

Package ‘hirisplexr’

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Type Package

Title From 'PLINK' to 'HirisPlex'

Version 0.1.0

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Description Read 'PLINK' 1.9 binary datasets (BED/BIM/FAM) and generate the CSV files required by the Erasmus MC 'HirisPlex' / 'HirisPlex-S' webtool <<https://hirisplex.erasmusmc.nl/>>. It maps 'PLINK' alleles to the webtool's required 'rsID_Allele' columns (0/1/2/NA). No external tools (e.g., 'PLINK CLI') are required.

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URL <https://github.com/adhikari-statgen-lab/hirisplexr>

BugReports <https://github.com/adhikari-statgen-lab/hirisplexr/issues>

Depends R (>= 3.6)

Imports BEDMatrix, data.table, utils

Suggests testthat (>= 3.0.0), knitr, rmarkdown

Config/testthat/edition 3

Encoding UTF-8

RoxygenNote 7.3.2

VignetteBuilder knitr

NeedsCompilation no

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Repository CRAN

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write_hirisplex_csv	<i>Write HIrisPlex / HIrisPlex-S CSV from a PLINK BED/BIM/FAM prefix</i>
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Description

Given a PLINK 1.9 binary dataset (prefix.bed/.bim/.fam), this function produces a CSV ready to upload to the HIrisPlex(-S) webtool.

Usage

```
write_hirisplex_csv(
  prefix,
  out,
  panel = c("hirisplexs", "hirisplex", "irisplex"),
  sample_id = c("IID", "FID_IID"),
  allow_strand_flip = TRUE
)
```

Arguments

prefix	Character. Path prefix to PLINK files, without extension.
out	Character. Output CSV path.
panel	Character. One of "hirisplexs" (default), "hirisplex", "irisplex".#
sample_id	Character. How to form 'SampleID': "IID" or "FID_IID". Default is "IID".
allow_strand_flip	Logical. If TRUE, attempt to match the required allele by allowing strand complements.

Details

Columns are 'SampleID' followed by one column per required SNP in the form rsID_Allele (e.g., rs12203592_T). Each cell contains 0/1/2 (count of the input allele) or NA when the SNP is missing. The column set and order are defined by the selected panel (IrisPlex, HIrisPlex, HIrisPlex-S).

Allele counting is based on the PLINK .bim alleles. Genotype dosage is read on demand from the .bed using `BEDMatrix::BEDMatrix()`, which encodes the dosage of the first allele in the .bim file (A1). If the webtool's required input allele equals A1, we use the dosage directly; if it equals A2, we use (2 - dosage). If `allow_strand_flip = TRUE`, we also reconcile complements (A<->T, C<->G) to account for strand orientation differences.

Value

(Invisibly) the output file path.

Examples

```
## Not run:  
write_hirisplex_csv("/path/to/prefix", tempfile(fileext = ".csv"), panel = "hirisplexs")  
  
## End(Not run)
```

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