# APIS: An Auto-Adaptive Parentage Inference Software tolerant to missing parents

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## Description

This package include all the functions to assign with APIS (Griot & al., 2019). Parentage assignment is widely used for farmed and natural populations. As most of the likelihood software are based on simulation, the estimation of the simulation parameters is a key point for assignment reliability. Among those parameters, the proportion of missing parent is one of the most important. To avoid estimation of missing parents, we developed APIS (Auto-Adaptive Parentage Inference Software), based on observed average Mendelian transmission probabilities. In this package, you will find all the functions to perform parentage assign with the method described in the paper.

## Install and load the package

```
devtools::install_github("rgriot/APIS")
library(APIS)
```

If an error message occurs during the installation, use the following command :

```
devtools::install_github("rgriot/APIS", args = "--no-multiarch")
library(APIS)
```

## Formate your data

APIS requires matrices of characters as inputs. Each matrix has individuals as rows, markers as columns. The individual labels are set as rownames. Each cell is the genotype of one marker, coded "All1/All2". For example "A/A", "A/B", "B/B" for bi-allelic markers and "NA/NA" for missing value. For multi-allelic markers, use the generic coding "All1/All2".

```
data("genotype_APIS")
head(off_full[,1:10])
#> V2 V3
                 V4
                      V5
                         V6
                                V7
                                      V8
                                            V9
                                                 V10
#> off_1 "A/B" "B/B" "A/A" "A/A" "B/B" "B/B" "NA/NA" "A/A" "A/A" "A/B"
#> off_2 "A/B" "B/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/A" "A/B"
                                           "A/A" "A/A" "A/A"
#> off_3 "A/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B"
#> off_4 "B/B" "B/B" "A/A" "A/A" "B/B" "B/B" "B/B" "A/A" "A/A" "A/A"
#> off_5 "B/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "A/B"
rownames(off_full[1:6,])
#> [1] "off_1" "off_2" "off_3" "off_4" "off_5" "off_6"
```

### Prepare the inputs

APIS main function requires 4 inputs:

- off.genotype = matrix of offspring genotypes coded as explained above
- sire.genotype = matrix of sires genotypes coded as explained above
- dam.genotype = matrix of dams genotypes coded as explained above
- error = accepted assignment error rate (What is the maximum error rate I accept in my assignment results?)

```
head(off_full[,1:10])
                                                            V10
        V2
               V3
                     V4
                           V5
                                 V6
#> off_1 "A/B" "B/B" "A/A" "A/A" "B/B" "B/B" "NA/NA" "A/A" "A/A" "A/B"
#> off_2 "A/B" "B/B" "A/B" "A/A" "B/B" "B/B" "B/B"
                                                    "A/A" "A/A" "A/B"
                                                    "A/A" "A/A" "A/A"
#> off_3 "A/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B"
#> off_4 "B/B" "B/B" "A/A" "A/A" "B/B" "B/B" "B/B"
                                                    "A/A" "A/B" "A/A"
#> off 5 "B/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/A" "A/B" "A/B"
#> off 6 "A/B" "B/B" "A/A" "A/A" "B/B" "B/B" "B/B" "A/A" "B/B" "A/B"
head(sire full[,1:10])
          V2
                V3
                      V4
                            V5
                                  V6
                                        V7
                                               V8
                                                     V9
                                                           V10
#> sire_1 "A/A" "A/B" "A/A" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "A/B"
#> sire_2 "B/B" "A/B" "B/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "B/B"
#> sire 3 "A/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "A/B"
#> sire 4 "A/B" "B/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "A/B"
#> sire 5 "B/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/A" "A/A" "A/B"
#> sire 6 "B/B" "A/A" "A/A" "A/A" "B/B" "B/B" "B/B" "A/A" "B/B" "A/A"
head(dam_full[,1:10])
                     V4
                           V5
                                 V6
                                       V\gamma
                                              V8
#> dam_1 "B/B" "A/B" "B/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/A" "B/B"
#> dam 2 "A/A" "B/B" "B/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/A" "B/B"
#> dam_3 "A/B" "A/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "B/B"
#> dam_4 "A/B" "B/B" "A/B" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "B/B"
#> dam_5 "A/B" "B/B" "A/A" "A/A" "B/B" "B/B" "B/B" "A/A" "A/B" "A/A"
#> dam_6 "A/A" "B/B" "B/B" "A/A" "B/B" "B/B" "B/B" "A/A" "B/B" "B/B"
error <- 0.05 #I accept 5% of errors in the results
```

## Running the assignment

The main function to perform parentage assignment with APIS the "APIS" function. Use the function as below, with default parameters for exclusion threshold and preselection of parents for maximizing the reliability.

## Analyse the results

APIS gives you 3 different outputs :

• pedigree

Pedigree header	Description
$\overline{off}$ $sire$ $dam$	Offspring ID Sire ID Dam ID

• log containing Mendelian transmission probabilities, mismatches and deltas for the first 3 parent pairs

Log header	Description
$\overline{offspring}$	offspring ID
$mrk\_genotype$	number of markers genotyped
sire1	ID of the most likely sire
dam1	ID of the most likely dam
mismatch1	number of mismatches for the most likely parent pair (sire1, dam1)
mendel1	average Mendelian transmission probability of the most likely parent pair (sire1,
	dam1)
sire2	ID of the second most likely sire
dam2	ID of the second most likely dam
mismatch 2	number of mismatches for the second most likely parent pair (sire2, dam2)
mendel 2	average Mendelian transmission probability of the second most likely parent pair
	(sire2, dam2)
$delta\_Pmendel12$	mendel1 - mendel2
sire3	ID of the third most likely sire
dam3	ID of the third most likely dam
mismatch 3	number of mismatches for the third most likely parent pair (sire3, dam3)
mendel 3	average Mendelian transmission probability of the third most likely parent pair
	(sire3, dam3)
$delta\_Pmendel23$	mendel2 - mendel3

• graphs of the distributions of deltas, Mendelian transmission probabilities and mismatches

According to the graphs, you can change the thresholds to improve your assignment.

If you want to set up your threshold on Mendelian probabilities, use :

If you want to set up your threshold on mismatches, use :

## Examples

#### Full data

This example uses the full data provided by the package (Figure 1).

```
result <- APIS(off.genotype = off_full,</pre>
                         sire.genotype = sire_full,
                         dam.genotype = dam_full,
                         error = 0.05)
                       number of individual
                             300
                                                                                                                    Delta
                             200
                                                                                                                        delta1
                             100
                                                                                                                        delta2
                               0 -
                                                                  0.10
                                                                                  0.15
                                                                                                 0.20
                                    0.00
                                                   0.05
                                                                      delta
                       number of individuals
                                                                                                         Mendelian probability
                             200
                                                                                                            P1
                             100
                                                                                                            P2
                                               average Mendelian tranmission probability
                       number of individuals
                             800 -
                             600
                                                                                                               missmatches
                                                                                                                   miss1
                             400 -
                                                                                                                   miss2
                             200
                                     0
                                                                                               30
                                                        10
```

Figure 1: APIS outputs calculated on 1000 offspring from 10 sires and 10 dams, genotyped on 400 markers. When I look at the mismatch distributions, I prefer to use exclusion and allow for 5 mismatches (Figure 2).

number of mismatches

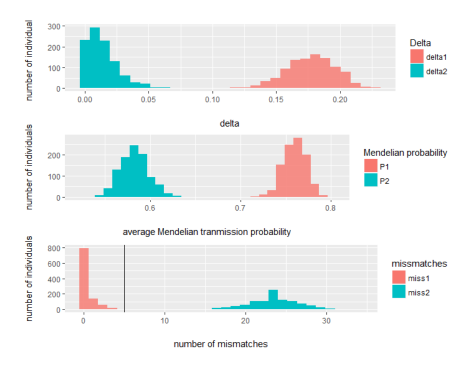


Figure 2: APIS outputs based on mismatches calculated on 1000 offspring from 10 sires and 10 dams, genotyped on 400 markers

#### Degraded data

This example uses the degraded data provided by the package (Figure 3).

In this situation, the theoretical assignment power is low and there are missing parents. The distribution graphs do not give you more information about a new threshold value.

Thus, the better answer is to keep APIS results.

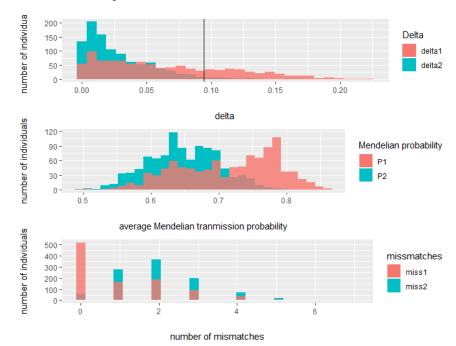


Figure 3: APIS outputs calculated on 1000 offspring from 5 sires (5 missing sires) and 10 dams, genotyped on 50 markers

# Other parameters of APIS function

APIS function can handle 2 other parameters :

- exclusion.threshold : For exclusion procedure, the number of mismatches allowed by the user. Use default value for Mendelian transmission probability procedure.
- preselect.Parent : Use of a parent preselection function based on Mendelian incompatibilities.
  - Default value : FALSE
  - One interger (n): Selection of n sires and n dams
  - Two integers (n, m): Selection of n sires and m dams

Use default value to get the most accurate results. If a value is specified, this will decrease computation time but can decrease assignment reliability.

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#### Annexe

```
print(sessionInfo(), locale=FALSE)
#> R version 3.4.1 (2017-06-30)
#> Platform: x86 64-w64-mingw32/x64 (64-bit)
#> Running under: Windows 7 x64 (build 7601) Service Pack 1
#> Matrix products: default
#>
#> attached base packages:
#> [1] parallel stats
                          graphics grDevices utils
                                                       datasets methods
#> [8] base
#>
#> other attached packages:
                        gridExtra_2.3
#> [1] APIS_0.1.0
                                          qqplot2_3.1.0
                                                             doParallel 1.0.14
#> [5] doSNOW_1.0.16
                        snow_0.4-3
                                           iterators_1.0.10 foreach_1.4.4
#>
#> loaded via a namespace (and not attached):
#> [1] tidyselect_0.2.5 xfun_0.5
                                           remotes_2.0.2
  [4] purrr_0.3.0
                         colorspace_1.4-0 usethis_1.4.0
#> [7] htmltools_0.3.6 yaml_2.2.0
                                           rlang_0.3.1
#> [10] pkqbuild 1.0.2
                         pillar 1.3.1
                                            qlue 1.3.0
#> [13] withr_2.1.2
                         sessioninfo_1.1.1 plyr_1.8.4
#> [16] stringr_1.4.0
                         munsell\_0.5.0
                                           gtable_0.2.0
#> [19] devtools_2.0.1
                         codetools_0.2-16 memoise_1.1.0
                                            callr_3.1.1
#> [22] evaluate_0.13
                         knitr_1.21
                                            backports_1.1.3
#> [25] ps_1.3.0
                         Rcpp_1.0.0
#> [28] scales_1.0.0
                         desc_1.2.0
                                           pkqload_1.0.2
#> [31] fs_1.2.6
                         digest_0.6.18
                                           stringi_1.3.1
#> [34] processx_3.2.1
                         dplyr_0.8.0.1
                                           rprojroot_1.3-2
#> [37] grid_3.4.1
                          cli_1.0.1
                                            tools_3.4.1
#> [40] magrittr_1.5
                         lazyeval_0.2.1
                                           tibble 2.0.1
#> [43] crayon_1.3.4
                         pkgconfig_2.0.2
                                            prettyunits_1.0.2
#> [46] assertthat_0.2.0 rmarkdown_1.11
                                           R6 2.4.0
#> [49] compiler_3.4.1
```