Supplementary Material: Code Documentation

The discrepancy between the high population size and the low genetic diversity in European grey seals

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This document illustrates the data processing and analysis pipeline used in my thesis project. For further information visit the main document. The headline numbering here matches that in the main document.

This document is highly inspired by Olkkonen & Löytynoja (2023) and by the material of the course EEB-210 at University of Helsinki offered by Dr. Ari Löytynoja (A.L.) in 10-12/2024.

All data processing and analysis work was performed on *puhti* from CSC - IT Center of Science, Finland. Large data was stored on *Allas* from CSC.

2. Materials and Methods

2.1. Data

The data consisted of 140 whole-genome sequences from two seal species, grey seals (*Halichoerus grypus*) and ringed seals (*Pusa hispida*), as well as Saimaa ringed seals (*Pusa saimensis*), originating from two different data sources (CPH and HEL).

2.2 Data processing

Set up

In this document, all the steps are performed in the same working directory (\$work). For simplicity this documentation assumes that all tools are available in the bin directory. In reality many of the tools used were available on *puhti*, which were loaded with module load tool. The rest of the tools were made available by A.L.

```
work=$(pwd)
tools=$work/bin
bwa=$tools/bwa
samtools=$tools/samtools
gatk3=$tools/gatk3
gatk=$tools/gatk
bcftools=$tools/bcftools
```

```
rmask=$tools/RepeatMasker
seqbility=$tools/seqbility
bedtools=$tools/bedtools
plink=$tools/plink
vcftools=$tools/vcftools
miniprot=$tools/miniprot
busco=$tools/busco
beagle=$tools/beagle
bedextract=$tools/bedextract
genmulti=$tools/generate_multihetsep.py
msmc2=$tools/msmc2
combinecc=$tools/combineCrossCoal.py
msmc_im=$tools/MSMC_IM.py
```

Note: Seqbility contains splitfa, gen_raw_mask.pl, gen_mask, and makeMappabilityMask.py.

Preparation of the reference genome

The ribbon seal reference genome was available in FASTA format. The reference genome was unzipped and indexed.

```
ref=$work/reference/ribbon
gunzip $ref.fa.gz
$samtools faidx $ref.fa
$samtools dict $ref.fa > ref.dict
$bwa index $ref.fa
```

Read mapping

Here and in the following steps (Realignment around indels and Variant calling) the pipeline is illustrated for one sample (sample-1); in my thesis these steps were performed for all 140 grey and ringed seal samples.

The grey and ringed seal FASTQ reads were mapped to the reference genome resulting in BAM files. fastq and bams were the directories, where the reads and mapped files were stored, respectively. Mapping required defining a "readgroup" (rgroup) (Further information: https://gatk.broadinstitute.org/hc/en-us/articles/360035890671-Read-groups).

Realignment around indels

A list of regions to realign was created and the BAM files were realigned around the target. The duplicates were marked and the resulting file was indexed.

```
$gatk3 -T RealignerTargetCreator \
   -R ${ref} -I $bams/${smp}_bwa.bam -o $bams/$smp.intervals
$gatk3 -T IndelRealigner \
   -R ${ref} -I $bams/${smp}_bwa.bam -targetIntervals $bams/$smp.intervals \
   -o $bams/${smp}_real.bam
$samtools markdup $bams/${smp}_real.bam $bams/${smp}_mdup.bam
$samtools index $bams/${smp}_mdup.bam
```

Variant calling

Variants were called and the GVCF files were stored in the gvcfs directory.

```
mkdir $work/gvcfs
gvcfs=$work/gvcfs
gatk HaplotypeCaller \
    -R $ref.fa -I $bams/${smp}_mdup.bam \
    -0 $gvcfs/$smp.gvcf.gz -ERC GVCF
```

Genotype calling

Assuming the last three steps were performed for all samples, there should now be 140 *GVCF* files in the <code>gvcfs</code> directory. For simplicity, the next step is illustrated with only 8 samples, i.e. two HEL grey seals <code>HEL_grey_1.gvcf.gz</code> & <code>HEL_grey_2.gvcf.gz</code>, two CPH grey seals <code>CPH_grey_1.gvcf.gz</code> & <code>CPH_grey_2.gvcf.gz</code>, two HEL ringed seals <code>HEL_ringed_1.gvcf.gz</code> & <code>HEL_ringed_2.gvcf.gz</code>, two CPH ringed seals <code>CPH_ringed_2.gvcf.gz</code>.

In reality, **genotype calling**, **joint calling**, and **SNP filtering** was done in 19 batches, for which the reference genotype was split into 19 parts with *AWK*. These batches were then called separately by specifying the respective reference genome batch with the -0 flag. The batches were then concatenated using bcftools concat from samtools.

For simplicity, this documentation illustrates the processing of the entire genome (rather than the batches). The *GVCF* files were combined in three groups resulting in three combined *GVCF* files in the gvcf directory: (i) all grey seal samples (HEL and CPH) grey.gvcf.gz, (ii) HEL ringed seals ringed_HEL.gvcf.gz, (iii) CPH ringed seals ringed_CPH.gvcf.gz.

```
mkdir $work/gvcf

# all grey seals
smp_HEL_grey_1=$gvcfs/HEL_grey_1.gvcf.gz
smp_HEL_grey_2=$gvcfs/HEL_grey_2.gvcf.gz
smp_CPH_grey_1=$gvcfs/CPH_grey_1.gvcf.gz
smp_CPH_grey_2=$gvcfs/CPH_grey_2.gvcf.gz
$mp_CPH_grey_2=$gvcfs/CPH_grey_2.gvcf.gz
$gatk CombineGVCFs \
    -R $ref.fa \
    -V $smp_HEL_grey_1 \
    -V $smp_HEL_grey_2 \
```

```
-V $smp_CPH_grey_1 \
 -V $smp_CPH_grey_2 \
-0 $gvcf/grey.gvcf.gz
# HEL ringed seals
smp_HEL_ringed_1=$gvcfs/HEL_ringed_1.gvcf.gz
smp_HEL_ringed_2=$gvcfs/HEL_ringed_2.gvcf.gz
$gatk CombineGVCFs \
-R $ref.fa \
-V $smp_HEL_ringed_1 \
-V $smp_HEL_ringed_2 \
-0 $gvcf/ringed_HEL.gvcf.gz
# CPH ringed seals
smp_CPH_ringed_1=$gvcfs/CPH_ringed_1.gvcf.gz
smp_CPH_ringed_2=$gvcfs/CPH_ringed_2.gvcf.gz
$gatk CombineGVCFs \
-R $ref.fa \
-V $smp_CPH_ringed_1 \
-V $smp_CPH_ringed_2 \
-0 $gvcf/ringed_CPH.gvcf.gz
```

Joint calling

As a next step each of the three combined GVCF files was joint called resulting in three VCF files stored in the vcf directory.

```
mkdir $work/vcf
vcf=$work/vcf
# all grey seals
$gatk --java-options "-Xmx4g" GenotypeGVCFs \
-R $ref.fa \
-V $gvcf/grey.gvcf.gz \
-0 $vcf/grey.vcf.gz
# HEL ringed seals
gatk --java-options "-Xmx4g" GenotypeGVCFs \setminus
-R $ref.fa \
-V $gvcf/ringed_HEL.gvcf.gz \
-0 $vcf/ringed_HEL.vcf.gz
# CPH ringed seals
$gatk --java-options "-Xmx4g" GenotypeGVCFs \
 -R $ref.fa \
-V $gvcf/ringed_CPH.gvcf.gz \
-0 $vcf/ringed_CPH.vcf.gz
```

SNP filtering

The VCF files were then filtered to remove all non-binary and fixed SNPs.

```
$bcftools view -v snps -m2 -M2 -q0.001:minor $vcf/grey.vcf.gz > $vcf/grey_snp.vcf.gz
$bcftools view -v snps -m2 -M2 -q0.001:minor $vcf/ringed_HEL.vcf.gz > $vcf/ringed_HEL_snp.vcf.gz
$bcftools view -v snps -m2 -M2 -q0.001:minor $vcf/ringed_CPH.vcf.gz > $vcf/ringed_CPH_snp.vcf.gz
```

Joint masking

The repetitive elements in the reference genome were identified using RepeatMasker and dog repeat libraries resulting in the *GFF* file \$ref.fa.out.gff

```
mkdir $work/RM
RM=$work/RM
ncpu=40
$rmask -xsmall -gff --engine rmblast -dir $RM -pa $ncpu -species "Canis familiaris" $ref.fa
```

To create the positive mask, the reference genome was split into 75 bp fragments with splitfa and realign back with bwa to the reference genome. Based on that a FASTA sequence was created with gen_raw_mask.pl and gen_mask. The FASTA sequence was then converted into BED format using makeMappabilityMask.py.

```
mkdir $work/maskdir
maskdir=$work/maskdir
cd $maskdir
$seqbility/splitfa $refs.fa 75 | split -1 20000000
for i in (ls x??);
  do $bwa aln -R 1000000 -0 3 -E 3 $refs.fa $i > $i.sai
done
for i in (ls x??);
 do $bwa samse $refs.fa $i.sai $i > $i.sam
done
cat x??.sam | $seqbility/gen_raw_mask.pl > rawMask.fa
$seqbility/gen_mask -1 35 -r 0.5 rawMask.fa > mask.fa
python $seqbility/makeMappabilityMask.py
out=$maskdir/posmask.bed.gz
for i in $(ls $maskdir/posmask);
  do zcat $maskdir/posmask/$i | bgzip -c >> $out
done
```

Repeat mask and positive mask were then combined into a joint mask, excluding the repeat regions and including the positive mask.

```
$bedtools subtract -a $maskdir/posmask.bed -b $RM/$ref.fa.out.gff > $maskdir/joint_mask.bed
```

Then the grey and ringed seal *VCF* files were filtered based on the joint mask.

```
mask=$maskdir/joint_mask.bed
$bcftools view -T $mask $vcf/grey_snp.vcf.gz -Oz -o $vcf/grey_snp_mask.vcf.gz
$bcftools view -T $mask $vcf/ringed_HEL_snp.vcf.gz -Oz -o $vcf/ringed_HEL_snp_mask.vcf.gz
$bcftools view -T $mask $vcf/ringed_CPH_snp.vcf.gz -Oz -o $vcf/ringed_CPH_snp_mask.vcf.gz
```

The length of the whole genome, and the joint mask was determined with AWK, using the reference genome index and the mask BED file, respectively.

```
awk '{s+=$2}END{print s}' $ref.fa.fai > ${ref}_length.txt
awk '{s+=$3-$2}END{print s}' $mask > $maskdir/joint_mask_length.txt
```

Merge CPH and HEL ringed seals

Next, the HEL and CPH ringed seal data was merged.

```
$bcftools merge $vcf/ringed_HEL_snp_mask.vcf.gz $vcf/ringed_CPH_snp_mask.vcf.gz \
-0z -o $vcf/ringed_snp_mask.vcf.gz
```

Sequencing depth

The sequencing depth of the first ten million sites was estimated with samtools depth for all grey seals and HEL ringed seals, and with bcftools query for CPH ringed seals. The samtools approach was done for all samples individually and is here illustrated for one sample sample-1. The bcftools is performed a VCF file and outputs the estimated sequencing depth for each samples.

2.3. Population Structure

PCA and IBS distance analysis were performed with plink. PCA was performed for all grey seals and all ringed seals, as well as for CPH Atlantic and CPH Baltic grey seals. The output of the PCA is an eigenvec and an eigenval file. IBS distances were derived for CPH grey seals and HEL ringed seals, outputting a mdist.id and a mdist.gz file. The respective output files were visualized in R. The respective subgroups were filtered from the VCF files using bcftools view and referring to the metadata.tsv, that holds the sample name in the first row.

Note: PCA and IBS can also be run in one go by adding both the --pca and the --distance 1-ibs square gz flag.

2.4. Nucleotide diversity (π) and heterozygosity (H)

Pi and H were calculated with vcftools --site-pi. For H, this was done for each sample individually. For π , this was done for a group of samples.

 π

The π calculation is illustrated for CPH Atlantic grey seals, the π calculations for other study units were done in the same way.

\mathbf{H}

This illustrates the H calculation for grey seals, the calculation for ringed seals was done in the same way.

```
for file in $samples; do
    numnan=$(cat $file | tail -n+2 | grep nan | wc -1)
    corlength=$(( $lenth-$numnan ))
    cat $file | tail -n+2 | grep -v nan | awk '{s+=$3}END{print s/"'"$corlength"'"}' > z$file.txt
done

# combine H estimates of all individuals
awk 'FNR==1 {print FILENAME "\t" $0}' zH_grey_* | sort | sed 's/zH_grey_//' \
    | sed 's/.sites.pi.txt//' > H_grey.tsv
```

The averaged H per study unit was calculated in R.

2.5. Nucleotide diversity (π) and heterozygosity (H) in genomic windows

For that the .sites.pi files from 2.4. were used. The π and H in windows were calculated in R. The script was heavily inspired by Emmi Olkkonen's script. This documentation refers to a source script window_function.R found in the same GitHub repository. Make sure to specify the correct joint mask and reference index in the source script.

This documentation illustrates the calculations for CPH Atlantic grey seals.

```
setwd("work") # set the working directory accordingly
source("window function.R") # qet the functions "windows" and "bins.plot"
# pi
# read in data
Pi_grey_CPH_Atl <- as.data.frame(read.table("Pi_H/Pi_grey_CPH_Atl.sites.pi", sep = "\t", header = T))
Pi_grey_CPH_Atl_windows <- windows(Pi_grey_CPH_Atl) # calculate windows
rm(Pi_grey_CPH_Atl) # delete to save memory
Pi_grey_CPH_Atl_plot <- bins.plot(Pi_grey_CPH_Atl_windows) # counts the windows in 0.0001 pi bins
bin \leftarrow c(seq(0,0.00519,0.0001),0.025)
Pi <- as.data.frame(cbind(bin, Pi_grey_CPH_Atl_plot)) # add other study units here if needed
colnames(Pi) <- c("bins", "Pi_grey_CPH_Atl")</pre>
Pi.long <- gather(Pi, study units, count, -bins) # this can then be plotted with x=bins and y=count
# H
# select the correct samples here
H_grey_CPH_Atl_files <- c("H_grey_smp1.sites.pi", "H_grey_smp2.sites.pi", "H_grey_smp3.sites.pi")</pre>
H grey CPH Atl windows <- list()</pre>
a <- 1
for (i in H_grey_CPH_Atl_files) {
  filename <- paste0("Pi_H/", i)</pre>
  data <- as.data.frame(read.table(filename, sep = "\t", header = T)) # read in data</pre>
  H_grey_CPH_Atl_windows[[a]] <- windows(data)</pre>
                                                                          # calculate windows
  rm(data)
                                                                          # delete to save memory
  print(i)
  a <- a+1
H_grey_CPH_Atl_names1 <- gsub("^H_grey_", "", H_grey_CPH_Atl_files)</pre>
H grey CPH Atl names <- gsub("^.sites.pi", "", H grey CPH Atl names1)
H_grey_CPH_Atl_bins <- bin</pre>
```

```
for (a in 1:length(H_grey_CPH_Atl_windows)) {
    # counts the windows in 0.0001 pi bins
    temp.bins <- bins.plot(H_grey_CPH_Atl_windows[[a]])
    H_grey_CPH_Atl_bins <- cbind(H_grey_CPH_Atl_bins, temp.bins)
}
colnames(H_grey_CPH_Atl_bins) <- c("bins", H_grey_CPH_Atl_names)
# add other study units here if needed
H <- as.data.frame(cbind(H_grey_CPH_Atl_bins, study_unit[,-1]))
H.long <- gather(H, smp, count, -bins) # this can then be plotted with x=bins and y=count</pre>
```

2.6. Nucleotide diversity (π) in genes

CDS

To be able to determine coding site (CDS) regions in grey and ringed seals, the reference genome was annotated. This was done with the northern elephant seal (NES) protein sequence using miniprot.

First, get the NES protein sequence from NCBI, and rename it.

```
mkdir $work/annotation
annotation=$work/annotation

cd $annotation
wget https://ftp.ncbi.nlm.nih.gov/genomes/all/annotation_releases/9716/GCF_029215605.1-RS_2024_04 \
/GCF_029215605.1_mMirAng1.0.hap1_protein.faa.gz
mv GCF_029215605.1_mMirAng1.0.hap1_protein.faa.gz elephant_seal_protein.faa.gz
cd ..
```

Second, annotate the reference genome, select only CDS regions and make a TAB and a BED file.

BUSCO genes

Find carnivore BUSCO genes in the reference genome with busco and take the intersect between the CDS regions and the BUSCO regions.

immune genes

Immune genes from the ferret were identified by Dr. Mia Valtonen and Dr. Outi Hallikas from the Ensemble database as \$annotation/ferret_immune.txt. To be able to use those, the reference genome was annotated with the ferret protein sequence from Ensemble.

First, get the NES protein sequence from NCBI, and rename it.

```
cd $annotation
wget https://ftp.ensembl.org/pub/release-113/fasta/ \
   mustela_putorius_furo/pep/Mustela_putorius_furo.MusPutFur1.0.pep.all.fa.gz
mv Mustela_putorius_furo.MusPutFur1.0.pep.all.fa.gz ferret_protein.fa.gz
cd ..
```

Second, annotate the reference genome.

```
protein=$annotation/ferret_protein.fa.gz
$miniprot -t8 --gff $ref.fa.gz $protein > $annotation/annotation_ferret.gff
```

Third, find the immune regions in the ferret annotation.

Forth, overlap the ferret immune genes with the NES CDS regions.

filter my data

Filter the VCF files based on my annotations. This is illustrated for grey seal, and was done for ringed seals accordingly.

```
vcf_grey=$vcf/grey_snp_mask.vcf.gz

CDS=$annotation/elephant_seal_CDS.bed
CDS_BUSCO=$annotatio/elephant_seal_BUSCO.bed
CDS_immune=$annotation/elephant_seal_immune.bed

vcf_CDS_grey=$annotation/grey_snp_mask_CDS.vcf.gz
vcf_CDS_BUSCO_grey=$annotatio/grey_snp_mask_CDS_BUSCO.vcf.gz
vcf_CDS_immune_grey=$annotatio/grey_snp_mask_CDS_immune.vcf.gz

$bcftools view -T $CDS $vcf_grey -Oz -o $vcf_CDS_grey
$bcftools view -T $CDS_BUSCO $vcf_grey -Oz -o $vcf_CDS_BUSCO_grey
$bcftools view -T $CDS_BUSCO $vcf_grey -Oz -o $vcf_CDS_busco_grey
$bcftools view -T $CDS_immune $vcf_grey -Oz -o $vcf_CDS_immune_grey
```

Then, π was calculated for different study units for the three categories (CDS, BUSCO, and immune), as illustrated in 2.4. The relative π (i.e. π in CDS divided by π in the full genome) was done in R.

2.7. Demographic and isolation-migration inference

For this analysis only CPH data was used.

imputation and phasing

MSMC2 requires phased data, imputation was done to improve data quality. This was done for CPH Atlantic grey seals, excluding Russia and Norway, CPH Baltic grey seals and CPH Arctic ringed seals, separately. To achieve those sets the *VCF* files were subsetted with bcftools view, as illustrated before leading to the three files: grey_snp_mask_CPH_Atl.vcf.gz, grey_snp_mask_CPH_Bal.vcf.gz, ringed_snp_mask_CPH.vcf.gz.

beagle is run twice, once to impute and once to phase. This is illustrated for CPH Atlantic grey seals only, and was done accordingly with the other two sets.

```
mkdir $work/phasing
phasing=$work/phasing

$beagle_tool="java -Xmx145G -jar $beagle"
$beagle nthreads=5 ne=100000 gl=$vcf/grey_snp_mask_CPH_Atl.vcf.gz out=$phasing/grey_Atl_imp
$beagle nthreads=5 ne=100000 gt=$phasing/grey_Atl_imp.vcf.gz out=$phasing/grey_Atl_phased
```

Baltic and Atlantic grey seal data were then merged with bcftools merge.

```
$bcftools index $phasing/grey_Atl_phased.vcf.gz
$bcftools index $phasing/grey_Bal_phased.vcf.gz
$bcftools merge $phasing/grey_Atl_phased.vcf.gz $phasing/grey_Bal_phased.vcf.gz -0z -o \
$phasing/grey_phased.vcf.gz
$bcftools index $phasing/grey_phased.vcf.gz
```

This resulted in the two VCF files \$phasing/grey_phased.vcf.gz and \$phasing/ringed_phased.vcf.gz.

create multiHetSep files

I chose 6 samples, based on sequencing depth, from each study unit to include in the MSMC2 analysis. For the estimation of the sequencing depth see 2.2.

First, the joint mask was split into contigs.

```
mkdir $maskdir/contigs
contigs=$maskdir/contigs

for chr in {001..499}; do
   CHR=Hfa$chr
   $bedextract $CHR $mask > $contigs/$CHR.mask.bed
   gzip $contigs/$CHR.mask.bed
done
```

Then multiHetSep files for grey and ringed seals were created. This step is illustrated from grey seals only.

```
mkdir $work/msmc
msmc=$work/msmc
mkdir $msmc/ctgs_grey
mkdir $msmc/multi_grey
module load python-data # needed to run the $genmulti python script
SAMPLES="smp1 smp2 smp3 smp4 smp5 smp6" # add all 12 (2*6) samples here
VCF=$phasing/grey_phased.vcf.gz
FAI=$ref.fa.fai
for chr in {001..499}; do
CHR=Hfa$chr
len=`grep -w $CHR $FAI | cut -f2`
 if [ $len -gt 100000 ]; then
 list=""
 for SMP in $SAMPLES; do
   $bcftools view -s $SMP -r $CHR $VCF -Oz -o $msmc/ctgs_grey/$SMP.$chr.vcf.gz
  list=$list" "$msmc/ctgs_grey/$SMP.$chr.vcf.gz
  $genmulti --mask $contigs/$CHR.mask.bed.gz $list \
    > $msmc/multi_grey/chr${chr}.multihetsep.txt 2> $msmc/multi_grey/log.txt
  rm $list
fi
done
```

demographic analysis

Then the MSMC2 analysis was run for Baltic and Atlantic grey seals and Arctic ringed seals, each analysis was done with 4 samples at a time. The first 6 samples in $msmc/multi_grey/chr*.multihetsep.txt$ files refer to the Baltic samples and the last 6 to the Atlantic samples. As this step is computationally quite intensive, the individual msmc2 analyses were submitted as separate jobs or in groups of 2-3. The sample names are arbitrary.

Baltic grey seals:

Atlantic grey seals:

```
samples=(T Y N W I U)
for i in $(seq 6 10); do
       for j in $(seq $(( $i + 1 )) 11); do
                for k in $(seq $(( $j + 1 )) 11); do
                        for 1 in $(seq $(( $k + 1)) 11); do
                                a=$i-6
                                b=$j-6
                                c = k - 6
                                d=$1-6
                                file=A${samples[a]}${samples[b]}${samples[d]}
                                msmc2 - t 12 - p 27*1+1*2+1*3 - I $(( $i*2 )),$(( $i*2+1 )),$(( $j*2 ))
                                  ,$(( $j*2+1 )),$(( $k*2 )),$(( $k*2+1 )),$(( $l*2 )),$(( $l*2+1 )) \
                                    -o $msmc/output_grey/$file $msmc/multi_grey/chr*.multihetsep.txt \
                                    $> $msmc/output_grey/$file.runlog
                        done
                done
        done
done
```

The same was done for ringed seals.

isolation-migration inference

The MSMC-IM analysis was performed on rCCR outputs of MSMC2 analysis. This was only done for CPH grey seals. For that MSMC2 analyses were performed with 2 instead of 4 samples.

Baltic grey seals:

Atlantic grey seals:

Then the relative Cross-Coalescence-Rate (rCCR) was calculated. As this step is computationally quite intensive, the individual rCCR analyses were submitted as separate jobs or in groups of 2-3.

Then the respective MSMC2 and rCCR analysis were combined, so that the samples match.

Based on these combined files, MSMC-IM was run.

convert time boundatries and coalescence rates

The MSMC2 and MSMC-IM output data was then prosessed in R.

Demographic analysis:

This analysis is illustrated for Atlantic grey seals only.

```
setwd("work") # set the working directory accordingly
library(dplyr) # load library
# set mutation rate and generation time
mu=1.826e-8
gt = 10
# read in the data
grey_atl <- c("ANWIU", "ATNIU", "ATNWU", "ATNWU", "ATWIU", "ATYIU", "ATYIU", "ATYNI", \</pre>
                "ATYNU", "ATYNW", "ATYWI", "ATYWU", "AYNIU", "AYNWI", "AYNWU", "AYWIU")
dat_grey_atl <- c()</pre>
for(p in grey_atl) {
 tmp <- read.table(paste0("msmc/output_grey/",p,".final.txt"),head=T)</pre>
 dat_grey_atl <- rbind.data.frame(dat_grey_atl, cbind.data.frame(pop=p,tmp))</pre>
# convert time boundaries
dat_grey_atl <- dat_grey_atl %% mutate(time=left_time_boundary/mu*gt,ne=(1/lambda)/(2*mu))
# combine with other study units
dat.msmc <- rbind(dat_grey_atl, dat_grey_bal, dat_ring_arc) # this can then be used for plotting
```

Isolation-Migration analysis:

```
# read in the data
dat.im <- c()
for(file in dir("msmc/output_grey/","*MSMC_IM.estimates.txt")){
   tmp <- read.table(paste0("msmc/output_grey/",file),head=T)
   pair <- substr(file,1,13)
   tmp$pair <- pair
   tmp$right_time_boundary = c(tmp$left_time_boundary[2:length(tmp$left_time_boundary)],Inf)
   dat.im <- rbind.data.frame(dat.im,tmp)
}

# cut 'm' at 0.999
dat.im$m_prime <- if_else(dat.im$M <= 0.999, dat.im$m, 1e-30)

# dat.im can then be used for plotting</pre>
```

References

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