

Package ‘synthesisr’

October 14, 2022

Type Package

Title Import, Assemble, and Deduplicate Bibliographic Datasets

Version 0.3.0

Description A critical first step in systematic literature reviews and mining of academic texts is to identify relevant texts from a range of sources, particularly databases such as 'Web of Science' or 'Scopus'. These databases often export in different formats or with different metadata tags. 'synthesisr' expands on the tools outlined by Westgate (2019) [doi:10.1002/jrsm.1374](https://doi.org/10.1002/jrsm.1374) to import bibliographic data from a range of formats (such as 'bibtex', 'ris', or 'ciw') in a standard way, and allows merging and deduplication of the resulting dataset.

Depends R (>= 3.5.0)

Imports stringdist

Suggests knitr, rmarkdown, testthat

Date 2020-05-18

License GPL-3

LazyData true

RoxygenNote 7.1.0

VignetteBuilder knitr

Encoding UTF-8

NeedsCompilation no

Author Martin Westgate [aut, cre] (<https://orcid.org/0000-0003-0854-2034>),
Eliza Grames [aut] (<https://orcid.org/0000-0003-1743-6815>)

Maintainer Martin Westgate <martinjwestgate@gmail.com>

Repository CRAN

Date/Publication 2020-06-03 16:30:03 UTC

R topics documented:

add_line_breaks	2
bibliography-class	3
clean_	4
code_lookup	5
deduplicate	5
detect_	7
extract_unique_references	8
find_duplicates	9
format_citation	11
fuzz_	12
merge_columns	13
override_duplicates	14
parse_	15
read_refs	16
review_duplicates	17
string_	18
synthesisr	19
write_bib	20
Index	22

add_line_breaks	<i>Add line breaks to one or more strings</i>
-----------------	-----------------------------------------------

Description

This function takes a vector of strings and adds line breaks every `n` characters. Primarily built to be called internally by `format_citation`, this function has been made available as it can be useful in other contexts.

Usage

```
add_line_breaks(x, n = 50, max_n = 80, html = FALSE, max_time = 60)
```

Arguments

<code>x</code>	Either a string or a vector; if the vector is not of class <code>character</code> it will be coerced to one using <code>as.character</code> .
<code>n</code>	Numeric: The desired number of characters that should separate consecutive line breaks.
<code>max_n</code>	Numeric: The maximum number of characters that may separate consecutive line breaks.
<code>html</code>	logical: Should the line breaks be specified in <code>html</code> ?
<code>max_time</code>	Numeric: What is the maximum amount of time (in seconds) allowed to adjust groups until character thresholds are reached?

Details

Line breaks are only added between words, so the value of `n` is actually a threshold value rather than being matched exactly. `max_n` is matched exactly if a limit is set and `max_time` is not reached finding new break points between words.

Value

Returns the input vector unaltered except for the addition of line breaks.

Examples

```
add_line_breaks(c("On the Origin of Species"), n = 10)
```

bibliography-class	<i>bibliography-class</i>
--------------------	---------------------------

Description

This is a small number of standard methods for interacting with class 'bibliography'. More may be added later.

Usage

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

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

## S3 method for class 'bibliography'
x[n]

## S3 method for class 'bibliography'
c(...)

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

as.bibliography(x, ...)
```

Arguments

<code>object</code>	An object of class 'bibliography'
<code>...</code>	Any further information
<code>x</code>	An object of class 'bibliography'
<code>n</code>	Number of items to select/print

Details

Methods for class bibliography

clean_	<i>Clean a data.frame or vector</i>
--------	-------------------------------------

Description

Cleans column and author names

Usage

```
clean_df(data)
```

```
clean_authors(x)
```

```
clean_colnames(x)
```

Arguments

data	A data.frame with bibliographic information.
x	A vector of strings

Value

Returns the input, but cleaner.

Examples

```
df <- data.frame(  
  X..title. = c(  
    "EviAtlas: a tool for visualising evidence synthesis databases",  
    "revtools: An R package to support article screening for evidence synthesis",  
    "An automated approach to identifying search terms for systematic reviews",  
    "Reproducible, flexible and high-throughput data extraction from primary literature"),  
  YEAR = c("2019", "2019", "2019", "2019"),  
  authors = c(  
    "Haddaway et al",  
    "Westgate",  
    "EM Grimes AND AN Stillman & MW Tingley and CS Elphick",  
    "Pick et al")  
)  
  
clean_df(df)  
  
# or use sub-functions  
colnames(df) <- clean_colnames(df)  
# colnames(df) <- clean_colnames(colnames(df)) # also works  
df$author <- clean_authors(df$author)
```

code_lookup	<i>Bibliographic code lookup for search results assembly</i>
-------------	--------------------------------------------------------------

Description

A data frame that can be used to look up common codes for different bibliographic fields across databases and merge them to a common format.

Usage

```
code_lookup
```

Format

A data frame with 226 obs of 12 variables

code code used in search results

order the order in which to rank fields in assembled results

category_description type of bibliographic data

entry_description description of field

field bibliographic field that codes correspond to

ris_generic logical: If the code is used in generic ris files

ris_wos logical: If the code is used in Web of Science ris files

ris_pubmed logical: If the code is used in PubMed ris files

ris_scopus logical: If the code is used in Scopus ris files

ris_asp logical: If the code is used in Academic Search Premier ris files

ris_ovid logical: If the code is used in Ovid ris files

ris_synthesisr logical: If the code used in synthesisr imports & exports

deduplicate	<i>Remove duplicates from a bibliographic data set</i>
-------------	--------------------------------------------------------

Description

Removes duplicates using sensible defaults

Usage

```
deduplicate(data, match_by, method, type = "merge", ...)
```

Arguments

<code>data</code>	A <code>data.frame</code> containing bibliographic information.
<code>match_by</code>	Name of the column in <code>data</code> where duplicates should be sought.
<code>method</code>	The duplicate detection function to use; see <code>link{string_}</code> or <code>link{fuzz_}</code> for examples. Passed to <code>find_duplicates</code> .
<code>type</code>	How should entries be selected? Default is "merge" which selected the entries with the largest number of characters in each column. Alternatively "select" returns the row with the highest total number of characters.
<code>...</code>	Arguments passed to <code>find_duplicates</code> .

Details

This is a wrapper function to [find_duplicates](#) and `extract_unique_references`, which tries to choose some sensible defaults. Use with care.

Value

A `data.frame` containing data identified as unique.

See Also

[find_duplicates](#) and [extract_unique_references](#) for underlying functions.

Examples

```
my_df <- data.frame(
  title = c(
    "EviAtlas: a tool for visualising evidence synthesis databases",
    "revtools: An R package to support article screening for evidence synthesis",
    "An automated approach to identifying search terms for systematic reviews",
    "Reproducible, flexible and high-throughput data extraction from primary literature",
    "eviatlas:tool for visualizing evidence synthesis databases.",
    "REVTTOOLS a package to support article-screening for evidence synthesis"
  ),
  year = c("2019", "2019", "2019", "2019", NA, NA),
  authors = c("Haddaway et al", "Westgate",
             "Grames et al", "Pick et al", NA, NA),
  stringsAsFactors = FALSE
)

# run deduplication
dups <- find_duplicates(
  my_df$title,
  method = "string_osa",
  rm_punctuation = TRUE,
  to_lower = TRUE
)

extract_unique_references(my_df, matches = dups)
```

```
# or, in one line:
deduplicate(my_df, "title",
  method = "string_osa",
  rm_punctuation = TRUE,
  to_lower = TRUE)
```

detect_ *Detect file formatting information*

Description

Bibliographic data can be stored in a number of different file types, meaning that detecting consistent attributes of those files is necessary if they are to be parsed accurately. These functions attempt to identify some of those key file attributes. Specifically, `detect_parser` determines which `parse_` function to use; `detect_delimiter` and `detect_lookup` identify different attributes of RIS files; and `detect_year` attempts to fill gaps in publication years from other information stored in a `data.frame`.

Usage

```
detect_parser(x)
```

```
detect_delimiter(x)
```

```
detect_lookup(tags)
```

```
detect_year(df)
```

Arguments

<code>x</code>	A character vector containing bibliographic data
<code>tags</code>	A character vector containing RIS tags.
<code>df</code>	a <code>data.frame</code> containing bibliographic data

Value

`detect_parser` and `detect_delimiter` return a length-1 character; `detect_year` returns a character vector listing estimated publication years; and `detect_lookup` returns a `data.frame`.

Examples

```
revtools <- c(
  "",
  "PMID- 31355546",
  "VI - 10",
  "IP - 4",
  "DP - 2019 Dec",
  "TI - revtools: An R package to support article
```

```

        screening for evidence synthesis.",
    "PG - 606-614",
    "LID - 10.1002/jrsm.1374 [doi]",
    "AU - Westgate MJ",
    "LA - eng",
    "PT - Journal Article",
    "JT - Research Synthesis Methods",
    ""
)

# detect basic attributes of ris files
detect_parser(revtools)
detect_delimiter(revtools)

# determine which tag format to use
tags <- trimws(unlist(lapply(
  strsplit(revtools, "- "),
  function(a){a[1]}
)))
pubmed_tag_list <- detect_lookup(tags[!is.na(tags)])

# find year data in other columns
df <- as.data.frame(parse_pubmed(revtools))
df$year <- detect_year(df)

```

```
extract_unique_references
```

Remove duplicates from a bibliographic data set

Description

Given a list of duplicate entries and a data set, this function extracts only unique references.

Usage

```
extract_unique_references(data, matches, type = "merge")
```

Arguments

data	A data.frame containing bibliographic information.
matches	A vector showing which entries in data are duplicates.
type	How should entries be selected to retain? Default is "merge" which selects the entries with the largest number of characters in each column. Alternatively "select" which returns the row with the highest total number of characters.

Value

Returns a data.frame of unique references.

See Also[find_duplicates](#), [deduplicate](#)**Examples**

```
my_df <- data.frame(
  title = c(
    "EviAtlas: a tool for visualising evidence synthesis databases",
    "revtools: An R package to support article screening for evidence synthesis",
    "An automated approach to identifying search terms for systematic reviews",
    "Reproducible, flexible and high-throughput data extraction from primary literature",
    "eviatlas:tool for visualizing evidence synthesis databases.",
    "REVTTOOLS a package to support article-screening for evidence synthesis"
  ),
  year = c("2019", "2019", "2019", "2019", NA, NA),
  authors = c("Haddaway et al", "Westgate",
             "Grames et al", "Pick et al", NA, NA),
  stringsAsFactors = FALSE
)

# run deduplication
dups <- find_duplicates(
  my_df$title,
  method = "string_osa",
  rm_punctuation = TRUE,
  to_lower = TRUE
)

extract_unique_references(my_df, matches = dups)

# or, in one line:
deduplicate(my_df, "title",
  method = "string_osa",
  rm_punctuation = TRUE,
  to_lower = TRUE)
```

`find_duplicates`*Detect duplicate values*

Description

Identifies duplicate bibliographic entries using different duplicate detection methods.

Usage

```
find_duplicates(
  data,
  method = "exact",
  group_by,
```

```

    threshold,
    to_lower = FALSE,
    rm_punctuation = FALSE
  )

```

Arguments

data	A character vector containing duplicate bibliographic entries.
method	A string indicating how matching should be calculated. Either "exact" for exact matching (the default), or the name of a function for calculating string distance.
group_by	An optional vector, data.frame or list containing data to use as 'grouping' variables; that is, categories within which duplicates should be sought. Defaults to NULL, in which case all entries are compared against all others. Ignored if method = "exact".
threshold	Numeric: the cutoff threshold for deciding if two strings are duplicates. Sensible values depend on the method chosen. Defaults to 5 if method = "string_osa" and must be specified in all other instances except method = "exact" (where no threshold is required).
to_lower	Logical: Should all entries be converted to lower case before calculating string distance? Defaults to FALSE.
rm_punctuation	Logical: Should punctuation should be removed before calculating string distance? Defaults to FALSE.

Value

Returns a vector of duplicate matches, with attributes listing methods used.

See Also

[string_](#) or [fuzz_](#) for suitable functions to pass to methods; [extract_unique_references](#) and [deduplicate](#) for higher-level functions.

Examples

```

my_df <- data.frame(
  title = c(
    "EviAtlas: a tool for visualising evidence synthesis databases",
    "revtools: An R package to support article screening for evidence synthesis",
    "An automated approach to identifying search terms for systematic reviews",
    "Reproducible, flexible and high-throughput data extraction from primary literature",
    "eviatlas:tool for visualizing evidence synthesis databases.",
    "REVT00LS a package to support article-screening for evidence synthesis"
  ),
  year = c("2019", "2019", "2019", "2019", NA, NA),
  authors = c("Haddaway et al", "Westgate",
             "Grames et al", "Pick et al", NA, NA),
  stringsAsFactors = FALSE
)

```

```
# run deduplication
dups <- find_duplicates(
  my_df$title,
  method = "string_osa",
  rm_punctuation = TRUE,
  to_lower = TRUE
)

extract_unique_references(my_df, matches = dups)

# or, in one line:
deduplicate(my_df, "title",
  method = "string_osa",
  rm_punctuation = TRUE,
  to_lower = TRUE)
```

format_citation	<i>Format a citation</i>
-----------------	--------------------------

Description

This function takes an object of class data.frame, list, or bibliography and returns a formatted citation.

Usage

```
format_citation(
  data,
  details = TRUE,
  abstract = FALSE,
  add_html = FALSE,
  line_breaks = FALSE,
  ...
)
```

Arguments

data	An object of class data.frame, list, or or bibliography.
details	Logical: Should identifying information such as author names & journal titles be displayed? Defaults to TRUE.
abstract	Logical: Should the abstract be shown (if available)? Defaults to FALSE.
add_html	Logical: Should the journal title be italicized using html codes? Defaults to FALSE.
line_breaks	Either logical, stating whether line breaks should be added, or numeric stating how many characters should separate consecutive line breaks. Defaults to FALSE.
...	any other arguments.

Value

Returns a string of length equal to `length(data)` that contains formatted citations.

Examples

```
roses <- c("@article{haddaway2018,
  title={ROSES Reporting standards for Systematic Evidence Syntheses:
  pro forma, flow-diagram and descriptive summary of the plan and
  conduct of environmental systematic reviews and systematic maps},
  author={Haddaway, Neal R and Macura, Biljana and Whaley, Paul and Pullin, Andrew S},
  journal={Environmental Evidence},
  volume={7},
  number={1},
  pages={7},
  year={2018},
  publisher={Springer}
}")

tmp <- tempfile()
writeLines(roses, tmp)

citation <- read_ref(tmp)
format_citation(citation)
```

 fuzz_

Calculate similarity between two strings

Description

These functions duplicate the approach of the 'fuzzywuzzy' Python library for calculating string similarity.

Usage

```
fuzzdist(
  a,
  b,
  method = c("fuzz_m_ratio", "fuzz_partial_ratio", "fuzz_token_sort_ratio",
    "fuzz_token_set_ratio")
)

fuzz_m_ratio(a, b)

fuzz_partial_ratio(a, b)

fuzz_token_sort_ratio(a, b)

fuzz_token_set_ratio(a, b)
```

Arguments

a	A character vector of items to match to b.
b	A character vector of items to match to a.
method	The method to use for fuzzy matching.

Value

Returns a score of same length as b, giving the proportional dissimilarity between a and b.

Note

fuzz_m_ratio is a measure of the number of letters that match between two strings. It is calculated as one minus two times the number of matched characters, divided by the number of characters in both strings.

fuzz_partial_ratio calculates the extent to which one string is a subset of the other. If one string is a perfect subset, then this will be zero.

fuzz_token_sort_ratio sorts the words in both strings into alphabetical order, and checks their similarity using fuzz_m_ratio.

fuzz_token_set_ratio is similar to fuzz_token_sort_ratio, but compares both sorted strings to each other, and to a third group made of words common to both strings. It then returns the maximum value of fuzz_m_ratio from these comparisons.

fuzzdist is a wrapper function, for compatibility with stringdist.

Examples

```
fuzzdist("On the Origin of Species",
         "Of the Original Specs",
         method = "fuzz_m_ratio")
```

merge_columns

Bind two or more data.frames with different columns

Description

Takes two or more data.frames with different column names or different column orders and binds them to a single data.frame.

Usage

```
merge_columns(x, y)
```

Arguments

x	Either a data.frame or a list of data.frames.
y	A data.frame, optional if x is a list.

Value

Returns a single data.frame with all the input data frames merged.

Examples

```
df_1 <- data.frame(
  title = c(
    "EviAtlas: a tool for visualising evidence synthesis databases",
    "revtools: An R package to support article screening for evidence synthesis"
  ),
  year = c("2019", "2019")
)

df_2 <- data.frame(
  title = c(
    "An automated approach to identifying search terms for systematic reviews",
    "Reproducible, flexible and high-throughput data extraction from primary literature"
  ),
  authors = c("Grames et al", "Pick et al")
)

merge_columns(df_1, df_2)
```

override_duplicates *Manually override duplicates*

Description

Re-assign group numbers to text that was classified as duplicated but is unique.

Usage

```
override_duplicates(matches, overrides)
```

Arguments

matches	Numeric: a vector of group numbers for texts that indicates duplicates and unique values returned by the find_duplicates function.
overrides	Numeric: a vector of group numbers that are not true duplicates.

Value

The input matches vector with unique group numbers for members of groups that the user overrides.

parse_ *Parse bibliographic text in a variety of formats*

Description

Text in standard formats - such as imported via [readLines](#) - can be parsed using a variety of standard formats. Use [detect_parser](#) to determine which is the most appropriate parser for your situation.

Usage

```
parse_pubmed(x)

parse_ris(x, tag_naming = "best_guess")

parse_bibtex(x)

parse_csv(x)

parse_tsv(x)
```

Arguments

x	A character vector containing bibliographic information in ris format.
tag_naming	What format are ris tags in? Defaults to "best_guess" See read_refs for a list of accepted arguments.

Value

Returns an object of class bibliography (ris, bib, or pubmed formats) or data.frame (csv or tsv).

Examples

```
eviatlas <- c(
  "TY - JOUR",
  "AU - Haddaway, Neal R.",
  "AU - Feierman, Andrew",
  "AU - Grainger, Matthew J.",
  "AU - Gray, Charles T.",
  "AU - Tanriver-Ayder, Ezgi",
  "AU - Dhaubanjari, Sanita",
  "AU - Westgate, Martin J.",
  "PY - 2019",
  "DA - 2019/06/04",
  "TI - EviAtlas: a tool for visualising evidence synthesis databases",
  "JO - Environmental Evidence",
  "SP - 22",
  "VL - 8",
```

```

"IS - 1",
"SN - 2047-2382",
"UR - https://doi.org/10.1186/s13750-019-0167-1",
"DO - 10.1186/s13750-019-0167-1",
"ID - Haddaway2019",
"ER - "
)

detect_parser(eviatlas) # = "parse_ris"
df <- as.data.frame(parse_ris(eviatlas))
ris_out <- write_refs(df, format = "ris", file = FALSE)

```

read_refs

Import bibliographic search results

Description

Imports common bibliographic reference formats (i.e. .bib, .ris, or .txt).

Usage

```

read_refs(
  filename,
  tag_naming = "best_guess",
  return_df = TRUE,
  verbose = FALSE
)

read_ref(
  filename,
  tag_naming = "best_guess",
  return_df = TRUE,
  verbose = FALSE
)

```

Arguments

filename	A path to a filename or vector of filenames containing search results to import.
tag_naming	Either a length-1 character stating how should ris tags be replaced (see details for a list of options), or an object inheriting from class <code>data.frame</code> containing user-defined replacement tags.
return_df	If TRUE (default), returns a <code>data.frame</code> ; if FALSE, returns a list.
verbose	If TRUE, prints status updates (defaults to FALSE).

Details

The default for argument `tag_naming` is "best_guess", which estimates what database has been used for ris tag replacement, then fills any gaps with generic tags. Any tags missing from the database (i.e. `code_lookup`) are passed unchanged. Other options are to use tags from Web of Science ("wos"), Scopus ("scopus"), Ovid ("ovid") or Academic Search Premier ("asp"). If a `data.frame` is given, then it must contain two columns: "code" listing the original tags in the source document, and "field" listing the replacement column/tag names. The `data.frame` may optionally include a third column named "order", which specifies the order of columns in the resulting `data.frame`; otherwise this will be taken as the row order. Finally, passing "none" to `replace_tags` suppresses tag replacement.

Value

Returns a `data.frame` or list of assembled search results.

Functions

- `read_ref`: Import a single file

Examples

```
litsearchr <- c(
  "@article{grames2019,
  title={An automated approach to identifying search terms for
  systematic reviews using keyword co-occurrence networks},
  author={Grames, Eliza M and Stillman, Andrew N and Tingley, Morgan W and Elphick, Chris S},
  journal={Methods in Ecology and Evolution},
  volume={10},
  number={10},
  pages={1645--1654},
  year={2019},
  publisher={Wiley Online Library}
}"
)

tmp <- tempfile()

writeLines(litsearchr, tmp)

df <- read_refs(tmp, return_df = TRUE, verbose = TRUE)
```

review_duplicates

Manually review potential duplicates

Description

Allows users to manually review articles classified as duplicates.

Usage

```
review_duplicates(text, matches)
```

Arguments

`text` A character vector of the text that was used to identify potential duplicates.
`matches` Numeric: a vector of group numbers for texts that indicates duplicates and unique values returned by the `find_duplicates` function.

Value

A data.frame of potential duplicates grouped together.

string_ *Calculate similarity between two strings*

Description

These functions each access a specific "methods" argument provided by `stringdist`, and are provided for convenient calling by `find_duplicates`. They do not include any new functionality beyond that given by `stringdist`, which you should use for your own analyses.

Usage

```
string_osa(a, b)
string_lv(a, b)
string_dl(a, b)
string_hamming(a, b)
string_lcs(a, b)
string_qgram(a, b)
string_cosine(a, b)
string_jaccard(a, b)
string_jw(a, b)
string_soundex(a, b)
```

Arguments

`a` A character vector of items to match to `b`.
`b` A character vector of items to match to `a`.

Value

Returns a score of same length as b, giving the dissimilarity between a and b.

synthesisr	<i>synthesisr: Import, assemble, and deduplicate bibliographic datasets</i>
------------	-----------------------------------------------------------------------------

Description

Systematic review searches include multiple databases that export results in a variety of formats with overlap in coverage between databases. To streamline the process of importing, assembling, and deduplicating results, `synthesisr` recognizes bibliographic files exported from databases commonly used for systematic reviews and merges results into a standardized format.

Import & Export

The key task performed by `synthesisr` is flexible import and presentation of bibliographic data. This is typically achieved by `read_refs`, which can import multiple files at once and link them together into a single `data.frame`. Conversely, export is via `write_refs`. Users that require more detailed control can use the following functions:

- `detect_` Detect file attributes
- `parse_` Parse a vector containing bibliographic data
- `clean_` Cleaning functions for author and column names
- `code_lookup` A dataset of potential ris tags

Data formatting

- `bibliography-class` Methods for class 'bibliography'
- `merge_columns` rbind two `data.frames` with different numbers of columns
- `format_citation` Return a clean citation from a bibliography or `data.frame`
- `add_line_breaks` Set a maximum character width for strings

Deduplication

When importing from multiple databases, it is likely that there will be duplicates in the resulting dataset. The easiest way to deal with this problem in `synthesisr` is using the `deduplicate` command; but this can be risky, particularly if there are no DOIs in the dataset. To get finer control of the deduplication process, consider using the sub-functions:

- `find_duplicates` Locate potentially duplicated references
- `extract_unique_references` Return a `data.frame` with only 'unique' references
- `review_duplicates` Manually review potential duplicates
- `override_duplicates` Manually override identified duplicates
- `fuzz_` Fuzzy string matching c/o 'fuzzywuzzy'
- `string_` Fuzzy string matching c/o stringdist

`write_bib`*Export data to a bibliographic format*

Description

This function exports data.frames containing bibliographic information to either a .ris or .bib file.

Usage

```
write_bib(x)
```

```
write_ris(x, tag_naming = "synthesisr")
```

```
write_refs(x, format = "ris", tag_naming = "synthesisr", file = FALSE)
```

Arguments

<code>x</code>	Either a data.frame containing bibliographic information or an object of class <code>bibliography</code> .
<code>tag_naming</code>	what naming convention should be used to write RIS files? See details for options.
<code>format</code>	What format should the data be exported as? Options are <code>ris</code> or <code>bib</code> .
<code>file</code>	Either logical indicating whether a file should be written (defaulting to <code>FALSE</code>), or a character giving the name of the file to be written.

Value

Returns a character vector containing bibliographic information in the specified format if `file` is `FALSE`, or saves output to a file if `TRUE`.

Functions

- `write_bib`: Format a bib file for export
- `write_ris`: Format a ris file for export

Examples

```
eviatlas <- c(
  "TY - JOUR",
  "AU - Haddaway, Neal R.",
  "AU - Feierman, Andrew",
  "AU - Grainger, Matthew J.",
  "AU - Gray, Charles T.",
  "AU - Tanriver-Ayder, Ezgi",
  "AU - Dhaubanjari, Sanita",
  "AU - Westgate, Martin J.",
  "PY - 2019",
```

```
"DA - 2019/06/04",  
"TI - EviAtlas: a tool for visualising evidence synthesis databases",  
"JO - Environmental Evidence",  
"SP - 22",  
"VL - 8",  
"IS - 1",  
"SN - 2047-2382",  
"UR - https://doi.org/10.1186/s13750-019-0167-1",  
"DO - 10.1186/s13750-019-0167-1",  
"ID - Haddaway2019",  
"ER - "  
)  
  
detect_parser(eviatlas) # = "parse_ris"  
df <- as.data.frame(parse_ris(eviatlas))  
ris_out <- write_refs(df, format = "ris", file = FALSE)
```

Index

- * **datasets**
 - code_lookup, 5
- [.bibliography (bibliography-class), 3
- add_line_breaks, 2, 19
- as.bibliography (bibliography-class), 3
- as.data.frame.bibliography (bibliography-class), 3
- bibliography-class, 3
- c.bibliography (bibliography-class), 3
- c.bibliography, (bibliography-class), 3
- clean_, 4, 19
- clean_authors (clean_), 4
- clean_colnames (clean_), 4
- clean_df (clean_), 4
- code_lookup, 5, 19
- deduplicate, 5, 9, 10, 19
- detect_, 7, 19
- detect_delimiter (detect_), 7
- detect_lookup (detect_), 7
- detect_parser, 15
- detect_parser (detect_), 7
- detect_year (detect_), 7
- extract_unique_references, 6, 8, 10, 19
- find_duplicates, 6, 9, 9, 14, 18, 19
- format_citation, 11, 19
- fuzz_, 10, 12, 19
- fuzz_m_ratio (fuzz_), 12
- fuzz_partial_ratio (fuzz_), 12
- fuzz_token_set_ratio (fuzz_), 12
- fuzz_token_sort_ratio (fuzz_), 12
- fuzzdist (fuzz_), 12
- merge_columns, 13, 19
- override_duplicates, 14, 19
- parse_, 7, 15, 19
- parse_bibtex (parse_), 15
- parse_csv (parse_), 15
- parse_pubmed (parse_), 15
- parse_ris (parse_), 15
- parse_tsv (parse_), 15
- print.bibliography (bibliography-class), 3
- print.bibliography, (bibliography-class), 3
- read_ref (read_refs), 16
- read_refs, 15, 16, 19
- readLines, 15
- review_duplicates, 17, 19
- string_, 10, 18, 19
- string_cosine (string_), 18
- string_dl (string_), 18
- string_hamming (string_), 18
- string_jaccard (string_), 18
- string_jw (string_), 18
- string_lcs (string_), 18
- string_lv (string_), 18
- string_osa (string_), 18
- string_qgram (string_), 18
- string_soundex (string_), 18
- summary.bibliography (bibliography-class), 3
- summary.bibliography, (bibliography-class), 3
- synthesisr, 19
- write_bib, 20
- write_refs, 19
- write_refs (write_bib), 20
- write_ris (write_bib), 20