

Package ‘gaawr2’

July 22, 2025

Title Genetic Association Analysis

Version 0.0.3

Date 2025-3-24

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Description It gathers information, meta-data and scripts in a two-part Henry-Stewart talk by Zhao (2009, <[doi:10.69645/DCRY5578](https://doi.org/10.69645/DCRY5578)>), which showcases analysis in aspects such as testing 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 annotation. It also covers R integration with the Linux environment, GitHub, package creation and web applications.

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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, Cairo, EnsDb.Hsapiens.v75, ensemblldb, GMMAT, HardyWeinberg, haplo.stats, httr, httpuv, jsonlite, kableExtra, knitr, MCMCglmm, plumber, powerEQTL, R2jags, regress, seqminer, SNPassoc, testthat, tidy

RoxygenNote 7.3.2

NeedsCompilation no

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Repository CRAN

Date/Publication 2025-03-24 15:00:09 UTC

Contents

welcome	2
Index	3

welcome	<i>An enhanced welcome</i>
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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)

Index

welcome, [2](#)