BreedR Overview

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Intro

What is breedR

- R-package implementing statistical models specifically suited for forest genetic resources analysts.
- Ultimately Mixed Models, but not necessarily easy to implement and use
- breedR acts as an interface which provides the means to:
 - 1. Combine any number of these models as components of a larger model
 - 2. Compute automatically incidence and covariance matrices from a few input parameters
 - 3. **Fit** the model
 - 4. Plot data and results, and perform model diagnostics

Installation

- Project web page http://famuvie.github.io/breedR/
 - Set up this URL as a package repository in .Rprofile (detailed instructions on the web)
 - install.packages('breedR')
 - Not possible to use CRAN due to closed-source BLUPF90 programs
- GitHub dev-site https://github.com/famuvie/breedR
 - if(!require(devtools)) install.packages('devtools')

- devtools::install_github('famuvie/breedR')

Where to find help

- Package's help: help(package = breedR)
 - Help pages ?remlf90
 - Code demos demo(topic, package = 'breedR') (omit topic for a list)
 - Vignettes vignette(package = 'breedR') (pkg and wiki)
- Wiki pages
 - Guides, tutorials, FAQ
- Mailing list http://groups.google.com/group/breedr
 - Questions and debates about usage and interface
- Issues page
 - Bug reports
 - Feature requests

License



Figure 1: GPL-3

- breedR is FOSS. Licensed GPL-3
 - RShowDoc('LICENSE', package = 'breedR')
- ullet You can **use** and **distribute breedR** for any purpose
- You can **modify** it to suit your needs
 - we encourage to!
 - please consider contributing your improvements
 - you can **distribute** your modified version under the GPL
- However, **breedR** makes (intensive) use of the BLUPF90 suite of Fortran programs, which are for *free* but not **free** (remember CRAN?)

Roadmap | Future developments

- Bayesian inference
- Multi-trait support
- $\bullet \ \ {\rm Genotype}{\times} {\rm Environment\ interaction}$
- Support for longitudinal data

Functionality

Inference

Frequentist

- Currently, only **frequentist inference** is supported via REML estimation of variance components.
- The function remlf90(), provides an interface to both REMLF90 and AIREMLF90 functions in the BLUPF90 suite of Fortran programs.
- Type ?remlf90 for details on the syntax

Bayesian

- It's on the roadmap for the next year
- Will use a gibbs sampler from BLUPF90, and possibly also INLA
- The interface will change a bit, separating the model specification from the fit

Linear Mixed Models with unstructured random effects

Example dataset

self	dad	mum	gen	gg	bl	phe_X	X	У	fam
69	0	64	1	14	13	15.756	0	0	64
70	0	41	1	4	13	11.141	3	0	41
71	0	56	1	14	13	19.258	6	0	56
72	0	55	1	14	13	4.775	9	0	55
73	0	22	1	8	13	19.099	12	0	22
74	0	50	1	14	13	19.258	15	0	50

```
1021 obs. of 10 variables:
   $ self : int 69 70 71 72 73 74 75 76 77 78 ...
   $ dad : int 000000004 ...
   $ mum : int 64 41 56 55 22 50 67 59 49 8 ...
   $ gen : Factor w/ 1 level "1": 1 1 1 1 1 1 1 1 1 1 ...
          : Factor w/ 14 levels "1","2","3","4",..: 14 4 14 14 8 14 14 14 14 11 ...
          : Factor w/ 15 levels "1","2","3","4",..: 13 13 13 13 13 13 13 13 9 9 ...
   $ phe X: num
                 15.76 11.14 19.26 4.78 19.1 ...
##
   $ x
          : int
                 0 3 6 9 12 15 18 21 24 27 ...
                 0 0 0 0 0 0 0 0 0 0 ...
          : int
   $ fam : Factor w/ 63 levels "6","7","8","9",...: 59 36 51 50 17 45 62 54 44 3 ...
```

A simple Provenance Test

Specify the genetic group gg as an unstructured random effect using the standard formulas in R

```
\begin{aligned} \text{phe}_X = & \mu + Z \text{gg} + \varepsilon \\ \text{gg} \sim & N(0, \sigma_{\text{gg}}^2) \\ & \varepsilon \sim & N(0, \sigma_{\varepsilon}^2) \end{aligned}
```

```
## Using default initial variances given by default_initial_variance()
## See ?breedR.getOption.
```

Initial variances specification

To avoid the notification, initial values for *all* the variance components must be made explicit using the argument var.ini:

Although in most cases the results will not change at all, we encourage to give explicit initial values for variance components. Specially when some estimate can be artifact. This is also useful for checking sensitivity to initial values.

Exploring the results

```
summary(res)
```

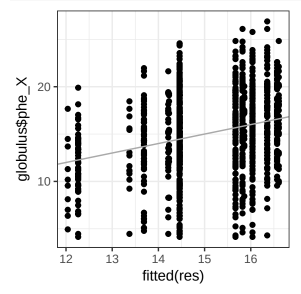
```
## Formula: phe_X ~ 0 + Intercept + gg
##
     Data: globulus
    AIC BIC logLik
##
##
  5864 5874 -2930
##
## Parameters of special components:
##
##
## Variance components:
##
          Estimated variances
                          2.857 1.3584
## gg
## Residual
                         17.695 0.7888
##
## Fixed effects:
              value
## Intercept 14.799 0.4911
```

- Note that AI-REML has been used by default.
- You can also specify method = 'em'.
- Learn about the difference.

Further extractor functions

```
fixef(res)
## $Intercept
##
       value
                   s.e.
## 1 14.79913 0.4910931
ranef(res)
## $gg
##
           value
                      s.e.
## 1 -1.1113031 0.6582245
## 2 -0.5850024 0.8241561
## 3
      1.2381743 0.6017957
## 4 -2.5360692 0.7047331
## 5
      1.0223492 0.6298409
     -2.7605955 1.0884704
## 6
     -0.5691183 0.9776411
## 7
     0.8700425 0.5933964
## 8
## 9
       1.5572484 0.6381498
## 10 -1.4262287 0.9961138
## 11 1.7715256 0.6527002
## 12 1.8079958 0.8241561
## 13 1.0604393 0.9776411
## 14 -0.3394577 0.5380184
```

Further extractor functions



```
str(resid(res))
   Named num [1:1021] 1.3 -1.12 4.8 -9.68 3.43 ...
   - attr(*, "names")= chr [1:1021] "1" "2" "3" "4" ...
extractAIC(res)
## [1] 5863.716
logLik(res)
## 'log Lik.' -2929.858 (df=2)
```

Hierarchical and Factorial models

- In globulus, the **family** (mum) is nested within the **provenance** (gg)
- This is a matter of codification:

Nested factors

	ľ
A 1	
A 2	
В 3	
B 4	

Crossed factors

gg	mum
A	1
A	2
В	1
В	2

Model specification

• Otherwise, in both cases we specify the model in the same way:

```
random = ~ gg + factor(mum) # note that mum is numeric
```

• Furthermore, this approach can handle unbalanced and mixed designs

Interactions

• Standard R notation:

```
random = ~ gg * factor(mum)
```

- Not available yet (feature request?)
- Workaround: build the interaction variable manually
- Example: gg and block are crossed factors

Exercise | Hierarchical and Factorial models

- 1. Use remlf90() and the globulus dataset to fit
 - a hierarchical model using mum within gg
 - a factorial model using gg and bl
- 2. Explore the results with summary()
 - is the family (mum) effect relevant?
 - is there any evidence of interaction between gg and bl?

Hierarchical and Factorial models #1 | Fitting models

Hierarchical and Factorial models #2 | Hierarchical model

- The family effect is not very **important**, in terms of explained variance
- However, the model is a bit better with it (AIC, logLik)

summary(res)

```
## Formula: phe_X ~ 0 + Intercept + gg
##
      Data: globulus
##
    AIC BIC logLik
##
   5864 5874 -2930
## Parameters of special components:
##
##
## Variance components:
##
           Estimated variances
                                  S.E.
## gg
                         2.857 1.3584
## Residual
                        17.695 0.7888
## Fixed effects:
              value
                    s.e.
## Intercept 14.799 0.4911
```

```
summary(res.h)
## Formula: phe_X ~ 0 + Intercept + factor(mum) + gg
##
      Data: globulus
     AIC BIC logLik
##
   5857 5872 -2926
##
## Parameters of special components:
##
##
## Variance components:
               Estimated variances
                            0.8955 0.4177
## factor(mum)
                           2.0540 1.1706
## gg
## Residual
                           17.0770 0.7819
##
## Fixed effects:
              value
## Intercept 14.973 0.4702
```

Hierarchical and Factorial models #3 | Factorial model

- Looks like the interaction between **block** and **provenance** is negligible
- (apart from the fact that it makes no sense at all, and shuld not have been even considered in the first place)
- compare with the model without interaction

```
summary(res.f)
```

```
## Formula: phe_X ~ 0 + Intercept + gg + bl + gg_bl
     Data: globulus.f
##
##
    AIC BIC logLik
##
   5752 5772 -2872
## Parameters of special components:
##
##
## Variance components:
          Estimated variances S.E.
##
## gg
                       3.10970 1.4329
## bl
                       2.57280 1.0606
                       0.02912 0.2713
## gg_bl
## Residual
                       15.19800 0.7159
##
## Fixed effects:
             value s.e.
## Intercept 14.764 0.653
## result without interaction
res.f0 <- remlf90(fixed = phe_X ~ 1,
                  random = ~gg + bl,
                  data = globulus)
paste('AIC:', round(extractAIC(res.f0)),
     'logLik:', round(logLik(res.f0)))
```

Additive Genetic Effect

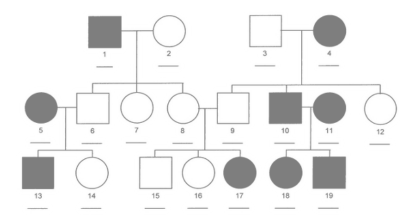


Figure 2: pedigree

What is an additive genetic effect

- Random effect at individual level
- Based on a **pedigree**
- BLUP of Breeding Values from own and relatives' phenotypes
- Represents the additive component of the genetic value
- More general:
- family effect is a particular case
- accounts for more than one generation
- mixed relationships
- More flexible: allows to select individuals within families

Specifying a pedigree

- A 3-column data.frame or matrix with the codes for each individual and its parents
- A **family** effect is easily translated into a pedigree:
 - use the **family code** as the identification of a fictitious **mother**
 - use 0 or NA as codes for the unknown fathers

self	dad	mum
69	0	64
70	0	41
71	0	56
72	0	55
73	0	22
74	0	50

Fitting an animal model

Animal model: results

- gg explains almost the same amount of phenotypic variability
- The (additive) genetic effect explains part of the formerly residual variance
- The **heritability** is computed automatically as

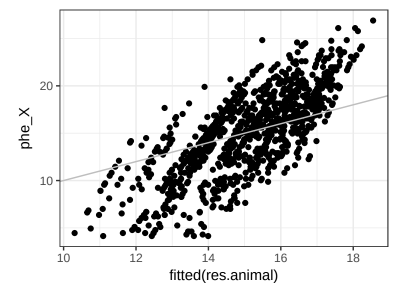
$$h^2 = \frac{\sigma_a^2}{\sigma_a^2 + \sigma_{qq}^2 + \sigma^2}$$

```
summary(res.animal)
```

```
## Formula: phe_X ~ 0 + Intercept + gg + pedigree
##
      Data: globulus
##
    AIC BIC logLik
   5857 5872 -2926
##
##
## Parameters of special components:
##
##
## Variance components:
          Estimated variances S.E.
##
## gg
                          2.356 1.249
## genetic
                          3.632 1.649
## Residual
                         14.271 1.561
##
##
               Estimate
                            S.E.
## Heritability 0.1795 0.08253
##
## Fixed effects:
##
              value s.e.
## Intercept 14.797 0.47
```

Extracting Predicted Breeding Values

```
## Predicted Breeding Values
# for the full pedigree first, and for the observed individuals
# by matrix multiplication with the incidence matrix
PBV.full <- ranef(res.animal)$genetic
PBV <- model.matrix(res.animal)$genetic %*% PBV.full
# Predicted genetic values vs.</pre>
```



Handling pedigrees

- The pedigree needs to meet certain conditions
- If it does not, breedR automatically completes, recodes and sorts
- If recoding is necessary, breedR issues a warning because you need to be careful when retrieving results
- See this guide for more details

Spatial autocorrelation

What is spatial autocorrelation

- The **residuals** of any LMM must be **noise**
- However, most times there are environmental factors that affect the response
- This causes that observations that are close to each other **tend** to be more similar that observations that are far away
- This is called spatial autocorrelation
- It may affect both the estimations and their accuracy
- This is why experiments are randomized into spatial ${\bf blocks}$

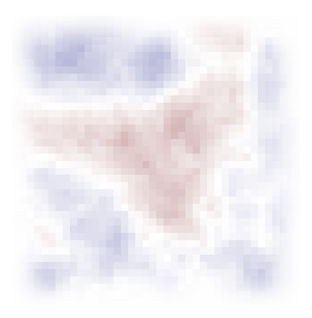
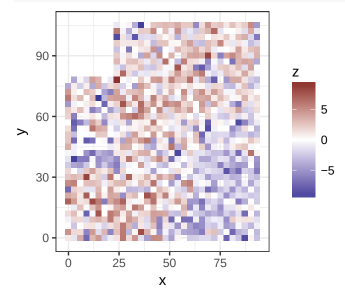


Figure 3: spatial

Diagnosing spatial autocorrelation | residuals spatial plot

- You can plot() the spatial arrangement of various model components (e.g. residuals)
- Look like **independent** gaussian observations (i.e. noise)?
- Do you see any **signal** in the background?

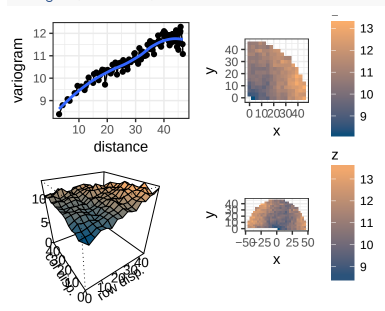
```
## Since coordinates have not
## been passed before they
## must be provided explicitly.
coordinates(res.animal) <-
    globulus[, c('x', 'y')]
plot(res.animal, 'resid')</pre>
```



Diagnosing spatial autocorrelation | variograms of residuals

• Plot the variogram of residuals with variogram()

variogram(res.animal)

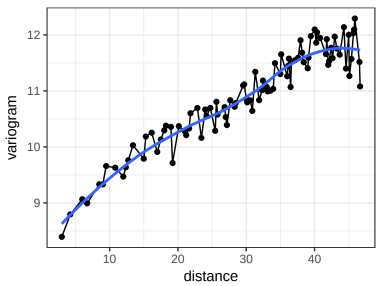


Interpreting the variograms

• Isotropic variogram:

$$\gamma(h) = \frac{1}{2}V[Z(\mathbf{u}) - Z(\mathbf{v})], \quad \text{dist}(\mathbf{u}, \mathbf{v}) = h$$

The **empirical** isotropic variogram is built by aggregating all the pairs of points separated by h, no matter the direction.

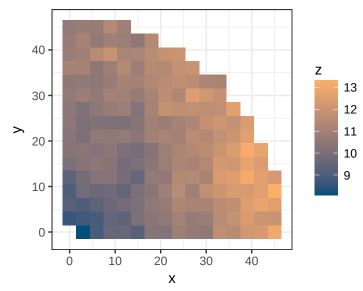


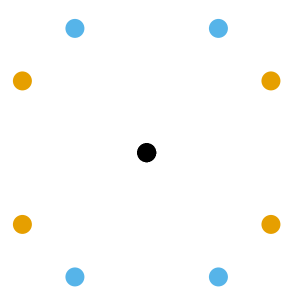
Interpreting the variograms

• Row/Column variogram:

$$\gamma(x,y) = \frac{1}{2}V[Z(\mathbf{u}) - Z(\mathbf{v})], \text{ dist}(\mathbf{u}, \mathbf{v}) = (x, y)$$

The **empirical** row/col variogram is built by aggregating **all the pairs** of points separated by exactly x rows and y columns.



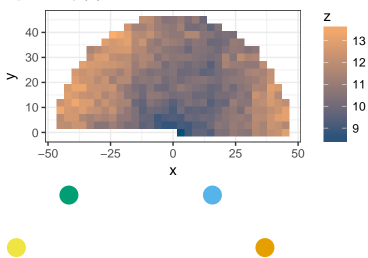


Interpreting the variograms

• Anisotropic variogram:

$$\gamma(\mathbf{x}) = \frac{1}{2}V[Z(\mathbf{u}) - Z(\mathbf{v})], \quad \mathbf{u} = \mathbf{v} \pm \mathbf{x}$$

The **empirical** anisotropic variogram is built by aggregating **all the pairs** of points **in the same direction** separated by $|\mathbf{x}|$.



Accounting for spatial autocorrelation

- Include an explicit spatial effect in the model
- I.e., a **random effect** with a specific covariance structure that reflects the spatial relationship between individuals
- The **block** effect, is a very particular case:
 - It is designed from the begining, possibly using prior knowledge
 - Introduces **independent** effects between blocks
 - Most neighbours are within the same block (i.e. share the same effect)

The blocks model

- The blocks spatial model is equivalent to random = ~ bl, but:
 - specifying coord is convenient for plotting (remember?)
 - blocks behaves as expected, even if bl is not a factor

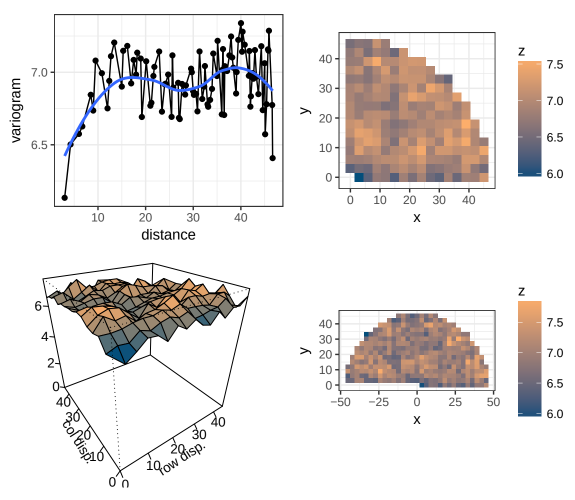
Animal-spatial model: results

```
summary(res.blk)
## Formula: phe_X ~ 0 + Intercept + gg + pedigree + spatial
     Data: globulus
##
##
    AIC BIC logLik
##
   5734 5753 -2863
##
## Parameters of special components:
## spatial: n.blocks: 15
##
## Variance components:
##
           Estimated variances S.E.
## gg
                          2.385 1.274
                          5.275 1.836
## genetic
## spatial
                          2.650 1.081
## Residual
                         10.279 1.601
##
##
                Estimate
                            S.E.
## Heritability 0.2556 0.08989
```

```
##
## Fixed effects:
## value s.e.
## Intercept 14.762 0.6342
```

• Now the additive-genetic variance and the heritability have increased! (3.6 and 0.18 before)

Variogram of residuals



• There seems to remain some intra-block spatial autocorrelation

B-Splines model

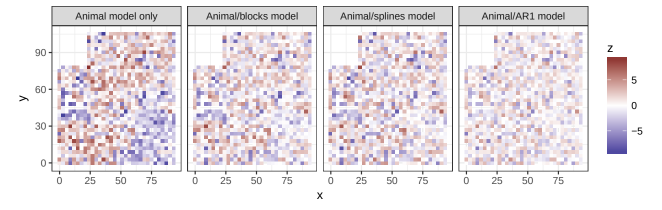
- A continuous and smooth spatial surface built from a linear combination of basis functions
- The coefficients are modelled as a random effect

Autoregressive model

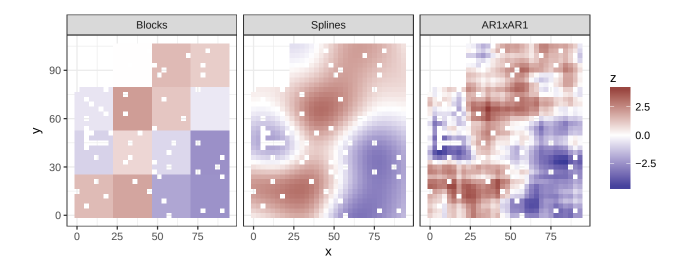
• A separable kronecker product of First order Autoregressive processes on the rows and the colums

Change in model residuals

• We preserve the scale by using compare.plots()

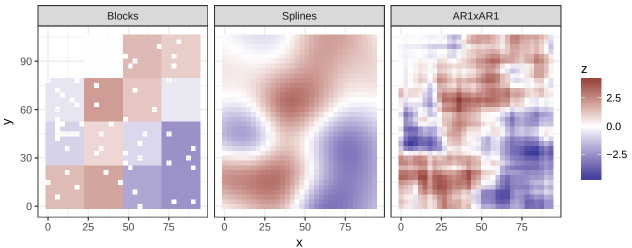


Comparison of spatial components



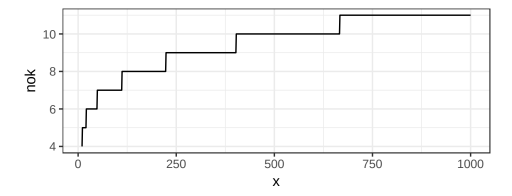
Prediction of the spatial effect in unobserved locations

- The type fullspatial fills the holes (when possible)
- See ?plot.remlf90



Spatial parameters | Number of knots of a splines model

- The smoothness of the spatial surface can be controlled modifying the number of base functions
- This is, directly determined by the **number of knots** (nok) in each dimension
- n.knots can be used as an additional argument in the spatial effect as a numeric vector of size 2.
- Otherwise, is determined by the function given in breedR.getOption('splines.nok')



Spatial parameters | Autoregressive parameters of a AR model

- Analogously, the patchiness of the AR effects can be controlled by the autoregressive parameter for each dimension
- rho can be given as an additional argument in the spatial effect as a numeric vector of size 2
- By default, breedR runs all the combinations in the grid produced by the values from breedR.getOption('ar.eval') and returns the one with largest likelihood
- It returns also the full table of combinations and likelihoods in res\$rho

Exercise | Tuning spatial parameters

- Tuning parameters:
 - model splines: n.knots
 - model AR: rho
- 1. Increase the number of knots in the splines model and see if it improves the fit
- 2. Visualize the log-likelihood of the fitted AR models
- 3. Refine the grid around the most likely values, and refit using rho = rho.grid, where

- What are now the most likely parameters?

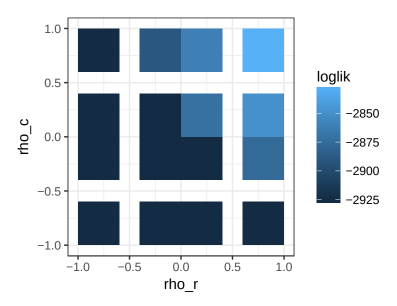
Spatial #1 | B-splines model with increased nok

• nok were (6, 6) by default (see summary())

```
## Formula: phe_X ~ 0 + Intercept + gg + pedigree + spatial
##
      Data: globulus
            BIC logLik
     AIC
##
## 5685 unknown -2838
## Parameters of special components:
## spatial: n.knots: 12 12
## Variance components:
##
        Estimated variances
## gg
                          2.568
                          4.498
## genetic
## spatial
                          4.199
                         10.070
## Residual
##
## Fixed effects:
##
              value
## Intercept 14.479 0.9163
summary(res.spl99)
## Formula: phe_X ~ 0 + Intercept + gg + pedigree + spatial
      Data: globulus
##
     AIC
            BIC logLik
## 5681 unknown -2836
##
## Parameters of special components:
## spatial: n.knots: 15 15
##
## Variance components:
##
          Estimated variances
## gg
                         2.509
## genetic
                         4.651
## spatial
                          3.490
## Residual
                          9.552
##
## Fixed effects:
              value
                     s.e.
## Intercept 14.611 0.6947
```

Spatial #2 | Visualize log-likelihoods

```
qplot(rho_r, rho_c,
    fill = loglik,
    geom = 'tile',
    data = res.ar1$rho)
```



rho_r	rho_c	loglik
-0.8	-0.8	-2925.648
-0.2	-0.8	-2925.647
0.2	-0.8	-2925.645
0.8	-0.8	-2925.636
-0.8	-0.2	-2925.647
-0.2	-0.2	-2925.645
0.2	-0.2	-2925.023
0.8	-0.2	-2876.893
-0.8	0.2	-2925.645
-0.2	0.2	-2925.645
0.2	0.2	-2871.691
0.8	0.2	-2849.814
-0.8	0.8	-2925.645
-0.2	0.8	-2890.606
0.2	0.8	-2860.981
0.8	0.8	-2828.017

Spatial #3 | Refine grid

Formula: phe_X ~ 0 + gg + pedigree + spatial

```
##
     Data: globulus
##
     AIC BIC logLik
   5603 5617 -2798
##
##
## Parameters of special components:
## spatial: rho: 0.8666667 0.7833333
## Variance components:
            Estimated variances S.E.
##
## genetic
                          5.090 1.715
## spatial
                          4.984 1.053
## Residual
                          7.583 1.499
##
##
                Estimate
                            S.E.
                  0.2878 0.09383
## Heritability
##
## Fixed effects:
         value
                  s.e.
## gg.1 13.351 0.7195
## gg.2 14.331 0.9112
## gg.3 15.945 0.7698
## gg.4 11.585 0.9394
## gg.5
        15.913 0.8200
         9.593 1.6964
## gg.6
        13.761 1.5681
## gg.7
## gg.8 15.521 0.7486
        16.302 0.8260
## gg.9
## gg.10 12.684 1.1531
## gg.11 16.459 0.9849
## gg.12 16.801 1.1412
## gg.13 15.783 1.5665
## gg.14 14.211 0.6486
```

Competition

Theoretical model

- Each individual have two (unknown) Breeding Values (BV)
- The direct BV affects its **own** phenotype, while the competition BV affects its **neghbours**' (as the King moves)
- The effect of the neighbouring competition BVs is given by their sum weighted by $1/d^{\alpha}$, where α is a tuning parameter called decay
- Each set of BVs is modelled as a zero-mean random effect with structure matrix given by the **pedigree** and independent variances σ_a^2 and σ_c^2
- Both random effects are modelled jointly with correlation ρ

Permanent Environmental Effect (pec)

• Optional effect with environmental (rather than genetic) basis

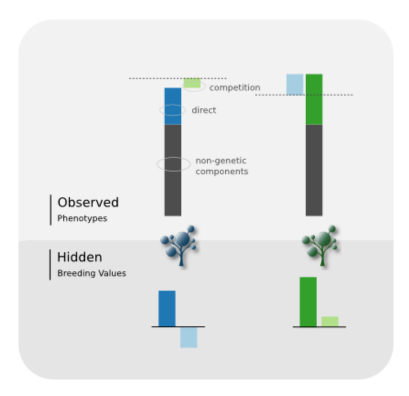


Figure 4: Competition model

• Modelled as an individual **independent** random effect that affects **neighbouring** trees in the same (weighted) way

Simulation of data

 \mathbf{breedR} implements a convenient dataset $\mathbf{simulator}$ which keeps a similar syntax.

• See ?simulation for details on the syntax

```
# Simulation parameters
grid.size <- c(x=20, y=25) # cols/rows
coord <- expand.grid(sapply(grid.size,</pre>
                              seq))
Nobs <- prod(grid.size)</pre>
Nparents \leftarrow c(mum = 20, dad = 20)
sigma2_a <- 2 # direct add-gen var
sigma2_c <- 1 # compet add-gen var
        <- -.7 # gen corr dire-comp
sigma2_s <- 1 # spatial variance</pre>
sigma2_p <- .5 # pec variance</pre>
sigma2 <- .5 # residual variance
S <- matrix(c(sigma2_a,</pre>
               rho*sqrt(sigma2_a*sigma2_c),
               rho*sqrt(sigma2_a*sigma2_c),
               sigma2_c),
             2, 2)
```

```
set.seed(12345)
simdat <-
  breedR.sample.phenotype(
   fixed = c(beta = 10),
   genetic = list(model = 'competition',
                   Nparents = Nparents,
                   sigma2_a = S,
                   check.factorial=FALSE,
                   pec = sigma2_p),
   spatial = list(model = 'AR',
                   grid.size = grid.size,
                   rho = c(.3, .8),
                   sigma2_s = sigma2_s),
   residual.variance = sigma2
## Remove founders
dat <- subset(simdat,</pre>
              !(is.na(simdat$sire)
                & is.na(simdat$dam)))
```

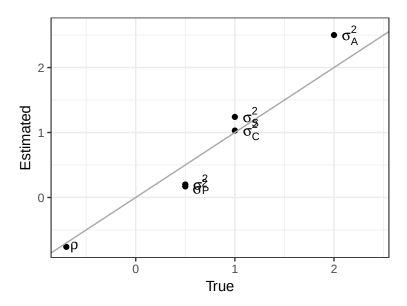
Fitting a competition model

81.045

True vs. estimated parameters

0.296 81.337

	True	Estimated
direct	2.0	2.50
compet.	1.0	1.03
correl.	-0.7	-0.76
spatial	1.0	1.24
pec	0.5	0.20
residual	0.5	0.17



Exercise | Competition models

- 1. Plot the true vs predicted:
 - direct and competition Breeding Values
 - spatial effects
 - pec effects
- 2. Plot the residuals and their variogram
 - Do you think the residuals are independent?
 - How would you improve the analysis?

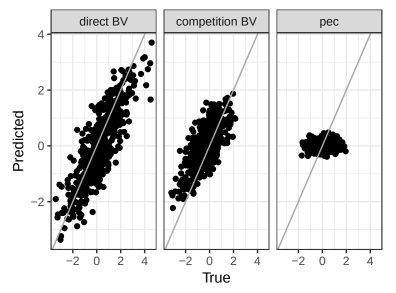
Competition #1 | True vs. predicted components

```
## compute the predicted effects for the observations
## by matrix multiplication of the incidence matrix and the BLUPs
pred <- list()
Zd <- model.matrix(res.comp)$'genetic_direct'
pred$direct <- Zd %*% ranef(res.comp)$'genetic_direct'

## Watch out! for the competition effects you need to use the incidence
## matrix of the direct genetic effect, to get their own value.
## Otherwise, you get the predicted effect of the neighbours on each
## individual.
pred$comp <- Zd %*% ranef(res.comp)$'genetic_competition'
pred$pec <- model.matrix(res.comp)$pec %*% ranef(res.comp)$pec</pre>
```

Competition #1 | True vs. predicted components

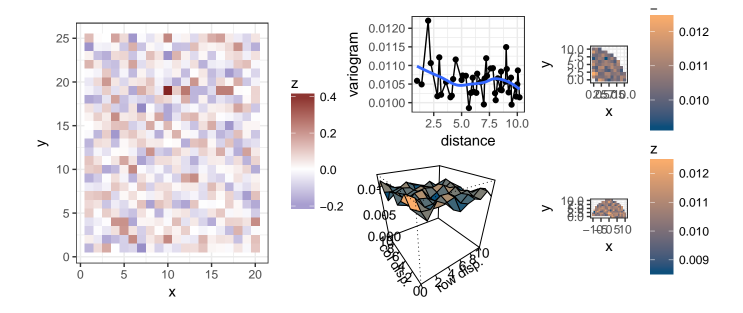
```
comp.pred <-
rbind(
  data.frame(
    Component = 'direct BV',
    True = dat$BV1,</pre>
```



The predition of the Permanent Environmental Competition effect is not precisely great...

Competition $#2 \mid$ Map of residuals and their variogram

```
plot(res.comp, type = 'resid')
variogram(res.comp)
```



Generic component

The Generic model

This additional component allows to introduce a random effect ψ with **arbitrary** incidence and covariance matrices Z and Σ :

$$y = \mu + X\beta + Z\psi + \varepsilon$$
$$\psi \sim N(0, \sigma_{\psi}^{2} \Sigma_{\psi})$$
$$\varepsilon \sim N(0, \sigma_{\varepsilon}^{2})$$

Implementation of the generic component

Example of result

```
## Formula: phe_X ~ 0 + gg
## Data: globulus
## AIC BIC logLik
## 5691 5701 -2844
##
## Parameters of special components:
##
```

```
##
## Variance components:
##
          Estimated variances
                        2.592 1.0640
## block
## Residual
                        15.208 0.6825
##
## Fixed effects:
##
         value
                  s.e.
## gg.1 13.534 0.6222
## gg.2 14.030 0.8464
## gg.3 16.106 0.5513
## gg.4 11.854 0.6824
## gg.5 15.883 0.5863
## gg.6 10.220 1.3041
## gg.7 13.995 1.0894
## gg.8 15.728 0.5410
## gg.9 16.478 0.5969
## gg.10 12.843 1.1225
## gg.11 16.744 0.6151
## gg.12 17.002 0.8464
## gg.13 16.297 1.0894
## gg.14 14.429 0.4730
```

Prediction

Predicting values for unobserved trees

- You can predict the Breeding Value of an unmeasured tree
- Or the expected phenotype of a death tree (or an hypothetical scenario)
- Information is gathered from the covariates, the spatial structure and the pedigree
- Simply include the individual in the dataset with the response set as NA

Leave-one-out cross-validation

- Re-fit the simulated competition data with one measurement removed
- Afterwards, compare the predicted values for the **unmeasured** individuals with their true simulated values

	True	Pred.loo
direct BV	-1.48	0.11
competition BV	0.46	0.36
exp. phenotype	6.80	9.90

Exercise | Cross validation

- 1. Extend the last table to include the predicted values with the full dataset
- 2. Remove 1/10th of the phenotypes randomly, and predict their expected phenotype
 - Have the parameter estimations changed too much?
- 3. Compute the Root Mean Square Error (RMSE) of Prediction with respect to the true values

Cross-validation #1 | Include prediction with full data

	True	Pred.full	Pred.loo
direct BV	-1.48	-1.30	0.11
competition BV	0.46	0.99	0.36
exp. phenotype	6.80	7.29	9.90

Cross-validation #2 | Perform cross-validation on 1/10th of the observations

```
rm.idx <- sample(nrow(dat), nrow(dat)/10)
dat.cv <- dat
dat.cv[rm.idx, 'phenotype'] <- NA
## Re-fit the model and build table</pre>
```

	Fully.estimated	CV.estimated
direct	2.50	2.69
compet.	1.03	0.86
correl.	-0.76	-0.80
spatial	1.24	1.03
pec	0.20	0.51
residual	0.17	0.14

Cross-validation #3 | MSE of Prediction

```
true.exp.cv <- with(dat[rm.idx, ], phenotype - resid)
round(sqrt(mean((fitted(res.comp.cv)[rm.idx] - true.exp.cv)^2)), 2)
## [1] 1.5</pre>
```

Multiple traits

breedR provides a basic interface for multi-trait models which only requires specifying the different traits in the main formula using cbind().

```
## Filter site and select relevant variables
dat <-
  droplevels(
    douglas[douglas$site == "s3",
            names(douglas)[!grepl("HO[^4]|AN|BR|site", names(douglas))]]
  )
res <-
  remlf90(
    fixed = cbind(H04, C13) ~ orig,
    # random = ~ block,
    genetic = list(
      model = 'add_animal',
      pedigree = dat[, 1:3],
      id = 'self'),
    data = dat
  )
## Warning in build_pedigree(1:3, data = ped.df): The pedigree has been
## recoded. Check attr(ped, 'map').
## Using default initial variances given by default_initial_variance()
## See ?breedR.getOption.
```

A full covariance matrix across traits is estimated for each random effect, and all results, including heritabilities, are expressed effect-wise:

```
## Formula: cbind(H04, C13) ~ 0 + orig + pedigree
##
     Data: dat
##
      AIC
           BIC logLik
##
   30968 31010 -15476
## Parameters of special components:
##
##
## Variance components:
                                         Estimated variances
##
                                                              S.E.
## genetic.direct.H04
                                                       918.1 438.6
## genetic.direct.HO4_genetic.direct.C13
                                                      1872.4 824.0
## genetic.direct.C13
                                                      5827.6 1829.6
## Residual.H04
                                                      8373.7 461.7
## Residual.H04_Residual.C13
                                                     10922.0 755.3
## Residual.C13
                                                     18439.0 1484.2
##
##
                                S.E.
                    Estimate
## Heritability:H04 0.0990 0.04589
## Heritability:C13
                      0.2391 0.07036
##
## Fixed effects:
                value
                         s.e.
## orig.H04.pA 352.00 6.2389
```

```
## orig.H04.pB 370.90 10.7947
## orig.H04.pC 346.93 13.0788
## orig.H04.pF 339.66 6.2268
## orig.H04.pG 313.00 24.0430
## orig.H04.pH 305.39 19.9334
## orig.H04.pI 323.29 20.0946
## orig.H04.pJ 343.87 19.8567
## orig.H04.pK 335.48 19.6409
## orig.C13.pA 460.01 13.6444
## orig.C13.pB 494.58 19.8635
## orig.C13.pC 430.86 25.5477
## orig.C13.pF 429.48 12.5501
## orig.C13.pG 376.42 48.3133
## orig.C13.pH 376.98 43.4266
## orig.C13.pI 404.62 43.6194
## orig.C13.pJ 418.91 43.2856
## orig.C13.pK 441.99 43.0567
Although the results are summarized in tabular form, the covariance matrices can be recovered directly:
res$var[["genetic", "Estimated variances"]]
              direct.H04 direct.C13
## direct.H04
                  918.08
                              1872.4
                  1872.40
                              5827.6
## direct.C13
## Use cov2cor() to compute correlations
cov2cor(res$var[["genetic", "Estimated variances"]])
##
              direct.H04 direct.C13
## direct.H04
              1.0000000 0.8094938
## direct.C13 0.8094938 1.0000000
Estimates of fixed effects and BLUPs of random effects can be recovered with fixef() and ranef() as usual.
The only difference is that they will return a list of matrices rather than vectors, with one column per trait.
The standard errors are given as attributes, and are displayed in tabular form whenever the object is printed.
fixef(res)
                  ## printed in tabular form, but...
## $orig
##
      value.H04 value.C13 s.e..H04 s.e..C13
## pA
                 460.0097
                            6.238914 13.64437
       352.0025
## pB
       370.8997
                 494.5846 10.794693 19.86351
## pC
       346.9318
                 430.8644 13.078774 25.54773
## pF
       339.6614
                  429.4795 6.226796 12.55013
## pG
       313.0000
                 376.4231 24.043034 48.31334
                  376.9779 19.933367 43.42664
## pH
       305.3889
## pI
       323.2885
                  404.6216 20.094619 43.61939
                  418.9064 19.856683 43.28562
## pJ
       343.8727
                 441.9861 19.640911 43.05671
## pK 335.4828
unclass(fixef(res)) ## actually a matrix of estimates with attribute "se"
## $orig
##
           H04
                     C13
## pA 352.0025 460.0097
```

pB 370.8997 494.5846

```
## pC 346.9318 430.8644
## pF 339.6614 429.4795
## pG 313.0000 376.4231
## pH 305.3889 376.9779
## pI 323.2885 404.6216
## pJ 343.8727 418.9064
## pK 335.4828 441.9861
## attr(, "se")
##
            H04
                     C13
## pA 6.238914 13.64437
## pB 10.794693 19.86351
## pC 13.078774 25.54773
## pF 6.226796 12.55013
## pG 24.043034 48.31334
## pH 19.933367 43.42664
## pI 20.094619 43.61939
## pJ 19.856683 43.28562
## pK 19.640911 43.05671
str(ranef(res))
## List of 1
   $ genetic: num [1:1525, 1:2] -6.02 -12.93 -10.16 33.51 6.77 ...
##
##
     ..- attr(*, "dimnames")=List of 2
     ....$ : chr [1:1525] "19" "21" "23" "25" ...
##
##
     ....$ : chr [1:2] "H04" "C13"
     ..- attr(*, "se")= num [1:1525, 1:2] 23 22.8 23.6 23 23.6 ...
##
##
     ...- attr(*, "dimnames")=List of 2
     ....$ : chr [1:1525] "19" "21" "23" "25" ...
##
##
     ....$ : chr [1:2] "H04" "C13"
##
     ..- attr(*, "names")= chr [1:3050] "1" "2" "3" "4" ...
   - attr(*, "class")= chr [1:2] "ranef.breedR" "breedR_estimates"
head(ranef(res)$genetic)
##
             H04
                         C13
## 19 -6.016271
                   40.093547
## 21 -12.925035 -108.673107
## 23 -10.164449
                   23.276658
## 25
       33.507715
                   80.855347
## 27
        6.768289
                   -5.018311
      22.201575
## 29
                   32.078520
```

Recovering the breeding values for each observation in the original dataset follows the same procedure as for one trait: multiply the incidence matrix by the BLUP matrix. The result, however, will be a matrix with one column per trait.

head(model.matrix(res)\$genetic %*% ranef(res)\$genetic)

```
## H04 C13
## 151 5.923689 -6.612036
## 153 7.760706 22.486000
## 155 -7.414378 -38.978615
## 157 7.894009 3.756494
## 159 3.536361 -10.654445
## 161 12.431919 12.736590
```

Some more features

Metagene interface

- We have used simulated data from the metagene software
- If you simulate data, import the results with read.metagene()
- Use several common methods with a metagene object:

```
- summary(), plot(), as.data.frame()
```

- Plus some more specific metagene functions:
 - b.values(), get.ntraits(), ngenerations(), nindividuals(), get.pedigree()
- And specific functions about spatial arrangement:
 - coordinates() extract coordinates
 - sim.spatial() simulates some spatial autocorrelation

Simulation framework

- The function breedR.sample.phenotype() simulates datasets from all the model structures available in breedR
- Limitation: only one generation, with random matings of founders
- See ?simulation for details

Remote computation

If you have access to a Linux server through SSH, you can perform computations remotely

- Take advantage of more **memory** or **faster** processors
- Parallelize jobs
- Free local resources while fitting models
- See ?remote for details

Package options

- breedR features a list of configurable options
- Use breedR.setOption(...) for changing an option during the current sesion
- Set options permanently in the file \$HOME/.breedRrc
- see ?breedR.option for details

breedR.getOption()

```
## $ar.eval
## [1] -0.8 -0.2 0.2 0.8
##
## $breedR.bin
## [1] "/home/facu/Work/Proyectos/2013.T4F/bin/PROGSF90/linux/32bit"
##
```

```
## $splines.nok
## determine.n.knots
## $default.initial.variance
## default_initial_variance
##
## $col.seq
## [1] "#034E7B" "#FDAE6B"
##
## $col.div
## [1] "#3A3A98FF" "#832424FF"
## $cygwin
## [1] "C:/cygwin"
##
## $cygwin.home
## [1] "/home/facu"
##
## $ssh.auth.sock
## [1] "/tmp/ssh-auth-sock-facu"
##
## $remote.host
## [1] "elnuevo"
## $remote.user
## [1] "fmunoz"
##
## $remote.port
## [1] 22
##
## $remote.bin
## [1] "/usr/local/lib/R/site-library/breedR/bin/linux"
##
## $ssh.options
## [1] "-x -o BatchMode=yes -o TCPKeepAlive=yes -e none"
```