# Package 'bumblebee'

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Title Quantify Disease Transmission Within and Between Population Groups

Version 0.1.0

**Description** A simple tool to quantify the amount of transmission of an infectious disease of interest occurring within and between population groups. 'bumblebee' uses counts of observed directed transmission pairs, identified phylogenetically from deep-sequence data or from epidemiological contacts, to quantify transmission flows within and between population groups accounting for sampling heterogeneity. Population groups might include: geographical areas (e.g. communities, regions), demographic groups (e.g. age, gender) or arms of a randomized clinical trial. See the 'bumblebee' website for statistical theory, documentation and examples <https://magosil86.github.io/bumblebee/>.

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URL https://magosil86.github.io/bumblebee/

BugReports https://github.com/magosil86/bumblebee/issues

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counts\_hiv\_transmission\_pairs

Observed HIV transmission pairs

# Description

Counts of directed HIV transmission pairs observed within and between intervention and control communities in the 30-community BCPP/Ya Tsie HIV prevention trial in Botswana (2013-2018). The Botswana -Ya Tsie trial was a pair-matched community randomized trial that evaluated the effect of a universal HIV test and treat intervention in reducing population-level incidence. For further details see references and: https://magosil86.github.io/bumblebee/.

# Usage

counts\_hiv\_transmission\_pairs

# Format

A data frame:

H1\_group Name of population group 1

H2\_group Name of population group 1

num\_linked\_pairs\_observed Number of observed directed transmission pairs between samples from population groups 1 and 2

# Source

https://magosil86.github.io/bumblebee/

#### References

Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial – BCPP/ Ya Tsie trial. To submit for publication, 2021.

#### Description

Estimated HIV transmissions within and betweeen intervention and control communities in the BCPP/Ya Tsie trial population adjusted for variability in sampling.

#### Usage

estimated\_hiv\_transmission\_flows

# Format

A data frame:

H1\_group Name of population group 1

H2\_group Name of population group 2

number\_hosts\_sampled\_group\_1 Number of individuals sampled from population group 1

**number\_hosts\_sampled\_group\_2** Number of individuals sampled from population group 2

**number\_hosts\_population\_group\_1** Estimated number of individuals in population group 1

number\_hosts\_population\_group\_2 Estimated number of individuals in population group 2

- max\_possible\_pairs\_in\_sample Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed Number of observed directed transmission pairs between samples from population groups 1 and 2
- p\_hat Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked
- est\_linkedpairs\_in\_population Estimated transmission pairs between population groups 1 and 2
- **theta\_hat** Estimated transmission flows or relative probability of transmission within and between population groups 1 and 2 adjusted for sampling heterogeneity. More precisely, the conditional probability that a pair of pathogen sequences is from a specific population group pairing given that the pair is linked.
- **obs\_trm\_pairs\_est\_goodman** Point estimate, Goodman method Confidence intervals for observed transmission pairs

obs\_trm\_pairs\_lwr\_ci\_goodman Lower bound of Goodman confidence interval

- obs\_trm\_pairs\_upr\_ci\_goodman Upper bound of Goodman confidence interval
- est\_goodman Point estimate, Goodman method Confidence intervals for estimated transmission flows
- lwr\_ci\_goodman Lower bound of Goodman confidence interval
- upr\_ci\_goodman Upper bound of Goodman confidence interval
- prob\_group\_pairing\_and\_linked Probability that a pair of pathogen sequences is from a specific
  population group pairing and is linked
- **c\_hat** Probability that a randomly selected pathogen sequence in one population group links to at least one pathogen sequence in another population group i.e. probability of clustering
- est\_goodman\_cc Point estimate, Goodman method Confidence intervals with continuity correction
- lwr\_ci\_goodman\_cc Lower bound of Goodman confidence interval
- upr\_ci\_goodman\_cc Upper bound of Goodman confidence interval

est\_sisonglaz Point estimate, Sison-Glaz method Confidence intervals

- lwr\_ci\_sisonglaz Lower bound of Sison-Glaz confidence interval
- upr\_ci\_sisonglaz Upper bound of Sison-Glaz confidence interval
- est\_qhurst\_acswr Point estimate, Queensbury-Hurst method Confidence intervals via ACSWR r package
- lwr\_ci\_qhurst\_acswr Lower bound of Queensbury-Hurst confidence interval
- upr\_ci\_qhurst\_acswr Upper bound of Queensbury-Hurst confidence interval
- est\_qhurst\_coinmind Point estimate, Queensbury-Hurst method Confidence intervals via Coin-MinD r package
- lwr\_ci\_qhurst\_coinmind Lower bound of Queensbury-Hurst confidence interval
- upr\_ci\_qhurst\_coinmind Upper bound of Queensbury-Hurst confidence interval
- lwr\_ci\_qhurst\_adj\_coinmind Lower bound of Queensbury-Hurst confidence interval adjusted
- upr\_ci\_qhurst\_adj\_coinmind Upper bound of Queensbury-Hurst confidence interval adjusted

#### Source

https://magosil86.github.io/bumblebee/

#### References

Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial – BCPP/ Ya Tsie trial. To submit for publication, 2021.

estimate\_c\_hat

# Description

This function estimates c\_hat, the probability that a randomly selected pathogen sequence in one population group links to at least one pathogen sequence in another population group.

#### Usage

```
estimate_c_hat(df_counts_and_p_hat, ...)
## Default S3 method:
estimate_c_hat(df_counts_and_p_hat, ...)
```

## Arguments

# Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2
- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2
- p\_hat, Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked
- c\_hat, Probability that a randomly selected pathogen sequence in one population group links to at least one pathogen sequence in another population group i.e. probability of clustering

#### Methods (by class)

• default: Estimates probability of clustering

#### References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- Carnegie, N.B., et al., Linkage of viral sequences among HIV-infected village residents in Botswana: estimation of linkage rates in the presence of missing data. PLoS Computational Biology, 2014. 10(1): p. e1003430.

# See Also

See estimate\_p\_hat to prepare input data to estimate c\_hat

# Examples

library(bumblebee)
library(dplyr)

# Estimate the probability of clustering between individuals from two population groups of interest

# We shall use the data of HIV transmissions within and between intervention and control # communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data # ?counts\_hiv\_transmission\_pairs, ?sampling\_frequency and ?estimated\_hiv\_transmission\_flows

```
# Load and view data
#
# The input data comprises counts of observed directed HIV transmission pairs within and
# between intervention and control communities in the BCPP/Ya Tsie trial, sampling
# information and the probability of linkage between individuals sampled from
# intervention and control communities (i.e. \code{p_hat})
#
# See ?estimate_p_hat() for details on estimating p_hat
results_estimate_p_hat <- estimated_hiv_transmission_flows[, c(1:10)]
results_estimate_p_hat
# Estimate c_hat
# Estimate_c_hat <- estimate_c_hat(df_counts_and_p_hat = results_estimate_p_hat)
# View results
results_estimate_c_hat</pre>
```

estimate\_multinom\_ci estimate\_multinom\_ci Estimates confidence intervals for transmission flows

# Description

This function computes simultaneous confidence intervals at the 5% significance level for estimated transmission flows. Available methods for computing confidence intervals are: Goodman, Goodman with a continuity correction, Sison-Glaz and Queensbury-Hurst.

#### Usage

```
estimate_multinom_ci(df_theta_hat, ...)
```

```
## Default S3 method:
estimate_multinom_ci(df_theta_hat, detailed_report = FALSE, ...)
```

#### Arguments

df_theta_hat	A data.frame returned by the function: <pre>estimate_theta_hat()</pre>
	Further arguments.
detailed_report	

A boolean value to produce detailed output of the analysis. (Default is FALSE)

#### Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2
- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2
- p\_hat, Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked
- est\_linkedpairs\_in\_population, Estimated transmission pairs between population groups 1 and 2
- theta\_hat, Estimated transmission flows or relative probability of transmission within and between population groups 1 and 2 adjusted for sampling heterogeneity. More precisely, the conditional probability that a pair of pathogen sequences is from a specific population group pairing given that the pair is linked.
- obs\_trm\_pairs\_est\_goodman, Point estimate, Goodman method Confidence intervals for observed transmission pairs

- obs\_trm\_pairs\_lwr\_ci\_goodman, Lower bound of Goodman confidence interval
- obs\_trm\_pairs\_upr\_ci\_goodman, Upper bound of Goodman confidence interval
- est\_goodman, Point estimate, Goodman method Confidence intervals for estimated transmission flows
- lwr\_ci\_goodman, Lower bound of Goodman confidence interval
- upr\_ci\_goodman, Upper bound of Goodman confidence interval

The following additional fields are returned if the detailed\_report flag is set

- est\_goodman\_cc, Point estimate, Goodman method Confidence intervals with continuity correction
- lwr\_ci\_goodman\_cc, Lower bound of Goodman confidence interval
- upr\_ci\_goodman\_cc, Upper bound of Goodman confidence interval
- est\_sisonglaz, Point estimate, Sison-Glaz method Confidence intervals
- · lwr\_ci\_sisonglaz, Lower bound of Sison-Glaz confidence interval
- upr\_ci\_sisonglaz, Upper bound of Sison-Glaz confidence interval
- est\_qhurst\_acswr, Point estimate, Queensbury-Hurst method Confidence intervals via AC-SWR r package
- · lwr\_ci\_qhurst\_acswr, Lower bound of Queensbury-Hurst confidence interval
- upr\_ci\_qhurst\_acswr, Upper bound of Queensbury-Hurst confidence interval
- est\_qhurst\_coinmind, Point estimate, Queensbury-Hurst method Confidence intervals via Coin-MinD r package
- lwr\_ci\_qhurst\_coinmind, Lower bound of Queensbury-Hurst confidence interval
- upr\_ci\_qhurst\_coinmind, Upper bound of Queensbury-Hurst confidence interval
- lwr\_ci\_qhurst\_adj\_coinmind, Lower bound of Queensbury-Hurst confidence interval adjusted
- upr\_ci\_qhurst\_adj\_coinmind, Upper bound of Queensbury-Hurst confidence interval adjusted

#### Methods (by class)

• default: Estimates confidence intervals for transmission flows

# References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- Goodman, L. A. On Simultaneous Confidence Intervals for Multinomial Proportions Technometrics, 1965. 7, 247-254.
- 3. Cherry, S., A Comparison of Confidence Interval Methods for Habitat Use-Availability Studies. The Journal of Wildlife Management, 1996. 60(3): p. 653-658.
- Sison, C.P and Glaz, J. Simultaneous confidence intervals and sample size determination for multinomial proportions. Journal of the American Statistical Association, 1995. 90:366-369.

- 5. Glaz, J., Sison, C.P. Simultaneous confidence intervals for multinomial proportions. Journal of Statistical Planning and Inference, 1999. 82:251-262.
- May, W.L., Johnson, W.D. Constructing two-sided simultaneous confidence intervals for multinomial proportions for small counts in a large number of cells. Journal of Statistical Software, 2000. 5(6). Paper and code available at https://www.jstatsoft.org/v05/i06.
- Carnegie, N.B., et al., Linkage of viral sequences among HIV-infected village residents in Botswana: estimation of linkage rates in the presence of missing data. PLoS Computational Biology, 2014. 10(1): p. e1003430.
- 8. Ratmann, O., et al., Inferring HIV-1 transmission networks and sources of epidemic spread in Africa with deep-sequence phylogenetic analysis. Nature Communications, 2019. 10(1): p. 1411.
- Wymant, C., et al., PHYLOSCANNER: Inferring Transmission from Within- and Between-Host Pathogen Genetic Diversity. Molecular Biology and Evolution, 2017. 35(3): p. 719-733.

#### See Also

See estimate\_theta\_hat to prepare input data to estimate confidence intervals.

To learn more about the Goodman and Sison-Glaz confidence interval methods see \code{\link[DescTools]{MultinomCI}}. For Queensbury-Hurst confidence intervals see \code{\link[ACSWR]{QH\_CI}} and \code{\link[CoinMinD]{QH}}

#### Examples

```
library(bumblebee)
library(dplyr)
```

- # Compute confidence intervals for estimated transmission flows
- # We shall use the data of HIV transmissions within and between intervention and control # communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data
- # ?counts\_hiv\_transmission\_pairs and ?sampling\_frequency

```
# Load and view data
```

#

# The data comprises counts of observed directed HIV transmission pairs between individuals

# sampled from intervention and control communities (i.e. num\_linked\_pairs\_observed);

```
\ensuremath{\texttt{\#}} and the estimated HIV transmissions within and between intervention and control
```

# communities in the BCPP/Ya Tsie trial population adjusted for sampling heterogneity

```
\# (i.e. \code{est_linkedpairs_in_population}). See ?estimate_theta_hat() for details on
```

```
# computing \code{est_linkedpairs_in_population} and \code{theta_hat}.
```

results\_estimate\_theta\_hat <- estimated\_hiv\_transmission\_flows[, c(1:13)]</pre>

```
results_estimate_theta_hat
```

```
# Compute Goodman confidence intervals (Default)
results_estimate_multinom_ci <- estimate_multinom_ci(
    df_theta_hat = results_estimate_theta_hat,</pre>
```

```
detailed_report = FALSE)
# View results
results_estimate_multinom_ci
# Compute Goodman, Sison-Glaz and Queensbury-Hurst confidence intervals
results_estimate_multinom_ci_detailed <- estimate_multinom_ci(
    df_theta_hat = results_estimate_theta_hat,
    detailed_report = TRUE)
# View results
results_estimate_multinom_ci_detailed</pre>
```

estimate\_prob\_group\_pairing\_and\_linked estimate\_prob\_group\_pairing\_and\_linked Estimates joint prob-ability of linkage

# Description

This function computes the joint probability that a pair of pathogen sequences is from a specific population group pairing and linked.

#### Usage

```
estimate_prob_group_pairing_and_linked(
    df_counts_and_p_hat,
    individuals_population_in,
    ...
)
## Default S3 method:
estimate_prob_group_pairing_and_linked(
    df_counts_and_p_hat,
    individuals_population_in,
    verbose_output = FALSE,
    ...
)
```

#### Arguments

#### Details

For a population group pairing (u, v), the joint probability that a pair is from groups (u, v) and is linked is computed as

$$(N_u v/N_c hoose_2) * p_h at_u v,$$

where,

- $N_u = N_u * N_v$ : maximum distinct possible (u, v) pairs in population
- p\_hat\_uv: probability of linkage between two individuals randomly sampled from groups u and v
- N choose 2 or (N \* (N 1))/2: all distinct possible pairs in population.

See bumblebee website for more details https://magosil86.github.io/bumblebee/.

# Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2
- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2
- p\_hat, Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked
- prob\_group\_pairing\_and\_linked, Probability that a pair of pathogen sequences is from a specific population group pairing and is linked

#### Methods (by class)

default: Estimates joint probability of linkage

#### References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- Carnegie, N.B., et al., Linkage of viral sequences among HIV-infected village residents in Botswana: estimation of linkage rates in the presence of missing data. PLoS Computational Biology, 2014. 10(1): p. e1003430.

# See Also

See estimate\_p\_hat to prepare input data to estimate prob\_group\_pairing\_and\_linked

#### Examples

```
library(bumblebee)
library(dplyr)
# Estimate joint probability that a pair is from a specific group pairing and linked
# We shall use the data of HIV transmissions within and between intervention and control
# communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data
# ?counts_hiv_transmission_pairs and ?sampling_frequency
# Load and view data
# The input data comprises counts of observed directed HIV transmission pairs
# within and between intervention and control communities in the BCPP/Ya Tsie
# trial, sampling information and the probability of linkage between individuals
# sampled from intervention and control communities (i.e. \code{p_hat})
#
# See ?estimate_p_hat() for details on estimating p_hat
results_estimate_p_hat <- estimated_hiv_transmission_flows[, c(1:10)]</pre>
results_estimate_p_hat
# Estimate prob_group_pairing_and_linked
results_prob_group_pairing_and_linked <- estimate_prob_group_pairing_and_linked(
    df_counts_and_p_hat = results_estimate_p_hat,
    individuals_population_in = sampling_frequency$number_population)
# View results
results_prob_group_pairing_and_linked
```

estimate_p_hat	estimate_p_hat Estimates probability of linkage between two indi-
	viduals

# Description

This function computes the probability that pathogen sequences from two individuals randomly sampled from their respective population groups (e.g. communities) are linked.

#### Usage

```
estimate_p_hat(df_counts, ...)
## Default S3 method:
estimate_p_hat(df_counts, ...)
```

#### Arguments

df_counts	A data.frame returned by the function: prep_p_hat()
	Further arguments.

## Details

For a population group pairing (u, v), p\_hat is computed as the fraction of distinct possible pairs between samples from groups u and v that are linked. Note: The number of distinct possible (u, v)pairs in the sample is the product of sampled individuals in groups u and u. If u = v, then the distinct possible pairs is the number of individuals sampled in population group u choose 2. See bumblebee website for more details https://magosil86.github.io/bumblebee/.

# Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2
- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2
- p\_hat, Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked

## Methods (by class)

• default: Estimates probability of linkage between two individuals

# References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- Carnegie, N.B., et al., Linkage of viral sequences among HIV-infected village residents in Botswana: estimation of linkage rates in the presence of missing data. PLoS Computational Biology, 2014. 10(1): p. e1003430.

# See Also

See prep\_p\_hat to prepare input data to estimate p\_hat

# Examples

```
library(bumblebee)
library(dplyr)
# Estimate the probability of linkage between two individuals randomly sampled from
# two population groups of interest.
# We shall use the data of HIV transmissions within and between intervention and control
# communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data
# ?counts_hiv_transmission_pairs and ?sampling_frequency
# Prepare input to estimate p_hat
# View counts of observed directed HIV transmissions within and between intervention
# and control communities
counts_hiv_transmission_pairs
# View the estimated number of individuals with HIV in intervention and control
# communities and the number of individuals sampled from each
sampling_frequency
results_prep_p_hat <- prep_p_hat(group_in = sampling_frequency$population_group,</pre>
                             individuals_sampled_in = sampling_frequency$number_sampled,
                       individuals_population_in = sampling_frequency$number_population,
                                 linkage_counts_in = counts_hiv_transmission_pairs,
                                 verbose_output = FALSE)
# View results
results_prep_p_hat
# Estimate p_hat
results_estimate_p_hat <- estimate_p_hat(df_counts = results_prep_p_hat)
# View results
results_estimate_p_hat
```

estimate_theta_hat	estimate_theta_hat Estimates	conditional probability of	of linkage
	(transmission flows)		

#### Description

This function estimates theta\_hat, the relative probability of transmission within and between population groups accounting for variable sampling rates among population groups. This relative probability is also refferred to as transmission flows.

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estimate\_theta\_hat

# Usage

```
estimate_theta_hat(df_counts_and_p_hat, ...)
```

## Default S3 method: estimate\_theta\_hat(df\_counts\_and\_p\_hat, ...)

Further arguments.

#### Arguments

df\_counts\_and\_p\_hat

A data.frame returned by the function: estimate\_p\_hat()

. . .

# Details

For a population group pairing (u, v), the estimated transmission flows within and between population groups u and v, are represented by the vector theta\_hat,

$$\hat{\theta} = (\hat{\theta}_{uu}, \hat{\theta}_{uv}, \hat{\theta}_{vu}, \hat{\theta}_{vv}),$$

and are computed as

$$\hat{\theta_{ij}} = \frac{N_{ij}p_{ij}}{\sum_{m}\sum_{n\geq m}N_{mn}p_{mn}}, where i = u, vand j = u, v,$$

See bumblebee website for more details https://magosil86.github.io/bumblebee/.

#### Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2
- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2

- p\_hat, Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked
- est\_linkedpairs\_in\_population, Estimated transmission pairs between population groups 1 and 2
- theta\_hat, Estimated transmission flows or relative probability of transmission within and between population groups 1 and 2 adjusted for sampling heterogeneity. More precisely, the conditional probability that a pair of pathogen sequences is from a specific population group pairing given that the pair is linked.

# Methods (by class)

• default: Estimates conditional probability of linkage (transmission flows)

#### References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- Carnegie, N.B., et al., Linkage of viral sequences among HIV-infected village residents in Botswana: estimation of linkage rates in the presence of missing data. PLoS Computational Biology, 2014. 10(1): p. e1003430.

#### See Also

See estimate\_p\_hat to prepare input data to estimate theta\_hat

# Examples

```
library(bumblebee)
library(dplyr)
```

# Estimate transmission flows within and between population groups accounting for variable
# sampling among population groups

# We shall use the data of HIV transmissions within and between intervention and control # communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data

```
# ?counts_hiv_transmission_pairs and ?sampling_frequency
```

```
# Load and view data
#
# The input data comprises counts of observed directed HIV transmission pairs within
# and between intervention and control communities in the BCPP/Ya Tsie trial,
# sampling information and the probability of linkage between individuals sampled
# from intervention and control communities (i.e. \code{p_hat})
#
# See ?estimate_p_hat() for details on estimating p_hat
results_estimate_p_hat <- estimated_hiv_transmission_flows[, c(1:10)]</pre>
```

```
results_estimate_p_hat
```

```
# Estimate theta_hat
results_estimate_theta_hat <- estimate_theta_hat(df_counts_and_p_hat = results_estimate_p_hat)</pre>
```

# View results
results\_estimate\_theta\_hat

estimate\_transmission\_flows\_and\_ci

estimate\_transmission\_flows\_and\_ci *Estimates transmission flows and corresponding confidence intervals* 

# Description

This function estimates transmission flows or the relative probability of transmission within and between population groups accounting for variable sampling among population groups.

Corresponding confidence intervals are provided with the following methods: Goodman, Goodman with a continuity correction, Sison-Glaz and Queensbury-Hurst.

#### Usage

```
estimate_transmission_flows_and_ci(
  group_in,
  individuals_sampled_in,
  individuals_population_in,
  linkage_counts_in,
  . . .
)
## Default S3 method:
estimate_transmission_flows_and_ci(
  group_in,
  individuals_sampled_in,
  individuals_population_in,
  linkage_counts_in,
  detailed_report = FALSE,
  verbose_output = FALSE,
)
```

# Arguments

group\_in,

A character vector indicating population groups/strata (e.g. communities, agegroups, genders or trial arms) between which transmission flows will be evaluated, individuals\_sampled\_in,

A numeric vector indicating the number of individuals sampled per population group,

individuals\_population\_in,

A numeric vector of the estimated number of individuals per population group,

linkage\_counts\_in

A data.frame of counts of linked pairs identified between samples of each population group pairing of interest.

The data.frame should contain the following three fields:

- H1\_group (character) Name of population group 1
- H2\_group (character) Name of population group 2
- number\_linked\_pairs\_observed (numeric) Number of observed directed transmission pairs between samples from population groups 1 and 2

Further arguments.

detailed\_report,

A boolean value to produce detailed output of the analysis

verbose\_output,

A boolean value to display intermediate output (Default is FALSE)

# Details

Counts of observed directed transmission pairs can be obtained from deep-sequence phylogenetic data (via phyloscanner) or from known epidemiological contacts. Note: Deep-sequence data is also commonly referred to as high-throughput or next-generation sequence data. See references to learn more about phyloscanner.

The estimate\_transmission\_flows\_and\_ci() function is a wrapper function that calls the following functions:

- 1. The prep\_p\_hat() function to determine all possible combinations of the population groups/strata provided by the user. Type ?prep\_p\_hat() at R prompt to learn more.
- The estimate\_p\_hat() function to compute the probability of linkage between pathogen sequences from two individuals randomly sampled from their respective population groups. Type ?estimate\_p\_hat() at R prompt to learn more.
- 3. The estimate\_theta\_hat() function that uses p\_hat estimates to compute the conditional probability of linkage that a pair of pathogen sequences is from a specific population group pairing given that the pair is linked. The conditional probability, theta\_hat represents transmission flows or the relative probability of transmission within and between population groups adjusted for variable sampling among population groups. Type ?estimate\_theta\_hat() at R prompt to learn more.
- 4. The estimate\_multinom\_ci() function to estimate corresponding confidence intervals for the computed transmission flows.

Further to estimating transmission flows and corresponding confidence intervals the estimate\_transmission\_flows\_and\_confidence intervals the estimate for:

 prob\_group\_pairing\_and\_linked, the joint probability that a pair of pathogen sequences is from a specific population group pairing and linked. Type ?estimate\_prob\_group\_pairing\_and\_linked() at R prompt to learn more.  c\_hat, the probability of clustering that a pathogen sequence from a population group of interest is linked to one or more pathogen sequences in another population group of interest. Type ?estimate\_c\_hat() at R prompt to learn more.

# Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2
- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2
- p\_hat, Probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked
- est\_linkedpairs\_in\_population, Estimated transmission pairs between population groups 1 and 2
- theta\_hat, Estimated transmission flows or relative probability of transmission within and between population groups 1 and 2 adjusted for sampling heterogeneity. More precisely, the conditional probability that a pair of pathogen sequences is from a specific population group pairing given that the pair is linked.
- obs\_trm\_pairs\_est\_goodman, Point estimate, Goodman method Confidence intervals for observed transmission pairs
- obs\_trm\_pairs\_lwr\_ci\_goodman, Lower bound of Goodman confidence interval
- obs\_trm\_pairs\_upr\_ci\_goodman, Upper bound of Goodman confidence interval
- est\_goodman, Point estimate, Goodman method Confidence intervals for estimated transmission flows
- lwr\_ci\_goodman, Lower bound of Goodman confidence interval
- upr\_ci\_goodman, Upper bound of Goodman confidence interval

The following additional fields are returned if the detailed\_report flag is set

- prob\_group\_pairing\_and\_linked, Probability that a pair of pathogen sequences is from a specific population group pairing and is linked
- c\_hat, Probability that a randomly selected pathogen sequence in one population group links to at least one pathogen sequence in another population group i.e. probability of clustering

- est\_goodman\_cc, Point estimate, Goodman method Confidence intervals with continuity correction
- lwr\_ci\_goodman\_cc, Lower bound of Goodman confidence interval
- upr\_ci\_goodman\_cc, Upper bound of Goodman confidence interval
- est\_sisonglaz, Point estimate, Sison-Glaz method Confidence intervals
- · lwr\_ci\_sisonglaz, Lower bound of Sison-Glaz confidence interval
- upr\_ci\_sisonglaz, Upper bound of Sison-Glaz confidence interval
- est\_qhurst\_acswr, Point estimate, Queensbury-Hurst method Confidence intervals via AC-SWR r package
- lwr\_ci\_qhurst\_acswr, Lower bound of Queensbury-Hurst confidence interval
- upr\_ci\_qhurst\_acswr, Upper bound of Queensbury-Hurst confidence interval
- est\_qhurst\_coinmind, Point estimate, Queensbury-Hurst method Confidence intervals via Coin-MinD r package
- lwr\_ci\_qhurst\_coinmind, Lower bound of Queensbury-Hurst confidence interval
- upr\_ci\_qhurst\_coinmind, Upper bound of Queensbury-Hurst confidence interval
- lwr\_ci\_qhurst\_adj\_coinmind, Lower bound of Queensbury-Hurst confidence interval adjusted
- upr\_ci\_qhurst\_adj\_coinmind, Upper bound of Queensbury-Hurst confidence interval adjusted

#### Methods (by class)

• default: Estimates transmission flows and accompanying confidence intervals

## References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- Carnegie, N.B., et al., Linkage of viral sequences among HIV-infected village residents in Botswana: estimation of linkage rates in the presence of missing data. PLoS Computational Biology, 2014. 10(1): p. e1003430.
- 3. Cherry, S., A Comparison of Confidence Interval Methods for Habitat Use-Availability Studies. The Journal of Wildlife Management, 1996. 60(3): p. 653-658.
- 4. Ratmann, O., et al., Inferring HIV-1 transmission networks and sources of epidemic spread in Africa with deep-sequence phylogenetic analysis. Nature Communications, 2019. 10(1): p. 1411.
- 5. Wymant, C., et al., PHYLOSCANNER: Inferring Transmission from Within- and Between-Host Pathogen Genetic Diversity. Molecular Biology and Evolution, 2017. 35(3): p. 719-733.
- Goodman, L. A. On Simultaneous Confidence Intervals for Multinomial Proportions Technometrics, 1965. 7, 247-254.
- 7. Sison, C.P and Glaz, J. Simultaneous confidence intervals and sample size determination for multinomial proportions. Journal of the American Statistical Association, 1995. 90:366-369.
- Glaz, J., Sison, C.P. Simultaneous confidence intervals for multinomial proportions. Journal of Statistical Planning and Inference, 1999. 82:251-262.

 May, W.L., Johnson, W.D. Constructing two-sided simultaneous confidence intervals for multinomial proportions for small counts in a large number of cells. Journal of Statistical Software, 2000. 5(6). Paper and code available at https://www.jstatsoft.org/v05/i06.

# See Also

estimate\_theta\_hat and estimate\_multinom\_ci to learn more about estimation of transmission
flows and confidence intervals.

#### Examples

library(bumblebee)
library(dplyr)

# Estimate transmission flows and confidence intervals

# We shall use the data of HIV transmissions within and between intervention and control # communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data # ?counts\_hiv\_transmission\_pairs and ?sampling\_frequency

# View counts of observed directed HIV transmissions within and between intervention
# and control communities
counts\_hiv\_transmission\_pairs

```
# View the estimated number of individuals with HIV in intervention and control
# communities and the number of individuals sampled from each
sampling_frequency
```

# Estimate transmission flows within and between intervention and control communities # accounting for variable sampling among population groups.

```
# Basic output
```

```
results_estimate_transmission_flows_and_ci <- estimate_transmission_flows_and_ci(
    group_in = sampling_frequency$population_group,
    individuals_sampled_in = sampling_frequency$number_sampled,
    individuals_population_in = sampling_frequency$number_population,
    linkage_counts_in = counts_hiv_transmission_pairs)</pre>
```

```
# View results
results_estimate_transmission_flows_and_ci
```

# Retrieve dataset of estimated transmission flows
dframe <- results\_estimate\_transmission\_flows\_and\_ci\$flows\_dataset</pre>

```
# Detailed output
```

```
results_estimate_transmission_flows_and_ci_detailed <- estimate_transmission_flows_and_ci(
    group_in = sampling_frequency$population_group,
    individuals_sampled_in = sampling_frequency$number_sampled,
    individuals_population_in = sampling_frequency$number_population,
    linkage_counts_in = counts_hiv_transmission_pairs,
    detailed_report = TRUE)</pre>
```

```
# View results
results_estimate_transmission_flows_and_ci_detailed
# Retrieve dataset of estimated transmission flows
dframe <- results_estimate_transmission_flows_and_ci_detailed$flows_dataset
# Options:
# To show intermediate output set verbose_output = TRUE
# Basic output
results_estimate_transmission_flows_and_ci <- estimate_transmission_flows_and_ci(
   group_in = sampling_frequency$population_group,
   individuals_sampled_in = sampling_frequency$number_sampled,
   individuals_population_in = sampling_frequency$number_population,
  linkage_counts_in = counts_hiv_transmission_pairs,
  verbose_output = TRUE)
# View results
results_estimate_transmission_flows_and_ci
# Detailed output
results_estimate_transmission_flows_and_ci_detailed <- estimate_transmission_flows_and_ci(
   group_in = sampling_frequency$population_group,
   individuals_sampled_in = sampling_frequency$number_sampled,
   individuals_population_in = sampling_frequency$number_population,
  linkage_counts_in = counts_hiv_transmission_pairs,
  detailed_report = TRUE,
   verbose_output = TRUE)
# View results
results_estimate_transmission_flows_and_ci_detailed
```

```
prep_p_hat
```

prep\_p\_hat Prepares input data to estimate p\_hat

## Description

This function generates variables required for estimating p\_hat, the probability that pathogen sequences from two individuals randomly sampled from their respective population groups are linked. For a population group pairing (u, v), prep\_p\_hat determines all possible group pairings i.e. (uu, uv, vu, vv).

# Usage

```
prep_p_hat(
  group_in,
  individuals_sampled_in,
  individuals_population_in,
  linkage_counts_in,
```

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```
...
)
## Default S3 method:
prep_p_hat(
  group_in,
  individuals_sampled_in,
  individuals_population_in,
  linkage_counts_in,
  verbose_output = FALSE,
  ...
)
```

# Arguments

group_in	A character vector indicating population groups/strata (e.g. communities, age- groups, genders or trial arms) between which transmission flows will be evalu- ated,	
individuals_sampled_in		
	A numeric vector indicating the number of individuals sampled per population	
	group,	
individuals_population_in		
	A numeric vector of the estimated number of individuals per population group,	
linkage_counts_in		
0	A data.frame of counts of linked pairs identified between samples of each pop- ulation group pairing of interest.	
	The data.frame should contain the following three fields:	
	• H1_group (character) Name of population group 1	
	• H2_group (character) Name of population group 2	
	<ul> <li>number_linked_pairs_observed (numeric) Number of observed directed trans mission pairs between samples from population groups 1 and 2</li> </ul>	
	Further arguments.	
verbose_output	A boolean value to display intermediate output. (Default is FALSE)	

# Details

Counts of observed directed transmission pairs can be obtained from deep-sequence phylogenetic data (via phyloscanner) or from known epidemiological contacts. Note: Deep-sequence data is also commonly referred to as high-throughput or next-generation sequence data. See references to learn more about phyloscanner.

# Value

Returns a data.frame containing:

- H1\_group, Name of population group 1
- H2\_group, Name of population group 2

- number\_hosts\_sampled\_group\_1, Number of individuals sampled from population group 1
- number\_hosts\_sampled\_group\_2, Number of individuals sampled from population group 2
- number\_hosts\_population\_group\_1, Estimated number of individuals in population group 1
- number\_hosts\_population\_group\_2, Estimated number of individuals in population group 2
- max\_possible\_pairs\_in\_sample, Number of distinct possible transmission pairs between individuals sampled from population groups 1 and 2
- max\_possible\_pairs\_in\_population, Number of distinct possible transmission pairs between individuals in population groups 1 and 2
- num\_linked\_pairs\_observed, Number of observed directed transmission pairs between samples from population groups 1 and 2

# Methods (by class)

default: Prepares input data to estimate p\_hat

#### References

- 1. Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial BCPP/ Ya Tsie trial. To submit for publication, 2021.
- 2. Ratmann, O., et al., Inferring HIV-1 transmission networks and sources of epidemic spread in Africa with deep-sequence phylogenetic analysis. Nature Communications, 2019. 10(1): p. 1411.
- 3. Wymant, C., et al., PHYLOSCANNER: Inferring Transmission from Within and Between-Host Pathogen Genetic Diversity. Molecular Biology and Evolution, 2017. 35(3): p. 719-733.

#### See Also

#### estimate\_p\_hat

# Examples

library(bumblebee)
library(dplyr)

# Prepare input to estimate p\_hat

# We shall use the data of HIV transmissions within and between intervention and control

- # communities in the BCPP/Ya Tsie HIV prevention trial. To learn more about the data
- # ?counts\_hiv\_transmission\_pairs and ?sampling\_frequency

# View counts of observed directed HIV transmissions within and between intervention # and control communities counts\_hiv\_transmission\_pairs

# View the estimated number of individuals with HIV in intervention and control # communities and the number of individuals sampled from each sampling\_frequency

# View results
results\_prep\_p\_hat

sampling\_frequency Sampling fequency

### Description

Estimated number of individuals with HIV in intervention and control communities of the BCPP/Ya Tsie trial, and the number of individuals sampled from each for HIV viral phylogenetic analysis.

# Usage

sampling\_frequency

## Format

A data frame:

population\_group Population group
number\_sampled Number of individuals sampled per population group
number\_population Estimated number of individuals in each population group

## Source

https://magosil86.github.io/bumblebee/

#### References

Magosi LE, et al., Deep-sequence phylogenetics to quantify patterns of HIV transmission in the context of a universal testing and treatment trial – BCPP/ Ya Tsie trial. To submit for publication, 2021.

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