Handling pedigrees

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Contents

What is 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
- A pedigree sintetizes any kind of (genetic) relationship between individuals from one or more generations

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

Checking pedigrees

- For computational reasons, the pedigree needs to meet certain conditions:
 - Completness: all the individuals (also parents) must have an entry
 - * with possibly unknown parents (code 0 or NA)
 - The offspring must follow the parents
 - The codes must be sorted increasingly
 - The codes must be consecutive
- So, not every 3-column data.frame or matrix with codes is a proper pedigree:

```
## full_ped offsp_follows codes_sorted codes_consecutive
## FALSE FALSE FALSE FALSE
```

Building pedigrees

[24] 13 15 14 11 NA 12 9

- breedR implements a pedigree constructor that completes, sorts and recodes as necessary
- The resulting object, of class pedigree is guranteed to meet the conditions

self	sire	dam
9	2	20
23	13	30
12	21	22
24	27	29
25	10	26

self	sire	dam
1	NA	NA
2	NA	NA
3	1	2
4	NA	NA
5	NA	NA
6	NA	NA
7	5	6
8	NA	NA
9	NA	NA
10	8	9
11	NA	NA
12	NA	NA
13	11	12
14	NA	NA
15	4	14

Using a pedigree in an additive genetic effect

• just include your original pedigree information and let breedR fix it for you

```
test.dat <- data.frame(ped.nightmare, y = rnorm(n.ped))</pre>
res.raw <- remlf90(fixed = y ~ 1,
                   genetic = list(model = 'add_animal',
                                  pedigree = ped.nightmare,
                                # pedigree = test.dat[, 1:3], # same thing
                                  var.ini = 1,
                                  id = 'self'),
                   var.ini = list(resid = 1),
                   data
                         = test.dat)
## Warning in build_pedigree(1:3, data = ped.df): The pedigree has been
## recoded. Check attr(ped, 'map').
## No specification of initial variances.
        Using default value of 1 for all variance components.
##
##
        See ?breedR.getOption.
## pedigree has been recoded!
length(ranef(res.raw)$genetic)
## [1] 15
## The pedigree used in the model matches the one manually built
identical(ped.fix, get_pedigree(res.raw))
## [1] TRUE
```

Recovering Breeding Values in the original coding

	value	s.e.
9	0.21	0.89
23	-1.29	0.89
12	0.42	0.89
24	-0.18	0.89
25	0.84	0.89