

Package ‘couplr’

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Title Optimal Pairing and Matching via Linear Assignment

Version 1.5.3

Description Solves optimal pairing and matching problems using linear assignment algorithms. Provides implementations of the Hungarian method (Kuhn 1955) <doi:10.1002/nav.3800020109>, Jonker-Volgenant shortest path algorithm (Jonker and Volgenant 1987) <doi:10.1007/BF02278710>, Auction algorithm (Bertsekas 1988) <doi:10.1007/BF02186476>, cost-scaling (Goldberg and Kennedy 1995) <doi:10.1007/BF01585996>, scaling algorithms (Gabow and Tarjan 1989) <doi:10.1137/0218069>, push-relabel (Goldberg and Tarjan 1988) <doi:10.1145/48014.61051>, and Sinkhorn entropy-regularized transport (Cuturi 2013) <doi:10.48550/arxiv.1306.0895>. Designed for matching plots, sites, samples, or any pairwise optimization problem. Supports rectangular matrices, forbidden assignments, data frame inputs, batch solving, k-best solutions, and pixel-level image morphing for visualization. Includes automatic preprocessing with variable health checks, multiple scaling methods (standardized, range, robust), greedy matching algorithms, and comprehensive balance diagnostics for assessing match quality using standardized differences and distribution comparisons.

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animated_methods	<i>List methods that currently support animation</i>
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Description

Returns the character vector of method strings for which a trace function has been registered. Use any of these with `lap_animate()`.

Usage

```
animated_methods()
```

Value

Character vector of registered method names.

assignment	<i>Linear assignment solver</i>
------------	---------------------------------

Description

Solve the linear assignment problem (minimum- or maximum-cost matching) using several algorithms. Forbidden edges can be marked as NA or Inf.

Usage

```
assignment(
  cost,
  maximize = FALSE,
  method = c("auto", "jv", "hungarian", "munkres", "auction", "auction_gs",
    "auction_scaled", "sap", "ssp", "csflow", "hk01", "bruteforce", "ssap_bucket",
    "cycle_cancel", "gabow_tarjan", "lapmod", "csa", "ramshaw_tarjan", "push_relabel",
    "orlin", "network_simplex"),
  auction_eps = NULL,
  eps = NULL,
  memory_mode = "auto"
)
```

Arguments

cost Numeric matrix; rows = tasks, columns = agents. NA or Inf entries are treated as forbidden assignments.

maximize Logical; if TRUE, maximizes the total cost instead of minimizing.

method Character string indicating the algorithm to use. Options:

General-purpose solvers:

- "auto" — Automatic selection based on problem characteristics (default)
- "jv" — 'Jonker-Volgenant', fast general-purpose $O(n^3)$ with warm-start
- "hungarian" — Classic 'Hungarian' (shortest augmenting path) $O(n^3)$
- "munkres" — Matrix-form 'Kuhn-Munkres' $O(n^4)$, reference implementation

Auction-based solvers:

- "auction" — 'Bertsekas' auction with adaptive epsilon
- "auction_gs" — 'Gauss-Seidel' variant, good for spatial structure

- "auction_scaled" — 'Epsilon-scaling', fastest for large dense problems

Specialized solvers:

- "sap" / "ssp" — Shortest augmenting path, handles sparsity well
- "lapmod" — Sparse JV variant, faster when $>50\%$
- "hk01" — 'Hopcroft-Karp' for binary (0/1) costs only
- "ssap_bucket" — 'Dial' algorithm for integer costs
- "line_metric" — $O(n \log n)$ for 1D assignment problems
- "bruteforce" — Exact enumeration for tiny problems ($n \leq 8$)

Advanced solvers:

- "csa" — 'Goldberg-Kennedy' cost-scaling, often fastest for medium-large
- "gabow_tarjan" — 'Gabow-Tarjan' bit-scaling with complementary slackness $O(n^3 \log C)$
- "cycle_cancel" — Cycle-canceling with 'Karp' algorithm
- "csflow" — Cost-scaling network flow
- "network_simplex" — 'Network simplex' with spanning tree representation
- "orlin" — 'Orlin-Ahuja' scaling $O(\sqrt{n} * m * \log(nC))$
- "push_relabel" — 'Push-relabel' max-flow based solver
- "ramshaw_tarjan" — 'Ramshaw-Tarjan', optimized for rectangular matrices ($n \neq m$)

auction_eps	Optional numeric epsilon for the 'Auction'/'Auction-GS' methods. If NULL, an internal default (e.g., $1e-9$) is used.
eps	Deprecated. Use auction_eps. If provided and auction_eps is NULL, its value is used for auction_eps.
memory_mode	One of "auto" (default), "dense", or "lazy". cost is already a materialized matrix by the time it reaches assignment(), so "auto" here is diagnostic only: it warns if the matrix is large relative to free system RAM (nothing else can be done post-hoc once the matrix already exists – build it via compute_distances(memory_mode = ...) instead to avoid materializing it in the first place). No lazy solver exists yet, so memory_mode = "lazy" currently always errors.

Details

method = "auto" selects an algorithm based on problem size/shape and data characteristics:

- Very small ($n \leq 8$ and $m \leq 8$): "bruteforce" — exact enumeration
- Binary/constant costs: "hk01" — specialized for 0/1 costs
- Sparse ($>50\%$)
- Very rectangular ($m \geq 3n$): "sap" — handles rectangular well
- Otherwise: "jv" — fastest general-purpose solver at every size

The other solvers are available by naming them explicitly.

Value

An object of class `lap_solve_result`, a list with elements:

- `match` — integer vector of length `min(nrow(cost), ncol(cost))` giving the assigned column for each row (0 if unassigned).
- `total_cost` — numeric scalar, the objective value.
- `status` — character scalar, e.g. "optimal".
- `method_used` — character scalar, the algorithm actually used.

See Also

- `lap_solve()` — Tidy interface returning tibbles
- `lap_solve_kbest()` — Find k-best assignments ('Murty' algorithm)
- `assignment_duals()` — Extract dual variables for sensitivity analysis
- `bottleneck_assignment()` — Minimize maximum edge cost (minimax)
- `sinkhorn()` — Entropy-regularized optimal transport

Examples

```
cost <- matrix(c(4,2,5, 3,3,6, 7,5,4), nrow = 3, byrow = TRUE)
res <- assignment(cost)
res$match; res$total_cost
```

assignment_duals

Solve assignment problem and return dual variables

Description

Solves the linear assignment problem and returns dual potentials (u, v) in addition to the optimal matching. The dual variables provide an optimality certificate and enable sensitivity analysis.

Usage

```
assignment_duals(cost, maximize = FALSE)
```

Arguments

<code>cost</code>	Numeric matrix; rows = tasks, columns = agents. NA or Inf entries are treated as forbidden assignments.
<code>maximize</code>	Logical; if TRUE, maximizes the total cost instead of minimizing.

Details

The dual variables satisfy the complementary slackness conditions:

- For minimization: $u[i] + v[j] \leq \text{cost}[i, j]$ for all (i, j)
- For any assigned pair (i, j) : $u[i] + v[j] = \text{cost}[i, j]$

This implies that $\text{sum}(u) + \text{sum}(v) = \text{total_cost}$ (strong duality).

Applications of dual variables:

- **Optimality verification:** Check that duals satisfy constraints
- **Sensitivity analysis:** Reduced cost $c[i, j] - u[i] - v[j]$ shows how much an edge cost must decrease before it enters the solution
- **Pricing in column generation:** Use duals to price new columns
- **Warm starting:** Reuse duals when costs change slightly

Value

A list with class "assignment_duals_result" containing:

- `match` - integer vector of column assignments (1-based)
- `total_cost` - optimal objective value
- `u` - numeric vector of row dual variables (length `n`)
- `v` - numeric vector of column dual variables (length `m`)
- `status` - character, e.g. "optimal"

See Also

[assignment\(\)](#) for standard assignment without duals

Examples

```
cost <- matrix(c(4, 2, 5, 3, 3, 6, 7, 5, 4), nrow = 3, byrow = TRUE)
result <- assignment_duals(cost)

# Check optimality: u + v should equal cost for assigned pairs
for (i in 1:3) {
  j <- result$match[i]
  cat(sprintf("Row %d -> Col %d: u + v = %.2f, cost = %.2f\n",
              i, j, result$u[i] + result$v[j], cost[i, j]))
}

# Verify strong duality
cat("sum(u) + sum(v) =", sum(result$u) + sum(result$v), "\n")
cat("total_cost =", result$total_cost, "\n")

# Reduced costs (how much must cost decrease to enter solution)
reduced <- outer(result$u, result$v, "+")
reduced_cost <- cost - reduced
print(round(reduced_cost, 2))
```

`as_assignment_matrix` *Convert assignment result to a binary matrix*

Description

Turns a tidy assignment result back into a 0/1 assignment matrix.

Usage

```
as_assignment_matrix(x, n_sources = NULL, n_targets = NULL)
```

Arguments

<code>x</code>	An assignment result object of class <code>lap_solve_result</code>
<code>n_sources</code>	Number of source nodes, optional
<code>n_targets</code>	Number of target nodes, optional

Value

Integer matrix with 0 and 1 entries

`as_matchit` *Convert couplr Result to matchit Object*

Description

Constructs a `matchit`-class S3 object from a `couplr` result, enabling use with any function that accepts **MatchIt** objects (e.g., **cobalt**, **marginaleffects**).

Usage

```
as_matchit(
  result,
  left,
  right,
  formula = NULL,
  left_id = "id",
  right_id = "id",
  ...
)
```

Arguments

<code>result</code>	A couplr result object (<code>matching_result</code> , <code>full_matching_result</code> , <code>cem_result</code> , or <code>subclass_result</code>)
<code>left</code>	Data frame of left (treated) units
<code>right</code>	Data frame of right (control) units
<code>formula</code>	Optional formula used for matching. If not provided, a default formula is constructed from <code>result\$info\$vars</code> .
<code>left_id</code>	Name of ID column in left (default: "id")
<code>right_id</code>	Name of ID column in right (default: "id")
<code>...</code>	Additional arguments (ignored)

Value

An S3 object of class "matchit" with fields:

match.matrix Match matrix (treated x controls)

treat Named treatment vector (1/0)

weights Matching weights

X Covariate matrix

call Original call

info Metadata from couplr

Examples

```
## Not run:
left <- data.frame(id = 1:5, age = c(25, 35, 45, 55, 65))
right <- data.frame(id = 6:15, age = runif(10, 20, 70))
result <- match_couples(left, right, vars = "age")
mi <- as_matchit(result, left, right)
# Now use with cobalt:
cobalt::bal.tab(mi)

## End(Not run)
```

augment

Generic Augment Function

Description

S3 generic for augmenting model results with original data.

Usage

```
augment(x, ...)
```

Arguments

x	An object to augment
...	Additional arguments passed to methods

Value

Augmented data (depends on method)

augment.matching_result

Augment Matching Results with Original Data (broom-style)

Description

S3 method for augmenting matching results following the broom package conventions. This is a thin wrapper around `join_matched()` with sensible defaults for quick exploration.

Usage

```
## S3 method for class 'matching_result'  
augment(x, left, right, ...)
```

Arguments

x	A matching_result object
left	The original left dataset
right	The original right dataset
...	Additional arguments passed to <code>join_matched()</code>

Details

This method follows the `augment()` convention from the broom package, making it easy to integrate couplr into tidymodels workflows. It's equivalent to calling `join_matched()` with default parameters.

If the broom package is not loaded, you can use `couplr::augment()` to access this function.

Value

A tibble with matched pairs and original data (see `join_matched()`)

Examples

```

left <- data.frame(
  id = 1:5,
  treatment = 1,
  age = c(25, 30, 35, 40, 45)
)

right <- data.frame(
  id = 6:10,
  treatment = 0,
  age = c(24, 29, 36, 41, 44)
)

result <- match_couples(left, right, vars = "age")
couplr::augment(result, left, right)

```

autoplot.balance_diagnostics

ggplot2 autoplot for balance diagnostics

Description

Produces ggplot2-based balance assessment plots. Returns a ggplot object.

Usage

```

## S3 method for class 'balance_diagnostics'
autoplot(
  object,
  type = c("love", "histogram", "variance"),
  threshold = 0.1,
  ...
)

```

Arguments

object	A balance_diagnostics object
type	Type of plot: "love" (default), "histogram", or "variance"
threshold	Threshold for standardized differences (default: 0.1)
...	Additional arguments (ignored)

Value

A ggplot object

Examples

```

if (requireNamespace("ggplot2", quietly = TRUE)) {
  set.seed(42)
  left <- data.frame(id = 1:10, age = rnorm(10, 45, 10),
                     income = rnorm(10, 50000, 15000))
  right <- data.frame(id = 11:30, age = rnorm(20, 47, 10),
                     income = rnorm(20, 52000, 15000))
  result <- match_couples(left, right, vars = c("age", "income"))
  bal <- balance_diagnostics(result, left, right, vars = c("age", "income"))
  ggplot2::autoplot(bal)
}

```

autoplot.matching_result

ggplot2 autoplot for matching results

Description

Produces ggplot2-based visualizations of matching distance distributions. Returns a ggplot object that can be further customized.

Usage

```

## S3 method for class 'matching_result'
autoplot(object, type = c("histogram", "density", "ecdf"), ...)

```

Arguments

object	A matching_result object
type	Type of plot: "histogram" (default), "density", or "ecdf"
...	Additional arguments (ignored)

Details

Use plot() for base graphics or autoplot() for ggplot2 output. The ggplot2 package must be installed.

Value

A ggplot object

Examples

```

if (requireNamespace("ggplot2", quietly = TRUE)) {
  left <- data.frame(id = 1:5, x = c(1, 2, 3, 4, 5))
  right <- data.frame(id = 6:10, x = c(1.1, 2.2, 3.1, 4.2, 5.1))
  result <- match_couples(left, right, vars = "x")
  ggplot2::autoplot(result)
  ggplot2::autoplot(result, type = "density")
}

```

autoplot.sensitivity_analysis

ggplot2 autoplot for sensitivity analysis

Description

Plots p-value upper bounds against sensitivity parameter Gamma.

Usage

```

## S3 method for class 'sensitivity_analysis'
autoplot(object, alpha = 0.05, ...)

```

Arguments

object	A sensitivity_analysis object
alpha	Significance level (default: 0.05)
...	Additional arguments (ignored)

Value

A ggplot object

bal.tab.matching_result

Balance Table for Matching Results (cobalt integration)

Description

S3 method enabling `cobalt::bal.tab()` on `couplr` result objects. Requires the **cobalt** package to be installed.

Usage

```

bal.tab.matching_result(x, left, right, ...)

bal.tab.full_matching_result(x, left, right, ...)

bal.tab.cem_result(x, left, right, ...)

bal.tab.subclass_result(x, data = NULL, ...)

```

Arguments

x	A couplr result object
left	Data frame of left (treated) units
right	Data frame of right (control) units
...	Additional arguments passed to <code>cobalt::bal.tab()</code>
data	Data frame used for subclassification (for <code>subclass_result</code> only)

Details

These methods convert couplr results to the format cobalt expects (a `matchit`-class object) and then delegate to cobalt's own `bal.tab.matchit()` method. The **cobalt** package must be installed but is not required for couplr to function.

Value

A cobalt balance table object

balance_diagnostics	<i>Balance Diagnostics for Matched Pairs</i>
---------------------	--

Description

Computes comprehensive balance statistics comparing the distribution of matching variables between left and right units in the matched sample.

Usage

```

balance_diagnostics(result, ...)

## S3 method for class 'matching_result'
balance_diagnostics(
  result,
  left,
  right,
  vars = NULL,
  left_id = "id",

```

```

    right_id = "id",
    ...
)

## S3 method for class 'full_matching_result'
balance_diagnostics(
  result,
  left,
  right,
  vars = NULL,
  left_id = "id",
  right_id = "id",
  ...
)

## S3 method for class 'cem_result'
balance_diagnostics(
  result,
  left,
  right,
  vars = NULL,
  left_id = "id",
  right_id = "id",
  ...
)

## S3 method for class 'subclass_result'
balance_diagnostics(result, data = NULL, vars = NULL, ...)

```

Arguments

result	A matching result object from <code>match_couples()</code> , <code>full_match()</code> , <code>cem_match()</code> , or <code>subclass_match()</code>
...	Additional arguments passed to methods
left	Data frame of left units
right	Data frame of right units
vars	Character vector of variable names to check balance for. Defaults to the variables used in matching (if available in result).
left_id	Character, name of ID column in left data (default: "id")
right_id	Character, name of ID column in right data (default: "id")
data	Data frame used for subclassification (when result is a <code>subclass_result</code>)

Details

This function computes several balance metrics:

Standardized Difference: The difference in means divided by the pooled standard deviation. Values less than 0.1 indicate excellent balance, 0.1-0.25 good balance.

Variance Ratio: The ratio of variances (left/right). Values close to 1 are ideal; the conventional acceptable range is 0.5 to 2. For weighted estimators (full matching, CEM, subclassification) the means and variances are computed using the stratum weights.

KS Statistic: Kolmogorov-Smirnov test statistic comparing distributions. Lower values indicate more similar distributions.

Overall Metrics include mean absolute standardized difference across all variables, proportion of variables with large imbalance (lstd diff > 0.25), and maximum standardized difference.

Value

An S3 object of class `balance_diagnostics` containing:

var_stats Tibble with per-variable balance statistics

overall List with overall balance metrics

pairs Tibble of matched pairs with variables

n_matched Number of matched pairs

n_unmatched_left Number of unmatched left units

n_unmatched_right Number of unmatched right units

method Matching method used

has_blocks Whether blocking was used

block_stats Per-block statistics (if blocking used)

Examples

```
# Create sample data
set.seed(123)
left <- data.frame(
  id = 1:10,
  age = rnorm(10, 45, 10),
  income = rnorm(10, 50000, 15000)
)
right <- data.frame(
  id = 11:30,
  age = rnorm(20, 47, 10),
  income = rnorm(20, 52000, 15000)
)

# Match
result <- match_couples(left, right, vars = c("age", "income"))

# Get balance diagnostics
balance <- balance_diagnostics(result, left, right, vars = c("age", "income"))
print(balance)

# Get balance table
balance_table(balance)
```

balance_table	Create Balance Table
---------------	----------------------

Description

Formats balance diagnostics into a clean table for display or export.

Usage

```
balance_table(balance, digits = 3)
```

Arguments

balance	A balance_diagnostics object from balance_diagnostics()
digits	Number of decimal places for rounding (default: 3)

Value

A tibble with formatted balance statistics

bottleneck_assignment	Solve the Bottleneck Assignment Problem
-----------------------	---

Description

Finds an assignment that minimizes (or maximizes) the maximum edge cost in a perfect matching. Unlike standard LAP which minimizes the sum of costs, BAP minimizes the maximum (bottleneck) cost.

Usage

```
bottleneck_assignment(cost, maximize = FALSE)
```

Arguments

cost	Numeric matrix; rows = tasks, columns = agents. NA or Inf entries are treated as forbidden assignments.
maximize	Logical; if TRUE, maximizes the minimum edge cost instead of minimizing the maximum (maximin objective). Default is FALSE (minimax).

Details

The Bottleneck Assignment Problem (BAP) is a variant of the Linear Assignment Problem where instead of minimizing the sum of assignment costs, we minimize the maximum cost among all assignments (minimax objective).

Algorithm: Uses binary search on the sorted unique costs combined with 'Hopcroft-Karp' bipartite matching to find the minimum threshold that allows a perfect matching.

Complexity: $O(E * \sqrt{V} * \log(\text{unique costs}))$ where E = edges, V = vertices.

Applications:

- Task scheduling with deadline constraints (minimize latest completion)
- Resource allocation (minimize maximum load/distance)
- Network routing (minimize maximum link utilization)
- Fair division problems (minimize maximum disparity)

Value

A list with class "bottleneck_result" containing:

- match - integer vector of length `nrow(cost)` giving the assigned column for each row (1-based indexing)
- bottleneck - numeric scalar, the bottleneck (max/min edge) value
- status - character scalar, e.g. "optimal"

See Also

`assignment()` for standard LAP (sum objective), `lap_solve()` for tidy LAP interface

Examples

```
# Simple example: minimize max cost
cost <- matrix(c(1, 5, 3,
                2, 4, 6,
                7, 1, 2), nrow = 3, byrow = TRUE)
result <- bottleneck_assignment(cost)
result$bottleneck # Maximum edge cost in optimal assignment

# Maximize minimum (fair allocation)
profits <- matrix(c(10, 5, 8,
                  6, 12, 4,
                  3, 7, 11), nrow = 3, byrow = TRUE)
result <- bottleneck_assignment(profits, maximize = TRUE)
result$bottleneck # Minimum profit among all assignments

# With forbidden assignments
cost <- matrix(c(1, NA, 3,
                2, 4, Inf,
                5, 1, 2), nrow = 3, byrow = TRUE)
result <- bottleneck_assignment(cost)
```

cardinality_match	<i>Cardinality Matching</i>
-------------------	-----------------------------

Description

Maximizes the number of matched pairs subject to balance constraints. Uses iterative pruning: starts with a full match, then removes pairs that contribute most to imbalance until all variables satisfy the standardized difference threshold.

Usage

```
cardinality_match(
  left,
  right,
  vars,
  max_std_diff = 0.1,
  distance = "euclidean",
  weights = NULL,
  scale = FALSE,
  auto_scale = FALSE,
  method = "auto",
  max_iter = 100L,
  batch_fraction = 0.1
)
```

Arguments

left	Data frame of "left" units
right	Data frame of "right" units
vars	Character vector of matching variable names
max_std_diff	Maximum allowed absolute standardized difference (default: 0.1, corresponding to "excellent" balance)
distance	Distance metric (default: "euclidean")
weights	Optional named vector of variable weights
scale	Scaling method (default: FALSE)
auto_scale	If TRUE, automatically select scaling (default: FALSE)
method	LAP solver method (default: "auto")
max_iter	Maximum pruning iterations (default: 100)
batch_fraction	Fraction of worst pairs to remove per iteration (default: 0.1). Larger values speed up convergence but may over-prune.

Details

Cardinality matching (Zubizarreta 2014) finds the largest matched sample that satisfies pre-specified balance constraints. This implementation uses an iterative pruning heuristic:

1. Run full optimal matching via `match_couples()`
2. Compute balance diagnostics
3. While any `lstd_diff` exceeds `max_std_diff`:
 - Identify the variable with worst balance
 - Remove the batch of pairs contributing most to that imbalance
 - Recompute balance

Value

A `matching_result` object with additional cardinality info: `result$info$pruning_iterations`, `result$info$pairs_removed`, `result$info$final_balance`.

Examples

```
set.seed(42)
left <- data.frame(id = 1:20, x = rnorm(20, 0, 1), y = rnorm(20, 0, 1))
right <- data.frame(id = 21:50, x = rnorm(30, 0.5, 1), y = rnorm(30, 0.3, 1))
result <- cardinality_match(left, right, vars = c("x", "y"), max_std_diff = 0.2)
print(result)
```

cem_match

Coarsened Exact Matching

Description

Coarsens continuous variables into bins, then performs exact matching on the coarsened values. Units in strata containing both left and right units are kept; others are pruned. Matched units receive weights inversely proportional to stratum sizes to maintain balance.

Usage

```
cem_match(
  left,
  right,
  vars,
  cutpoints = NULL,
  n_bins = "sturges",
  grouping = NULL,
  keep = "all",
  left_id = "id",
  right_id = "id"
)
```

Arguments

<code>left</code>	Data frame of left (treated) units
<code>right</code>	Data frame of right (control) units
<code>vars</code>	Character vector of variable names to coarsen and match on
<code>cutpoints</code>	Named list of break vectors per variable. If NULL, automatic binning is used.
<code>n_bins</code>	Binning method when cutpoints is NULL: "sturges" (default), "fd" (Freedman-Diaconis), "scott", or an integer specifying the number of bins for all variables.
<code>grouping</code>	Character vector of variable names to match exactly (without coarsening). These are typically categorical variables.
<code>keep</code>	Which units to return: "all" (default) returns all units with weight 0 for unmatched, "matched" drops unmatched units.
<code>left_id</code>	Name of ID column in left (default: "id")
<code>right_id</code>	Name of ID column in right (default: "id")

Details

CEM algorithm:

1. Coarsen each numeric variable using `cut` with either user-specified breakpoints or automatic binning (Sturges, FD, or Scott rule)
2. Categorical variables in `grouping` are kept as-is
3. Create strata by concatenating all coarsened values
4. Drop strata with 0 left or 0 right units
5. Compute CEM weights: left units get weight 1, right units get weight $n_{\text{left_in_stratum}} / n_{\text{right_in_stratum}}$ so that the total weight of right units in each stratum equals the number of left units

Value

An S3 object of class `c("cem_result", "couplr_result")` containing:

matched Tibble with columns `id`, `side`, `stratum`, `weight`

strata_summary Tibble with per-stratum counts

info List with `n_strata`, `n_matched_left`, `n_matched_right`, `n_pruned_left`, `n_pruned_right`, `method`, `vars`

Examples

```
set.seed(42)
left <- data.frame(
  id = 1:20, age = rnorm(20, 40, 10),
  income = rnorm(20, 50000, 10000)
)
right <- data.frame(
  id = 21:60, age = rnorm(40, 42, 10),
  income = rnorm(40, 52000, 10000)
```

```

)
result <- cem_match(left, right, vars = c("age", "income"))
print(result)

```

compute_distances

Compute and Cache Distance Matrix for Reuse

Description

Precomputes a distance matrix between left and right datasets, allowing it to be reused across multiple matching operations with different constraints. This is particularly useful when exploring different matching parameters (max_distance, calipers, methods) without recomputing distances.

Usage

```

compute_distances(
  left,
  right,
  vars,
  distance = "euclidean",
  weights = NULL,
  scale = FALSE,
  auto_scale = FALSE,
  left_id = "id",
  right_id = "id",
  block_id = NULL,
  memory_mode = "auto"
)

```

Arguments

left	Left dataset (data frame)
right	Right dataset (data frame)
vars	Character vector of variable names to use for distance computation
distance	Distance metric (default: "euclidean")
weights	Optional numeric vector of variable weights
scale	Scaling method: FALSE, "standardize", "range", or "robust"
auto_scale	Apply automatic preprocessing (default: FALSE)
left_id	Name of ID column in left (default: "id")
right_id	Name of ID column in right (default: "id")
block_id	Optional block ID column name for blocked matching
memory_mode	One of "auto" (default), "dense", or "lazy". "auto" warns (or, for a built-in distance metric, switches) when the dense matrix would consume a large fraction of free system RAM. "lazy" returns a lazy_cost_spec instead of a matrix (built-in distance metrics only); pass it straight to match_couples() the same way as a dense cost_matrix. "dense" skips the RAM check entirely.

Details

This function computes distances once and stores them in a reusable object. The resulting `distance_object` can be passed to `match_couples()` (optionally with `method = "greedy"`) instead of providing datasets and variables.

Benefits:

- **Performance:** Avoid recomputing distances when trying different constraints
- **Exploration:** Quickly test `max_distance`, `calipers`, or `methods`
- **Consistency:** Ensures same distances used across comparisons

With the default `memory_mode = "auto"/"dense"`, the stored `cost_matrix` is a plain dense matrix; `couplr` has no sparse matrix representation, so forbidden pairs (`Inf`) still occupy a cell rather than being dropped from storage. With `memory_mode = "lazy"`, `cost_matrix` is instead a `lazy_cost_spec` that recomputes distances on demand from the underlying feature data.

The `distance_object` stores the original datasets, allowing downstream functions like `join_matched()` to work seamlessly.

Value

An S3 object of class `"distance_object"` containing:

- `cost_matrix`: Numeric matrix of distances
- `left_ids`: Character vector of left IDs
- `right_ids`: Character vector of right IDs
- `block_id`: Block ID column name (if specified)
- `metadata`: List with computation details (`vars`, `distance`, `scale`, etc.)
- `original_left`: Original left dataset (for later joining)
- `original_right`: Original right dataset (for later joining)

Examples

```
# Compute distances once
left <- data.frame(id = 1:5, age = c(25, 30, 35, 40, 45), income = c(45, 52, 48, 61, 55) * 1000)
right <- data.frame(id = 6:10, age = c(24, 29, 36, 41, 44), income = c(46, 51, 47, 60, 54) * 1000)

dist_obj <- compute_distances(
  left, right,
  vars = c("age", "income"),
  scale = "standardize"
)

# Reuse for different matching strategies
result1 <- match_couples(dist_obj, max_distance = 0.5)
result2 <- match_couples(dist_obj, max_distance = 1.0)
result3 <- match_couples(dist_obj, method = "greedy", strategy = "sorted")

# All use the same precomputed distances
```

diagnose_distance_matrix	<i>Diagnose distance matrix and suggest fixes</i>
--------------------------	---

Description

Comprehensive diagnostics for a distance matrix with actionable suggestions.

Usage

```
diagnose_distance_matrix(  
  cost_matrix,  
  left = NULL,  
  right = NULL,  
  vars = NULL,  
  warn = TRUE  
)
```

Arguments

cost_matrix	Numeric matrix of distances
left	Left dataset (for variable checking)
right	Right dataset (for variable checking)
vars	Variables used for matching
warn	If TRUE, issue warnings

Value

List with diagnostic results and suggestions

example_costs	<i>Example cost matrices for assignment problems</i>
---------------	--

Description

Small example datasets for demonstrating couplr functionality across different assignment problem types: square, rectangular, sparse, and binary.

Usage

```
example_costs
```

Format

A list containing four example cost matrices:

simple_3x3 A 3x3 cost matrix with costs ranging from 2-7. Optimal assignment: row 1 -> col 2 (cost 2), row 2 -> col 1 (cost 3), row 3 -> col 3 (cost 4). Total optimal cost: 9.

rectangular_3x5 A 3x5 rectangular cost matrix demonstrating assignment when rows < columns. Each of 3 rows is assigned to one of 5 columns; 2 columns remain unassigned. Costs range 1-6.

sparse_with_na A 3x3 matrix with NA values indicating forbidden assignments. Use this to test algorithms' handling of constraints. Position (1,3), (2,2), and (3,1) are forbidden.

binary_costs A 3x3 matrix with binary (0/1) costs, suitable for testing the HK01 algorithm. Diagonal entries are 0 (preferred), off-diagonal entries are 1 (penalty).

Details

These matrices are designed to test different aspects of LAP solvers:

simple_3x3: Basic functionality test. Any correct solver should find total cost = 9.

rectangular_3x5: Tests handling of non-square problems. The optimal solution assigns all 3 rows with minimum total cost.

sparse_with_na: Tests constraint handling. Algorithms must avoid NA positions while finding an optimal assignment among valid entries.

binary_costs: Tests specialized binary cost algorithms. The optimal assignment uses all diagonal entries (total cost = 0).

See Also

[lap_solve](#), [example_df](#)

Examples

```
# Simple 3x3 assignment
result <- lap_solve(example_costs$simple_3x3)
print(result)
# Optimal: sources 1,2,3 -> targets 2,1,3 with cost 9

# Rectangular problem (3 sources, 5 targets)
result <- lap_solve(example_costs$rectangular_3x5)
print(result)
# All 3 sources assigned; 2 targets unassigned

# Sparse problem with forbidden assignments
result <- lap_solve(example_costs$sparse_with_na)
print(result)
# Avoids NA positions

# Binary costs - test HK01 algorithm
result <- lap_solve(example_costs$binary_costs, method = "hk01")
print(result)
```

```
# Finds diagonal assignment (cost = 0)
```

example_df

Example assignment problem data frame

Description

A tidy data frame representation of assignment problems, suitable for use with grouped workflows and batch solving. Contains two independent 3x3 assignment problems in long format.

Usage

```
example_df
```

Format

A tibble with 18 rows and 4 columns:

sim Simulation/problem identifier. Integer with values 1 or 2, distinguishing two independent assignment problems. Use with `group_by(sim)` for grouped solving.

source Source node index. Integer 1-3 representing the row (source) in each 3x3 cost matrix.

target Target node index. Integer 1-3 representing the column (target) in each 3x3 cost matrix.

cost Cost of assigning source to target. Numeric values ranging from 1-7. Each source-target pair has exactly one cost entry.

Details

This dataset demonstrates couplr's data frame interface for LAP solving. The long format (one row per source-target pair) is converted internally to a cost matrix for solving.

Simulation 1: Costs from `example_costs$simple_3x3`

- Optimal assignment: (1->2, 2->1, 3->3)
- Total cost: 9

Simulation 2: Different cost structure

- Optimal assignment: (1->1, 2->3, 3->3) or equivalent
- Total cost: 4

See Also

[lap_solve](#), [lap_solve_batch](#), [example_costs](#)

Examples

```
library(dplyr)

# Solve both problems with grouped workflow
example_df |>
  group_by(sim) |>
  lap_solve(source, target, cost)

# Batch solving for efficiency
example_df |>
  group_by(sim) |>
  lap_solve_batch(source, target, cost)

# Inspect the data structure
example_df |>
  group_by(sim) |>
  summarise(
    n_pairs = n(),
    min_cost = min(cost),
    max_cost = max(cost)
  )
```

full_match

Full Matching

Description

Assigns every unit (left and right) to a matched group with variable ratios (1:k or k:1). Unlike 1:1 matching, full matching does not discard units, producing matched groups where each group contains at least one left and one right unit.

Usage

```
full_match(
  left,
  right,
  vars,
  distance = "euclidean",
  min_controls = 1,
  max_controls = Inf,
  caliper = NULL,
  caliper_sd = NULL,
  weights = NULL,
  scale = FALSE,
  auto_scale = FALSE,
  sigma = NULL,
  left_id = "id",
```

```

    right_id = "id",
    method = "optimal",
    memory_mode = "auto"
)

```

Arguments

left	Data frame of left (treated) units
right	Data frame of right (control) units
vars	Character vector of variable names to match on
distance	Distance metric: "euclidean" (default), "mahalanobis", "manhattan", or a custom function
min_controls	Minimum number of right units per group (default: 1)
max_controls	Maximum number of right units per group (default: Inf)
caliper	Maximum allowable distance for a match. Units with no eligible partner within the caliper are left unmatched.
caliper_sd	If not NULL, caliper is expressed in standard deviations of the pooled distance distribution rather than absolute units.
weights	Named numeric vector of variable weights
scale	Scaling method: FALSE (default), "robust", "standardize", or "range"
auto_scale	If TRUE, automatically preprocess and scale variables
sigma	Optional covariance matrix for Mahalanobis distance
left_id	Name of ID column in left (default: "id")
right_id	Name of ID column in right (default: "id")
method	Matching algorithm: "optimal" (default) uses min-cost max-flow to find the globally optimal group assignment minimizing total distance; "greedy" uses a fast two-pass heuristic.
memory_mode	One of "auto" (default) or "dense". "auto" warns if the dense cost matrix would consume a large fraction of free system RAM. full_match() uses a different (min-cost-flow) solver backend than match_couples()/assignment(), so memory_mode = "lazy" is not available here yet and errors if requested; "dense" skips the RAM check entirely.

Details

Full matching creates matched groups of variable size. Two algorithms are available:

Optimal (method = "optimal", default): Solves a min-cost max-flow problem that minimizes total distance across all group assignments simultaneously. Each left unit becomes a group center absorbing 1 to max_controls right units, with the globally optimal assignment found via Dijkstra's algorithm with Johnson potentials. When n_left > n_right, roles are transposed automatically.

Greedy (method = "greedy"): A fast two-pass heuristic:

1. Each left unit picks its nearest eligible right unit
2. Remaining right units are assigned to their nearest already-matched left unit, respecting max_controls

This is faster but does not guarantee globally optimal results.

Weights are computed so that within each group, the total weight of right units equals the total weight of left units (which is 1). For a group with 1 left and k right units, each right unit receives weight $1/k$.

Value

An S3 object of class `c("full_matching_result", "couplr_result")` containing:

groups Tibble with columns `group_id`, `id`, `side` ("left"/"right"), and `weight`

info List with `n_groups`, `n_left`, `n_right`, `n_unmatched_left`, `n_unmatched_right`, `method`, `vars`

unmatched List of unmatched left and right IDs (if caliper excludes units)

Examples

```
set.seed(42)
left <- data.frame(id = 1:5, age = c(25, 35, 45, 55, 65))
right <- data.frame(id = 6:20, age = runif(15, 20, 70))
result <- full_match(left, right, vars = "age")
print(result)
```

get_method_used

Extract method used from assignment result

Description

Extract method used from assignment result

Usage

```
get_method_used(x)
```

Arguments

`x` An assignment result object

Value

Character string indicating method used

get_total_cost	<i>Extract total cost from assignment result</i>
----------------	--

Description

Extract total cost from assignment result

Usage

```
get_total_cost(x)
```

Arguments

`x` An assignment result object

Value

Numeric total cost

hospital_staff	<i>Hospital staff scheduling example dataset</i>
----------------	--

Description

A comprehensive example dataset for demonstrating couplr functionality across vignettes. Contains hospital staff scheduling data with nurses, shifts, costs, and preference scores suitable for assignment problems, as well as nurse characteristics for matching workflows.

Usage

```
hospital_staff
```

Format

A list containing eight related datasets:

basic_costs A 10x10 numeric cost matrix for assigning 10 nurses to 10 shifts. Values range from approximately 1-15, where lower values indicate better fit (less overtime, matches skills, respects preferences). Use with `lap_solve()` for basic assignment.

preferences A 10x10 numeric preference matrix on a 0-10 scale, where higher values indicate stronger nurse preference for a shift. Use with `lap_solve(..., maximize = TRUE)` to optimize preferences rather than minimize costs.

schedule_df A tibble with 100 rows (10 nurses x 10 shifts) in long format for data frame workflows:

nurse_id Integer 1-10. Unique identifier for each nurse.

shift_id Integer 1-10. Unique identifier for each shift.
cost Numeric. Assignment cost (same values as basic_costs).
preference Numeric 0-10. Nurse preference score.
skill_match Integer 0/1. Binary indicator: 1 if nurse skills match shift requirements, 0 otherwise.

nurses A tibble with 10 rows describing nurse characteristics:

nurse_id Integer 1-10. Links to schedule_df and basic_costs rows.
experience_years Numeric 1-20. Years of nursing experience.
department Character. Primary department: "ICU", "ER", "General", or "Pediatrics".
shift_preference Character. Preferred shift type: "day", "evening", or "night".
certification_level Integer 1-3. Certification level where 3 is highest (e.g., 1=RN, 2=BSN, 3=MSN).

shifts A tibble with 10 rows describing shift requirements:

shift_id Integer 1-10. Links to schedule_df and basic_costs cols.
department Character. Department needing coverage.
shift_type Character. Shift type: "day", "evening", or "night".
min_experience Numeric. Minimum years of experience required.
min_certification Integer 1-3. Minimum certification level.

weekly_df A tibble for batch solving with 500 rows (5 days x 10 nurses x 10 shifts):

day Character. Day of week: "Mon", "Tue", "Wed", "Thu", "Fri".
nurse_id Integer 1-10. Nurse identifier.
shift_id Integer 1-10. Shift identifier.
cost Numeric. Daily assignment cost (varies by day).
preference Numeric 0-10. Daily preference score.
 Use with group_by(day) for solving each day's schedule.

nurses_extended A tibble with 200 nurses for matching examples, representing a treatment group (e.g., full-time nurses):

nurse_id Integer 1-200. Unique identifier.
age Numeric 22-65. Nurse age in years.
experience_years Numeric 0-40. Years of nursing experience.
hourly_rate Numeric 25-75. Hourly wage in dollars.
department Character. Primary department assignment.
certification_level Integer 1-3. Certification level.
is_fulltime Logical. TRUE for full-time status.

controls_extended A tibble with 300 potential control nurses (e.g., part-time or registry nurses) for matching. Same structure as nurses_extended. Designed to have systematic differences from nurses_extended (older, less experience on average) to demonstrate matching's ability to create comparable groups.

Details

This dataset is used throughout the couplr documentation to provide a consistent, realistic example that evolves in complexity. It supports three use cases: (1) basic LAP solving with cost matrices, (2) batch solving across multiple days, and (3) matching workflows comparing nurse groups.

The dataset is designed to demonstrate progressively complex scenarios:

Basic LAP (`vignette("getting-started")`):

- `basic_costs`: Simple 10x10 assignment
- `preferences`: Maximization problem
- `schedule_df`: Data frame input, grouped workflows
- `weekly_df`: Batch solving across days

Algorithm comparison (`vignette("algorithms")`):

- Use `basic_costs` to compare algorithm behavior
- Modify with NA values for sparse scenarios

Matching workflows (`vignette("matching-workflows")`):

- `nurses_extended`: Treatment group (full-time nurses)
- `controls_extended`: Control pool (part-time/registry nurses)
- Match on age, experience, department for causal analysis

See Also

[lap_solve](#) for basic assignment solving, [lap_solve_batch](#) for batch solving, [match_couples](#) for matching workflows, `vignette("getting-started")` for introductory tutorial

Examples

```
# Basic assignment: assign nurses to shifts minimizing cost
lap_solve(hospital_staff$basic_costs)

# Maximize preferences instead
lap_solve(hospital_staff$preferences, maximize = TRUE)

# Data frame workflow
library(dplyr)
hospital_staff$schedule_df |>
  lap_solve(nurse_id, shift_id, cost)

# Batch solve weekly schedule
hospital_staff$weekly_df |>
  group_by(day) |>
  lap_solve(nurse_id, shift_id, cost)

# Matching workflow: match full-time to part-time nurses
match_couples(
  left = hospital_staff$nurses_extended,
```

```

    right = hospital_staff$controls_extended,
    vars = c("age", "experience_years", "certification_level"),
    auto_scale = TRUE
  )

```

is_distance_object	<i>Check if Object is a Distance Object</i>
--------------------	---

Description

Check if Object is a Distance Object

Usage

```
is_distance_object(x)
```

Arguments

x	Object to check
---	-----------------

Value

Logical: TRUE if x is a distance_object

Examples

```

left <- data.frame(id = 1:3, x = c(1, 2, 3))
right <- data.frame(id = 4:6, x = c(1.1, 2.1, 3.1))
dist_obj <- compute_distances(left, right, vars = "x")
is_distance_object(dist_obj) # TRUE
is_distance_object(list()) # FALSE

```

is_lap_solve_batch_result	<i>Check if object is a batch assignment result</i>
---------------------------	---

Description

Check if object is a batch assignment result

Usage

```
is_lap_solve_batch_result(x)
```

Arguments

x Object to test

Value

Logical indicating if x is a batch assignment result

is_lap_solve_kbest_result
<i>Check if object is a k-best assignment result</i>

Description

Check if object is a k-best assignment result

Usage

is_lap_solve_kbest_result(x)

Arguments

x Object to test

Value

Logical indicating if x is a k-best assignment result

is_lap_solve_result	<i>Check if object is an assignment result</i>
---------------------	--

Description

Check if object is an assignment result

Usage

is_lap_solve_result(x)

Arguments

x Object to test

Value

Logical indicating if x is an assignment result

join_matched	<i>Join Matched Pairs with Original Data</i>
--------------	--

Description

Creates an analysis-ready dataset by joining matched pairs with variables from the original left and right datasets. This eliminates the need for manual joins and provides a convenient format for downstream analysis.

Usage

```
join_matched(result, ...)

## S3 method for class 'matching_result'
join_matched(
  result,
  left,
  right,
  left_vars = NULL,
  right_vars = NULL,
  left_id = "id",
  right_id = "id",
  suffix = c("_left", "_right"),
  include_distance = TRUE,
  include_pair_id = TRUE,
  include_block_id = TRUE,
  ...
)

## S3 method for class 'full_matching_result'
join_matched(result, left, right, left_id = "id", right_id = "id", ...)

## S3 method for class 'cem_result'
join_matched(result, left, right, left_id = "id", right_id = "id", ...)

## S3 method for class 'subclass_result'
join_matched(result, data = NULL, ...)
```

Arguments

result	A result object from <code>match_couples()</code> , <code>full_match()</code> , or <code>cem_match()</code>
...	Additional arguments passed to methods
left	The original left dataset
right	The original right dataset
left_vars	Character vector of variable names to include from left. If NULL (default), includes all variables except the ID column.

<code>right_vars</code>	Character vector of variable names to include from right. If NULL (default), includes all variables except the ID column.
<code>left_id</code>	Name of the ID column in left dataset (default: "id")
<code>right_id</code>	Name of the ID column in right dataset (default: "id")
<code>suffix</code>	Character vector of length 2 specifying suffixes for left and right variables (default: <code>c("_left", "_right")</code>)
<code>include_distance</code>	Include the matching distance in output (default: TRUE)
<code>include_pair_id</code>	Include <code>pair_id</code> column (default: TRUE)
<code>include_block_id</code>	Include <code>block_id</code> if blocking was used (default: TRUE)
<code>data</code>	Data frame used for subclassification

Details

This function simplifies the common workflow of joining matched pairs with original data. Instead of manually merging `result$pairs` with left and right datasets, `join_matched()` handles the joins automatically and applies consistent naming conventions.

When variables appear in both left and right datasets, suffixes are appended to distinguish them (e.g., "age_left" and "age_right"). This makes it easy to compute differences or use both values in models.

Value

A tibble with one row per matched pair, containing:

- `pair_id`: Sequential pair identifier (if `include_pair_id = TRUE`)
- `left_id`: ID from left dataset
- `right_id`: ID from right dataset
- `distance`: Matching distance (if `include_distance = TRUE`)
- `block_id`: Block identifier (if blocking used and `include_block_id = TRUE`)
- Variables from left dataset (with left suffix)
- Variables from right dataset (with right suffix)

Examples

```
# Basic usage
left <- data.frame(
  id = 1:5,
  treatment = 1,
  age = c(25, 30, 35, 40, 45),
  income = c(45000, 52000, 48000, 61000, 55000)
)

right <- data.frame(
```

```
    id = 6:10,
    treatment = 0,
    age = c(24, 29, 36, 41, 44),
    income = c(46000, 51500, 47500, 60000, 54000)
  )

result <- match_couples(left, right, vars = c("age", "income"))
matched_data <- join_matched(result, left, right)
head(matched_data)

# Specify which variables to include
matched_data <- join_matched(
  result, left, right,
  left_vars = c("treatment", "age", "income"),
  right_vars = c("age", "income"),
  suffix = c("_treated", "_control")
)

# Without distance or pair_id
matched_data <- join_matched(
  result, left, right,
  include_distance = FALSE,
  include_pair_id = FALSE
)
```

lap_animate*Animate an assignment algorithm step-by-step*

Description

Produce an interactive bipartite-graph animation showing how a chosen linear-assignment algorithm transforms the matching over time. The result is an `htmlwidget` suitable for use in R Markdown, Quarto, pkgdown vignettes, Shiny apps, or standalone HTML output.

Usage

```
lap_animate(
  cost,
  method = "hungarian",
  maximize = FALSE,
  width = NULL,
  height = NULL,
  elementId = NULL,
  ...
)
```

Arguments

cost	Numeric cost matrix. Rows = workers/sources, columns = jobs/targets. NA or Inf entries mark forbidden assignments.
method	Character; the algorithm to animate. Must match one of the methods registered for animation (see <code>animated_methods()</code> for the current list). Method names are the same strings used by <code>assignment()</code> , e.g. "hungarian", "jv", "auction", "gabow_tarjan".
maximize	Logical; if TRUE, animate the maximization variant.
width, height	Optional explicit widget dimensions (pixels or CSS units).
elementId	Optional DOM id for the widget container.
...	Algorithm-specific extra arguments forwarded to the trace function (e.g. <code>auction_eps</code>).

Details

This is a *teaching* interface. It runs a slower R reference implementation that emits a state trace at every step, then plays it back in the browser. For production solving, use `assignment()` or `lap_solve()` which call the fast C++ backends.

Value

An `htmlwidget` object.

Animated methods

Animation support is added incrementally. Call `animated_methods()` to see which method strings currently have a registered trace.

See Also

`assignment()` for production solving, `lap_solve()` for the tidy interface.

Examples

```
## Not run:
cost <- matrix(c(4, 2, 5,
                 3, 3, 6,
                 7, 5, 4), nrow = 3, byrow = TRUE)
lap_animate(cost, method = "hungarian")

## End(Not run)
```

lap_solve	<i>Solve linear assignment problems</i>
-----------	---

Description

Provides a tidy interface for solving the linear assignment problem using 'Hungarian' or 'Jonker-Volgenant' algorithms. Supports rectangular matrices, NA/Inf masking, and data frame inputs.

Usage

```
lap_solve(
  x,
  source = NULL,
  target = NULL,
  cost = NULL,
  maximize = FALSE,
  method = "auto",
  forbidden = NA
)
```

Arguments

x	Cost matrix, data frame, or tibble. If a data frame/tibble, must include columns specified by source, target, and cost.
source	Column name for source/row indices (if x is a data frame)
target	Column name for target/column indices (if x is a data frame)
cost	Column name for costs (if x is a data frame)
maximize	Logical; if TRUE, maximizes total cost instead of minimizing (default: FALSE)
method	Algorithm to use. One of: "auto" (default): automatically selects best algorithm "jv": 'Jonker-Volgenant' algorithm (general purpose, fast) "hungarian": Classic 'Hungarian' algorithm "auction": 'Bertsekas' auction algorithm (good for large dense problems) "sap": Sparse assignment (good for sparse/rectangular problems) "hk01": 'Hopcroft-Karp' for binary/uniform costs
forbidden	Value to mark forbidden assignments (default: NA). Can also use Inf.

Value

A tibble with columns:

- source: row/source indices
- target: column/target indices
- cost: cost of each assignment
- total_cost: total cost (attribute)

Examples

```

# Matrix input
cost <- matrix(c(4, 2, 5, 3, 3, 6, 7, 5, 4), nrow = 3)
lap_solve(cost)

# Data frame input
library(dplyr)
df <- tibble(
  source = rep(1:3, each = 3),
  target = rep(1:3, times = 3),
  cost = c(4, 2, 5, 3, 3, 6, 7, 5, 4)
)
lap_solve(df, source, target, cost)

# With NA masking (forbidden assignments)
cost[1, 3] <- NA
lap_solve(cost)

# Grouped data frames
df <- tibble(
  sim = rep(1:2, each = 9),
  source = rep(1:3, times = 6),
  target = rep(1:3, each = 3, times = 2),
  cost = runif(18, 1, 10)
)
df |> group_by(sim) |> lap_solve(source, target, cost)

```

lap_solve_batch

*Solve multiple assignment problems efficiently***Description**

Solve many independent assignment problems at once. Supports lists of matrices, 3D arrays, or grouped data frames. Optional parallel execution via `n_threads`.

Usage

```

lap_solve_batch(
  x,
  source = NULL,
  target = NULL,
  cost = NULL,
  maximize = FALSE,
  method = "auto",
  n_threads = 1,
  forbidden = NA
)

```

Arguments

x	One of: List of cost matrices, 3D array, or grouped data frame
source	Column name for source indices (if x is a grouped data frame)
target	Column name for target indices (if x is a grouped data frame)
cost	Column name for costs (if x is a grouped data frame)
maximize	Logical; if TRUE, maximizes total cost (default: FALSE)
method	Algorithm to use (default: "auto"). See lap_solve for options.
n_threads	Number of threads for parallel execution (default: 1). Set to NULL to use all available cores.
forbidden	Value to mark forbidden assignments (default: NA)

Value

A tibble with columns:

- problem_id: identifier for each problem
- source: source indices for assignments
- target: target indices for assignments
- cost: cost of each assignment
- total_cost: total cost for each problem
- method_used: algorithm used for each problem

Examples

```
# List of matrices
costs <- list(
  matrix(c(1, 2, 3, 4), 2, 2),
  matrix(c(5, 6, 7, 8), 2, 2)
)
lap_solve_batch(costs)

# 3D array
arr <- array(runif(2 * 2 * 10), dim = c(2, 2, 10))
lap_solve_batch(arr)

# Grouped data frame
library(dplyr)
df <- tibble(
  sim = rep(1:5, each = 9),
  source = rep(1:3, times = 15),
  target = rep(1:3, each = 3, times = 5),
  cost = runif(45, 1, 10)
)
df |> group_by(sim) |> lap_solve_batch(source, target, cost)

# Parallel execution (requires n_threads > 1)
lap_solve_batch(costs, n_threads = 2)
```

lap_solve_kbest	<i>Find k-best optimal assignments</i>
-----------------	--

Description

Returns the top k optimal (or near-optimal) assignments using 'Murty' algorithm. Useful for exploring alternative optimal solutions or finding robust assignments.

Usage

```
lap_solve_kbest(
  x,
  k = 3,
  source = NULL,
  target = NULL,
  cost = NULL,
  maximize = FALSE,
  method = "murty",
  single_method = "jv",
  forbidden = NA
)
```

Arguments

x	Cost matrix, data frame, or tibble. If a data frame/tibble, must include columns specified by source, target, and cost.
k	Number of best solutions to return (default: 3)
source	Column name for source/row indices (if x is a data frame)
target	Column name for target/column indices (if x is a data frame)
cost	Column name for costs (if x is a data frame)
maximize	Logical; if TRUE, finds k-best maximizing assignments (default: FALSE)
method	Algorithm for each sub-problem (default: "murty"). Future versions may support additional methods.
single_method	Algorithm used for solving each node in the search tree (default: "jv")
forbidden	Value to mark forbidden assignments (default: NA)

Value

A tibble with columns:

- rank: ranking of solutions (1 = best, 2 = second best, etc.)
- solution_id: unique identifier for each solution
- source: source indices
- target: target indices
- cost: cost of each edge in the assignment
- total_cost: total cost of the complete solution

Examples

```
# Matrix input - find 5 best solutions
cost <- matrix(c(4, 2, 5, 3, 3, 6, 7, 5, 4), nrow = 3)
lap_solve_kbest(cost, k = 5)

# Data frame input
library(dplyr)
df <- tibble(
  source = rep(1:3, each = 3),
  target = rep(1:3, times = 3),
  cost = c(4, 2, 5, 3, 3, 6, 7, 5, 4)
)
lap_solve_kbest(df, k = 3, source, target, cost)

# With maximization
lap_solve_kbest(cost, k = 3, maximize = TRUE)
```

lap_solve_line_metric *Solve 1-D Line Assignment Problem*

Description

Solves the linear assignment problem when both sources and targets are ordered points on a line. Uses efficient $O(n*m)$ dynamic programming for rectangular problems and $O(n)$ sorting for square problems.

Usage

```
lap_solve_line_metric(x, y, cost = "L1", maximize = FALSE)
```

Arguments

x	Numeric vector of source positions (will be sorted internally)
y	Numeric vector of target positions (will be sorted internally)
cost	Cost function for distance. Either: • "L1" (default): absolute distance ('Manhattan' distance) • "L2": squared distance (squared 'Euclidean' distance) Can also use aliases: "abs", "manhattan" for L1; "sq", "squared", "quadratic" for L2
maximize	Logical; if TRUE, maximizes total cost instead of minimizing (default: FALSE)

Details

This is a specialized solver that exploits the structure of 1-dimensional assignment problems where costs depend only on the distance between points on a line. It is much faster than general LAP solvers for this special case.

The algorithm works as follows:

Square case ($n == m$): Both vectors are sorted and matched in order: $x[1] \rightarrow y[1]$, $x[2] \rightarrow y[2]$, etc. This is optimal for any metric cost function on a line.

Rectangular case ($n < m$): Uses dynamic programming to find the optimal assignment that matches all n sources to a subset of the m targets, minimizing total distance. The DP recurrence is:

$$dp[i][j] = \min(dp[i][j-1], dp[i-1][j-1] + \text{cost}(x[i], y[j]))$$

This finds the minimum cost to match the first i sources to the first j targets.

Complexity:

- Time: $O(n*m)$ for rectangular, $O(n \log n)$ for square
- Space: $O(n*m)$ for DP table

Value

A list with components:

- `match`: Integer vector of length n with 1-based column indices
- `total_cost`: Total cost of the assignment

Examples

```
# Square case: equal number of sources and targets
x <- c(1.5, 3.2, 5.1)
y <- c(2.0, 3.0, 5.5)
result <- lap_solve_line_metric(x, y, cost = "L1")
print(result)

# Rectangular case: more targets than sources
x <- c(1.0, 3.0, 5.0)
y <- c(0.5, 2.0, 3.5, 4.5, 6.0)
result <- lap_solve_line_metric(x, y, cost = "L2")
print(result)

# With unsorted inputs (will be sorted internally)
x <- c(5.0, 1.0, 3.0)
y <- c(4.5, 0.5, 6.0, 2.0, 3.5)
result <- lap_solve_line_metric(x, y, cost = "L1")
print(result)
```

matchmaker

Create blocks for stratified matching

Description

Constructs blocks (strata) for matching, using either grouping variables or clustering algorithms. Returns the input data frames with block IDs assigned, along with block summary statistics.

Usage

```
matchmaker(
  left,
  right,
  block_type = c("none", "group", "cluster"),
  block_by = NULL,
  block_vars = NULL,
  block_method = "kmeans",
  n_blocks = NULL,
  min_left = 1,
  min_right = 1,
  drop_imbalanced = FALSE,
  imbalance_threshold = Inf,
  return_dropped = TRUE,
  ...
)
```

Arguments

<code>left</code>	Data frame of "left" units (e.g., treated, cases)
<code>right</code>	Data frame of "right" units (e.g., control, controls)
<code>block_type</code>	Type of blocking to use: • "none": No blocking (default) • "group": Block by existing categorical variable(s) • "cluster": Block using clustering algorithm
<code>block_by</code>	Variable name(s) for grouping (if <code>block_type = "group"</code>)
<code>block_vars</code>	Variable names for clustering (if <code>block_type = "cluster"</code>)
<code>block_method</code>	Clustering method (if <code>block_type = "cluster"</code>): • "kmeans": K-means clustering • "hclust": Hierarchical clustering
<code>n_blocks</code>	Target number of blocks (for clustering)
<code>min_left</code>	Minimum number of left units per block
<code>min_right</code>	Minimum number of right units per block
<code>drop_imbalanced</code>	Drop blocks with extreme imbalance
<code>imbalance_threshold</code>	Maximum allowed $ n_left - n_right / \max(n_left, n_right)$
<code>return_dropped</code>	Include dropped blocks in output
<code>...</code>	Additional arguments passed to clustering function

Details

This function does NOT perform matching - it only creates the block structure. Use [match_couples\(\)](#) to perform matching within blocks.

Value

A list with class "matchmaker_result" containing:

- left: Left data frame with block_id column added
- right: Right data frame with block_id column added
- block_summary: Summary statistics for each block
- dropped: Information about dropped blocks (if any)
- info: Metadata about blocking process

Examples

```
# Group blocking
left <- data.frame(id = 1:10, region = rep(c("A", "B"), each = 5), x = rnorm(10))
right <- data.frame(id = 11:20, region = rep(c("A", "B"), each = 5), x = rnorm(10))
blocks <- matchmaker(left, right, block_type = "group", block_by = "region")
print(blocks$block_summary)

# Clustering
blocks <- matchmaker(left, right, block_type = "cluster",
                     block_vars = "x", n_blocks = 3)
```

match_couples

Match two datasets into couples

Description

Performs one-to-one matching between two datasets. Supports blocking, distance constraints, and various distance metrics.

Usage

```
match_couples(
  left,
  right = NULL,
  vars = NULL,
  distance = "euclidean",
  weights = NULL,
  scale = FALSE,
  auto_scale = FALSE,
  max_distance = Inf,
  calipers = NULL,
  block_id = NULL,
  ignore_blocks = FALSE,
  require_full_matching = FALSE,
  method = "auto",
```

```

strategy = c("row_best", "sorted", "pq"),
return_unmatched = TRUE,
return_diagnostics = FALSE,
parallel = FALSE,
replace = FALSE,
ratio = 1L,
check_costs = TRUE,
sigma = NULL,
memory_mode = "auto"
)

```

Arguments

left	Data frame of "left" units (e.g., treated, cases)
right	Data frame of "right" units (e.g., control, controls)
vars	Variable names to use for distance computation
distance	Distance metric: "euclidean", "manhattan", "mahalanobis", or a custom function
weights	Optional named vector of variable weights
scale	Scaling method: FALSE (none), "standardize", "range", or "robust"
auto_scale	If TRUE, automatically check variable health and select scaling method (default: FALSE)
max_distance	Maximum allowed distance (pairs exceeding this are forbidden)
calipers	Named list of per-variable maximum absolute differences
block_id	Column name containing block IDs (for stratified matching)
ignore_blocks	If TRUE, ignore block_id even if present
require_full_matching	If TRUE, error if any units remain unmatched
method	Matching method. A LAP solver for optimal matching ("auto", "hungarian", "jv", "gabow_tarjan", ...), or "greedy" for fast approximate matching (see strategy).
strategy	Greedy strategy, used only when method = "greedy". All three strategies solve the same full cost matrix already built by match_couples(); none of them reduce the $O(n*m)$ memory that matrix takes. • "row_best": for each row, take its best available column (default). The only strategy that needs no extra storage beyond the cost matrix. • "sorted": collect every valid pair, sort by distance, greedily assign • "pq": collect every valid pair into a heap and pop the smallest first. Avoids the upfront sort but holds the same number of candidate pairs as "sorted", so it is not more memory-efficient
return_unmatched	Include unmatched units in output
return_diagnostics	Include detailed diagnostics in output
parallel	Enable parallel processing for blocked matching. Requires 'future' and 'future.apply' packages. Can be:

	• FALSE: Sequential processing (default) • TRUE: Auto-configure parallel backend • Character: Specify future plan (e.g., "multisession", "multicore")
replace	If TRUE, allow matching with replacement (same right unit can be matched to multiple left units). Default: FALSE.
ratio	Integer, number of right units to match per left unit. Default: 1 (one-to-one matching). For k:1 matching, set ratio = k.
check_costs	If TRUE, check distance distribution for potential problems and provide helpful warnings before matching (default: TRUE)
sigma	Optional covariance matrix for Mahalanobis distance. If NULL (default), the pooled sample covariance is used. Only relevant when distance = "mahalanobis".
memory_mode	One of "auto" (default), "dense", or "lazy". "auto" warns (or, when method is "jv"/"auction" with a built-in distance metric, switches) when the dense cost matrix would consume a large fraction of free system RAM. "lazy" computes each pairwise distance from the underlying feature data as the solver needs it, instead of allocating the full n_left x n_right matrix; supported for method = "jv"/"auction" with a built-in distance metric, and not yet for replace = TRUE, ratio > 1, method = "greedy", or custom distance functions (blocking via block_id is the other option that reduces memory, by solving smaller sub-problems). "dense" skips the RAM check entirely.

Details

With method set to a LAP solver (the default "auto", or "jv", "hungarian", ...) it finds the matching that minimizes total distance among all feasible matchings. With method = "greedy" it uses a fast greedy strategy (selected by strategy) that does not guarantee the optimal total distance but scales to very large datasets.

Value

A list with class "matching_result" containing:

- pairs: Tibble of matched pairs with distances
- unmatched: List of unmatched left and right IDs
- info: Matching diagnostics and metadata

Examples

```
# Basic matching
left <- data.frame(id = 1:5, x = c(1, 2, 3, 4, 5), y = c(2, 4, 6, 8, 10))
right <- data.frame(id = 6:10, x = c(1.1, 2.2, 3.1, 4.2, 5.1), y = c(2.1, 4.1, 6.2, 8.1, 10.1))
result <- match_couples(left, right, vars = c("x", "y"))
print(result$pairs)

# With constraints
result <- match_couples(left, right, vars = c("x", "y"),
                        max_distance = 1,
                        calipers = list(x = 0.5))
```

```
# With blocking
left$region <- c("A", "A", "B", "B", "B")
right$region <- c("A", "A", "B", "B", "B")
blocks <- matchmaker(left, right, block_type = "group", block_by = "region")
result <- match_couples(blocks$left, blocks$right, vars = c("x", "y"))

# Fast greedy matching for large datasets
result <- match_couples(left, right, vars = c("x", "y"),
                        method = "greedy", strategy = "sorted")
```

match_data

Extract Analysis-Ready Data from Matching Results

Description

A generic function that converts any couplr matching result into a single analysis-ready data frame with weights, subclass, and distance columns. This is the couplr equivalent of MatchIt's `match.data()`.

Usage

```
match_data(result, ...)

## S3 method for class 'matching_result'
match_data(result, left, right, left_id = "id", right_id = "id", ...)

## S3 method for class 'full_matching_result'
match_data(result, left, right, left_id = "id", right_id = "id", ...)

## S3 method for class 'cem_result'
match_data(result, left, right, left_id = "id", right_id = "id", ...)

## S3 method for class 'subclass_result'
match_data(result, data = NULL, ...)
```

Arguments

<code>result</code>	A couplr result object (<code>matching_result</code> , <code>full_matching_result</code> , <code>cem_result</code> , or <code>subclass_result</code>)
<code>...</code>	Additional arguments passed to methods
<code>left</code>	Data frame of left (treated) units
<code>right</code>	Data frame of right (control) units
<code>left_id</code>	Name of ID column in left (default: "id")
<code>right_id</code>	Name of ID column in right (default: "id")
<code>data</code>	Data frame containing all units (for CEM and subclassification, left and right are not always needed separately)

Details

The output format is compatible with downstream packages like **cobalt**, **WeightIt**, and **marginal-effects**. The stacked (long) format with treatment and weights columns is the standard layout expected by these tools.

Value

A tibble with all original variables plus standardized columns:

id Unit identifier

treatment 1 for left/treated, 0 for right/control

weights Matching weights

subclass Matched group/stratum identifier

distance Matching distance (where applicable)

Examples

```
set.seed(42)
left <- data.frame(id = 1:5, age = c(25, 35, 45, 55, 65))
right <- data.frame(id = 6:15, age = runif(10, 20, 70))
result <- match_couples(left, right, vars = "age")
md <- match_data(result, left, right)
head(md)
```

pixel_morph

Pixel-level image morphing (final frame only)

Description

Computes optimal pixel assignment from A to B and returns the final transported frame (without intermediate animation frames).

Usage

```
pixel_morph(
  imgA,
  imgB,
  n_frames = 16L,
  mode = c("color_walk", "exact", "recursive", "color_match"),
  lap_method = "jv",
  maximize = FALSE,
  quantize_bits = 5L,
  downscale_steps = 0L,
  alpha = 1,
  beta = 0,
```

```

    patch_size = 1L,
    upscale = 1,
    show = interactive()
)

```

Arguments

imgA	Source image (file path or magick image object)
imgB	Target image (file path or magick image object)
n_frames	Internal parameter for rendering (default: 16)
mode	Assignment algorithm: "color_walk" (default), "exact", "recursive", or "color_match" (spatially match pixels of identical quantized colour, filling the remainder by identity)
lap_method	LAP solver method (default: "jv")
maximize	Logical, maximize instead of minimize cost (default: FALSE)
quantize_bits	Color quantization for "color_walk" mode (default: 5)
downscale_steps	Number of 2x reductions before computing assignment (default: 0)
alpha	Weight for color distance in cost function (default: 1)
beta	Weight for spatial distance in cost function (default: 0)
patch_size	Tile size for tiled modes (default: 1)
upscale	Post-rendering upscaling factor (default: 1)
show	Logical, display result in viewer (default: interactive())

Details

Transport-Only Semantics:

This function returns a SHARP, pixel-perfect transport of A's pixels to positions determined by the assignment to B.

Key Points:

- Assignment computed using: $\text{cost} = \alpha * \text{color_dist} + \beta * \text{spatial_dist}$
- B's COLORS influence assignment but DO NOT appear in output
- Result has A's colors arranged to match B's layout
- No motion blur (unlike intermediate frames in animation)

See [pixel_morph_animate](#) for detailed explanation of assignment vs rendering semantics.

Permutation Warnings:

Assignment is guaranteed to be a bijection (permutation) ONLY when:

- `downscale_steps = 0` (no resolution changes)
- `mode = "exact"` with `patch_size = 1`

With downscaling or tiled modes, assignment may have:

- **Overlaps:** Multiple source pixels map to same destination (last write wins)
- **Holes:** Some destinations never filled (remain transparent)

If assignment is not a bijection (due to downscaling or tiling), a warning will be issued. The result may contain:

- Overlapped pixels (multiple sources -> one destination)
- Transparent holes (some destinations unfilled)

For guaranteed pixel-perfect results, use:

```
pixel_morph(A, B, mode = "exact", downscale_steps = 0)
```

Value

magick image object of the final transported frame

See Also

[pixel_morph_animate](#) for animated version

Examples

```
if (requireNamespace("magick", quietly = TRUE)) {
  imgA <- system.file("extdata/icons/circleA_40.png", package = "couplr")
  imgB <- system.file("extdata/icons/circleB_40.png", package = "couplr")
  if (nzchar(imgA) && nzchar(imgB)) {
    result <- pixel_morph(imgA, imgB, n_frames = 4, show = FALSE)
  }
}
```

pixel_morph_animate *Pixel-level image morphing (animation)*

Description

Creates an animated morph by computing optimal pixel assignment from image A to image B, then rendering intermediate frames showing the transport.

Usage

```
pixel_morph_animate(
  imgA,
  imgB,
  n_frames = 16L,
  fps = 10L,
  format = c("gif", "webp", "mp4"),
  outfile = NULL,
  show = interactive(),
  mode = c("color_walk", "exact", "recursive", "color_match"),
  lap_method = "jv",
  maximize = FALSE,
```

```

    quantize_bits = 5L,
    downscale_steps = 0L,
    alpha = 1,
    beta = 0,
    patch_size = 1L,
    upscale = 1
)

```

Arguments

imgA	Source image (file path or magick image object)
imgB	Target image (file path or magick image object)
n_frames	Integer number of animation frames (default: 16)
fps	Frames per second for playback (default: 10)
format	Output format: "gif", "webp", or "mp4"
outfile	Optional output file path
show	Logical, display animation in viewer (default: interactive())
mode	Assignment algorithm: "color_walk" (default), "exact", "recursive", or "color_match" (spatially match pixels of identical quantized colour, filling the remainder by identity)
lap_method	LAP solver method (default: "jv")
maximize	Logical, maximize instead of minimize cost (default: FALSE)
quantize_bits	Color quantization for "color_walk" mode (default: 5)
downscale_steps	Number of 2x reductions before computing assignment (default: 0)
alpha	Weight for color distance in cost function (default: 1)
beta	Weight for spatial distance in cost function (default: 0)
patch_size	Tile size for tiled modes (default: 1)
upscale	Post-rendering upscaling factor (default: 1)

Details

Assignment vs Rendering Semantics:

CRITICAL: This function has two separate phases with different semantics:

Phase 1 - Assignment Computation:

The assignment is computed by minimizing:

$$\text{cost}(i, j) = \alpha * \text{color_distance}(A[i], B[j]) + \beta * \text{spatial_distance}(\text{pos}_i, \text{pos}_j)$$

This means B's COLORS influence which pixels from A map to which positions.

Phase 2 - Rendering (Transport-Only):

The renderer uses ONLY A's colors:

- Intermediate frames: A's pixels move along paths with motion blur

- Final frame: A's pixels at their assigned positions (sharp, no blur)
- B's colors NEVER appear in the output

Result: You get A's colors rearranged to match B's geometry/layout.

What This Means:

- B influences WHERE pixels go (via similarity in cost function)
- B does NOT determine WHAT COLORS appear in output
- Final image has A's palette arranged to mimic B's structure

Parameter Guidance:

For pure spatial rearrangement (ignore B's colors in assignment):

```
pixel_morph_animate(A, B, alpha = 0, beta = 1)
```

For color-similarity matching (default):

```
pixel_morph_animate(A, B, alpha = 1, beta = 0)
```

For hybrid (color + spatial):

```
pixel_morph_animate(A, B, alpha = 1, beta = 0.2)
```

Permutation Guarantees:

Assignment is guaranteed to be a bijection (permutation) ONLY when:

- `downscale_steps = 0` (no resolution changes)
- `mode = "exact"` with `patch_size = 1`

With downscaling or tiled modes, assignment may have:

- **Overlaps:** Multiple source pixels map to same destination (last write wins)
- **Holes:** Some destinations never filled (remain transparent)

A warning is issued if overlaps/holes are detected in the final frame.

Value

Invisibly returns a list with animation object and metadata:

<code>animation</code>	magick animation object
<code>width</code>	Image width in pixels
<code>height</code>	Image height in pixels
<code>assignment</code>	Integer vector of 1-based assignment indices (R convention)
<code>n_pixels</code>	Total number of pixels
<code>mode</code>	Mode used for matching
<code>upscale</code>	Upscaling factor applied

Examples

```

if (requireNamespace("magick", quietly = TRUE)) {
  imgA <- system.file("extdata/icons/circleA_40.png", package = "couplr")
  imgB <- system.file("extdata/icons/circleB_40.png", package = "couplr")
  if (nzchar(imgA) && nzchar(imgB)) {
    outfile <- tempfile(fileext = ".gif")
    pixel_morph_animate(imgA, imgB, outfile = outfile, n_frames = 4, show = FALSE)
  }
}

```

plot.balance_diagnostics

Plot method for balance diagnostics

Description

Produces a Love plot (dot plot) of standardized differences.

Usage

```

## S3 method for class 'balance_diagnostics'
plot(x, type = c("love", "histogram", "variance"), threshold = 0.1, ...)

```

Arguments

x	A balance_diagnostics object
type	Type of plot: "love" (default), "histogram", or "variance"
threshold	Threshold line for standardized differences (default: 0.1)
...	Additional arguments passed to plotting functions

Value

The balance_diagnostics object (invisibly)

`plot.matching_result` *Plot method for matching results*

Description

Produces a histogram of pairwise distances from a matching result.

Usage

```
## S3 method for class 'matching_result'
plot(x, type = c("histogram", "density", "ecdf"), ...)
```

Arguments

<code>x</code>	A <code>matching_result</code> object
<code>type</code>	Type of plot: "histogram" (default), "density", or "ecdf"
<code>...</code>	Additional arguments passed to plotting functions

Value

The `matching_result` object (invisibly)

`plot.sensitivity_analysis`
Plot method for sensitivity analysis (base graphics)

Description

Plot method for sensitivity analysis (base graphics)

Usage

```
## S3 method for class 'sensitivity_analysis'
plot(x, alpha = 0.05, ...)
```

Arguments

<code>x</code>	A <code>sensitivity_analysis</code> object
<code>alpha</code>	Significance level (default: 0.05)
<code>...</code>	Additional arguments passed to plot

Value

The `sensitivity_analysis` object (invisibly)

`preprocess_matching_vars`*Preprocess matching variables with automatic checks and scaling*

Description

Main preprocessing function that orchestrates variable health checks, categorical encoding, and automatic scaling selection.

Usage

```
preprocess_matching_vars(  
  left,  
  right,  
  vars,  
  auto_scale = TRUE,  
  scale_method = "auto",  
  check_health = TRUE,  
  remove_problematic = TRUE,  
  verbose = TRUE  
)
```

Arguments

<code>left</code>	Data frame of left units
<code>right</code>	Data frame of right units
<code>vars</code>	Character vector of variable names
<code>auto_scale</code>	Logical, whether to perform automatic preprocessing (default: TRUE)
<code>scale_method</code>	Scaling method: "auto", "standardize", "range", "robust", or FALSE
<code>check_health</code>	Logical, whether to check variable health (default: TRUE)
<code>remove_problematic</code>	Logical, automatically exclude constant/all-NA variables (default: TRUE)
<code>verbose</code>	Logical, whether to print warnings (default: TRUE)

Value

A list with class "preprocessing_result" containing:

- `left`: Preprocessed left data frame
- `right`: Preprocessed right data frame
- `vars`: Final variable names (after exclusions)
- `health`: Variable health diagnostics
- `scaling_method`: Selected scaling method
- `excluded_vars`: Variables that were excluded
- `warnings`: List of warnings issued

```
print.balance_diagnostics
```

Print Method for Balance Diagnostics

Description

Print Method for Balance Diagnostics

Usage

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

Arguments

x	A balance_diagnostics object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
print.cem_result
```

Print Method for CEM Results

Description

Print Method for CEM Results

Usage

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

Arguments

x	A cem_result object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

`print.distance_object` *Print Method for Distance Objects*

Description

Print Method for Distance Objects

Usage

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

Arguments

<code>x</code>	A <code>distance_object</code>
<code>...</code>	Additional arguments (ignored)

Value

Invisibly returns the input object `x`.

`print.full_matching_result`
Print Method for Full Matching Results

Description

Print Method for Full Matching Results

Usage

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

Arguments

<code>x</code>	A <code>full_matching_result</code> object
<code>...</code>	Additional arguments (ignored)

Value

Invisibly returns the input object `x`.

```
print.lap_solve_batch_result
```

Print method for batch assignment results

Description

Prints a summary and the table of results for a batch of assignment problems solved with `lap_solve_batch()`.

Usage

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

Arguments

`x` A `lap_solve_batch_result` object.
`...` Additional arguments passed to `print()`. Currently ignored.

Value

Invisibly returns the input object `x`.

```
print.lap_solve_kbest_result
```

Print method for k-best assignment results

Description

Print method for k-best assignment results

Usage

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

Arguments

`x` A `lap_solve_kbest_result`.
`...` Additional arguments passed to `print()`. Ignored.

Value

Invisibly returns the input object `x`.

```
print.lap_solve_result
```

Print method for assignment results

Description

Nicely prints a lap_solve_result object, including the assignments, total cost, and method used.

Usage

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

Arguments

x	A lap_solve_result object.
...	Additional arguments passed to print(). Currently ignored.

Value

Invisibly returns the input object x.

```
print.matching_result
```

Print method for matching results

Description

Print method for matching results

Usage

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

Arguments

x	A matching_result object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
print.matchmaker_result
```

Print method for matchmaker results

Description

Print method for matchmaker results

Usage

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

Arguments

x	A matchmaker_result object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
print.preprocessing_result
```

Print method for preprocessing result

Description

Print method for preprocessing result

Usage

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

Arguments

x	A preprocessing_result object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
print.sensitivity_analysis
```

Print method for sensitivity analysis

Description

Print method for sensitivity analysis

Usage

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

Arguments

x	A sensitivity_analysis object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
print.subclass_result
```

Print Method for Subclassification Results

Description

Print Method for Subclassification Results

Usage

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

Arguments

x	A subclass_result object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
print.variable_health Print method for variable health
```

Description

Print method for variable health

Usage

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

Arguments

x	A variable_health object
...	Additional arguments (ignored)

Value

Invisibly returns the input object x.

```
ps_match Propensity Score Matching
```

Description

Matches treated and control units based on estimated propensity scores. Fits a logistic regression model (or accepts a pre-fitted one), computes logit propensity scores, and calls [match_couples\(\)](#) with a caliper on the logit scale.

Usage

```
ps_match(
  formula = NULL,
  data = NULL,
  treatment = NULL,
  ps_model = NULL,
  caliper_sd = 0.2,
  method = "auto",
  replace = FALSE,
  ratio = 1L,
  ...
)
```

Arguments

formula	Formula for propensity score model (treatment ~ covariates). Required if ps_model is NULL.
data	Combined dataset containing both treated and control units
treatment	Name of the binary treatment column (0/1 or logical)
ps_model	Pre-fitted glm object (alternative to formula). If provided, formula is ignored.
caliper_sd	Caliper width in standard deviations of logit(PS). Default: 0.2 (Rosenbaum and Rubin recommendation).
method	LAP solver method (default: "auto")
replace	If TRUE, match with replacement (default: FALSE)
ratio	Integer k for k:1 matching (default: 1)
...	Additional arguments passed to match_couples()

Details

The propensity score is the probability of treatment assignment conditional on observed covariates. Matching is performed on the logit of the propensity score (Rosenbaum and Rubin 1985), which provides better distributional properties than matching on the raw probability scale.

The default caliper of 0.2 SD of logit(PS) is recommended by Austin (2011) as removing approximately 98% of bias.

Value

A `matching_result` object with additional propensity score info in `result$info$ps_model` and `result$info$caliper_value`.

Examples

```
set.seed(42)
n <- 100
data <- data.frame(
  id = seq_len(n),
  treated = rbinom(n, 1, 0.4),
  age = rnorm(n, 50, 10),
  income = rnorm(n, 50000, 15000)
)
result <- ps_match(treated ~ age + income, data = data, treatment = "treated")
print(result)
```

sensitivity_analysis *Rosenbaum Sensitivity Analysis*

Description

Assesses how sensitive a matched comparison is to hidden bias using Rosenbaum bounds on the Wilcoxon signed-rank statistic.

Usage

```
sensitivity_analysis(
  result,
  left,
  right,
  outcome_var,
  gamma = seq(1, 3, by = 0.25),
  alternative = c("greater", "less", "two.sided"),
  left_id = "id",
  right_id = "id"
)
```

Arguments

result	A <code>matching_result</code> object from <code>match_couples()</code>
left	Original left (treated) dataset
right	Original right (control) dataset
outcome_var	Name of the outcome column in <code>left</code> and <code>right</code>
gamma	Numeric vector of sensitivity parameters (default: <code>seq(1, 3, by = 0.25)</code>). Gamma = 1 means no hidden bias.
alternative	Direction of the test: "greater" (default), "less", or "two.sided"
left_id	Name of ID column in left (default: "id")
right_id	Name of ID column in right (default: "id")

Details

Rosenbaum (2002, Chapter 4) bounds quantify how much hidden bias (an unobserved confounder) would be needed to explain away the observed treatment effect. The sensitivity parameter Gamma represents the maximum ratio of treatment odds between two matched units:

- Gamma = 1: No hidden bias (standard Wilcoxon test)
- Gamma = 2: One unit could be twice as likely to receive treatment due to an unobserved factor

The function computes upper and lower bounds on the p-value of the Wilcoxon signed-rank test under each level of hidden bias. A finding is "insensitive to bias" if `p_upper` remains below 0.05 even at large Gamma.

Value

An S3 object of class `sensitivity_analysis` containing:

results Tibble with columns: `gamma`, `t_stat`, `p_upper`, `p_lower`

n_pairs Number of matched pairs analyzed

critical_gamma Smallest `gamma` at which `p_upper` > 0.05

alternative Direction of test

References

Rosenbaum, P.R. (2002). *Observational Studies*, 2nd edition. Springer.

Examples

```
set.seed(42)
left <- data.frame(id = 1:20, x = rnorm(20), outcome = rnorm(20, 1, 1))
right <- data.frame(id = 21:40, x = rnorm(20), outcome = rnorm(20, 0, 1))
result <- match_couples(left, right, vars = "x")
sens <- sensitivity_analysis(result, left, right, outcome_var = "outcome")
print(sens)
```

sinkhorn

'Sinkhorn-Knopp' optimal transport solver

Description

Compute an entropy-regularized optimal transport plan using the 'Sinkhorn-Knopp' algorithm. Unlike other LAP solvers that return a hard 1-to-1 assignment, this returns a soft assignment (doubly stochastic matrix).

Usage

```
sinkhorn(
  cost,
  lambda = 10,
  tol = 1e-09,
  max_iter = 1000,
  r_weights = NULL,
  c_weights = NULL
)
```

Arguments

<code>cost</code>	Numeric matrix of transport costs. NA or Inf entries are treated as very high cost (effectively forbidden).
<code>lambda</code>	Regularization parameter (default 10). Higher values produce sharper (more deterministic) transport plans; lower values produce smoother distributions. Typical range: 1-100.
<code>tol</code>	Convergence tolerance (default 1e-9).
<code>max_iter</code>	Maximum iterations (default 1000).
<code>r_weights</code>	Optional numeric vector of row marginals (source distribution). Default is uniform. Will be normalized to sum to 1.
<code>c_weights</code>	Optional numeric vector of column marginals (target distribution). Default is uniform. Will be normalized to sum to 1.

Details

The 'Sinkhorn-Knopp' algorithm solves the entropy-regularized optimal transport problem:

$$P^* = \arg \min_P \langle C, P \rangle - \frac{1}{\lambda} H(P)$$

subject to row sums = `r_weights` and column sums = `c_weights`.

The entropy term $H(P)$ encourages spread in the transport plan. As $\lambda \rightarrow \infty$, the solution approaches the standard (unregularized) optimal transport.

Key differences from standard LAP solvers:

- Returns a soft assignment (probabilities) not a hard 1-to-1 matching
- Supports unequal marginals (weighted distributions)
- Differentiable, making it useful in ML pipelines
- Very fast: $O(n^2)$ per iteration with typically $O(1/\text{tol}^2)$ iterations

Use `sinkhorn_to_assignment()` to round the soft assignment to a hard matching.

Value

A list with elements:

- `transport_plan` — numeric matrix, the optimal transport plan P . Row sums approximate `r_weights`, column sums approximate `c_weights`.
- `cost` — the transport cost $\langle C, P \rangle$ (without entropy term).
- `u, v` — scaling vectors ($P = \text{diag}(u) * K * \text{diag}(v)$ where $K = \exp(-\lambda * C)$).
- `converged` — logical, whether the algorithm converged.
- `iterations` — number of iterations used.
- `lambda` — the regularization parameter used.

References

Cuturi, M. (2013). 'Sinkhorn Distances': Lightspeed Computation of Optimal Transport. *Advances in Neural Information Processing Systems*, 26.

See Also

[assignment\(\)](#) for hard 1-to-1 matching, [sinkhorn_to_assignment\(\)](#) to round soft assignments.

Examples

```
cost <- matrix(c(1, 2, 3, 4, 5, 6, 7, 8, 9), nrow = 3, byrow = TRUE)

# Soft assignment with default parameters
result <- sinkhorn(cost)
print(round(result$transport_plan, 3))

# Sharper assignment (higher lambda)
result_sharp <- sinkhorn(cost, lambda = 50)
print(round(result_sharp$transport_plan, 3))

# With custom marginals (more mass from row 1)
result_weighted <- sinkhorn(cost, r_weights = c(0.5, 0.25, 0.25))
print(round(result_weighted$transport_plan, 3))

# Round to hard assignment
hard_match <- sinkhorn_to_assignment(result)
print(hard_match)
```

sinkhorn_to_assignment

Round 'Sinkhorn' transport plan to hard assignment

Description

Convert a soft transport plan from [sinkhorn\(\)](#) to a hard 1-to-1 assignment using greedy rounding.

Usage

```
sinkhorn_to_assignment(result)
```

Arguments

result Either a result from [sinkhorn\(\)](#) or a transport plan matrix.

Details

Greedy rounding iteratively assigns each row to its most probable column, ensuring no column is assigned twice. This may not give the globally optimal hard assignment; for that, use the transport plan as a cost matrix with [assignment\(\)](#).

Value

Integer vector of column assignments (1-based), same format as `assignment()`.

See Also

`sinkhorn()`

Examples

```
cost <- matrix(c(1, 2, 3, 4, 5, 6, 7, 8, 9), nrow = 3, byrow = TRUE)
result <- sinkhorn(cost, lambda = 20)
hard_match <- sinkhorn_to_assignment(result)
print(hard_match)
```

subclass_match	<i>Subclassification on Propensity Score</i>
----------------	--

Description

Divides units into K strata based on quantiles of the propensity score, then computes within-stratum weights for treatment effect estimation. This is a simple, transparent approach to propensity score adjustment that allows visual inspection of balance within each subclass.

Usage

```
subclass_match(
  formula = NULL,
  data = NULL,
  treatment = NULL,
  n_subclasses = 5L,
  ps = NULL,
  ps_model = NULL,
  estimand = "ATT"
)
```

Arguments

formula	Formula for propensity score model (e.g., <code>treatment ~ age + income</code>). Ignored if <code>ps</code> is provided.
data	Data frame containing all variables
treatment	Character, name of the binary treatment column (0/1)
n_subclasses	Integer, number of subclasses to create (default: 5). Cochran (1968) showed that 5 subclasses removes over 90\ a single covariate.
ps	Optional pre-computed numeric vector of propensity scores (one per row in data). If NULL, a logistic regression model is fit using formula.
ps_model	Optional pre-fitted glm object for propensity scores
estimand	Target estimand: "ATT" (default), "ATE", or "ATC"

Details

The algorithm:

1. Estimate propensity scores via logistic regression (or use pre-computed scores)
2. Divide the propensity score distribution into K quantile-based strata
3. For each stratum, check overlap (both treated and control units present)
4. Compute within-stratum weights based on the target estimand:
 - **ATT**: Treated units get weight 1; control units get weight $n_{\text{treated_in_stratum}} / n_{\text{control_in_stratum}}$
 - **ATE**: Both groups get weight proportional to stratum size relative to total sample
 - **ATC**: Control units get weight 1; treated units get weight $n_{\text{control_in_stratum}} / n_{\text{treated_in_stratum}}$

Value

An S3 object of class `c("subclass_result", "couplr_result")` containing:

matched Tibble with columns `id`, `side`, `subclass`, `ps`, `weight`

subclass_summary Tibble with per-subclass statistics: counts, mean PS, and overlap status

info List with `n_subclasses`, `estimand`, `n_left`, `n_right`, `method`, `vars`

Examples

```
set.seed(42)
n <- 200
data <- data.frame(
  id = 1:n,
  age = rnorm(n, 40, 10),
  income = rnorm(n, 50000, 15000)
)
data$treatment <- rbinom(n, 1, plogis(-2 + 0.05 * data$age))
result <- subclass_match(treatment ~ age + income, data, treatment = "treatment")
print(result)
```

```
summary.balance_diagnostics
```

Summary method for balance diagnostics

Description

Summary method for balance diagnostics

Usage

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

Arguments

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

Value

A list containing summary statistics (invisibly)

summary.distance_object

Summary Method for Distance Objects

Description

Summary Method for Distance Objects

Usage

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

Arguments

object	A distance_object
...	Additional arguments (ignored)

Value

Invisibly returns the input object.

summary.lap_solve_kbest_result

Get summary of k-best results

Description

Extract summary information from k-best assignment results.

Usage

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

Arguments

object	An object of class lap_solve_kbest_result.
...	Additional arguments (unused).

Value

A tibble with one row per solution containing:

- rank: solution rank
- solution_id: solution identifier
- total_cost: total cost of the solution
- n_assignments: number of assignments in the solution

`summary.matching_result`*Summary method for matching results*

Description

Summary method for matching results

Usage

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

Arguments

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

Value

A list containing summary statistics (invisibly)

```
summary.sensitivity_analysis
```

Summary method for sensitivity analysis

Description

Summary method for sensitivity analysis

Usage

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

Arguments

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

Value

A list containing summary statistics (invisibly)

```
update_constraints
```

Update Constraints on Distance Object

Description

Apply new constraints to a precomputed distance object without recomputing the underlying distances. This is useful for exploring different constraint scenarios quickly.

Usage

```
update_constraints(dist_obj, max_distance = Inf, calipers = NULL)
```

Arguments

dist_obj	A distance_object from compute_distances()
max_distance	Maximum allowed distance (pairs with distance > max_distance become Inf)
calipers	Named list of per-variable calipers

Details

This function creates a new `distance_object` with modified constraints applied to the cost matrix. The original `distance_object` is not modified.

Constraints:

- `max_distance`: Sets cost to Inf for pairs exceeding this threshold
- `calipers`: Per-variable restrictions (e.g., `calipers = list(age = 5)`)

The function returns a new object rather than modifying in place, following R's copy-on-modify semantics.

Value

A new `distance_object` with updated `cost_matrix`

Examples

```
left <- data.frame(id = 1:5, age = c(25, 30, 35, 40, 45))
right <- data.frame(id = 6:10, age = c(24, 29, 36, 41, 44))
dist_obj <- compute_distances(left, right, vars = "age")

# Apply constraints
constrained <- update_constraints(dist_obj, max_distance = 2)
result <- match_couples(constrained)
```

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