



# The synbreed R package

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## Introduction

In many **plant and livestock breeding** programs dense genome-wide markers are being used to increase the genetic gain. The prediction of the genetic potential of individuals from their DNA sequence is called **genomic prediction**. Different statistical models and software packages have been used for this purpose. However, there is no comprehensive **software** available where all required data analysis steps including data processing and visualization have been implemented. We present a novel **R package** named synbreed which implements methods to build an **analysis pipeline** of genomic prediction data within the freely available **open source** software R [1].

An **unified data object** for genomic prediction or association analysis is created to merge phenotypic, genotypic, marker map and pedigree data. Thereby, the implementation is flexible with respect to data structure and fast with respect to data size. Within this framework, the evaluation of data from a **breeding program** of both plant and animal breeding is feasible and can be **automatized** to a large extend. The synbreed package is hosted and developed on

https://r-forge.r-project.org/projects/synbreed/

In 2011, the package synbreed will be released to CRAN together with a detailed user manual.

# Structure

The class gpData is used for data storage of all required data sources:

pheno data.frame for one or more traits (with or without replications)

matrix with genotypic data in arbitrary coding

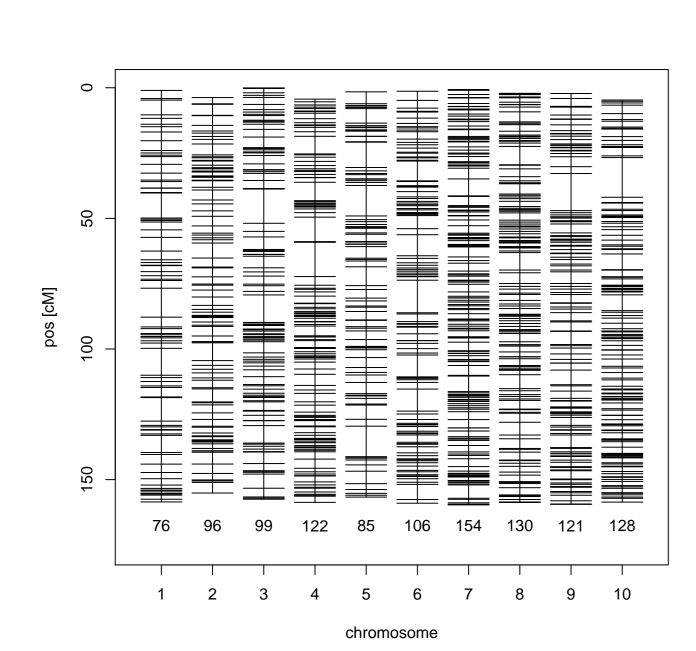
(by genotypes or by alleles)

data.frame with chromosome number and position within each chromosome (genetic or physical distance) for markers in geno

chromosome (genetic or physical distance) for markers in geno pedigree object of class pedigree with pedigree structure of individuals

covar data.frame with covariates for individuals

Unique row names/column names are used for individuals/markers to have clear **data queries** and **data merges**. Further methods of the package are implemented based on this structure.



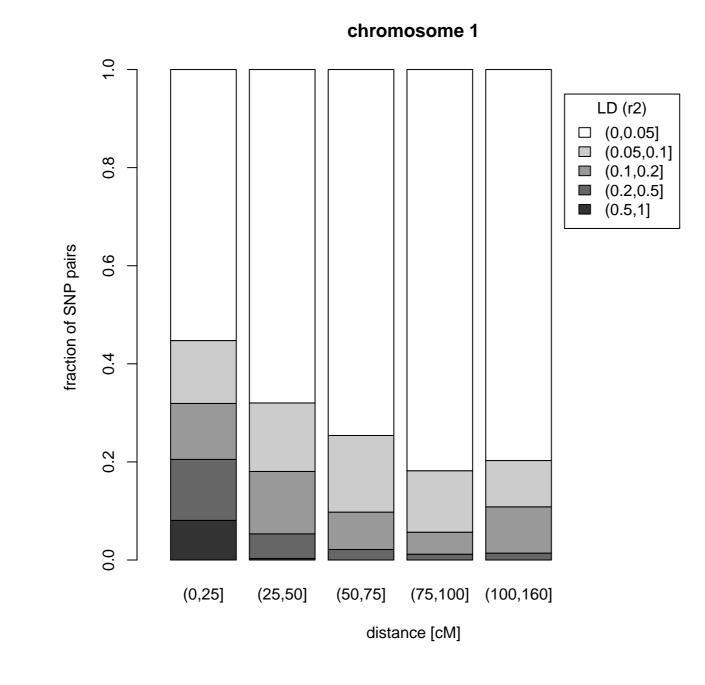
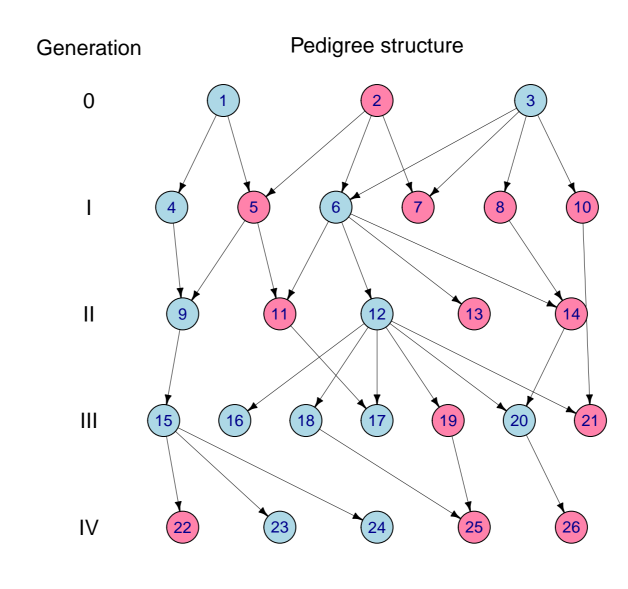


Figure 1: Simulated maize data: Marker map for data set (left) and LD decay visualization for 76 markers on the first chromosome (right).



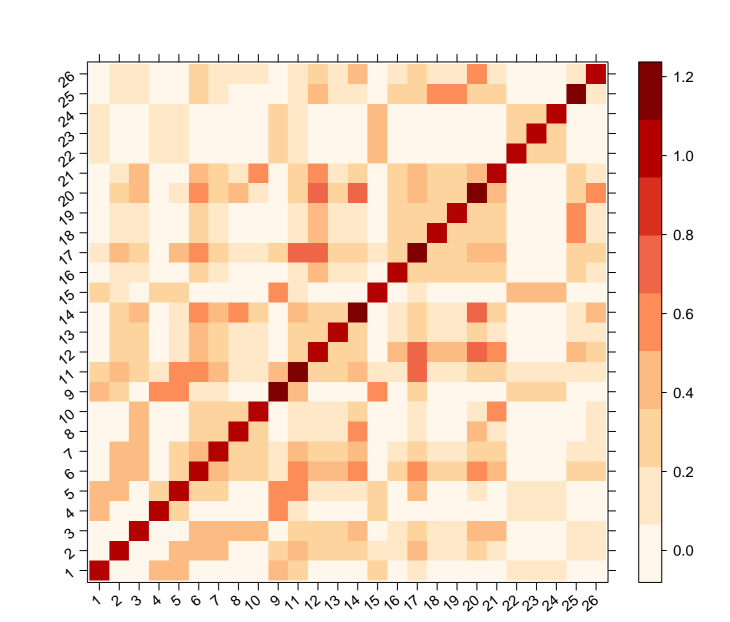


Figure 2: Pedigree structure (left) for 26 individuals (blue=sire, red=dam) and heatmap of the corresponding estimated numerator relationship matrix based on pedigree (right)

# **Functions**

#### **Data processing**

- Combining raw data sources to a gpData object
- Conversion from and to class cross in package qtl
- Coding marker data into number of copies of the minor allele
- Preselection of markers according to MAF, % missing values and LD
- Imputation of missing genotypes by fixed value, marginal allele distribution or by family structure for fully homozygous inbred individuals

## Data visualization and analysis

- Summary method for classes gpData, pedigree and relationshipMatrix
- Marker map representation for low and high density maps (Figure 1)
- -LD computation as  $r^2$  and LD decay visualization as as scatterplot or stacked histogram (Figure 1)
- Pedigree tree and kinship visualization of relatedness between individuals (Figure 2)

## Statistical models

- Estimation of pedigree based relationship (additive and dominance)
- Marker based relationship (according to vanRaden [2] or Rogers' [3] distance)
- Cross-validation for BLUP, Ridge Regression and Bayesian methods

# Data sets

maize Simulated data for a maize breeding program using 1250 doubled haploids (DH) with phenotypes and genotypes (Figure 1)

mice Publicly available data set of a heterogeneous mice stock population from http://mus.well.ox.ac.uk/GSCAN recently analyzed by e.g. [4] comprising 1940 individuals genotyped with 12545 SNP markers and 2527 individuals phenotyped for the traits weight and growth slope

## References

[1] R Development Core Team (2010), http://www.R-project.org/

[2] vanRaden (2008) J Dairy Sci 91:4414-4423.

[3] Rogers (1972) In: Studies in genetics VII, Univ. of Texas, pp 145-153

[4] Legarra et al. (2008) Genetics 180:611-618

