# Package 'poth'

November 1, 2024

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Title Precision of Treatment Hierarchy (POTH)					
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Maintainer Augustine Wigle <amhwigle@uwaterloo.ca></amhwigle@uwaterloo.ca>					
<b>Description</b> Calculate POTH for treatment hierarchies from frequentist and Bayesian network meta- analysis. POTH quantifies the certainty in a treatment hierarchy. Subset POTH, POTH residu- als, and cumulative POTH can also be calculated to improve interpretation of treatment hierar- chies.					
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poth-package

poth: Brief overview of methods and general hints

# **Description**

R package **poth** allows to calculate the precision of treatment hierarchy (POTH) metric to quantify the uncertainty in a treatment hierarchy in network meta-analysis (Wigle et al., 2024).

#### **Details**

R package **poth** provides the following methods:

- Calculate the separation in ranking metric (poth)
- Conduct leave-one-out analysis (loo.poth)

Type help(package = "poth") for a listing of R functions available in **poth**.

Type citation("poth") on how to cite **poth** in publications.

The development version of **poth** is available on GitHub https://github.com/augustinewigle/poth.

# Author(s)

Augustine Wigle <amhwigle@uwaterloo.ca>, Guido Schwarzer <guido.schwarzer@uniklinik-freiburg.de>

#### References

Wigle, A., Béliveau, A., Salanti, G., Rücker, G., Schwarzer, G., Mavridis, D., Nikolakopoulou, A. (2024): Precision of Treatment Hierarchy: A Metric for Quantifying Uncertainty in Treatment Hierarchies in Network Meta-Analysis

#### See Also

Useful links:

- https://github.com/augustinewigle/poth
- Report bugs at https://github.com/augustinewigle/poth/issues

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cumul.poth	Cumulative method for precision of treatment hierarchy (POTH) metric
	ric

#### **Description**

Cumulative method for precision of treatment hierarchy (POTH) metric

# Usage

```
## S3 method for class 'poth'
cumul(x, sort = TRUE, ...)
## S3 method for class 'cumul.poth'
print(x, digits = 3, legend = TRUE, ...)
cumul(x, ...)
```

# Arguments

X	An R object of class poth.
sort	A logical indicating whether results should be sorted by decreasing ranking metric.
	Additional arguments.
digits	Minimal number of significant digits, see print.default.
legend	A logical indicating whether a legend should be printed.

#### Value

A data frame with additional class cumul.poth and the following variables:

trt Name of added treatment.
rank Treatment rank (global).
score Ranking metric (global).
poth\_cum Cumulative POTH.

# **Examples**

```
library("netmeta")
data(Senn2013)
net1 <- netmeta(TE, seTE, treat1.long, treat2.long, studlab,
   data = Senn2013, sm = "MD", random = FALSE)

# Cumulative method
c1 <- cumul(poth(net1))
c1</pre>
```

loo.poth

```
plot(c1)
plot(c1, labels = TRUE)
c2 <- cumul(poth(net1), sort = FALSE)
c2
plot(c2)
plot(c2, labels = TRUE)</pre>
```

loo.poth

Leave-one-out method for precision of treatment hierarchy (POTH) metric

#### **Description**

Leave-one-out method for precision of treatment hierarchy (POTH) metric

#### Usage

```
## S3 method for class 'poth'
loo(x, ...)
loo(x, ...)
## S3 method for class 'loo.poth'
print(x, digits = 3, legend = TRUE, ...)
```

#### **Arguments**

An R object of class poth.
 Additional arguments.
 Minimal number of significant digits, see print.default.
 legend A logical indicating whether a legend should be printed.

# Value

A data frame with additional class loo.poth and the following variables:

trt Treatment names.
rank Treatment rank (global).
score Ranking metric (global).
poth\_loo Leave-one-out POTH.

resid Residuals (global POTH minus leave-one-out POTH.
ratio Ratio of residual devided by absolute sum of residuals.

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#### **Examples**

```
library("netmeta")
data(smokingcessation)
p1 <- pairwise(list(treat1, treat2, treat3),
    event = list(event1, event2, event3), n = list(n1, n2, n3),
    data = smokingcessation, sm = "OR")
net1 <- netmeta(p1, random = FALSE)

# Leave-one-out method
loo1 <- loo(poth(net1))
loo1</pre>
```

plot.cumul.poth

Plot cumulative precision of treatment hierarchy (POTH) values

#### **Description**

Plot cumulative precision of treatment hierarchy (POTH) values

#### Usage

```
## S3 method for class 'cumul.poth'
plot(x, labels = FALSE, nchar.trts = 4, digits = 3, ...)
```

#### **Arguments**

x	R object of class cumul.poth.
labels	A logical indicating whether treatment names should be shown in the plot.
nchar.trts	Number of characters to keep for each treatment name if labels = TRUE.
digits	Minimal number of significant digits for cumulative POTH, see print. default.
	Additional arguments (ignored).

#### **Details**

Plot results of cumulative method for precision of treatment hierarchy (POTH) metric (Wigle et al., 2024).

#### Value

A ggplot2 object.

### Author(s)

Augustine Wigle <amhwigle@uwaterloo.ca>, Guido Schwarzer <guido.schwarzer@uniklinik-freiburg.de>

plot.loo.poth

#### References

Wigle, A., Béliveau, A., Salanti, G., Rücker, G., Schwarzer, G., Mavridis, D., Nikolakopoulou, A. (2024): Precision of Treatment Hierarchy: A Metric for Quantifying Uncertainty in Treatment Hierarchies in Network Meta-Analysis

#### **Examples**

```
library("netmeta")
data(Senn2013)
net1 <- netmeta(TE, seTE, treat1.long, treat2.long, studlab,
    data = Senn2013, sm = "MD", random = FALSE)

# Cumulative method
c1 <- cumul(poth(net1))
c1
plot(c1)
plot(c1, labels = TRUE)
c2 <- cumul(poth(net1), sort = FALSE)
c2
plot(c2)
plot(c2, labels = TRUE)</pre>
```

plot.loo.poth

Plot results of leave-one-out method

# **Description**

Plot results of leave-one-out method for precision of treatment hierarchy (POTH) metric

# Usage

```
## S3 method for class 'loo.poth'
plot(x, labels = TRUE, digits = 3, ...)
```

# **Arguments**

Χ	R object of class poth.
labels	A logical indicating whether treatment names should be shown in the plot.
digits	Minimal number of significant digits for global POTH, see print.default.
	Additional arguments (ignored).

#### **Details**

Plot results of leave-one-out method for precision of treatment hierarchy (POTH) metric (Wigle et al., 2024).

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#### Value

A ggplot2 object.

#### Author(s)

Augustine Wigle <amhwigle@uwaterloo.ca>, Guido Schwarzer <guido.schwarzer@uniklinik-freiburg.de>

#### References

Wigle, A., Béliveau, A., Salanti, G., Rücker, G., Schwarzer, G., Mavridis, D., Nikolakopoulou, A. (2024): Precision of Treatment Hierarchy: A Metric for Quantifying Uncertainty in Treatment Hierarchies in Network Meta-Analysis

#### **Examples**

```
library("netmeta")
data(smokingcessation)
p1 <- pairwise(list(treat1, treat2, treat3),</pre>
  event = list(event1, event2, event3), n = list(n1, n2, n3),
  data = smokingcessation, sm = "OR")
net1 <- netmeta(p1, random = FALSE)</pre>
# Leave-one-out method
loo1 <- loo(poth(net1))</pre>
1001
plot(loo1)
data(Senn2013)
net2 <- netmeta(TE, seTE, treat1.long, treat2.long, studlab,</pre>
                 data = Senn2013, sm = "MD", random = FALSE)
# Leave-one-out method (without sorting by ranking metric)
loo2 <- loo(poth(net2), sort = FALSE)</pre>
1002
plot(loo2)
```

poth

Calculate precision of treatment hierarchy (POTH) metric

#### **Description**

Precision of treatment hierarchy (POTH) is a metric to quantify the uncertainty in a treatment hierarchy in network meta-analysis

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#### Usage

```
poth(x, se = NULL, small.values, pooled, trts = NULL)
## S3 method for class 'poth'
print(x, sort = TRUE, digits = 3, ...)
## S3 method for class 'poth'
summary(object, ...)
## S3 method for class 'summary.poth'
print(x, sort = TRUE, digits = 3, ...)
```

#### **Arguments**

X	Mandatory argument with suitable information on the treatment hierarchy (see Details).
se	Matrix of estimated standard errors for relative effects.
small.values	A character string specifying whether small outcome values indicate a beneficial ("desirable") or harmful ("undesirable") effect, can be abbreviated.
pooled	A character string indicating whether the treatment hierarchy is based on a common or random effects model. Either "common" or "random", can be abbreviated.
trts	An optional vector with treatment names. Must match the order of treatments provided for argument x.
sort	A logical indicating whether printout should be sorted by decreasing ranking metric.
digits	Minimal number of significant digits, see print.default.
	Additional arguments (ignored).
object	An object of class summary.poth.

#### **Details**

This function calculates the precision of treatment hierarchy (POTH) metric to quantify the uncertainty in a treatment hierarchy in network meta-analysis (Wigle et al., 2024).

Argument x providing information on the treatment hierarchy is the only mandatory argument. The following input formats can be provided:

- 1. vector representing a ranking metric, i.e., SUCRAs or P-scores,
- 2. square matrix with the probabilities for each possible rank (with treatments in rows and ranks in columns),
- 3. MCMC samples (with samples in rows and treatments in columns),
- 4. relative effect matrix,
- 5. R object created with netmeta, netrank, or rankogram object from R package netmeta.

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Argument se must be provided if argument x is a matrix with relative effects. Otherwise, argument se is ignored.

Argument small.values must be provided if argument x contains MCMC samples, relative effects, or is an object created with netmeta. This argument can be provided for an R object created with netrank or rankogram and is ignored otherwise.

Argument trts is ignored for netmeta, netrank, and rankogram objects.

#### Value

An object of class poth with corresponding print function. The object is a list containing the following components:

poth Separation in ranking metric.

ranking A named numeric vector with rankings, i.e., SUCRAs or P-scores.

ranking.matrix A square matrix with the probabilities for each possible rank (if information is

available).

pooled As defined above.

#### Author(s)

Augustine Wigle <amhwigle@uwaterloo.ca>, Guido Schwarzer <guido.schwarzer@uniklinik-freiburg.de>

#### References

Wigle, A., Béliveau, A., Salanti, G., Rücker, G., Schwarzer, G., Mavridis, D., Nikolakopoulou, A. (2024): Precision of Treatment Hierarchy: A Metric for Quantifying Uncertainty in Treatment Hierarchies in Network Meta-Analysis

#### **Examples**

```
library("netmeta")
data(smokingcessation)
p1 <- pairwise(list(treat1, treat2, treat3),
    event = list(event1, event2, event3), n = list(n1, n2, n3),
    data = smokingcessation, sm = "OR")
net1 <- netmeta(p1, random = FALSE)

# Calculate probabilities for each possible rank
set.seed(1909) # make results reproducible
rg1 <- rankogram(net1)
rg1

# Calculate POTH
s1 <- poth(rg1)
s1

# Also print probabilities for each possible rank
summary(s1)

# Use SUCRAs to calculate POTH</pre>
```

pscores pscores

pscores

Calculate P-scores from a set of relative effects and standard errors

# Description

Calculate P-scores from a set of relative effects and standard errors

# Usage

```
pscores(TE, seTE, small.values = "desirable", trts = NULL)
```

# Arguments

TE	Matrix of relative effects
seTE	Matrix of estimated standard errors for relative effects
small.values	A character string specifying whether small outcome values indicate a beneficial ("desirable") or harmful ("undesirable") effect, can be abbreviated.
trts	optional; vector of treatment names matching order in TE and sds

#### Value

named vector of P-scores

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rankMCMC Calculate a ranking probabilities matrix from MCMC samples	rankMCMC	Calculate a ranking probabilities matrix from MCMC samples
---	----------	--

# Description

Calculate a ranking probabilities matrix from MCMC samples

# Usage

```
rankMCMC(x, small.values = "desirable", trts = NULL)
```

# Arguments

Х	a matrix or data.frame of MCMC samples, where rows are MCMC samples and columns are relative effects (relative to anchor) for treatments. must have column names that are the name of each treatment.
small.values	A character string specifying whether small outcome values indicate a beneficial ("desirable") or harmful ("undesirable") effect, can be abbreviated.
trts	character vector of treatment names, optional if samples has column names

# Value

A matrix of ranking probabilities where rows are treatments and columns are ranks

subset.poth	Calculate the local POTH for a subset of treatments	

# Description

Calculate the local POTH for a subset of treatments

# Usage

```
## S3 method for class 'poth'
subset(x, subset, top, bottom, ...)
```

# Arguments

x	An object of class poth.
subset	A character vector of treatment names to consider as the set of competing treatments.
top	A single integer to define the number of treatments with the largest ranking metric to consider in subset.
bottom	A single integer to define the number of treatments with the smallest ranking metric to consider in subset.
	Additional arguments (ignored).

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#### Value

An R object of class poth.

#### **Examples**

```
library("netmeta")
data(smokingcessation)
p1 <- pairwise(list(treat1, treat2, treat3),
    event = list(event1, event2, event3), n = list(n1, n2, n3),
    data = smokingcessation, sm = "OR")
net1 <- netmeta(p1, random = FALSE)

# Use P-scores to calculate local POTH for treatments "A" and "C"
subset(poth(net1), subset = c("A", "C"))

# Use P-scores to calculate local POTH for first three treatments
subset(poth(net1), top = 3)

# Use P-scores to calculate local POTH for first three treatments
subset(poth(net1), bottom = 3)</pre>
```

Xu2018

Safety of Immune Checkpoint Inhibitors in Cancer

# **Description**

Data from a network meta-analysis on immune checkpoint inhibitors (ICIs) to assess the safety of ICI drugs as cancer treatment (Xu et al., 2018).

# Format

A data frame with the following columns:

studyID study id
 treatment treatment name
 adverse number of adverse events
 n group sample size

#### **Details**

Data were obtained from Rosenberger et al. (2021), who re-analysed the data. There are seven treatments and 23 studies. The outcome of interest is the number of treatment-related adverse events, so smaller values indicate a safer treatment.

#### Source

Rosenberger, K.J., Duan, R., Chen, Y. et al. (2021): Predictive P-score for treatment ranking in Bayesian network meta-analysis. *BMC Med Res Methodol* **21**, 213. doi:10.1186/s12874021013975

Xu2018

# See Also

netmeta

# Examples

```
data(Xu2018)
head(Xu2018)

library("netmeta")
pw <- pairwise(treat = treatment, event = adverse, n = n,
    studlab = studyID, data = Xu2018, sm = "OR")
#
net <- netmeta(pw, small.values = "desirable", method.tau = "REML",
    common = FALSE)
#
poth(net)</pre>
```

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