# Package 'SNPtools'

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check.sample.call.rate

Check Sample Call Rate

# Description

Identifies samples with call rate below a given threshold.

#### Usage

```
check.sample.call.rate(sample.summary, min.call.rate)
```

#### Arguments

```
sample.summary A data frame with a "Call.rate" column for each sample.
min.call.rate Minimum acceptable call rate (between 0 and 1).
```

#### Value

A character vector with the names of samples to remove.

combinarSNPData

Combine multiple SNPDataLong objects

#### **Description**

This function merges a list of SNPDataLong objects, typically representing different SNP panels or datasets, into a single unified SNPDataLong object. It ensures that all genotype matrices have the same set of SNPs (filling missing SNPs with NA), and merges the marker map information while removing duplicate SNP entries.

#### Usage

```
combinarSNPData(lista)
```

#### **Arguments**

lista

A list of SNPDataLong objects to be combined.

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#### **Details**

The function performs the following steps internally:

- 1. Computes the union of all SNPs across input objects.
- 2. Fills missing SNP columns in each genotype matrix with NA-coded columns.
- 3. Combines genotype matrices by rows (individuals).
- 4. Merges marker maps, removing duplicates (retaining the first occurrence).
- 5. Creates and returns a new SNPDataLong object with the combined data.

#### Value

A single SNPDataLong object containing the combined genotype matrix, merged map, and a concatenated path string.

#### **Examples**

```
## Not run:
# Example usage:
combined_data <- combinarSNPData(list(snp_data1, snp_data2, snp_data3))
## End(Not run)</pre>
```

FImputeRunner

Build FImputeRunner object

#### **Description**

A convenience function to construct a 'FImputeRunner' object from basic inputs.

#### Usage

```
FImputeRunner(object, path, exec_path = "FImpute3", name = "data")
```

# Arguments

path A character string indicating the directory to save FImpute files.

exec\_path Path to the FImpute executable (default = "FImpute3").

Name for the dataset (used internally, default = "gen\_data").

geno A SnpMatrix object.

map A data.frame with SNP metadata (columns: Name, Chromosome, Position).

#### Value

An object of class 'FImputeRunner'.

4 importAllGenos

getGeno	Flexible and efficient genotype file reading with autodetection using fread

#### **Description**

This generic and method allow flexible import of SNP genotype data from Illumina FinalReport files, supporting fast initial column detection using data.table::fread, followed by full genotype matrix construction via snpStats::read.snps.long.

#### Usage

```
getGeno(...)
```

#### **Arguments**

fields A list specifying column indices for sample, SNP, allele1, allele2, and confi-

dence

codes A character vector with allele codes (e.g., c("A", "B"))

threshold Confidence threshold for genotype calling

sep Field separator used in the files

skip Number of lines to skip at the start of the file verbose Logical; if TRUE, displays progress messages every Frequency of progress update (number of SNPs)

#### Value

An SNPDataLong object containing the genotype matrix and map, or NULL if an error occurs

importAllGenos	
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#### **Description**

This generic and method import genotype data from multiple configurations defined in an SNPImportList object, then combine them into a single unified SNPDataLong object.

#### Usage

```
importAllGenos(object)
```

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#### **Arguments**

object

An object of class SNPImportList containing import configurations

#### Value

A single combined SNPDataLong object

import\_geno\_list

Import multiple genotype datasets from a list of configurations

#### **Description**

This function iterates over a list of configuration lists (each specifying parameters such as path, fields, separators, etc.), imports each genotype dataset using getGeno(), and then combines them into a single SNPDataLong object.

#### Usage

```
import_geno_list(config_list)
```

#### **Arguments**

config\_list A list of configuration lists. Each element must include at least path and fields. Optional elements include codes, threshold, sep, skip, and verbose.

#### Value

A unified SNPDataLong object containing combined genotype data from all configurations.

```
## Not run:
configs <- list(
   list(path = "panel1", fields = list(sample = 2, snp = 1, allele1 = 7, allele2 = 8, confidence = 9)),
   list(path = "panel2", fields = list(sample = 2, snp = 1, allele1 = 7, allele2 = 8, confidence = 9), threshold = 0.10
)
combined_data <- import_geno_list(configs)
## End(Not run)</pre>
```

6 qcSamples

qcSamples

Quality control on samples

#### **Description**

Applies quality control (QC) procedures to samples in a 'SNPDataLong' object, based on heterozygosity and call rate thresholds.

#### Usage

```
qcSamples(x, ...)
## S4 method for signature 'SNPDataLong'
qcSamples(
    x,
    heterozygosity = NULL,
    smp_cr = NULL,
    action = c("report", "filter", "both")
)
```

#### **Arguments**

x An object of class 'SNPDataLong'.

heterozygosity A numeric threshold or range for heterozygosity. Samples outside this threshold

are removed.

smp\_cr Minimum acceptable sample call rate (between 0 and 1). Samples below this

value are removed.

action Character string indicating the action to perform. One of: - "report": only

returns a list of samples to remove and those kept; - "filter": returns a filtered object without reporting; - "both": performs filtering and returns the filtered

object.

#### Value

Depending on the 'action' argument: - '"report"': returns a list with removed and kept samples; - '"filter"': returns a new 'SNPDataLong' object with filtered genotypes; - '"both"': returns a list with: - 'filtered': the filtered 'SNPDataLong' object; - 'report': a list of removed and kept samples.

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qcSNPs	Quality Control for SNPDataLong with optional criteria

# Description

Allows applying genotypic quality filters with user-defined criteria, including call rate, MAF, HWE, monomorphism, chromosome filtering, and removal of SNPs at the same genomic position.

#### Usage

```
qcSNPs(x, ...)
```

# Arguments

X	An object of class SNPDataLong.
missing_ind	Maximum allowed proportion of missing data per individual (optional). *[Currently not implemented in this function]*
missing_snp	Maximum allowed proportion of missing data per SNP (optional). *[Currently not implemented in this function]*
min_snp_cr	Minimum acceptable call rate for SNPs.
min_maf	Minimum minor allele frequency allowed for SNPs (optional).
hwe	p-value threshold for Hardy-Weinberg equilibrium test (optional).
snp_position	Logical. If TRUE, removes SNPs mapped to the same position, keeping the one with the highest MAF.
snp_mono	Logical. If TRUE, identifies and removes monomorphic SNPs.
remove_chr	Character vector of chromosomes to exclude (optional).
action	One of "report" (returns a list of removed SNPs), "filter" (returns a filtered SNPDataLong object), or "both" (returns both).

#### Value

Depending on the action argument: - "report": list with SNPs removed by each criterion and SNPs kept. - "filter": filtered SNPDataLong object. - "both": list with the filtered object and detailed report.

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```
qcSNPs(x, min_snp_cr = 0.8, min_maf = 0.05, snp_mono = TRUE, snp_position = TRUE, action = "report")
## End(Not run)
```

qc\_header

Formatted header message

#### **Description**

Prints a formatted message with a border for section titles in the console.

#### Usage

```
qc_header(title)
```

#### **Arguments**

title

Character string to be printed inside the header box.

#### Value

No return value. Used for side effects (message).

# **Examples**

```
qc_header("Quality Control on Samples")
```

read.fimpute

Read imputed genotypes from FImpute output and return SNPData-Long object

#### **Description**

Reads imputed genotypes and SNP information from FImpute output, builds a SnpMatrix and a corresponding map, and returns an SNPDataLong object.

#### Usage

```
read.fimpute(file)
```

#### **Arguments**

file

Character. Path to the FImpute output directory (usually "output\_fimpute").

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#### Value

An object of class SNPDataLong containing the imputed genotypes and SNP map.

#### **Examples**

```
## Not run:
snp_long <- read.fimpute("output_fimpute")
## End(Not run)</pre>
```

runFImpute

Run FImpute from a FImputeRunner object

# Description

This function runs the external FImpute software using a 'FImputeRunner' object, ensuring that all required input files are present and the results are imported.

#### Usage

```
runFImpute(object, verbose = TRUE)
## S4 method for signature 'FImputeRunner'
runFImpute(object, verbose = TRUE)
```

#### Arguments

object An object of class 'FImputeRunner'.

verbose Logical. If TRUE (default), FImpute output will be printed to the console.

#### Value

An updated 'FImputeRunner' object with the 'results' slot populated (SnpMatrix).

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saveFImpute

Save genotype and map files in FImpute format

#### **Description**

S4 method to export genotype ('.gen'), map ('.map'), and parameter ('data.par') files compatible with the [FImpute](https://www.aps.uoguelph.ca/~msargol/fimpute/) software.

#### Usage

```
saveFImpute(object, ...)
## S4 method for signature 'FImputeExport'
saveFImpute(object)
## S4 method for signature 'SNPDataLong'
saveFImpute(object, path = NULL)
```

#### **Arguments**

object An object of class 'FImputeExport' or 'SNPDataLong'.

... Further arguments passed to methods.

path Character. Output directory where files will be written (only for 'SNPDataLong' method; default = '"fimpute\_run"').

#### Value

No return value. Files are saved to disk.

```
## Not run:
if (requireNamespace("snpStats", quietly = TRUE)) {
   mat <- matrix(sample(c(0L, 1L, 2L), 50, replace = TRUE), nrow = 5)
   colnames(mat) <- paste0("snp", 1:10)
   rownames(mat) <- paste0("ind", 1:5)

sm <- new("SnpMatrix", data = as.raw(mat))</pre>
```

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```
map <- data.frame(Name = colnames(mat), Chromosome = 1, Position = 1:10)
x <- new("SNPDataLong", geno = sm, map = map)
saveFImpute(x, path = tempdir())
}
## End(Not run)</pre>
```

saveFImputeRaw

Export genotypes and map using basic arguments

#### **Description**

Convenience function to export FImpute files directly from a 'SnpMatrix' and map 'data.frame'.

#### Usage

```
saveFImputeRaw(geno, map, path)
```

#### Arguments

geno A 'SnpMatrix' object (from the 'snpStats' package).

Map A 'data.frame' with columns 'Name', 'Chromosome', and 'Position'.

Path where the files will be saved.

# Value

No return value. Files are saved to disk.

savePlink

Save SNPDataLong object to PLINK format

# Description

Saves genotype and map data from an SNPDataLong object in PLINK format (.ped/.map and optionally binary files).

# Usage

```
savePlink(
  object,
  path = "plink_out",
  name = "plink_data",
  run_plink = TRUE,
  chunk_size = 1000
)
```

#### **Arguments**

object An object of class SNPDataLong.

path Character. Directory where files will be saved.

name Character. Base name for PLINK output files.

run\_plink Logical. If TRUE (default), runs PLINK1 to convert to binary files. If FALSE,

only .ped and .map files are saved.

chunk\_size Integer. Number of individuals per chunk for writing .ped file (default: 1000).

#### Value

No return value. Files are saved to disk.

#### **Examples**

```
## Not run:
savePlink(genotypes_qc, path = "plink_out", name = "nelore_qc", run_plink = TRUE, chunk_size = 2000)
## End(Not run)
```

 $\verb|summary,SNPD| ataLong-method|$ 

Summary for SNPDataLong objects

# Description

Provides a detailed summary of an SNPDataLong object, including sample and SNP counts, proportion of missing data, and SNP distribution by chromosome if mapping information is available.

#### Usage

```
## S4 method for signature 'SNPDataLong'
summary(object, ...)
```

#### **Arguments**

object An object of class SNPDataLong.

#### Value

Prints a summary to the console. Returns NULL (invisible).

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