# Pipeline for "Recombination suppression and selection affect local ancestries in genomes of a migratory songbird"

#### Local PCA

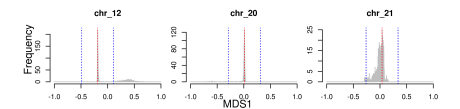
```
Here I show how local PCA was run with three example chromosomes, 12, 20,
and 21, which are listed in local_PCA/list/chromosomes/list
dirbase=$PWD/local_PCA
dirin=$dirbase/input
dirvcf=$dirin/vcf
dirtab=$dirin/table
dirlist=$dirbase/list
dirout=$dirbase/output
dirlocalpca=$dirout/local_PCA
dirlocalpcamds=$dirout/local_PCA_MDS
dirscripts=$dirbase/scripts
Download filtered VCF files for chromosomes 12, 20, 21.
Make list of sites from VCF.
while read chr
do
        bcftools query -f '%POS\n' $dirvcf/$chr.vcf.gz > $dirtab/$chr.sites.list
done <$dirlist/chromosomes.list</pre>
Make genotype table from VCF.
while read chr
        sbatch $dirscripts/vcf2table.sh $dirvcf/$chr.vcf.gz $dirtab/$chr.table
done <$dirlist/chromosomes.list</pre>
Run local PCA. local_PCA/scripts/local_PCA.sh submits local_PCA/scripts/local_PCA.R
via slurm. Check local_PCA/scripts/local_PCA.sh and local_PCA/scripts/local_PCA.R
for detail.
module load R/3.5.3
while read chr
do
        sbatch $dirscripts/local_PCA.sh $dirtab $dirtab $dirlocalpca $dirlocalpcamds $dirli;
```

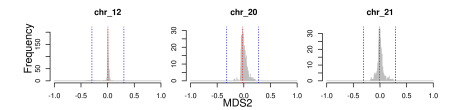
```
done <$dirlist/chromosomes.list</pre>
```

Concatenate the output for the three chromosomes.

Get threshold of MDS values and plot MDS distribution.

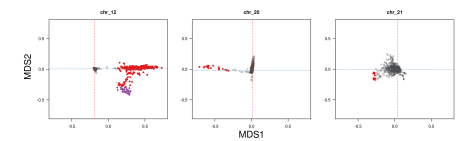
Rscript \$dirscripts/plot\_local\_PCA\_MDS\_distribution.R --input \$dirlocalpcamds/local\_PCA\_MDS\_





Plot MDS1 vs MDS2.

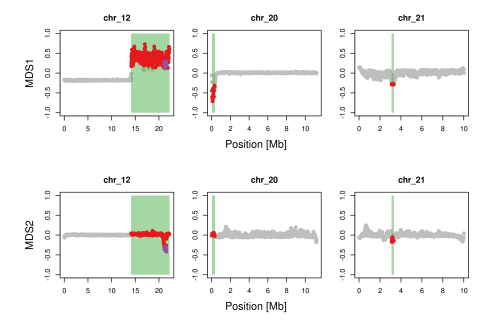
Rscript \$dirscripts/plot\_local\_PCA\_MDS\_MDS1-vs-MDS2.R --input \$dirlocalpcamds/local\_PCA\_MDS\_MDS1-vs-MDS2.R --input \$dirlocalpcamds/local\_PCA\_MDS1-vs-MDS2.R --input \$dirlocalpcamds/localpcamds/local\_PCA\_MDS1-vs-MDS2.R --input \$dirlocalpcamds/l



Get coordinates of outlier windows.

Rscript \$dirscripts/getOutliers.R --input \$dirlocalpcamds/local\_PCA\_MDS\_3\_chromosomes.txt -- Make Manhattan plots.

Rscript \$dirscripts/plot\_local\_PCA\_MDS\_manhattan.R --input \$dirlocalpcamds/local\_PCA\_MDS\_3\_o



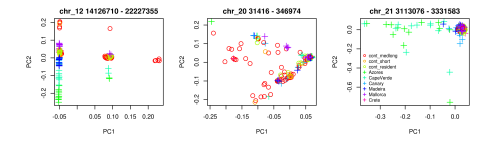
### PCA in local PCA outliers

Here I show how PCA was run for three local PCA outlier regions of chromosomes 12, 20, and 21, which represent class-1, 2, and 3 outliers.

dirbase=\$PWD/PCA
dirin=\$dirbase/input
dirvcf=\$dirin/vcf

```
dirlist=$dirbase/list
dirout=$dirbase/output
dirscripts=$dirbase/scripts
Link VCF files from local_PCA/input/vcf and index them.
ln $dirbase/../local_PCA/input/vcf/*vcf.gz $dirvcf
while read chr
        echo $chr
        bcftools index $dirvcf/$chr.vcf.gz
done<$dirlist/chromosomes.list</pre>
Copy BED file for coordinates of local PCA outliers of chromosomes 12, 20, and
21.
ln $dirbase/../local_PCA/output/local_PCA_MDS/local_PCA_MDS_outlier.bed $dirlist
Extract SNPs within the local PCA outlier regions.
while read chr from to
do
        bcftools view -O z -r $chr:$from-$to -S $dirlist/blackcap_id.list $dirvcf/$chr.vcf.
done < $dirlist/local_PCA_MDS_outlier.bed</pre>
Run PLINK for PCA.
while read chr from to
do
        sbatch $dirscripts/plink_pca.sh $dirvcf/${chr}_${from}_$to $dirout/${chr}_${from}_$
done < $dirlist/local_PCA_MDS_outlier.bed</pre>
Plot PCA results.
```

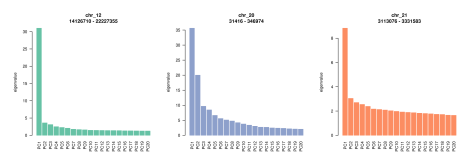
Rscript \$dirscripts/plot\_pca\_per\_outlier.R --dirpca \$dirout --poplist \$dirlist/id\_spp\_pop\_satisfies the control of the control



Get Eigenvalues.

Plot Eigenvalues.

Rscript \$dirscripts/plot\_eigenval.R --dirpca \$dirout



#### Class-1 genomic islands

```
dirbase=$PWD/class-1
dirin=$dirbase/input
dirvcf=$dirin/vcf
dirlist=$dirbase/list
dirout=$dirbase/output
dirscripts=$dirbase/scripts
```

#### Heterozygosity

Make link to VCF of chromosome 12, as an example chromosome harbouring a class-1 genomic island.

```
ln $dirbase/../PCA/input/vcf/chr_12.vcf.gz $dirvcf
ln $dirbase/../PCA/input/vcf/chr_12.vcf.gz.csi $dirvcf
```

Get coordinates of class-1 genomic island of chromosome 12.

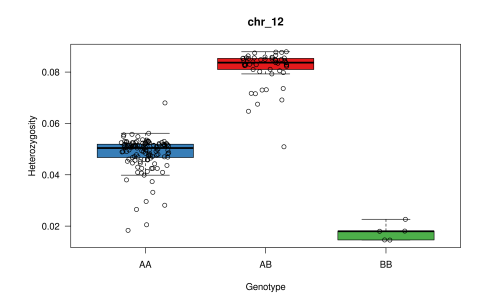
```
awk -v OFS="\t" '$1=="chr_12"{$1=$1;print $0}' $dirbase/../local_PCA/output/local_PCA_MDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_NDS/local_PCA_
```

Make list of individuals with AA, AB, and BB.

awk '\$3<0{print \$1}' \$dirbase/../PCA/output/chr\_12\_14126710\_22227355.eigenvec > \$dirlist/chrawk '\$3>0.15{print \$1}' \$dirbase/../PCA/output/chr\_12\_14126710\_22227355.eigenvec > \$dirlist/chrawk '\$1\$' \$dirbase/../PCA/output/chrawk '\$1\$' \$dirbase/../PCA/output/chrawk '\$1\$' \$dirbase/../PCA/o

Plot heterozygosity for AA, AB, and BB.

Rscript \$dirscripts/plot\_het.R --dir \$dirout



## FST, dXY and

FST, dXY

 $F_{ST}$   $\pi$