# Package 'rrbgen'

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Type Package	
Title Robbie's lightweight limited functionality R bgen read/write library	
Version 0.0.3	
<b>Date</b> 2018-07-11	
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<b>Description</b> A lightweight R bgen library with limited functionality	
Imports Rcpp, RcppArmadillo	
LinkingTo Rcpp, RcppArmadillo	
License GPL   file LICENSE	
SystemRequirements C++11	
NeedsCompilation yes	
Suggests testthat	
RoxygenNote 6.0.1	
R topics documented:	
Tropies documented.	
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rrbgen_load Load entire bgen file	
	_
Usage	

rrbgen\_load(bgen\_file, gp\_names\_col = "varid", vars\_to\_get = NULL,

samples\_to\_get = NULL)

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## **Arguments**

bgen\_file Path to bgen file to load Which column to use to label the variants in the first dimension of the genotype gp\_names\_col probabilities. Note that bgen has two variant ID columns, a variant identifier (here labelled varid) and rsid (here labelled rsid) Only load these variants from file (from varid column). Note that this requires vars\_to\_get a complete pass over the data, although this should be efficient, as nothing is decompressed samples\_to\_get Only load these samples from file. Note that this requires a complete pass over

the data, although this should be efficient, as nothing is decompressed

#### Value

gp, sample\_names and var\_info

## Author(s)

Robert Davies

rrbgen\_load\_samples Load samples names only from a bgen file

#### Usage

```
rrbgen_load_samples(bgen_file)
```

## Arguments

bgen\_file Path to bgen file to load

#### Value

sample\_names

#### Author(s)

Robert Davies

```
rrbgen_load_variant_info
```

Load variant information only from a bgen file

#### Usage

```
rrbgen_load_variant_info(bgen_file)
```

## **Arguments**

bgen\_file Path to bgen file to load

#### Value

var\_info

#### Author(s)

Robert Davies

rrbgen\_write

Write variant information to a bgen file

#### Usage

```
rrbgen_write(bgen_file, sample_names = NULL, var_info = NULL, gp = NULL,
  list_of_gp_raw_t = NULL, free = NULL, Layout = 2,
  CompressedSNPBlocks = 1, B_bit_prob = 8, close_bgen_file = TRUE,
  add_to_bgen_connection = FALSE, bgen_file_connection = NULL,
  previous_offset = NULL, header_M = NULL)
```

## **Arguments**

bgen\_file Path to bgen file to be written to sample\_names character vector of sample names

var\_info var\_info (6 first columns of gen file, or chr, varid, rsid, position (1-based), ref,

alt)

gp genotype probabilities as 3-dimensional array, where dim1 = variants, dim2 =

samples, dim3 = hom ref, het, hom alt

list\_of\_gp\_raw\_t

list of raw data. bespoke format. to understand, investigate the tests

free free text for header. Should probably remain NULL. Not read if written in rrb-

gen\_read

Layout Only supported 2 (see bgen spec)

CompressedSNPBlocks

Only supported 1. Whether to (1) or not to (0) compress genotype probabilities

B\_bit\_prob How many bits to use to encode genotype probabilities (only supported 8, 16,

24, 32)

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# Author(s)

Robert Davies

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