solareclipser

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Introduction and Installation

solareclipser is an R package to interface with <u>SOLAR-Eclipse</u>. solareclipser requires <u>solar</u>.

Install solareclipser from GitHub using devtools with the following:

```
install.packages("devtools")
devtools::install_github("txbrain-org/solareclipser")
```

Usage

solareclipser defaults to using a tmp directory for the output files. However, this can be overridden by passing settings to the Solar class.

Solar S6 class

Set the settings for Solar class.

```
library(solareclipser)

settings <- list(
  output = list(
    dir = "tests/output/solar",
    tcl = FALSE,
    stdout_and_stderr = FALSE
)
)</pre>
```

Note: setting dir to a directory from a previous run will allow not needing to load the pedigree and phenotypes again.

Models

Polygenic

FPHI

```
solar$cmd$fphi(evd_data = "evd_data")
solar$run()
#> Executing solar command... Done.
```

SolarFiles

SolarFiles class can be used to get all/individual output files.

```
sf <- SolarFiles$new(settings = settings)</pre>
sf$set_polygenic_trait_d(trait d = "CC")
sfc <- SolarFilesController$new(sf = sf)</pre>
sfc$set polygenic trait d files()
trait_d_files <- sfc$get_polygenic_trait_d_files()</pre>
mod_files <- sfc$get_mod_files()</pre>
out files <- sfc$get out files()</pre>
stat_files <- sfc$get_stats_files()</pre>
summary <- readLines(out_files[stringr::str_detect(out_files, "polygenic.out")])</pre>
cat(stringr::str_trim(summary), sep = "\n")
#> Pedigree:
                HCP_imputed_filtered_ped.csv empirical
#> Phenotypes: HCP WM ave norm.csv
#> Trait:
                CC
                                        Individuals: 999
#>
\#> H2r \ is \ 0.9344430 \ p = 8.148806e-70 \ (Significant)
#> H2r Std. Error: 0.0104121
#>
#> Warning. Unexpectedly high heritabilities might result from
#> numerical problems, especially if mztwins are present.
#>
#>
#> Loglikelihoods and chi's are in CC/polygenic.logs.out
#> Best model is named poly and null0
#> Final models are named poly, spor
#> Residual Kurtosis is -0.0900, within normal range
```

Additional Examples

Using settings to set tcl and stdout and stderr to TRUE.

```
library(solareclipser)
settings <- list(
 output = list(
   dir = "tests/output/solar",
   tcl = TRUE.
   stdout and stderr = TRUE
solar <- Solar$new(settings = settings)</pre>
solar$cmd$load(obj = "pedigree",
             fpath = "tests/input/solar/HCP_imputed_filtered_ped.csv",
             cond = "-t 0")
solar$cmd$load(obj = "phenotypes",
              fpath = "tests/input/solar/HCP_WM_ave_norm.csv")
solar$cmd$trait("CC")$polygenic()
solarsrun()
#>
#> proc file772770d4455e {} {
   load pedigree HCP_imputed_filtered_ped.csv -t 0
#>
#>
    load phenotypes HCP_WM_ave_norm.csv
    trait CC
#>
#>
    polygenic
#> }
#> ==:
#> ==========> stdout and stderr <=======
#> Unloading current pedigree data ...
#> Loading Empirical Pedigree...
#> There are 2284 people. If not correct, be sure ID's are unique.
#> Determining families (use -1 option to skip this)
#> Warning if number of families is greater than one it may
#> cause an issue when loading phi2 matrix. Use -1 option to
#> avoid any potential issues.
```

```
#> * Maximize sporadic model *
#> *********************
#>
#>
     *** Loglikelihood of sporadic model is -487.520945
#>
#> *******************
#> * Maximize polygenic model
#>
#>
     *** Loglikelihood of polygenic model is -332.229219
    *** H2r in polygenic model is 0.9344430
#>
#>
     *** Determining significance of H2r
#>
    *** Comparing polygenic and sporadic models
#>
    *** chi = 310.5835, deg = 1, p = 8.148806e-70
#>
#>
#> *
                       Summary of Results
#> **************************
#>
#> Pedigree: HCP_imputed_filtered_ped.csv empirical
#> Phenotypes: HCP_WM_ave_norm.csv
#> Trait: CC
                              Individuals: 999
#>
         H2r is 0.9344430 p = 8.148806e-70 (Significant)
#>
#>
       H2r Std. Error: 0.0104121
#>
#> Warning. Unexpectedly high heritabilities might result from
#>
  numerical problems, especially if mztwins are present.
#>
#>
#> Output files and models are in directory CC/
#> Summary results are in CC/polygenic.out
#> Loglikelihoods and chi's are in CC/polygenic.logs.out
#> Best model is named poly and null0 (currently loaded)
#> Final models are named poly, spor
#>
#> Residual Kurtosis is -0.0900, within normal range
```

Additional References

- education solar-eclipse-genetics.org
- solareclipse.pdf solar-eclipse-genetics.org
- www.nitrc.org solar documents