

Package: diyar (via r-universe)

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Type Package

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Description An R package for iterative and batched record linkage, and applying epidemiological case definitions. 'diyar' can be used for deterministic and probabilistic record linkage, or multistage record linkage combining both approaches. It features the implementation of nested match criteria, and mechanisms to address missing data and conflicting matches during stepwise record linkage. Case definitions are implemented by assigning records to groups based on match criteria such as person or place, and overlapping time or duration of events e.g. sample collection dates or periods of hospital stays. Matching records are assigned a unique group ID. Index and duplicate records are removed or further analyses as required.

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Encoding UTF-8

LazyData true

Imports methods, utils, ggplot2, rlang

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attr_eval	<i>Sub-criteria attributes.</i>
-----------	---------------------------------

Description

Recursive evaluation of a function (func) on each attribute (vector) in a [sub_criteria](#).

Usage

```
attr_eval(x, func = length, simplify = TRUE)
```

Arguments

x	[sub_criteria]
func	[function]
simplify	If TRUE (default), coerce to a vector.

Value

vector; list

Examples

```
x <- sub_criteria(rep(1, 5), rep(5 * 10, 5))
attr_eval(x)
attr_eval(x, func = max)
attr_eval(x, func = max, simplify = FALSE)
attr_eval(sub_criteria(x, x), func = max, simplify = FALSE)
```

bys_funcs	<i>Vectorised approach to group operations.</i>
-----------	---

Description

Vectorised approach to group operations.

Usage

```
bys_count(by)

bys_rank(..., by = NULL, from_last = FALSE)

bys_position(val, by = NULL, from_last = FALSE, ordered = TRUE)

bys_val(..., val, by = NULL, from_last = FALSE)
```

```

bys_nval(..., val, by = NULL, from_last = FALSE, n = 1, nmax = FALSE)

bys_min(val, by = NULL, na.rm = TRUE)

bys_max(val, by = NULL, na.rm = TRUE)

bys_sum(val, by = NULL, na.rm = TRUE)

bys_prod(val, by = NULL, na.rm = TRUE)

bys_cummin(val, by = NULL, na.rm = TRUE)

bys_cummax(val, by = NULL, na.rm = FALSE)

bys_cumsum(val, by = NULL, na.rm = TRUE)

bys_cumprod(val, by = NULL, na.rm = TRUE)

bys_lag(val, by = NULL, n = 1)

bys_lead(val, by = NULL, n = 1)

```

Arguments

by	[atomic]. Groups.
...	[atomic]. Sort levels
from_last	[logical] Sort order - TRUE (descending) or FALSE (ascending).
val	[atomic]. Value
ordered	If TRUE, values are sequential.
n	[integer] Position.
nmax	[logical] If TRUE, use length([by]) when n is greater than the number of records in a group.
na.rm	If TRUE, remove NA values

Value

[atomic]

Examples

```

x <- data.frame(
  group = c(2, 2, 1, 2, 1, 1, 1, 2, 1, 1),
  value = c(13, 14, 20, 9, 2, 1, 8, 18, 3, 17))

bys_count(x$group)
bys_position(x$value, by = x$group, from_last = TRUE)
bys_rank(by = x$group, val = x$value, from_last = TRUE)
bys_val(x$value, by = x$group, val = x$value, from_last = TRUE)

```

```
bys_nval(x$value, by = x$group, val = x$value, from_last = TRUE, n = 2)
bys_min(by = x$group, val = x$value)
bys_max(by = x$group, val = x$value)
bys_sum(by = x$group, val = x$value)
bys_prod(by = x$group, val = x$value)
bys_cummin(by = x$group, val = x$value)
bys_cummax(by = x$group, val = x$value)
bys_cumsum(by = x$group, val = x$value)
bys_cumprod(by = x$group, val = x$value)
bys_lag(by = x$group, val = x$value)
bys_lead(by = x$group, val = x$value)
```

combi

Vector combinations

Description

Numeric codes for unique combination of vectors.

Usage

```
combi(...)
```

Arguments

```
... [atomic]
```

Value

numeric

Examples

```
x <- c("A", "B", "A", "C", "B", "B")
y <- c("X", "X", "Z", "Z", "X", "Z")
combi(x, y)
```

```
# The code above is equivalent to but quicker than the one below.
```

```
z <- paste0(y, "-", x)
z <- match(z, z)
z
```

custom_sort	<i>Nested sorting</i>
-------------	-----------------------

Description

Returns a sort order after sorting by a vector within another vector.

Usage

```
custom_sort(..., decreasing = FALSE, unique = FALSE)
```

Arguments

...	Sequence of atomic vectors. Passed to order .
decreasing	Sort order. Passed to order .
unique	If FALSE (default), ties get the same rank. If TRUE, ties are broken.

Value

numeric sort order.

Examples

```
a <- c(1, 1, 1, 2, 2)
b <- c(2, 3, 2, 1, 1)

custom_sort(a, b)
custom_sort(b, a)
custom_sort(b, a, unique = TRUE)
```

delink	<i>Unlink group identifiers</i>
--------	---------------------------------

Description

Unlink records from an episode ([epid](#)), record group ([pid](#)) or pane ([pane](#)) object.

Usage

```
delink(x, lgk, ...)  
  
## S3 method for class 'epid'  
delink(x, lgk, ...)  
  
## S3 method for class 'pane'  
delink(x, lgk, ...)  
  
## S3 method for class 'pid'  
delink(x, lgk, ...)
```

Arguments

x	[epid pid pane]
lgk	[logical]. Subset of records to unlink.
...	Other arguments.

Value

[epid](#); [pid](#); [pane](#)

Examples

```
ep <- episodes(1:8)  
unlinked_ep <- delink(ep, ep@sn %in% c(3, 8))  
ep; unlinked_ep  
  
pn <- partitions(1:8, length.out = 2, separate = TRUE)  
unlinked_pn <- delink(pn, pn@.Data == 5)  
pn; unlinked_pn  
  
pd <- links(list(c(1, 1, 1, NA, NA),  
                c(NA, NA, 2, 2, 2)))  
unlinked_pd <- delink(pd, pd@pid_cri == 1)  
pd; unlinked_pd  
  
# A warning is given if an index record is unlinked as this will lead to seemly impossible links.  
ep2 <- episodes(1:8, 2, episode_type = "rolling")  
unlinked_ep2 <- delink(ep2, ep2@sn %in% c(3, 5))  
schema(ep2, custom_label = decode(ep2@case_nm), seed = 2)  
schema(unlinked_ep2, custom_label = decode(unlinked_ep2@case_nm), seed = 2)
```

d_report	<i>d_report</i>
----------	-----------------

Description

d_report

Usage

```
## S3 method for class 'd_report'
plot(
  x,
  ...,
  metric = c("cumulative_duration", "duration", "max_memory", "records_checked",
            "records_skipped", "records_assigned")
)

## S3 method for class 'd_report'
as.list(x, ...)

## S3 method for class 'd_report'
as.data.frame(x, ...)
```

Arguments

x	[d_report].
...	Arguments passed to other methods
metric	Report information

encode	<i>Labelling in diyar</i>
--------	---------------------------

Description

Encode and decode character and numeric values.

Usage

```
encode(x, ...)

decode(x, ...)

## Default S3 method:
encode(x, ...)
```



```
## S3 method for class 'd_label'  
encode(x, ...)  
  
## Default S3 method:  
decode(x, ...)  
  
## S3 method for class 'd_label'  
decode(x, ...)  
  
## S3 method for class 'd_label'  
rep(x, ...)  
  
## S3 method for class 'd_label'  
x[i, ..., drop = TRUE]  
  
## S3 method for class 'd_label'  
x[[i, ..., drop = TRUE]]
```

Arguments

x	[d_label atomic]
...	Other arguments.
i	i
drop	drop

Details

To minimise memory usage, most components of [pid](#), [epid](#) and [pane](#) are integer objects with labels. `encode()` and `decode()` translates these codes and labels as required.

Value

d_label; atomic

Examples

```
cds <- encode(rep(LETTERS[1:5], 3))  
cds  
  
nms <- decode(cds)  
nms
```

epid-class

epid *object*

Description

S4 objects storing the result of [episodes](#).

Usage

```
is.epid(x)

as.epid(x, ...)

## S3 method for class 'epid'
format(x, ...)

## S3 method for class 'epid'
unique(x, ...)

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

## S3 method for class 'epid_summary'
print(x, ...)

## S3 method for class 'epid'
as.data.frame(x, ..., decode = TRUE)

## S3 method for class 'epid'
as.list(x, ..., decode = TRUE)

## S4 method for signature 'epid'
show(object)

## S4 method for signature 'epid'
rep(x, ...)

## S4 method for signature 'epid'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'epid'
x[[i, j, ..., exact = TRUE]]

## S4 method for signature 'epid'
c(x, ...)
```

Arguments

x	x
...	...
object	object
decode	If TRUE, data is decoded
i	i
j	j
drop	drop
exact	exact

Slots

sn Unique record identifier.

.Data Unique episode identifier.

wind_id Unique reference ID for each match.

wind_nm Type of window i.e. "Case" or "Recurrence".

case_nm Record type in regards to case assignment.

dist_wind_index Unit difference between each record and its window's reference record.

dist_epid_index Unit difference between each record and its episode's reference record.

epid_dataset Data sources in each episode.

epid_interval The start and end dates of each episode. A [number_line](#) object.

epid_length The duration or length of (epid_interval).

epid_total The number of records in each episode.

iteration The iteration when a record was matched to it's group (.Data).

options Some options passed to the instance of [episodes](#).

Examples

```
# A test for `epid` objects
ep <- episodes(date = 1)
is.epid(ep); is.epid(2)

ep <- episodes(date = 1)
is.epid(ep); is.epid(2)
```

episodes *Group dated events into episodes.*

Description

Dated events (records) within a certain duration of an index event are assigned to a unique group. Each group has unique ID and are described as "episodes". "episodes" can be "fixed" or "rolling" ("recurring"). Each episodes has a "Case" and/or "Recurrent" record while all other records within the group are either "Duplicates" of the "Case" or "Recurrent" event.

Usage

```
episodes(
  date,
  case_length = Inf,
  episode_type = "fixed",
  recurrence_length = case_length,
  episode_unit = "days",
  strata = NULL,
  sn = NULL,
  episodes_max = Inf,
  rolls_max = Inf,
  case_overlap_methods = 8,
  recurrence_overlap_methods = case_overlap_methods,
  skip_if_b4_lengths = FALSE,
  data_source = NULL,
  data_links = "ANY",
  custom_sort = NULL,
  skip_order = Inf,
  reference_event = "last_record",
  case_for_recurrence = FALSE,
  from_last = FALSE,
  group_stats = c("case_nm", "wind", "epid_interval"),
  display = "none",
  case_sub_criteria = NULL,
  recurrence_sub_criteria = case_sub_criteria,
  case_length_total = 1,
  recurrence_length_total = case_length_total,
  skip_unique_strata = TRUE,
  splits_by_strata = 1,
  batched = "semi"
)

links_wf_episodes(
  date,
  case_length = Inf,
  episode_type = "fixed",
```

```

    strata = NULL,
    sn = NULL,
    display = "none"
)

episodes_af_shift(
  date,
  case_length = Inf,
  sn = NULL,
  strata = NULL,
  group_stats = FALSE,
  episode_type = "fixed",
  data_source = NULL,
  episode_unit = "days",
  data_links = "ANY",
  display = "none"
)

```

Arguments

date [date|datetime|integer|[number_line](#)]. Record date or period.

case_length [integer|[number_line](#)]. Duration from an index event distinguishing one "Case" from another.

episode_type [character]. Options are "fixed" (default) or "rolling". See Details.

recurrence_length [integer|[number_line](#)]. Duration from an index event distinguishing a "Recurrent" event from its "Case" or prior "Recurrent" event.

episode_unit [character]. Unit of time for case_length and recurrence_length. Options are "seconds", "minutes", "hours", "days" (default), "weeks", "months" or "years". See `diyar::episode_unit`.

strata [atomic]. Subsets of the dataset. Episodes are created separately by each strata.

sn [integer]. Unique record ID.

episodes_max [integer]. Maximum number of episodes permitted within each strata.

rolls_max [integer]. Maximum number of times an index event can recur. Only used if episode_type is "rolling".

case_overlap_methods [character|integer]. Specific ways a period (record) most overlap with a "Case" event. See ([overlaps](#)).

recurrence_overlap_methods [character|integer]. Specific ways a period (record) most overlap with a "Recurrent" event. See ([overlaps](#)).

skip_if_b4_lengths [logical]. If TRUE (default), events before a lagged case_length or recurrence_length are skipped.

data_source	[character]. Source ID for each record. If provided, a list of all sources in each episode is returned. See epid_dataset_slot .
data_links	[list character]. data_source required in each epid . An episode without records from these data_sources will be unlinked . See Details.
custom_sort	[atomic]. Preferential order for selecting index events. See custom_sort .
skip_order	[integer]. End episode tracking in a strata when the an index event's custom_sort order is greater than the supplied skip_order.
reference_event	[character]. Specifies which of the records are used as index events. Options are "last_record" (default), "last_event", "first_record" or "first_event".
case_for_recurrence	[logical]. If TRUE, a case_length is applied to both "Case" and "Recurrent" events. If FALSE (default), a case_length is applied to only "Case" events.
from_last	[logical]. Track episodes beginning from the earliest to the most recent record (FALSE) or vice versa (TRUE).
group_stats	[character]. A selection of group metrics to return for each episode. Most are added to slots of the epid object. Options are NULL or any combination of "case_nm", "wind" and "epid_interval".
display	[character]. Display progress update and/or generate a linkage report for the analysis. Options are; "none" (default), "progress", "stats", "none_with_report", "progress_with_report" or "stats_with_report".
case_sub_criteria	[sub_criteria]. Additional nested match criteria for events in a case_length.
recurrence_sub_criteria	[sub_criteria]. Additional nested match criteria for events in a recurrence_length.
case_length_total	[integer number_line]. Minimum number of matched case_lengths required for an episode.
recurrence_length_total	[integer number_line]. Minimum number of matched recurrence_lengths required for an episode.
skip_unique_strata	[logical]. If TRUE, a strata with a single event is skipped.
splits_by_strata	[integer]. Split analysis into n parts. This typically lowers max memory usage but increases run time.
batched	[character]. Create and compare records in batches. Options are "yes", "no", and "semi". typically, the ("semi") option will have a higher max memory and shorter run-time while ("no") will have a lower max memory but longer run-time

Details

`episodes()` links dated records (events) that are within a set duration of each other in iterations. Every record is linked to a unique group (episode; [epid](#) object). These episodes represent occurrences of interest as specified by function's arguments and defined by a case definition.

Two main type of episodes are possible;

- "fixed" - An episode where all events are within a fixed duration of an index event.
- "rolling" - An episode where all events are within a recurring duration of an index event.

Every record in each episode is categorised as one of the following;

- "Case" - Index event of the episode (without a nested match criteria).
- "Case_CR" - Index event of the episode (with a nested match criteria).
- "Duplicate_C" - Duplicate of the index event.
- "Recurrent" - Recurrence of the index event (without a nested match criteria).
- "Recurrent_CR" - Recurrence of the index event (with a nested match criteria).
- "Duplicate_R" - Duplicate of the recurrent event.
- "Skipped" - Skipped records.

If `data_links` is supplied, every element of the list must be named "l" (links) or "g" (groups). Unnamed elements are assumed to be "l".

- If named "l", groups without records from every listed `data_source` will be unlinked.
- If named "g", groups without records from any listed `data_source` will be unlinked.

All records with a missing (NA) `strata` or `date` are skipped.

Wrapper functions or alternative implementations of `episodes()` for specific use cases or benefits:

- `episodes_wf_repeats()` - Identical records are excluded from the main analysis.
- `episodes_af_shift()` - A mostly vectorised approach.
- `links_wf_episodes()` - The same functionality achieved with [links](#).

See `vignette("episodes")` for further details.

Value

`epid`; list

See Also

[episodes_wf_repeats](#); [custom_sort](#); [sub_criteria](#); [epid_length](#); [epid_window](#); [partitions](#); [links](#); [overlaps](#);

Examples

```
data(infections)
data(hospital_admissions)

# One 16-day (15-day difference) fixed episode per type of infection
episodes(date = infections$date,
         strata = infections$infection,
         case_length = 15,
         episodes_max = 1,
```

```

        episode_type = "fixed")

# Multiple 16-day episodes with an 11-day recurrence period
episodes(date = infections$date,
         strata = NULL,
         case_length = 15,
         episodes_max = Inf,
         episode_type = "rolling",
         recurrence_length = 10)

# Overlapping periods of hospital stays
dfr <- hospital_admissions[2:3]

dfr$admin_period <-
  number_line(dfr$admin_dt, dfr$discharge_dt)

dfr$ep <-
  episodes(date = dfr$admin_period,
         strata = NULL,
         case_length = index_window(dfr$admin_period),
         case_overlap_methods = "inbetween")

dfr
as.data.frame(dfr$ep)

```

episodes_wf_repeats *Link events to chronological episodes.*

Description

episodes_wf_repeats is a wrapper function of [episodes](#). It's designed to be more efficient with larger datasets. Duplicate records which do not affect the case definition are excluded prior to episode tracking. The resulting episode identifiers are then recycled for the duplicate records.

Usage

```
episodes_wf_repeats(..., duplicates_recovered = "ANY")
```

Arguments

...	Arguments passed to episodes .
duplicates_recovered	[character]. Determines which duplicate records are recycled. Options are "ANY" (default), "without_sub_criteria", "with_sub_criteria" or "ALL". See Details.
reframe	[logical]. Determines if the duplicate records in a sub_criteria are re-framed (TRUE) or excluded (FALSE).

Details

`episodes_wf_repeats()` reduces or re-frames a dataset to the minimum datasets required to implement a case definition. This leads to the same outcome but with the benefit of a shorter processing time.

The `duplicates_recovered` argument determines which identifiers are recycled. Selecting the `"with_sub_criteria"` option will force only identifiers created resulting from a matched `sub_criteria` (`"Case_CR"` and `"Recurrent_CR"`) are recycled. However, if `"without_sub_criteria"` is selected then only identifiers created that do not result from a matched `sub_criteria` (`"Case"` and `"Recurrent"`) are recycled. Excluded duplicates of `"Duplicate_C"` and `"Duplicate_R"` are always recycled.

The `reframe` argument will either `reframe` or subset a `sub_criteria`. Both will require slightly different functions for `match_funcs` or `equal_funcs`.

Value

`epid`; list

See Also

`episodes`; `sub_criteria`

Examples

```
# With 2,000 duplicate records of 20 events,
# `episodes_wf_repeats()` will take less time than `episodes()`
dates <- seq(from = as.Date("2019-04-01"), to = as.Date("2019-04-20"), by = 1)
dates <- rep(dates, 2000)

system.time(
  ep1 <- episodes(dates, 1)
)
system.time(
  ep2 <- episodes_wf_repeats(dates, 1)
)

# Both leads to the same outcome.
all(ep1 == ep2)
```

links

Multistage record linkage

Description

Assign records to unique groups based on an ordered set of match criteria.

Usage

```

links(
  criteria,
  sub_criteria = NULL,
  sn = NULL,
  strata = NULL,
  data_source = NULL,
  data_links = "ANY",
  display = "none",
  group_stats = FALSE,
  expand = TRUE,
  shrink = FALSE,
  recursive = "none",
  check_duplicates = FALSE,
  tie_sort = NULL,
  batched = "yes",
  repeats_allowed = FALSE,
  permutations_allowed = FALSE,
  ignore_same_source = FALSE
)

```

Arguments

criteria	[list atomic]. Ordered list of attributes to be compared. Each element of the list is a stage in the linkage process. See Details.
sub_criteria	[list sub_criteria]. Nested match criteria. This must be paired to a stage of the linkage process (criteria). See sub_criteria
sn	[integer]. Unique record ID.
strata	[atomic]. Subsets of the dataset. Record-groups are created separately for each strata. See Details.
data_source	[character]. Source ID for each record. If provided, a list of all sources in each record-group is returned. See pid_dataset slot .
data_links	[list character]. data_source required in each pid. A record-group without records from these data_sources will be unlinked . See Details.
display	[character]. Display progress update and/or generate a linkage report for the analysis. Options are; "none" (default), "progress", "stats", "none_with_report", "progress_with_report" or "stats_with_report".
group_stats	[logical]. If TRUE (default), return group specific information like record counts for each pid .
expand	[logical]. If TRUE, a record-group gains new records if a match is found at the next stage of the linkage process. <i>Not interchangeable with</i> shrink.
shrink	[logical]. If TRUE, a record-group loses existing records if no match is found at the next stage of the linkage process. <i>Not interchangeable with</i> expand.
recursive	[logical]. If TRUE, within each iteration of the process, a match can spawn new matches. Ignored when batched is "no".

check_duplicates	[logical]. If TRUE, within each iteration of the process, duplicates values of an attributes are not checked. The outcome of the logical test on the first instance of the value will be recycled for the duplicate values. Ignored when batched is "no".
tie_sort	[atomic]. Preferential order for breaking match ties within an iteration of record linkage.
batched	[character] Determines if record-pairs are created and compared in batches. Options are "yes", "no" or "semi".
repeats_allowed	[logical] If TRUE, pairs made up of repeat records are not created and compared. Only used when batched is "no".
permutations_allowed	[logical] If TRUE, permutations of record-pairs are created and compared. Only used when batched is "no".
ignore_same_source	[logical] If TRUE, only records-pairs from a different data_source are created and compared.

Details

The priority of matches decreases with each subsequent stage of the linkage process. Therefore, the attributes in `criteria` should be in an order of decreasing relevance.

Records with missing data (NA) for each `criteria` are skipped at the respective stage, while records with missing data strata are skipped from every stage.

If a record is skipped from a stage, another attempt will be made to match the record at the next stage. If a record is still unmatched by the last stage, it is assigned a unique group ID.

A `sub_criteria` adds nested match criteria to each stage of the linkage process. If used, only records with a matching `criteria` and `sub_criteria` are linked.

In `links`, each `sub_criteria` must be linked to a `criteria`. This is done by adding each `sub_criteria` to a named element of a list - "cr" concatenated with the corresponding stage's number. For example, 3 `sub_criteria` linked to `criteria` 1, 5 and 13 will be;

```
list(cr1 = sub_criteria(...), cr5 = sub_criteria(...), cr13 = sub_criteria(...))
```

Any unlinked `sub_criteria` will be ignored.

Every element in `data_links` must be named "l" (links) or "g" (groups). Unnamed elements of `data_links` will be assumed to be "l".

- If named "l", groups without records from every listed `data_source` will be unlinked.
- If named "g", groups without records from any listed `data_source` will be unlinked.

See vignette("links") for more information.

Value

`pid`; list

See Also

[links_af_probabilistic](#); [episodes](#); [predefined_tests](#); [sub_criteria](#)

Examples

```

data(patient_records)
dfr <- patient_records
# An exact match on surname followed by an exact match on forename
stages <- as.list(dfr[c("surname", "forename")])
p1 <- links(criteria = stages)

# An exact match on forename followed by an exact match on surname
p2 <- links(criteria = rev(stages))

# Nested matches
# Same sex OR birth year
m.cri.1 <- sub_criteria(
  format(dfr$dateofbirth, "%Y"), dfr$sex,
  operator = "or")

# Same middle name AND a 10 year age difference
age_diff <- function(x, y){
  diff <- abs(as.numeric(x) - as.numeric(y))
  wgt <- diff %in% 0:10 & !is.na(diff)
  wgt
}
m.cri.2 <- sub_criteria(
  format(dfr$dateofbirth, "%Y"), dfr$middlename,
  operator = "and",
  match_funcs = c(age_diff, exact_match))

# Nested match criteria 'm.cri.1' OR 'm.cri.2'
n.cri <- sub_criteria(
  m.cri.1, m.cri.2,
  operator = "or")

# Record linkage with additional match criteria
p3 <- links(
  criteria = stages,
  sub_criteria = list(cr1 = m.cri.1,
                     cr2 = m.cri.2))

# Record linkage with additional nested match criteria
p4 <- links(
  criteria = stages,
  sub_criteria = list(cr1 = n.cri,
                     cr2 = n.cri))

dfr$p1 <- p1; dfr$p2 <- p2
dfr$p3 <- p3; dfr$p4 <- p4

head(dfr)

```

link_wf	<i>Record linkage</i>
---------	-----------------------

Description

Deterministic and probabilistic record linkage Assign unique identifiers to records based on partial, nested or calculated probabilities.

Usage

```
links_af_probabilistic(  
  attribute,  
  blocking_attribute = NULL,  
  cmp_func = diyar::exact_match,  
  attr_threshold = 1,  
  probabilistic = TRUE,  
  m_probability = 0.95,  
  u_probability = NULL,  
  score_threshold = 1,  
  repeats_allowed = FALSE,  
  permutations_allowed = FALSE,  
  data_source = NULL,  
  ignore_same_source = TRUE,  
  display = "none"  
)
```

```
links_wf_probabilistic(  
  attribute,  
  blocking_attribute = NULL,  
  cmp_func = diyar::exact_match,  
  attr_threshold = 1,  
  probabilistic = TRUE,  
  m_probability = 0.95,  
  u_probability = NULL,  
  score_threshold = 1,  
  id_1 = NULL,  
  id_2 = NULL,  
  return_weights = FALSE,  
  ...  
)
```

```
prob_score_range(attribute, m_probability = 0.95, u_probability = NULL)
```

Arguments

attribute	[atomic list data.frame matrix d_attribute]. Attributes to compare.
blocking_attribute	[atomic]. Passed to criteria in links .
cmp_func	[list function]. String comparators for each attribute. See Details.
attr_threshold	[list numeric number_line]. Weight-thresholds for each cmp_func. See Details.
probabilistic	[logical]. If TRUE, scores are assigned base on Fellegi-Sunter model for probabilistic record linkage. See Details.
m_probability	[list numeric]. The probability that a matching records are the same entity.
u_probability	[list numeric]. The probability that a matching records are not the same entity.
score_threshold	[numeric number_line]. Score-threshold for linked records. See Details.
repeats_allowed	[logical] Passed to repeats_allowed in links .
permutations_allowed	[logical] Passed to permutations_allowed in links .
data_source	[character]. Passed to data_source in links .
ignore_same_source	[logical] Passed to ignore_same_source in links .
display	[character]. Passed to display in links .
id_1	[list numeric]. Record id or index of one half of a record-pair.
id_2	[list numeric]. Record id or index of one half of a record-pair.
return_weights	If TRUE, returns the match-weights and score-thresholds for record pairs.
...	Arguments passed to links

Details

links_wf_probabilistic() - A wrapper function of [links](#) with a with a specific [sub_criteria](#) and to achieve to achieve probabilistic record linkage It excludes functionalities for the nested and multi-stage linkage. links_wf_probabilistic() requires a score_threshold in advance. To help with this, prob_score_range() can be used to return the range of scores attainable for a given set of attribute, m and u-probabilities. Additionally, id_1 and id_2 can be used to link specific records pairs, aiding the review of potential scores.

links_af_probabilistic() - A simpler version of [links](#). It excludes functionalities for the batched, nested and multi-stage linkage. links_af_probabilistic() requires a score_threshold in advance, however, since it exports the match weights, the score_threshold can be changed after the analysis.

Value

[pid](#); list


```

recursive = FALSE,
check_duplicates = FALSE)

# Each implementation can lead to different results
summary(pids_1a$pid)
summary(pids_1b$pid)
summary(pids_1c$pid)

## End(Not run)

# Weighted (non-probabilistic) comparison of forename, middlename and age difference
criteria_2 <- as.list(patient_records[c("forename", "middlename", "dateofbirth")])
age_diff <- function(x, y){
  diff <- abs(as.numeric(x) - as.numeric(y))
  wgt <- diff %in% 0:(365 * 10) & !is.na(diff)
  wgt
}

pids_2a <- links_af_probabilistic(attribute = criteria_2,
                                blocking_attribute = patient_records$surname,
                                cmp_func = c(exact_match, exact_match, age_diff),
                                score_threshold = number_line(3, 5),
                                probabilistic = FALSE,
                                display = "stats")

# Larger weights can be assigned to particular attributes through `cmp_func`
# For example, a smaller age difference can contribute a higher score (e.g 0 to 3)
age_diff_2 <- function(x, y){
  diff <- as.numeric(abs(x - y))
  wgt <- diff %in% 0:(365 * 10) & !is.na(diff)
  wgt[wgt] <- match(as.numeric(cut(diff[wgt], 3)), 3:1)
  wgt
}
pids_2b <- links_af_probabilistic(attribute = criteria_2,
                                blocking_attribute = patient_records$surname,
                                cmp_func = c(exact_match, exact_match, age_diff_2),
                                score_threshold = number_line(3, 5),
                                probabilistic = FALSE,
                                display = "stats")

head(pids_2a$pid_weights, 10)
head(pids_2b$pid_weights, 10)

```

listr

Grammatical lists.

Description

A convenience function to format atomic vectors as a written list.

Usage

```
listr(x, sep = ", ", conj = " and ", lim = Inf)
```

Arguments

x atomic vector.
 sep Separator.
 conj Final separator.
 lim Elements to include in the list. Other elements are abbreviated to " ...".

Value

character.

Examples

```
listr(1:5)
listr(1:5, sep = "; ")
listr(1:5, sep = "; ", conj = " and")
listr(1:5, sep = "; ", conj = " and", lim = 2)
```

make_ids	<i>Convert an edge list to record identifiers.</i>
----------	--

Description

Convert an edge list to record identifiers.

Usage

```
make_ids(x_pos, y_pos, id_length = max(x_pos, y_pos))
```

Arguments

x_pos [integer]. Index of first half of a record-pair.
 y_pos [integer]. Index of second half of a record-pair.
 id_length Length of the record identifier.

Details

Record groups from non-recursive links have the lowest record ID (sn) in the set as their group ID.

Value

list

Examples

```
make_ids(x_pos = rep(7, 7), y_pos = 1:7)
make_ids(x_pos = c(1, 6), y_pos = 6:7)
make_ids(x_pos = 1:5, y_pos = c(1, 1, 2, 3, 4))
```

make_pairs	<i>Combinations and permutations of record-sets.</i>
------------	--

Description

Combinations and permutations of record-sets.

Usage

```
sets(n, r, permutations_allowed = TRUE, repeats_allowed = TRUE)
```

```
make_sets(
  x,
  r,
  strata = NULL,
  permutations_allowed = TRUE,
  repeats_allowed = TRUE
)
```

```
make_pairs(
  x,
  strata = NULL,
  repeats_allowed = TRUE,
  permutations_allowed = FALSE
)
```

```
make_pairs_wf_source(..., data_source = NULL)
```

Arguments

n	[integer]. Size of Vector.
r	[integer]. Number of elements in a set.
permutations_allowed	[logical]. If TRUE, permutations of the same set are included.
repeats_allowed	[logical]. If TRUE, repeat values are included in each set.
x	[atomic]. Vector.
strata	Subsets of x. Blocking attribute. Limits the creation of combinations or permutations to those from the same strata.
...	Arguments passed to make_pairs .
data_source	[character]. Data source identifier. Limits the creation of combinations or permutations to those from a different data_source

Details

set() - Create r-set combinations or permutations of n observations.

make_set() - Create r-set combinations or permutations of vector x.

make_pairs() - Create 2-set combinations or permutations of vector x.

make_pairs_wf_source() - Create 2-set combinations or permutations of vector x that are from different sources (data_source).

Value

A list of a vector's elements and corresponding indexes.

See Also

[eval_sub_criteria](#)

Examples

```
sets(4, 2)
sets(4, 2, repeats_allowed = FALSE, permutations_allowed = FALSE)
make_sets(month.abb[1:4], 2)
make_sets(month.abb[1:4], 3)

make_pairs(month.abb[1:4])
make_pairs(month.abb[1:4], strata = c(1, 1, 2, 2))
make_pairs_wf_source(month.abb[1:4], data_source = c(1, 1, 2, 2))
```

make_s4_ids

Create epid and pid objects with index of matching records

Description

Create epid and pid objects with index of matching records

Usage

```
make_episodes(
  x_pos,
  y_pos,
  x_val,
  date,
  case_nm,
  wind_id,
  wind_nm,
  from_last,
  data_source,
  data_links,
```

```

    iteration,
    options,
    episode_unit
)

make_pids(
    x_pos,
    y_pos,
    x_val,
    link_id,
    pid_cri,
    data_source,
    data_links,
    iteration
)

```

Arguments

x_pos	[integer]. Index of one half of a record pair.
y_pos	[integer]. Index of one half of a record pair.
x_val	[integer]. Value of one half of a record pair.
date	[date datetime integer number_line]. Record date or period.
case_nm	[integer character] Record type in regards to case assignment (sub_criteria [Encoded]).
wind_id	[integer]. Unique reference ID for each match.
wind_nm	[list]. Type of window i.e. "Case" or "Recurrence".
from_last	[logical]. Chronological order of episode tracking i.e. ascending (TRUE) or descending (FALSE).
data_source	[character]. Source ID for each record.
data_links	[list character]. <code>data_source</code> required in each record-group. A record-group without records from these <code>data_sources</code> will be unlinked .
iteration	The iteration when a record was matched to it's group (.Data).
options	[list]. Some options passed to the instance of episodes .
episode_unit	[character]. Time unit for <code>case_length</code> and <code>recurrence_length</code> . See episodes
link_id	[integer]. Unique reference ID for each match.
pid_cri	Match stage of the step-wise linkage.

merge_identifiers *Merge group identifiers*

Description

Consolidate two group identifiers.

Usage

```
merge_ids(...)

## Default S3 method:
merge_ids(id1, id2, tie_sort = NULL, expand = TRUE, shrink = FALSE, ...)

## S3 method for class 'pid'
merge_ids(id1, id2, tie_sort = NULL, expand = TRUE, shrink = FALSE, ...)

## S3 method for class 'epid'
merge_ids(id1, id2, tie_sort = NULL, expand = TRUE, shrink = FALSE, ...)

## S3 method for class 'pane'
merge_ids(id1, id2, tie_sort = NULL, expand = TRUE, shrink = FALSE, ...)
```

Arguments

...	Other arguments
id1	[integer epid pid pane].
id2	[integer epid pid pane].
tie_sort	[atomic]. Preferential order for breaking tied matches.
expand	[logical]. If TRUE, id1 gains new records if id2 indicates a match. <i>Not interchangeable with shrink.</i>
shrink	[logical]. If TRUE, id1 loses existing records id2 does not indicate a match. <i>Not interchangeable with expand.</i>

Details

Groups in id1 are expanded or shrunk by groups in id2.

A unique group with only one record is considered a non-matching record.

Note that the expand and shrink features are not interchangeable. The outcome when shrink is TRUE is not the same when expand is FALSE. See Examples.

See Also

[links](#); [links_af_probabilistic](#)

Examples

```
id1 <- rep(1, 5)
id2 <- c(2, 2, 3, 3, 3)
merge_ids(id1, id2, shrink = TRUE)

id1 <- c(rep(1, 3), 6, 7)
id2 <- c(2,2,3,3,3)
merge_ids(id1, id2, shrink = TRUE)
merge_ids(id1, id2, expand = FALSE)
```

```
id1 <- rep(1, 5)
id2 <- c(1:3, 4, 4)
merge_ids(id1, id2, shrink = TRUE)
merge_ids(id1, id2, expand= FALSE)

data(missing_staff_id)
dfr <- missing_staff_id
id1 <- links(dfr[[5]])
id2 <- links(dfr[[6]])
merge_ids(id1, id2)
```

number_line	number_line
-------------	-------------

Description

A range of numeric values.

Usage

```
number_line(l, r, id = NULL, gid = NULL)

as.number_line(x)

is.number_line(x)

left_point(x)

left_point(x) <- value

right_point(x)

right_point(x) <- value

start_point(x)

start_point(x) <- value

end_point(x)

end_point(x) <- value

number_line_width(x)

reverse_number_line(x, direction = "both")

shift_number_line(x, by = 1)
```

```

expand_number_line(x, by = 1, point = "both")

invert_number_line(x, point = "both")

number_line_sequence(
  x,
  by = NULL,
  length.out = 1,
  fill = TRUE,
  simplify = FALSE
)

```

Arguments

l	[numeric-based]. Left point of the number_line.
r	[numeric-based]. Right point of the number_line. Must be able to be coerced to a numeric object.
id	[integer]. Unique element identifier. Optional.
gid	[integer]. Unique group identifier. Optional.
x	[number_line]
value	[numeric based]
direction	[character]. Type of number_line reverse. Options are; "increasing", "decreasing" or "both" (default).
by	[integer]. Increment or decrement. Passed to seq() in number_line_sequence().
point	[character]. "start", "end", "left" or "right" point.
length.out	[integer]. Number of splits. For example, 1 for two parts or 2 for three parts. Passed to seq().
fill	[logical]. Retain (TRUE) or drop (FALSE) the remainder of an uneven split.
simplify	[logical]. If TRUE, returns a sequence of finite numbers.

Details

A number_line object represents a range of numbers. It is made up of a start and end point as the lower and upper ends of the range respectively. The location of the start point - left or right, determines whether it is an "increasing" or "decreasing" number_line. This is the direction of the number_line.

reverse_number_line() - reverse the direction of a number_line. A reversed number_line has its left and right points swapped. The direction argument specifies which type of number_line will be reversed. number_line with non-finite start or end points (i.e. NA, NaN and Inf) can't be reversed.

shift_number_line() - Shift a number_line towards the positive or negative end of the number line.

expand_number_line() - Increase or decrease the width of a number_line.

`invert_number_line()` - Change the left or right points from a negative to positive value or vice versa.

`number_line_sequence()` - Split a `number_line` into equal parts (`length.out`) or by a fixed recurring width (`by`).

Value

`number_line`

See Also

[overlaps](#); [set_operations](#); [episodes](#); [links](#)

Examples

```
number_line(-100, 100)

# Also compatible with other numeric based object classes
number_line(as.POSIXct("2019-05-15 13:15:07", tz = "UTC"),
            as.POSIXct("2019-05-15 15:17:10", tz = "UTC"))

# Coerce compatible object classes to `number_line` objects
as.number_line(5.1); as.number_line(as.Date("2019-10-21"))

# A test for number_line objects
a <- number_line(as.Date("2019-04-25"), as.Date("2019-01-01"))
is.number_line(a)

# Structure of a number_line object
left_point(a); right_point(a); start_point(a); end_point(a)

# Reverse number_line objects
reverse_number_line(number_line(as.Date("2019-04-25"), as.Date("2019-01-01")))
reverse_number_line(number_line(200, -100), "increasing")
reverse_number_line(number_line(200, -100), "decreasing")

c <- number_line(5, 6)
# Shift number_line objects towards the positive end of the number line
shift_number_line(x = c(c, c), by = c(2, 3))
# Shift number_line objects towards the negative end of the number line
shift_number_line(x = c(c, c), by = c(-2, -3))

# Change the duration, width or length of a number_line object
d <- c(number_line(3, 6), number_line(6, 3))

expand_number_line(d, 2)
expand_number_line(d, -2)
expand_number_line(d, c(2,-1))
expand_number_line(d, 2, "start")
expand_number_line(d, 2, "end")

# Invert `number_line` objects
```



```
e <- c(number_line(3, 6), number_line(-3, -6), number_line(-3, 6))
e
invert_number_line(e)
invert_number_line(e, "start")
invert_number_line(e, "end")

# Split number line objects
x <- number_line(Sys.Date() - 5, Sys.Date())
x
number_line_sequence(x, by = 2)
number_line_sequence(x, by = 4)
number_line_sequence(x, by = 4, fill = FALSE)
number_line_sequence(x, length.out = 2)
```

number_line-class	number_line object
-------------------	--------------------

Description

S4 objects representing a range of numeric values

Usage

```
## S4 method for signature 'number_line'
show(object)

## S4 method for signature 'number_line'
rep(x, ...)

## S4 method for signature 'number_line'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'number_line'
x[[i, j, ..., exact = TRUE]]

## S4 replacement method for signature 'number_line'
x[i, j, ...] <- value

## S4 replacement method for signature 'number_line'
x[[i, j, ...]] <- value

## S4 method for signature 'number_line'
x$name

## S4 replacement method for signature 'number_line'
x$name <- value

## S4 method for signature 'number_line'
```

```

c(x, ...)

## S3 method for class 'number_line'
unique(x, ...)

## S3 method for class 'number_line'
seq(x, precision = NULL, fill = FALSE, ...)

## S3 method for class 'number_line'
sort(x, decreasing = FALSE, ...)

## S3 method for class 'number_line'
format(x, ...)

## S3 method for class 'number_line'
as.list(x, ...)

## S3 method for class 'number_line'
as.data.frame(x, ...)

```

Arguments

object	object
x	x
...	...
i	i
j	j
drop	drop
exact	exact
value	value
name	slot name
precision	Round precision
fill	[logical]. Retain (TRUE) or drop (FALSE) the remainder of an uneven split.
decreasing	If TRUE, sort in descending order.

Slots

start First value in the range.

id Unique element id. Optional.

gid Unique group id. Optional.

.Data Length, duration or width of the range.

overlaps	<i>Overlapping number line objects</i>
----------	--

Description

Identify overlapping [number_line](#) objects

Usage

`overlaps(x, y, methods = 8)`

`overlap(x, y)`

`none(x, y)`

`exact(x, y)`

`across(x, y)`

`x_across_y(x, y)`

`y_across_x(x, y)`

`chain(x, y)`

`x_chain_y(x, y)`

`y_chain_x(x, y)`

`aligns_start(x, y)`

`x_aligns_start_y(x, y)`

`y_aligns_start_x(x, y)`

`aligns_end(x, y)`

`x_aligns_end_y(x, y)`

`y_aligns_end_x(x, y)`

`inbetween(x, y)`

`x_inbetween_y(x, y)`

`y_inbetween_x(x, y)`

```

overlap_method(x, y)

include_overlap_method(methods)

exclude_overlap_method(methods)

overlap_method_codes(methods)

overlap_method_names(methods)

```

Arguments

x	[number_line]
y	[number_line]
methods	[character integer]. Type of overlap. See <code>as.data.frame(diyar::overlap_methods\$options)</code> for options.

Details

There are 6 mutually exclusive types of overlap;

- `exact()` - identical `start_point` and `end_point` points.
- `inbetween()` - Both `start_point` and `end_point` of one `number_line` object are within the `start_point` and `end_point` of another.
- `across()` - Only the `start_point` or `end_point` of one `number_line` object is in between the `start_point` and `end_point` of another.
- `chain()` - `end_point` of one `number_line` object is identical to the `start_point` of another.
- `aligns_start()` - identical `start_point` only.
- `aligns_end()` - identical `end_point` only.

Except `exact()`, each type of overlap has two variations;

- `x_‘method’_y()` - `number_line-x` starts before `number_line-y`.
- `y_‘method’_x()` - `number_line-y` starts before `number_line-x`.

There are two mutually inclusive types of overlap;

- `overlap()` - a convenient option to select "ANY" and "ALL" type of overlap.
- `none()` - a convenient option to select "NO" type of overlap.

Selecting multiple types of overlap;

- `overlaps()` - select specific type(s) of overlap.
- `overlap_method()` - return the type of overlap for a pair of `number_line` objects.
- `overlap_method_codes()` - return the corresponding overlap method code for a specific type(s) of overlap.
- `overlap_method_names()` - return the corresponding type(s) of overlap for a specific overlap code.

- `include_overlap_method()` - return a character(1) value for specified type(s) of overlap.
- `exclude_overlap_method()` - return a character(1) value for all type(s) of overlap except those specified.

Value

logical; character

See Also

[number_line](#); [set_operations](#)

Examples

```
a <- number_line(-100, 100)
g <- number_line(100, 100)
overlaps(a, g)

# It's neither an "exact" or "chain"-overlap
overlaps(a, g, methods = "exact|chain")

# It's an "aligns_end"-overlap
overlap_method(a, g)
overlaps(a, g, methods = "exact|chain|x_aligns_end_y")

# Corresponding overlap code
overlap_method_codes("exact|chain|x_aligns_end_y")
include_overlap_method(c("exact", "chain", "x_aligns_end_y"))

# Corresponding overlap name
overlap_method_names(overlap_method_codes("exact|chain|x_aligns_end_y"))

# Every other type overlap
exclude_overlap_method(c("exact", "chain", "x_aligns_end_y"))
overlap_method_names(exclude_overlap_method(c("exact", "chain", "x_aligns_end_y")))

# All the above is based on tests for each specific type of overlap as seen below
none(a, g)
exact(a, g)
across(a, g)
x_across_y(a, g)
y_across_x(a, g)
chain(a, g)
x_chain_y(a, g)
y_chain_x(a, g)
inbetween(a, g)
x_inbetween_y(a, g)
y_inbetween_x(a, g)
aligns_start(a, g)
x_aligns_start_y(a, g)
y_aligns_start_x(a, g)
aligns_end(a, g)
```

```
x_aligns_end_y(a, g)
y_aligns_end_x(a, g)
```

pane-class	pane <i>object</i>
------------	--------------------

Description

S4 objects storing the result of [partitions](#).

Usage

```
is.pane(x)

as.pane(x)

## S3 method for class 'pane'
format(x, ...)

## S3 method for class 'pane'
unique(x, ...)

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

## S3 method for class 'pane_summary'
print(x, ...)

## S3 method for class 'pane'
as.data.frame(x, ..., decode = TRUE)

## S3 method for class 'pane'
as.list(x, ..., decode = TRUE)

## S4 method for signature 'pane'
show(object)

## S4 method for signature 'pane'
rep(x, ...)

## S4 method for signature 'pane'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'pane'
x[[i, j, ..., exact = TRUE]]
```

```
## S4 method for signature 'pane'
c(x, ...)
```

Arguments

x	x
...	...
object	object
decode	If TRUE, data is decoded
i	i
j	j
drop	drop
exact	exact

Slots

sn Unique record identifier.

.Data Unique pane identifier.

case_nm Record type in regards to index assignment.

window_list A list of considered windows for each pane.

dist_pane_index The difference between each event and it's index event.

pane_dataset Data sources in each pane.

pane_interval The start and end dates of each pane. A [number_line](#) object.

pane_length The duration or length of (pane_interval).

pane_total The number of records in each pane.

options Some options passed to the instance of [partitions](#).

window_matched A list of matched windows for each pane.

Examples

```
# A test for pane objects
pn <- partitions(date = 1, by = 1)
is.pane(pn); is.pane(2)
```

partitions	<i>Distribute events into specified intervals.</i>
------------	--

Description

Distribute events into groups defined by time or numerical intervals. Each set of linked records are assigned a unique identifier with relevant group-level data.

Usage

```
partitions(
  date,
  window = NULL,
  windows_total = 1,
  separate = FALSE,
  sn = NULL,
  strata = NULL,
  data_links = "ANY",
  custom_sort = NULL,
  group_stats = FALSE,
  data_source = NULL,
  by = NULL,
  length.out = NULL,
  fill = TRUE,
  display = "none",
  precision = 1
)
```

Arguments

date	[date datetime integer number_line]. Event date or period.
window	[integer number_line]. Numeric or time intervals.
windows_total	[integer number_line]. Minimum number of matched windows required for a pane. See details
separate	[logical]. If TRUE, events matched to different windows are not linked.
sn	[integer]. Unique record identifier. Useful for creating familiar pane identifiers.
strata	[atomic]. Subsets of the dataset. Panes are created separately for each strata.
data_links	[list character]. A set of data_sources required in each pane . A pane without records from these data_sources will be unlinked. See Details.
custom_sort	[atomic]. Preferred order for selecting "index" events.
group_stats	[logical]. If TRUE (default), the returned pane object will include group specific information like panes start and end dates.
data_source	[character]. Unique data source identifier. Adds the list of datasets in each pane to the pane . Useful when the data is from multiple sources.

by	[integer]. Width of splits.
length.out	[integer]. Number of splits.
fill	[logical]. Retain (TRUE) or drop (FALSE) the remainder of an uneven split.
display	[character]. Display a status update. Options are; "none" (default), "progress" or "stats".
precision	Round precision

Details

Each assigned group is referred to as a [pane](#). A [pane](#) consists of events within a specific time or numerical intervals (window).

Each window must cover a separate interval. Overlapping windows are merged before events are distributed into panes. Events that occur over two windows are assigned to the last one listed.

Alternatively, you can create windows by splitting a period into equal parts (`length.out`), or into a sequence of intervals with fixed widths (`by`).

By default, the earliest event is taken as the "Index" event of the [pane](#). An alternative can be chosen with `custom_sort`. Note that this is simply a convenience option because it has no bearing on how groups are assigned.

`partitions()` will categorise records into 3 types;

- "Index" - Index event/record of the pane.
- "Duplicate_I" - Duplicate of the "Index" record.
- "Skipped" - Records that are not assigned to a pane.

Every element in `data_links` must be named "l" (links) or "g" (groups). Unnamed elements of `data_links` will be assumed to be "l".

- If named "l", only groups with records from every listed `data_source` will be retained.
- If named "g", only groups with records from any listed `data_source` will be retained.

NA values in strata excludes records from the partitioning process.

See `vignette("episodes")` for more information.

Value

[pane](#)

See Also

[pane](#); [number_line_sequence](#); [episodes](#); [links](#); [overlaps](#); [number_line](#); [schema](#)

Examples

```

events <- c(30, 2, 11, 10, 100)
windows <- number_line(c(1, 9, 25), c(3, 12, 35))

events
partitions(date = events, length.out = 3, separate = TRUE)
partitions(date = events, by = 10, separate = TRUE)
partitions(date = events, window = windows, separate = TRUE)
partitions(date = events, window = windows, separate = FALSE)
partitions(date = events, window = windows, separate = FALSE, windows_total = 4)

```

pid-class

pid *objects*

Description

S4 objects storing the result of [links](#).

Usage

```

is.pid(x)

as.pid(x, ...)

## S3 method for class 'pid'
format(x, ...)

## S3 method for class 'pid'
unique(x, ...)

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

## S3 method for class 'pid_summary'
print(x, ...)

## S3 method for class 'pid'
as.data.frame(x, ..., decode = TRUE)

## S3 method for class 'pid'
as.list(x, ..., decode = TRUE)

## S4 method for signature 'pid'
show(object)

## S4 method for signature 'pid'

```

```

rep(x, ...)

## S4 method for signature 'pid'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'pid'
x[[i, j, ..., exact = TRUE]]

## S4 method for signature 'pid'
c(x, ...)

```

Arguments

x	x
...	...
object	object
decode	If TRUE, data is decoded
i	i
j	j
drop	drop
exact	exact

Slots

sn Unique record identifier.

.Data Unique group identifier.

link_id Unique reference ID for each match.

pid_cri Match stage of the step-wise linkage.

pid_dataset Data sources in each group.

pid_total The number of records in each group.

iteration The iteration when a record was matched to it's group (.Data).

Examples

```

# A test for pid objects
pd <- links(criteria = 1)
is.pid(pd); is.pid(2)

```

```
predefined_tests      Predefined logical tests in diyar
```

Description

A collection of predefined logical tests used with [sub_criteria](#) objects

Usage

```
exact_match(x, y)

range_match(x, y, range = 10)

prob_link(
  x,
  y,
  cmp_func,
  attr_threshold,
  score_threshold,
  probabilistic,
  return_weights = FALSE
)

true(x, y)

false(x, y)
```

Arguments

<code>x</code>	Attribute(s) to be compared against.
<code>y</code>	Attribute(s) to be compared by.
<code>range</code>	Difference between <code>y</code> and <code>x</code> .
<code>cmp_func</code>	Logical tests such as string comparators. See links_wf_probabilistic .
<code>attr_threshold</code>	Matching set of weight thresholds for each result of <code>cmp_func</code> . See links_wf_probabilistic .
<code>score_threshold</code>	Score threshold determining matched or linked records. See links_wf_probabilistic .
<code>probabilistic</code>	If TRUE, matches determined through a score derived base on Fellegi-Sunter model for probabilistic linkage. See links_wf_probabilistic .
<code>return_weights</code>	If TRUE, returns the match-weights and score-thresholds for record pairs.

Details

```
exact_match() - test that x == y
range_match() - test that x ≤ y ≤ (x + range)
```

`prob_link()` - Test that a record-pair relate to the same entity based on Fellegi and Sunter (1969) model for deciding if two records belong to the same entity.

In summary, record-pairs are created and categorised as matches and non-matches (`attr_threshold`) with user-defined functions (`cmp_func`). If `probabilistic` is TRUE, two probabilities (`m` and `u`) are used to calculate weights for matches and non-matches. The `m`-probability is the probability that matched records are actually from the same entity i.e. a true match, while `u`-probability is the probability that matched records are not from the same entity i.e. a false match. Record-pairs whose total score are above a certain threshold (`score_threshold`) are assumed to belong to the same entity.

Agreement (match) and disagreement (non-match) scores are calculated as described by Asher et al. (2020).

For each record pair, an agreement for attribute i is calculated as;

$$\log_2(m_i/u_i)$$

For each record pair, a disagreement score for attribute i is calculated as;

$$\log_2((1 - m_i)/(1 - u_i))$$

where m_i and u_i are the `m` and `u`-probabilities for each value of attribute i .

Note that each probability is calculated as a combined probability for the record pair. For example, if the values of the record-pair have `u`-probabilities of 0.1 and 0.2 respectively, then the `u`-probability for the pair will be 0.02.

Missing data (NA) are considered non-matches and assigned a `u`-probability of 0.

Examples

```
`exact_match`
exact_match(x = 1, y = 1)
exact_match(x = 1, y = 2)

`range_match`
range_match(x = 10, y = 16, range = 6)
range_match(x = 16, y = 10, range = 6)
```

reframe

Modify sub_criteria objects

Description

Modify the attributes of a `sub_criteria` object.

Usage

```
reframe(x, ...)

## S3 method for class 'sub_criteria'
reframe(x, func = identity, ...)
```

Arguments

```
x           [sub_criteria].
...         Arguments passed to methods.
func       [function]. Transformation function.
```

See Also

[sub_criteria](#); [eval_sub_criteria](#); [attr_eval](#)

Examples

```
s_cri <- sub_criteria(month.abb, month.name)
reframe(s_cri, func = function(x) x[12])
reframe(s_cri, func = function(x) x[12:1])
reframe(s_cri, func = function(x) attrs(x[1:6], x[7:12]))
```

schema

Schema diagram for group identifiers

Description

Create schema diagrams for [number_line](#), [epid](#), [pid](#) and [pane](#) objects.

Usage

```
schema(x, ...)

## S3 method for class 'number_line'
schema(x, show_labels = c("date", "case_overlap_methods"), ...)

## S3 method for class 'epid'
schema(
  x,
  title = NULL,
  show_labels = c("length_arrow"),
  show_skipped = TRUE,
  show_non_finite = FALSE,
  theme = "dark",
  seed = NULL,
  custom_label = NULL,
```

```

    ...
  )

## S3 method for class 'pane'
schema(
  x,
  title = NULL,
  show_labels = c("window_label"),
  theme = "dark",
  seed = NULL,
  custom_label = NULL,
  ...
)

## S3 method for class 'pid'
schema(
  x,
  title = NULL,
  show_labels = TRUE,
  theme = "dark",
  orientation = "by_pid",
  seed = NULL,
  custom_label = NULL,
  ...
)

```

Arguments

x	[number_line epid pid pane]
...	Other arguments.
show_labels	[logical character]. Show/hide certain parts of the schema. See Details.
title	[character]. Plot title.
show_skipped	[logical]. Show/hide "Skipped" records.
show_non_finite	[logical]. Show/hide records with non-finite date values.
theme	[character]. Options are "dark" or "light".
seed	[integer]. See <code>set.seed</code> . Used to get a consistent arrangement of items in the plot.
custom_label	[character]. Custom label for each record of the identifier.
orientation	[character]. Show each record of a pid object within its group id ("by_pid") or its pid_cri ("by_pid_cri")

Details

A visual aid to describe the data linkage ([links](#)), episode tracking ([episodes](#)) or partitioning process ([partitions](#)).

show_labels **options (multi-select)**

- schema.epid - **TRUE, FALSE**, "sn", "epid", "date", "case_nm", "wind_nm", "length", "length_arrow", "case_overlap_methods" or "recurrence_overlap_methods"
- schema.pane - **TRUE, FALSE**, "sn", "pane", "date", "case_nm" or "window_label"
- schema.pid - **TRUE, FALSE**, "sn" or "pid"

Value

ggplot objects

Examples

```
schema(number_line(c(1, 2), c(2, 1)))

schema(episodes(1:10, 2))

schema(partitions(1:10, by = 2, separate = TRUE))

schema(links(list(c(1, 1, NA, NA), c(NA, 1, 1, NA))))
```

set_operations	<i>Set operations on number line objects</i>
----------------	--

Description

Perform set operations on a pair of [[number_line](#)]s.

Usage

```
union_number_lines(x, y)

intersect_number_lines(x, y)

subtract_number_lines(x, y)
```

Arguments

```
x           [number\_line]
y           [number\_line]
```

Details

`union_number_lines()` - Combined the range of x and that of y

`intersect_number_line()` - Subset of x that overlaps with y and vice versa

`subtract_number_lines()` - Subset of x that does not overlap with y and vice versa.

The direction of the returned [[number_line](#)] will be that of the widest one (x or y). If x and y have the same length, it'll be an "increasing" direction.

If x and y do not overlap, NA ("NA ?? NA") is returned.

Value

```
[number_line]; list
```

See Also

```
number_line; overlaps
```

Examples

```
n1_1 <- c(number_line(1, 5), number_line(1, 5), number_line(5, 9))
n1_2 <- c(number_line(1, 2), number_line(2, 3), number_line(0, 6))

# Union
n1_1; n1_2; union_number_lines(n1_1, n1_2)

n1_3 <- number_line(as.Date(c("01/01/2020", "03/01/2020", "09/01/2020"), "%d/%m/%Y"),
  as.Date(c("09/01/2020", "09/01/2020", "25/12/2020"), "%d/%m/%Y"))

n1_4 <- number_line(as.Date(c("04/01/2020", "01/01/2020", "01/01/2020"), "%d/%m/%Y"),
  as.Date(c("05/01/2020", "05/01/2020", "03/01/2020"), "%d/%m/%Y"))

# Intersect
n1_3; n1_4; intersect_number_lines(n1_3, n1_4)

# Subtract
n1_3; n1_4; subtract_number_lines(n1_3, n1_4)
```

staff_records

Datasets in diyar package

Description

Datasets in diyar package

Usage

```
data(staff_records)

data(missing_staff_id)

data(infections)

data(infections_2)

data(infections_3)
```

```
data(infections_4)
data(hospital_admissions)
data(patient_list)
data(patient_list_2)
data(hourly_data)
data(0pes)
data(episode_unit)
data(overlap_methods)
data(patient_records)
```

Format

```
data.frame
data.frame
data.frame
data.frame
data.frame
data.frame
data.frame
data.frame
data.frame
An object of class data . frame with 5 rows and 4 columns.
data.frame
data.frame
list
list
data.frame
```

Details

staff_records - Staff record with some missing data
missing_staff_id - Staff records with missing staff identifiers
infections, infections_2, infections_3 and infections_4 - Reports of bacterial infections
hospital_admissions - Hospital admissions and discharges
patient_list & patient_list_2 - Patient list with some missing data
Hourly data

Opes - List of individuals with the same name
 Duration in seconds for each 'episode_unit'
 Permutations of [number_line](#) overlap methods

Examples

```
data(staff_records)
data(missing_staff_id)
data(infections)
data(infections_2)
data(infections_3)
data(infections_4)
data(hospital_admissions)
data(patient_list)
data(patient_list_2)
data(hourly_data)
data(Opes)
data(episode_unit)
data(overlap_methods)
data(patient_records)
```

sub_criteria

Match criteria

Description

Match criteria for record linkage with [links](#) and [episodes](#)

Usage

```
sub_criteria(
  ...,
  match_funcs = c(exact = diyar::exact_match),
  equal_funcs = c(exact = diyar::exact_match),
  operator = "or"
)

attrs(..., .obj = NULL)

eval_sub_criteria(x, ...)

## S3 method for class 'sub_criteria'
print(x, ...)

## S3 method for class 'sub_criteria'
format(x, show_levels = FALSE, ...)

## S3 method for class 'sub_criteria'
```

```
eval_sub_criteria(
  x,
  x_pos = seq_len(max(attr_eval(x))),
  y_pos = rep(1L, length(x_pos)),
  check_duplicates = TRUE,
  depth = 0,
  ...
)
```

Arguments

...	[atomic] Attributes passed to or <code>eval_sub_criteria()</code> or <code>eval_sub_criteria()</code> Arguments passed to methods for <code>eval_sub_criteria()</code>
match_funcs	[function]. User defined logical test for matches.
equal_funcs	[function]. User defined logical test for identical record sets (all attributes of the same record).
operator	[character]. Options are "and" or "or".
.obj	[data.frame list]. Attributes.
x	[sub_criteria]. Attributes.
show_levels	[logical]. If TRUE, show recursive depth for each logic statement of the match criteria.
x_pos	[integer]. Index of one half of a record pair.
y_pos	[integer]. Index of one half of a record pair.
check_duplicates	[logical]. If FALSE, does not check duplicate values. The result of the initial check will be recycled.
depth	[integer]. First order of recursion.

Details

`sub_criteria()` - Create a match criteria as a `sub_criteria` object. A `sub_criteria` object contains attributes to be compared, logical tests for the comparisons (see [predefined_tests](#) for examples) and another set of logical tests to determine identical records.

`attrs()` - Create a `d_attribute` object - a collection of atomic objects that can be passed to `sub_criteria()` as a single attribute.

`eval_sub_criteria()` - Evaluates a `sub_criteria` object.

At each iteration of [links](#) or [episodes](#), record-pairs are created from each attribute of a `sub_criteria` object. `eval_sub_criteria()` evaluates each record-pair using the `match_funcs` and `equal_funcs` functions of a `sub_criteria` object. See [predefined_tests](#) for examples of `match_funcs` and `equal_funcs`.

User-defined functions are also permitted as `match_funcs` and `equal_funcs`. Such functions must meet three requirements:

1. It must be able to compare the attributes.

2. It must have two arguments named ``x`` and ``y``, where ``y`` is the value for one observation being compared against all other observations (``x``).
3. It must return a logical object i.e. TRUE or FALSE.

`attrs()` is useful when the match criteria requires an interaction between the multiple attributes. For example, attribute 1 + attribute 2 > attribute 3.

Every attribute, including those in `attrs()`, must have the same length or a length of 1.

Value

`sub_criteria`

See Also

[predefined_tests](#); [links](#); [episodes](#); [eval_sub_criteria](#)

Examples

```
# Attributes
attr_1 <- c(30, 28, 40, 25, 25, 29, 27)
attr_2 <- c("M", "F", "U", "M", "F", "U", "M")

# A match criteria
## Example 1 - A maximum difference of 10 in attribute 1
s_cri1 <- sub_criteria(attr_1, match_funcs = range_match)
s_cri1

# Evaluate the match criteria
## Compare the first element of 'attr_1' against all other elements
eval_sub_criteria(s_cri1)
## Compare the second element of 'attr_1' against all other elements
x_pos_val <- seq_len(max(attr_eval(s_cri1)))
eval_sub_criteria(s_cri1,
                  x_pos = x_pos_val,
                  y_pos = rep(2, length(x_pos_val)))

## Example 2 - `s_cri1` AND an exact match on attribute 2
s_cri2 <- sub_criteria(
  s_cri1,
  sub_criteria(attr_2, match_funcs = exact_match),
  operator = "and")
s_cri2

## Example 3 - `s_cri1` OR an exact match on attribute 2
s_cri3 <- sub_criteria(
  s_cri1,
  sub_criteria(attr_2, match_funcs = exact_match),
  operator = "or")
s_cri3

# Evaluate the match criteria
eval_sub_criteria(s_cri2)
```

```

eval_sub_criteria(s_cri3)

# Alternatively, using `attr()`
AND_func <- function(x, y) range_match(x$a1, y$a1) & x$a2 == y$a2
OR_func <- function(x, y) range_match(x$a1, y$a1) | x$a2 == y$a2

## Create a match criteria
s_cri2b <- sub_criteria(attrs(.obj = list(a1 = attr_1, a2 = attr_2)),
                       match_funcs = AND_func)
s_cri3b <- sub_criteria(attrs(.obj = list(a1 = attr_1, a2 = attr_2)),
                       match_funcs = OR_func)

# Evaluate the match criteria
eval_sub_criteria(s_cri2b)
eval_sub_criteria(s_cri3b)

```

windows

Windows and lengths

Description

Covert windows to and from `case_lengths` and `recurrence_lengths`.

Usage

```
epid_windows(date, lengths, episode_unit = "days")
```

```
epid_lengths(date, windows, episode_unit = "days")
```

```
index_window(date, from_last = FALSE)
```

Arguments

<code>date</code>	As used in episodes .
<code>lengths</code>	The duration (lengths) between a date and window.
<code>episode_unit</code>	Time unit of lengths. Options are "seconds", "minutes", "hours", "days", "weeks", "months" or "years". See <code>diyar::episode_unit</code>
<code>windows</code>	The range (windows) relative to a date for a given duration (length).
<code>from_last</code>	As used in episodes .

Details

`epid_windows` - returns the corresponding window for a given a date, and `case_length` or `recurrence_length`.

`epid_lengths` - returns the corresponding `case_length` or `recurrence_length` for a given date and window.

`index_window` - returns the corresponding `case_length` or `recurrence_length` for the date only.

`index_window(date = x)` is a convenience function for `epid_lengths(date = x, window = x)`.

Value

`number_line`.

Examples

```
# Which `window` will a given `length` cover?  
date <- Sys.Date()  
epid_windows(date, 10)  
epid_windows(date, number_line(5, 10))  
epid_windows(date, number_line(-5, 10))  
epid_windows(date, -5)
```

```
# Which `length` is required to cover a given `window`?  
date <- number_line(Sys.Date(), Sys.Date() + 20)  
epid_lengths(date, Sys.Date() + 30)  
epid_lengths(date, number_line(Sys.Date() + 25, Sys.Date() + 30))  
epid_lengths(date, number_line(Sys.Date() - 10, Sys.Date() + 30))  
epid_lengths(date, Sys.Date() - 10)
```

```
# Which `length` is required to cover the `date`?  
index_window(20)  
index_window(number_line(15, 20))
```

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