The R-Package 'synbreed'

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Abstract

This document gives an introduction to the R-package synbreed. The package contains tools for plant and animal breeding utilizing quantitative genetics and statistical methods. The goal is to create an analysis pipeline for genomic selection. This comprises tools for genotypic, phenotypic and pedigree data. The steps of a typical analysis are presented in this document. This starts with the coding of the marker data, followed by the estimation of relatdness according to pedigree or molecular marker data, e.g. according to vanRaden (2008). Finally the estimation of breeding values and estimation of variance components using mixed models is described. All steps are illustrated using simulated data for maize.

Keywords: synergistic plant and animal breeding, simulation, pedigree, genomic marker data, mixed models, genomic selection

1 Introduction

The R-package synbreed aims to provide the tools that are necessary to analyze data of breeding programs and estimate (genomic) breeding values. Of course, there exists already software for this purpose. In R, package genetics contains classes and methods for handling genetic data (Warnes, 2003). Note that R-Genetics project has developed an set of enhanced genetics packages to replace genetics (http://rgenetics.org). Package qtl could be used for QTL analysis in experimental crosses (Broman and Churchill, 2003). Library GenABEL is designed for genome-wide association analysis and effective SNP data storage and manipulation (YS et al., 2007).

The idea of this package is to collect the methods in one package, so that analysis can be performed in one software with just a few steps as described in this document. Additional, this package takes care of special

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problems of modern breeding programs as the use of doubled haploid (DH) lines in plant breeding. To our knowledge, there is no package in R which provides comparable features. Most of packages source code is written in R, so that methods could easily be adopted for own purposes. Package synbreed makes no stringent restriction concerning input data format to allow for a wide range of possible data sources.

Modern breeding programs use genomic information of individuals. On the genomic level, individuals could be distinguished by alleles which are different states at a particular gene locus. In diploid species, every individuals has two sets of chromosomes and thus two copies of each allele at a locus. If both alleles are the same, the individual is homozygous for this locus, otherwise it heterozygous. For many species molecular markers are available and used to detect SNP (single nucleotide polymorphism) variation which occurs when a single nucleotide (A, T, C, or G) differs between individuals. In this document, the term genotype refers to an individual's set of alleles read by molecular markers and is used as a synonym for an individual. On the other hand, phenotype denotes the observed and measured value of a genotype, i.e. a trait of commercial interest. It is assumed that the phenotype is determined by a certain degree the genotype and by the environment.

The idea to use molecular markers in breeding is to predict genomic breeding values for the individuals based on marker information. In case a dense marker map is available, all quantitative trait loci (QTL) are assumed to be in linkage with at least one marker. To obtain genetic breeding values, Meuwissen et al. (2001) proposed to regress the phenotype on the markers (genotype). Once the model is available, individuals with a favorable set of genes are selected for the next cycle in breeding program. This is called genomic selection.

The remainder of this document is structured as follows. In section 2 a simulated data set is presented which is used to illustrate the methods in this document. Section 3 describes the coding of marker data and imputing for missing genotypic data. In Section 4 is shown how Linkage Disequilibrium (LD) between markers could be computed and visualized using synbreed package. Section 5 shows how to utilize pedigree information. In Section 6 basic concepts of quantitative genetics are introduced. Section 7 presents several methods to estimate relatedness between individuals. In section 8 the use of mixed models to estimate variance components and breeding values is illustrated. Section 9 presents full analysis pipeline for genomic selection comparing different models using simulated data.

2 Example data

In this document the steps of an analysis pipeline for genotypic and phenotypic data in plant or animal breeding with the R-package synbreed are

presented. For illustration, a the package contains a simulated data set for maize, called maize. This data set could be used to test performance of methods because estimated values could easily be compared with specified parameters of the simulation (position of QTL, size of marker effects, true breeding values for individuals). To load maize data, use

- > library(symbreed)
- > data(maize)

This data set contains genotypic and phenotypic data, as well as pedigree information up to grandparents for 1250 doubled haploid (DH) lines of maize. Performance of DH lines was evaluated in test crosses with a common single cross tester. All Dh lines were genotyped for 696 single nucleotide polymorphisms (SNP). When loading maize data, the following four data sets are loaded into workspace

maize.geno This is a data.frame containing the marker data of 696 biallelic SNP markers for the 1250 genotypes. The first column contains the ID to identify the genotypes. This variable should be used for the merge with the phenotypic data. The following columns contain data for each SNP. The marker data is coded with 0/1 with no missing values. Note that the coding does not contain any information about allele frequencies, thus 1 could be minor or major allele. As all genotyped individuals are fully inbred, no heterozygous genotypes are present.

maize.pheno This is a data.frame with column ID and column Trait containing the measured phenotypic trait (higher values indicate better performance). The order of the genotypes is the same as the order of rows in maize.geno.

maize.ped This data.frame contains the pedigree information of 1301 genotypes (1250 lines and 51 ancestors).

maize.marker.pos This data.frame contains additional information for the SNP markers. The first column pos gives the position of the marker on the chromosome in cM. Markers are order by their position within one chromosome. The second column chr sepecifies to which of the 10 chromosomes of maize (linkage group) the marker belongs (every marker is assigned to one linkage group). The order of the markers is the same as the order of columns in maize.geno.

3 Marker data

In the first step, genotypes have to be coded in a way that it could be used for the construction of genomic relationship matrices. The convention used in this package is to code genotypes by the number of minor alleles, i.e. 0 for one homozygous genotypes and 2 for the other and 1 for heterozygous. This task is done by the function codeGeno. If no missing values and no heterozygous genotypes for any loci are present and all markers should be used in the following analyses, this function does simply recode the alleles to 0 (major) and 2 (minor) as mentioned above. For the maize data, use

```
> marker <- codeGeno(maize.geno[, -1])</pre>
```

```
step 1 : No markers removed due to fraction of missing values
step 2 : Recoding alleles
step 5 : No markers discarded due to minor allele frequency
step 6 : No duplicated markers discarded
step 7 : Restoring original data format
```

to obtain an object marker which contains the recoded marker data. Note that the first column is not used because it contains the ID. Now, the minor allele frequencies are easily obtained by dividing the column means of marker by 2. A kernel density estimate of minor allele frequency (MAF) is shown in Figure 1.

```
> plot(density(colMeans(marker)/2), xlab = "MAF", main = "")
```

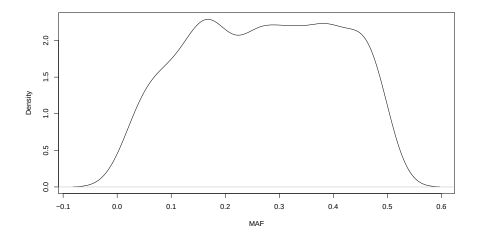


Figure 1: Kernel density estimate of the minor allele frequency of the 696 SNP markers in maize data.

In experimental data usually missing values occur in genotypic data due to different reasons. The function codeGeno could be used to impute missing values by chance or according to family structure using the following rules:

with population structure Suppose an observation i is missing (NA) for a marker j in population k. If marker j is fixed in population k, the imputed value will be the fixed allele. If marker j is segregating in population k, the value is 0 with probability 0.5 and 2 with probability 0.5.

without population structure The missing values for a marker j are sampled from the allele distribution of marker j.

An alternative is to treat missing values as heteroyzgous genotypes using argument replace.value of function codeGeno. A complete data set of genotypes is important if marker data should be used to estimate relatedness between individuals. To illustrate imputing of missing values, 200 entries of the marker matrix are selected, the values saved and these entries are coded as NA.

```
> marker <- as.matrix(marker)
> ind1 <- sample(1:nrow(marker), 200)
> ind2 <- sample(1:ncol(marker), 200)
> posNA <- cbind(ind1, ind2)
> original <- marker[posNA]
> marker[posNA] <- NA</pre>
```

Note that missing values in the marker data must be coded as NA. The 1250 genotypes in the maize data comprise 25 half sib families with 50 genotypes in each family. The genotypes are ordered by family, thus population structure simply is

```
> pop <- rep(1:25, each = 50)
```

Recoding of the marker data and imputing of the missing values using population structure is done as follows

```
> marker1 <- codeGeno(marker, impute = TRUE, pop)</pre>
step 1 : No markers removed due to fraction of missing values
step 2 : Recoding alleles
step 3: Imputing of missing values by population structure
... approximative run time 2.97 seconds ...
step 4 : No recoding of alleles necessary after imputation
step 5 : No markers discarded due to minor allele frequency
step 6 : No duplicated markers discarded
step 7 : Restoring original data format
Summary of imputation
  total number of missing values
                                                : 200
 number of imputations by family structure
                                               : 123
 number of random imputations
                                                : 77
```

approximate fraction of correct imputations

: 0.808

A report is printed on the screen which informs about the number of imputations performed either according to family structure n_F or chance n_R . The approximate fraction of correct imputations is $\frac{n_F+0.5n_R}{n_F+n_R}$. For simulated data original values are known. The quality of the classification of the missing values is judged in the following cross-table of imputed and original values

The fraction of correct replacements is

```
> sum(diag(t1))/sum(t1)
[1] 0.79
```

which is much higher than the expected fraction of correct imputations without family structure which equals 0.5. At the moment there is no possibility to impute missing values using information of other (flanking) markers. This is the best alternative if no population structure is given. For this purpose, package function fill.geno of package qtl could be used.

In an analysis of genotypic data molecular markers with a small minor allele frequency and/or many missing values are discarded using arguments maf and nmiss of function codeGeno. In this case all markers with more than nmiss·100% missing values are discarded before recoding and after recoding only markers with a minor allele frequency > maf are returned by the function. By default, no markers are selected by one of both criteria, thus maf=nmiss=0.

4 Linkage Disequilibrium

Linkage Disequilibrium (LD) is defined as a non-random association between polymorphisms of two or more molecular markers (usually part of same linkage group). It appears because many individuals can inherit a linked allele pair from an ancestor without any recombination event. Average LD over the whole genome is important as it influences prospects of genomic selection. It is calculated as the difference between observed and expected (assuming random distributions) allelic frequencies. There are many possibilities to compute LD from genotypic data, see Foulkes (2009) using genetics package. In synbreed package, LD between two loci i and j denoted as LD_{ij} is computed as coefficient of determination R^2 of a regression of $\mathbf{x_i}$ on $\mathbf{x_j}$,

where $\mathbf{x_i}$ and $\mathbf{x_j}$ are a *n*-dimensional vectors containing marker data of *n* individuals. This equals squared correlation coefficient of both data vectors, thus

$$LD_{ij} = r(\mathbf{x_i}, \mathbf{x_j})^2.$$

Relationship between markers is assumed to decrease as their distance on chromosome increases. To visualize this decline, the LD is plotted against the distance on the chromosome. For maize we plot the LD for the marker located on the first (out of 10) chromosomes using function LDDist. This function provides several possibilities for graphical visualisation. To visualize LD decay on first chromosome of maize data, use

```
> data(maize)
> marker <- codeGeno(maize.geno[, -1])
> LDDist(marker, maize.marker.pos$chr, maize.marker.pos$pos,
+ chr = 1)
```

Figure 2(a) and 2(b) present two possibilites for graphical visulisation using different values for argument type.

As genotypes are effected by selection due in simulation progress for many generations, LD above 0.6 is only observable up to a distance of 10 cM for the first chromosome, see Figure 2(b). Another type of graphic is a heatmap of LD for markers located on the same chromosome. LD heatmap for pairwise LD of markers on the first chromosome is obtained by function LDMap which is simply a wrapper for funtion LDheatmap of package LDheatmap.

Strong LD could be observed between the markers on the "left" margin of Chromosome 1, compare Figure 3.

5 Pedigree

An important source of information in breeding programs is pedigree information. Especially in animal breeding, pedigree is recorded over many generations. The pedigree usually consists of a list of individuals (animals or plants) of the current generation which is the subject of analysis and their ancestors (for which often no phenotypic data is available). The pedigree is sorted by generation, beginning with the individuals with unknown parents. An example for a pedigree with five individuals belonging to 4 generations is given below. Parents of A and B are unknown.

In synbreed class "pedigree" should be used for handling pedigree information. An object of class "pedigree" consists of a data.frame with at least variables ID, Par1, Par2 and gener. The function create.pedigree creates an object of class "pedigree" for a given set of individuals and the pair of parents. Note that unknown parents are coded as "0" in synbreed package and generation starts with 0. The generation can be specified by the user or optional computed by the function.

ID	Par1	Par2	gener
A	-	-	0
В	-	-	0
\mathbf{C}	A	В	1
D	A	\mathbf{C}	2
E	D	В	3

Suppose we have the pedigree structure of the example. This structure is carried into synbreed package with the following commands

```
> id <- c("A", "B", "C", "D", "E")
> par1 <- c(0, 0, "A", "A", "D")
> par2 <- c(0, 0, "B", "C", "B")
> ped <- create.pedigree(id, par1, par2)
> ped
  ID Par1 Par2 gener
        0
             0
1
  Α
2
  B
        0
             0
                   0
3
   C
             В
                   1
        Α
             C
4 D
                   2
        Α
5 E
```

An object of class "pedigree" could be visualized with generic plotting function for S3 class "pedigree".

It is possible to simulate a pedigree structure with function simul.pedigree. As arguments, the number of generations to simulate and the number of individuals in each generation has to be specified. By default, random mating is assumed in each generation. As there are no further restrictions, it is possible that inbreeds could be generated when parent 1 equals parent 2. To simulate a pedigree with 6 generations and 4, 6, 7, 9, 10 and 10 individuals in each generation, use

```
> set.seed(123)
> ped.simul <- simul.pedigree(gener = 6, ids = c(4, 6,
+ 7, 9, 10, 10))</pre>
```

The resulting pedigree is visualized in Figure 4. A basic summary of the pedigree is given by the generic summary method for class "pedigree".

```
# individuals : 46
# Par1 (sire) : 27
# Par2 (dam) : 25
# generations : 6
# unknow parents : 8
# inbred : 5
```

> summary(ped.simul)

6 Genotypic Means and Variances

This section summarazies some theory of quantitative genetics which is necessarry for the following sections. Those readers who are familiar with the concepts of genetics as presented in Falconer and Mackay (1996) or Bernardo (2002) may skip this section. All formulas are presented for the one-locus model in a breeding population.

We consider a single locus with two alleles A_1 and A_2 so that genotypes A_1A_1 , A_1A_2 and A_2A_2 are possible. The phenotypic mean G_{ij} of individuals having the A_iA_j genotype can be expressed as

$$P_{ij} = G_{ij} + e_{ij}$$

where G_{ij} is the genotypic value of A_iA_j and e_{ij} is the nongenetic residual which is assumed to be unrelated to genotypic value. The genotypic value can be further divided into

$$G_{ij} = \mu + \alpha_i + \alpha_j + \delta_{ij}$$

where μ is the population mean, α_i is the effect of the A_i allele which is defined as the average effect of an allele. This is the difference of the mean of those individuals who received A_i allele from one parent and the other allele at random from the population. The effect α_j is defined in a similar way. The breeding value of the A_iA_j genotype is defined as the sum of α_i and α_j . The term δ_{ij} is the dominance deviation of the A_iA_j genotype. Dominance deviation is due to the interaction of both alleles at one locus.

On a population level, we consider the variance of the effects mentioned above. Phenotypic variance V_P can be partioned into genotypic variance V_G and non genetic variance V_E . The genetic variance within one locus is divided into additive variance and dominance variance, thus $V_G = V_A + V_D$. If more than one locus is considered, additional epistatic variance V_I occurs which is due to interactions between different loci. Epistatic variance consists in two locus model of additive-additive variance V_{AA} , additive-dominace variance V_{AD} and dominance-dominance variance V_{DD} due to the two-way interactions of additive and dominance effects.

An important concept in breeding is the covariance between relatives which is a function of the probability that two alleles are identical by descent and genetic variance components. Suppose we have the pedigree structure as shown in Figure 5.

Covariance between relatives is due to alleles that are identical by descent (IBD), denoted by the \equiv symbol. There are four possibilities that two alleles in X and Y are IBD and covariance due to breeding values is given by

$$Cov_{\alpha} = P(A_i \equiv A_k)Cov(\alpha_i, \alpha_k) + P(A_i \equiv A_l)Cov(\alpha_i, \alpha_l)$$

$$+ P(A_j \equiv A_k)Cov(\alpha_j, \alpha_k) + P(A_j \equiv A_l)Cov(\alpha_j, \alpha_l)$$

$$= 2f_{XY}V_A$$

where f_{XY} is the coefficient of coancestry (Wright, 1922) which is defined as

$$f_{XY} = P(X \equiv Y)$$

$$= \frac{1}{4} [P(x_1 \equiv y_1) + P(x_1 \equiv y_2) + P(x_2 \equiv y_1) + P(x_2 \equiv y_2)]. (1)$$

Covariance between relatives according to dominance and epistatic variances can be derived in a similar way, see Bernardo (2002) and next section.

7 Estimation of Relatedness

This section presents methods to estimate relatedness between a set of individuals. The goal is to set up a variance-covariance matrix for the individuals. Relatedness between individuals refers to their covariance due to some kind of genetic effect. In classical breeding programs, pedigree information is used to compute *expected* relatedness. Thus the entries of the variance-covariance matrix are derived by the formulas of the previous section. If molecular markers are available, *observed* relatedness between individuals could be estimated using similarity measures for genotypic data.

7.1 Based on Pedigree

The computation of the pedigree based relatedness in **synbreed** starts with the gametic relationship. A gamete is the genetic unit that an individual passes to its offspring. The genotype of a diploid individual at one locus consists of two alleles. Suppose there is an individual C with parents A and B. Individual C has to alleles C_1 and C_2 . The source of allele C_1 is Parent A, thus allele C_1 could either be IBD to A_1 or A_2 which are the alleles (and possible gametes) of A. Allele C_2 was inherited of parent B, thus it could be IBD to B_1 or B_2 which are the alleles of B. To compute the gametic relationship start with an expanded table with two alleles for each individual.

This table is converted into the gametic relationship G matrix which is of order 2n, if the number of individuals is n. The entires of G are the probability that two alleles A_1 and A_2 are identical by descent (IBD), denoted as $P(A_1 \equiv A_2)$. Thus diagonal values are always 1. If parents are unknown, they are assumed as progeny of a random mating population. In this case the off-diagonals are zero. The gametic relationship matrix is constructed recursively, starting with the first generation in pedigree. The combination of $2^2 = 4$ alleles that describe the relationship of progeny A

ID	Allele	Par1	Par2
A	A_1	-	-
A	A_2	-	-
В	B_1	-	-
В	B_2	-	-
\mathbf{C}	C_1	A_1	A_2
\mathbf{C}	C_2	B_1	B_2
D	D_1	A_1	A_2
D	D_2	C_1	C_2
\mathbf{E}	E_1	D_1	D_2
E	E_2	B_1	B_2

with parent C are computed as follows

$$P(A_1 \equiv C_1) = 0.5 \cdot [P(A_1 \equiv A_1) + P(A_1 \equiv A_2)]$$

$$P(A_1 \equiv C_2) = 0.5 \cdot [P(A_1 \equiv B_1) + P(A_1 \equiv B_2)]$$

$$P(A_2 \equiv C_1) = 0.5 \cdot [P(A_2 \equiv A_1) + P(A_2 \equiv A_2)]$$

$$P(A_2 \equiv C_2) = 0.5 \cdot [P(A_2 \equiv B_1) + P(A_2 \equiv B_2)]$$

The only nonzero probability in the equations above is $P(A_1 \equiv A_1)$ which is 1. The gametic relationship for all individuals of a given pedigree is obtained as follows

```
> G <- kinship(ped, ret = "gam")
> G
```

```
B_2 C_1 C_2
                                              D_2
                                                    E_1
            A_2
                  B_{\perp}1
                                       D_1
                                                          E_2
A_1 1.000 0.000 0.000 0.000 0.5 0.00 0.500 0.250 0.375 0.000
A_2 0.000 1.000 0.000 0.000 0.5 0.00 0.500 0.250 0.375 0.000
B_1 0.000 0.000 1.000 0.000 0.0 0.50 0.000 0.250 0.125 0.500
B_2 0.000 0.000 0.000 1.000 0.0 0.50 0.000 0.250 0.125 0.500
C_1 0.500 0.500 0.000 0.000 1.0 0.00 0.500 0.500 0.500 0.000
C_2 0.000 0.000 0.500 0.500 0.0 1.00 0.000 0.500 0.250 0.500
D_1 0.500 0.500 0.000 0.000 0.5 0.00 1.000 0.250 0.625 0.000
D_2 0.250 0.250 0.250 0.250 0.50 0.50 0.250 1.000 0.625 0.250
E_1 0.375 0.375 0.125 0.125 0.5 0.25 0.625 0.625 1.000 0.125
E_2 0.000 0.000 0.500 0.500 0.0 0.50 0.000 0.250 0.125 1.000
attr(,"class")
[1] "relationshipMatrix"
```

The resulting object G is of class "relationshipMatrix" which is the general class for all variance-covariance matrices due to genetic effects (additive, dominance or epistatic). To derive the variance-covariance matrix due to an genetic effect, one has to multiply this matrix with the corresponding variance component $(V_A, V_D \text{ or } V_I)$ as shown below. Once the gametic

relationship is computed, it could be converted in the additive numerator relationship matrix \mathbf{A} which is due to additive genetic effects (breeding values) or the dominance relationship matrix \mathbf{D} which is due to dominance effects

In case of inbreeding, the entry in G of allele A_1 and allele A_2 of an individual X equals his inbreeding coefficient

$$F_X = P(A_1 \equiv A_2).$$

For example, the inbreeding coefficient of individual D is

```
> as.numeric(G["D_1", "D_2"])
```

[1] 0.25

which is nonzero because individuals A and C, which are the parents of D, are relatives.

The additive relationship between individuals X and Y is given by

$$A_{XY} = \begin{cases} 1 + F_X, & X = Y \\ 2f_{XY}, & X \neq Y \end{cases}$$

thus off-diagonal entries equal twice teh kinship coefficient. The additive numerator relationship is of order n and derived for a given pedigree as

```
> A <- kinship(ped, ret = "add")
> A
```

```
A B C D E
A 1.000 0.000 0.500 0.75 0.375
B 0.000 1.000 0.500 0.25 0.625
C 0.500 0.500 1.000 0.75 0.625
D 0.750 0.250 0.750 1.25 0.750
E 0.375 0.625 0.625 0.75 1.125
attr(,"class")
[1] "relationshipMatrix"
```

The relationship between individuals A and C is 0.5 which is the expected value for parent-offspring relation (Bernardo, 2002). Note that the diagonals of **A** are $1+F_i$ for i=1,...,n. Inbreeding coefficients could easily be obtained as

```
> diag(A) - 1
```

```
A B C D E
0.000 0.000 0.000 0.250 0.125
```

Sometimes the kinship matrix is required, which is half of the additive numerator relationship matrix. It is obtained by

> K <- kinship(ped, ret = "kin")</pre>

Dominance covariance matrix is computed if argument ret="dom" is used. Variance-covariance matrices for epistatic effects as additive-additive (AA), additive-dominance (AD) or dominance-dominance (DD) are the products of the corresponding variance-covariance matrices for additice and dominace effects. Models for genomic selection can be extended by these effects, but the contribution of three-way or higher interactions is usually small (Bernardo, 2002).

7.2 Based on marker data

Relatedness based on marker is an alternative to estimate relationship instead of using additive numerator relationship matrix based on pedigree. Marker data could be used to estimate relationship between relatives more precise than the numerator relationship based on pedigree as it takes into account the Mendelian sampling effect. Two methods for the construction of a relationship matrix based on marker data are implemented in the synbreed package: genomic relationship according to vanRaden (vanRaden, 2008) and according to Roger's distance (Rogers, 1972). Both methods are kinds of similarity measures which compare the number of alleles that two individuals share.

For vanRaden, the SNP genotypes are coded as the number of copies of the minor allele, i.e., 0, 1 or 2 (no missing values allowed). Thus the marker data could be the result of a call of codeGeno when imputing for the missing values was performed or the missing values were replaced with the value 1. The genomic relationship matrix according to vanRaden for n individuals and p molecular markers is computed as

$$\frac{\mathbf{Z}\mathbf{Z}'}{2\sum_{i=1}^{p} p_i (1-p_i)},\tag{2}$$

where $\mathbf{Z} = \mathbf{M} - \mathbf{P}$ and \mathbf{M} is the $n \times p$ marker matrix and \mathbf{P} contains the allele frequencies multiplied by 2. In (2) p_i is the allele frequency of marker i. As an example we look at the marker data of 6 individuals genotyped with 8 SNP markers. Let

$$\mathbf{M} = \begin{pmatrix} 2 & 0 & 0 & 2 & 2 & 0 & 0 & 0 \\ 2 & 0 & 2 & 2 & 2 & 0 & 2 & 2 \\ 2 & 0 & 2 & 2 & 0 & 0 & 2 & 0 \\ 0 & 0 & 2 & 2 & 0 & 0 & 2 & 0 \\ 0 & 0 & 2 & 0 & 0 & 0 & 2 & 0 \\ 2 & 2 & 2 & 2 & 0 & 0 & 0 & 2 \end{pmatrix},$$

then it holds that

$$\mathbf{P} = \begin{pmatrix} 1.33 & 0.33 & 1.67 & 1.67 & 0.67 & 0 & 1.33 & 0.67 \\ 1.33 & 0.33 & 1.67 & 1.67 & 0.67 & 0 & 1.33 & 0.67 \\ 1.33 & 0.33 & 1.67 & 1.67 & 0.67 & 0 & 1.33 & 0.67 \\ 1.33 & 0.33 & 1.67 & 1.67 & 0.67 & 0 & 1.33 & 0.67 \\ 1.33 & 0.33 & 1.67 & 1.67 & 0.67 & 0 & 1.33 & 0.67 \\ 1.33 & 0.33 & 1.67 & 1.67 & 0.67 & 0 & 1.33 & 0.67 \end{pmatrix}$$

$$\mathbf{Z} = \begin{pmatrix} 0.67 & -0.33 & -1.67 & 0.33 & 1.33 & 0.00 & -1.33 & -0.67 \\ 0.67 & -0.33 & 0.33 & 0.33 & 1.33 & 0.00 & 0.67 & 1.33 \\ 0.67 & -0.33 & 0.33 & 0.33 & -0.67 & 0.00 & 0.67 & -0.67 \\ -1.33 & -0.33 & 0.33 & 0.33 & -0.67 & 0.00 & 0.67 & -0.67 \\ -1.33 & -0.33 & 0.33 & -1.67 & -0.67 & 0.00 & 0.67 & -0.67 \\ 0.67 & 1.67 & 0.33 & 0.33 & -0.67 & 0.00 & -1.33 & 1.33 \end{pmatrix}$$

and

$$\mathbf{ZZ'} = \begin{pmatrix} 7.44 & 0.11 & -1.22 & -2.56 & -3.22 & -0.56 \\ 0.11 & 4.78 & -0.56 & -1.89 & -2.56 & 0.11 \\ -1.22 & -0.56 & 2.11 & 0.78 & 0.11 & -1.22 \\ -2.56 & -1.89 & 0.78 & 3.44 & 2.78 & -2.56 \\ -3.22 & -2.56 & 0.11 & 2.78 & 6.11 & -3.22 \\ -0.56 & 0.11 & -1.22 & -2.56 & -3.22 & 7.44 \end{pmatrix}$$

with the denominator being $2\sum_{i=1}^{p}p_i(1-p_i)=2.611$. Correcting with the allele frequencies gives more weight to individuals that share a rare allele. To compute the genomic relationship according to vanRaden, matrix **M** is passed to the function vanRaden

```
2, 2, 0, 2, 2, 2, 0, 2, 2, 0, 0, 2, 0, 0, 0, 2, 2,
     0, 0, 2, 0, 0, 0, 2, 0, 0, 0, 2, 0, 2, 2, 2, 2, 0,
     0, 0, 2), nrow = 6, byrow = TRUE)
> vR <- vanRaden(M)
> round(vR, 3)
      [,1]
             [,2]
                   [,3]
                          [,4]
                                [,5]
[1,]
            0.043 -0.468 -0.979 -1.234 -0.213
     2.851
[2,] 0.043
           1.830 -0.213 -0.723 -0.979 0.043
[3,] -0.468 -0.213  0.809  0.298  0.043 -0.468
[4,] -0.979 -0.723 0.298
                       1.319
                               1.064 -0.979
[5,] -1.234 -0.979 0.043 1.064 2.340 -1.234
attr(,"class")
[1] "relationshipMatrix"
```

 $> M \leftarrow matrix(data = c(2, 0, 0, 2, 2, 0, 0, 0, 2, 0, 2,$

Note the object vR is again of class "relationshipMatrix". Negative values indicate pairs of individuals sharing fewer alleles than expected by the allele frequencies.

Another possibility is to compute the genomic relationship matrix according to Roger's distance. Roger's distance is computed as

$$d = \frac{1}{p} \sum_{i=1}^{p} \sqrt{1/2 \sum_{j=1}^{n_i} (p_{ij} - q_{ij})^2}$$
 (3)

where p is the number of markers and n_i is the number of alleles for marker i. Let p_{ij} and q_{ij} denote the allele frequencies of allele j for marker i respectively. Note that marker data should be coded -1 and 1 for homozygous genotypes and 0 for heterozygous. If marker data is coded as 0/1/2, data is transformed by function rogers, which computes relationship based on Roger's distance. Using transformation of Hayes and Goddard (2008) rogers distance is related to relationship as

$$f = \frac{s - s_{min}}{1 - s_{min}},$$

with similarity measure s = 1 - d and s_{min} minimum of all $\frac{n}{2}(n+1)$ values for s. Using rogers distance to compute relationship based on marker data gives

```
> round(ro, 3)
     [,1] [,2] [,3] [,4] [,5] [,6]
    2.0 0.8 0.8 0.4 0.0
    0.8 2.0
               1.2
[3,] 0.8 1.2
               2.0
                   1.6
[4,] 0.4 0.8
              1.6
                  2.0
[5,] 0.0 0.4
              1.2 1.6 2.0 0.0
[6,] 0.4 0.8
              0.8 0.4 0.0
attr(, "class")
[1] "relationshipMatrix"
```

> ro <- rogers(M, correction = "Hayes")</pre>

Note that function rogers returns 2f which is the relationship between individuals.

7.3 Doubled haploid lines

In modern plant breeding programs in maize, doubled haploid (DH) lines are used as parents in crosses. DH lines are fully inbred and thus have an inbreeding coefficient of 1. This has to be taken into account, when the relationship matrix in a pedigree with DH lines is computed. As an example the maize data is taken.

```
> data(maize)
> head(maize.ped)
```

```
ID Par1 Par2 DH
  1
       0
           0
              1
2
 2
            0 1
       0
3 3
       0
            0 1
 4
            0 1
5
 5
       0
            0 1
6
  6
       0
            0
              1
```

Last column is a logical that indicates DH lines. First, the additive numerator relationship matrix is computed. There are 1276 DH lines and 25 non DH lines in the pedigree. For DH lines special treatment is necessary, as the inbreeding coefficient must be 1 and no heterozygous genotypes occur. An argument DH is available for function kinship where for each individual in the pedigree it specified whether this is a DH line or not. This information is available for the maize data. To obtain the additive numerator relationship matrix, use

```
> ped.maize <- create.pedigree(maize.ped$ID, maize.ped$Par1,
+ maize.ped$Par2)
> A.maize <- kinship(ped.maize, DH = maize.ped$DH, ret = "add")
> dim(A.maize)

[1] 1301 1301
All inbreeding coefficients of DH lines are now one
> head(diag(A.maize) - 1)

1 2 3 4 5 6
1 1 1 1 1 1
```

7.4 Visualization of relationship matrices

In most cases a relationship matrix is too big to print on screen. There are two possibilities for visualization of an object of class "relationshipMatrix" in synbreed package. A generic summary method is defined which gives the important characteristics of a relationship matrix. Use

```
Dimension : 1301 x 1301
Rank : 1276
Range : 0 -- 2
# of unique values: 6
```

> summary(A.maize)

to get the summary for the pedigree based additive relationship matrix of the maize data. Another possibility is the plot method which could be applied to an object of class "relationshipMatrix". This gives a heatmap of the entries of the relationship matrix

Note that objects of class "relationshipMatrix" can be written to input files appropriate for Mixed Model software as WOMBAT (Meyer, 2006) or ASReml (Gilmour et al., 2000) using function write.relationshipMatrix. File formats are *.gin for WOMBAT and *.grm or *.giv for ASReml.

8 Mixed Models

The ultimate aim in the analysis of a breeding program is to estimate genetic effects and variance components. The basic statistical model for this purpose is a linear mixed model (Henderson, 1984) denoted by

$$y = Xb + Zu + e, (4)$$

where \mathbf{y} is the $n \times 1$ vector of phenotypic records, \mathbf{b} a $t \times 1$ vector of fixed effects and \mathbf{u} a $m \times 1$ vector of random effects. \mathbf{X} and \mathbf{Z} are the corresponding design matrices with dimension $n \times t$ and $n \times m$ respectively. The random genetic effect has distribution

$$\mathbf{u} \sim \mathrm{N}(\mathbf{0}, \mathbf{G}\sigma_{\mathbf{g}}^2)$$

where **G** is a genetic variance-covariance matrix. In (4) **e** denotes the $n \times 1$ vector of residuals with $\mathbf{e} \sim \mathrm{N}(\mathbf{0}, \mathbf{I_n} \sigma^2)$ and $\mathbf{I_n}$ is the *n*-dimensional identity matrix.

If **G** equals the additive numerator relationship matrix **A**, model (4) is called *animal model*. Here the random effect is usually denoted as **a** and $\mathbf{a} \sim \mathrm{N}(\mathbf{0}, \mathbf{A}\sigma_{\mathbf{a}}^2)$. This model is used to estimate additive genetic effects (breeding values) based on phenotypic records and pedigree information. As an example we will simulate plant breeding data. A pedigree with 5 generations and 20 individuals in each generation is simulated. Phenotypic data was measured in a field trial consisting of 5 locations with two replications (blocks) within locations for each of the n=100 genotypes.

The simulation of phenotypes is done with the function simul.phenotype which simulates records based on model (4) with an overall mean as fixed effect and random effects for genotype, location and block. Random effects for location, block nested in location and residuals are i.i.d. normal with mean zero and given variance component. Random effects for genotype are taken from a multivariate normal distribution $N(\mathbf{0}, \mathbf{A}\sigma_{\mathbf{a}}^2)$, where \mathbf{A} is the numerator relationship matrix obtained by pedigree information. The additive genetic variance σ_a^2 and variance components for location, block and residual are specified by the user. Variance components are specified in a

separate list and values used for simulation are $\sigma_a^2 = 10$, $\sigma^2 = 15$ and 0 for location and block nested in location. Simulated data is obtained as follows

Variance components for location and block are set to zero as only additive genetic effects should be considered in this example. The simulated random effects for genotype are called true breeding values (TBV) and returned by function simul.phenotype. Note that TBV for each ID is the same for all replications across locations and blocks.

Estimation of variance components with REML and prediction of random effects in model (4) could be done with function regress in package regress (Clifford and McCullagh, 2006). This package allows arbitrary variance-covariance matrices of random effects. Solutions for animal model with overall mean as fixed effect are obtained as

```
> library(regress)
> A <- kinship(ped, ret = "add")
> A <- A %x% matrix(1, 10, 10)
> mod <- regress(Trait ~ 1, Vformula = ~A, data = dat)
> summary(mod)
Maximised Residual Log Likelihood is -1950.051
Linear Coefficients:
            Estimate Std. Error
 (Intercept) 99.897
                          0.762
Variance Coefficients:
            Estimate Std. Error
              10.349 1.938
         In
               15.544
                          0.731
```

Note that variance-covariance matrix must be of same dimension as y. This could easily be obtained by using the Kronecker product to enlarge relationship matrix as data is sorted by individuals. This is shown in the code

above. Estimated variance components could be used to estimate narrowsense heritability as in Piepho and Möhring (2007) as

$$\hat{h}^2 = \frac{\hat{\sigma}_a^2}{\hat{\sigma}_a^2 + \hat{\sigma}^2} = \frac{10.35}{10.35 + 15.54} = 0.4.$$

The fitted model contains only one random effect, thus estimated breeding values are obtained as

```
> ebv <- mod$predicted - mod$fitted
```

as predicted equals $\hat{\mathbf{y}}$ and fitted contains estimated overall mean $\hat{\mathbf{m}}\mathbf{u}$ and $\hat{\mathbf{a}} = \hat{\mathbf{y}} - \hat{\mathbf{m}}\mathbf{u}$. Correlation between observed and estimated phenotypes (also called *predictive ability* of the model, see Legarra et al. (2008)) is

```
> y <- dat$Trait
> cor(ebv, y)

[,1]
[1,] 0.5823288
```

As true breeding values are known in simulate, also correlation between estimated and true genetic effect (called *accuracy* of the model) is available

```
> tbv <- dat$TBV
> cor(ebv, tbv)

[,1]
[1,] 0.9067806
```

9 Genomic Selection

In this section, all steps that are necessary to derive (genomic) breeding values out of genotypic and phenotypic data are presented using maize data. Different models that make use of pedigree and/or marker data are compared due to estimated effects and variance components.

First, additive relationship matrix based on pedigree is created

```
> data(maize)
> ped.maize <- create.pedigree(maize.ped$ID, maize.ped$Par1,
+ maize.ped$Par2)
> A <- kinship(ped.maize, DH = maize.ped$DH, ret = "add")
> summary(A)
```

Dimension : 1301 x 1301
Rank : 1276
Range : 0 -- 2
of unique values: 6

Additive relationship matrix was constructed using full pedigree, but first 51 individuals have no phenotypes and should not be used for further analysis, i.e. breeding values for parents are not of interest. For further analysis, only those elements which belong to phenotypes are used.

```
> A \leftarrow A[-c(1:51), -c(1:51)]
```

Marker data is coded by the number of minor alleles, i.e. 0, 1 and 2. As simulated data contains no missing values no further preparation to estimate variance-covariance matrix using method of vanRaden is necessarry

```
> marker <- codeGeno(maize.geno[, -1])</pre>
```

```
step 1 : No markers removed due to fraction of missing values
step 2 : Recoding alleles
step 5 : No markers discarded due to minor allele frequency
step 6 : No duplicated markers discarded
step 7 : Restoring original data format

> G <- vanRaden(marker)
> summary(G)

Dimension : 1250 x 1250
Rank : 692
Range : -0.8276297 -- 2.469210
# of unique values: 639886
```

Both pedigree based and marker based variance-covariance matrices were used in the following models

```
\begin{array}{lll} \text{Model 1 : } \mathbf{y} &=& \mu + \mathbf{Z}\mathbf{a} + \mathbf{e} \\ \text{Model 2 : } \mathbf{y} &=& \mu + \mathbf{Z}\mathbf{u} + \mathbf{e} \\ \text{Model 3 : } \mathbf{y} &=& \mu + \mathbf{Z}\mathbf{a} + \mathbf{Z}\mathbf{u} + \mathbf{e} \end{array}
```

Additive genetic effect due to pedigree has distribution $\mathbf{a} \sim \mathrm{N}(\mathbf{0}, \mathbf{A}\sigma_{\mathbf{a}}^2)$, genetic effect due to marker data hat $\mathbf{u} \sim \mathrm{N}(\mathbf{0}, \mathbf{G}\sigma_{\mathbf{g}}^2)$ and residual term in all models is $\mathbf{e} \sim \mathrm{N}(\mathbf{0}, \sigma^2)$. Estimation of breeding values and variance components is performed using function regress

Model	LogL	$\hat{\mu}$	$\hat{\sigma}_a^2$	$\hat{\sigma}_g^2$	$\hat{\sigma}^2$
1	-3139.6	1193.7	24.3		12
2	-3028.2	1194.2		16.7	37.1
3	-3028.2	1194.2	1.3	16.6	34.8

Table 1: Comparision of Models for Genomic Selection in maize data.

```
> library(regress)
> y <- maize.pheno$Trait
> mod1 <- regress(y ~ 1, Vformula = ~A)
> mod2 <- regress(y ~ 1, Vformula = ~G)
> mod3 <- regress(y ~ 1, Vformula = ~A + G)</pre>
```

Estimates for variance components and fixed effects are shown in the summary of the models. In Table 1 the three models are compared.

Model 2 results in a better model adaption than Model 1 regarding the Likelihood. Thus using genomic relationship instead of additive relationship based on pedigree as covariance matrix for breeding values results in a better model fit. Combining both information in Model 3 does not further improve model adaption. Correlation between estimated breeding value and phenotype is 0.998 for Model 1, 0.693 for Model 2 and 0.75 for Model 3 respectively.

In maize data, each genotype has only one replication. In most breeding programs each genotype is planted in several locations and perhaps replicated in each location. In this case, there are two possibilities to analyze data:

- Two-stage: Reduce data to one phenotype for each genotype by using adjusted genotypw means. Evalution of adjusted means can be carried out as shown in the example
- One-stage: Use all observation in one model and extend the mixed model by fixed or random effects for location and block

10 Summary

The preceding examples illustrated, how genomic selection could be applied to a breeding program where genotypic and phenotypic data is available. As shown in simulated data, genomic selection is expected to outperform classial selection based on pedigree and phenotypes. The main steps of analysis are as follows

 Code genotypic data as the number of minor alleles, i.e. 0, 1 and 2 for diploid species and impute missing genotypic data using function codeGeno.

- 2. Estimate reletedness based on marker data using functions vanRaden or rogers.
- 3. Set uo mixed models and estimate variance components and genomic breeding values using function regress.

Future work comprises mothods for phenotypic analysis and expanded usage, i.e. for animal breeding.

11 Acknowledgements

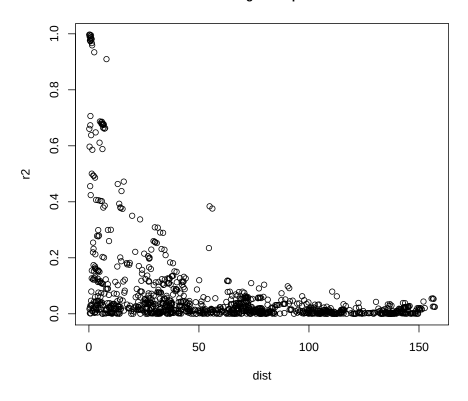
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References

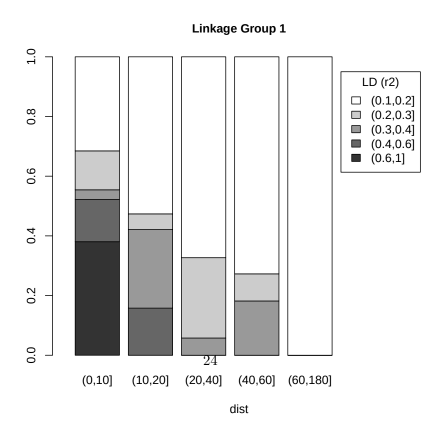
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Linkage Group 1



(a) Scatterplot using type="p"



(b) Stacked histogram using type="bars"

Figure 2: LD vs distance for first chromosome of maize data.

> LDMap(marker[, chr1], maize.marker.pos\$chr[chr1], maize.marker.pos\$pos[chr1])

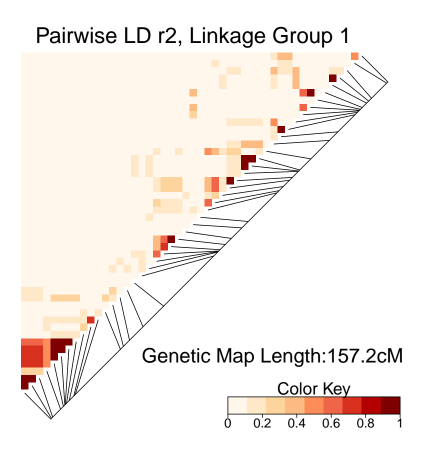
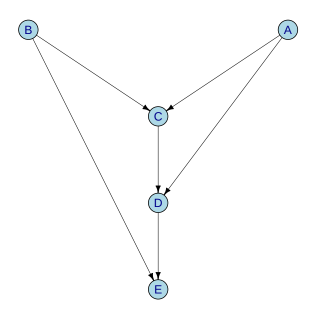


Figure 3: LD heatmap for markers on first chromosome of maize data.

> plot(ped)



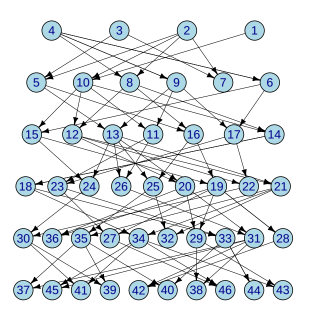


Figure 4: Simulated pedigree structure

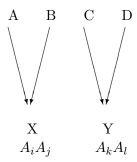


Figure 5: Pedigree of X and Y.