



The synbreed R package: A framework for genomebased prediction using R

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Motivation

The synbreed R package [1] is designed to derive genome-based predictions from high-throughput genotyping and large scale phenotyping data. It contains a **comprehensive collection** of functions required to fit and cross-validate genome-based prediction models. All functions are embedded within the framework of a single, unified data object. Thereby a versatile genomic prediction analysis pipeline covering data processing, visualization, and analysis is established within **one software package**. The implementation is flexible with respect to a wide range of data formats and models. The package fills an existing gap in the availability of **user-friendly software** for next-generation genetics research and education.

Availability

The synbreed package is open-source and available through CRAN (see QR code below):

http://cran.r-project.org/web/packages/synbreed

The latest development version is available from R-Forge:

http://synbreed.r-forge.r-project.org

The package synbreed is released with a vignette (available using vignette("IntroSyn")), a manual and three large-scale example data sets from maize, cattle and mice (in package synbreedData, also on CRAN).

Example

Data: Simulated maize breeding program with 1250 doubled haploid (DH) lines fingerprinted for 1117 polymorphic SNPs and a quantitative trait evaluated in testcrosses of DH lines with one common tester.

Step 1 (Load data):

R> library(synbreed)

R> data(maize)

Step 2 (processing & filtering): Recoding SNP marker genotypes to the number of copies of the minor allele, i.e. 0, 1 and 2 and preselection of SNPs with a minor allele frequency (MAF) \geq 0.05 is conducted using

R> maizeC <- codeGeno(maize, maf=0.05)</pre>

Recoded marker genotypes were used to estimate pairwise linkage disequilibrium (LD) measured as r^2 by all marker pairs on chromosome 1 using

R> LD1 <- pairwiseLD(maizeC,chr=1)</pre>

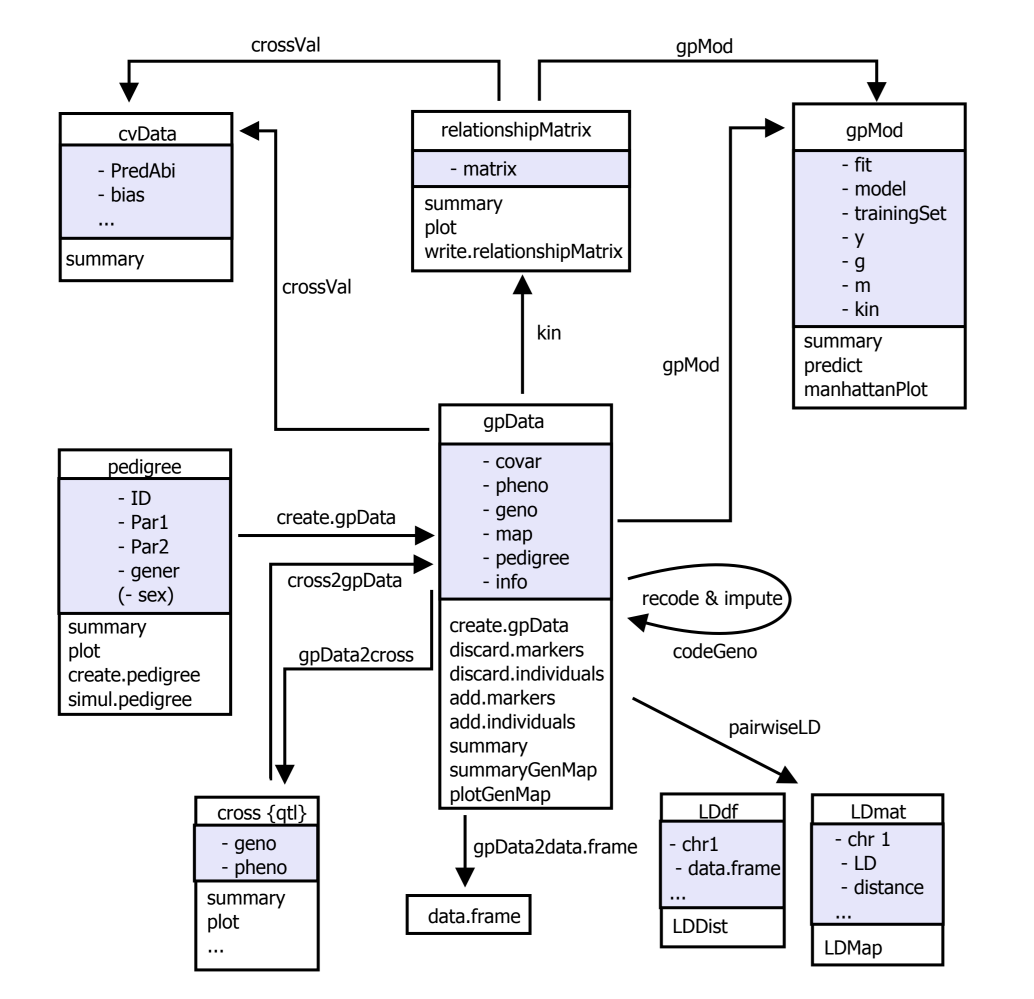
The extent of LD and LD decay is visualized (Figure 2) using

R> LDDist(LD1,type="bars",breaks=list(dist=c(0,10,20,40,60,180), + r2=c(1,0.6,0.4,0.3,0.1,0)))

Step 3 (kinship coefficients): The remaining 995 SNPs were used to estimate the realized relationship matrix for the 1250 DH lines based on the

Overview

The data flow in synbreed is guided by a single, unified data object of class gpData ("genomic prediction Data") which is used for storage of multiple data sources (see Figure 1).



recoded marker genotypes:

R> U <- kin(maizeC, ret="realized")</pre>

A heatmap visualization is available by using plot(U) (see Figure 3).

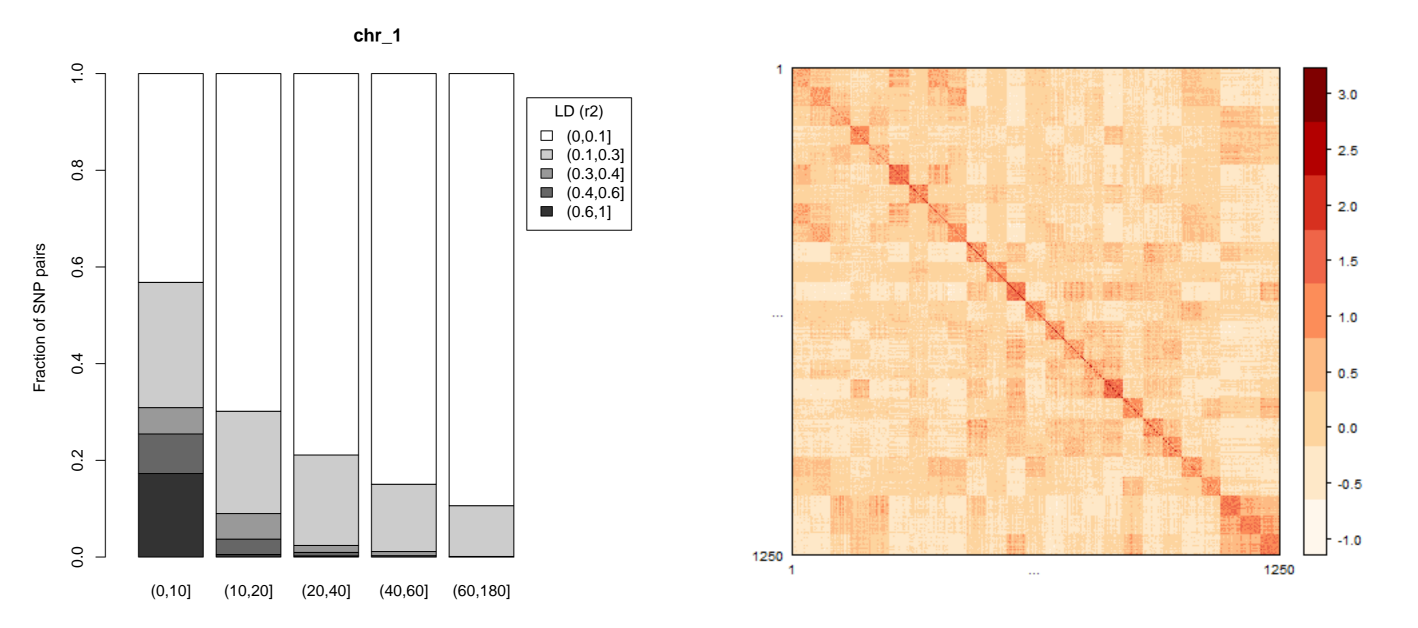


Figure 2: LD decay visualization for chromosome 1 **Figure 3:** Heatmap of the marker-based relationship matrix

Step 4 (prediction model): A GBLUP model for the testcross values is developed using the realized relationship matrix from step 3. For the prediction of testcross values, the relationship matrix must be replaced by the kinship matrix, i.e. divided by 2.

Figure 1: Overview of object classes, methods and functions within the synbreed package. Each box indicates a class together with the class name, the elements and the available functions and methods. The arrows indicate the data flow. The origin indicates the input argument and the head is the return value of the function.

References

[1] Wimmer V, Albrecht T, Auinger HJ, Schön CC (2012) synbreed: A framework for the analysis of genomic prediction data using R. Bioinformatics

R> GBLUP <- gpMod(maizeC, mod="BLUP", kin=U/2)</pre>

Step 5 (model validation): Finally, we estimate the predictive ability of GBLUP using 2-fold cross-validation with 5 replications each with a random assignment into estimation set (ES) and test set. The estimated variance components are committed from step 4 and used to build a prediction model within every ES:

R> cv <- crossVal(maizeC, k = 2, Rep = 5,

cov.matrix = list(U/2), varComp = GBLUP\$fit\$sigma, Seed=1) +

By using summary(cv) we obtain an average predictive ability of 0.48 with a range from 0.44 to 0.52.

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