

# Package: vennDiagramLab (via r-universe)

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**Type** Package

**Title** Headless Venn Diagram Analysis and Rendering

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**Description** Headless companion to the 'Venn Diagram Lab' web tool (<<https://www.venndiagramlab.org/>>). Build, render, and statistically analyze Venn / 'UpSet' diagrams from 'CSV' / 'TSV' / 'GMT' / 'GMX' inputs. Provides the same 44 SVG models, intersection / 'Jaccard' / hypergeometric statistics, and PDF report layout as the web tool, with byte-equivalent 'TSV' exports (parity-tested against the published Python package). Integrates with 'ggplot2', 'tidygraph', and 'broom'.

**License** MIT + file LICENSE

**Language** en-US

**URL** <https://zoliqua.github.io/Venn-Diagram-Lab/r/>,  
<https://github.com/ZoliQua/Venn-Diagram-Lab>,  
<https://www.venndiagramlab.org/>

**BugReports** <https://github.com/ZoliQua/Venn-Diagram-Lab/issues>

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 'io.R' 'proportional.R' 'region-accessors.R'  
 'region-expression.R' 'render-enrichment-bar.R'  
 'render-enrichment-lollipop.R' 'render-network.R'  
 'render-pdf.R' 'render-share-distribution.R'  
 'render-svg-items.R' 'render-svg.R' 'render-upset.R'  
 'samples.R' 'share-distribution.R' 'statistics.R'  
 'tsv-export.R' 'zip-report.R' 'zzz.R'

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analyze	<i>Analyze a Venn diagram dataset</i>
---------	---------------------------------------

---

## Description

Compute the Venn region map for a [`VennDataset-class`] and bind it to a model.

## Usage

```
analyze(dataset, model = "auto")
```

## Arguments

dataset	A [ <code>VennDataset-class</code> ] (from one of the <code>load_*</code> functions).
model	Model identifier. <code>"auto"</code> picks the canonical model for the dataset's set count (alphabetical first match), e.g. 4 sets -> <code>'venn-4-set'</code> . <code>"proportional"</code> requests an area-proportional layout (only supports 2-3 sets, added in Phase 3). Otherwise pass an explicit name from <code>[list_models()]</code> .

**Value**

A [`'RegionResult-class'`] with the per-region item membership, set sizes, and (lazily) `'statistics(result)'`.

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
result@model

ds <- load_sample("dataset_real_cancer_drivers_4")
result <- analyze(ds, model = "auto")
result@model
```

---

augment.RegionResult *Augment method for RegionResult (broom-compatible)*

---

**Description**

Returns one row per item in the dataset's universe, with boolean columns indicating set membership and a `'region_label'` column naming the exact region (e.g. `"A"`, `"AB"`, `"ABC"`) the item belongs to. Item ordering follows `'dataset@item_order'` (first-seen across all sets, JS Set/Map semantics).

**Usage**

```
## S3 method for class 'RegionResult'
augment(x, ...)
```

**Arguments**

```
x          A ['RegionResult-class'].
...        Unused (broom convention).
```

**Details**

Region labels use the package's positional letter convention (A-I), matching the labels in `'RegionResult@regions'` and the bundled SVG models, regardless of the dataset's `'set_names'`.

**Value**

A tibble (or data.frame fallback) with `'nrow(out) == length(x@dataset@item_order)'` and columns: `'item'` (character), one logical column per set (named after the set), `'region_label'` (character).

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
if (requireNamespace("broom", quietly = TRUE)) broom::augment(result)

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
broom::augment(result)
```

---

bh\_fdr

*Benjamini-Hochberg FDR adjustment*

---

**Description**

Wraps 'stats::p.adjust(p, method = "BH)'. Returns adjusted p-values in the same order as the input. Empty input -> empty output.

**Usage**

```
bh_fdr(p_values)
```

**Arguments**

p\_values      Numeric vector of raw p-values in [0, 1].

**Value**

Numeric vector of adjusted p-values, same length as input.

**Examples**

```
bh_fdr(c(0.001, 0.01, 0.05, 0.5))
```

---

circle\_intersection\_area

*Lens-shaped intersection area of two circles*

---

### Description

Mirrors 'src/utils/proportionalLayout.ts circleIntersectionArea' (web tool) and Python 'circle\_intersection\_area' byte-for-byte.

### Usage

```
circle_intersection_area(r1, r2, d)
```

### Arguments

r1	Radius of circle 1 (positive numeric).
r2	Radius of circle 2 (positive numeric).
d	Distance between centers (non-negative numeric).

### Value

Numeric: 0 if circles are disjoint,  $\pi * \min(r1,r2)^2$  if fully nested, else the lens-shaped intersection area.

### Examples

```
circle_intersection_area(1, 1, 1) # ~ 1.228
circle_intersection_area(1, 1, 3) # 0 (disjoint)
```

---

cluster\_set\_order

*Hierarchical clustering on a symmetric distance matrix.*

---

### Description

Wraps `stats::hclust` to produce a normalized output that mirrors the webtool's pure-JS `clusterSetOrder` and Python `cluster_set_order`.

### Usage

```
cluster_set_order(D, linkage = c("average", "complete", "single"))
```

### Arguments

D	Symmetric numeric distance matrix (NxN, zero diagonal).
linkage	One of "average" (UPGMA), "complete", "single".

**Value**

A list with:

- `leaf_order`: 1-based integer vector — left-to-right ordering of original indices. At each internal node the subtree whose minimum original leaf index is smaller is placed on the left (deterministic; mirrors the webtool / Python convention).
- `merges`: data.frame with columns `left`, `right` (0-based cluster ids matching the webtool/Python convention; leaves are 0..N-1, internal nodes are N..2N-2), `height` (linkage distance), `size` (number of leaves in the merged cluster).

**Examples**

```
D <- matrix(c(0, 0.2, 0.9, 0.2, 0, 0.85, 0.9, 0.85, 0), nrow = 3)
cluster_set_order(D, linkage = "average")
```

---

compute_pairwise	<i>Compute all 5 pairwise statistical tables</i>
------------------	--

---

**Description**

Orchestrator that returns a [`'StatisticsResult-class'`] populated with Jaccard, Dice, Overlap Coefficient, Fold Enrichment (square NxN matrices) plus a long-form hypergeometric table with BH-FDR adjustment.

**Usage**

```
compute_pairwise(
  set_names,
  inclusive_sizes,
  pairwise_intersections,
  universe_size
)
```

**Arguments**

`set_names` Ordered character vector of set identifiers (e.g. `c("A","B","C")`).

`inclusive_sizes` Named integer vector of inclusive set sizes (`'names(inclusive_sizes)'` matches `'set_names'`).

`pairwise_intersections` Named list of pair intersection counts. Keys are `"set_a|set_b"` with `set_a` appearing earlier in `'set_names'` than `set_b`.

`universe_size` Hypergeometric universe N (population size). Integer  $\geq 1$ .

**Value**

A [`'StatisticsResult-class'`] object.

**Examples**

```
compute_pairwise(
  set_names = c("A", "B"),
  inclusive_sizes = c(A = 10L, B = 8L),
  pairwise_intersections = list("A|B" = 5L),
  universe_size = 100L
)
```

---

dice	<i>Sorensen-Dice coefficient</i>
------	----------------------------------

---

**Description**

Computes  $2 * |A \text{ intersection } B| / (|A| + |B|)$ . Returns 0 if both sets are empty (matches web tool convention).

**Usage**

```
dice(size_a, size_b, intersection)
```

**Arguments**

size_a	Inclusive size of set A (integer $\geq 0$ ).
size_b	Inclusive size of set B (integer $\geq 0$ ).
intersection	Inclusive intersection size $ A \text{ intersection } B $ .

**Value**

Numeric in  $[0, 1]$ .

**Examples**

```
dice(10, 10, 5)
```

---

effective_universe	<i>Effective hypergeometric universe size for a RegionResult</i>
--------------------	--

---

**Description**

Returns the universe  $N$  consistent with the web tool. Binary CSV/TSV datasets get 'dataset@universe\_size' (= csv.rows.length, includes all-zero rows); aggregated/GMT/GMX datasets fall back to 'length(item\_order)' (= lunion of items).

**Usage**

```
effective_universe(result)

## S4 method for signature 'RegionResult'
effective_universe(result)
```

**Arguments**

result            A [`'RegionResult-class'`].

**Value**

Integer, the universe size N.

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
effective_universe(result)

ds <- load_sample("dataset_real_cancer_drivers_4")
result <- analyze(ds)
effective_universe(result) # 20000 for binary cancer drivers sample
```

---

exclusive_items	<i>Items exclusive to a specific set combination</i>
-----------------	--

---

**Description**

Returns items in EXACTLY the combination of sets listed in `'sets'` — that is, in every set in `'sets'` AND in none of the other sets in the dataset.

**Usage**

```
exclusive_items(result, sets)
```

**Arguments**

result            A [`'RegionResult-class'`] from `[analyze()]`.

sets              Character vector of set letters (`"A"`, `"B"`, ...) or display names (values from `'result@dataset@set_names'`). May mix both.

**Value**

A character vector of item IDs. Empty character if none.

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B", "C"),
  items = list(A = c("x", "y"), B = c("y", "z"), C = c("z", "w")),
  item_order = c("x", "y", "z", "w"),
  universe_size = 10L, source_path = NULL, format = "csv")
res <- analyze(ds)
exclusive_items(res, c("A", "B")) # "y" (in A and B, not in C)
```

---

<code>fold_enrichment</code>	<i>Fold enrichment (observed / expected ratio)</i>
------------------------------	--

---

**Description**

Computes  $(k * N) / (K * n)$ . Returns 0.0 if any denominator is zero (matches web tool convention).

**Usage**

```
fold_enrichment(N, K, n, k)
```

**Arguments**

<code>N</code>	Population size (total items in the universe). Integer $\geq 1$ .
<code>K</code>	Number of success states in the population (e.g. inclusive  A ). Integer $\geq 0$ .
<code>n</code>	Number of draws (e.g. inclusive  B ). Integer $\geq 0$ .
<code>k</code>	Observed successes (e.g.  A intersection B ). Integer $\geq 0$ .

**Value**

Numeric ( $\geq 0$ ; can exceed 1 for over-representation).

**Examples**

```
fold_enrichment(20000, 138, 581, 126)
```

---

`generate_proportional_svg`*Generate an area-proportional SVG for a 2- or 3-set RegionResult*

---

## Description

Circle sizes and inter-circle distances are solved analytically (2-set, exact) or by triangulation (3-set, approximate) so that overlap areas match the requested intersection counts. The returned SVG matches the 44-model schema: ShapeA-I, NameA-I, Count\_\*, CountSUM\_\*, Bullet\* elements are all present and addressable via xml2.

## Usage

```
generate_proportional_svg(  
  result,  
  width = .PROP_DEFAULT_WIDTH,  
  height = .PROP_DEFAULT_HEIGHT  
)
```

## Arguments

<code>result</code>	A [ <code>'RegionResult-class'</code> ].
<code>width</code>	Canvas width in pixels (default 600).
<code>height</code>	Canvas height in pixels (default 600).

## Value

A `'character'` (length 1) with the raw SVG.

## Examples

```
tmp <- tempfile(fileext = ".tsv")  
writeLines(c("Gene\tSetA\tSetB",  
            "GENE1\t1\t0",  
            "GENE2\t1\t1",  
            "GENE3\t0\t1"), tmp)  
ds <- load_tsv(tmp, binary = TRUE)  
res <- analyze(ds, model = "proportional")  
svg <- generate_proportional_svg(res)  
nchar(svg) > 0
```

geom\_venn

*Embed a rendered Venn diagram as a ggplot2 layer***Description**

Returns a list of ggplot2 layers that draw ‘data’ (a [‘RegionResult-class’]) as a rasterized Venn diagram on a unit-square coordinate system, ready to compose with other ggplot2 elements (titles, themes, additional annotations).

**Usage**

```
geom_venn(
  mapping = NULL,
  data = NULL,
  stat = "identity",
  position = "identity",
  ...,
  width_px = .GEOM_VENN_DEFAULT_WIDTH
)
```

**Arguments**

mapping	Accepted for ggplot2 layer-signature consistency. Currently ignored (the Venn diagram is rendered from ‘data’, not from aesthetic mappings). Reserved for a future Stat-based extension.
data	A [‘RegionResult-class’] (required). The Venn diagram to embed.
stat	Accepted for signature consistency; currently ignored.
position	Accepted for signature consistency; currently ignored.
...	Forwarded to ‘ggplot2::annotation_custom()’ (e.g. ‘xmin’, ‘xmax’, ‘ymin’, ‘ymax’ to position the venn on a non-unit coordinate system).
width_px	Raster width in pixels (default 800). Larger values give sharper output at the cost of memory.

**Details**

This is a NEW capability – the Python package has no equivalent. It uses the same rasterization pipeline as [to\_pdf\_report()]: render the SVG via [render\_venn\_svg()], rasterize via ‘rsvg::rsvg\_nativeraster()’, and wrap as a ‘grid::rasterGrob()’ inside ‘ggplot2::annotation\_custom()’.

**Value**

A list of ggplot2 layers: an ‘annotation\_custom’ carrying the rasterized Venn, a ‘geom\_blank’ establishing ‘[0, 1] x [0, 1]’ limits, and a ‘coord\_fixed(ratio = 1)’ so the diagram remains square. Note that ‘coord\_fixed’ will override any coordinate system the user has already added; add ‘geom\_venn()’ before other coord layers to avoid a warning.

**Examples**

```

ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
if (getRversion() >= "4.6") {
  p <- ggplot2::ggplot() + geom_venn(data = result) + ggplot2::theme_void()
  inherits(p, "ggplot")
}

if (getRversion() >= "4.6") {
  result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
  library(ggplot2)
  ggplot() +
    geom_venn(data = result) +
    theme_void() +
    ggtitle("Cancer driver overlap (4 sources)")
}

```

---

glance.RegionResult    *Glance method for RegionResult (broom-compatible)*

---

**Description**

Returns a 1-row tibble summarizing the analysis: number of sets, number of non-empty regions, total unique items, hypergeometric universe size, resolved model name, whether the layout is approximate (proportional 3-set), and the count of statistically significant / highly significant pairs (FDR-adjusted  $q < 0.05$  /  $< 0.001$ ).

**Usage**

```
## S3 method for class 'RegionResult'
glance(x, ...)
```

**Arguments**

x                    A [`'RegionResult-class'`].

...                    Unused (broom convention).

**Value**

A 1-row tibble (or data.frame fallback) with columns: `'n_sets'`, `'n_regions'`, `'n_items'`, `'universe_size'`, `'model'`, `'is_approximate'`, `'n_significant'`, `'n_highly_significant'`.

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
if (requireNamespace("broom", quietly = TRUE)) broom::glance(result)

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
broom::glance(result)
```

---

hypergeometric\_p\_value

*One-sided hypergeometric p-value (over-representation)*

---

**Description**

Computes  $P(X \geq k)$  where  $X \sim \text{Hypergeometric}(N, K, n)$ . Returns 1.0 for invalid inputs so the metric is safe to feed into BH-FDR without filtering.

**Usage**

```
hypergeometric_p_value(N, K, n, k)
```

**Arguments**

N	Population size (total items in the universe). Integer $\geq 1$ .
K	Number of success states in the population (e.g. inclusive  A ). Integer $\geq 0$ .
n	Number of draws (e.g. inclusive  B ). Integer $\geq 0$ .
k	Observed successes (e.g.  A intersection B ). Integer $\geq 0$ .

**Details**

Maps to R's 'phyper(k - 1, K, N - K, n, lower.tail = FALSE)'. Note that R's phyper parameter convention differs from Python's scipy: R uses 'm' for success-in-population and 'n' for failure-in-population ( $= N - K$ ), where Python uses 'N' for total population.

**Value**

Numeric in [0, 1].

**Examples**

```
hypergeometric_p_value(20000, 138, 581, 126)
```

---

intersection\_items      *Items in the intersection of the named sets*

---

**Description**

Returns items that appear in every set listed in ‘sets’, regardless of whether they also appear in other (unlisted) sets. For "items in this specific combination only", see [exclusive\_items()].

**Usage**

```
intersection_items(result, sets)
```

**Arguments**

**result**            A [‘RegionResult-class’] from [analyze()].

**sets**              Character vector of set letters (“A”, “B”, ...) or display names (values from ‘result@dataset@set\_names’). May mix both.

**Value**

A character vector of item IDs. Empty character if none.

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B", "C"),
  items = list(A = c("x", "y"), B = c("y", "z"), C = c("z", "w")),
  item_order = c("x", "y", "z", "w"),
  universe_size = 10L, source_path = NULL, format = "csv")
res <- analyze(ds)
intersection_items(res, c("A", "B")) # "y"
```

---

item\_share\_distribution

*Item Share Distribution*

---

**Description**

Per-membership-count item totals.

**Usage**

```
item_share_distribution(matrix)
```

**Arguments**

**matrix**            Binary item × set matrix.

**Details**

Given a binary item  $\times$  set matrix (rows = items, columns = sets, cells in  $\{0, 1\}$ ), returns a named integer vector keyed by  $k = 1..n\_sets$  giving the number of items belonging to exactly  $k$  sets. All bins are present even when their count is zero. Rows that sum to zero are skipped (universe-rule violation; defensive).

**Value**

Named integer vector with names "1", "2", ..., "n\_sets".

**Examples**

```
m <- matrix(c(
  1, 0, 0,
  1, 1, 0,
  1, 1, 1
), ncol = 3, byrow = TRUE)
item_share_distribution(m)
```

---

jaccard

*Jaccard similarity index*


---

**Description**

Computes  $|A \cap B| / |A \cup B|$ . Matches the web tool's convention of returning 0 when both sets are empty (NaN-safe).

**Usage**

```
jaccard(size_a, size_b, intersection)
```

**Arguments**

size_a	Inclusive size of set A (integer $\geq 0$ ).
size_b	Inclusive size of set B (integer $\geq 0$ ).
intersection	Inclusive intersection size $ A \cap B $ .

**Value**

Numeric in  $[0, 1]$ .

**Examples**

```
jaccard(10, 10, 5)
jaccard(0, 0, 0)
```

---

list_models	<i>List all bundled Venn diagram models</i>
-------------	---

---

**Description**

Returns metadata for the 44 bundled SVG model templates plus the ‘proportional’ synthetic generator (added in Phase 3). Read from JSON region files in ‘inst/extdata/models/json/’.

**Usage**

```
list_models()
```

**Value**

A ‘data.frame’ with columns ‘name’ (filename stem), ‘set\_count’ (2-9), and ‘display\_name’ (from the JSON ‘name’ field). Sorted by ‘(set\_count, name)’.

**Examples**

```
head(list_models())
```

---

list_samples	<i>List bundled sample dataset names</i>
--------------	--

---

**Description**

Returns the names of the 5 bundled sample datasets, sorted alphabetically. Use [load\_sample()] to load one.

**Usage**

```
list_samples()
```

**Value**

Character vector of 5 sample identifiers.

**Examples**

```
list_samples()
```

---

load_csv	<i>Load a delimited file (CSV/TSV) into a [<code>VennDataset-class</code>]</i>
----------	--

---

### Description

Supports two layouts: \* Binary mode (default): one row per item, with 0/1 columns marking membership in each set. The first `prefix_cols` columns are item metadata; remaining columns are sets.  
 \* Aggregated mode (`binary = FALSE`): each column is a set, and cells contain item identifiers. Empty cells are ignored.

### Usage

```
load_csv(path, binary = TRUE, delimiter = NULL, prefix_cols = 1L)
```

### Arguments

path	Path to the file.
binary	'TRUE' for binary 0/1 mode (default), 'FALSE' for aggregated.
delimiter	Explicit delimiter override. 'NULL' auto-detects from ',', ';', tab, and space.
prefix_cols	Number of leading metadata columns in binary mode (default 1). Ignored when <code>binary = FALSE</code> .

### Value

A [`VennDataset-class`].

### Examples

```
tmp <- tempfile(fileext = ".csv")
writeLines(c("Gene,SetA,SetB", "G1,1,0", "G2,1,1", "G3,0,1"), tmp)
ds <- load_csv(tmp, binary = TRUE)
ds@set_names
```

---

load_gmt	<i>Load a GMT (Gene Matrix Transposed) file into a [<code>VennDataset-class</code>]</i>
----------	---

---

### Description

Each line is one set: `'set_name<TAB>description<TAB>item1<TAB>item2<TAB>...'`. Lines with fewer than 3 tab-separated columns or empty set names are skipped.

### Usage

```
load_gmt(path)
```

**Arguments**

path                    Path to the .gmt file.

**Value**

A [`'VennDataset-class'`].

**Examples**

```
tmp <- tempfile(fileext = ".gmt")
writeLines(c("SetA\tdesc\tGENE1\tGENE2\tGENE3",
            "SetB\tdesc\tGENE2\tGENE3\tGENE4"), tmp)
ds <- load_gmt(tmp)
ds@set_names
```

---

load\_gmx

*Load a GMX file (transposed GMT) into a [`'VennDataset-class'`]*

---

**Description**

Row 0 = set names, row 1 = descriptions, rows 2+ = items column-aligned.

**Usage**

```
load_gmx(path)
```

**Arguments**

path                    Path to the .gmx file.

**Value**

A [`'VennDataset-class'`].

**Examples**

```
tmp <- tempfile(fileext = ".gmx")
writeLines(c("SetA\tSetB",
            "desc_A\tdesc_B",
            "GENE1\tGENE2",
            "GENE2\tGENE3"), tmp)
ds <- load_gmx(tmp)
length(ds@items)
```

---

load_sample	<i>Load a bundled sample dataset by name</i>
-------------	--

---

### Description

Sample datasets ship inside the package under ‘inst/extdata/samples/’ and cover biological (cancer drivers, MSigDB pathways) and mock (streaming platforms, gene sets) use cases. Use [list\_samples()] to enumerate.

### Usage

```
load_sample(name)
```

### Arguments

name                    Sample identifier from [list\_samples()].

### Value

A [‘VennDataset-class’] with the appropriate format and mode applied.

### Examples

```
ds <- load_sample("dataset_mock_gene_sets")
length(ds@set_names)

ds <- load_sample("dataset_real_cancer_drivers_4")
analyze(ds)@model
```

---

load_tsv	<i>Load a tab-separated file into a [‘VennDataset-class’]</i>
----------	---

---

### Description

Equivalent to ‘load\_csv(path, binary = binary, delimiter = "\t", prefix\_cols = prefix\_cols)’.

### Usage

```
load_tsv(path, binary = TRUE, prefix_cols = 1L)
```

### Arguments

path                    Path to the file.  
binary                   ‘TRUE’ for binary 0/1 mode (default), ‘FALSE’ for aggregated.  
prefix\_cols            Number of leading metadata columns in binary mode (default 1). Ignored when ‘binary = FALSE’.

**Value**

A [`'VennDataset-class'`].

**Examples**

```
tmp <- tempfile(fileext = ".tsv")
writeLines(c("Gene\tSetA\tSetB", "G1\t1\t0", "G2\t1\t1", "G3\t0\t1"), tmp)
ds <- load_tsv(tmp, binary = TRUE)
ds@universe_size
```

---

overlap\_coefficient     *Szymkiewicz-Simpson overlap coefficient*

---

**Description**

Computes  $|A \cap B| / \min(|A|, |B|)$ . Useful when one set is much smaller than the other.

**Usage**

```
overlap_coefficient(size_a, size_b, intersection)
```

**Arguments**

`size_a`            Inclusive size of set A (integer  $\geq 0$ ).

`size_b`            Inclusive size of set B (integer  $\geq 0$ ).

`intersection`    Inclusive intersection size  $|A \cap B|$ .

**Value**

Numeric in  $[0, 1]$ .

**Examples**

```
overlap_coefficient(10, 5, 3)
```

---

 parse\_region\_expression

*Parse a Boolean region expression into bitmasks*


---

## Description

Translates an expression like "A & B + !C" into a sorted integer vector of region bitmasks (each  $1..2^{n\_sets - 1}$ ). The output is intended for composition with [render\_venn\_svg()] ('highlight = ...') and with the region-accessor family in user code.

## Usage

```
parse_region_expression(expr, n_sets)
```

## Arguments

expr	Character scalar; a Boolean expression in the grammar above.
n_sets	Integer; number of sets in the diagram (2..9).

## Details

Grammar:

- 'A', 'B', ..., 'I' — set atoms (uppercase ASCII).
- '&' — intersection.
- '|' or '+' — union.
- '~' or '!' — complement (unary, against  $1..2^{n\_sets - 1}$ ).
- '(', ')' — grouping.

Precedence (highest first): '~ !', '&', '| +'. Binary operators are left-associative.

## Value

Sorted integer vector of region bitmasks. Empty integer vector when the expression is satisfiable by no region (e.g. "A & ~A").

## Examples

```
parse_region_expression("A & B", n_sets = 3L)      # c(3L, 7L)
parse_region_expression("A + B", n_sets = 3L)     # c(1, 2, 3, 5, 6, 7)
parse_region_expression("A & ~B", n_sets = 3L)   # c(1L, 5L)
parse_region_expression("(A | B) & C", n_sets = 3L) # c(5L, 6L, 7L)
```

---

RegionData-class      *RegionData: one region of a Venn diagram*

---

### Description

Returned as elements of 'RegionResult@regions'. Bitmask convention: bit 'i' set means "in set with index 'i' in 'dataset@set\_names'".

### Slots

bitmask Region bitmask (1 to  $2^n - 1$ ).

label Human-readable label like "AB" or "ABC".

set\_indices Integer vector of 0-based set indices in this region.

set\_names Names of the sets in this region.

exclusive\_items Items present in exactly these sets.

inclusive\_items Items present in at least these sets.

---

RegionResult-class      *RegionResult: result of analyze()*

---

### Description

Bundles the input dataset, chosen model, region map, set sizes, and a lazy ['StatisticsResult-class'] accessible via 'statistics(result)'.

### Slots

dataset The input ['VennDataset-class'].

model Resolved model name (e.g. "venn-4-set" or "proportional").

regions Named list keyed by 'as.character(bitmask)', each value a ['RegionData-class']. Only non-empty regions are stored (sparse for high set counts with few overlaps).

set\_sizes Named integer vector: set name -> inclusive size.

is\_approximate 'TRUE' for the proportional 3-set layout where exact areas can't be achieved with circles.

---

render\_cluster\_heatmap

*Render a cluster-ordered Jaccard similarity heatmap*


---

### Description

Mirrors the webtool's cluster-aware buildEnrichmentHeatmapSvg path. Distance  $D = 1 - \text{Jaccard}$  is fed to [cluster\_set\_order()]; the resulting leaf\_order permutes both axes, and merge heights drive the L-shaped overlays emitted under <g class="hm-dendro-col"> (top band) and <g class="hm-dendro-row"> (left band).

### Usage

```
render_cluster_heatmap(
  result,
  linkage = c("average", "complete", "single"),
  show_row_dendrogram = TRUE,
  show_col_dendrogram = TRUE,
  dendrogram_fraction = 0.12,
  style = NULL
)
```

### Arguments

result	A ['RegionResult-class'] from [analyze()].
linkage	Linkage method passed to [cluster_set_order()]: one of "average" (UPGMA, default), "complete", "single".
show_row_dendrogram	Logical, default TRUE. When FALSE, the row band is omitted from layout entirely (no reserved gap).
show_col_dendrogram	Logical, default TRUE. Same semantics for the column band.
dendrogram_fraction	Fraction of the grid extent reserved per dendrogram band (default 0.12, minimum effective height 20 pixels). Matches the webtool's dendrogramFraction option.
style	Optional named list of style overrides (reserved for v2.3+; currently ignored).

### Details

Pure string construction – no **xml2**.

### Value

An ['SvgImage-class'] with 'content', 'width', 'height' set from the computed layout extents.

**Examples**

```
ds <- load_sample("dataset_real_cancer_drivers_4")
res <- analyze(ds)
img <- render_cluster_heatmap(res, linkage = "average")
nchar(slot(img, "content")) > 0
```

---

render\_enrichment\_bar *Render the pairwise enrichment bar chart*

---

**Description**

One bar per pairwise statistic, ordered by `(set_a_index, set_b_index)`. Bar height is proportional to the chosen `metric`:

- `"neglog10fdr"` (default):  $-\log_{10}(\text{BH-FDR})$ , floor  $1e-300$ .
- `"foldEnrichment"`:  $(k \cdot N) / (K \cdot n)$  from `[StatisticsResult-class]`'s `fold_enrichment` slot.

**Usage**

```
render_enrichment_bar(
  result,
  metric = c("neglog10fdr", "foldEnrichment"),
  width = 560L,
  height = 240L
)
```

**Arguments**

<code>result</code>	A <code>[RegionResult-class]</code> from <code>[analyze()]</code> .
<code>metric</code>	Bar-height metric – <code>"neglog10fdr"</code> or <code>"foldEnrichment"</code> .
<code>width, height</code>	Output SVG dimensions in pixels.

**Details**

Bars use `#2e7d32` for significant pairs ( $\text{FDR} < 0.05$ ) and `#888888` otherwise. Significance markers `****` ( $< 0.001$ ), `***` ( $< 0.01$ ), and `**` ( $< 0.05$ ) appear above each bar.

Pure string construction – no **xml2**.

**Value**

An `[SvgImage-class]` with `content`, `width`, `height`.

## Examples

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
res <- analyze(ds)
img <- render_enrichment_bar(res)
nchar(slot(img, "content")) > 0
```

---

render\_enrichment\_lollipop

*Render the pairwise enrichment lollipop chart*

---

## Description

Same data and significance scheme as [render\_enrichment\_bar()] but as a stem-and-dot plot: a vertical line rises from the baseline to the metric value, capped by a filled circle whose radius scales with  $\sqrt{\text{intersection} / \text{max\_intersection}}$  (range 2.5–8 px).

## Usage

```
render_enrichment_lollipop(
  result,
  metric = c("neglog10fdr", "foldEnrichment"),
  width = 560L,
  height = 240L
)
```

## Arguments

result	A [ <code>'RegionResult-class'</code> ] from [analyze()].
metric	Stem-height metric – <code>"neglog10fdr"</code> or <code>"foldEnrichment"</code> .
width, height	Output SVG dimensions in pixels.

## Details

Pure string construction – no **xml2**.

## Value

An [`'SvgImage-class'`] with `'content'`, `'width'`, `'height'`.

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
res <- analyze(ds)
img <- render_enrichment_lollipop(res)
nchar(slot(img, "content")) > 0
```

render\_network

*Render a force-directed network plot from a RegionResult***Description**

Builds a ggraph plot where nodes are sets (sized by inclusive cardinality) and edges are pairwise overlaps (thickness proportional to the chosen metric; blue for FDR-significant edges below ‘significance\_threshold’, grey otherwise). Layout uses the deterministic ‘stress’ algorithm from graphlayouts.

**Usage**

```
render_network(
  result,
  edge_metric = "intersection",
  seed = 42L,
  significance_threshold = 0.05,
  node_color_map = NULL
)
```

**Arguments**

result	A [‘RegionResult-class’].
edge_metric	One of “intersection”, “jaccard”, “fold_enrichment” (capped at 20.0), “overlap_coefficient”.
seed	Retained for API compatibility; currently unused. The ‘stress’ layout algorithm is fully deterministic and does not rely on a random seed.
significance_threshold	FDR p_adjusted threshold below which edges are colored as significant (default 0.05).
node_color_map	Optional named character vector mapping letters (“A”, “B”, ...) to fill hex colors. Unspecified letters default to yellow (“#FFF200”).

**Details**

Idiomatic R port of Python ‘render\_network’ – same parameter contract, but renders via ggraph + tidygraph instead of networkx + matplotlib.

**Value**

A ‘ggplot’ (ggraph subclass).

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
p <- render_network(result)
inherits(p, "ggplot")

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
p <- render_network(result, edge_metric = "jaccard")
ggplot2::ggsave(tempfile(fileext = ".png"), p, width = 7, height = 7)
```

---

render\_share\_distribution

*Render the Item Share Distribution histogram*

---

**Description**

Vertical bar chart with N bars (k = 1..N), tier-gradient fill from #ffe4b5 (k=1) to #7e14ff (k=N). Mirrors the webtool’s buildShareDistributionSvg layout (480x280 viewBox, Tahoma typography, sd-bar CSS class on every rect) so downstream CSS, PDF embed, and cross-package parity assertions can key on the same structure.

**Usage**

```
render_share_distribution(dataset, style = NULL)
```

**Arguments**

dataset	A [‘VennDataset-class’].
style	Optional named list of style overrides (reserved for v2.3+; currently ignored).

**Details**

Pure string construction – no **xml2** – so the output is byte-stable.

**Value**

An [‘SvgImage-class’] with ‘content’, ‘width = 480’, ‘height = 280’.

**Examples**

```
ds <- load_sample("dataset_real_cancer_drivers_4")
img <- render_share_distribution(ds)
nchar(slot(img, "content")) > 0
```

render\_upset

*Render an UpSet plot from a RegionResult***Description**

Builds a ComplexUpset ggplot showing intersection sizes (top bars), set membership matrix (middle dot grid), and per-set sizes (left bars). Idiomatic R port of Python ‘render\_upset’ – same parameter contract, but renders via ComplexUpset (ggplot2) instead of matplotlib (not a 1:1 port).

**Usage**

```
render_upset(
  result,
  max_columns = 20L,
  sort_by = c("size", "degree"),
  threshold = 0L,
  color_mode = c("depth", "heatmap", "custom"),
  colors = NULL
)
```

**Arguments**

result	A [ <code>'RegionResult-class'</code> ].
max_columns	Maximum number of intersections to display (default 20). Top-N by the active sort.
sort_by	<code>"size"</code> (default – descending) or <code>"degree"</code> (membership count ascending then alphabetical).
threshold	Exclude intersections with size strictly below this value (default 0L = no filter).
color_mode	<code>"depth"</code> (default – viridis on degree), <code>"heatmap"</code> (Reds on size), or <code>"custom"</code> (use the <code>'colors'</code> mapping).
colors	Named character vector mapping intersection LABELS (e.g. <code>"AB"</code> ) to fill hex colors when <code>'color_mode = "custom"</code> . Unspecified labels fall back to <code>"#ccc-ccc"</code> .

**Value**

A ‘ggplot’ object (saveable via `'ggplot2::ggsave()'`).

**Examples**

```

ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
if (getRversion() >= "4.6") {
  p <- render_upset(result)
  inherits(p, "ggplot")
}

if (getRversion() >= "4.6") {
  result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
  p <- render_upset(result, sort_by = "degree", color_mode = "heatmap")
  ggplot2::ggsave(tempfile(fileext = ".png"), p, width = 8, height = 5)
}

```

---

render_venn_svg	<i>Render a RegionResult onto its model SVG and return the raw SVG string</i>
-----------------	---

---

**Description**

Loads the bundled SVG template for ‘result@model’ (or the explicit ‘model’ override), walks the DOM via xml2 to overwrite text content (‘Name\*’, ‘Count\_\*’, ‘CountSUM\_\*’, ‘Title’) and inline ‘fill:’ colors (‘Shape\*’, ‘Shape\*2’ for Euler extras, ‘Bullet\*’), and serializes back to a string.

**Usage**

```

render_venn_svg(
  result,
  model = NULL,
  set_names = NULL,
  colors = NULL,
  title = NULL,
  show_names = TRUE,
  show_counts = TRUE,
  show_items = FALSE,
  item_options = NULL,
  highlight = NULL
)

```

**Arguments**

result	A [‘RegionResult-class’].
model	Optional model id override (filename stem). Default = ‘result@model’.

set_names	Optional named character vector mapping letters ("A", "B", ...) to display names. Unspecified letters fall back to 'result@dataset@set_names'.
colors	Optional named character vector mapping letters to fill hex colors. Applies to 'BulletX', 'ShapeX', and 'ShapeX2' (Euler extra shapes).
title	Optional title override. If 'NULL', the template's default title text is preserved.
show_names	If 'FALSE', blanks every 'NameA-I' element.
show_counts	If 'FALSE', blanks every 'Count_*' and 'CountSUM_*' element.
show_items	If 'TRUE', replace the per-region count text with the actual item identifiers (rendered as '<tspan>' lines inside each 'Count_*' text node). Default 'FALSE'.
item_options	Named list of overrides for the item-text engine. Recognised keys: 'max_items_per_region' (default 20), 'ncol_items' (default 1), 'truncate_long_names' (default 12; 0 disables), 'line_height' (default 10), 'font_size' (default 8), 'show_counts_with_items' (default 'FALSE'), 'ellipsis' (default "..."). Unknown keys raise a warning.
highlight	Character vector of region labels (e.g. 'c("AB", "ABC)") or an integer vector of region bitmasks (e.g. the output of [parse_region_expression()). When set, only the listed regions keep their original fill colour; all other set-shapes are desaturated to '#cccccc' at 25% opacity. Default 'NULL' (no spotlight).

### Details

For 'model = "proportional"', delegates to [generate\_proportional\_svg()].

Mirrors Python 'render\_venn\_svg' byte-for-byte except for: (a) the return type is 'character' instead of an 'SvgImage' wrapper class; (b) xml2 may emit slightly different whitespace/attribute ordering than lxml. Functional content (text, fill colors, structure) is identical.

### Value

A 'character' (length 1) with the raw SVG.

### Examples

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
svg <- render_venn_svg(result)
nchar(svg) > 0

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
svg <- render_venn_svg(result)
nchar(svg) > 0
```

---

`solve_2set`*Area-proportional 2-set circle layout*

---

### Description

Solves for two circles whose areas equal 'a\_only + ab' and 'b\_only + ab' and whose intersection area equals 'ab', by analytical bisection on the inter-center distance. Always exact (returns 'is\_approximate = FALSE').

### Usage

```
solve_2set(a_only, b_only, ab)
```

### Arguments

<code>a_only</code>	Items in A only (integer $\geq 0$ ).
<code>b_only</code>	Items in B only (integer $\geq 0$ ).
<code>ab</code>	Items in A intersection B (integer $\geq 0$ ).

### Details

Mirrors Python 'solve\_2set' byte-for-byte.

### Value

A named list with elements:

**circles** Length-2 list of 'list(cx, cy, r)' named lists.

**error** Relative error of the achieved area fit (typically  $< 1e-4$ ).

**is\_approximate** Always 'FALSE' for 2-set.

### Examples

```
solve_2set(a_only = 30L, b_only = 30L, ab = 10L)
```

---

solve_3set	<i>Area-proportional 3-set circle layout (Wilkinson 2012-style triangulation)</i>
------------	---

---

### Description

Computes pairwise inter-center distances via bisection on the lens intersection area, then places circle C via barycentric triangulation against AB. Always sets 'is\_approximate = TRUE' because perfect 3-circle area-proportional fits don't always exist mathematically.

### Usage

```
solve_3set(regions)
```

### Arguments

regions	Named list keyed by 'as.character(bitmask)' (1..7) -> exclusive count. Missing keys are treated as 0.
---------	---

### Details

Mirrors Python 'solve\_3set' byte-for-byte (including the 'error = NaN' deliberate sentinel - 3-set fit error is not measured in v0.1).

### Value

A named list with elements 'circles' (length 3), 'error' (always NaN in v0.1), 'is\_approximate' (always TRUE).

### Examples

```
solve_3set(list("1" = 100L, "2" = 80L, "3" = 30L,
               "4" = 60L, "5" = 20L, "6" = 15L, "7" = 5L))
```

---

statistics	<i>Lazy pairwise statistics for a RegionResult</i>
------------	--

---

### Description

Computes (and on subsequent calls re-computes) the ['StatisticsResult-class'] for the pairwise metric tables. R has no built-in 'cached\_property' equivalent for S4 slots, so this is recomputed each call. Cache externally via 'stats <- statistics(result)' if you need to access it many times.

**Usage**

```

statistics(result)

## S4 method for signature 'RegionResult'
statistics(result)

```

**Arguments**

result            A [`'RegionResult-class'`].

**Value**

A [`'StatisticsResult-class'`].

**Examples**

```

ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
stats <- statistics(result)
stats@jaccard["A", "B"]

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
stats <- statistics(result)
stats@jaccard
stats@hypergeometric

```

---

StatisticsResult-class

*StatisticsResult: container for pairwise statistical metric tables*

---

**Description**

Returned by [`compute_pairwise()`] and (lazily) by `'statistics()'` on a `'RegionResult'`. Holds five tables:

**Slots**

`jaccard` NxN named matrix of Jaccard indices.  
`dice` NxN named matrix of Sorensen-Dice coefficients.  
`overlap_coefficient` NxN named matrix of Szymkiewicz-Simpson overlap coefficients.  
`fold_enrichment` NxN named matrix of fold-enrichment values.  
`hypergeometric` Long-form data.frame (one row per set pair) with columns: `set_a`, `set_b`, `intersection`, `expected`, `p_value`, `p_adjusted`, `significant`, `highly_significant`.

---

SvgImage-class	<i>SvgImage: wrapper for rendered SVG output</i>
----------------	--

---

**Description**

Returned by [render\_share\_distribution()] and [render\_cluster\_heatmap()]. Mirrors the Python ‘SvgImage’ dataclass so cross-package parity tests can use a uniform attribute name (‘content’). Note that [render\_venn\_svg()] still returns a plain ‘character’ for backward compatibility with the v2.0.x public API; the new plot renderers return ‘SvgImage’ so callers can introspect explicit ‘width’ / ‘height’ extents.

**Slots**

content The SVG document as a string.  
width Pixel width of the rendered SVG.  
height Pixel height of the rendered SVG.

---

tidy.RegionResult	<i>Tidy method for RegionResult (broom-compatible)</i>
-------------------	--

---

**Description**

Returns a long-form table with one row per ordered set pair, combining the five pairwise statistical metrics (Jaccard, Dice, overlap coefficient, fold enrichment, hypergeometric p-value + BH-FDR-adjusted q-value). Pair ordering is ‘(set\_a, set\_b)’ with ‘set\_a’ appearing earlier in ‘result@dataset@set\_names’.

**Usage**

```
## S3 method for class 'RegionResult'
tidy(x, ...)
```

**Arguments**

x A [‘RegionResult-class’].  
... Unused (broom convention).

**Value**

A tibble (or data.frame if ‘tibble’ is not installed) with columns ‘set\_a’, ‘set\_b’, ‘intersection’, ‘expected’, ‘jaccard’, ‘dice’, ‘overlap\_coefficient’, ‘fold\_enrichment’, ‘p\_value’, ‘p\_adjusted’, ‘significant’, ‘highly\_significant’. One row per unordered pair, so ‘n\*(n-1)/2’ rows for an ‘n’-set dataset.

**Examples**

```

ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
if (requireNamespace("broom", quietly = TRUE)) broom::tidy(result)

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
broom::tidy(result)

```

---

to\_excel\_workbook

*Write a 3-sheet Excel workbook matching the webtool's ZIP bundle*


---

**Description**

Sheets:

- Jaccard – NxN matrix of Jaccard indices.
- Sorensen-Dice (the actual sheet title uses the o-with-stroke character) – NxN matrix of Dice coefficients.
- Enrichment – long-form (set\_a, set\_b, intersection, union, expected, fold\_enrichment, p\_value, fdr, significant).

**Usage**

```
to_excel_workbook(result, path)
```

**Arguments**

result	A [ <code>'RegionResult-class'</code> ] from <code>[analyze()]</code> .
path	Output xlsx file path.

**Details**

Mirrors the Python `'to_excel_workbook()'` byte-for-byte in sheet titles, column order, and 4-decimal numeric formatting. Uses **openxlsx** (pure R, no Java dependency).

**Value**

Invisibly returns `'NULL'`.

**Examples**

```

ds <- load_sample("dataset_real_cancer_drivers_4")
res <- analyze(ds)
to_excel_workbook(res, tempfile(fileext = ".xlsx"))

```

---

`to_matrix_tsv`*Write the Item Matrix TSV*

---

## Description

Mirrors the React webapp's "Export Matrix" button + Python's `RegionResult.to_matrix_tsv()` byte-for-byte.

## Usage

```
to_matrix_tsv(result, path)

## S4 method for signature 'RegionResult'
to_matrix_tsv(result, path)
```

## Arguments

<code>result</code>	A [ <code>'RegionResult-class'</code> ].
<code>path</code>	Destination file path.

## Details

Columns: Item, `<SetName1>`, `<SetName2>`, ..., Region. Rows: one per item. Iteration order: mask  $1..(2^n - 1)$ ; within each mask, items in `'dataset@item_order'`.

## Value

Invisibly returns `'path'`.

## Examples

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
to_matrix_tsv(result, tempfile(fileext = ".tsv"))

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
to_matrix_tsv(result, tempfile(fileext = ".tsv"))
```

---

to\_pdf\_report

*Compose a multi-page PDF report from a RegionResult*


---

### Description

Writes a US Letter landscape PDF with overview, venn+upset, statistics tables, and (by default) network and methodology pages. Each page has a footer with package version, generation timestamp, and page number.

### Usage

```
to_pdf_report(
  result,
  path,
  title = NULL,
  include_network = TRUE,
  include_about = TRUE,
  include_share = TRUE,
  include_cluster = FALSE
)
```

### Arguments

result	A [ <code>'RegionResult-class'</code> ].
path	Output PDF file path.
title	Optional title override for the overview page.
include_network	If <code>'TRUE'</code> (default), include the network page.
include_about	If <code>'TRUE'</code> (default), include the methodology page.
include_share	If <code>'TRUE'</code> (default), include the Item Share Distribution page.
include_cluster	If <code>'TRUE'</code> , include the Cluster Heatmap page (default <code>'FALSE'</code> — opt-in like Python's <code>'cluster_heatmap=True'</code> ).

### Value

Invisibly returns `'NULL'`. The PDF is written to `'path'`.

### Examples

```
if (getRversion() >= "4.6") {
  result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
  to_pdf_report(result, tempfile(fileext = ".pdf"))
}
```

---

to\_region\_summary\_tsv *Write the Region Summary TSV*

---

## Description

Mirrors the React webapp's "Export Region Summary" button + Python's `RegionResult.to_region_summary_tsv()` byte-for-byte.

## Usage

```
to_region_summary_tsv(result, path)

## S4 method for signature 'RegionResult'
to_region_summary_tsv(result, path)
```

## Arguments

result	A [ <code>'RegionResult-class'</code> ].
path	Destination file path.

## Details

Columns: Region, Sets, Depth, Exclusive\_Count, Inclusive\_Count, Exclusive\_Pct, Items. Rows: every region ( $1..2^n - 1$ ). Sorted by (Depth ASC, Region label ASC). Items: semicolon-joined, ordered by `'dataset@item_order'`. Cells starting with `'=/+/'-/'@'` (after optional leading whitespace) are escape-prefixed with a single quote.

## Value

Invisibly returns `'path'`.

## Examples

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
to_region_summary_tsv(result, tempfile(fileext = ".tsv"))

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
to_region_summary_tsv(result, tempfile(fileext = ".tsv"))
```

---

to\_statistics\_tsv      *Write the pairwise Statistics TSV*

---

### Description

Mirrors the React webapp's DataSummaryPanel "Export Statistics" button + Python's 'RegionResult.to\_statistics\_tsv()' byte-for-byte.

### Usage

```
to_statistics_tsv(result, path)

## S4 method for signature 'RegionResult'
to_statistics_tsv(result, path)
```

### Arguments

result            A ['RegionResult-class'].  
 path             Destination file path.

### Details

Columns: Set\_A, Set\_B, Name\_A, Name\_B, Size\_A, Size\_B, Intersection, Union, Jaccard, Overlap\_Coeff, Dice, Expected, Fold\_Enrichment, P\_value, FDR, Significant. Float formatting: \* Jaccard / Overlap\_Coeff / Dice: 4 decimals via [.js\_to\_fixed()] \* Expected: 2 decimals \* Fold\_Enrichment: 3 decimals \* P\_value / FDR: scientific (JS toExponential(2)) if '< 0.001', else 6 decimals \* Significant: one of "\*\*\*\*", "\*\*\*", "\*\*", "ns" keyed off FDR thresholds (0.001, 0.01, 0.05).

Rows are sorted by P\_value ascending (matches the underlying StatisticsResult).

### Value

Invisibly returns 'path'.

### Examples

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B"),
  items = list(A = c("x", "y"), B = c("y", "z")),
  item_order = c("x", "y", "z"),
  universe_size = 10L, source_path = NULL, format = "csv")
result <- analyze(ds)
to_statistics_tsv(result, tempfile(fileext = ".tsv"))

result <- analyze(load_sample("dataset_real_cancer_drivers_4"))
to_statistics_tsv(result, tempfile(fileext = ".tsv"))
```

---

to_zip_report	<i>Write a Full Report ZIP bundle</i>
---------------	---------------------------------------

---

### Description

Bundles the multi-page PDF report alongside the supporting SVGs (Venn, UpSet, Network, Share Distribution), TSVs (Region Summary, Item Matrix, Statistics), the 3-sheet xlsx workbook, and a README.txt provenance header into a single ZIP. Mirrors the webtool's \*Download Everything\* button and Python's `vd1 report zip`.

### Usage

```
to_zip_report(result, path, include_share = TRUE, include_cluster = FALSE)
```

### Arguments

result	A ['RegionResult-class'] from [analyze()].
path	Output '.zip' file path.
include_share	Passed through to [to_pdf_report()] (default 'TRUE').
include_cluster	Passed through to [to_pdf_report()] (default 'FALSE').

### Value

Invisibly returns 'NULL'. The ZIP is written to 'path'.

### Examples

```
if (getRversion() >= "4.6") {
  ds <- load_sample("dataset_real_cancer_drivers_4")
  res <- analyze(ds)
  to_zip_report(res, tempfile(fileext = ".zip"))
}
```

---

union_items	<i>Items in the union of the named sets</i>
-------------	---

---

### Description

Returns items that appear in ANY of the sets listed in 'sets'. Equivalent to 'unique(c(items\_in\_A, items\_in\_B, ...))'.

### Usage

```
union_items(result, sets)
```

**Arguments**

`result` A ['RegionResult-class'] from [analyze()].

`sets` Character vector of set letters ("A", "B", ...) or display names (values from 'result@dataset@set\_names'). May mix both.

**Value**

A character vector of item IDs (deduplicated, first-seen order).

**Examples**

```
ds <- methods::new("VennDataset",
  set_names = c("A", "B", "C"),
  items = list(A = c("x", "y"), B = c("y", "z"), C = c("z", "w")),
  item_order = c("x", "y", "z", "w"),
  universe_size = 10L, source_path = NULL, format = "csv")
res <- analyze(ds)
union_items(res, c("A", "B")) # "x", "y", "z"
```

---

vdl\_version

*Get the vennDiagramLab package version*


---

**Description**

Returns the installed version of vennDiagramLab as a character string. This trivial function exists so the Phase 0 skeleton has one public export; Phase 1 introduces the real analyze() / load\_\*() API.

**Usage**

```
vdl_version()
```

**Value**

Character string, the package version (e.g. "2.0.0").

**Examples**

```
vdl_version()
```

---

VennDataset-class	<i>VennDataset: in-memory representation of a Venn-diagram input</i>
-------------------	--

---

**Description**

Returned by the 'load\_\*()' family and consumed by [analyze()]. Holds the deduplicated set members, first-seen item ordering for byte-equivalent TSV output, and source metadata (path, format, optional hypergeometric universe size).

**Slots**

`set_names` Ordered character vector of set identifiers (length 2-9).

`items` Named list ('names(items) == set\_names') of character vectors, each containing the deduplicated members of the corresponding set.

`item_order` First-seen insertion order of all items across all sets, matching JS Set/Map semantics. Used by TSV writers (Phase 2) for byte-equivalent output to the web tool.

`universe_size` Hypergeometric universe N (population size) from the source file, when known. Binary CSV/TSV loaders set this to the row count (matching the web tool's 'csv.rows.length'); other formats leave it 'NULL', signaling "compute as length(item\_order)" downstream.

`source_path` Original file path if loaded from disk; 'NULL' for in-memory datasets.

`format` Source format: one of "csv", "tsv", "gmt", "gmx".

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