Computing Inbreeding and Additive Relationships using the pedigreeTools R-package

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```
knitr::opts_chunk$set(echo = TRUE)
```

The Pedigree Tools R-package (available at Github) can be used to edit, sort and handle complete pedigrees.

Installing pedigreeTools from GitHub

```
remotes:::install_github('https://github.com/Rpedigree/pedigreeTools/') #3# install pedigre
```

Creating a pedigree object

A pedigree consist of the indivudal ID as well as the IDs of the father/mother (refered ast to sire and dam in animal breeding).

```
## sire dam
## 1 <NA> <NA>
## 2 1 <NA>
## 3 2 <NA>
## 4 2 3
## 5 2 <NA>
```

All the functions that ork on pedigrees require the pedigree to be complete. A complete pedigree is one where each sire and dam also appear as a label. Most often, the pedigrees that we have are not complete. To complete it, and to get the generation number you can use the editPed() function. This is illustrated in the following script.

```
PED=PED[-1,] # keeping in the pedigree only the individuals with at least one known parent (this is typrint(PED)
```

```
## 3 3
          2 <NA>
## 4 4
          2
## 5 5
          2 <NA>
# This fails because the pedigree is incomplete
 # ped <- with(PED, pedigree(label=id, sire=sire, dam=dam))</pre>
PED=editPed(label=PED$id,sire=PED$sire,dam=PED$dam)
print(PED)
    label sire dam generation
## 1
        1 <NA> <NA>
             1 <NA>
## 2
        2
                              1
                              2
## 3
        3 2 <NA>
## 5
        5
           2 <NA>
                              2
## 4
                              3
class(PED)
## [1] "data.frame"
# Now we use the complete pedigree to createa a 'pedigree object'
PED=pedigree(label=PED$label, sire=PED$sire, dam=PED$dam)
class(PED)
## [1] "pedigree"
## attr(,"package")
## [1] "pedigreeTools"
Computing inbreeding and additive relationships
F=inbreeding(PED)
## [1] 0.00 0.00 0.00 0.00 0.25
A=as.matrix(getA(PED)) # A has 1+F in the diagonal, the offdiagonals are 2*coancestry
A # additive relationship matrix
##
         1
              2
                  3
                         5
## 1 1.000 0.50 0.25 0.250 0.375
## 2 0.500 1.00 0.50 0.500 0.750
## 3 0.250 0.50 1.00 0.250 0.750
## 5 0.250 0.50 0.25 1.000 0.375
## 4 0.375 0.75 0.75 0.375 1.250
A/2 # co-ancestry
## 1 0.5000 0.250 0.125 0.1250 0.1875
## 2 0.2500 0.500 0.250 0.2500 0.3750
## 3 0.1250 0.250 0.500 0.1250 0.3750
```

id sire dam

1 <NA>

2 2

```
## 5 0.1250 0.250 0.125 0.5000 0.1875
## 4 0.1875 0.375 0.375 0.1875 0.6250
```

cbind(round(diag(A)-1,4),F) # Inbreeding

```
## F
## 1 0.00 0.00
## 2 0.00 0.00
## 3 0.00 0.00
## 5 0.00 0.00
## 4 0.25 0.25
```

heatmap(A,Rowv = FALSE,Colv=F)

