Package 'Pv3Rs'

November 20, 2025

Title Estimate the Cause of Recurrent Vivax Malaria using Genetic Data Version 1.0.0 **Description** Plot malaria parasite genetic data on two or more episodes. Compute per-person posterior probabilities that each Plasmodium vivax (Pv) recurrence is a recrudescence, relapse, or reinfection (3Rs) using per-person P. vivax genetic data on two or more episodes and a statistical model described in Taylor, Foo and White (2022) <doi:10.1101/2022.11.23.22282669>. Plot per-recurrence posterior probabilities. **License** GPL (>= 3) URL https://aimeertaylor.github.io/Pv3Rs/, https://github.com/aimeertaylor/Pv3Rs BugReports https://github.com/aimeertaylor/Pv3Rs/issues **Depends** R (>= 3.5) Imports dplyr, fields, grDevices, igraph, matrixStats, methods, multicool, partitions, purrr, RColorBrewer Suggests codetools, gtools, knitr, Matrix, plyr, R.rsp, rmarkdown, testthat (>= 3.0.0), tictoc, utils VignetteBuilder knitr Config/Needs/website rmarkdown Config/testthat/edition 3 **Encoding UTF-8** LazyData true RoxygenNote 7.3.2 NeedsCompilation no **Author** Aimee Taylor [aut, cre] (ORCID: <https://orcid.org/0000-0002-2337-8992>), Yong See Foo [aut] (ORCID: https://orcid.org/0000-0003-3010-9106), Tymoteusz Kwiecinski [com] (compiled included macros.Rd), Duncan Murdoch [ctb] (author of included macros.Rd),

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Date/Publication 2025-11-20 13:10:02 UTC
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compu	e_posterior Compute posterior probabilities of P. vivax recurrence states	

Description

Computes per-person posterior probabilities of *P. vivax* recurrence states — recrudescence, relapse, reinfection — using per-person genetic data on two or more episodes. For usage, see **Examples** below and Demonstrate Pv3Rs usage. For a more complete understanding of compute_posterior output, see Understand posterior probabilities.

Note: The progress bar may increment non-uniformly (see **Details**); it may appear stuck when computations are ongoing.

Usage

```
compute_posterior(
  y,
  fs,
  prior = NULL,
  MOIs = NULL,
  return.RG = FALSE,
  return.logp = FALSE,
  progress.bar = TRUE
)
```

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Arguments

List of lists encoding allelic data. The outer list contains episodes in chrono-У logical order. The inner list contains named markers per episode. Marker names must be consistent across episodes. NA indicates missing marker data; otherwise, specify a per-marker vector of distinct alleles detected (presently, compute_posterior() does not support data on the proportional abundance of detected alleles). Repeat alleles and NA entries within allelic vectors are ignored. Allele names are arbitrary, allowing for different data types, but must correspond with frequency names. fs List of per-marker allele frequency vectors, with names matching marker names in y. Per-marker alleles frequencies mut contain one frequency per named allele, with names matching alleles in y. Per-marker frequencies must sum to one. Matrix of prior probabilities of recurrence states per episode, with rows as episodes prior in chronological order, and columns named "C", "L", and "I" for recrudescence, relapse and reinfection, respectively. Row names are ignored. If NULL (default), per-episode recurrence states are assumed equally likely a priori. MOIs Vector of per-episode multiplicities of infection (MOIs); because the Pv3Rs model assumes no genotyping errors, MOIs must be greater than or equal to the most parsimonious MOI estimates compatible with the data; see determine_MOIs(y). These are the estimates used when MOIs = NULL (default). Logical; returns the relationship graphs (default FALSE). Automatically set to return.RG TRUE if return. logp = TRUE. Logical; returns the log-likelihood for each relationship graph (default FALSE). return.logp Setting TRUE disables permutation symmetry optimisation and thus increases runtime, especially when MOIs are large. Does not affect the output of the posterior probabilities; for an an example of permutation symmetry, see Exploration of relationship graphs in Demonstrate Pv3Rs usage. Logical; show progress bars (default TRUE). Note that the progress bar may upprogress.bar date non-uniformly.

Details

compute_posterior() computes posterior probabilities proportional to the likelihood multiplied by the prior. The likelihood sums over:

- ways to phase allelic data onto haploid genotypes
- graphs of relationships between haploid genotypes
- · ways to partition alleles into clusters of identity-by-descent

We enumerate all possible relationship graphs between haploid genotypes, where pairs of genotypes can either be clones, siblings, or strangers. The likelihood of a sequence of recurrence states can be determined from the likelihood of all relationship graphs compatible with said sequence. More details on the enumeration of relationship graphs can be found in enumerate_RGs. For each relationship graph, the model sums over all possible identity-by-descent partitions. Because some graphs are compatible with more partitions than others, the log p(YlRG) progress bar may advance non-uniformly. We do not recommend running 'compute_posterior() when the total genotype count (sum of MOIs) exceeds eight because there are too many relationship graphs.

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Notable model assumptions and limitations:

- · All siblings are regular siblings
- Recrudescent parasites derive only from the immediately preceding episode
- Recrudescence, relapse and reinfection are mutually exclusive
- Undetected alleles, genotyping errors, and de novo mutations are not modelled
- Population structure and various other complexities that confound molecular correction are not modelled

Value

List containing:

marg Matrix of marginal posterior probabilities for each recurrence, with rows as recurrences and columns as "C" (recrudescence), "L" (relapse), and "I" (reinfection). Each marginal probability sums over a subset of joint probabilities. For example, the marginal probability of "C" at the first of two recurrences sums over the joint probabilities of "CC", "CL", and "CI".

joint Vector of joint posterior probabilities for each recurrence state sequence; within a sequence "C", "L", and "I" are used as above.

RGs List of lists encoding relationship graphs; returned only if return.RG = TRUE (default FALSE), and with log-likelihoods if return.logp = TRUE (default FALSE). A relationship graph encoded as a list can be converted into a igraph object using RG_to_igraph and thus plotted using plot_RG. For more details on relationship graphs, see enumerate_RGs.

Examples

```
# Running example (runs across compute_posterior, plot_data and plot_simplex)
# based on real data from chloroquine-treated participant 52 of the Vivax
# History Trial (Chu et al. 2018a, https://doi.org/10.1093/cid/ciy319)
y <- ys_VHX_BPD[["VHX_52"]] # y is a list of length two (two episodes)
compute_posterior(y, fs_VHX_BPD, progress.bar = FALSE)
# Numerically named alleles
y \leftarrow list(enrol = list(m1 = c('3', '2'), m2 = c('1', '2')),
          recur1 = list(m1 = c('1', '4'), m2 = c('1')),
          recur2 = list(m1 = c('1'), m2 = NA))
fs \leftarrow list(m1 = c('1' = 0.78, '2' = 0.14, '3' = 0.07, '4' = 0.01),
           m2 = c('1' = 0.27, '2' = 0.73))
compute_posterior(y, fs, progress.bar = FALSE)
# Arbitrarily named alleles, plotting per-recurrence posteriors
y <- list(enrolment = list(marker1 = c("Tinky Winky", "Dipsy"),
                          marker2 = c("Tinky Winky", "Laa-Laa", "Po")),
          recurrence = list(marker1 = "Tinky Winky",
                          marker2 = "Laa-Laa"))
fs <- list(marker1 = c("Tinky Winky" = 0.4, "Dipsy" = 0.6),
           marker2 = c("Tinky Winky" = 0.1, "Laa-Laa" = 0.1, "Po" = 0.8))
plot_simplex(p.coords = compute_posterior(y, fs, progress.bar = FALSE)$marg)
```

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```
# Episode names are cosmetic: "r1_prior" is returned for "r2"
y \leftarrow list(enrol = list(m1 = NA), r2 = list(m1 = NA), r1 = list(m1 = NA))
prior <- matrix(c(0.6,0.7,0.2,0.3,0.2,0), ncol = 3,
                dimnames = list(c("r1_prior", "r2_prior"), c("C", "L", "I")))
suppressMessages(compute_posterior(y, fs = list(m1 = c(a = 1)), prior))
prior
# Prior is returned when all data are missing
y_missing <- list(enrol = list(m1 = NA), recur = list(m1 = NA))</pre>
suppressMessages(compute_posterior(y_missing, fs = list(m1 = c("A" = 1))))
# Return of the prior re-weighted to the exclusion of recrudescence:
suppressMessages(compute_posterior(y_missing, fs = list(m1 = c("A" = 1)),
                 MOIs = c(1,2))
# (Recrudescing parasites are clones of previous blood-stage parasites. The
# Pv3R model assumes no within-host de-novo mutations and perfect allele
# detection. As such, recrudescence is incompatible with an MOI increase on
# the preceding infection.)
# Beware provision of unpaired data: the prior is not necessarily returned;
# for more details, see link above to "Understand posterior estimates"
y \leftarrow list(list(m1 = c('1', '2')), list(m1 = NA))
fs \leftarrow list(m1 = c('1' = 0.5, '2' = 0.5))
suppressMessages(compute_posterior(y, fs))$marg
```

determine_MOIs

Determine multiplicities of infection (MOIs)

Description

Returns a MOI estimate for each episode based on allelic diversity across markers.

Usage

```
determine_MOIs(y, return.names = FALSE)
```

Arguments

У

List of lists encoding allelic data; see compute_posterior for more details. The outer list contains episodes in chronological order. The inner list contains named markers per episode. For each marker, one must specify an allelic vector: a set of distinct alleles detected at that marker; or NA if marker data are missing.

return.names

Logical; if TRUE and y has named episodes, episode names are returned.

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Details

A true MOI is a number of genetically distinct groups of clonal parasites within an infection. Give or take *de novo* mutations, all parasites within a clonal group share the same DNA sequence, which we call a genotype. As such, MOIs are distinct parasite genotype counts. Under the Pv3Rs model assumption that there are no genotyping errors, the true MOI of an episode is greater than or equal to the maximum distinct allele count for any marker in the data on that episode. In other words, under the assumption of no genotyping errors, maximum distinct allelic counts are the most parsimonious MOI estimates compatible with the data. By default, these MOI estimates are used by compute_posterior.

Value

Numeric vector containing one MOI estimate per episode, each estimate representing the maximum number of distinct alleles observed at any marker per episode.

Examples

enumerate_RGs

Enumerate relationship graphs (RGs)

Description

An RG is a graph over all per-person parasite genotypes (each as a vertex), with edges between clone and sibling genotypes. Valid RGs satisfy:

- Subgraphs induced by clone edges are cluster graphs.
- Subgraphs induced by clone plus sibling edges are cluster graphs.
- Clone edges only link genotypes from different episodes.

Usage

```
enumerate_RGs(MOIs, igraph = TRUE, progress.bar = TRUE)
```

Arguments

MOIs Vector of per-episode multiplicities of infection (MOIs), i.e., numbers of per-

episode genotypes / vertices.

igraph Logical; returns RGs as igraph objects (default TRUE).

progress.bar Logical; show progress bar (default TRUE).

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Details

RGs are generated by enumerating nested set partitions under specific constraints; see Understand graph and partition enumeration. Each nested set partition is an RG. Clone edges induce a cluster graph, equivalent to a partition of genotypes, with no intra-episode clones allowed. Sibling edges refine the clone partition, with no constraints (intra-episode siblings allowed). Each nested set partition is encoded as a list. Each partition is represented by a list of vectors (either clone or sib) and by a membership vector (either clone.vec or sib.vec). By default, an RG encoded as a list is converted to an igraph object.

Value

A list of RGs. If igraph = FALSE, each RG is a list of length four with:

clone List of vectors encoding genotypes per clonal cell.

clone.vec Numeric vector with the clonal cell membership of each genotype.

sib List of vectors encoding clonal cells per sibling cell.

sib.vec Numeric vector with the sibling cell membership of each clonal cell.

If igraph = TRUE (default), each RG is encoded as an igraph object (see RG_to_igraph).

Examples

```
graphs <- enumerate_RGs(c(2, 1, 2), progress.bar = FALSE) # nine graphs</pre>
```

fs_VHX_BPD

Allele frequencies computed using example Plasmodium vivax data

Description

The posterior mean of a multinomial-Dirichlet model with uniform prior fit to data on allele prevalence in initial episodes of ys_VHX_BPD. Because the model is fit to allele prevalence (observed) not allele frequency (requires integrating-out unknown multiplicities of infection) it is liable to underestimate the frequencies of common alleles and overestimate those of rare but detected alleles.

Usage

fs_VHX_BPD

Format

A list of nine markers; for each marker a named vector of allele frequencies that sum to one.

Source

- MS_data_PooledAnalysis.RData downloaded from https://zenodo.org/records/3368828
- https://github.com/aimeertaylor/Pv3Rs/blob/main/data-raw/fs_VHX_BPD.R

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plot_data

Plots the data

Description

Plots allelic data as a grid of coloured rectangles.

Usage

```
plot_data(
  ys,
  fs = NULL,
  person.vert = FALSE,
 mar = c(1.5, 3.5, 1.5, 1),
  gridlines = TRUE,
 palette = RColorBrewer::brewer.pal(12, "Paired"),
 marker.annotate = TRUE,
  legend.lab = "Allele frequencies",
  legend.line = 0.2,
  legend.ylim = c(0.05, 0.2),
  cex.maj = 0.7,
  cex.min = 0.5,
  cex.text = 0.5,
  x.line = 0.2,
 y.line = 2.5
)
```

Arguments

fs

ys	Nested list of per-person, per-episode, per-marker allelic data; see Examples
	and compute_posterior() for the expected per-person structure.

A per-marker list of numeric vectors of allele frequencies. If NULL (default), only the alleles present in ys are shown in the the legend, with all per-marker alleles represented equally. Because the colour scheme is adaptive, the same allele may have different colours given different ys. If fs is specified, all alleles in fs feature in the legend with areas proportional to allele frequencies, so that common alleles occupy larger areas than rarer alleles. Specify fs to fix the allele colour scheme across plots of different ys.

colour scheme across plots of different ys.

person.vert Logical. If TRUE (default), person IDs are printed vertically; otherwise, they are

printed horizontally.

mar Vector of numbers of lines of margin for the main plot; see mar entry of par.

gridlines Logical. If true (default), white grid lines separating people and markers are

drawn.

palette Colour palette for alleles, see the **Value** section of brewer.pal. Generally, colours are interpolated: if a marker has d possible alleles, then the colours used

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	are the $1/(d+1)$,, $d/(d+1)$ quantiles of the palette to ensure that markers with different allele counts use different colours.			
marker.annotate				
	Logical. If true (default), the names of the alleles are printed on top of their colours in the legend.			
legend.lab	Label for the axis of the legend. Defaults to "Allele frequencies". Set to NA to omit the label; if so, consider adjusting legend. ylim to use more plotting space.			
legend.line	Distance (in character heights) from the colour bar to the legend label (defaults to 1.5).			
legend.ylim	Vector specifying the y-coordinate limits of the legend in device coordinates (between 0 and 1). Defaults to $c(0.05,0.2)$.			
cex.maj	Numeric; font scaling of major axis labels.			
cex.min	Numeric; font scaling of minor axis labels.			
cex.text	Numeric; font scaling of the allele labels.			
x.line	Distance between top x-axis and x-axis label, defaults to 0.2.			
y.line	Distance between left y-axis and y-axis label, defaults to 2.5.			

Details

This function plots alleles (colours), which are observed in different episodes (columns), on different markers (rows), with episodes grouped by person. Per-person episodes are plotted from left to right in chronological order. If multiple alleles are detected for a marker within an episode, the corresponding grid element is subdivided vertically into different colours.

By default, markers are ordered lexicographically. If fs is provided, markers are ordered to match the order within fs.

The legend depicts the alleles for each marker in the same vertical order as the main plot. The default colour scheme is adaptive, designed to visually differentiate the alleles as clearly as possible by maximizing hue contrast within a qualitative palette. Interpolation is used to make different colour palettes for markers with different numbers of possible alleles. The names of the alleles are printed on top of their colours if marker.annotate is set to TRUE.

Value

None

Examples

```
oldpar <- par(no.readonly = TRUE) # Store user's options before plotting

# Running example (runs across compute_posterior, plot_data and plot_simplex)
# based on real data from chloroquine-treated participant 52 of the Vivax
# History Trial (Chu et al. 2018a, https://doi.org/10.1093/cid/ciy319)
ys <- ys_VHX_BPD["VHX_52"] # ys is a list of length one (one participant)
plot_data(ys, fs = fs_VHX_BPD, marker.annotate = FALSE)

# Full data set:
mar <- c(2, 3.5, 1.5, 1) # extra vertical margin for vertical person labels</pre>
```

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plot_RG

Plot a relationship graph (RG)

Description

This function is a wrapper around plot.igraph, written to group parasite genotypes by episode both spatially and using vertex colour (specifically, parasite genotypes within episodes are vertically distributed with some horizontal jitter when layout.by.group = TRUE (default), and equicolored), and to ensure clone and sibling edges are plotted using different line types.

Usage

```
plot_RG(
   RG,
   layout.by.group = TRUE,
   vertex.palette = "Set2",
   edge.lty = c(sibling = "dashed", clone = "solid"),
   edge.col = c(sibling = "black", clone = "black"),
   edge.width = 1.5,
   ...
)
```

Arguments

RG igraph object encoding an RG; see RG_to_igraph .

layout.by.group

Logical; if TRUE (default) overrides the default layout of plot.igraph so that vertices that represent parasite genotypes from different episodes are distributed horizontally and vertices that represent genotypes within episodes are distributed vertically.

vertex.palette A character string specifying an RColorBrewer palette. Overrides the default palette of plot.igraph.

edge.lty Named vector of edge line types corresponding to different relationships.

edge.col Named vector of edge colours corresponding to different relationships.

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```
edge.width Overrides the default edge.width of plot.igraph.... Additional arguments to pass to plot.igraph, e.g., edge.curved.
```

Details

To see how to plot relationship graphs outputted by compute_posterior, please refer to Exploration of relationship graphs in Demonstrate Pv3Rs usage.

Value

None

Provenance

This function was adapted from plot_Vivax_model at https://github.com/jwatowatson/RecurrentVivax/blob/master/Genetic_Model/iGraph_functions.R.

Examples

```
RGs <- enumerate_RGs(c(2, 1, 1), progress.bar = FALSE)
oldpar <- par(no.readonly = TRUE) # record user's options
par(mfrow = c(3, 4), mar = c(0.1, 0.1, 0.1, 0.1))
for (i in 12:23) {
   plot_RG(RGs[[i]],
   edge.col = c(sibling = "gray", clone = "black"),
   edge.lty = c(sibling = "dotted", clone = "solid"),
   edge.curved = 0.1)
   box()
}
par(oldpar) # restore user's options</pre>
```

plot_simplex

Plots a 2D simplex

Description

Plots a 2D simplex (a triangle with unit sides centered at the origin) onto which per-recurrence posterior probabilities of recrudescence, relapse, reinfection (or any other probability triplet summing to one) can be projected; see project2D() and **Examples** below.

Usage

```
plot_simplex(
  v.labels = c("Recrudescence", "Relapse", "Reinfection"),
  v.cutoff = 0.5,
  v.colours = c("yellow", "purple", "red"),
  plot.tri = TRUE,
  lim.mar = 0.1,
```

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```
p.coords = NULL,
p.labels = rownames(p.coords),
p.labels.pos = 3,
p.labels.cex = 1,
...
)
```

Arguments

v.labels	Vertex labels anticlockwise from top (default: "Recrudescence", "Relapse", "Reinfection"). If NULL, vertices are not labelled.
v.cutoff	Number between 0.5 and 1 that separates lower vs higher probability regions. Use with caution for recrudescence and reinfection classification; see Understand posterior probabilities.
v.colours	Vertex colours anticlockwise from top.
plot.tri	Logical; draws the triangular boundary if TRUE (default).
lim.mar	Margin away from simplex for axes limits; defaults to 0.1.
p.coords	Matrix of 3D simplex coordinates (e.g., per-recurrence probabilities of recrude-scence, relapse and reinfection), one vector of 3D coordinates per row, each row is projected onto 2D coordinates using project2D() and plotted as a single simplex point using graphics::points(). If the user provides a vector encoding a probability triplet summing to one, it is converted to a matrix with one row.
p.labels	Labels of points in p. coords (default row names of p. coords) No labels if NA.
p.labels.pos	Position of p.labels: $1 = below$, $2 = left$, $3 = above$ (default) and $4 = right$. Can be a single value or a vector.
p.labels.cex	Size expansion of p.labels passed to text.
	Additional parameters passed to graphics::points().

Value

None

Examples

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```
# compute the posterior probabilities of the 3Rs and plot the deviation of the
# posterior from the prior
# Some data:
y \leftarrow list(list(m1 = c('a', 'b'), m2 = c('c', 'd')), # Enrollment episode
          list(m1 = c('a'), m2 = c('c'))) # Recurrent episode
# Some allele frequencies:
fs \leftarrow list(m1 = setNames(c(0.4, 0.6), c('a', 'b')),
           m2 = setNames(c(0.2, 0.8), c('c', 'd')))
# A vector of prior probabilities:
prior <- array(c(0.2, 0.3, 0.5), dim = c(1,3),
               dimnames = list(NULL, c("C", "L", "I")))
# Compute posterior probabilities
post <- compute_posterior(y, fs, prior, progress.bar = FALSE)</pre>
# Plot simplex with the prior and posterior
plot_simplex(p.coords = rbind(prior, post$marg),
             p.labels = c("Prior", "Posterior"),
             pch = 20)
# Add the deviation between the prior and posterior: requires obtaining 2D
# coordinates manually
xy_prior <- project2D(as.vector(prior))</pre>
xy_post <- project2D(as.vector(post$marg))</pre>
arrows(x0 = xy_prior["x"], x1 = xy_post["x"],
       y0 = xy_prior["y"], y1 = xy_post["y"], length = 0.1)
```

project2D

Project 3D probability coordinates onto 2D simplex coordinates

Description

Project three probabilities that sum to one (e.g., per-recurrence probabilities of recrudescence, relapse and reinfection) onto the coordinates of a 2D simplex centred at the origin (i.e., a triangle centred at (0,0) with unit-length sides).

Usage

```
project2D(v)
```

Arguments

٧

A numeric vector of three numbers in zero to one that sum to one.

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Details

The top, left, and right vertices of the 2D simplex correspond with the first, second and third entries of v, respectively. Each probability is proportional to the distance from the point on the simplex to the side opposite the corresponding probability; see **Examples** below and plot_simplex() for more details.

Value

A numeric vector of two coordinates that can be used to plot the probability vector v on the origincentred 2D simplex.

Examples

```
probabilities_of_v1_v2_v3 <- c(0.75,0.20,0.05)
coordinates <- project2D(v = probabilities_of_v1_v2_v3)

# Plot probability vector on 2D simplex
plot_simplex(v.labels = c("v1", "v2", "v3"))
points(x = coordinates[1], y = coordinates[2], pch = 20)

# Plot the distances that represent probabilities
# get vertices, get points on edges by orthogonal projection, plot arrows
v <- apply(matrix(c(1,0,0,0,1,0,0,0,1), nrow = 3), 1, project2D)
p3 <- v[,1] + sum((coordinates - v[,1]) * (v[,2] - v[,1])) * (v[,2] - v[,1])
p1 <- v[,2] + sum((coordinates - v[,2]) * (v[,3] - v[,2])) * (v[,3] - v[,2])
p2 <- v[,3] + sum((coordinates - v[,3]) * (v[,1] - v[,3])) * (v[,1] - v[,3])
arrows(x0 = coordinates[1], y0 = coordinates[2], x1 = p1[1], y1 = p1[2], length = 0.1)
arrows(x0 = coordinates[1], y0 = coordinates[2], x1 = p3[1], y1 = p3[2], length = 0.1)</pre>
```

RG_to_igraph

Converts a relationship graph (RG) encoded as a list to an igraph object

Description

Converts an RG encoded as a list to an igraph object, which requires more memory allocation but can be plotted using plot_RG.

Usage

```
RG_to_igraph(RG, MOIs)
```

Arguments

RG List encoding an RG; see **Value** of enumerate_RGs when igraph = FALSE.

MOTS Vector of per-episode multiplicities of infection (MOIs), i.e., numbers of

Vector of per-episode multiplicities of infection (MOIs), i.e., numbers of per-episode genotypes / vertices; adds to the graph an attribute that is used by plot_RG to group genotypes / vertices by episode.

sample_RG

Value

A weighted graph whose edge weights 1 and 0.5 encode clonal and sibling relationships, respectively.

Examples

```
MOIs <- c(3,2)
set.seed(6)
RG_as_list <- sample_RG(MOIs, igraph = FALSE)
RG_as_igraph <- RG_to_igraph(RG_as_list, MOIs)
# RG encoded as a list requires less memory allocation
utils::object.size(RG_as_list)
utils::object.size(RG_as_igraph)
# RG encoded as an igraph object can be plotted using plot_RG() and
# manipulated using igraph functions
plot_RG(RG_as_igraph, margin = rep(0,4), vertex.label = NA)
# Edge weights 1 and 0.5 encode clonal and sibling relationships
igraph::E(RG_as_igraph)$weight
# Vertex attribute group encodes episode membership
igraph::V(RG_as_igraph)$group</pre>
```

sample_RG

Sample a relationship graph (RG)

Description

Uses the techniques in <code>enumerate_RGs</code> to sample a single RG uniformly. All clonal partitions are generated, each weighted by its number of consistent sibling partitions. A clonal partition is sampled proportional to its weight, then a consistent sibling partition is drawn uniformly. The resulting nested partition represents the RG; see <code>enumerate_RGs</code> for details.

Usage

```
sample_RG(MOIs, igraph = TRUE)
```

Arguments

MOIs Vector of per-episode multiplicities of infection (MOIs), i.e., numbers of per-

episode genotypes / vertices.

igraph Logical; if TRUE (default), returns the RG as an igraph object.

Value

An RG encoded either as an igraph object (default), or as a list; see enumerate_RGs for details.

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Examples

```
set.seed(1)
RG <- sample_RG(c(3, 2))
plot_RG(RG, vertex.label = NA)</pre>
```

ys_VHX_BPD

Example Plasmodium vivax data

Description

Previously-published microsatellite data on *P. vivax* parasites extracted from study participants enrolled in the Best Primaquine Dose (BPD) and Vivax History (VHX) trials; see Taylor & Watson et al. 2019 (doi:10.1038/s4146701913412x) for more details of the genetic data; for more details of the VHX and BPD trials, see Chu et al. 2018a (doi:10.1093/cid/ciy319) and Chu et al. 2018b (doi:10.1093/cid/ciy735).

Usage

```
ys_VHX_BPD
```

Format

A list of 217 study participants; for each study participant, a list of one or more episodes; for each episode, a list of three or more microsatellite markers; for each marker, a vector of observed alleles (repeat lengths). For example:

BPD_103 Study participant identifier: study participant 103 in the BPD trial

BPD_103_1 Episode identifier: episode one of study participant 103 in the BPD trial

PV.3.27 Marker identifier: *P. vivax* 3.27 **18** Allele identifier: 18 repeat lengths

Source

- MS_data_PooledAnalysis.RData downloaded from https://zenodo.org/records/3368828
- https://github.com/aimeertaylor/Pv3Rs/blob/main/data-raw/ys_VHX_BPD.R

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