Genomic projects

Stefano Scansani

March 1, 2024

Abstract

So far this is the \LaTeX versin of the README.md.

Contents

1	Genomic projects tutorials			
	1.1	Tools	· · · · · · · · · · · · · · · · · · ·	2
	1.2	Resou	urces & Data	3
		of the working environment	3	
			Get PLINK working in Linux	
			Get TASSEL (GUI) on Linux	
		1.3.3	Get rTASSEL working in Linux	4
			Get GEMMA	
2	-		NP Profiling of worldwide goat populations	5 5
3	Methods			
		3.0.1	Data import	5
4	Results			
		4.0.1	Multidimensional scaling	5
_	_			
L	ist (of Fig	gures	
	1	1 Multidimensional Scaling of the goat dataset		
	2 Multidimensional Scaling of the goat dataset, subset of 6 goat bre			

List of Tables

1 Genomic projects tutorials

!!! warning "This repository is under construction"

This repository contains a collection of genomic projects that I am working on. GitHub repository of bioinformatic projects recolving around genomics using different tools like Plink through plinkr R package, rTASSEL and TASSEL 5 (GUI),

GEMMA for mixed models analysis in R, SAMtools from the command line to analyze BAM files, gBLUP coming soon.

The repository has been created for self-teaching purposes of biological concept and bioinformatic tools, and make use of other repositories, scripts and data sources, taken or modified as such.

1. SNP profiling of goat breeds.

- Data source: Colli et al. (2018) https://doi.org/10.1186/s12711-018-0422-x

<small>Multidimensional Scaling (MDS) Plot of a population of 4,653 Individuals from 169 Goat Breeds genotyped with 49,953 SNPs.

<small>The MDS plot visualizes genetic relationships among 4,653 individuals from 169 goat breeds. Genetic distances were computed using PLINK to generate the kinship matrix, and MDS analysis was conducted with the cmdscale function based on genotyping data from 49,953 SNPs. Each point represents a goat, and spatial arrangement reflects genetic dissimilarities. This exploratory analysis offers insights into genetic diversity, population structure, and relatedness. </small>

1. a. Manhattan plot of a GWAS on dog population for deafness. *Data source*: **Hayward et al.** (2020) https://doi.org/10.1371/journal.pone.0232900

 <small>Manhattan plots showing the genome wide association (GWA) between dog deafness and their genotype. The plot displays the genomic positions of single nucleotide polymorphisms (SNPs) across the genome on the x-axis, with the corresponding -log₁₀ transformed P-values indicating the strength of association with the trait on the y-axis. </small>

1. b. Plot of the top significant SNPs identified by the above GWAS.

and a zoom in the chromosome 3 above LOD 3.5. <img src="Figures/SNPchr3top.png"
height="350px">

1.1 Tools

- PLINK 1.90 https://www.cog-genomics.org/plink2/
- plinkr R package repository documentation. https://github.com/AJResearchGroup/ plinkr
- TASSEL 5 https://www.maizegenetics.net/tassel. Bradbury et al., (2007) TASSEL: software for association mapping of complex traits in diverse samples, Bioinformatics, Volume 23, Issue 19, Pages 26332635 https://doi.org/10.1093/bioinformatics/btm308
- rTASSEL R package repository documentation.

 rtassel.maizegenetics.net/index.html, Repository: https://github.com/maize-genetics/rTASSEL. Monier et al., (2022). rTASSEL: An R interface to TASSEL for analyzing genomic diversity. Journal of Open Source Software, 7(76), 4530, https://doi.org/10.21105/joss.04530

• GEMMA Genome-wide Efficient Mixed Model Association https://github.com/genetics-statistics/GEMMA. Xiang Zhou and Matthew Stephens (2012). Genome-wide efficient mixed-model analysis for association studies. Nature Genetics 44, 821824.

1.2 Resources & Data

- Marees et al. (2018) A tutorial on conducting genome-wide association studies: Quality control and statistical analysis. *Int J Methods Psychiatr Res.* 27:e1608. https://doi.org/10.1002/mpr.1608
- Marees et al. (2018) tutorial https://github.com/MareesAT/GWA_tutorial
- **Gábor Mészáros** (2021) Genomic Boot Camp Book https://genomicsbootcamp.github.io/book/
- Gábor Mészáros video tutorials https://www.youtube.com/c/GenomicsBootCamp
- **Colli et al.** (2018) Genome-wide SNP profiling of worldwide goat populations reveals strong partitioning of diversity and highlights post-domestication migration routes. *Genet Sel Evol* 50, 58. https://doi.org/10.1186/s12711-018-0422-x
- DATA: Colli et al. (2020). Signatures of selection and environmental adaptation across the goat genome post-domestication [Dataset]. *Dryad*. https://doi.org/10.5061/dryad.v8g21pt
- Decker et al. (2014) Worldwide Patterns of Ancestry, Divergence, and Admixture in Domesticated Cattle. PLOS Genetics 10(3): e1004254.https://doi.org/10.1371/journal.pgen.1004254¹,
- DATA: **Decker et al.** (2015) Worldwide patterns of ancestry, divergence, and admixture in domesticated cattle [Dataset]. Dryad. https://doi.org/10.5061/dryad.th092

1.3 Setup of the working environment

Install R: The Comprehensive R Archive Network (CRAN)² IDE:VSCode³/RStudio⁴^{*} Install Python: Miniconda 3⁵^{*} OS: Linux^{*}/WSL

<small>^{*}Suggested</small>

¹https://journals.plos.org/plosgenetics/article?id=10.1371/journal.pgen.

²https://cran.r-project.org/

³https://code.visualstudio.com/

⁴https://posit.co/download/

⁵https://docs.anaconda.com/free/miniconda/index.html

1.3.1 Get PLINK working in Linux

- 1. Download PLINK 1.90 Linux 64-bit6
- 2. Install PLINK cd Downloads/ sudo unzip plink_linux_x86_64_20200616.zip
 -d plink_install
- 3. PLINK in usr/local/bin

cd plink_install sudo cp plink /usr/local/bin sudo chmod 755
/usr/local/bin/plink

4. Add PLINK to PATH

with bash/zsh/...

sudo nano ~/.bashrc

adn include the line:

export PATH=/usr/local/bin:\$PATH

Save and exit. Refresh the terminal and you should be able to call plink from the terminal at any user position in the system.

source ~/.bashrc plink -help

1.3.2 Get TASSEL (GUI) on Linux

- 1. Go on the website https://www.maizegenetics.net/tassel and download the last UNIX verison.
- 2. Download the TASSEL_{xxx}_unix.sh and make it executable chmod +x ~/Downloads/TASSEL_{xxx}_unix.sh
- 3. Run the TASSEL installer ~/Downloads/TASSEL_{xxx}_unix.sh

1.3.3 Get rTASSEL working in Linux

1. rJava installation

sudo apt install default-jdk sudo R CMD javareconf R install.packages("rJava")

2. Installation in R

```
if (!require("devtools")) install.packages("devtools") devtools::install_github(
repo = "maize-genetics/rTASSEL", ref = "master", build_vignettes =
TRUE, dependencies = TRUE )
```

- 3. Run rTASSEL
- Allocate job's memory¹ and start the logger (here at the root of the project):

¹"-Xmx50g" and "-Xms50g", "50g" represents 50 Gigabytes of memory.

!! Choose an appropriate value that fits your machine !!

options(java.parameters = c("-Xmx50g", "-Xms50g")) rTASSEL::startLogger(fullPath
= NULL, fileName = NULL)

⁶https://s3.amazonaws.com/plink1-assets/plink_linux_x86_64_20231211.zip

· Run & infos

library(rTASSEL) ??rTASSEL

Useful resource for rTASSEL are the vignettes and tutorials at https://rtassel. maizegenetics.net/index.html

1.3.4 Get GEMMA

GEMMA can be installed from source at the GitHub repo, but is also available through Bioconda http://www.ddocent.com/bioconda/. To install is suggested to have miniconda installed and working, and then added the channel for Bioconda, you should already have defaults and conda-forge.

conda config -add channels defaults conda config -add channels
conda-forge conda config -add channels biocond conda install gemma
And use GEMMA with

gemma -h

2 Project: SNP Profiling of worldwide goat populations

2.1 Data

Data source: Colli et al. (2018) https://doi.org/10.1186/s12711-018-0422-x

3 Methods

3.0.1 Data import

4 Results

4.0.1 Multidimensional scaling

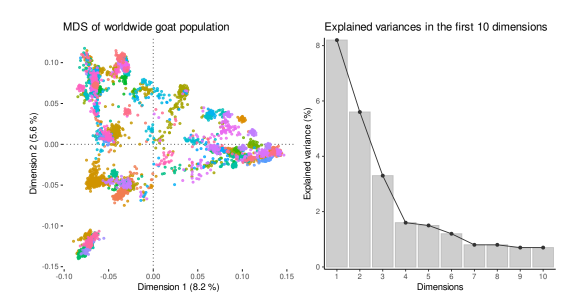


Figure 1: Multidimensional Scaling of the goat dataset

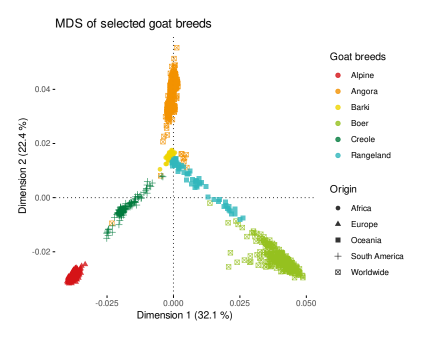


Figure 2: Multidimensional Scaling of the goat dataset, subset of 6 goat breeds.