# How\_to\_use\_EFGLmh

**Thomas Delomas** 

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## Introduction

This will walk through the functions in the EFGLmh package. This package was written as a replacement for IDFGEN mainly motivated by the need to work with microhaps. It has been written to function for any codominant diploid marker, but is written and tested with SNPs and microhaps (SNPs are really just a subcategory of microhaps) in mind.

If you want to see all the options for a function, the manual has this information in a quicker-to-find format than this vignette.

One of the main differences between IDFGEN and EFGLmh is that IDFGEN keeps data in a separate environment, as if the data is somewhere in the ether until using an IDFGEN function to access it. This (mostly) prevents users from accidentally modifying things, but it also causes some bugginess, makes it hard for users to purposefully modify objects, and can cause some issues when loading in data from multiple data files. EFGLmh instead holds data as objects of a new class, called EFGLdata. As such, your dataset will have a variable name associated with it.

Now, let's walk through the functions in the package.

# **Getting data into EFGLmh**

First, install if needed, and load the package.

```
# install tidyverse if you haven't already
install.packages("tidyverse")

# install the package if needed
devtools::install_github("delomast/EFGLmh")

# and load the package
library(EFGLmh)
# and load the tidyverse for the examples here
library(tidyverse)
options(tibble.max_extra_cols = 10) # one of my preferred options for tidyverse
```

## Loading in data

Now, let's first load in our data. We can either read our data into as a dataframe, matrix, or tibble, perhaps do some modifications, and then hand it over to EFGLmh, or we can load it directly from a tab separated file into EFGLmh.

If we read it in separately, the dataframe, matrix, or tibble (below, variable exampleData) must have a column of population (pedigree) names, a column of unique individual names, optional metadata columns, and then

genotype columns in a two column per call format. The pedigree and individual names can be anywhere, if their locations are specified. The genotype columns must be consecutive and on the right hand side. The start of the genotype columns can be specified, but if not, it will look for the first column with a column name ending in ".A1", ".a1", "-A1", or "-a1". Locus names are pulled from the first column for each locus.

```
# example of loading from file outside of EFGLmh
exampleData <- readr::read_tsv("example_snp_mh.txt", guess_max = 1e4)</pre>
```

```
print(exampleData)
#> # A tibble: 4,881 x 744
     Pedigree `Individual Nam~ Gender DateSampled LengthFork1 FieldID1 GenMa GenPa
                    <chr> <date>
     <chr> <chr>
                                                     <dbl> <chr>
                                                                    <chr> <chr>
#> 1 OmyOXBO~ OmyOXBO195 0001 F
                                    2019-03-11
                                                      710 0001
                                                                    <NA> <NA>
                                    2019-03-11
#> 2 OmyOXBO~ OmyOXBO19S 0002 F
                                                       680 0002
                                                                    OmyO~ OmyO~
#> 3 OmyOXBO~ OmyOXBO195 0003 F
                                                                    OmyO~ OmyO~
                                    2019-03-11
                                                       690 0003
#> 4 OmyOXBO~ OmyOXBO19S 0004 F
                                    2019-03-11
                                                                    OmyO~ OmyO~
                                                       700 0004
                                    2019-03-11
#> 5 OmyOXBO~ OmyOXBO19S 0005 F
                                                       700 0005
                                                                    <NA> <NA>
#> 6 OmyOXBO~ OmyOXBO19S 0006 F
                                    2019-03-11
                                                                    OmyO~ OmyO~
                                                       680 0006
#> 7 OmyOXBO~ OmyOXBO19S 0007 F
                                    2019-03-11
                                                       650 0007
                                                                    OmyO~ OmyO~
#> 8 OmyOXBO~ OmyOXBO195 0008 F
                                    2019-03-11
                                                                    OmyO~ OmyO~
                                                       730 0008
#> 9 OmyOXBO~ OmyOXBO195 0009 F
                                    2019-03-11
                                                                    <NA> <NA>
                                                       660 0009
#> 10 OmyOXBO~ OmyOXBO19S 0010 F
                                    2019-03-14
                                                       740 0010
                                                                    OmyO~ OmyO~
#> # ... with 4,871 more rows, and 736 more variables: `OMS00039 mh-A1` <chr>,
    `OMS00039_mh-A2` <chr>, `OMS00052_mh-A1` <chr>, `OMS00052_mh-A2` <chr>,
     `OMS00077 mh-A1` <chr>, `OMS00077 mh-A2` <chr>, `OMS00101 mh-A1` <chr>,
     `OMS00101 mh-A2` <chr>, `OMS00116 mh-A1` <chr>, `OMS00116 mh-A2` <chr>, ...
# example of modifying in r, here we are just selecting a few populations
t1 <- exampleData %>% filter(Pedigree %in% c("OmyOXBO19S", "OmyEFSW19S"))
# or, without dplyr:
# t1 <- t[t$Pedigree %in% c("OmyOXBO19S", "OmyEFSW19S"),]
# and now we pass the data to EFGLmh
data1 <- readInData(t1)</pre>
```

If we want to read data directly from a file, we just specify the file name as the first argument. The file must have the same structure as described above: a column of population (pedigree) names, a column of unique individual names, optional metadata columns, and then genotype columns in a two column per call format. The pedigree and individual names can be anywhere, if their locations are specified. The genotype columns must be consecutive and on the right hand side. The start of the genotype columns can be specified, but if not, it will look for the first column with a column name ending in ".A1", ".a1", "-A1", or "-a1". Locus names are pulled from the first column for each locus. This has been chosen to work directly with Progeny outputs.

```
data_direct_from_file <- readInData("example_snp_mh.txt")
# metadata columns withs lots of blanks can give parsing errors.
# this can usually be solved with a larger number for the guess_max argument:
# data_direct_from_file <- readInData("example_snp_mh.txt", guess_max = 1e5)</pre>
```

There are many optional arguments for readInData:

- genotypeStart: you can specify the first genotype column if you don't want it to be auto-detected.
- pedigreeColumn: The column number that contains population (pedigree) names. Default is 1
- nameColumn: The column number that contains individual names. Default is 2

- convertNames: TRUE to convert genotype and pedigree names in the same way that IDFGEN does (remove special characters from both and remove "." from genotype names). Default is TRUE
- convertMetaDataNames: TRUE to remove special characters and spaces from metadata column names. This makes accessing them easier. Default is TRUE
- missingAlleles: a vector of values to treat as missing alleles. Default is c("0", "00", "000")
- guess\_max: when reading from a file, this is passed to read\_tsv. Parsing errors due to colun data types
  can sometimes be fixed by increasing this. Default is 1e4

The readInData function creates object of class EFGLdata. When we print them, we just a list of the population names and the number of loci.

```
data1
#> Populations:
#> [1] "OmyEFSW19S" "OmyOXB019S"
#>
#> 368 Loci
```

## Structure of EFGLdata objects

The underlying structure of an EFGLdata object is just a list of two tibbles. The first entry is named "genotypes" and contains... metadata! No, it contains population names, individual names, and genotypes in a two column per call format (missing genotype is NA). The second entry is named "metadata" and contains population names, individual names, and metadata.

```
names(data1)
#> [1] "genotypes" "metadata"
data1$genotypes
#> # A tibble: 501 x 738
     Pop Ind OMS00039 mh.A1 OMS00039 mh.A2 OMS00052 mh.A1 OMS00052 mh.A2
     <chr> <chr> <chr> <chr>
                                               <chr>>
                                                              <chr>
#> 1 OmyE~ OmyE~ GGC
                                GGC
                                               GG
                                                              TG
#> 2 OmyE~ OmyE~ GAC
                                GAC
                                               GG
                                                              GA
#> 3 OmyE~ OmyE~ GGC
                                GGC
                                               GG
                                                              TG
#> 4 OmyE~ OmyE~ GGC
                                GAC
                                               GG
#> 5 OmyE~ OmyE~ GGC
                                GAC
                                               GG
#> 6 OmyE~ OmyE~ GGC
                                GAC
                                               GG
                                                              GG
#> 7 OmyE~ OmyE~ GGC
                                GAC
                                               GG
                                                              TG
#> 8 OmyE~ OmyE~ GGC
                                GAC
                                               GG
                                                              TG
#> 9 OmyE~ OmyE~ GGC
                                GAC
                                               TG
                                                              TG
#> 10 OmyE~ OmyE~ GGC
                                GGC
                                               GG
                                                              GG
#> # ... with 491 more rows, and 732 more variables: OMS00077 mh.A1 <chr>,
#> # OMS00077 mh.A2 <chr>, OMS00101 mh.A1 <chr>, OMS00101 mh.A2 <chr>,
      OMS00116 mh.A1 <chr>, OMS00116 mh.A2 <chr>, OMS00118 mh.A1 <chr>,
#> # OMS00118 mh.A2 <chr>, OMS00120 mh.A1 <chr>, OMS00120 mh.A2 <chr>, ...
data1$metadata
#> # A tibble: 501 x 8
     Pop Ind Gender DateSampled LengthFork1 FieldID1 GenMa
                                                                         GenPa
     <chr> <chr> <chr> <chr> <chr> <date> <dbl> <chr> <dr>
                                                                         <chr>>
#> 1 OmyEFS~ OmyEFSW1~ F 2019-04-07
                                                710 0001 <NA>
                                                                         <NA>
#> 2 OmyEFS~ OmyEFSW1~ M
#> 2 OmyEFS~ OmyEFSW1~ M 2019-04-07

#> 3 OmyEFS~ OmyEFSW1~ M 2019-04-10

#> 4 OmyEFS~ OmyEFSW1~ F 2019-04-12
                              2019-04-07
                                                570 0003
                                                               <NA>
                                                                         <NA>
                              2019-04-10
                                                  640 0004
                                                               <NA>
                                                                         <NA>
                                                            OmyEFSW1~ OmyEFSW1~
                                                  680 0007
```

```
#> 5 OmyEFS~ OmyEFSW1~ M
                                         520 0011
                         2019-04-12
                                                    OmyEFSW1~ OmyEFSW1~
#> 6 OmyEFS~ OmyEFSW1~ M
                         2019-04-15
                                         #> 7 OmyEFS~ OmyEFSW1~ F
                         2019-04-15
                                         710 0017
                                                   OmyEFSW1~ OmyEFSW1~
#> 8 OmyEFS~ OmyEFSW1~ F
                         2019-04-15
                                         #> 9 OmyEFS~ OmyEFSW1~ M
                         2019-04-17
                                         760 0020
                                                    OmyEFSW1~ OmyEFSW1~
#> 10 OmyEFS~ OmyEFSW1~ M
                         2019-04-20
                                         580 0022
                                                    OmyEFSW1~ OmyEFSW1~
#> # ... with 491 more rows
```

This (hopefully) makes it simple for you to pull out or modify data manually if there is not a specially written function in the package addressing what you want to accomplish. Simply refer to data1\$genotypes or data1\$metadata and treat it as you would treat any dataframe or tibble. One thing to remember if you modify things manually: the genotypes and metadata tibbles in an EFGLdata object should have the same individuals in the same order. You can check that you haven't messed things up by using the construct\_EFGLdata function.

```
# do some modification on data1
# ...
# and then run some error checking
data1 <- construct_EFGLdata(data1)</pre>
```

# Summarizing data

There are some common data summaries you may want from your data that EFGLmh has special functions to address. We'll walk through them here.

## Just accessing data

Get the names of all populations in an EFGLdata object

```
pop_names <- getPops(data1)
pop_names
#> [1] "OmyEFSW19S" "OmyOXB019S"
```

Get the names of all individuals, or all individuals in a subset of populations

```
# all inds in data1
all_inds <- getInds(data1)
# just Look at first 20
all_inds[1:20]
#> [1] "OmyEFSW19S_0001" "OmyEFSW19S_0003" "OmyEFSW19S_0004" "OmyEFSW19S_0007"
#> [5] "OmyEFSW19S_0011" "OmyEFSW19S_0016" "OmyEFSW19S_0017" "OmyEFSW19S_0018"
#> [9] "OmyEFSW19S_0020" "OmyEFSW19S_0022" "OmyEFSW19S_0023" "OmyEFSW19S_0024"
#> [13] "OmyEFSW19S_0026" "OmyEFSW19S_0032" "OmyEFSW19S_0034" "OmyEFSW19S_0035"
#> [17] "OmyEFSW19S_0042" "OmyEFSW19S_0052" "OmyEFSW19S_0053" "OmyEFSW19S_0054"

# only inds in one pop
subset_inds <- getInds(data1, pops = c("OmyOXB019S"))
# Looking at first 20
subset_inds[1:20]
#> [1] "OmyOXB019S_0001" "OmyOXB019S_0002" "OmyOXB019S_0003" "OmyOXB019S_0004"
```

```
#> [5] "OmyOXBO19S_0005" "OmyOXBO19S_0006" "OmyOXBO19S_0007" "OmyOXBO19S_0008"

#> [9] "OmyOXBO19S_0009" "OmyOXBO19S_0010" "OmyOXBO19S_0011" "OmyOXBO19S_0012"

#> [13] "OmyOXBO19S_0013" "OmyOXBO19S_0014" "OmyOXBO19S_0015" "OmyOXBO19S_0016"

#> [17] "OmyOXBO19S_0017" "OmyOXBO19S_0018" "OmyOXBO19S_0019" "OmyOXBO19S_0020"
```

#### Get the locus names

```
loci <- getLoci(data1)</pre>
# looking at first 20
loci[1:20]
#> [1] "OMS00039 mh"
                        "OMS00052 mh"
                                            "OMS00077 mh"
                                                              "OMS00101 mh"
#> [5] "OMS00116_mh"
                         "OMS00118 mh"
                                          "OMS00120 mh"
                                                              "OMS00128 mh"
                         "OMS00143_mh"
                                           "OMS00149_mh"
#> [9] "OMS00129_mh"
                                                              "OMS00151 mh"
#> [13] "OMS00175_mh"
                         "Omy_101832195_mh" "Omy_102867443_mh" "Omy_104519624_mh"
#> [17] "Omy 107336170 mh" "Omy 108007193 mh" "Omy 109243222 mh" "Omy 110201359 mh"
```

#### Get the metadata column names

```
meta_names <- getMeta(data1)
meta_names
#> [1] "Gender" "DateSampled" "LengthFork1" "FieldID1" "GenMa"
#> [6] "GenPa"
```

Get the number of individuals in all, or a subset of populations

```
countInds <- numInds(data1)
countInds
#> OmyEFSW19S OmyOXB019S
#> 25    476
countInds_1 <- numInds(data1, pops = c("OmyOXB019S"))
countInds_1
#> OmyOXB019S
#> 476
```

dumpTable from IDFGEN is also included, as it is commonly used

```
dumpTable(geno_success, "genotyping_success.txt")
```

### some common calculations

Calculate allelic richness. I recommend doing this if you load in a dataset of just SNPs to make sure every locus has  $\leq 2$  alleles, as expected.

```
allele_rich <- aRich(data1)
# this returns a tibble
allele_rich
#> # A tibble: 343 x 2
#> Locus aRich
#> <chr> <int>
```

```
#> 1 M09AAC055
                      2
#> 2 M09AAD076
#> 3 M09AAE082
                      2
#> 4 M09AAJ163
                      2
#> 5 Ocl gshpx357
                     1
#> 6 OMGH1PROM1SNP1
                     2
#> 7 OMS00002
                      2
#> 8 OMS00003
                      2
#> 9 OMS00006
                      2
#> 10 OMS00008
                      2
#> # ... with 333 more rows
# if you want to see counts of loci by allelic richness
allele_rich %>% count(aRich)
#> # A tibble: 5 x 2
#> aRich n
#>
   <int> <int>
#> 1 1 8
#> 2
      2 262
#> 4 4 18
      5
           1
#> 5
# same thing, without tidyverse
table(allele_rich$aRich)
#>
   1 2 3 4 5
#> 8 262 54 18 1
```

#### Genotyping success of loci, as a proportion

```
loci_success <- lociSuccess(data1)</pre>
loci_success
#> # A tibble: 368 x 2
#> Locus success
    <chr>
                <dbl>
#>
#> 1 M09AAC055
                 0.952
#> 2 M09AAD076
#> 3 M09AAE082
                 0.962
#> 4 M09AAJ163 0.904
#> 5 Ocl_gshpx357 0.962
#> 6 OMGH1PROM1SNP1 0.958
#> 7 OMS00002
                0.902
#> 8 OMS00003
                  0.768
#> 9 OMS00006
                   0.964
#> 10 OMS00008
                   0.878
#> # ... with 358 more rows
```

#### Genotyping success of individuals (more about removing failed individuals later)

```
#> <chr> <chr>
                                <dbl> <int>
#> 1 OmyEFSW19S OmyEFSW19S 0001
                                0.910
                                          33
#> 2 OmyEFSW19S OmyEFSW19S 0003
                               0.880
                                          44
#> 3 OmyEFSW19S OmyEFSW19S 0004
                               0.918
                                          30
#> 4 OmyEFSW19S OmyEFSW19S 0007
                               0.916
                                         31
#> 5 OmyEFSW19S OmyEFSW19S 0011 0.924
                                        28
#> 6 OmyEFSW19S OmyEFSW19S 0016 0.932
                                         25
#> 7 OmyEFSW19S OmyEFSW19S 0017 0.913
                                        32
#> 8 OmyEFSW19S OmyEFSW19S_0018 0.916
                                        31
#> 9 OmyEFSW19S OmyEFSW19S 0020 0.916
                                        31
#> 10 OmyEFSW19S OmyEFSW19S 0022 0.913
                                          32
#> # ... with 491 more rows
# or with just a subset of loci
subsetLoci <- getLoci(data1)</pre>
subsetLoci <- subsetLoci[!grepl("SEX", subsetLoci)]</pre>
geno success noSDY <- genoSuccess(data1, loci = subsetLoci)</pre>
geno_success_noSDY
#> # A tibble: 501 x 4
#> Pop Ind
                             success numFail
    <chr> <chr>
                              <dbl> <int>
#> 1 OmyEFSW19S OmyEFSW19S 0001 0.910
                                        33
#> 2 OmyEFSW19S OmyEFSW19S 0003 0.880
                                        44
#> 3 OmyEFSW19S OmyEFSW19S 0004 0.918
                                         30
#> 4 OmyEFSW19S OmyEFSW19S 0007 0.916
                                        31
#> 5 OmyEFSW19S OmyEFSW19S 0011 0.924
                                        28
#> 6 OmyEFSW19S OmyEFSW19S 0016 0.932
                                        25
#> 7 OmyEFSW19S OmyEFSW19S 0017 0.913
                                        32
#> 8 OmyEFSW19S OmyEFSW19S 0018 0.916
                                        31
#> 9 OmyEFSW19S OmyEFSW19S 0020 0.916
                                         31
#> 10 OmyEFSW19S OmyEFSW19S 0022 0.913
                                          32
#> # ... with 491 more rows
```

Calculate observed and expected heterozygosity within each population

```
heterozygosity <- calcHet(data1)</pre>
heterozygosity
#> # A tibble: 686 x 4
                    expHet obsHet
    Pop
        Locus
   <chr>
           <chr>
                      <dbl> <dbl>
#> 1 OmyEFSW19S M09AAC055
                      0.147 0.08
#> 4 OmyEFSW19S M09AAJ163
                      0.5 0.5
#> 5 OmyEFSW19S Ocl gshpx357 0 0
#> 6 OmyEFSW19S OMGH1PROM1SNP1 0.269 0.24
#> 7 OmyEFSW19S OMS00002 0.493 0.48
#> 8 OmyEFSW19S OMS00003
                      0.340 0.348
#> # ... with 676 more rows
```

# manipulating EFGLdata objects

These functions handle some common operations we perform on our datasets.

Combining EFGLdata objects. Let's say we have two data files to read in (e.g. one mixture and one baseline, or multiple PBT baselines). We then want to combine them for filtering, analysis, export, etc.

```
# creating a second input datafile
t2 <- exampleData %>% filter(!(Pedigree %in% c("OmyOXBO195", "OmyEFSW195")))

# and now creating a second EFGLdata object
data2 <- readInData(t2)

data2
#> PopuLations:
#> [1] "OmyDWOR195" "OmyLSCR195" "OmyLYON195" "OmyPAHH195" "OmySAWT195"

#> [6] "OmyWALL195"
#>
#> 368 loci
data1
#> PopuLations:
#> [1] "OmyEFSW195" "OmyOXBO195"
#>
#> 368 loci
```

So we have two EFGLdata objects, one with 6 populations and one with 2. Now, we combine them:

```
all_data <- combineEFGLdata(data1, data2, genoComb = "intersect", metaComb = "intersect")
all_data
#> Populations:
#> [1] "OmyDWOR19S" "OmyEFSW19S" "OmyLSCR19S" "OmyLYON19S" "OmyOXBO19S"
#> [6] "OmyPAHH19S" "OmySAWT19S" "OmyWALL19S"
#>
#> 368 Loci
```

We've now created a third EFGLdata object with all 8 populations. The arguments genoComb and metaComb tell the function how to combine loci and metadata if they are different. Options are "intersect" to only keep the loci or metadata that are in both, or "union" to keep the loci or metadata that are in either (missing loci/metadata will be given values of NA). Note that you can combine an unlimited number of EFGLdata objects in one command. For example, for four objects: all\_data <- combineEFGLdata(data1, data2, data3, data4, genoComb = "intersect", metaComb = "intersect"). Having many EFGLdata objects can use a lot of memory if you have large numbers of individuals/loci. So, if you are done with the original EFGLdata objects, you can remove them to free up some memory:

```
rm(data1)
rm(data2)
```

Moving all individuals in a subset of populations to a different (new or existing) population.

```
# say we want to combine OmyDWOR19S and OmyEFSW19S, and call it "newPop"
all_data <- movePops(all_data, pops = c("OmyDWOR19S", "OmyEFSW19S"), newName = "newPop")
all_data
#> Populations:
#> [1] "newPop" "OmyLSCR19S" "OmyLYON19S" "OmyOXBO19S" "OmyPAHH19S"
```

```
#> [6] "OmySAWT19S" "OmyWALL19S"
#>
#> 368 Loci
```

#### Moving a subset of individuals to a different (new or existing) population

```
numInds(all data)
     newPop OmyLSCR19S OmyLYON19S OmyOXBO19S OmyPAHH19S OmySAWT19S OmyWALL19S
                                          1278
       1395
            126
                         87 476
                                                     864
                                                            655
toMove <- c("OmyWALL19S 0696", "OmyWALL19S 0697", "OmyWALL19S 0698")
all_data <- moveInds(all_data, inds = toMove, newName = "specialInds")</pre>
numInds(all_data)
#>
    newPop OmyLSCR19S OmyLYON19S OmyOXBO19S OmyPAHH19S OmySAWT19S
      1395 126 87 476 1278 864
#> OmyWALL19S specialInds
#> 652 3
```

#### Remove loci

```
toRemove <- c("OMS00079", "Omy_Omyclmk43896")
all_data <- removeLoci(all_data, lociRemove = toRemove)
all_data
#> Populations:
#> [1] "newPop" "OmyLSCR19S" "OmyLYON19S" "OmyOXB019S" "OmyPAHH19S"
#> [6] "OmySAWT19S" "OmyWALL19S" "specialInds"
#>
#> 366 Loci
```

#### Remove individuals

```
numInds(all_data)
#>
      newPop OmyLSCR19S OmyLYON19S OmyOXBO19S OmyPAHH19S OmySAWT19S
      1395 126 87 476 1278 864
#> OmyWALL19S specialInds
       652
toRemove <- c("OmyWALL19S 0001", "OmyWALL19S 0002")
all data <- removeInds(all data, inds = toRemove)
numInds(all data)
#>
    newPop OmyLSCR19S OmyLYON19S OmyOXBO19S OmyPAHH19S OmySAWT19S
      1395 126
                     87 476
                                            1278
                                                       864
#> OmyWALL19S specialInds
#> 650
```

#### Remove populations

```
all_data
#> Populations:
#> [1] "newPop" "OmyLSCR19S" "OmyLYON19S" "OmyOXBO19S" "OmyPAHH19S"
#> [6] "OmySAWT19S" "OmyWALL19S" "specialInds"
#>
#> 366 Loci
```

```
all_data <- removePops(all_data, pops = c("OmyLSCR19S", "OmySAWT19S"))
all_data
#> Populations:
#> [1] "newPop" "OmyLYON19S" "OmyOXBO19S" "OmyPAHH19S" "OmyWALL19S"
#> [6] "specialInds"
#>
#> 366 Loci
```

# exporting data

These functions export data in formats used by other packages and programs. They are listed here mainly so you can have a list of the export functions in one place and see an example. For a full explanation of the options within each of these functions, consult the manual.

A rubias baseline

A rubias mixture

A gRandma baseline or mixture

In addition to exporting gRandma inputs, there is also a function to remove loci for gRandma inputs that either failed for all individuals, or have no variation.

```
# this creates a list with baseline and mixture
cleanInput <- cleanGrandma(baseline = gma_baseline, mixture = gma_mixture)
#> Removing Locus Ocl_gshpx357 for no variation.
#> Removing Locus Omy_myclarp404111 for no variation.
#> Removing Locus Omy_BAMBI4238 for no variation.
#> Removing Locus Omy_G3PD_2246 for no variation.
#> Removing Locus Omy_RAD10335945 for no variation.
gma_baseline <- cleanInput$baseline
gma_mixture <- cleanInput$mixture</pre>
```

A hierfstat input dataframe

```
hfstat_in <- exportHierFstat(all_data)
```

Write a GenePop file

```
exportGenePop(all_data, "genepop.txt", useIndNames = TRUE)
```

Write a GenAlEx file

```
exportGenAlEx(all_data, "genalexInput.txt")
```

Write a SNPPIT input file (only biallelic markers used)

Write PLINK input files (only biallelic markers used)

```
exportPlink(data1, "testPlink.ped", map = "testPlink.map")
```

# examples of common steps in data analysis

## remove poorly genotyping individuals

First, we determine genotyping success. We're using all loci, but remember genoSuccess can also use just a subset of loci if you input a vector of locus names.

```
geno_success <- genoSuccess(all_data)</pre>
geno_success
#> # A tibble: 3,889 x 4
#> Pop Ind
                        success numFail
    <chr> <chr> <dbl> <int>
#> 1 newPop OmyDWOR19S 0001 0.962 14
#> 2 newPop OmyDWOR19S 0002 0.995
                                      2
#> 3 newPop OmyDWOR19S 0003 0.978
                                     8
#> 4 newPop OmyDWOR19S_0004 0.995
                                     2
#> 5 newPop OmyDWOR19S 0005 0.989
                                     4
#> 6 newPop OmyDWOR19S 0006 0.992
                                     3
                                     3
#> 7 newPop OmyDWOR19S 0007 0.992
#> 8 newPop OmyDWOR19S 0008 0.989
                                      4
#> 9 newPop OmyDWOR19S 0009
                           0.997
                                      1
#> 10 newPop OmyDWOR19S 0010
                           0.989
                                      4
#> # ... with 3,879 more rows
```

Now we get a list of individual names to remove and then use the removeInds function. We can filter by the proportion success or by the number of missing loci.

```
# identify any under 90%
failedInds <- geno_success %>% filter(success < .9) %>% pull(Ind)
# example of code to filter by number of missing loci
# (remove any with more than 37 genotypes missing)
# failedInds <- geno_success %>% filter(numFail < 37) %>% pull(Ind)
# and remove
sum(numInds(all_data))
#> [1] 3889
all_data <- removeInds(all_data, inds = failedInds)
sum(numInds(all_data))
#> [1] 3448
```

And perhaps we want to save a list of the failed individuals and their populations

```
removeTable <- geno_success %>% filter(Ind %in% failedInds)
dumpTable(removeTable, "failed_inds.txt")
# or, to efficiently do it all in the tidyverse:
# geno_success %>% filter(Ind %in% failedInds) %>% dumpTable("failed_inds.txt")
```

## removing duplicates (with some outside help)

Just as we use GSI sim to quickly identify duplicate samples, we can use rubias.

```
rubiasIn <- exportRubias_baseline(all data, pops = c("OmyPAHH19S", "specialInds"),</pre>
                                  repunit = "Pop", collection = "Pop")
# require 70% genotypes successful in both (but we've already filtered), and
# requie 95% of genotypes to be the same
library(rubias)
dupTable <- close_matching_samples(rubiasIn, gen start col = 5, min frac non miss = .7,</pre>
         min frac matching = .95)
#> Summary Statistics:
#> 1009 Individuals in Sample
#> 366 Loci: M09AAC055.A1, M09AAD076.A1, M09AAE082.A1, M09AAJ163.A1, Ocl_gshpx357.A1,
         OMGH1PROM1SNP1.A1, OMS00002.A1, OMS00003.A1, OMS00006.A1, OMS00008.A1, OMS00013.A1,
         OMS00014.A1, OMS00015.A1, OMS00017.A1, OMS00018.A1, OMS00024.A1, OMS00030.A1,
         OMS00039_mh.A1, OMS00041.A1, OMS00048.A1, OMS00052_mh.A1, OMS00053.A1, OMS00056.A1,
         OMS00057.A1, OMS00058.A1, OMS00061.A1, OMS00062.A1, OMS00064.A1, OMS00068.A1, OMS00070.A1,
         OMS00071.A1, OMS00072.A1, OMS00074.A1, OMS00077_mh.A1, OMS00078.A1, OMS00087.A1,
         OMS00089.A1, OMS00090.A1, OMS00092.A1, OMS00095.A1, OMS00096.A1, OMS00101_mh.A1,
         OMS00103.A1, OMS00105.A1, OMS00106.A1, OMS00111.A1, OMS00112.A1, OMS00114.A1,
         OMS00116_mh.A1, OMS00118_mh.A1, OMS00119.A1, OMS00120_mh.A1, OMS00121.A1, OMS00127.A1,
         OMS00128_mh.A1, OMS00129_mh.A1, OMS00132.A1, OMS00133.A1, OMS00134.A1, OMS00138.A1,
         OMS00143_mh.A1, OMS00149_mh.A1, OMS00151_mh.A1, OMS00153.A1, OMS00154.A1, OMS00156.A1,
         OMS00164.A1, OMS00169.A1, OMS00173.A1, OMS00174.A1, OMS00175_mh.A1, OMS00176.A1,
         OMS00179.A1, OMS00180.A1, Omy_1004.A1, Omy_101554306.A1, Omy_101832195_mh.A1,
         Omy_101993189.A1, Omy_102505102.A1, Omy_102867443_mh.A1, Omy_103705558.A1,
         Omy_104519624_mh.A1, Omy_104569114.A1, Omy_105075162.A1, Omy_105105448.A1, Omy_105385406.A1,
         Omy_105714265.A1, Omy_107031704.A1, Omy_10728569.A1, Omy_107336170_mh.A1, Omy_10780634.A1,
         Omy_108007193_mh.A1, Omy_109243222_mh.A1, Omy_109525403.A1, Omy_109894185.A1,
         Omy_110064419.A1, Omy_110201359_mh.A1, Omy_110362585.A1, Omy_110689148.A1, Omy_111084526.A1,
         Omy_11138351_mh.A1, Omy_111666301.A1, Omy_112301202_mh.A1, Omy_11282082_mh.A1,
         Omy_113490159.A1, Omy_114315438.A1, Omy_114587480.A1, Omy_114976223.A1, Omy_116733349.A1,
         Omy_116938264.A1, Omy_117286374.A1, Omy_117370400.A1, Omy_117540259_mh.A1, Omy_11781581.A1,
         Omy_118175396.A1, Omy_118205116.A1, Omy_11865491.A1, Omy_120255332_mh.A1, Omy_128693455.A1,
```

UMY\_128923433\_mn.A1, UMY\_128996481.A1, UMY\_1298/0/56.A1, UMY\_130524160\_mn.A1,

```
Omy 131460646 mh.A1, Omy 187760385 mh.A1, Omy 96222125 mh.A1, Omy 9707773 mh.A1,
         Omy_97660230.A1, Omy_97865196.A1, Omy_97954618_mh.A1, Omy_98683165.A1, Omy_99300202_mh.A1,
         Omy ada1071.A1, Omy anp17 mh.A1, Omy aromat280 mh.A1, Omy arp630.A1, Omy aspAT123 mh.A1,
         Omy b1266.A1, Omy b9164.A1, Omy BACB4324.A1, Omy BACF5284 mh.A1, Omy BAMBI2312.A1,
         Omy_BAMBI4238.A1, Omy_bcAKala380rd.A1, Omy_ca05064_mh.A1, Omy_carban1264_mh.A1,
         Omy_cd28130.A1, Omy_cd59206.A1, Omy_cd59b112.A1, Omy_cin172.A1, Omy_cox1221.A1,
         Omy_cox2335_mh.A1, Omy_crb106.A1, Omy_cyp17153_mh.A1, Omy_e1147_mh.A1, Omy_ftzf1217.A1,
         Omy_g1103.A1, Omy_g1282_mh.A1, Omy_G3PD_2246.A1, Omy_G3PD_2371.A1, Omy_gadd45332.A1,
         Omy_gdh271.A1, Omy_GH1P1_2.A1, Omy_gh475.A1, Omy_GHSR121_mh.A1, Omy_gluR79.A1,
         Omy_GREB1_05.A1, Omy_GREB1_09.A1, Omy_hsc71580_mh.A1, Omy_hsf1b241.A1, Omy_hsf2146.A1,
         Omy hsp4786.A1, Omy hsp70aPro329.A1, Omy hsp90BA193.A1, Omy hus152.A1, Omy IL17185.A1,
         Omy Il1b 028.A1, Omy IL1b163 mh.A1, Omy IL6320.A1, Omy impa155.A1, Omy inos97.A1,
         Omy LDHB1 i2.A1, Omy LDHB2 e5.A1, Omy LDHB2 i6.A1, Omy Lpl220 mh.A1, Omy mapK3103.A1,
         Omy mcsf268 mh.A1, Omy metA161 mh.A1, Omy metB138 mh.A1, Omy MYC 2 mh.A1,
         Omy myclarp404111.A1, Omy_myoD178.A1, Omy_nach200_mh.A1, Omy_NaKATPa350.A1, Omy_ndk152.A1,
         Omy nips299.A1, Omy nkef241 mh.A1, Omy ntl27.A1, Omy nxt2273 mh.A1, Omy Ogo4212.A1,
         Omy_OmyP9180.A1, Omy_Ots249227.A1, Omy_oxct85_mh.A1, Omy_p53262.A1, Omy_pad196_mh.A1,
         Omy ppie232 mh.A1, Omy RAD10335945.A1, Omy RAD1073310.A1, Omy RAD11659.A1, Omy RAD118659.A1,
         Omy RAD1243964.A1, Omy RAD1256614.A1, Omy RAD1303467.A1, Omy RAD1307316.A1,
         Omy RAD1349913.A1, Omy RAD1403346.A1, Omy RAD1570953.A1, Omy RAD1610420 mh.A1,
         Omy RAD1763223.A1, Omy RAD1784916.A1, Omy RAD1890348 mh.A1, Omy RAD191922.A1,
         Omy RAD1934024.A1, Omy RAD1957859.A1, Omy RAD2091711.A1, Omy RAD2212369.A1,
         Omy RAD2357743.A1, Omy RAD2389458 mh.A1, Omy RAD2428774.A1, Omy RAD2504268.A1,
         Omy RAD25678 mh.A1, Omy RAD2608069 mh.A1, Omy RAD2669136 mh.A1, Omy RAD2774055.A1,
         Omy RAD2823638.A1, Omy RAD2970018 mh.A1, Omy RAD297626 mh.A1, Omy RAD3039217.A1,
         Omy RAD3061961.A1, Omy RAD3140867.A1, Omy RAD320910.A1, Omy RAD3213958.A1,
         Omy RAD3312247.A1, Omy RAD3379824.A1, Omy RAD3500513.A1, Omy RAD351499.A1, Omy RAD354179.A1,
         Omy RAD365148.A1, Omy RAD3667.A1, Omy RAD368487.A1, Omy RAD3695253.A1, Omy RAD3781668.A1,
         Omy RAD3840619.A1, Omy RAD3915633.A1, Omy RAD392622 mh.A1, Omy RAD4013255.A1,
         Omy RAD4052048.A1, Omy RAD4064158.A1, Omy RAD4159434.A1, Omy RAD4246532.A1,
         Omy RAD4279359.A1, Omy RAD4357337.A1, Omy RAD4361242 mh.A1, Omy RAD4369441.A1,
         Omy RAD4510418.A1, Omy RAD4631435.A1, Omy RAD4645251.A1, Omy RAD4667227.A1,
         Omy RAD4708054.A1, Omy RAD4744453 mh.A1, Omy RAD4795551.A1, Omy RAD484814.A1,
         Omy RAD4879969.A1, Omy RAD4911135 mh.A1, Omy RAD5063221.A1, Omy RAD5245817 mh.A1,
         Omy RAD5281228 mh.A1, Omy RAD537456 mh.A1, Omy RAD5540454.A1, Omy RAD5599710.A1,
         Omy RAD5791629.A1, Omy RAD5821370 mh.A1, Omy RAD5883515 mh.A1, Omy RAD5975841.A1,
         Omy RAD5995044.A1, Omy RAD6013512.A1, Omy RAD61959.A1, Omy RAD6259638.A1, Omy RAD6580868.A1,
         Omy_RAD6595969.A1, Omy_RAD6640236.A1, Omy_RAD6683417.A1, Omy_RAD6863440.A1,
         Omy_RAD701631_mh.A1, Omy_RAD72108_mh.A1, Omy_RAD7252844_mh.A1, Omy_RAD7320463_mh.A1,
         Omy_RAD738450_mh.A1, Omy_RAD73959.A1, Omy_RAD7396373_mh.A1, Omy_RAD7606020.A1,
         Omy_RAD7657062_mh.A1, Omy_RAD7778954.A1, Omy_RAD7814727.A1, Omy_RAD7850257.A1,
         Omy_RAD7877610.A1, Omy_RAD7931458_mh.A1, Omy_RAD8513135.A1, Omy_RAD8670672_mh.A1,
         Omy_RAD880287.A1, Omy_RAD8812232.A1, Omy_RAD900413_mh.A1, Omy_RAD9248564_mh.A1,
         Omy_RAD9358037.A1, Omy_RAD9871553.A1, Omy_rapd167.A1, Omy_rbm4b203.A1, Omy_redd1410.A1,
         Omy_sast264_mh.A1, Omy_SECC22b88_mh.A1, Omy_srp0937.A1, Omy_sSOD1.A1, Omy_star206.A1,
         Omy_stat3273.A1, Omy_sys1188_mh.A1, Omy_tlr3377.A1, Omy_tlr5205_mh.A1, Omy_txnip343_mh.A1,
         Omy_u0779166.A1, Omy_u0953469.A1, Omy_u0954311.A1, Omy_u0956119_mh.A1, Omy_u0961043.A1,
         Omy_U11_2b154_mh.A1, Omy_UBA3b.A1, Omy_UT16_2173.A1, Omy_vamp5303.A1, Omy_vatf406.A1,
         Omy_zg5791.A1, OMY1011SNP.A1, Omy25_61284413.A1, Omy25_61285646.A1, Omy25_61286316.A1,
         Omy25_61287415.A1, Omy25_61294400.A1, Omy25_61316270.A1, Omy25_61317685.A1,
         Omy25_61317777.A1, Omy25_61318852.A1, Omy25_61322413.A1, Omy28_11607954.A1,
         Omy28_11625241.A1, Omy28_11632591.A1, Omy28_11658853.A1, Omy28_11667578.A1,
         Omy28_11671116.A1, Omy28_11676622.A1, Omy28_11683204.A1, Omy28_11773194.A1, OmyR14589.A1,
         OmyR19198.A1, OmyR24370.A1, OmyR33562.A1, OmyR40252.A1, OmyR40319.A1, OmyY1 2SEXY.A1
#> 2 Reporting Units: OmyPAHH19S, specialInds
#>
#> 2 Collections: OmyPAHH19S, specialInds
#> 8.85% of allelic data identified as missing
dupTable
#> # A tibble: 2 x 10
#> num_non_miss num_match indiv_1 indiv_2 collection_1 collection_2 sample_type_1
                      <int> <chr> <chr>
#>
            <int>
                                                         <chr>>
                                                                      <chr>>
#> 1
             335
                       335 OmyPAH~ OmyPAH~ OmyPAHH19S OmyPAHH19S
                                                                      reference
```

So we've found two pairs of dulicates, now we want to keep the ones with more genotypes. We can use the genotyping success calculated earlier to choose which one to remove. There is a special function to identify the one with lower genotyping success.

```
toRemove <- whichLower(dupTable, geno_success)
all_data <- removeInds(all_data, inds = toRemove)</pre>
```

And let's write a table of the duplicates

```
dupTable %>% select(1:6) %>% dumpTable("duplicates.txt")
```