

Package ‘tidypmc’

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Type Package

Title Parse Full Text XML Documents from PubMed Central

Version 1.7

Description Parse XML documents from the Open Access subset of Europe PubMed Central <<https://europepmc.org>>
including section paragraphs, tables, captions and references.

URL <https://github.com/cstubben/tidypmc>

BugReports <https://github.com/ropensci/tidypmc/issues>

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Encoding UTF-8

VignetteBuilder knitr

Imports xml2, tokenizers, stringr, tibble, dplyr, readr

Suggests europepmc, tidytext, rmarkdown, knitr, testthat, covr

RoxygenNote 6.1.1

NeedsCompilation no

Author Chris Stubben [aut, cre]

Maintainer Chris Stubben <chris.stubben@hci.utah.edu>

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collapse_rows*Collapse a list of PubMed Central tables***Description**

Collapse rows into a semi-colon delimited list with column names and cell values

Usage

```
collapse_rows(pmcs, na.string)
```

Arguments

<code>pmcs</code>	a list of tables, usually from pmc_table
<code>na.string</code>	additional cell values to skip, default is NA and ""

Value

A tibble with table and row number and collapsed text

Author(s)

Chris Stubben

Examples

```
x <- data.frame(
  genes = c("aroB", "glnP", "ndhA", "pyrF"),
  fold_change = c(2.5, 1.7, -3.1, -2.6)
)
collapse_rows(list(`Table 1` = x))
```

<code>pmc_caption</code>	<i>Split captions into sentences</i>
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Description

Split figure, table and supplementary material captions into sentences

Usage

```
pmc_caption(doc)
```

Arguments

doc xml_document from PubMed Central

Value

a tibble with tag, label, sentence number and text

Author(s)

Chris Stubben

Examples

```
# doc <- pmc_xml("PMC2231364") # OR
doc <- xml2::read_xml(system.file("extdata/PMC2231364.xml",
  package = "tidypmc"
))
x <- pmc_caption(doc)
x
dplyr::filter(x, sentence == 1)
```

<code>pmc_metadata</code>	<i>Get article metadata</i>
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Description

Get a list of journal and article metadata in /front tag

Usage

```
pmc_metadata(doc)
```

Arguments

doc xml_document from PubMed Central

Value

a list

Author(s)

Chris Stubben

Examples

```
# doc <- pmc_xml("PMC2231364") # OR
doc <- xml2::read_xml(system.file("extdata/PMC2231364.xml",
  package = "tidypmc"
))
pmc_metadata(doc)
```

pmc_reference	<i>Format references cited</i>
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Description

Format references cited

Usage

```
pmc_reference(doc)
```

Arguments

doc	xml_document from PubMed Central
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Value

a tibble with id, pmid, authors, year, title, journal, volume, pages, and doi.

Note

Mixed citations without any child tags are added to the author column.

Author(s)

Chris Stubben

Examples

```
# doc <- pmc_xml("PMC2231364")
doc <- xml2::read_xml(system.file("extdata/PMC2231364.xml",
  package = "tidypmc"
))
x <- pmc_reference(doc)
x
```

pmc_table	<i>Convert table nodes to tibbles</i>
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Description

Convert PubMed Central table nodes into a list of tibbles

Usage

```
pmc_table(doc)
```

Arguments

doc	xml_document from PubMed Central
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Value

a list of tibbles

Note

Saves the caption and footnotes as attributes and collapses multiline headers, expands all rowspan and colspan attributes and adds subheadings to column one.

Author(s)

Chris Stubben

Examples

```
# doc <- pmc_xml("PMC2231364")
doc <- xml2::read_xml(system.file("extdata/PMC2231364.xml",
  package = "tidypmc"
))
x <- pmc_table(doc)
sapply(x, dim)
x
attributes(x[[1]])
```

pmc_text*Split section paragraphs into sentences***Description**

Split section paragraph tags into a table with subsection titles and sentences using `tokenize_sentences`

Usage

```
pmc_text(doc)
```

Arguments

<code>doc</code>	xml_document from PubMed Central
------------------	----------------------------------

Value

a tibble with section, paragraph and sentence number and text

Note

Subsections may be nested to arbitrary depths and this function will return the entire path to the subsection title as a delimited string like "Results; Predicted functions; Pathogenicity". Tables, figures and formulas that are nested in section paragraphs are removed, superscripted references are replaced with brackets, and any other superscripts or subscripts are separated with ^ and _.

Author(s)

Chris Stubben

Examples

```
# doc <- pmc_xml("PMC2231364")
doc <- xml2::read_xml(system.file("extdata/PMC2231364.xml",
  package = "tidypmc"
))
txt <- pmc_text(doc)
txt
dplyr::count(txt, section, sort = TRUE)
```

`pmc_xml`

Download XML from PubMed Central

Description

Download XML from PubMed Central

Usage

```
pmc_xml(id)
```

Arguments

<code>id</code>	a PMC id starting with 'PMC'
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Value

```
xml_document
```

Source

<https://europepmc.org/RestfulWebService>

Examples

```
## Not run:  
doc <- pmc_xml("PMC2231364")  
  
## End(Not run)
```

`separate_genes`

Separate genes and operons into multiple rows

Description

Separate genes and operons mentioned in full text into multiple rows

Usage

```
separate_genes(txt, pattern = "\\\b[A-Za-z][a-z]{2}[A-Z0-9]+\\\b",  
genes, operon = 6, column = "text")
```

Arguments

txt	a table
pattern	regular expression to match genes, default is to match microbial genes like AbcD, default [A-Za-z][a-z]2[A-Z0-9]+
genes	an optional vector of genes, set pattern to NA to only match this list.
operon	operon length, default 6. Split genes with 6 or more letters into separate genes, for example AbcDEF is split into abcD, abcE and abcF.
column	column name to search, default "text"

Value

a tibble with gene name, matching text and rows.

Note

Check for genes in italics using `xml_text(xml_find_all(doc, "//sec//p//italic"))` and update the pattern or add additional genes as an optional vector if needed

Author(s)

Chris Stubben

Examples

```
x <- data.frame(row = 1, text = "Genes like YacK, hmu and sufABC")
separate_genes(x)
separate_genes(x, genes = "hmu")
```

separate_refs *Separate references cited into multiple rows*

Description

Separates references cited in brackets or parentheses into multiple rows and splits the comma-delimited numeric strings and expands ranges like 7-9 into new rows

Usage

```
separate_refs(txt, column = "text")
```

Arguments

txt	a table
column	column name, default "text"

Value

a tibble

Author(s)

Chris Stubben

Examples

```
x <- data.frame(row = 1, text = "some important studies [7-9,15]")
separate_refs(x)
```

separate_tags

Separate locus tag into multiple rows

Description

Separates locus tags mentioned in full text and expands ranges like YPO1970-74 into new rows

Usage

```
separate_tags(txt, pattern, column = "text")
```

Arguments

txt	a table
pattern	regular expression to match locus tags like YPO[0-9-]+ or the locus tag prefix like YPO.
column	column name to search, default "text"

Value

a tibble with locus tag, matching text and rows.

Author(s)

Chris Stubben

Examples

```
x <- data.frame(row = 1, text = "some genes like YP01002 and YP01970-74")
separate_tags(x, "YPO")
```

separate_text	<i>Separate all matching text into multiple rows</i>
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Description

Separate all matching text into multiple rows

Usage

```
separate_text(txt, pattern, column = "text")
```

Arguments

txt	a tibble, usually results from pmc_text
pattern	either a regular expression or a vector of words to find in text
column	column name, default "text"

Value

a tibble

Note

passed to grep and str_extract_all

Author(s)

Chris Stubben

Examples

```
# doc <- pmc_xml("PMC2231364")
doc <- xml2::read_xml(system.file("extdata/PMC2231364.xml",
                                    package = "tidyxml"))
txt <- pmc_text(doc)
separate_text(txt, "[ATCGN]{5,}")
separate_text(txt, "\\([A-Z]{3,6}s?\\)")
# pattern can be a vector of words
separate_text(txt, c("hmu", "ybt", "yfe", "yfu"))
# wrappers for separate_text with extra step to expand matched ranges
separate.refs(txt)
separate.genes(txt)
separate.tags(txt, "YPO")
```

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