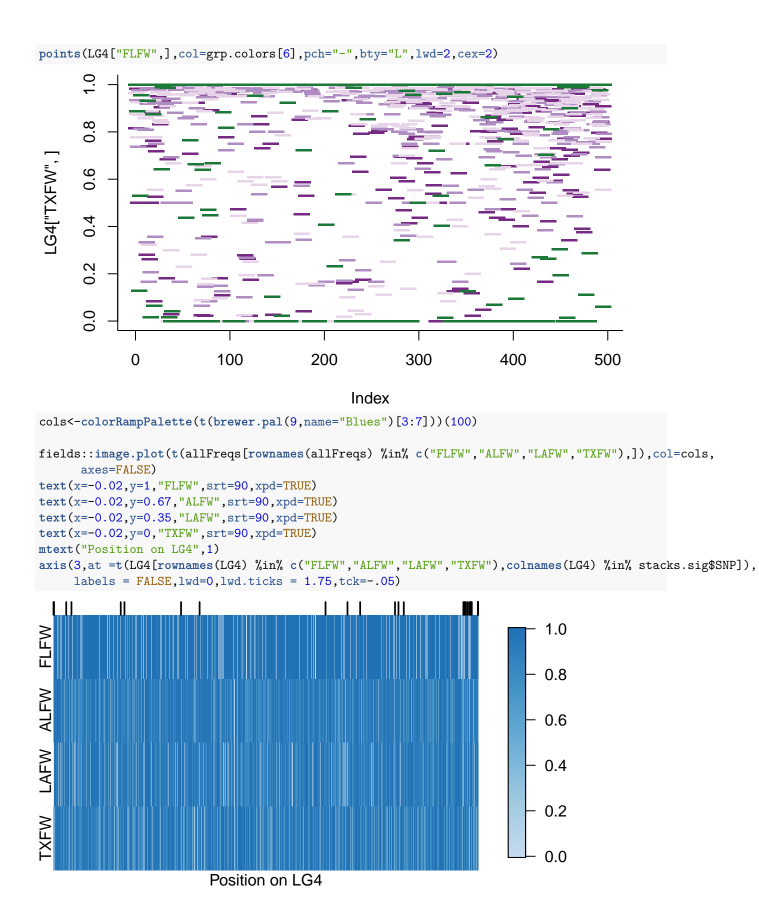
Once those are calculated, I can calculate the neutral F matrix, in addition to its determinant and inverse.

```
sampleSizes<-unlist(lapply(pop.list,function(pop){
   n<-length(grep(pop,colnames(vcf)))</pre>
```

```
numPops<-16
#calculate the neutral F matrix
saveRDS(sampleSizes,"dmc/sampleSizes.RDS")
neutralF filename<-"dmc/neutralF p4LG4 50" #for 50 positions
source("../programs/dmc-master/calcNeutralF.R")
F estimate <- read RDS ("dmc/neutralF p4LG4 50.RDS")
#calculate the determinant and inverse of the neutral F matrix
M <- numPops
Tmatrix \leftarrow matrix(data = rep(-1 / M, (M - 1) * M), nrow = M - 1, ncol = M)
diag(Tmatrix) = (M - 1) / M
sampleErrorMatrix = diag(1/sampleSizes, nrow = numPops, ncol = numPops)
det_FOmegas_neutral = det(Tmatrix %*% (F_estimate + sampleErrorMatrix) %*% t(Tmatrix))
saveRDS(det_FOmegas_neutral, "dmc/det_FOmegas_neutral_p4LG4_50.RDS")
inv_FOmegas_neutral = ginv(Tmatrix %*% (F_estimate + sampleErrorMatrix) %*% t(Tmatrix))
saveRDS(inv_FOmegas_neutral, "dmc/inv_FOmegas_neutral_p4LG4_50.RDS")
selSite = positions[seq(1, length(positions[positions<12000000]), length.out = 100)]
allFreqs<-readRDS("dmc/neutralAlleleFreqs_p4LG4.RDS")</pre>
#calculate the neutral F matrix
sampleSizes<-readRDS("dmc/sampleSizes.RDS")</pre>
neutralF_filename<-"dmc/neutralF_p4LG4_100" #for 50 positions
source("../programs/dmc-master/calcNeutralF.R")
F_estimate<-readRDS("dmc/neutralF_p4LG4_100.RDS")
#calculate the determinant and inverse of the neutral F matrix
M <- numPops
Tmatrix \leftarrow matrix(data = rep(-1 / M, (M - 1) * M), nrow = M - 1, ncol = M)
diag(Tmatrix) = (M - 1) / M
sampleErrorMatrix = diag(1/sampleSizes, nrow = numPops, ncol = numPops)
det FOmegas neutral = det(Tmatrix %*% (F estimate + sampleErrorMatrix) %*% t(Tmatrix))
saveRDS(det_FOmegas_neutral, "dmc/det_FOmegas_neutral_p4LG4_100.RDS")
inv_FOmegas_neutral = ginv(Tmatrix %*% (F_estimate + sampleErrorMatrix) %*% t(Tmatrix))
saveRDS(inv_F0megas_neutral, "dmc/inv_F0megas_neutral_p4LG4_100.RDS")
Note: I had to use to allow read/write in the scripts.
Let's look at the allele frequencies on LG4. This will help inform the hypotheses I test with dmc.
stacks.sig<-read.delim("p4.stacks.sig.snps.txt")</pre>
stacks.sig$SNP<-paste(stacks.sig$Chr,(stacks.sig$BP+1),sep=".")</pre>
LG4<-readRDS("dmc/selectedRegionAlleleFreqs_p4LG4.RDS")
allFreqs<-readRDS("dmc/neutralAlleleFreqs_p4LG4.RDS")
# LG4[rownames(LG4) %in% c("FLFW", "ALFW", "LAFW", "TXFW"), colnames(LG4) %in% stacks.sig$SNP]
# LG4[rownames(LG4) %in% c("FLFW", "ALFW", "LAFW", "TXFW"),
#
           which(colnames(LG4) %in% stacks.sig$SNP)[1]:
             which(colnames(LG4) %in% stacks.sig$SNP)[length(which(colnames(LG4) %in% stacks.sig$SNP))]
grp.colors<-c('#762a83','#af8dc3','#e7d4e8','#d9f0d3','#7fbf7b','#1b7837')</pre>
plot(LG4["TXFW",],col=grp.colors[1],pch="-",bty="L",lwd=2,cex=2)
points(LG4["LAFW",],col=grp.colors[2],pch="-",bty="L",lwd=2,cex=2)
```

return(2\*n) }))

points(LG4["ALFW",],col=grp.colors[3],pch="-",bty="L",lwd=2,cex=2)



This shows that the FLFW site has somewhat different allele frequencies than the other sites. This suggests that I should test to see whether there was an independent mutation in the FLFW population and the other three have a single sweep.

#### Calculate F(S) matrices

We generate matrices for the following five models: 1. All selected pops have independent mutations of beneficial allele 2. All selected pops share beneficial allele via migration 3. Beneficial allele was standing the ancestor of all selected pops 4. There was an independent mutation in FLFW and the other three share a sweep. \*5. FLFW has an independent mutation and the others share a beneficial allele via standing variation

I've written a giant function to run all five models and calculate their composite likelihoods. The composite likelihoods include knowing how far away we are from the proposed selected site. Therefore, the first step for calculating them is to randomize with respect to the reference allele.

The inverses and determinants generated can be used to calculate the log-likelihood of a site a given distance away from a proposed selected site. To do this, we sum the log-likelihoods across all SNPs in the window to obtain a composite log-likelihood under a given convergence model with a set of parameters for a proposed selected site.

```
run.dmc<-function(F_estimate,out_name,positions, sampleSizes,selSite=NA,nselsites=50,rec =2.17*10^-8,
                  Ne = 8.3*10^6, selPops = c(3,5,7,16), numBins = 1000, numPops = 16,
                  sels = c(1e-4, 1e-3, 0.01, seq(0.02, 0.14, by = 0.01), seq(0.15, 0.3, by = 0.05),
                           seq(0.4, 0.6, by = 0.1)),
                  times = c(0, 5, 25, 50, 100, 500, 1000, 1e4, 1e6),
                  migs = c(10^-(seq(5, 1, by = -2)), 0.5, 1), mod4_sets=list(c(3,5,7),16),
                  mod1=TRUE,mod2=TRUE,mod3=TRUE,mod4=TRUE,mod5=TRUE,complike=TRUE,
                  neutral det name="dmc/det FOmegas neutral p4LG4.RDS",
                  neutral inv name="dmc/inv FOmegas neutral p4LG4.RDS"){
  #make all the parameters global
  F_estimate << - F_estimate
  positions<<-positions
  sampleSizes<<-sampleSizes
  Ne<<-Ne
  rec<<-rec
  selPops<<-selPops
  numBins<<-numBins
  numPops<<-numPops
  sels<<-sels
  times<<-times
  migs<<-migs
  neutral det name <<-neutral det name
  neutral_inv_name<<-neutral_inv_name
  print(paste(neutral_det_name,neutral_inv_name,sep=","))
  #set up parameters
  M <<- numPops
  Tmatrix << matrix(data = rep(-1 / M, (M - 1) * M), nrow = M - 1, ncol = M)
  diag(Tmatrix) = (M - 1) / M
  gs << -c(1/(2*Ne), 10^-(4:1))
  sampleErrorMatrix <<- diag(1/sampleSizes, nrow = numPops, ncol = numPops)</pre>
  if(is.na(selSite[1])) selSite=seq(min(positions), max(positions), length.out = nselsites)
  selSite<<-selSite
  sources <<- selPops
  sets<<-mod4_sets
```

```
#save params
params<-list(F_estimate,out_name,positions, sampleSizes,selSite,rec,</pre>
            Ne, selPops, numBins, numPops, sels, times, gs, migs, mod4 sets)
names(params)<-c("F_estimate","out_name","positions", "sampleSizes","selSite","rec",</pre>
                "Ne", "selPops", "numBins", "numPops", "sels", "times", "gs", "migs", "mod4_sets")
print(params)
source("../programs/dmc-master/genSelMatrices_individualModes.R")
if(mod1==TRUE){
 FOmegas_ind = lapply(sels, function(sel) {
   calcFOmegas_indSweeps(sel)
 })
 saveRDS(FOmegas_ind, paste("dmc/FOmegas_ind_",out_name,".RDS",sep=""))
  #model 1 determinant
 det_FOmegas_ind = lapply(FOmegas_ind, function(sel) {
   lapply(sel, function(dist) {
     det(dist)
   })
 })
 saveRDS(det_FOmegas_ind, paste("dmc/det_FOmegas_ind_",out_name,".RDS",sep=""))
  #model 1 inverse
 inv_FOmegas_ind = lapply(FOmegas_ind, function(sel) {
   lapply(sel, function(dist) {
     ginv(dist)
   })
 })
  saveRDS(inv_FOmegas_ind, paste("dmc/inv_FOmegas_ind_",out_name,".RDS",sep=""))
if(mod2==TRUE){
 FOmegas_mig = lapply(sels ,function(sel) {
   lapply(migs, function(mig) {
     lapply(sources, function(my.source) {
       calcFOmegas_mig(sel, mig, my.source)
     })
   })
 })
 saveRDS(FOmegas_mig, paste("dmc/FOmegas_mig_",out_name,".RDS",sep=""))
 det_FOmegas_mig = lapply(FOmegas_mig, function(sel) {
   lapply(sel, function(mig) {
     lapply(mig, function(source) {
       lapply(source, function(dist) {
         det(dist)
       })
     })
   })
 })
  saveRDS(det_FOmegas_mig, paste("dmc/det_FOmegas_mig_",out_name,".RDS",sep=""))
```

```
inv_FOmegas_mig = lapply(FOmegas_mig, function(sel) {
   lapply(sel, function(mig) {
     lapply(mig, function(source) {
       lapply(source, function(dist) {
         ginv(dist)
       })
     })
   })
 })
 saveRDS(inv_FOmegas_mig, paste("dmc/inv_FOmegas_mig_",out_name,".RDS",sep=""))
if(mod3==TRUE)
 FOmegas_sv = lapply(sels, function(sel) {
   lapply(gs, function(g) {
     lapply(times, function(time) {
       lapply(sources, function(my.source) {
         calcFOmegas_stdVar.source(sel, g, time, my.source)
       })
     })
   })
 })
 saveRDS(FOmegas_sv, paste("dmc/FOmegas_sv_",out_name,".RDS",sep=""))
  #determinant
 det_FOmegas_sv = lapply(FOmegas_sv, function(sel) {
   lapply(sel, function(g) {
     lapply(g, function(time) {
       lapply(time, function(my.source) {
         lapply(my.source, function(dist) {
           det(dist)
         })
       })
     })
   })
 saveRDS(det FOmegas sv, paste("dmc/det FOmegas sv ",out name,".RDS",sep=""))
  #inverse
 inv_FOmegas_sv = lapply(FOmegas_sv, function(sel) {
   lapply(sel, function(g) {
     lapply(g, function(time) {
       lapply(time, function(my.source) {
         lapply(my.source, function(dist) {
           ginv(dist)
         })
       })
     })
   })
 })
  saveRDS(inv_F0megas_sv, paste("dmc/inv_F0megas_sv_",out_name,".RDS",sep=""))
}
```

```
if(mod4==TRUE){
 source("../programs/dmc-master/genSelMatrices multipleModes.R")
 my.modes migInd=c("mig","ind")
 #the parameters time and g are not involved in the migration model so we only loop over
 ## the first element of these vectors
 FOmegas_mixed_migInd = lapply(sels ,function(sel) {
   lapply(gs[1], function(g) {
     lapply(times[1], function(time) {
       lapply(migs, function(mig) {
         lapply(sources, function(my.source) {
           calcFOmegas_mixed(sel, g, time, mig, my.source, my.modes_migInd)
         })
       })
     })
   })
 })
 saveRDS(FOmegas mixed migInd, paste("dmc/FOmegas mixed migInd ",out name,".RDS",sep=""))
 detFOmegas_mixed_migInd = lapply(FOmegas_mixed_migInd, function(sel) {
   lapply(sel, function(g) {
     lapply(g, function(time) {
       lapply(time, function(mig) {
         lapply(mig, function(source) {
           lapply(source, function(dist) {
             det(dist)
           })
         })
       })
     })
   })
 })
  saveRDS(detFOmegas_mixed_migInd, paste("dmc/det_FOmegas_mixed_migInd_",out_name,".RDS",sep=""))
 invFOmegas mixed migInd = lapply(FOmegas mixed migInd, function(sel) {
   lapply(sel, function(g) {
     lapply(g, function(time) {
       lapply(time, function(mig) {
         lapply(mig, function(source) {
           lapply(source, function(dist) {
             ginv(dist)
           })
         })
       })
     })
   })
 })
  saveRDS(invFOmegas_mixed_migInd, paste("dmc/inv_FOmegas_mixed_migInd_",out_name,".RDS",sep=""))
}
```

```
if(mod5==TRUE){
 my.modes svInd = c("sv", "ind")
 #the parameter mig is not involved in the standing variant model so we only loop over
 ## the first element of this vector
 FOmegas_mixed_svInd = lapply(sels ,function(sel) {
   lapply(gs, function(g) {
     lapply(times, function(time) {
       lapply(migs[1], function(mig) {
         lapply(sources, function(my.source) {
           calcFOmegas_mixed(sel, g, time, mig, my.source, my.modes_svInd)
         })
       })
     })
   })
 })
 saveRDS(FOmegas_mixed_svInd, paste("dmc/FOmegas_mixed_svInd_",out_name,".RDS",sep=""))
 detFOmegas mixed svInd = lapply(FOmegas mixed svInd, function(sel) {
   lapply(sel, function(g) {
     lapply(g, function(time) {
       lapply(time, function(mig) {
         lapply(mig, function(source) {
           lapply(source, function(dist) {
             det(dist)
           })
         })
       })
     })
   })
 saveRDS(detFOmegas_mixed_svInd, paste("dmc/det_FOmegas_mixed_svInd_",out_name,".RDS",sep=""))
 invFOmegas_mixed_svInd = lapply(FOmegas_mixed_svInd, function(sel) {
   lapply(sel, function(g) {
     lapply(g, function(time) {
       lapply(time, function(mig) {
         lapply(mig, function(source) {
           lapply(source, function(dist) {
             ginv(dist)
           })
         })
       })
     })
   })
 })
 saveRDS(invFOmegas_mixed_svInd, paste("dmc/inv_FOmegas_mixed_svInd_",out_name,".RDS",sep=""))
if(complike==TRUE){
  #randomize allele freqs
```

```
freqs_notRand = readRDS("dmc/selectedRegionAlleleFreqs_p4LG4.RDS")
randFreqs = apply(freqs_notRand, 2, function(my.freqs) {
  if(runif(1) < 0.5) {
    my.freqs = 1 - my.freqs
  }
  my.freqs
})
saveRDS(randFreqs, paste("dmc/selectedRegionAlleleFreqsRand ",out name,".RDS",sep=""))
freqs<<-randFreqs
#calc the likelihoods
source("../programs/dmc-master/calcCompositeLike.R")
## Neutral model
det_FOmegas_neutral = readRDS(neutral_det_name)
inv_FOmegas_neutral = readRDS(neutral_inv_name)
compLikelihood_neutral = lapply(1 : length(selSite), function(j) {
  calcCompLikelihood_neutral(j, det_FOmegas_neutral, inv_FOmegas neutral)
})
saveRDS(compLikelihood_neutral, paste("dmc/compLikelihood_neutral_",out_name,".RDS",sep=""))
## Model 1
det_FOmegas_ind = readRDS(paste("dmc/det_FOmegas_ind_",out_name,".RDS",sep=""))
inv FOmegas ind = readRDS(paste("dmc/inv FOmegas ind ",out name,".RDS",sep=""))
compLikelihood ind = lapply(1 : length(selSite), function(j) {
  lapply(1 : length(sels), function(sel) calcCompLikelihood_1par(j, det_FOmegas_ind,
                                                                 inv FOmegas ind, sel))
saveRDS(compLikelihood_ind, paste("dmc/compLikelihood_ind_",out_name,".RDS",sep=""))
## Model 2
det_FOmegas_mig = readRDS(paste("dmc/det_FOmegas_mig_",out_name,".RDS",sep=""))
inv_FOmegas_mig = readRDS(paste("dmc/inv_FOmegas_mig_",out_name,".RDS",sep=""))
compLikelihood_mig = lapply(1 : length(selSite), function(j) {
  lapply(1 : length(sels), function(sel) {
    lapply(1 : length(migs), function(mig) {
      lapply(1 : length(sources), function(my.source) {
        calcCompLikelihood_3par(j, det_FOmegas_mig, inv_FOmegas_mig, sel, mig,
                                my.source)
     })
   })
  })
saveRDS(compLikelihood_mig, paste("dmc/compLikelihood_mig_",out_name,".RDS",sep=""))
## Model 3
det_FOmegas_sv = readRDS(paste("dmc/det_FOmegas_sv_",out_name,".RDS",sep=""))
inv_FOmegas_sv = readRDS(paste("dmc/inv_FOmegas_sv_",out_name,".RDS",sep=""))
compLikelihood_sv = lapply(1 : length(selSite), function(j) {
  lapply(1 : length(sels), function(sel) {
    lapply(1 : length(gs), function(g) {
      lapply(1 : length(times), function(t) {
```

```
lapply(1: length(sources), function(my.source) {
          calcCompLikelihood_4par(j, det_FOmegas_sv, inv_FOmegas_sv, sel, g, t,
                                  my.source)
       })
     })
   })
 })
})
saveRDS(compLikelihood sv, paste("dmc/compLikelihood sv ",out name,".RDS",sep=""))
## Model 4
det_FOmegas_mixed_migInd = readRDS(paste("dmc/det_FOmegas_mixed_migInd_",out_name,".RDS",sep=""))
inv_FOmegas_mixed_migInd = readRDS(paste("dmc/inv_FOmegas_mixed_migInd_",out_name,".RDS",sep=""))
# same trick as above (the parameters time and q are not involved in the migration
## model so we only loop over the first element of these vectors)
# now save lists for each proposed selected site (may want to do this for other
## models/more elegantly depending on density of parameter space)
for(j in 1 : length(selSite)) {
  compLikelihood_mixed_migInd = lapply(1 : length(sels), function(sel) {
    lapply(1 : length(gs[1]), function(g) {
      lapply(1 : length(times[1]), function(t) {
        lapply(1 : length(migs), function (mig) {
          lapply(1: length(sources), function(my.source) {
            calcCompLikelihood_5par(j, det_FOmegas_mixed_migInd,
                                    inv FOmegas mixed migInd, sel, g, t, mig,
                                    my.source)
          })
        })
     })
   })
  })
  saveRDS(compLikelihood_mixed_migInd,
          paste("dmc/compLikelihood_mixed_migInd_",out_name,"_selSite", j, ".RDS",
                sep = "")
}
## Model 5
det_FOmegas_mixed_svInd = readRDS(paste("dmc/det_FOmegas_mixed_svInd_",out_name,".RDS",sep=""))
inv FOmegas mixed svInd = readRDS(paste("dmc/inv FOmegas mixed svInd ",out name,".RDS",sep=""))
#same trick as above (the parameter mig is not involved in the migration model so we
##only loop over the first element of this vector)
# now save lists for each proposed selected site (may want to do this for other
## models/more elegantly depending on density of parameter space)
for(j in 1 : length(selSite)) {
  compLikelihood_mixed_svInd = lapply(1 : length(sels), function(sel) {
    lapply(1 : length(gs), function(g) {
      lapply(1 : length(times), function(t) {
        lapply(1 : length(migs[1]), function (mig) {
          lapply(1: length(sources), function(my.source) {
            calcCompLikelihood_5par(j, det_FOmegas_mixed_svInd,
                                    inv_FOmegas_mixed_svInd, sel, g, t, mig,
```

```
my.source)
           })
         })
       })
     })
   })
    saveRDS(compLikelihood_mixed_svInd,
            paste("dmc/compLikelihood mixed svInd ",out name," selSite", j, ".RDS",
                  sep = "")
 }
  #combine models 4 and 5 output
 ## Model 4
  compLikelihood_mixed_migInd_all = lapply(1: length(selSite), function(i) {
    readRDS(paste("dmc/compLikelihood_mixed_migInd_",out_name,"_selSite", i, ".RDS",
                  sep = ""))
 })
 saveRDS(compLikelihood_mixed_migInd_all, paste("dmc/compLikelihood_mixed_migInd_",out_name,".RDS",s
 ## Model 5
  compLikelihood mixed svInd all = lapply(1: length(selSite), function(i) {
   readRDS(paste("dmc/compLikelihood_mixed_svInd_",out_name,"_selSite", i, ".RDS",
                  sep = "")
 })
  saveRDS(compLikelihood_mixed_svInd_all, paste("dmc/compLikelihood_mixed_svInd_",out_name,".RDS",sep
return(params)
```

I have to set some parameters before running the function, including the sets of parameters I want to try different settings for.

And I'm going to run it with a variety of effective population sizes and recombination rates to see how robust the models are to that parameter.

```
Nes <- c(8.3*10^3,8.3*10^4,8.3*10^5,8.3*10^6)
rs<-c(2.17*10^-8,2*10^-7,6*10^-6,5.6*10^-6)
dmc.out<-lapply(rs,function(r){</pre>
 rec<-r
 rname < -gsub("(\d).*(\d)$","\1_\2",as.character(r))
 lapply(Nes, function(ne){
   out_name<-paste("p4LG4_",ne,"_",rname,sep="")</pre>
   dir<-getwd()
   print(paste("running",out_name,"in",dir,sep=" "))
   p<-run.dmc(F_estimate = F_estimate,out_name = out_name,positions = positions,sampleSizes = sampleSizes
              selSite=selSite,rec =rec,
              Ne = ne,selPops = selPops,numBins = numBins,numPops = numPops,
              sels = sels, times = times,
              migs = migs,mod4_sets=mod4_sets,
              neutral_det_name="dmc/det_FOmegas_neutral_p4LG4_50.RDS",
              neutral_inv_name = "dmc/inv_FOmegas_neutral_p4LG4_50.RDS")
   return(p)
 })
})
print("Done running dmc")
```

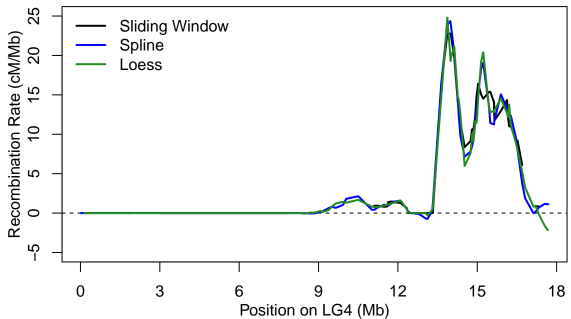
I'm running this one on abies using nohup Rscript ~/Projects/popgen/scripts/002\_run\_dmc.R >> ~/Projects/popgen/dmc\_ne.log 2>&1 &.

#### Estimate recombination rates

```
ssc.map<-read.delim("../../scovelli_genome/SSC_linkage_map_20160701.txt",header=T)
colnames(ssc.map)<-c("map","mkr","gen","scaffold","scaffold_orientation","scaffold.bp","phys","node")</pre>
ssc.map$set<-"Syngnathus.scovelli"
map<-ssc.map[,c("set","map","mkr","phys","gen")]</pre>
write.table(map,"../../scovelli_genome/ssc_map.txt",sep=" ",row.names=FALSE, col.names = TRUE)
library(MareyMap)
startMareyMapGUI()
#using the GUI, I ran all three types of recombination rate estimators and saved the MapSet
ssc.map<-read.delim("../../scovelli_genome/ssc_MareyMap.txt",comment.char = "#",sep=" ")
#genome-wide
r<-data.frame(Genome_wide=c(mean(ssc.map$slidingwindow,na.rm = TRUE)/1000000,
                            mean(ssc.map$spline,na.rm = TRUE)/1000000,
                            mean(ssc.map$loess,na.rm = TRUE)/1000000),
              LG4=c(mean(ssc.map$slidingwindow[ssc.map$map=="LG4"],na.rm = TRUE)/1000000,
                            mean(ssc.map$spline[ssc.map$map=="LG4"],na.rm = TRUE)/1000000,
                            mean(ssc.map$loess[ssc.map$map=="LG4"],na.rm = TRUE)/1000000),
              LG4_first12Mb=c(mean(ssc.map$slidingwindow[ssc.map$map=="LG4" & ssc.map$phys<12*10^6],na..
                            mean(ssc.map$spline[ssc.map$map=="LG4"& ssc.map$phys<12*10^6],na.rm = TRUE)
                            mean(ssc.map$loess[ssc.map$map=="LG4"& ssc.map$phys<12*10^6],na.rm = TRUE)/
              row.names = c("Sliding Window", "Spline", "Loess"))
kable(r,caption="Recombination rate estimates")
```

Table 1: Recombination rate estimates

	Genome_wide	LG4	LG4_first12Mb
Sliding Window	5.8e-06	3.7e-06	2e-07
Spline	5.6e-06	3.0e-06	2e-07
Loess	5.6e-06	3.0e-06	2e-07



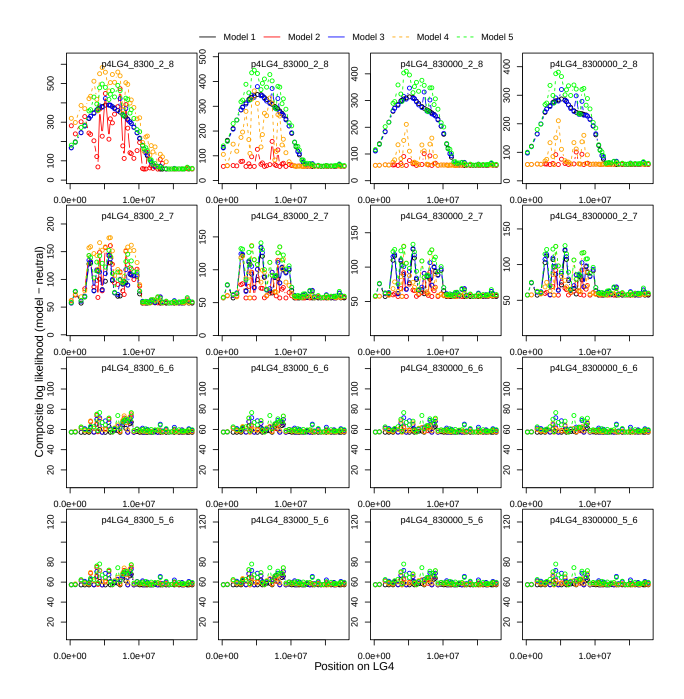
#### Evaluate the output of the initial dmc runs

Plot maximum composite likelihood ratios for models over proposed selected sites

```
calc.max.complike<-function(pattern,neutral_name=NA){
  #read in composite likelihood files and calculate max for all proposed selected sites
  if(is.na(neutral_name)){
    compLikelihood_neutral = readRDS(paste("dmc/compLikelihood_neutral_",pattern,".RDS",sep=""))
}else{</pre>
```

```
compLikelihood_neutral = readRDS(neutral_name)
  }
  compLikelihood_neutral_site = sapply(1 : length(compLikelihood_neutral), function(i) {
    max(unlist(compLikelihood_neutral[[i]]))
  })
  compLikelihood ind = readRDS(paste("dmc/compLikelihood ind ",pattern,".RDS",sep=""))
  compLikelihood_ind_site = sapply(1 : length(compLikelihood_ind), function(i) {
    max(unlist(compLikelihood ind[[i]]))
  })
  compLikelihood_mig = readRDS(paste("dmc/compLikelihood_mig_",pattern,".RDS",sep=""))
  compLikelihood_mig_site = sapply(1 : length(compLikelihood_mig), function(i) {
    max(unlist(compLikelihood_mig[[i]]))
  })
  compLikelihood_sv = readRDS(paste("dmc/compLikelihood_sv_",pattern,".RDS",sep=""))
  compLikelihood_sv_site = sapply(1 : length( compLikelihood_sv), function(i) {
    max(unlist(compLikelihood_sv[[i]]))
  })
  compLikelihood_mixed_migInd = readRDS(paste("dmc/compLikelihood_mixed_migInd_",pattern,".RDS",sep="")
  compLikelihood_mixed_migInd_site = sapply(1 : length(compLikelihood_mixed_migInd), function(i) {
    max(unlist(compLikelihood mixed migInd[[i]]))
  })
  compLikelihood_mixed_svInd = readRDS(paste("dmc/compLikelihood_mixed_svInd_",pattern,".RDS",sep=""))
  compLikelihood_mixed_svInd_site = sapply(1 : length(compLikelihood_mixed_svInd), function(i) {
    max(unlist(compLikelihood_mixed_svInd[[i]]))
  })
  return(list(max.likes=c(ind=(compLikelihood_ind_site - compLikelihood_neutral_site),
           mig=(compLikelihood_mig_site - compLikelihood_neutral_site),
           sv=(compLikelihood_sv_site - compLikelihood_neutral_site),
           migInd=(compLikelihood_mixed_migInd_site - compLikelihood_neutral_site),
           svInd=(compLikelihood_mixed_svInd_site - compLikelihood_neutral_site)),
         max.complikes=c(neutral=compLikelihood_neutral,ind=compLikelihood_ind,
           mig=compLikelihood_mig,
           sv=compLikelihood_sv,
           migInd=compLikelihood mixed migInd,
           svInd=compLikelihood_mixed_svInd)))
}
plot.complike<-function(pattern, selSite, leg=TRUE, lab=TRUE) {</pre>
  max.complikes<-calc.max.complike(pattern)</pre>
  max.likes<-max.complikes[[1]]</pre>
  plot_range = range(max.likes)
  if(lab==TRUE){
   x.lab<-"Proposed position selected site"
    y.lab<-"Composite log-likelihood (model - neutral)"</pre>
  } else{
   x.lab<-y.lab<-""
```

```
plot(selSite, max.likes[grep("ind",names(max.likes))], type = "b",
       ylim = c(plot_range[1] - 50, plot_range[2] + 50),
       xlab = x.lab,
       ylab = y.lab)
  lines(selSite, max.likes[grep("mig\\d+",names(max.likes))], col = "red",
        type = "b")
  lines(selSite, max.likes[grep("sv\\d+",names(max.likes))], col = "blue",
        type = "b")
  lines(selSite, max.likes[grep("migInd",names(max.likes))],
        col = "orange", lty = 2, type = "b")
  lines(selSite, max.likes[grep("svInd",names(max.likes))],
        col = "green", lty = 2, type = "b")
  legend("top",bty='n',legend = pattern)
  if(isTRUE(leg)){
    legend("topright", col = c("black", "red", "blue", "orange", "green"),
           lty = c(rep(1, 3), rep(2, 2)), sapply(1 : 5, function(i) paste("Model", i)))
  }
  return(list(max.complikes[[2]],max.likes,pattern))
patterns<-c("p4LG4_8300_2_8", "p4LG4_83000_2_8", "p4LG4_830000_2_8", "p4LG4_8300000_2_8",
            "p4LG4_8300_2_7", "p4LG4_83000_2_7", "p4LG4_830000_2_7", "p4LG4_830000_2_7",
            "p4LG4_8300_6_6", "p4LG4_83000_6_6", "p4LG4_830000_6_6", "p4LG4_8300000_6_6",
            "p4LG4_8300_5_6","p4LG4_83000_5_6","p4LG4_830000_5_6","p4LG4_830000_5_6")
selSite = positions[seq(1, length(positions), length.out = 50)]
par(mfrow=c(4,4), mar=c(1.5,1.5,0.5,0.5), oma=c(2,2,2,2))
ne.complikes<-lapply(patterns,plot.complike,leg=FALSE,lab=FALSE,selSite=selSite)</pre>
mtext("Position on LG4",1,outer=TRUE,cex=0.8,line=.5)
mtext("Composite log likelihood (model - neutral)",2,outer=T,cex=0.8,line=.5)
par(fig=c(0,1,0,1),oma=c(0,0,0,0),mar=c(0,0,0,0),new=TRUE)
plot(0,0,type='n',bty='n',xaxt='n',yaxt='n')
legend("top", col = c("black", "red", "blue", "orange", "green"),ncol=5,bty='n',
           lty = c(rep(1, 3), rep(2, 2)), sapply(1 : 5, function(i) paste("Model", i)))
```



#### Get maximum composite likelihood estimates

```
source("../programs/dmc-master/getMCLE.R")
  ## Model 1
  m1<-cbind(getMCLEind(max.complikes[grep("ind",names(max.complikes))], selSite, sels),
            maxG=NA,maxTime=NA,maxSource=NA)
  ## Model 2
  m2<-cbind(getMCLEmig(max.complikes[grep("mig\\d+",names(max.complikes))], selSite, sels, migs, source
            maxG=NA)
  ## Model 3
  m3<-getMCLEsv_source(max.complikes[grep("sv\\d+",names(max.complikes))], selSite, sels, gs, times, so
  ## Model 4
  m4<-getMCLEmixed(max.complikes[grep("migInd",names(max.complikes))], selSite, sels, gs[1], times[1],
  ## Model 5
  m5<-getMCLEmixed(max.complikes[grep("svInd",names(max.complikes))], selSite, sels, gs, times, migs[1]
  mcles < -rbind(m1, m2, m3, m4, m5)
  rownames(mcles)<-c("Model1", "Model2", "Model3", "Model4", "Model5")</pre>
  return(mcles)
}
rs<-c("2_8"=2.17*10^-8,"2_7"=2*10^-7,"6_6"=6*10^-6,"5_6"=5.6*10^-6)
ne.mcles<-lapply(ne.complikes, function(m.complikes){</pre>
  pattern<-m.complikes[[3]]</pre>
  print(pattern)
  Ne<-as.numeric(gsub("p4LG4_(\\d+)_.*","\\1",pattern))</pre>
  r<-gsub("p4LG4_\\d+_(\\d_\\d)","\\1",pattern)
  rec<-rs[r]
  max.comp<-m.complikes[[1]]</pre>
  mcle<-get.mcle(max.comp,Ne=Ne,rec=rec)</pre>
  return(list(mcle,Ne,rec))
})
## [1] "p4LG4_8300_2_8"
## [1] "p4LG4_83000_2_8"
## [1] "p4LG4_830000_2_8"
## [1] "p4LG4_8300000_2_8"
## [1] "p4LG4_8300_2_7"
## [1] "p4LG4_83000_2_7"
## [1] "p4LG4_830000_2_7"
## [1] "p4LG4_8300000_2_7"
## [1] "p4LG4_8300_6_6"
## [1] "p4LG4_83000_6_6"
## [1] "p4LG4_830000_6_6"
## [1] "p4LG4_8300000_6_6"
## [1] "p4LG4_8300_5_6"
## [1] "p4LG4_83000_5_6"
## [1] "p4LG4 830000 5 6"
## [1] "p4LG4_8300000_5_6"
```

```
lapply(ne.mcles,function(mcle){ kable(mcle[[1]]) })
## $p4LG4_8300_2_8
##
##
##
         maxLoc maxSel maxG maxTime maxSource
## ----- ----- -----
## Model1 5641459
                   0.6
                            NA
                                    NA
                                              NΑ
        7263580
                   0.6 1.00e-05
                                     3
## Model2
                                              NA
## Model3 5122070
                                   50
                   0.6 1.00e-04
                                               5
## Model4 4665911 0.6 6.02e-05
## Model5 4665911 0.6 1.00e-04
                                    0
                                               5
                                    5
                                               3
##
## $p4LG4_83000_2_8
##
##
##
         maxLoc maxSel maxG maxTime maxSource
## -----
## Model1 5122070 0.6 NA
                                 NA
                                           NA
        7263580
                   0.4 1e-05
                                  3
## Model2
                                           NA
## Model3 5122070
                   0.6 6e-06
                                 50
                                            5
## Model4 4665911
                  0.6 6e-06
                                  0
                                            3
## Model5 4665911
                   0.6 6e-06
                                  5
                                            3
##
## $p4LG4_830000_2_8
##
##
##
          maxLoc
                maxSel
                        maxG
                              maxTime
                                     maxSource
                 0.60 NA
## Model1 5122070
                                 NA
## Model2
        7263580
                0.09 1e+00
                                  5
                                           NA
## Model3
        5122070
                 0.60 6e-07
                                 50
                                            5
                                            7
## Model4 4665911 0.30 6e-07
                                  0
## Model5 4665911 0.60 6e-07
                                 25
##
## $p4LG4_8300000_2_8
##
##
##
          maxLoc maxSel maxG maxTime maxSource
## -----
                -----
                   0.6 NA
## Model1 5122070
                                 NA
## Model2 7263580
                  0.1 1e+00
                                  5
                                           NΑ
## Model3 5122070
                  0.6 1e-07
                                  50
                                            5
        4665911 0.3 1e-07
## Model4
                                  0
                                             7
## Model5 4665911
                   0.6 1e-07
                                 25
                                             3
## $p4LG4_8300_2_7
##
##
##
                maxSel maxG maxTime maxSource
          {\tt maxLoc}
## -----
## Model1 5641459
                    0.6
                             NA
                                     NA
                                              NA
## Model2 5846714
                  0.6 1.00e-05
                                    5
```

names(ne.mcles)<-patterns</pre>

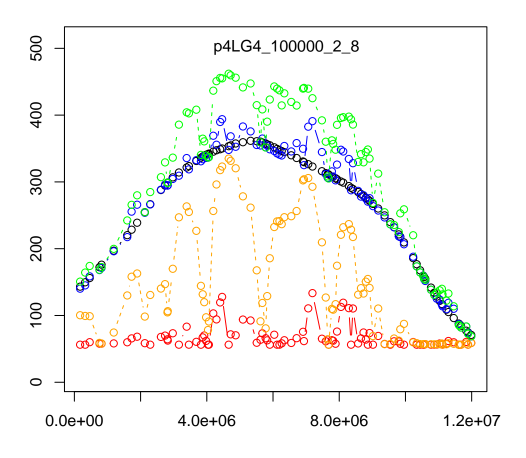
	Model3 Model4	5641459 5846714		1.00e-0		.00	3 5
		5641459		1.00e-0		25	5
##	Modelo	3041439	0.0	1.00e-0	) <del>4</del>	20	5
	\$n/11 C/1 S	33000_2_7					
##	ФРагаа-С	33000_2_1					
##							
##		maxLoc	maxSel	maxG	maxTime	maxSour	ce
		5641459	0.6	NA	NA	1	NA
		8097311		1e-05	5		NA
		5641459		6e-06			3
		4291898		6e-06	0		7
		5641459		6e-06	50		5
##							
##	\$p4LG4_8	330000_2_7					
##	-						
##							
##		maxLoc	maxSel	${\tt maxG}$	maxTime	maxSour	ce
##							
		5641459	0.6		NA	I	NΑ
		8097311		1e+00		I	NΑ
		5641459		6e-07	100		3
		4291898		6e-07	0		7
	Model5	5641459	0.6	6e-07	50		5
##							
	\$p4LG4_8	3300000_2_7					
##							
##		<b>.</b>	<b>a</b> 1	<b>a</b>	m ·	9	
##		maxLoc	maxSel	maxG	maxlime	maxSour	ce
##		E6/1/E0	0.6				 \T A
		5641459 8097311	0.6	NA 1e+00	NA 5		NA NA
		5641459		1e-07		1	3
		4291898		1e-07	0		7
	Model5			1e-07	50		5
##	1104010	0011100	0.0	10 01			Ü
	\$p4LG4_8	3300 6 6					
##	*F						
##							
##		maxLoc	maxSel	max	κG maxTi	me maxSo	ource
##							
##	Model1	7856956	0.6	ľ	JA	NA	NA
##	Model2	8889185	0.6	1.00e-0	)5	5	NA
##	Model3	8889185	0.6	1.00e-0	)4	5	3
##	Model4	8889185	0.6	6.02e-0	)5	0	3
##	Model5	4291898	0.6	1.00e-0	)1	0	5
##							
	\$p4LG4_8	3000_6_6					
##							
##		_	<u>.</u> -			2	
##						maxSour	ce
		7056056					
##	Mode11	7856956	0.6	NA	NA	I	NA

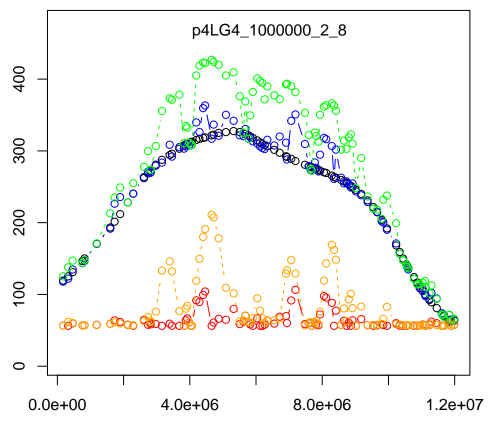
##	Model2	8889185	0.6	1e+00	3	NA	
##	Model3	4291898	0.6	1e-01	0	5	
##	Model4	8889185	0.6	6e-06	0	3	
##	Model5	4291898	0.6	1e-01	0	5	
##							
##	<b>\$p4LG4_8</b>	30000_6_6					
##							
##							
##		${\tt maxLoc}$	maxSel	maxG	maxTime	${\tt maxSource}$	
##		7054054					
	Model1	7856956	0.6	NA 4 · OO	NA	NA	
	Model2	8889185	0.6		3	NA	
	Model3	4291898	0.6		0	5	
		8889185		6e-07	0	3	
	Model5	4291898	0.6	1e-01	0	5	
##	<b>ተ</b> ъ/1፣ ሮ/ 0	200000 6 6					
##	φbaraa_o	300000_6_6					
##							
##		maxLoc	maxSel	maxG	maxTime	maxSource	
##							
##	Model1	7856956	0.6	NA	NA	NA	
##	Model2	8889185	0.6	1e+00	3	NA	
	Model3	4291898		1e-01	0	5	
##	Model4	8889185	0.6	1e-07	0	3	
	Model5	4291898	0.6	1e-01	0	5	
##	Φ 4Τ Ω4 Ο	200 5 6					
##	\$p4LG4_8	300_5_6					
##							
##		maxLoc	maxSel	ma <sup>.</sup>	xG maxTi	me maxSoui	cce
##							
	Model1	7856956	0.6	]	NA 1	NA	NA
##	Model2	8889185		1.00e-	05	5	NA
	Model3	8889185		1.00e-		5	3
##	Model4	8889185	0.6	6.02e-		0	3
##	Model5	4291898	0.6	1.00e-	01	0	5
##							
##	<b>\$p4LG4_8</b>	3000_5_6					
##							
##						_	
##		maxLoc	maxSel	maxG	maxTime	maxSource	
##	Model1	7856956	0.6	NA	NA	NA	
	Model2	8889185	0.6	1e+00	3	NA NA	
	Model3	4291898	0.5	1e-01	0	5	
	Model4	8889185	0.6	6e-06	0	3	
	Model5	4291898	0.6	1e-01	0	5	
##		1201000		10 01	•	J	
	\$p4LG4 8	30000_5_6					
##							
##							
##		maxLoc	maxSel	maxG	maxTime	maxSource	
##							

##	Model1	7856956	0.6	NA	NA	NA
##	Model2	8889185	0.6	1e+00	3	NA
##	Model3	4291898	0.5	1e-01	0	5
##	Model4	8889185	0.6	6e-07	0	3
##	Model5	4291898	0.6	1e-01	0	5
##						
##	\$p4LG4_	8300000_5_6				
##						
##						
## ##		maxLoc	maxSel	${\tt maxG}$	maxTime	maxSource
		maxLoc	maxSel	maxG	maxTime	maxSource
## ##	 Model1	maxLoc  7856956	maxSel 	maxG  NA	maxTime 	maxSource
## ## ##	Model1					
## ## ## ##		7856956	0.6	NA	NA	NA
## ## ## ##	Model2 Model3	7856956 8889185	0.6	NA 1e+00	NA 3	NA NA

## Model with chosen parameters

```
selSite = positions[seq(1, length(positions[positions<12000000]), length.out = 100)]
par(mfrow=c(2,1),oma=c(2,2,2,2),mar=c(2,2,2,2))
max.complikes1<-plot.complike("p4LG4_100000_2_8",leg=FALSE,selSite = selSite)
max.complikes2<-plot.complike("p4LG4_1000000_2_8",leg=FALSE,selSite = selSite)</pre>
```





```
mcle1<-get.mcle(max.complikes1[[1]],Ne=100000,rec=2*10^-8,selSite = selSite)
kable(mcle1)</pre>
```

	$\max Loc$	$\max Sel$	$\max G$	$\max Time$	maxSource
Model1	5322346	0.60	NA	NA	NA
Model2	7191007	0.25	1e-05	3	NA
Model3	4462715	0.60	5e-06	50	5
Model4	4661258	0.60	5e-06	0	7
Model5	4661258	0.60	5e-06	25	5

```
mcle2<-get.mcle(max.complikes2[[1]],Ne=1000000,rec=2*10^-8,selSite = selSite)
kable(mcle2)</pre>
```

	maxLoc	maxSel	maxG	maxTime	maxSource
Model1	5322346	0.60	NA	NA	NA
Model2	7191007	0.08	1e+00	5	NA
Model3	4462715	0.60	5e-07	50	3
Model4	4661258	0.25	5e-07	0	7
Model5	4661258	0.60	5e-07	25	3

The selection values are still the smallest ones. Kristin Lee indicated that this might occur if there aren't enough neutral SNPs - so I'll try re-calculating neutral SNPS

```
#calculate selected allele frequencies
LG4.sub<-calc.allFreqs(vcf[vcf$`#CHROM` == "LG4" & vcf$POS < 12*10^6,],pop.list,pop.labs)
saveRDS(LG4.sub, "dmc/selectedRegionAlleleFreqs_p4LG4_subset.RDS")
saveRDS(vcf[vcf$`#CHROM`=="LG4"& vcf$POS < 12*10^6,"POS"],"dmc/selectedRegionPositions_p4LG4_subset.RDS</pre>
#calculate neutral allele frequencies
all.out<-read.delim("all_outliers.txt",sep='\t',header=TRUE)</pre>
allFreqs<-calc.allFreqs(vcf[!vcf$^#CHROM^ %in% "LG4" & !vcf$SNP %in% all.out$SNP,],
                   pop.list,pop.labs)
saveRDS(allFreqs, "dmc/neutralAlleleFreqs_p4LG4_notSNPs.RDS")
neutralF_filename<-"dmc/neutralF_p4LG4_subset"</pre>
source("../programs/dmc-master/calcNeutralF.R")
F_estimate<-readRDS("dmc/neutralF_p4LG4_subset.RDS")
#calculate the determinant and inverse of the neutral F matrix
M <- numPops
Tmatrix \leftarrow matrix(data = rep(-1 / M, (M - 1) * M), nrow = M - 1, ncol = M)
diag(Tmatrix) = (M - 1) / M
sampleErrorMatrix = diag(1/sampleSizes, nrow = numPops, ncol = numPops)
det_FOmegas_neutral = det(Tmatrix %*% (F_estimate + sampleErrorMatrix) %*% t(Tmatrix))
saveRDS(det_FOmegas_neutral, "dmc/det_FOmegas_neutral_p4LG4_subset.RDS")
inv_FOmegas_neutral = ginv(Tmatrix %*% (F_estimate + sampleErrorMatrix) %*% t(Tmatrix))
saveRDS(inv_FOmegas_neutral, "dmc/inv_FOmegas_neutral_p4LG4_subset.RDS")
positions <- read RDS ("dmc/selected Region Positions p4LG4 subset.RDS")
F_estimate<-readRDS("dmc/neutralF_p4LG4_subset.RDS")
```

```
sampleSizes<-readRDS("dmc/sampleSizes.RDS")</pre>
neutral_det_name = "dmc/det_FOmegas_neutral_p4LG4_subset.RDS"
neutral_inv_name = "dmc/inv_F0megas_neutral_p4LG4_subset.RDS"
selSite = positions[seq(1, length(positions[positions<12000000]), length.out = 100)]</pre>
par(mfrow=c(1,1),oma=c(2,2,2,2),mar=c(2,2,2,2))
max.complikes1<-plot.complike("p4LG4_100000_2_8_sub",leg=TRUE,selSite = selSite)</pre>
                                          p4LG4_100000_2_8_sub
                                                                                            Model 1
-2300
                                                                                            Model 2
                                                                                            Model 3
                                                                                            Model 4
                                                                                            Model 5
-2400
-2600
                  2.0e+06
                                 4.0e+06
                                               6.0e+06
                                                                             1.0e+07
   0.0e+00
                                                              8.0e+06
                                                                                            1.2e+07
{\tt mcle.sub <- get.mcle(max.complikes1[[1]], Ne=100000, rec=2*10^-8,}
```

	maxLoc	maxSel	maxG	maxTime	maxSource
Model1	5322346	0.60	NA	NA	NA
Model2	7191007	0.09	1e-05	5	NA

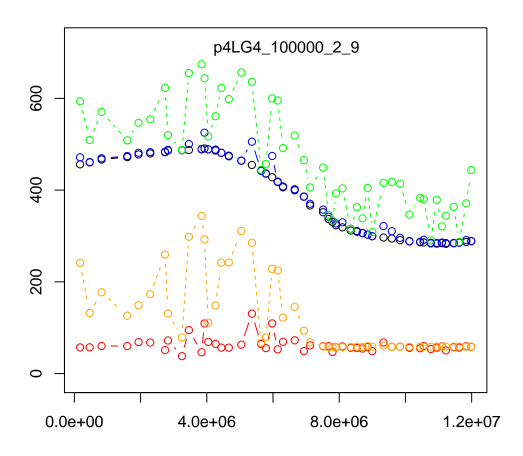
selSite=selSite)

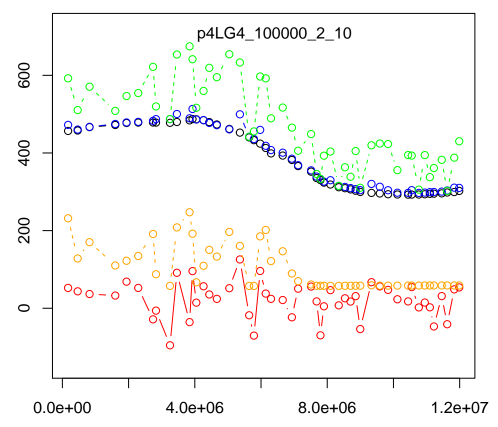
kable(mcle.sub)

	$\max Loc$	$\max Sel$	$\max G$	$\max Time$	maxSource
Model3	4462715	0.60	5e-06	50	5
Model4	4661258	0.50	5e-06	0	7
Model5	4661258	0.60	5e-06	25	5

### Let's try lower recombination rates

```
selSite = positions[seq(1, length(positions), length.out = 50)]
F_estimate<-readRDS("dmc/neutralF_p4LG4_50.RDS")
sampleSizes<-readRDS("dmc/sampleSizes.RDS")
par(mfrow=c(2,1),oma=c(2,2,2,2),mar=c(2,2,2,2))
max.complikes1<-plot.complike("p4LG4_100000_2_9",leg=FALSE,selSite = selSite)
max.complikes2<-plot.complike("p4LG4_100000_2_10",leg=FALSE,selSite = selSite)</pre>
```





# mcle1<-get.mcle(max.complikes1[[1]],Ne=100000,rec=2\*10^-9) kable(mcle1)</pre>

	$\max Loc$	$\max Sel$	$\max G$	$\max Time$	$\max Source$
Model1	3930682	0.15	NA	NA	NA
Model2	5362182	0.02	1e-05	3	NA
Model3	3930682	0.15	1e-04	500	5
Model4	3842610	0.06	5e-06	0	7
Model5	3842610	0.20	5e-06	50	3

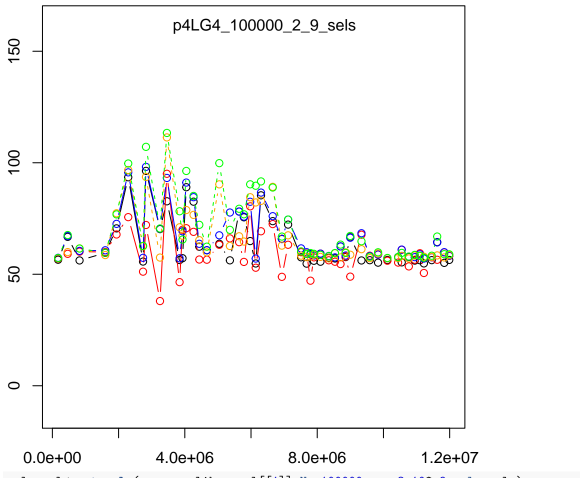
```
mcle2<-get.mcle(max.complikes2[[1]],Ne=100000,rec=2*10^-10)
kable(mcle2)</pre>
```

	maxLoc	maxSel	maxG	maxTime	maxSource
Model1	3930682	0.010	NA	NA	NA
Model2	5362182	0.001	1e-05	3	NA
Model3	3930682	0.020	5e-06	10000	5
Model4	3842610	0.010	5e-06	0	7
Model5	3842610	0.020	5e-06	500	3

#### Different selection parameters

Going below 1e-5 (i.e. to 1e-6) caused infinite values to occur

```
sels = c(9e-4,7.5e-4,5e-4, 2.5e-4,1e-4)
selSite = positions[seq(1, length(positions[positions<12000000]), length.out = 50)]
F_estimate<-readRDS("dmc/neutralF_p4LG4_50.RDS")
par(mfrow=c(1,1),oma=c(2,2,2,2),mar=c(2,2,2,2))
max.complikes.sel<-plot.complike("p4LG4_100000_2_9_sels",leg=FALSE,selSite = selSite)</pre>
```

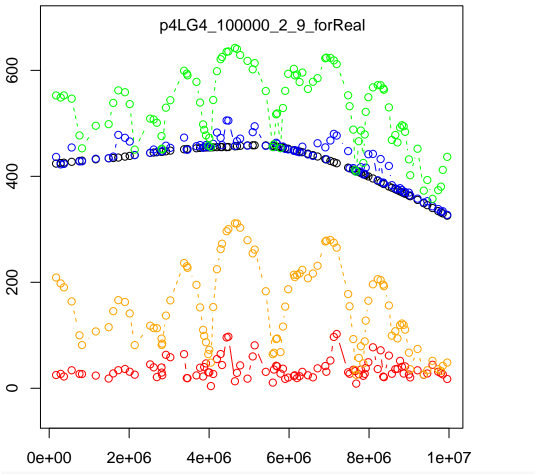


mcle.sel<-get.mcle(max.complikes.sel[[1]],Ne=100000,rec=2\*10^-9,sels=sels)
kable(mcle.sel)</pre>

	$\max Loc$	maxSel	$\max G$	$\max Time$	maxSource
Model1	2827629	0.00090	NA	NA	NA
Model2	3459260	0.00075	1e-05	7	NA
Model3	2827629	0.00090	1e-02	10000	5
Model4	3459260	0.00090	5e-06	0	7
Model5	3459260	0.00090	1e-01	100	5

#### Chosen parameter runs

Using recombination rate of 2e-9, Ne= 100000, with a wider range of selection values that go below 1e-4, and 100 selection sites from 0 to 10Mb.



mcle<-get.mcle(max.complikes[[1]],Ne=100000,rec=2\*10^-9,sels=sels,selSite=selSite)
kable(mcle)</pre>

	$\max Loc$	maxSel	$\max G$	$\max Time$	maxSource
Model1	5139485	0.15	NA	NA	NA
Model2	7191007	0.02	1e-05	3	NA
Model3	4437840	0.20	5e-06	500	5
Model4	4645842	0.06	5e-06	0	7
Model5	4645842	0.20	5e-06	50	3

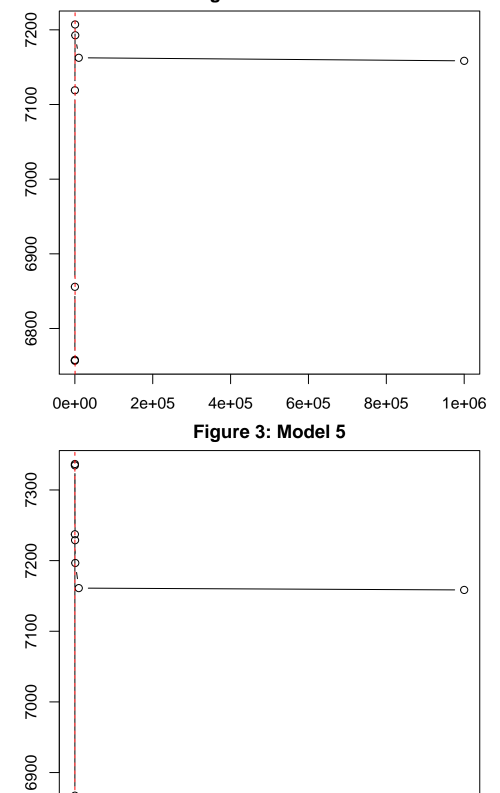
## Plot profile likelihood surfaces

Plot the profile likelihood surfaces for Models 3 and 5, where standing variation is present in all or one population, which have the highest composite log-likelihoods.

```
selSite = positions[seq(1, length(positions[positions<10000000]), length.out = 100)] \\ gs<-c(1/(2*Ne), 10^-(4:1)) \\ Ne=100000 \\ rec=2*10^-9 \\ sources <- c(3,5,7,16) \\ times = c(0, 5, 25, 50, 100, 500, 1000, 1e4, 1e6) \\ migs = c(10^-(seq(5, 1, by = -2)), 0.5, 1) \\ sels = c(9e-4,7.5e-4,5e-4, 2.5e-4,1e-4,1e-3, 0.01, 100000000]), length.out = 100)]
```

```
seq(0.02, 0.14, by = 0.01), seq(0.15, 0.3, by = 0.05))
par(mfrow=c(2,1),oma=c(2,2,2,2),mar=c(2,2,2,2))
## Model 3
compLikelihood_sv<-readRDS("dmc/compLikelihood_sv_p4LG4_100000_2_9_forReal.RDS")
names(compLikelihood sv)<-selSite</pre>
compLikelihood_sv_site = sapply(1 : length(selSite), function(i) {
  max(unlist(compLikelihood sv[[i]]))
})
mcle sv = getMCLEsv source(compLikelihood sv,selSite, sels, gs, times, sources)
mcle_sv_index<-which(mcle_sv[1]==selSite)</pre>
compLike_sv_byTime = lapply(1 : length(times), function(time) {
  sapply(1: length(sels), function(sel) {
    sapply(1 : length(gs), function(g) {
      compLikelihood_sv[[mcle_sv_index]][[sel]][[g]][[time]]
    })
 })
})
profileLike_time_sv = sapply(1: length(compLike_sv_byTime), function(i) {
  max(unlist(compLike_sv_byTime[[i]]))
})
plot(times, profileLike_time_sv, type = "b", xlab = "Time",
     ylab = "Profile composite log-likelihood", main = "Figure 2: Model 3")
abline(v = mcle sv[4], lty = 2, col = "red")
## Model 5
compLikelihood_mixed_svInd<-readRDS("dmc/compLikelihood_mixed_svInd_p4LG4_100000_2_9_forReal.RDS")
mcle_mixed_svInd = getMCLEmixed(compLikelihood_mixed_svInd, selSite, sels,
                                gs, times, migs[1], sources)
mcle_mixed_svInd_index<-which(mcle_sv[1]==selSite)</pre>
compLike_mixed_svInd_byTime = lapply(1 : length(times), function(time) {
  sapply(1: length(sels), function(sel) {
    sapply(1 : length(gs), function(g) {
      compLikelihood_mixed_svInd[[mcle_mixed_svInd_index]][[sel]][[g]][[time]]
   })
 })
})
profileLike_time_mixed_svInd = sapply(1: length(compLike_mixed_svInd_byTime), function(i) {
  max(unlist(compLike_mixed_svInd_byTime[[i]]))
})
plot(times, profileLike_time_mixed_svInd, type = "b", xlab = "Time",
     ylab = "Profile composite log-likelihood", main = "Figure 3: Model 5")
abline(v = mcle_mixed_svInd[4], lty = 2, col = "red")
```

Figure 2: Model 3



8e+05

1e+06

6e+05

0e+00

2e+05

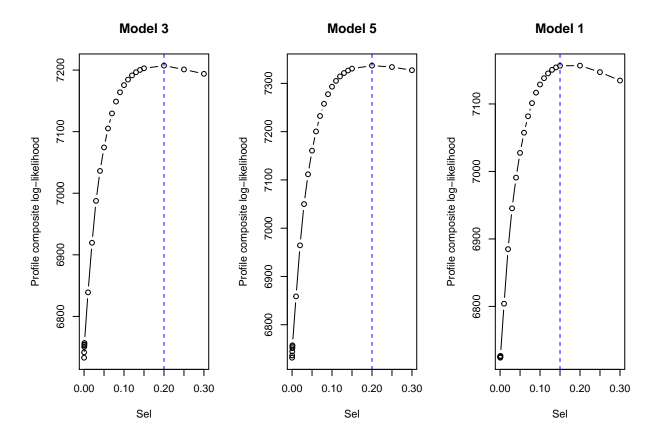
4e+05

We see little difference in the composite log-likelihoods of models 3 and 5, provides further evidence that independent mutations (or selection very old standing variation) in FLFW generated the patterns observed in the data. The point estimates of MCLE of t are marked by red lines in Figures 2 and 3.

#### MCLE for the strength of selection

The best fitting models are 5 (green), 3 (blue), and 1 (black). Models 3 and 5 estimate selection at 0.2 and model 1 estimates it at 0.15.

```
## Model 1
compLikelihood_ind<-readRDS("dmc/compLikelihood_ind_p4LG4_100000_2_9_forReal.RDS")
mcle_ind = getMCLEind(compLikelihood_ind, selSite, sels)
mcle_ind_index<-which(mcle_sv[1]==selSite)</pre>
profileLike_sel_ind = sapply(1: length(sels), function(i) {
  max(unlist(compLikelihood_ind[[mcle_ind_index]][[i]]))
})
## Model 3
profileLike_sel_sv = sapply(1: length(sels), function(i) {
 max(unlist(compLikelihood_sv[[mcle_sv_index]][[i]]))
})
## Model 5
profileLike_sel_mixed_svInd = sapply(1: length(sels), function(i) {
  max(unlist(compLikelihood_mixed_svInd[[mcle_mixed_svInd_index]][[i]]))
})
par(mfrow = c(1, 3))
plot(sels, profileLike_sel_sv, type = "b", xlab = "Sel",
     ylab = "Profile composite log-likelihood", main = "Model 3")
abline(v = mcle_sv[2], lty = 2, col = "blue")
plot(sels, profileLike_sel_mixed_svInd, type = "b", xlab = "Sel",
     ylab = "Profile composite log-likelihood", main = "Model 5")
abline(v = mcle_mixed_svInd[2], lty = 2, col = "blue")
plot(sels, profileLike_sel_ind, type = "b", xlab = "Sel",
     ylab = "Profile composite log-likelihood", main = "Model 1")
abline(v = mcle_ind[2], lty = 2, col = "blue")
```



#### Where is the selected site?

The selected site is somewhere between  $4.43784 \times 10^6$  and  $5.139485 \times 10^6$ , if we take the maximum estimates from models 1, 3, and 5.

```
par(mfrow=c(1,1),oma=c(2,2,2,2),mar=c(2,2,2,2))
max.complikes<-plot.complike("p4LG4_100000_2_9_forReal",leg=FALSE,selSite = selSite)
abline(v=mcle_ind[1],lty=2,lwd=2,col="black")
abline(v=mcle_sv[1],lty=2,lwd=2,col="blue")
abline(v=mcle_mixed_svInd[1],lty=2,lwd=2,col="green")</pre>
```

