Package 'reclin2'

January 7, 2022

Type Package

Version 0.1.1

Title Record Linkage Toolkit

```
estimating m- and u-probabilities
      (I. Fellegi & A. Sunter (1969) <doi:10.1080/01621459.1969.10501049>,
      T.N. Herzog, F.J. Scheuren, & W.E. Winkler (2007),
      "Data Quality and Record Linkage Techniques", ISBN:978-0-387-69502-0),
      forcing one-to-one matching. Can also be
      used for pre- and post-processing for machine learning methods for record
      linkage. Focus is on memory, CPU performance and flexibility.
BugReports https://github.com/djvanderlaan/reclin2/issues
URL https://github.com/djvanderlaan/reclin2
Depends data.table, R (>= 3.6.0)
Imports stringdist, stats, utils, lpSolve, Rcpp, parallel
Suggests simplermarkdown
LinkingTo Rcpp
VignetteBuilder simplermarkdown
SystemRequirements C++11
License GPL-3
LazyLoad yes
Encoding UTF-8
RoxygenNote 7.1.1
NeedsCompilation yes
Author Jan van der Laan [aut, cre] (<a href="https://orcid.org/0000-0002-0693-1514">https://orcid.org/0000-0002-0693-1514</a>)
Maintainer Jan van der Laan <r@eoos.dds.nl>
Repository CRAN
Date/Publication 2022-01-07 13:22:42 UTC
```

Description Functions to assist in performing probabilistic record linkage and deduplication: generating pairs, comparing records, em-algorithm for

2 add_from_x

R topics documented:

	add_rrom_x	2
	cluster_call	3
	cluster_collect	4
	cluster_modify_pairs	5
	cluster_pair	6
	cluster_pair_blocking	7
	cluster_pair_minsim	9
	compare_pairs.cluster_pairs	10
	compare_vars.cluster_pairs	12
	deduplicate_equivalence	13
	get_inspect_pairs	14
	greedy	15
	identical	15
	link	17
	linkexample1	18
	match_n_to_m	18
	pair	19
	pair_blocking	20
	pair_minsim	21
	predict.problink_em	22
	problink_em	24
	select_greedy.cluster_pairs	25
	select_threshold.cluster_pairs	28
	summary.problink_em	29
	tabulate_patterns.cluster_pairs	30
	town_names	31
Index		32

 add_from_x

Add a variable from one of the data sets to pairs

Description

Add a variable from one of the data sets to pairs

```
add_from_x(pairs, variable, new_variable = variable, ...)
add_from_y(pairs, variable, new_variable = variable, ...)
```

cluster_call 3

Arguments

pairs data.table with pairs. Should contain the columns .x and .y.

variable name of the variable that should be added

new_variable optional variable name of the new variable in pairs. When omitted variable

is used.

... other parameters are passed on to compare_vars. Especially inplace, x and y

might be of interest.

Value

Returns the pairs with the column added. When inplace = TRUE pairs is returned invisibly and the original pairs is modified.

cluster_call

Call a function on each of the worker nodes and pass it the pairs

Description

Call a function on each of the worker nodes and pass it the pairs

Usage

```
cluster_call(pairs, fun, ...)
```

Arguments

pairs an object or type cluster_pairs as created for example by cluster_pair.

fun a function to call on each of the worker nodes. See details on the arguments of

this function.

... additional arguments are passed on to fun.

Details

The function will have to accept the following arguments as its first three arguments:

pairs the data.table with the pairs of the worker node.

 \mathbf{x} a data. table with the portion of x present on the worker node.

y a data.table with y.

Value

The function will return a list with for each worker the result of the function call. When the functions return NULL the result is returned invisibly. Because the result is returned to main node, make sure you don't accidentally return all pairs. If you don't want to return anything end your function with NULL.

4 cluster_collect

Examples

```
# Generate some pairs
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)</pre>
pairs <- cluster_pair(cl, linkexample1, linkexample2)</pre>
compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))
# Add a new column to pairs
cluster_call(pairs, function(pairs, ...) {
  pairs[, name := firstname & lastname]
  # we don't want to return the pairs; so make sure to return something
  # else
  NULL
})
# Get the number of pairs on each node
lenghts <- cluster_call(pairs, function(pairs, ...) {</pre>
  nrow(pairs)
})
lengths <- unlist(lenghts)</pre>
lenghts
# Cleanup
stopCluster(cl)
```

cluster_collect

Collect pairs from cluster nodes

Description

Collect pairs from cluster nodes

Usage

```
cluster_collect(pairs, select = NULL, clear = FALSE)
```

Arguments

pairs an object or type cluster_pairs as created for example by cluster_pair.

select the name of a logical column that is used to select the pairs that will be collected remove the pairs from the cluster nodes

Value

Returns an object of type pairs which is a data.table. This object can be used as a regular (non-cluster) set of pairs

cluster_modify_pairs 5

Examples

```
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)

pairs <- cluster_pair(cl, linkexample1, linkexample2)
local_pairs <- cluster_collect(pairs, clear = FALSE)

compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))
model <- problink_em(~ lastname + firstname + address + sex, data = pairs)
predict(model, pairs, type = "mpost", add = TRUE, binary = TRUE)
# Select pairs with a mpost > 0.5
select_threshold(pairs, "selected", "mpost", 0.5)
# Collect the selected pairs
local_pairs <- cluster_collect(pairs, "selected")

stopCluster(cl)</pre>
```

Description

Call a function on each of the worker nodes to modify the pairs on the node

Usage

```
cluster_modify_pairs(pairs, fun, ..., new_name = NULL)
```

Arguments

pairs an object or type cluster_pairs as created for example by cluster_pair.

fun a function to call on each of the worker nodes. See details on the arguments of this function.

... additional arguments are passed on to fun.

new_name name of new object to assign the pairs to on the cluster nodes.

Details

The function will have to accept the following arguments as its first three arguments:

```
pairs the data.table with the pairs of the worker node.x a data.table with the portion of x present on the worker node.y a data.table with y.
```

6 cluster_pair

The function should either return a data.table with the new pairs, or NULL. When a data.table is returned this values will replace the pairs when new_name is missing or create new pairs in the environment new_name. When the function returns NULL it is assumed that the function modified the pairs by reference (e.g. using pairs[,new_var := new_val]). Note that this also means that new_name is ignored.

Value

Will return a cluster_pairs object. When new_name is not given it will return the input pairs invisibly. Otherwise it will return a new cluster_pairs object.

Examples

```
# Generate some pairs
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)
pairs <- cluster_pair(cl, linkexample1, linkexample2)
compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))

# Create a new set of pairs containing a random sample of the original
# pairs.
sample <- cluster_call(pairs, new_name = "sample", function(pairs, ...) {
    sel <- sample(nrow(pairs), round(nrow(pairs)*0.1))
    pairs[sel, ]
})

# Cleanup
stopCluster(cl)</pre>
```

cluster_pair

Generate all possible pairs using multiple processes

Description

Generates all combinations of records from x and y.

Usage

```
cluster_pair(cluster, x, y, deduplication = FALSE, name = "default")
```

Arguments

```
cluster a cluster object as created by makeCluster from parallel or makeCluster from snow.

x first data.frame
y second data.frame. Ignored when deduplication = TRUE.
deduplication generate pairs from only x. Ignore y. This is usefull for deduplication of x.
the name of the resulting object to create locally on the different R processes.
```

cluster_pair_blocking 7

Details

Generating (all) pairs of the records of two data sets, is usually the first step when linking the two data sets.

x is split into length{cluster} parts which are distributed over the worker nodes. y is copied to each of the nodes. On the nodes then pair is called. The pairs are stored in the global object reclin_env on the nodes in the variable name. The pairs can then be further processes using functions such as compare_pairs, and tabulate_patterns. The function cluster_collect collects the pairs from each of the nodes.

Value

A object of type cluster_pairs which is a list containing the cluster and the name of the pairs object on the cluster nodes. For the pairs objects created on the nodes see the documentation of pair.

See Also

cluster_pair_blocking and cluster_pair_minsim are other methods to generate pairs.

Examples

```
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)
pairs <- cluster_pair(cl, linkexample1, linkexample2)
stopCluster(cl)</pre>
```

cluster_pair_blocking Generate pairs using simple blocking using multiple processes

Description

Generates all combinations of records from x and y where the blocking variables are equal.

```
cluster_pair_blocking(
  cluster,
  x,
  y,
  on,
  deduplication = FALSE,
  name = "default"
)
```

Arguments

cluster	a cluster object as created by ${\tt makeCluster}$ from parallel or ${\tt makeCluster}$ from snow.
x	first data.frame
У	second data.frame. Ignored when deduplication = TRUE.
on	the variables defining the blocks or strata for which all pairs of x and y will be generated.
deduplication	generate pairs from only x . Ignore y . This is usefull for deduplication of x .
name	the name of the resulting object to create locally on the different R processes.

Details

Generating (all) pairs of the records of two data sets, is usually the first step when linking the two data sets. However, this often results in a too large number of records. Therefore, blocking is usually applied.

x is split into length{cluster} parts which are distributed over the worker nodes. y is copied to each of the nodes. On the nodes then pair_blocking is called. The pairs are stored in the global object reclin_env on the nodes in the variable name. The pairs can then be further processes using functions such as compare_pairs, and tabulate_patterns. The function cluster_collect collects the pairs from each of the nodes.

Value

A object of type cluster_pairs which is a list containing the cluster and the name of the pairs object on the cluster nodes. For the pairs objects created on the nodes see the documentation of pair.

See Also

cluster_pair and cluster_pair_minsim are other methods to generate pairs.

Examples

```
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)
pairs <- cluster_pair_blocking(cl, linkexample1, linkexample2, "postcode")
stopCluster(cl)</pre>
```

cluster_pair_minsim 9

cluster_pair_minsim

Generate pairs with a minimal similarity using multiple processes

Description

Generates all combinations of records from x and y where the blocking variables are equal.

Usage

```
cluster_pair_minsim(
  cluster,
  x,
  y,
  on,
  minsim = 0,
  comparators = list(default_comparator),
  default_comparator = identical(),
  keep_simsum = TRUE,
  deduplication = FALSE,
  name = "default"
)
```

Arguments

name

	cluster	a cluster object as created by ${\tt makeCluster}$ from parallel or ${\tt makeCluster}$ from snow.
	x	first data.frame
	У	second data.frame. Ignored when deduplication = TRUE.
	on	the variables defining the blocks or strata for which all pairs of x and y will be generated.
	minsim	minimal similarity score.
	comparators	named list of functions with which the variables are compared. This function should accept two vectors. Function should either return a vector or a data.table with multiple columns.
default_comparator		
		variables for which no comparison function is defined using comparators is compares with the function default_comparator.
	keep_simsum	add a variable minsim to the result with the similarity score of the pair.
	deduplication	generate pairs from only x. Ignore y. This is usefull for deduplication of x.

the name of the resulting object to create locally on the different R processes.

Details

Generating (all) pairs of the records of two data sets, is usually the first step when linking the two data sets. However, this often results in a too large number of records. pair_minsim will only keep pairs with a similarity score equal or larger than minsim. The similarity score is calculated by summing the results of the comparators for all variables of on.

x is split into length{cluster} parts which are distributed over the worker nodes. y is copied to each of the nodes. On the nodes then cluster_pair_minsim is called. The pairs are stored in the global object reclin_env on the nodes in the variable name. The pairs can then be further processes using functions such as compare_pairs, and tabulate_patterns. The function cluster_collect collects the pairs from each of the nodes.

Value

A object of type cluster_pairs which is a list containing the cluster and the name of the pairs object on the cluster nodes. For the pairs objects created on the nodes see the documentation of pair.

See Also

cluster_pair and cluster_pair_blocking are other methods to generate pairs.

Examples

```
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)
# Either address or postcode has to match to keep a pair
pairs <- cluster_pair_minsim(cl, linkexample1, linkexample2,
    on = c("postcode", "address"), minsim = 1)
stopCluster(cl)</pre>
```

```
compare_pairs.cluster_pairs
```

Compare pairs on a set of variables common in both data sets

Description

Compare pairs on a set of variables common in both data sets

```
## $3 method for class 'cluster_pairs'
compare_pairs(
  pairs,
  on,
  comparators = list(default_comparator),
```

```
default_comparator = identical(),
  new_name = NULL,
)
compare_pairs(
  pairs,
  on,
  comparators = list(default_comparator),
  default_comparator = identical(),
)
## S3 method for class 'pairs'
compare_pairs(
  pairs,
  on,
  comparators = list(default_comparator),
  default_comparator = identical(),
  x = attr(pairs, "x"),
  y = attr(pairs, "y"),
  inplace = FALSE,
)
```

Arguments

pairs data.table with pairs. Should contain the columns .x and .y.

on character vector of variables that should be compared.

comparators named list of functions with which the variables are compared. This func-

tion should accept two vectors. Function should either return a vector or a

data.table with multiple columns.

default_comparator

variables for which no comparison function is defined using comparators is

 $compares \ with \ the \ function \ default_comparator.$

new_name name of new object to assign the pairs to on the cluster nodes.

... Ignored for now

x data.table with one half of the pairs.
y data.table with the other half of the pairs.

inplace logical indicating whether pairs should be modified in place. When pairs is

large this can be more efficient.

Details

It is assumed the variables in on are present in both x and y. Variables with the same names are added to pairs. When the comparator returns a data.table multiple columns are added to pairs. The names of these columns are variable pasted together with the names of the data.table returned by comparator (separated by "_").

Value

Returns the data. table pairs with one or more columns added in case of compare_pairs.pairs.

In case of compare_pairs.cluster_pairs, compare_pair.pairs is called on each cluster node and the resulting pairs are assigned to new_name in the environment reclin_env. When new_name is not given (or equal to NULL) the original pairs on the nodes are overwritten.

```
compare_vars.cluster_pairs
```

Compare pairs on given variables

Description

Compare pairs on given variables

```
## S3 method for class 'cluster_pairs'
compare_vars(
  pairs,
  variable,
  on_x = variable,
  on_y = on_x,
  comparator = identical(),
  new_name = NULL,
)
compare_vars(
  pairs,
  variable,
 on_x = variable,
 on_y = on_x,
  comparator = identical(),
)
## S3 method for class 'pairs'
compare_vars(
  pairs,
  variable,
  on_x = variable,
  on_y = on_x,
  comparator = identical(),
  x = attr(pairs, "x"),
  y = attr(pairs, "y"),
  inplace = FALSE,
```

```
)
```

Arguments

pairs	data.table with pairs. Should contain the columns .x and .y.
variable	character vector with name of resulting column name that is added to pairs.
on_x	character vector with the column names from x on which to compare.
on_y	character vector with the column names from y on which to compare.
comparator	function with which the variables are compared. When on_x and on_y have length 1, this function should accept two vectors. Otherwise it will receive two data.tables. Function should either return a vector or a data.table with multiple columns.
new_name	name of new object to assign the pairs to on the cluster nodes.
	Used to pass additional arguments to methods
x	data.table with one half of the pairs.
у	data.table with the other half of the pairs.
inplace	logical indicating whether pairs should be modified in place. When pairs is large this can be more efficient.

Details

When comparator returns a data.table multiple columns are added to pairs. The names of these columns are variable pasted together with the names of the data.table returned by comparator (separated by "_").

Value

Returns the data. table pairs with one or more columns added.

```
deduplicate_equivalence
```

Deduplication using equivalence groups

Description

Deduplication using equivalence groups

```
deduplicate_equivalence(pairs, variable, selection, x = attr(pairs, "x"))
```

14 get_inspect_pairs

Arguments

pairs	a pairs object, such as generated by pair_blocking
variable	name of the variable to create in x that will contain the group labels.
selection	a logical variable with the same length as pairs has rows, or the name of such a variable in pairs. Pairs are only selected when select is TRUE. When missing it is assumed all pairs are selected.
X	the first data set; when missing attr(pairs, "x") is used.

Value

Returns x with a variable containing the group labels. Records with the same group label (should) correspond to the same entity.

get_inspect_pairs Get a subset of pairs to inspect

Description

Get a subset of pairs to inspect

Usage

```
get_inspect_pairs(
  pairs,
  variable,
  threshold,
  position = NULL,
  n = 11,
  x = attr(pairs, "x"),
  y = attr(pairs, "y")
)
```

Arguments

pairs	data.table with pairs.
variable	name of variable to base the selection on; should be a variable with the similarity score of the pairs.
threshold	the threshold around which to select pairs. Used when position is not given.
position	select pairs around this position (based on order of variable), e.g. position = 1 will select the pairs with the highest similarity score.
n	number of pairs to select. Pairs are selected symmetric around the theshold.
x	data.table with one half of the pairs.
У	data.table with the other half of the pairs.

greedy 15

Value

Returns a list with elements pairs with the selected pairs; x records from x corresponding to the pairs; y records from y corresponding to the pairs; position position of the selected pairs; index index of the pairs in pairs.

greedy

Greedy one-to-one matching of pairs

Description

Greedy one-to-one matching of pairs

Usage

```
greedy(x, y, weight)
```

Arguments

x id's of lhs of pairsy id's of rhs of pairsweight weight of pair

Details

Pairs with the highest weight are selected as long a neither the lhs as the rhs are already selected in a pair with a higher weight.

Value

A logical vector with the same length as x.

identical

Comparison functions

Description

Comparison functions

```
identical()
jaro_winkler(threshold = 0.95)
lcs(threshold = 0.8)
jaccard(threshold = 0.8)
```

16 identical

Arguments

threshold

threshold to use for the Jaro-Winkler string distance when creating a binary result.

Details

A comparison function should accept two arguments: both vectors. When the function is called with both arguments it should compare the elements in the first vector to those in the second. When called in this way, both vectors have the same length. What the function should return depends on the methods used to score the pairs. Usually the comparison functions return a similarity score with a value of 0 indication complete difference and a value > 0 indicating similarity (often a value of 1 will indicate perfect similarity).

Some methods, such as problink_em, can handle similarity scores, but also need binary values (0/FALSE = complete dissimilarity; 1/TRUE = complete similarity). In order to allow for this the comparison function is called with one argument.

When the comparison is called with one argument, it is passed the result of a previous comparison. The function should translate that result to a binary (TRUE/FALSE or 1/0) result. The result should not contain missing values.

The jaro_winkler, lcs and jaccard functions use the corresponding methods from stringdist except that they are transformed from a distance to a similarity score.

Value

The functions return a comparison function (see details).

Examples

link 17

link

Use the selected pairs to generate a linked data set

Description

Use the selected pairs to generate a linked data set

Usage

```
link(
  pairs,
  selection = NULL,
  all = FALSE,
  all_x = all,
  all_y = all,
  x = attr(pairs, "x"),
  y = attr(pairs, "y"),
  suffixes = c(".x", ".y"),
  keep_from_pairs = c(".x", ".y")
)
```

Arguments

pairs	a pairs object, such as generated by pair_blocking
selection	a logical variable with the same length as pairs has rows, or the name of such a variable in pairs. Pairs are only selected when select is TRUE. When missing attr(pairs, "selection") is used when available.
all	return all records from x and y; even those that don't match.
all_x	return all records from x.
all_y	return all records from y.
X	the first data set; when missing attr(pairs, "x") is used.
У	the second data set; when missing attr(pairs, "y") is used.
suffixes	a character vector of length 2 specifying the suffixes to be used for making unique the names of columns in the result.
keep_from_pairs	
	character vector with names of variables in pairs that should be included in the output.

Details

Uses the selected pairs to link the two data sets to each other. Renames variables that are in both data sets.

match_n_to_m

Value

Returns a data.table containing records from x and y and pairs. Columns that occur both in x and y gain a suffix indicating from which data set they are.

linkexample1

Tiny example dataset for probabilistic linkage

Description

Contains fictional records of 7 persons.

Format

Two data frames with resp. 6 and 5 records and 6 columns.

Details

- id the id of the person; this contains no errors and can be used to validate the linkage.
- lastname the last name of the person; contains errors.
- firstname the first name of the persons; contains errors.
- address the address; contains errors.
- sex the sex; contains errors and missing values.
- postcode the postcode; contains no errors.

match_n_to_m

Force n to m matching on a set of pairs

Description

Force n to m matching on a set of pairs

Usage

```
match_n_to_m(x, y, w, n = 1, m = 1)
```

Arguments

X	a vector of identifiers for each x in each pair This vector should have a unique value for each element in x.
У	a vector of identifiers for each y in each pair This vector should have a unique value for each element in y.
W	a vector with weights for each pair. The algorithm will try to maximise the total weight of the selected pairs.
n	an integer. Each element of x can be linked to at most n elements of y.
m	an integer. Each element of y can be linked to at most m elements of y.

pair 19

Details

The algorithm will try to select pairs in such a way each element of x is matched to at most n elements of y and that each element of y is matched at most m elements of x. It tries to select elements in such a way that the total weight w of the selected elements is maximised.

Value

A logical vector with the same length as x indicating the selected records.

Examples

```
d <- data.frame(x=c(1,1,1,2,2,3,3), y=c(1,2,3,4,5,6,7), w=1:7)
# One-to-one matching:
d[match_n_to_m(d$x, d$y, d$w), ]

# N-to-one matching:
d[match_n_to_m(d$x, d$y, d$w, n=999), ]

# One-to-m matching:
d[match_n_to_m(d$x, d$y, d$w, m=999), ]

# N-to-M matching, e.g. select all pairs
d[match_n_to_m(d$x, d$y, d$w, n=999, m=999), ]</pre>
```

pair

Generate all possible pairs

Description

Generates all combinations of records from x and y.

Usage

```
pair(x, y, deduplication = FALSE, add_xy = TRUE)
```

Arguments

```
x first data.frame
y second data.frame. Ignored when deduplication = TRUE.

deduplication generate pairs from only x. Ignore y. This is usefull for deduplication of x.

add_xy add x and y as attributes to the returned pairs. This makes calling some subsequent operations that need x and y (such as compare_pairs easier.
```

20 pair_blocking

Details

Generating (all) pairs of the records of two data sets, is usually the first step when linking the two data sets.

Value

A data.table with two columns, .x and .y, is returned. Columns .x and .y are row numbers from data.frames .x and .y respectively.

See Also

pair_blocking and pair_minsim are other methods to generate pairs.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair(linkexample1, linkexample2)</pre>
```

pair_blocking

Generate pairs using simple blocking

Description

Generates all combinations of records from x and y where the blocking variables are equal.

Usage

```
pair_blocking(x, y, on, deduplication = FALSE, add_xy = TRUE)
```

Arguments

X	first data.frame
у	second data.frame. Ignored when deduplication = TRUE.
on	the variables defining the blocks or strata for which all pairs of x and y will be generated.
deduplication	generate pairs from only x. Ignore y. This is usefull for deduplication of x.
add_xy	add x and y as attributes to the returned pairs. This makes calling some subse-

quent operations that need x and y (such as compare_pairs easier.

Details

Generating (all) pairs of the records of two data sets, is usually the first step when linking the two data sets. However, this often results in a too large number of records. Therefore, blocking is usually applied.

pair_minsim 21

Value

A data.table with two columns, .x and .y, is returned. Columns .x and .y are row numbers from data.frames .x and .y respectively.

See Also

pair and pair_minsim are other methods to generate pairs.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair_blocking(linkexample1, linkexample2, "postcode")</pre>
```

pair_minsim

Generate pairs with a minimal similarity

Description

Generates all combinations of records from x and y where the blocking variables are equal.

Usage

```
pair_minsim(
    x,
    y,
    on,
    minsim = 0,
    comparators = list(default_comparator),
    default_comparator = identical(),
    keep_simsum = TRUE,
    deduplication = FALSE,
    add_xy = TRUE
)
```

Arguments

X	first data.frame
у	second data.frame. Ignored when deduplication = TRUE.
on	the variables defining the blocks or strata for which all pairs of x and y will be generated.
minsim	minimal similarity score.
comparators	named list of functions with which the variables are compared. This function should accept two vectors. Function should either return a vector or a data.table with multiple columns.

22 predict.problink_em

default_comparator

variables for which no comparison function is defined using comparators is

 $compares \ with \ the \ function \ default_comparator.$

keep_simsum add a variable minsim to the result with the similarity score of the pair.

 $\label{eq:deduplication} \mbox{deduplication of x. Ignore y. This is usefull for deduplication of x.}$

add_xy add x and y as attributes to the returned pairs. This makes calling some subse-

quent operations that need x and y (such as compare_pairs easier.

Details

Generating (all) pairs of the records of two data sets, is usually the first step when linking the two data sets. However, this often results in a too large number of records. pair_minsim will only keep pairs with a similarity score equal or larger than minsim. The similarity score is calculated by summing the results of the comparators for all variables of on.

Value

A data.table with two columns, .x and .y, is returned. Columns .x and .y are row numbers from data.frames .x and .y respectively.

See Also

pair and pair_blocking are other methods to generate pairs.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair_minsim(linkexample1, linkexample2,
    on = c("postcode", "address"), minsim = 1)
# Either address or postcode has to match to keep a pair</pre>
```

predict.problink_em

Calculate weights and probabilities for pairs

Description

Calculate weights and probabilities for pairs

```
## S3 method for class 'problink_em'
predict(
  object,
  pairs = newdata,
  newdata = NULL,
  type = c("weights", "mpost", "probs", "all"),
```

predict.problink_em 23

```
binary = FALSE,
add = FALSE,
comparators,
new_name = NULL,
...
)
```

Arguments

object an object of type problink_em as produced by problink_em. a object with pairs for which to calculate weights. pairs newdata an alternative name for the pairs argument. Specify newdata or pairs. a character vector of length one specifying what to calculate. See results for type more information. binary convert comparison vectors to binary vectors using the comparison function in comparators. add add the predictions to the original pairs object. a list of comparison functions (see compare_pairs). When missing attr(pairs, 'comparators') comparators is used. new_name name of new object to assign the pairs to on the cluster nodes (only relevant when pairs is of type cluster_pairs.

Value

When pairs is of type pairs, returns a data.table with either the .x and .y columns from pairs (when add = FALSE) or all columns of pairs. To these columns are added:

- In case of type = "weights" a column weights with the calculated weights.
- In case of type = "mpost" a column mpost with the calculated posterior probabilities (probability that pair is a match given comparison vector.
- In case of type = "prob" the columns mprob and uprob with the m and u-probabilities and mpost and upost with the posterior m- and u-probabilities.
- In case of type = "all" all of the above.

unused.

In case of compare_pairs.cluster_pairs, compare_pair.pairs is called on each cluster node and the resulting pairs are assigned to new_name in the environment reclin_env. When new_name is not given (or equal to NULL) the original pairs on the nodes are overwritten.

24 problink_em

problink_em

Calculate EM-estimates of m- and u-probabilities

Description

Calculate EM-estimates of m- and u-probabilities

Usage

```
problink_em(
  formula,
  data,
  patterns,
  mprobs0 = list(0.95),
  uprobs0 = list(0.02),
  p0 = 0.05,
  tol = 1e-05,
  mprob_max = 0.999,
  uprob_min = 1e-04
)
```

Arguments

formula a formula object with the variables for which to calculate the m- and u-probabilities.

Should be of the form ~ var1 + var2.

data set with pairs on which to estimate the model. Alternatively one can use

the patterns argument.

patterns table of patterns (as output by tabulate_patterns).

mprobs0, uprobs0

initial values of the m- and u-probabilities. These should be lists with numeric

values. The names of the elements in the list should correspond to the names in

by_x in compare_pairs.

p0 the initial estimate of the probability that a pair is a match.

tol when the change in the m and u-probabilities is smaller than tol the algorithm

is stopped.

mprob_max maximum values of the estimated m-probabilities. Values equal to one can lead

to numerical instabilities.

uprob_min maximum values of the estimated m-probabilities. Values equal to zero can lead

to numerical instabilities.

Value

Returns an object of type problink_em. This is a list containing the estimated mprobs, uprobs and overall linkage probability p. It also contains the table of comparison patterns.

References

Fellegi, I. and A. Sunter (1969). "A Theory for Record Linkage", *Journal of the American Statistical Association*. 64 (328): pp. 1183-1210. doi: 10.2307/2286061.

Herzog, T.N., F.J. Scheuren and W.E. Winkler (2007). *Data Quality and Record Linkage Techniques*, Springer.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair_blocking(linkexample1, linkexample2, "postcode")
pairs <- compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))
model <- problink_em(~ lastname + firstname + address + sex, data = pairs)
summary(model)</pre>
```

```
select_greedy.cluster_pairs
```

Select matching pairs enforcing one-to-one linkage

Description

Select matching pairs enforcing one-to-one linkage

```
## S3 method for class 'cluster_pairs'
select_greedy(
 pairs,
 variable,
  score,
  threshold,
 preselect = NULL,
 id_x = NULL,
  id_y = NULL,
)
## S3 method for class 'cluster_pairs'
select_n_to_m(
 pairs,
 variable,
  score,
  threshold,
 preselect = NULL,
  id_x = NULL,
  id_y = NULL,
```

```
)
select_greedy(
  pairs,
  variable,
  score,
  threshold,
  preselect = NULL,
  id_x = NULL
 id_y = NULL,
)
## S3 method for class 'pairs'
select_greedy(
  pairs,
  variable,
  score,
  threshold,
  preselect = NULL,
  id_x = NULL,
  id_y = NULL,
 x = attr(pairs, "x"),
 y = attr(pairs, "y"),
  inplace = FALSE,
)
select_n_to_m(
 pairs,
  variable,
  score,
  threshold,
  preselect = NULL,
  id_x = NULL,
  id_y = NULL,
)
## S3 method for class 'pairs'
select_n_to_m(
 pairs,
 variable,
  score,
  threshold,
  preselect = NULL,
  id_x = NULL,
  id_y = NULL,
```

```
x = attr(pairs, "x"),
y = attr(pairs, "y"),
inplace = FALSE,
...
)
```

Arguments

pairs	a pairs object, such as generated by pair_blocking
variable	the name of the new variable to create in pairs. This will be a logical variable with a value of TRUE for the selected pairs.
score	name of the score/weight variable of the pairs. When not given and attr(pairs, "score") is defined, that is used.
threshold	the threshold to apply. Pairs with a score above the threshold are selected.
preselect	a logical variable with the same length as pairs has rows, or the name of such a variable in pairs. Pairs are only selected when preselect is TRUE. This interacts with threshold (pairs have to be selected with both conditions).
id_x	a integer vector with the same length a the number of rows in pairs, or the name of a column in x. This vector should identify unique objects in x. When not specified it is assumed that each element in x is unique.
id_y	a integer vector with the same length a the number of rows in pairs, or the name of a column in y. This vector should identify unique objects in y. When not specified it is assumed that each element in y is unique.
	Used to pass additional arguments to methods
x	data.table with one half of the pairs.
у	data.table with the other half of the pairs.
inplace	logical indicating whether pairs should be modified in place. When pairs is large this can be more efficient.

Details

Both methods force one-to-one matching. select_greedy uses a greedy algorithm that selects the first pair with the highest weight. select_n_to_m tries to optimise the total weight of all of the selected pairs. In general this will result in a better selection. However, select_n_to_m uses much more memory and is much slower and, therefore, can only be used when the number of possible pairs is not too large.

Value

Returns the pairs with the variable given by variable added. This is a logical variable indicating which pairs are selected a matches.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair_blocking(linkexample1, linkexample2, "postcode")</pre>
```

```
pairs <- compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))</pre>
model <- problink_em(~ lastname + firstname + address + sex, data = pairs)</pre>
pairs <- predict(model, pairs, type = "mpost", add = TRUE, binary = TRUE)</pre>
# Select pairs with a mpost > 0.5 and force one-to-one linkage
pairs <- select_n_to_m(pairs, "ntom", "mpost", 0.5)</pre>
pairs <- select_greedy(pairs, "greedy", "mpost", 0.5)</pre>
table(pairs$ntom, pairs$greedy)
# The same example as above using a cluster;
library(parallel)
cl <- makeCluster(2)</pre>
pairs <- cluster_pair_blocking(cl, linkexample1, linkexample2, "postcode")</pre>
compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))
model <- problink_em(~ lastname + firstname + address + sex, data = pairs)</pre>
predict(model, pairs, type = "mpost", add = TRUE, binary = TRUE)
\# Select pairs with a mpost > 0.5 and force one-to-one linkage
# select_n_to_m and select_greedy only work on pairs that are local;
# therefore we first collect the pairs
select_threshold(pairs, "selected", "mpost", 0.5)
local_pairs <- cluster_collect(pairs, "selected")</pre>
local_pairs <- select_n_to_m(local_pairs, "ntom", "mpost", 0.5)</pre>
local_pairs <- select_greedy(local_pairs, "greedy", "mpost", 0.5)</pre>
table(local_pairs$ntom, local_pairs$greedy)
stopCluster(cl)
```

select_threshold.cluster_pairs

Select matching pairs with a score above a threshold

Description

Select matching pairs with a score above a threshold

Usage

```
## S3 method for class 'cluster_pairs'
select_threshold(pairs, variable, score, threshold, new_name = NULL, ...)
select_threshold(pairs, variable, score, threshold, ...)
## S3 method for class 'pairs'
select_threshold(pairs, variable, score, threshold, inplace = FALSE, ...)
```

Arguments

pairs a pairs object, such as generated by pair_blocking

summary.problink_em 29

variable	the name of the new variable to create in pairs. This will be a logical variable with a value of TRUE for the selected pairs.
score	name of the score/weight variable of the pairs. When not given and attr(pairs, "score") is defined, that is used.
threshold	the threshold to apply. Pairs with a score above the threshold are selected.
new_name	name of new object to assign the pairs to on the cluster nodes.
• • •	ignored
inplace	logical indicating whether pairs should be modified in place. When pairs is large this can be more efficient.

Value

Returns the pairs with the variable given by variable added. This is a logical variable indicating which pairs are selected a matches.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair_blocking(linkexample1, linkexample2, "postcode")</pre>
pairs <- compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))</pre>
model <- problink_em(~ lastname + firstname + address + sex, data = pairs)</pre>
pairs <- predict(model, pairs, type = "mpost", add = TRUE, binary = TRUE)</pre>
# Select pairs with a mpost > 0.5
select_threshold(pairs, "selected", "mpost", 0.5, inplace = TRUE)
# Example using cluster;
# In general the syntax is exactly the same except for the first call to
# to cluster_pair. Note the in general `inplace = TRUE` is implied when
# working with a cluster; therefore the assignment back to pairs can be
# omitted (also not a problem if it is not).
library(parallel)
data("linkexample1", "linkexample2")
cl <- makeCluster(2)</pre>
pairs <- cluster_pair(cl, linkexample1, linkexample2)</pre>
compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))
model <- problink_em(~ lastname + firstname + address + sex, data = pairs)</pre>
predict(model, pairs, type = "mpost", add = TRUE, binary = TRUE)
# Select pairs with a mpost > 0.5
# Unlike the regular pairs: inplace = TRUE is implied here
select_threshold(pairs, "selected", "mpost", 0.5)
stopCluster(cl)
```

summary.problink_em
Summarise the results from problink_em

Description

Summarise the results from problink_em

Usage

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

Arguments

```
object the problink_em object.
... ignored;
```

Value

Returns the original object with a data. frame with the patterns and corresponding m-, u-probabilities and weights added.

```
tabulate_patterns.cluster_pairs

Create a table of comparison patterns
```

Description

Create a table of comparison patterns

Usage

```
## S3 method for class 'cluster_pairs'
tabulate_patterns(pairs, on, comparators, complete = TRUE, ...)
tabulate_patterns(pairs, on, comparators, complete = TRUE, ...)
## S3 method for class 'pairs'
tabulate_patterns(pairs, on, comparators, complete = TRUE, ...)
```

Arguments

pairs	a pairs object, such as generated by pair_blocking
on	variables from pairs defining the comparison patterns. When missing names (comparators) is used.
comparators	a list with comparison functions for each of the columns. When missing or NULL, the function looks for columns in pairs with a comparator attribute.
complete	add patterns that do not occur in the dataset to the result (with $n = 0$).
	passed on to other methods.

town_names 31

Details

Since comparison vectors can contain continuous numbers (usually between 0 and 1), this could result in a very large number of possible comparison vectors. Therefore, the comparison vectors are passed on to the comparators in order to threshold them. This usually results in values 0 or 1. Missing values are usually codes as 0. However, this all depends on the comparison functions used. For more information see the documentation on the comparison functions.

Value

Returns a data.frame with all unique comparison patterns that exist in pairs, with a column n added with the number of times each pattern occurs.

Examples

```
data("linkexample1", "linkexample2")
pairs <- pair_blocking(linkexample1, linkexample2, "postcode")
pairs <- compare_pairs(pairs, c("lastname", "firstname", "address", "sex"))
tabulate_patterns(pairs)</pre>
```

town_names

Spelling variations of a set of town names

Description

Contains spelling variations found in various files of a set of town/village names. Names were selected that contain 'rdam' or 'rdm'. The correct/official names are also given. This data set can be used as an example data set for deduplication

Format

Data frames with 584 records and two columns.

Details

- name the name of the town/village as found in the files
- · official_name the official/correct name

Index

* datasets	match_n_to_m, 18
linkexample1, 18	
town_names, 31	pair, 7, 8, 10, 19, 21, 22
	pair_blocking, 8, 14, 17, 20, 20, 22, 27, 28,
add_from_x, 2	30
add_from_y (add_from_x), 2	pair_minsim, 20, 21, 21
	<pre>predict.problink_em, 22</pre>
cluster_call, 3	problink_em, 16, 23, 24, 29, 30
cluster_collect, 4, 7, 8, 10	
cluster_modify_pairs, 5	select_greedy
cluster_pair, $3-5$, 6 , 8 , 10	<pre>(select_greedy.cluster_pairs),</pre>
cluster_pair_blocking, 7, 7, 10	25
cluster_pair_minsim, 7 , 8 , 9 , 10	select_greedy.cluster_pairs,25
compare_pairs, 7, 8, 10, 19, 20, 22-24	select_n_to_m
compare_pairs	<pre>(select_greedy.cluster_pairs),</pre>
<pre>(compare_pairs.cluster_pairs),</pre>	25
10	select_threshold
compare_pairs.cluster_pairs, 10	<pre>(select_threshold.cluster_pairs),</pre>
compare_vars	28
<pre>(compare_vars.cluster_pairs),</pre>	$select_threshold.cluster_pairs, 28$
12	stringdist, <i>16</i>
<pre>compare_vars.cluster_pairs, 12</pre>	<pre>summary.problink_em, 29</pre>
comparison functions, 31	
	tabulate_patterns, 7 , 8 , 10 , 24
data.table, 3, 11, 13, 14, 20-22	tabulate_patterns
deduplicate_equivalence, 13	<pre>(tabulate_patterns.cluster_pairs), 30</pre>
<pre>get_inspect_pairs, 14</pre>	tabulate_patterns.cluster_pairs,30
greedy, 15	town_names, 31
89,	
identical, 15	
jaccard(identical), 15	
jaro_winkler(identical), 15	
lcs (identical), 15	
link, 17	
linkexample1, 18	
linkexample2(linkexample1), 18	
makeCluster. 6. 8. 9	