Package 'gaawr2'

February 21, 2025

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Title Genetic Association Analysis
Version 0.0.2
Date 2025-2-20
Maintainer Jing Hua Zhao <jinghuazhao@hotmail.com>
Description It gathers information, meta-data and scripts in a two-part Henry-Stewart talk by
     Zhao (2009, <doi:10.69645/DCRY5578>), which showcases analysis in aspects such as test-
     ing of polymorphic
     variant(s) for Hardy-
     Weinberg equilibrium, association with trait using genetic and statistical models
     as well as Bayesian implementation, power calculation in study design and genetic annota-
     tion. It also
     covers R integration with the Linux environment, GitHub, package creation and web applications.
License MIT + file LICENSE
URL https://jinghuazhao.github.io/gaawr2/,
     https://github.com/jinghuazhao/gaawr2
BugReports https://github.com/jinghuazhao/gaawr2/issues
Encoding UTF-8
Depends R (>= 3.5.0)
Imports dplyr, gap, gap.datasets, ggplot2, survival, Rdpack
RdMacros Rdpack
LazyData Yes
LazyLoad Yes
LazyDataCompression xz
VignetteBuilder knitr
Suggests BLR, BGLR, biomaRt, bookdown, EnsDb.Hsapiens.v75, ensembldb,
     GMMAT, HardyWeinberg, haplo.stats, httr, httpuv, jsonlite,
     kableExtra, knitr, MCMCglmm, plumber, powerEQTL, R2jags,
     regress, seqminer, SNPassoc, testthat, tidyr
RoxygenNote 7.3.2
NeedsCompilation no
```

2 welcome

Author Jing Hua Zhao [aut, cre] (https://orcid.org/0000-0003-4930-3582),
Benjamin Altmann [ctb]

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welco	ome	An enhanced welcome			

Description

It prints a welcome message, saying number of times.

Usage

welcome(n)

Arguments

n

The number of times (>1 integer) to welcome the user.

Value

Prints a welcome message to the console.

Examples

welcome(3)

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