

# Package: clusteredMSM (via r-universe)

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**Type** Package

**Title** Nonparametric Analysis of Clustered Multistate Processes

**Version** 0.1.0

**Description** Nonparametric estimation of population-averaged transition probabilities, with cluster-bootstrap pointwise confidence intervals, simultaneous confidence bands, and two-sample Kolmogorov-Smirnov-type tests for clustered or independent multistate process data. Estimation follows Bakoyannis (2021) <[doi:10.1111/biom.13327](https://doi.org/10.1111/biom.13327)>; two-sample inference for the cluster-randomized and independent-samples designs follows Bakoyannis and Bandyopadhyay (2022) <[doi:10.1007/s10463-021-00819-x](https://doi.org/10.1007/s10463-021-00819-x)>. Both methods use the working-independence Aalen-Johansen estimator. The package supports both progressive (acyclic) and non-monotone (e.g., illness-death with recovery) multistate processes, right censoring, left truncation, and informative cluster size. The user supplies data in interval format (one row per mutually-exclusive time interval per subject) and interacts with the package through a single formula-based function, `patp()`.

**License** GPL-3

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ci_cloglog	<i>Pointwise Confidence Intervals via Complementary Log-Log Transformation</i>
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## Description

Computes pointwise confidence intervals for a transition probability estimate using the cloglog transformation  $g(p) = \log(-\log p)$  with a delta-method standard error scaling. The bootstrap is run on the original probability scale; the cloglog SE is then derived analytically.

## Usage

```
ci_cloglog(point, se, level = 0.95)
```

## Arguments

point	Numeric vector of point estimates in (0, 1), length T.
se	Numeric vector of bootstrap standard errors of point on the probability scale, same length as point (i.e. <code>apply(boot_matrix, 1, sd, na.rm = TRUE)</code> ).
level	Confidence level. Default 0.95.

**Details**

By the delta method,  $SE(g(\hat{P})) = SE(\hat{P})/|\hat{P} \log \hat{P}|$  (here  $|g'(p)| = 1/|p \log p|$  on  $(0,1)$ ). The interval is built symmetrically on the cloglog scale and back-transformed via  $p = \exp(-\exp(\cdot))$ . Because  $g$  is monotone decreasing on  $(0, 1)$ , the upper end on the cloglog scale becomes the *lower* end on the probability scale, and vice versa.

For point estimates exactly equal to 0 or 1 the cloglog is undefined and NA is returned at those positions.

**Value**

A list with elements ll and ul: numeric vectors of lower and upper confidence limits, the same length as point.

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cluster_boot	<i>Generic Cluster Bootstrap Engine</i>
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**Description**

Performs the nonparametric cluster bootstrap: resamples whole clusters with replacement and applies a user-supplied statistic function to each bootstrap replicate.

**Usage**

```
cluster_boot(
  data,
  cluster,
  B,
  fn,
  ...,
  strata = NULL,
  seed = NULL,
  verbose = FALSE
)
```

**Arguments**

data	A data frame containing the cluster ID column named by cluster. All other columns are passed through unchanged to fn.
cluster	Character. Name of the cluster ID column.
B	Integer. Number of bootstrap replications.
fn	Function. Called as <code>fn(boot_data, ...)</code> once per bootstrap replicate. Must return a numeric vector of fixed length (the length must be the same across replicates – typically achieved by evaluating the statistic on a fixed grid of times).
...	Additional arguments passed to fn.

strata	Optional named vector mapping each unique cluster ID (as character) to a stratum label. When supplied, resampling is stratified: in each replicate, the original number of clusters in each stratum is drawn with replacement from the clusters in that stratum (so per-stratum cluster counts are preserved across replicates). When NULL (default), unstratified resampling is used.
seed	Optional integer. If non-NULL, set .seed(seed) is called before bootstrapping for reproducibility.
verbose	Logical. If TRUE, prints a progress message every 100 replicates. Default FALSE.

### Details

The bootstrap is nonparametric and at the cluster level: in each replicate,  $n$  clusters are sampled with replacement (where  $n$  is the number of unique values of `data[[cluster]]`), and the resulting cluster-bound rows are stacked into a new data frame.

Resampled clusters are re-IDed (1, 2, ...,  $n$ ) so that downstream code treating cluster IDs as distinct labels (e.g., `tapply` aggregations) works correctly even when the same original cluster appears multiple times in a replicate.

Stratified resampling (`strata`) supports cluster-randomized designs in which each cluster belongs to a single group (case ii of Bakoyannis & Bandyopadhyay 2022): the per-stratum cluster counts are fixed across replicates, so a replicate can never wipe out one of the groups.

The bootstrap is statistic-agnostic: the same engine is used for point-estimator standard errors, two-sample tests, and confidence bands. Whatever `fn` returns is what gets bootstrapped.

### Value

A numeric matrix with `length(fn(data, ...))` rows and `B` columns. Column `b` is the result of `fn` applied to bootstrap replicate `b`.

### Examples

```
data(example_msm)

# Cluster-bootstrap a simple summary: the mean of Tstop across
# rows. The user-supplied fn can return any fixed-length numeric
# vector; the typical use case is a transition probability curve
# evaluated on a fixed time grid (which is exactly what patp() does
# under the hood).
boot <- cluster_boot(
  data = example_msm, cluster = "cluster", B = 50,
  fn = function(d) mean(d$Tstop),
  seed = 1
)
c(point = mean(boot, na.rm = TRUE),
  se = stats::sd(boot, na.rm = TRUE))
```

---

confidence\_band      *Simultaneous Confidence Band via Cluster-Bootstrap Quantile*

---

### Description

Constructs a simultaneous (uniform-over-time) confidence band for a transition probability curve. The band is built on the cloglog scale: for each bootstrap replicate, the standardized supremum  $\sup_t |g(\hat{P}_b(t)) - g(\hat{P}(t))| / SE_g(t)$  is computed, where  $SE_g(t)$  is the delta-method SE on the cloglog scale. The level quantile of those suprema is the critical value, and the band is back-transformed to the probability scale.

### Usage

```
confidence_band(point, boot, times, level = 0.95, trim = c(0.05, 0.95))
```

### Arguments

point	Numeric vector of point estimates over a time grid.
boot	Numeric matrix of bootstrap replicates of point on the original probability scale: rows correspond to time points (matching point), columns to bootstrap replicates.
times	Numeric vector of times matching the rows of boot and the entries of point.
level	Confidence level. Default 0.95.
trim	Numeric vector of length 2. Lower and upper quantiles of the jump-time distribution at which to clip the band (avoids the "fans out" behavior near the extremes of follow-up). Default $c(0.05, 0.95)$ .

### Details

Only one bootstrap on the original (probability) scale is required:  $SE_g(t)$  is obtained by the delta method from  $SE(\hat{P}(t)) = sd(\hat{P}^*(t))$ , and the cloglog values  $g(\hat{P}_b(t))$  are computed by transforming the existing replicates pointwise – no separate cloglog-scale bootstrap is run.

### Value

A list with elements `ll.band` and `ul.band`: numeric vectors of band limits, with NA outside the trimmed range or where point is at the boundary 0/1.

### Note

This version uses an equal-precision-type band on the cloglog scale, in which the bootstrap deviation is standardized pointwise by the delta-method standard error of  $P(t)$ . By the self-cancellation of  $\sqrt{n}$  factors, this is asymptotically equivalent to standardizing the limit process by its asymptotic standard error on the cloglog scale, which is the standard equal-precision construction. This construction is asymptotically valid under conditions C1-C6 of Bakoyannis (2021), but differs from the Hall-Wellner-type weight proposed in Section 2.3 of that paper. Both constructions yield asymptotically valid simultaneous bands. The paper's Hall-Wellner-type construction will be added as a `band_method` option in a future release and will become the default.

## References

Bakoyannis, G. (2021). Nonparametric analysis of nonhomogeneous multistate processes with clustered observations. *Biometrics*, 77(2), 533-546. doi:10.1111/biom.13327

---

cut\_at\_lm

*Truncate a Long-Format Multistate Dataset at a Landmark Time*

---

## Description

Restricts a long-format multistate dataset to follow-up after a landmark time  $s$ . Intervals ending before  $s$  are removed; intervals straddling  $s$  are truncated to start at  $s$ .

## Usage

```
cut_at_lm(data, s)
```

## Arguments

data	A long-format data frame, typically produced by <code>intervals_to_long()</code> , with columns <code>Tstart</code> , <code>Tstop</code> , <code>from</code> , <code>to</code> , <code>trans</code> , <code>status</code> , <code>id</code> , and (optionally) <code>cluster</code> .
s	Numeric scalar. The landmark time.

## Details

This function is used in the landmark Aalen-Johansen estimator, which relaxes the Markov assumption by re-fitting hazards on the cohort still under observation at time  $s$ .

Important: this function only truncates the time axis. To compute the landmark estimator, you must also restrict to subjects who are in the relevant starting state at time  $s$  – use `state_at()` for that step.

Intervals where  $Tstop == s$  exactly are dropped (the subject has already transitioned by  $s$ ); intervals where  $Tstart >= s$  are kept unchanged; intervals straddling  $s$  ( $Tstart < s < Tstop$ ) have their  $Tstart$  reset to  $s$ . If a straddling interval was going to end in an event (`status == 1`) at  $Tstop > s$ , the event is preserved.

## Value

A data frame of the same shape as `data`, restricted and truncated as described above.

## References

Putter H, Spitoni C (2018). Non-parametric estimation of transition probabilities in non-Markov multi-state models: the landmark Aalen-Johansen estimator. *Statistical Methods in Medical Research* 27(7):2081-2092. doi:10.1177/0962280216674497

Bakoyannis G (2021). Nonparametric analysis of nonhomogeneous multistate processes with clustered observations. *Biometrics* 77(2):533-546. doi:10.1111/biom.13327

## Examples

```
data(example_msm)
tmat <- trans_mat(list(c(2, 3), c(1, 3), integer(0)),
                  names = c("Healthy", "Ill", "Dead"))
long <- intervals_to_long(example_msm, tmat)

# Restrict follow-up to after the landmark time s = 1.5
long_lm <- cut_at_lm(long, s = 1.5)
head(long_lm)
```

---

example\_msm

*Synthetic Clustered Illness-Death-with-Recovery Data*

---

## Description

A synthetic interval-format multistate dataset used in package examples, the README, and the vignette. The data are generated from an illness-death-with-recovery process with cluster-level frailty, so they exercise the package's headline feature: support for non-monotone (cyclic) transitions.

## Usage

```
example_msm
```

## Format

A data frame with 81 rows (one row per mutually-exclusive time interval per subject) and 7 columns:

**id** Integer subject identifier (40 subjects, ids 1..40).

**cluster** Integer cluster identifier (8 clusters, 5 subjects each).

**treatment** Binary treatment indicator (0 or 1), balanced 20/20 across subjects. Treatment 1 has a higher illness hazard than treatment 0.

**Tstart** Numeric start time of the interval.

**Tstop** Numeric end time of the interval.

**Sstart** Integer state during the interval: 1 = Healthy, 2 = Ill, 3 = Dead (absorbing).

**Sstop** Integer state at Tstop, or equal to Sstart on a final, censored row.

## Details

The transition structure is `trans_mat(list(c(2, 3), c(1, 3), integer(0)))`, i.e. Healthy <-> Ill, Healthy -> Dead, Ill -> Dead. Within each subject rows are temporally and spatially contiguous; censoring is encoded as `Sstart == Sstop` on the final row.

The same data are also shipped as a CSV at `system.file("extdata", "example_data.csv", package = "clusteredMSM")` for users who prefer to mimic loading their own file.

The generation script lives in `data-raw/example_msm.R` and is fully reproducible (`set.seed(2026)`).

**Source**

Simulated. See data-raw/example\_msm.R in the package sources.

**Examples**

```
data(example_msm)
head(example_msm)
table(example_msm$Sstart, example_msm$Sstop)
```

---

fit\_chaz

*Fit Cox Model Stratified by Transition and Return Cumulative Hazards*


---

**Description**

Fits a Cox proportional hazards model stratified by transition type and returns the Breslow-estimated cumulative transition hazards in long format.

**Usage**

```
fit_chaz(data, tmat, weights = NULL)
```

**Arguments**

data	A data frame in long multistate format with columns Tstart, Tstop, status, and trans. Typically the output of <code>intervals_to_long()</code> .
tmat	A K x K transition matrix from <code>trans_mat()</code> . Used only to validate that all transition IDs in data are defined in tmat; not used in the fit itself.
weights	Optional numeric vector of length <code>nrow(data)</code> with observation weights. Pass 1 / <code>data\$clust.size</code> for the inverse-cluster-size-weighted estimator. NULL (default) means unweighted.

**Details**

The model fitted is `coxph(Surv(Tstart, Tstop, status) ~ strata(trans), method = "breslow")`. This is the Andersen-Gill counting-process formulation, which handles multiple intervals per subject – including non-monotone processes with recovery – provided the input data was correctly constructed (one row per sojourn x candidate transition).

**Value**

A data frame with columns:

- time: jump time of the transition.
- Haz: Breslow estimate of the cumulative transition hazard at time.
- trans: integer transition ID (matches tmat).

Sorted by trans then time.

## Examples

```
data(example_msm)
tmat <- trans_mat(list(c(2, 3), c(1, 3), integer(0)),
                  names = c("Healthy", "Ill", "Dead"))

# fit_chaz takes long-format data; intervals_to_long() converts
# the interval format users supply. Most users reach fit_chaz
# indirectly via patp() -- this example shows the manual pipeline.
long <- intervals_to_long(example_msm, tmat)
haz <- fit_chaz(long, tmat)
head(haz)
```

---

intervals_to_long	<i>Convert Validated Interval Data to Long Multistate Format</i>
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---

## Description

Expands each row of an interval-format multistate dataset into one row per allowed transition out of the interval's starting state, with `status = 1` on the row whose destination matches the actual transition (and `0` on the others). The result is the long format expected by `fit_chaz` and other counting-process style routines.

## Usage

```
intervals_to_long(data, tmat)
```

## Arguments

<code>data</code>	A validated interval data frame with columns <code>id</code> , <code>Tstart</code> , <code>Tstop</code> , <code>Sstart</code> , <code>Sstop</code> , plus optional <code>cluster</code> and <code>group</code> columns. Run <code>validate_intervals()</code> first to confirm the data satisfy the package's contiguity and absorbing-state rules.
<code>tmat</code>	A $K \times K$ transition matrix from <code>trans_mat()</code> .

## Details

Most users do not need to call this function directly – `patp()` runs the full pipeline (parse, validate, expand, fit). It is exported so advanced users can compose the lower-level building blocks (`fit_chaz`, `prodint_AJ`, `cluster_boot`) into custom analyses.

## Value

A data frame in long multistate format with columns `id`, `Tstart`, `Tstop`, `from`, `to`, `trans`, `status`, plus `cluster` and `group` if present in `data`.

**Examples**

```
data(example_msm)
tmat <- trans_mat(list(c(2, 3), c(1, 3), integer(0)),
                  names = c("Healthy", "Ill", "Dead"))
long <- intervals_to_long(example_msm, tmat)
head(long)
```

ks\_pvalue

*Two-Sample Kolmogorov-Smirnov-Type Test via Cluster Bootstrap***Description**

Computes a p-value for the null hypothesis that a transition probability curve is equal across two groups, based on the supremum of the absolute difference between group-specific Aalen-Johansen estimators and a cluster-bootstrap reference distribution.

**Usage**

```
ks_pvalue(diff_point, diff_boot, scale)
```

**Arguments**

diff_point	Numeric vector. Observed group difference of the point estimator ( $\hat{P}_1 - \hat{P}_0$ ) on a fixed time grid.
diff_boot	Numeric matrix. Cluster bootstrap replicates of the group difference: rows = time points (matching diff_point), columns = replicates.
scale	Numeric scalar. Asymptotic scaling factor applied to both the observed sup-statistic and the bootstrap analogue. Use $\sqrt{n}$ ( $n$ = total clusters) for the dependent-groups design (Bakoyannis 2021, Theorem 3); use $\sqrt{n_1 n_2 / (n_1 + n_2)}$ for the cluster-randomized design (Bakoyannis & Bandyopadhyay 2022, Theorem 2).

**Details**

The test statistic is  $T = c_n \sup_t |\hat{P}_1(t) - \hat{P}_0(t)|$  with  $c_n = \text{scale}$ , and the null distribution is approximated by the bootstrap analogue applied to centered bootstrap differences. The empirical p-value is invariant to scale (it appears on both sides of the comparison); the role of scale is to put the reported statistic on the correct asymptotic scale per the relevant theorem.

This v0.1 implementation uses a unit weight: every time point on the grid contributes equally to the supremum. Bakoyannis (2021) Section 2.5 instead recommends a time-varying weight  $W(t) = \prod_p Y_p(t) / \sum_p Y_p(t)$  (the harmonic mean of the per-group at-risk processes), which down-weights regions where one group's at-risk set is small and naturally tames tail instability of the difference estimator. A weighted variant following Bakoyannis (2021) Section 2.5 and Bakoyannis & Bandyopadhyay (2022) is planned for v0.2.

**Value**

A list with elements `statistic` (the observed sup-statistic, on the chosen scale) and `p.value`.

---

msm *Construct a Multistate Interval Object for Use in patp() Formulas*

---

### Description

Packages four columns – interval start time, interval end time, starting state, and ending state – into a single object suitable for use as the response in a [patp](#) formula. Conceptually analogous to [Surv\(\)](#) in survival analysis: the user calls it inside a formula (`msm(Tstart, Tstop, Sstart, Sstop) ~ ...`).

### Usage

```
msm(Tstart, Tstop, Sstart, Sstop)
```

### Arguments

Tstart	Numeric vector. Start time of each interval.
Tstop	Numeric vector. End time of each interval.
Sstart	Integer-valued numeric vector. State occupied at Tstart. States must be whole numbers; the matrix backing stores them as double (matrices have a single storage mode), but values are validated to be integer-valued.
Sstop	Integer-valued numeric vector. State occupied at Tstop. Same storage caveat as Sstart. If Sstart == Sstop, the row represents a censored interval; this is permitted only on the final row of a subject's record.

### Details

Used inside a model formula passed to [patp](#), e.g. `patp(msm(Tstart, Tstop, Sstart, Sstop) ~ 1, ...)`. The four arguments can be named anything in the user's data – the formula machinery looks them up in the supplied data.

### Value

A four-column matrix of class "msm" with columns Tstart, Tstop, Sstart, Sstop.

### See Also

[patp](#), [validate\\_intervals](#).

### Examples

```
Tstart <- c(0, 1.5, 0, 0)
Tstop <- c(1.5, 3.0, 2.0, 1.2)
Sstart <- c(1, 2, 1, 1)
Sstop <- c(2, 3, 3, 1) # last row censored (S unchanged)
obj <- msm(Tstart, Tstop, Sstart, Sstop)
head(obj)
```

---

patp *Population-Averaged Transition Probabilities for Multistate Process Data*

---

### Description

Computes the working-independence Aalen-Johansen estimator of the transition probability  $P(X(t) = j \mid X(s) = h)$  for clustered or independent multistate process data, with cluster-bootstrap standard errors, pointwise confidence intervals, and (optionally) simultaneous confidence bands. If a grouping variable is supplied on the right-hand side of the formula, also conducts a two-sample Kolmogorov-Smirnov-type test of curve equality.

### Usage

```
patp(
  formula,
  data,
  tmat,
  id,
  cluster = NA,
  h,
  j,
  s = 0,
  weighted = FALSE,
  LMAJ = FALSE,
  B = 1000,
  cband = FALSE,
  design = c("auto", "shared", "cluster_random", "indep_random"),
  level = 0.95,
  seed = NULL
)
```

### Arguments

formula	A formula of the form $\text{msm}(T_{\text{start}}, T_{\text{stop}}, S_{\text{start}}, S_{\text{stop}}) \sim 1$ for a one-sample analysis, or $\text{msm}(T_{\text{start}}, T_{\text{stop}}, S_{\text{start}}, S_{\text{stop}}) \sim \text{group}$ for a two-sample analysis (estimate + test).
data	A data frame containing the variables in formula plus the id (and optional cluster) columns.
tmat	A $K \times K$ transition matrix, typically built with <code>trans_mat()</code> .
id	Character. Name of the subject ID column. Required.
cluster	Character or NA. Name of the cluster ID column. When NA (default), the bootstrap resamples individuals; when supplied, it resamples whole clusters.
h	Integer in 1..K. Starting state.
j	Integer in 1..K. Ending state.

s	Numeric scalar. Conditioning time. Default 0.
weighted	Logical. Inverse-cluster-size weighting to account for potentially informative cluster size (requires <code>cluster</code> ). Default FALSE.
LMAJ	Logical. Use the landmark Aalen-Johansen estimator (recommended when $s > 0$ and the Markov assumption is questionable). Default FALSE.
B	Integer. Number of bootstrap replications. $B = 0$ skips inference and is permitted only for one-sample formulas. Default 1000.
cband	Logical. If TRUE, return simultaneous confidence band limits at the level set by <code>level</code> (recommended $B \geq 1000$ ).
design	Character, two-sample only. One of "auto" (default), "shared", "cluster_random", "indep_random". Selects the two-sample regime; see the <i>Two-sample designs</i> section. "auto" infers the regime from the cluster/group structure of the data, defaulting to "indep_random" when each cluster carries a single group (a safer default than assuming cluster randomization). When that fallback fires, a warning is emitted asking the user to set "cluster_random" explicitly if the per-group cluster counts were fixed by cluster randomization.
level	Confidence level. Default 0.95.
seed	Optional integer for reproducible bootstrap.

### Value

An S3 object of class `patp` containing:

- `call`, `formula`: the original call and formula.
- `curves`: a data frame with columns `time`, `P`, `group` (if two-sample), and (if  $B > 0$ ) `se`, `ll`, `ul` – and `ll.band`, `ul.band` if `cband = TRUE`.
- `test`: NULL (one-sample) or a list with the observed K-S statistic and bootstrap p-value (two-sample).
- `n_subjects`, `n_clusters`, `groups`, `h`, `j`, `s`, `B`.

### Two-sample designs

Three regimes are supported, each with its own bootstrap scheme and asymptotic scaling factor; the test statistic in every case is  $T = c_n \sup_t |\hat{P}_1(t) - \hat{P}_0(t)|$ .

"shared" (**case i): dependent groups (multicenter trial)**. Every cluster contributes observations from both groups – e.g., each hospital randomizes its patients to treatment or control. Unstratified cluster bootstrap;  $c_n = \sqrt{n}$  (Bakoyannis 2021, Theorem 3), with  $n$  the total number of clusters.

"cluster\_random" (**case ii.a): cluster-randomized trial**. Each cluster is assigned to exactly one group, with per-group cluster counts  $n_1, n_2$  fixed by the randomization. Stratified cluster bootstrap (resample within each group);  $c_n = \sqrt{n_1 n_2 / (n_1 + n_2)}$  (Bakoyannis & Bandyopadhyay 2022, Theorem 2). This regime must be opted into; "auto" will not select it.

"indep\_random" (**case ii.b): independent observational comparison**. Each cluster carries one group, but  $n_1, n_2$  are random (the population, not the analyst, decided who landed in which group). Unstratified cluster bootstrap;  $c_n = \sqrt{n_1 n_2 / (n_1 + n_2)}$ . The asymptotic regime is the same

two-independent-samples limit as "cluster\_random"; only the bootstrap differs. This is the "auto" default when each cluster carries a single group.

Mixed structures (some clusters carry both groups, some only one) are not supported in this version.

patp() validates the supplied data against the requested design and stops with an informative error if they disagree.

The K-S statistic in v0.1 uses unit weight at every time point. Weighted variants – following the harmonic-mean weight  $W(t) = \prod_p Y_p(t) / \sum_p Y_p(t)$  of Bakoyannis (2021) Section 2.5 and the related construction in Bakoyannis & Bandyopadhyay (2022) – are planned for v0.2.

### Standard errors and confidence intervals

The cluster bootstrap is run *once*, on the original probability scale, in line with Bakoyannis (2021) Theorem 2. The se column in curves is the bootstrap standard deviation of  $\hat{P}(t)$  on that scale, i.e. `apply(boot_matrix, 1, sd, na.rm = TRUE)`.

Pointwise CIs (l1, u1) are then built on the cloglog scale  $g(p) = \log(-\log p)$  using the delta-method standardization  $SE_g(t) = SE(\hat{P}(t)) / |\hat{P}(t) \log \hat{P}(t)|$ , and back-transformed via  $p = \exp(-\exp(\cdot))$ . Simultaneous bands (l1.band, u1.band) use the same  $SE_g$ ; the level-quantile of the supremum gives the critical value.

Because the cloglog transformation is nonlinear, se and the resulting CI widths are *not* equal in general – the CI is asymmetric on the probability scale. Report se for descriptive purposes; use l1/u1 for inference.

The simultaneous band is trimmed to the central 90% of observed jump times by default to avoid the band fanning out near the extremes of follow-up; outside that window l1.band and u1.band are NA. See `confidence_band()` for the trim argument.

Note: in this version the simultaneous band uses an equal-precision-type construction on the cloglog scale, which is asymptotically valid but differs from the Hall-Wellner-type construction of Bakoyannis (2021) Section 2.3 (planned for a future release). See `confidence_band()` for the full disclosure.

### References

- Bakoyannis, G. (2021). Nonparametric analysis of nonhomogeneous multistate processes with clustered observations. *Biometrics*, 77(2), 533-546. doi:10.1111/biom.13327
- Bakoyannis, G., & Bandyopadhyay, D. (2022). Nonparametric tests for multistate processes with clustered data. *Annals of the Institute of Statistical Mathematics*, 74(5), 837-867. doi:10.1007/s1046302100819x
- Putter H, Spitoni C (2018). Non-parametric estimation of transition probabilities in non-Markov multi-state models: the landmark Aalen-Johansen estimator. *Statistical Methods in Medical Research* 27(7):2081-2092. doi:10.1177/0962280216674497

### See Also

`msm`, `trans_mat`, `validate_intervals`.

**Examples**

```

data(example_msm)
tmat <- trans_mat(list(c(2, 3), c(1, 3), integer(0)),
                  names = c("Healthy", "Ill", "Dead"))

# One-sample: P(Ill at t | Healthy at 0). B is small here for speed;
# use B >= 1000 for reported results.
fit <- patp(msm(Tstart, Tstop, Sstart, Sstop) ~ 1,
           data = example_msm, tmat = tmat,
           id = "id", cluster = "cluster",
           h = 1, j = 2, s = 0,
           B = 50, seed = 1)

fit

# Two-sample (estimate + test in one call). Each cluster in
# example_msm carries both treatment levels, so design = "shared".
tt <- patp(msm(Tstart, Tstop, Sstart, Sstop) ~ treatment,
          data = example_msm, tmat = tmat,
          id = "id", cluster = "cluster",
          h = 1, j = 2, B = 50,
          design = "shared", seed = 1)

tt

```

---

print.patp

*Print Method for patp Objects*


---

**Description**

Concise summary of a [patp](#) fit. For two-sample fits, includes the K-S statistic and p-value.

**Usage**

```

## S3 method for class 'patp'
print(x, digits = max(3L, getOption("digits") - 3L), ...)

```

**Arguments**

x	A patp object.
digits	Integer. Number of significant digits for printing.
...	Additional arguments (ignored).

**Value**

The object x, invisibly.

prodint\_AJ

*Aalen-Johansen Product Integral for General Multistate Processes***Description**

Computes the row of the transition probability matrix

$$P(s, t) = \prod_{u \in (s, t]} (I + dA(u))$$

starting from state  $h$ , given Nelson-Aalen transition hazard estimates.

**Usage**

```
prodint_AJ(haz_df, tmat, predt = 0, h = 1)
```

**Arguments**

haz_df	Data frame with columns time, Haz, trans, typically the output of fit_chaz() or the \$Haz element of an mstate::msfit object. Haz is the cumulative (not incremental) hazard.
tmat	A K x K transition matrix from trans_mat(). Cyclic transitions (both [h, j] and [j, h] non-NA) are permitted.
predt	Numeric scalar. Starting time s. Probabilities are computed for t > predt. Default 0.
h	Integer in 1..K. Starting state.

**Details**

At each unique jump time  $u$ , the increment matrix  $dA(u)$  has off-diagonal entry  $(h, j)$  equal to the Nelson-Aalen increment of the cumulative hazard for transition  $h \rightarrow j$ , and diagonal entries equal to the negative row sums. The transition probability matrix is then updated by  $P \leftarrow P(I + dA(u))$ .

Computational complexity is  $O(J K^2)$  where J is the number of unique jump times and K is the number of states.

**Value**

A data frame with columns time, pstate1, ..., pstateK. Row k gives  $P(predt, time_k)[h, ]$ , the probabilities of being in each state at time\_k given the process was in state h at predt.

**Examples**

```
data(example_msm)
tmat <- trans_mat(list(c(2, 3), c(1, 3)), integer(0)),
                 names = c("Healthy", "Ill", "Dead"))

long <- intervals_to_long(example_msm, tmat)
```

```
haz <- fit_chaz(long, tmat)
P <- prodint_AJ(haz, tmat, predt = 0, h = 1)
head(P)
```

---

state\_at *Identify Each Subject's State at a Given Time*

---

### Description

Given a long-format multistate dataset, returns the state each subject occupies at time  $s$ . Used by the landmark Aalen-Johansen estimator to identify the risk set at a conditioning time.

### Usage

```
state_at(data, s, id = "id")
```

### Arguments

data	A data frame in long format with columns Tstart, Tstop, from, to, status, and the subject ID column named by id.
s	Numeric scalar. The time at which to evaluate states.
id	Character. Name of the subject ID column. Default "id".

### Details

A subject's state at time  $s$  is the from state of the interval that contains  $s$  (i.e.,  $T_{\text{start}} \leq s < T_{\text{stop}}$ ). Because the long format duplicates intervals across competing transitions, the result is deduplicated to one row per subject.

For exact ties ( $s == T_{\text{stop}}$ ), the subject is treated as having transitioned: they appear in their new state if a subsequent interval exists, otherwise they are excluded as absorbed/censored.

### Value

A data frame with columns <id> and state, one row per subject who is under observation at time  $s$ . Subjects whose follow-up has not yet started ( $\min(T_{\text{start}}) > s$ ) or who have already entered an absorbing state ( $\max(T_{\text{stop}}) \leq s$  with status == 1) are excluded.

### References

Putter H, Spitoni C (2018). Non-parametric estimation of transition probabilities in non-Markov multi-state models: the landmark Aalen-Johansen estimator. *Statistical Methods in Medical Research* 27(7):2081-2092. doi:10.1177/0962280216674497

**Examples**

```

data(example_msm)
tmat <- trans_mat(list(c(2, 3), c(1, 3), integer(0)),
                  names = c("Healthy", "Ill", "Dead"))
long <- intervals_to_long(example_msm, tmat)

# Each subject's state at the landmark time s = 1.5
head(state_at(long, s = 1.5, id = "id"))

```

---

summary.patp

*Summary Method for patp Objects*


---

**Description**

Returns the full curve(s) and (if applicable) the test result.

**Usage**

```

## S3 method for class 'patp'
summary(object, ...)

```

**Arguments**

object	A patp object.
...	Additional arguments (ignored).

**Value**

A list with named components for further inspection.

---

trans\_mat

*Build a Transition Matrix for a Multistate Process*


---

**Description**

Constructs a square transition matrix in the format expected by [patp](#) and [prodint\\_AJ](#). It supports cyclic transitions (e.g., illness-death with recovery).

**Usage**

```

trans_mat(x, names = NULL)

```

**Arguments**

x	A list of length K. Element k is an integer vector of states reachable directly from state k. Use <code>integer(0)</code> or <code>NULL</code> for absorbing states.
names	Optional character vector of length K with state names. If <code>NULL</code> , states are named "1", "2", ..., "K".

**Value**

A  $K \times K$  integer matrix. Entry  $[h, j]$  is the integer transition ID for the direct transition  $h \rightarrow j$ , or NA if no such transition exists. Transition IDs are assigned in row-major order (all transitions out of state 1 first, then state 2, etc.). The matrix has `dimnames list(from = names, to = names)`.

**Examples**

```
# Illness-death without recovery (progressive)
trans_mat(list(c(2, 3), 3, integer(0)),
          names = c("Healthy", "Ill", "Dead"))

# Illness-death WITH recovery (non-monotone)
trans_mat(list(c(2, 3), c(1, 3), integer(0)),
          names = c("Healthy", "Ill", "Dead"))

# Four-state multistate model with recovery from two intermediate states
trans_mat(list(c(2, 3, 4), c(1, 4), c(1, 4), integer(0)))
```

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