Package ‘quanteda’

August 17, 2021

Version 3.1.0

Title Quantitative Analysis of Textual Data

Description A fast, flexible, and comprehensive framework for quantitative text analysis in R. Provides functionality for corpus management, creating and manipulating tokens and ngrams, exploring keywords in context, forming and manipulating sparse matrices of documents by features and feature co-occurrences, analyzing keywords, computing feature similarities and distances, applying content dictionaries, applying supervised and unsupervised machine learning, visually representing text and text analyses, and more.

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Description

Functions for creating and managing textual corpora, extracting features from textual data, and analyzing those features using quantitative methods.

Details

quanteda makes it easy to manage texts in the form of a corpus, defined as a collection of texts that includes document-level variables specific to each text, as well as meta-data. quanteda includes tools to make it easy and fast to manipulate the texts in a corpus, by performing the most common natural language processing tasks simply and quickly, such as tokenizing, stemming, or forming ngrams. quanteda’s functions for tokenizing texts and forming multiple tokenized documents into a document-feature matrix are both extremely fast and very simple to use. quanteda can segment texts easily by words, paragraphs, sentences, or even user-supplied delimiters and tags.

Built on the text processing functions in the stringi package, which is in turn built on C++ implementation of the ICU libraries for Unicode text handling, quanteda pays special attention to fast and correct implementation of Unicode and the handling of text in any character set.

quanteda is built for efficiency and speed, through its design around three infrastructures: the stringi package for text processing, the Matrix package for sparse matrix objects, and computationally intensive processing (e.g. for tokens) handled in parallelized C++. If you can fit it into
memory, quanteda will handle it quickly. (And eventually, we will make it possible to process objects even larger than available memory.)

quanteda is principally designed to allow users a fast and convenient method to go from a corpus of texts to a selected matrix of documents by features, after defining what the documents and features. The package makes it easy to redefine documents, for instance by splitting them into sentences or paragraphs, or by tags, as well as to group them into larger documents by document variables, or to subset them based on logical conditions or combinations of document variables. The package also implements common NLP feature selection functions, such as removing stopwords and stemming in numerous languages, selecting words found in dictionaries, treating words as equivalent based on a user-defined "thesaurus", and trimming and weighting features based on document frequency, feature frequency, and related measures such as tf-idf.

Tools for working with dictionaries are one of quanteda’s principal strengths, and the package includes several core functions for preparing and applying dictionaries to texts, for example for lexicon-based sentiment analysis.

Once constructed, a quanteda document-feature matrix ("dfm") can be easily analyzed using either quanteda’s built-in tools for scaling document positions, or used with a number of other text analytic tools, such as: topic models (including converters for direct use with the topicmodels, LDA, and stm packages) document scaling (using the quanteda.textmodels package’s functions for the "wordfish" and "Wordscores" models, or direct use with the ca package for correspondence analysis), or machine learning through a variety of other packages that take matrix or matrix-like inputs.

quanteda includes functions for converting its core objects, but especially a dfm, into other formats so that these are easy to use with other analytic packages.

Additional features of quanteda include:

- powerful, flexible tools for working with dictionaries;
- the ability to identify keywords associated with documents or groups of documents;
- the ability to explore texts using key-words-in-context;
- quick computation of word or document similarities, for clustering or to compute distances for other purposes;
- a comprehensive suite of descriptive statistics on text such as the number of sentences, words, characters, or syllables per document; and
- flexible, easy to use graphical tools to portray many of the analyses available in the package.

Source code and additional information

https://github.com/quanteda/quanteda

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as.character.corpus

Description
Coercion functions to and from corpus objects, including conversion to a plain character object; and checks for whether an object is a corpus.

Usage
## S3 method for class 'corpus'
as.character(x, use.names = TRUE, ...)

is.corpus(x)

as.corpus(x)

Arguments
x object to be coerced or checked
use.names logical; preserve (document) names if TRUE
...

Value
as.character() returns the corpus as a plain character vector, with or without named elements.
is.corpus returns TRUE if the object is a corpus.
as.corpus() upgrades a corpus object to the newest format. object.

Note
as.character(x) where x is a corpus is equivalent to calling the deprecated texts(x).
### as.dfm

Coercion and checking functions for dfm objects

**Description**

Convert an eligible input object into a dfm, or check whether an object is a dfm. Current eligible inputs for coercion to a dfm are: matrix, (sparse) Matrix, TermDocumentMatrix and DocumentTermMatrix (from the tm package), data.frame, and other dfm objects.

**Usage**

```r
as.dfm(x)

is.dfm(x)
```

**Arguments**

- `x`  
  a candidate object for checking or coercion to dfm

**Value**

- `as.dfm` converts an input object into a dfm. Row names are used for docnames, and column names for featnames, of the resulting dfm.
- `is.dfm` returns TRUE if and only if its argument is a dfm.

**See Also**

- `as.data.frame.dfm`, `as.matrix.dfm`, `convert()`

### as.dictionary

Coercion and checking functions for dictionary objects

**Description**

Convert a dictionary from a different format into a quanteda dictionary, or check to see if an object is a dictionary.

**Usage**

```r
as.dictionary(x, format = c("tidytext"), separator = " ", tolower = FALSE)

is.dictionary(x)
```
Arguments

x a dictionary-like object to be coerced or checked

format input format for the object to be coerced to a dictionary; current legal values are a data.frame with the fields word and sentiment (as per the tidytext package)

separator the character in between multi-word dictionary values. This defaults to "."

tolower if TRUE, convert all dictionary values to lowercase

Value

as.dictionary returns a quanteda dictionary object. This conversion function differs from the dictionary() constructor function in that it converts an existing object rather than creates one from components or from a file.

is.dictionary returns TRUE if an object is a quanteda dictionary.

Examples

## Not run:
data(sentiments, package = "tidytext")
as.dictionary(subset(sentiments, lexicon == "nrc"))
as.dictionary(subset(sentiments, lexicon == "bing"))
# to convert AFINN into polarities - adjust thresholds if desired
datafinn <- subset(sentiments, lexicon == "AFINN")
datafinn[["sentiment"]]
  with(datafinn,
    sentiment <- ifelse(score < 0, "negative", ifelse(score > 0, "positive", "neutral")))
  )
with(datafinn, table(score, sentiment))
as.dictionary(datafinn)

dat <- data.frame(
  word = c("Great", "Horrible"),
  sentiment = c("positive", "negative")
  )
as.dictionary(dat)
as.dictionary(dat, tolower = FALSE)

## End(Not run)

is.dictionary(dictionary(list(key1 = c("val1", "val2"), key2 = "val3")))
# [1] TRUE
is.dictionary(list(key1 = c("val1", "val2"), key2 = "val3"))
# [1] FALSE
as.fcm

Coercion and checking functions for fcm objects

Description
Convert an eligible input object into a fcm, or check whether an object is a fcm. Current eligible inputs for coercion to a dfm are: matrix, (sparse) Matrix and other fcm objects.

Usage
as.fcm(x)

Arguments
x a candidate object for checking or coercion to dfm

Value
as.fcm converts an input object into a fcm.

as.list.tokens

Coercion, checking, and combining functions for tokens objects

Description
Coercion functions to and from tokens objects, checks for whether an object is a tokens object, and functions to combine tokens objects.

Usage
## S3 method for class 'tokens'
as.list(x, ...)

## S3 method for class 'tokens'
as.character(x, use.names = FALSE, ...)

is.tokens(x)
as.tokens(x, concatenator = "_", ...)

## S3 method for class 'spacyr_parsed'
as.tokens(
x, concatenator = "/", include_pos = c("none", "pos", "tag"), use_lemma = FALSE,


is.tokens(x)

Arguments

x 
object to be coerced or checked

... 
additional arguments used by specific methods. For c.tokens, these are the tokens objects to be concatenated.

use.names 
logical; preserve names if TRUE. For as.character and unlist only.

concatenator 
character between multi-word expressions, default is the underscore character. See Details.

include_pos 
character; whether and which part-of-speech tag to use: "none" do not use any part of speech indicator, "pos" use the pos variable, "tag" use the tag variable. The POS will be added to the token after "concatenator".

use_lemma 
logical; if TRUE, use the lemma rather than the raw token

Details

The concatenator is used to automatically generate dictionary values for multi-word expressions in tokens_lookup() and dfm_lookup(). The underscore character is commonly used to join elements of multi-word expressions (e.g. "piece_of_cake", "New_York"), but other characters (e.g. whitespace " " or a hyphen "-" ) can also be used. In those cases, users have to tell the system what is the concatenator in your tokens so that the conversion knows to treat this character as the inter-word delimiter, when reading in the elements that will become the tokens.

Value

as.list returns a simple list of characters from a tokens object.

as.character returns a character vector from a tokens object.

is.tokens returns TRUE if the object is of class tokens, FALSE otherwise.

as.tokens returns a quanteda tokens object.

is.tokens returns TRUE if the object is of class tokens, FALSE otherwise.

Examples

# create tokens object from list of characters with custom concatenator
dict <- dictionary(list(country = "United States",
                   sea = c("Atlantic Ocean", "Pacific Ocean")))
lis <- list(c("The", "United-States", "has", "the", "Atlantic-Ocean",
            "and", "the", "Pacific-Ocean", "."))
toks <- as.tokens(lis, concatenator = "-")
tokens_lookup(toks, dict)
as.matrix.dfm

Coerce a dfm to a matrix or data.frame

Description

Methods for coercing a dfm object to a matrix or data.frame object.

Usage

```r
## S3 method for class 'dfm'
as.matrix(x, ...)
```

Arguments

- `x` dfm to be coerced
- `...` unused

Examples

```r
# coercion to matrix
as.matrix(data_dfm_lbgexample[, 1:10])
```

as.yaml

Convert quanteda dictionary objects to the YAML format

Description

Converts a quanteda dictionary object constructed by the dictionary function into the YAML format. The YAML files can be edited in text editors and imported into quanteda again.

Usage

```r
as.yaml(x)
```

Arguments

- `x` a dictionary object

Value

`as.yaml` a dictionary in the YAML format, as a character object
Examples

```r
## Not run:
dict <- dictionary(list(one = c("a b", "c*"), two = c("x", "y", "z??"))
cat(yaml <- as.yaml(dict))
cat(yaml, file = (yamlfile <- paste0(tempfile(), ".yml")))
dictionary(file = yamlfile)
## End(Not run)
```

---

**bootstrap_dfm**

**Bootstrap a dfm**

Description

Create an array of resampled dfms.

Usage

`bootstrap_dfm(x, n = 10, ..., verbose = quanteda_options("verbose"))`

Arguments

- `x`  
  a character or corpus object
- `n`  
  number of resamples
- `...`  
  additional arguments passed to `dfm()`
- `verbose`  
  if TRUE print status messages

Details

Function produces multiple, resampled dfm objects, based on resampling sentences (with replacement) from each document, recombining these into new "documents" and computing a dfm for each. Resampling of sentences is done strictly within document, so that every resampled document will contain at least some of its original tokens.

Value

A named list of dfm objects, where the first, dfm_0, is the dfm from the original texts, and subsequent elements are the sentence-resampled dfms.

Author(s)

Kenneth Benoit
char_select

Examples

# bootstrapping from the original text
set.seed(10)
txt <- c(textone = "This is a sentence. Another sentence. Yet another.",
        texttwo = "Premiere phrase. Deuxieme phrase.")
bootstrap_dfm(txt, n = 3, verbose = TRUE)

char_select  Select or remove elements from a character vector

Description

These functions select or discard elements from a character object. For convenience, the functions
char_remove and char_keep are defined as shortcuts for char_select(x, pattern, selection = "remove")
and char_select(x, pattern, selection = "keep"), respectively.

These functions make it easy to change, for instance, stopwords based on pattern matching.

Usage

char_select(
  x,
  pattern,
  selection = c("keep", "remove"),
  valuetype = c("glob", "fixed", "regex"),
  case_insensitive = TRUE
)
char_remove(x, ...)
char_keep(x, ...)

Arguments

x an input character vector
pattern a character vector, list of character vectors, dictionary, or collocations object. See
        pattern for details.
selection whether to "keep" or "remove" the tokens matching pattern
valuetype the type of pattern matching: "glob" for "glob"-style wildcard expressions;
        "regex" for regular expressions; or "fixed" for exact matching. See valu-
        etype for details.

case_insensitive logical; if TRUE, ignore case when matching a pattern or dictionary values

... additional arguments passed by char_remove and char_keep to char_select.

Cannot include selection.
Value

A modified character vector.

Examples

```r
# character selection
mykeywords <- c("natural", "national", "denatured", "other")
char_select(mykeywords, "nat\*", valuetype = "glob")
char_select(mykeywords, "nat", valuetype = "regex")
char_select(mykeywords, c("nature", "other"), selection = "remove")

# character removal
char_remove(letters[1:5], c("a", "c", "x"))
words <- c("any", "and", "Anna", "as", "announce", "but")
char_remove(words, "an\*")
char_remove(words, "an\*", caseInsensitive = FALSE)
char_remove(words, "\^n.\+$", valuetype = "regex")

# remove some of the system stopwords
stopwords("en", source = "snowball")[1:6]
stopwords("en", source = "snowball")[1:6] %>%
  char_remove(c("me", "my\*"))

# character keep
char_keep(letters[1:5], c("a", "c", "x"))
```

Description

`char_tolower()` and `char_toupper()` are replacements for `base::tolower()` and `base::toupper()` based on the `stringi` package. The `stringi` functions for case conversion are superior to the `base` functions because they correctly handle case conversion for Unicode. In addition, the `_tolower()` functions provide an option for preserving acronyms.

Usage

```r
char_tolower(x, keep_acronyms = FALSE)
char_toupper(x)
```

Arguments

- `x` the input object whose character/tokens/feature elements will be case-converted
- `keep_acronyms` logical; if TRUE, do not lowercase any all-uppercase words (applies only to `_tolower()` functions)
Examples

txt1 <- c(txt1 = "b A A", txt2 = "C C a b B")
char_tolower(txt1)
char_toupper(txt1)

# with acronym preservation
txt2 <- c(text1 = "England and France are members of NATO and UNESCO",
         text2 = "NASA sent a rocket into space."
         )
char_tolower(txt2)
char_tolower(txt2, keep_acronyms = TRUE)
char_toupper(txt2)

convert
Convert quanteda objects to non-quanteda formats

Description
Convert a quanteda dfm or corpus object to a format useable by other packages. The general function convert provides easy conversion from a dfm to the document-term representations used in all other text analysis packages for which conversions are defined. For corpus objects, convert provides an easy way to make a corpus and its document variables into a data.frame.

Usage
convert(x, to, ...)

## S3 method for class 'dfm'
convert(
x,
to = c("lda", "tm", "stm", "austin", "topicmodels", "lsa", "matrix", "data.frame", "tripletlist"),
docvars = NULL,
omit_empty = TRUE,
docid_field = "doc_id",
    ...
)

## S3 method for class 'corpus'
convert(x, to = c("data.frame", "json"), pretty = FALSE, ...)

Arguments

x a dfm or corpus to be converted
to target conversion format, one of:
   "lda" a list with components "documents" and "vocab" as needed by the function lda.collapsed.gibbs.sampler from the {ida} package
"tm" a DocumentTermMatrix from the \texttt{tm} package. Note: The \texttt{tm} package version of \texttt{as.TermDocumentMatrix()} allows a weighting argument, which supplies a weighting function for \texttt{TermDocumentMatrix()}. Here the default is for term frequency weighting. If you want a different weighting, apply the weights after converting using one of the \texttt{tm} functions. For other available weighting functions from the \texttt{tm} package, see \texttt{TermDocumentMatrix}.

"stm" the format for the \texttt{stm} package

"austin" the \texttt{wfm} format from the \texttt{austin} package

"topicmodels" the "dm" format as used by the \texttt{topicmodels} package

"lsa" the "textmatrix" format as used by the \texttt{lsa} package

"data.frame" a data.frame of without row.names, in which documents are rows, and each feature is a variable (for a dfm), or each text and its document variables form a row (for a corpus)

"json" (corpus only) convert a corpus and its document variables into JSON format, using the format described in \texttt{jsonlite::toJSON()}

"tripletlist" a named "triplet" format list consisting of document, feature, and frequency

... unused directly

docvars optional data.frame of document variables used as the meta information in conversion to the \texttt{stm} package format. This aids in selecting the document variables only corresponding to the documents with non-zero counts. Only affects the "stm" format.

omit_empty logical; if \texttt{TRUE}, omit empty documents and features from the converted dfm. This is required for some formats (such as STM) that do not accept empty documents. Only used when \texttt{to = "lda"} or \texttt{to = "topicmodels"}. For \texttt{to = "stm"} format, \texttt{omit_empty} is always \texttt{TRUE}'.

docid_field character; the name of the column containing document names used when \texttt{to = "data.frame"}. Unused for other conversions.

pretty adds indentation whitespace to JSON output. Can be \texttt{TRUE/FALSE} or a number specifying the number of spaces to indent. See \texttt{prettify}

Value

A converted object determined by the value of \texttt{to} (see above). See conversion target package documentation for more detailed descriptions of the return formats.

Examples

```r
## convert a dfm
toks <- corpus_subset(data_corpus_inaugural, Year > 1970) %>%
  tokens()
dfmat1 <- dfm(toks)

# austin's wfm format
identical(dim(dfmat1), dim(convert(dfmat1, to = "austin")))
```
# stm package format
stmmat <- convert(dfmat1, to = "stm")
str(stmmat)

# triplet
tripletmat <- convert(dfmat1, to = "tripletlist")
str(tripletmat)

## Not run:
# tm
## Not run:
# tm's DocumentTermMatrix format
tmdfm <- convert(dfmat1, to = "tm")
str(tmdfm)

# topicmodels package format
str(convert(dfmat1, to = "topicmodels"))

# lda package format
str(convert(dfmat1, to = "lda"))

## End(Not run)

## convert a corpus into a data.frame
corp <- corpus(c(d1 = "Text one.", d2 = "Text two."),
   docvars = data.frame(dvar1 = 1:2, dvar2 = c("one", "two"),
                        stringsAsFactors = FALSE))
convert(corp, to = "data.frame")
convert(corp, to = "json")

---

## corpus

### Construct a corpus object

### Description

Creates a corpus object from available sources. The currently available sources are:

- a character vector, consisting of one document per element; if the elements are named, these names will be used as document names.
- a data.frame (or a tibble tbl_df), whose default document id is a variable identified by docid_field; the text of the document is a variable identified by text_field; and other variables are imported as document-level meta-data. This matches the format of data.frames constructed by the the readtext package.
- a kwic object constructed by kwic().
- a tm VCorpus or SimpleCorpus class object, with the fixed metadata fields imported as docvars and corpus-level metadata imported as meta information.
- a corpus object.
Usage

corpus(x, ...)

## S3 method for class 'corpus'
corpus(
  x,
  docnames = quanteda::docnames(x),
  docvars = quanteda::docvars(x),
  meta = quanteda::meta(x),
  ...
)

## S3 method for class 'character'
corpus(
  x,
  docnames = NULL,
  docvars = NULL,
  meta = list(),
  unique_docnames = TRUE,
  ...
)

## S3 method for class 'data.frame'
corpus(
  x,
  docid_field = "doc_id",
  text_field = "text",
  meta = list(),
  unique_docnames = TRUE,
  ...
)

## S3 method for class 'kwic'
corpus(x, split_context = TRUE, extract_keyword = TRUE, meta = list(), ...)

## S3 method for class 'Corpus'
corpus(x, ...)

Arguments

x a valid corpus source object

... not used directly

docnames Names to be assigned to the texts. Defaults to the names of the character vector (if any); doc_id for a data.frame; the document names in a \texttt{tm} corpus; or a vector of user-supplied labels equal in length to the number of documents. If none of these are round, then "text1", "text2", etc. are assigned automatically.

docvars a data.frame of document-level variables associated with each text
a named list that will be added to the corpus as corpus-level, user meta-data. This can later be accessed or updated using `meta()`.

`unique_docnames` logical; if TRUE, enforce strict uniqueness in docnames; otherwise, rename duplicated docnames using an added serial number, and treat them as segments of the same document.

`docid_field` optional column index of a document identifier; defaults to "doc_id", but if this is not found, then will use the rownames of the data.frame; if the rownames are not set, it will use the default sequence based on ([quanteda_options]("base_docname").

`text_field` the character name or numeric index of the source data.frame indicating the variable to be read in as text, which must be a character vector. All other variables in the data.frame will be imported as docvars. This argument is only used for data.frame objects (including those created by `readtext`).

`split_context` logical; if TRUE, split each kwic row into two "documents", one for "pre" and one for "post", with this designation saved in a new docvar context and with the new number of documents therefore being twice the number of rows in the kwic.

`extract_keyword` logical; if TRUE, save the keyword matching pattern as a new docvar keyword.

Details

The texts and document variables of corpus objects can also be accessed using index notation and the $ operator for accessing or assigning docvars. For details, see `[.corpus()`.

Value

A `corpus` class object containing the original texts, document-level variables, document-level metadata, corpus-level metadata, and default settings for subsequent processing of the corpus.

For `quanteda >= 2.0`, this is a specially classed character vector. It has many additional attributes but you should not access these attributes directly, especially if you are another package author. Use the extractor and replacement functions instead, or else your code is not only going to be uglier, but also likely to break should the internal structure of a corpus object change. Using the accessor and replacement functions ensures that future code to manipulate corpus objects will continue to work.

See Also

`corpus`, `docvars()`, `meta()`, `as.character.corpus()`, `ndoc()`, `docnames()`

Examples

# create a corpus from texts
corpus(data_char_ukimmig2010)

# create a corpus from texts and assign meta-data and document variables
summary(corpus(data_char_ukimmig2010,
    docvars = data.frame(party = names(data_char_ukimmig2010)))); 5)
# import a tm VCorpus
if (requireNamespace("tm", quietly = TRUE)) {
  data(crude, package = "tm")  # load in a tm example VCorpus
  vcorp <- corpus(crude)
  summary(vcorp)

  data(acq, package = "tm")
  summary(corpus(acq), 5)

  vcorp2 <- tm::VCorpus(tm::VectorSource(data_char_ukimmig2010))
  corp <- corpus(vcorp2)
  summary(corp)
}

# construct a corpus from a data.frame
dat <- data.frame(letter_factor = factor(rep(letters[1:3], each = 2)),
  some_ints = 1L:6L,
  some_text = paste0("This is text number ", 1:6, ", ", )
  stringsAsFactors = FALSE,
  row.names = paste0("fromDf_", 1:6))

  dat
  summary(corpus(dat, text_field = "some_text",
    meta = list(source = "From a data.frame called mydf.")))

# from a kwic
kw <- kwic(tokens(data_char_sampletext, remove_separators = FALSE),
  pattern = "econom*", separator = "")

summary(corpus(kw))
summary(corpus(kw, split_context = FALSE))
as.character(corpus(kw, split_context = FALSE))

---

corpus_group  

Combine documents in corpus by a grouping variable

Description

Combine documents in a corpus object by a grouping variable, by concatenating their texts in the order of the documents within each grouping variable.

Usage

corpus_group(x, groups = docid(x), fill = FALSE, concatenator = " ")

Arguments

x  corpus object
corpus_reshape

groups grouping variable for sampling, equal in length to the number of documents. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for groups. See news(Version >= "3.0",package = "quanteda") for details.

fill logical; if TRUE and groups is a factor, then use all levels of the factor when forming the new documents of the grouped object. This will result in a new "document" with empty content for levels not observed, but for which an empty document may be needed. If groups is a factor of dates, for instance, then fill = TRUE ensures that the new object will consist of one new "document" by date, regardless of whether any documents previously existed with that date. Has no effect if the groups variable(s) are not factors.

concatenator the concatenation character that will connect the grouped documents.

Value

da corpus object whose documents are equal to the unique group combinations, and whose texts are the concatenations of the texts by group. Document-level variables that have no variation within groups are saved in docvars. Document-level variables that are lists are dropped from grouping, even when these exhibit no variation within groups.

Examples

corp <- corpus(c("a a b", "a b c", "a c d d", "a c c d"),
docvars = data.frame(grp = c("grp1", "grp1", "grp2", "grp2")))
corpus_group(corp, groups = grp)
corpus_group(corp, groups = c(1, 1, 2, 2))
corpus_group(corp, groups = factor(c(1, 1, 2, 2), levels = 1:3))
# with fill
corpus_group(corp, groups = factor(c(1, 1, 2, 2), levels = 1:3), fill = TRUE)

corpus_reshape  Recast the document units of a corpus

Description

For a corpus, reshape (or recast) the documents to a different level of aggregation. Units of aggregation can be defined as documents, paragraphs, or sentences. Because the corpus object records its current "units" status, it is possible to move from recast units back to original units, for example from documents, to sentences, and then back to documents (possibly after modifying the sentences).

Usage

corpus_reshape(
  x,
  to = c("sentences", "paragraphs", "documents"),
  use_docvars = TRUE,
  ...
)
corpus_sample

 Arguments

 x corpus whose document units will be reshaped
 to new document units in which the corpus will be recast
 use_docvars if TRUE, repeat the docvar values for each segmented text; if FALSE, drop the
docvars in the segmented corpus. Dropping the docvars might be useful in order
to conserve space or if these are not desired for the segmented corpus.
...
additional arguments passed to tokens(), since the syntactic segmenter uses
this function)

 Value

 A corpus object with the documents defined as the new units, including document-level meta-data
identifying the original documents.

 Examples

 # simple example
corp1 <- corpus(c(textone = "This is a sentence. Another sentence. Yet another.",
textwo = "Premiere phrase. Deuxieme phrase."),
docvars = data.frame(country=c("UK", "USA"), year=c(1990, 2000)))
summary(corp1)
summary(corpus_reshape(corp1, to = "sentences"))

 # example with inaugural corpus speeches
(corp2 <- corpus_subset(data_corpus_inaugural, Year>2004))
corp2para <- corpus_reshape(corp2, to = "paragraphs")
corp2para
summary(corp2para, 50, showmeta = TRUE)
## Note that Bush 2005 is recorded as a single paragraph because that text
## used a single \n to mark the end of a paragraph.

---

corpus_sample Randomly sample documents from a corpus

 Description

 Take a random sample of documents of the specified size from a corpus, with or without replace-
ment, optionally by grouping variables or with probability weights.

 Usage

 corpus_sample(x, size = ndoc(x), replace = FALSE, prob = NULL, by = NULL)
corpus_sample

Arguments

x a corpus object whose documents will be sampled

size a positive number, the number of documents to select; when used with by, the number to select from each group or a vector equal in length to the number of groups defining the samples to be chosen in each category of by. By defining a size larger than the number of documents, it is possible to oversample when replace = TRUE.

replace if TRUE, sample with replacement

prob a vector of probability weights for obtaining the elements of the vector being sampled. May not be applied when by is used.

by optional grouping variable for sampling. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for by. See news(Version >= "2.9", package = "quanteda") for details.

Value

a corpus object (re)sampled on the documents, containing the document variables for the documents sampled.

Examples

set.seed(123)
# sampling from a corpus
summary(corpus_sample(data_corpus_inaugural, size = 5))
summary(corpus_sample(data_corpus_inaugural, size = 10, replace = TRUE))

# sampling with by
corp <- data_corpus_inaugural
corp$century <- paste(floor(corp$Year / 100) + 1)
corp$century <- paste0(corp$century, ifelse(corp$century < 21, "th", "st"))
corpus_sample(corp, size = 2, by = century) %>% summary()
# needs drop = TRUE to avoid empty interactions
corpus_sample(corp, size = 1, by = interaction(Party, century, drop = TRUE), replace = TRUE) %>% summary()

# sampling sentences by document
corp <- corpus(c(one = "Sentence one. Sentence two. Third sentence.",
two = "First sentence, doc2. Second sentence, doc2.");
docvars = data.frame(var1 = c("a", "a"), var2 = c(1, 2)))
corpus_reshape(corp, to = "sentences") %>%
corpus_sample(replace = TRUE, by = docid(.))

# oversampling
corpus_sample(corp, size = 5, replace = TRUE)
corpus_segment

Segment texts on a pattern match

Description
Segment corpus text(s) or a character vector, splitting on a pattern match. This is useful for breaking the texts into smaller documents based on a regular pattern (such as a speaker identifier in a transcript) or a user-supplied annotation.

Usage
corpus_segment(
  x,
  pattern = "##*",
  valuetype = c("glob", "regex", "fixed"),
  case_insensitive = TRUE,
  extract_pattern = TRUE,
  pattern_position = c("before", "after"),
  use_docvars = TRUE
)

char_segment(
  x,
  pattern = "##*",
  valuetype = c("glob", "regex", "fixed"),
  case_insensitive = TRUE,
  remove_pattern = TRUE,
  pattern_position = c("before", "after")
)

Arguments
- **x**: character or corpus object whose texts will be segmented
- **pattern**: a character vector, list of character vectors, dictionary, or collocations object. See pattern for details.
- **valuetype**: the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.
- **case_insensitive**: logical; if TRUE, ignore case when matching a pattern or dictionary values
- **extract_pattern**: extracts matched patterns from the texts and save in docvars if TRUE
- **pattern_position**: either "before" or "after", depending on whether the pattern precedes the text (as with a user-supplied tag, such as ##INTRO in the examples below) or follows the text (as with punctuation delimiters)
corpus_segment

use.docvars if TRUE, repeat the docvar values for each segmented text; if FALSE, drop the
docvars in the segmented corpus. Dropping the docvars might be useful in order
to conserve space or if these are not desired for the segmented corpus.

remove_pattern removes matched patterns from the texts if TRUE

Details

For segmentation into syntactic units defined by the locale (such as sentences), use corpus_reshape() instead. In cases where more fine-grained segmentation is needed, such as that based on commas or
semi-colons (phrase delimiters within a sentence), corpus_segment() offers greater user control
than corpus_reshape().

Value

corpus_segment returns a corpus of segmented texts
char_segment returns a character vector of segmented texts

Boundaries and segmentation explained

The pattern acts as a boundary delimiter that defines the segmentation points for splitting a text
into new "document" units. Boundaries are always defined as the pattern matches, plus the end and
beginnings of each document. The new "documents" that are created following the segmentation
will then be the texts found between boundaries.

The pattern itself will be saved as a new document variable named pattern. This is most useful
when segmenting a text according to tags such as names in a transcript, section titles, or user-
supplied annotations. If the beginning of the file precedes a pattern match, then the extracted
text will have a NA for the extracted pattern document variable (or when pattern_position =
"after", this will be true for the text split between the last pattern match and the end of the docu-
ment).

To extract syntactically defined sub-document units such as sentences and paragraphs, use corpus_reshape() instead.

Using patterns

One of the most common uses for corpus_segment is to partition a corpus into sub-documents
using tags. The default pattern value is designed for a user-annotated tag that is a term beginning
with double "hash" signs, followed by a whitespace, for instance as #INTRODUCTION The text.
Glob and fixed pattern types use a whitespace character to signal the end of the pattern.

For more advanced pattern matches that could include whitespace or newlines, a regex pattern type
can be used, for instance a text such as
Mr. Smith: Text
Mrs. Jones: More text
could have as pattern = "/[A-Z].+,\s[A-Z][a-z]+: ", which would catch the title, the
name, and the colon.

For custom boundary delimitation using punctuation characters that come come at the end of a
clause or sentence (such as , and., these can be specified manually and pattern_position set
to "after". To keep the punctuation characters in the text (as with sentence segmentation), set
extract_pattern = FALSE. (With most tag applications, users will want to remove the patterns from the text, as they are annotations rather than parts of the text itself.)

See Also
corpus_reshape(), for segmenting texts into pre-defined syntactic units such as sentences, paragraphs, or fixed-length chunks

Examples

## segmenting a corpus

# segmenting a corpus using tags
corp1 <- corpus(c("##INTRO This is the introduction.  
##DOC1 This is the first document. Second sentence in Doc 1.  
##DOC3 Third document starts here. End of third document.",  
"##INTRO Document ##NUMBER Two starts before ##NUMBER Three."")
corpseg1 <- corpus_segment(corp1, pattern = "##*")

# segmenting a corpus based on speaker identifiers
corp2 <- corpus("Mr. Smith: Text.  
Mrs. Jones: More text.  
Mr. Smith: I'm speaking, again.")
corpseg2 <- corpus_segment(corp2, pattern = "\b\[A-Z]+\s\[A-Z][a-z]+:",
valuetype = "regex")

# segmenting a corpus using crude end-of-sentence segmentation
corpseg3 <- corpus_segment(corp1, pattern = ".", valuetype = "fixed",
pattern_position = "after", extract_pattern = FALSE)

# segmenting a character vector

# segment into paragraphs and removing the "- " bullet points
cat(data_char_ukimmig2010[4])
char_segment(data_char_ukimmig2010[4],
    pattern = "\n\n(-\s){0,1}", valuetype = "regex",
    remove_pattern = TRUE)

# segment a text into clauses
txt <- c(d1 = "This, is a sentence? You: come here.", d2 = "Yes, yes okay.")
char_segment(txt, pattern = "\p{P}", valuetype = "regex",
    pattern_position = "after", remove_pattern = FALSE)
**corpus_trim**

**Description**

Removes sentences from a corpus or a character vector shorter than a specified length.

**Usage**

```r
corpus_trim(x, trim = c("sentences", "paragraphs", "documents"),
            min_n = 1,
            max_n = NULL,
            exclude_pattern = NULL)
```

---

**corpus_subset**

**Description**

Returns subsets of a corpus that meet certain conditions, including direct logical operations on doc-vars (document-level variables). `corpus_subset` functions identically to `subset.data.frame()`, using non-standard evaluation to evaluate conditions based on the docvars in the corpus.

**Usage**

```r
corpus_subset(x, subset, drop_docid = TRUE, ...)
```

**Arguments**

- `x`: corpus object to be subsetted
- `subset`: logical expression indicating the documents to keep: missing values are taken as false
- `drop_docid`: if TRUE, docid for documents are removed as the result of subsetting.
- `...`: not used

**Value**

corpus object, with a subset of documents (and docvars) selected according to arguments

**See Also**

`subset.data.frame()`

**Examples**

```r
summary(corpus_subset(data_corpus_inaugural, Year > 1980))
summary(corpus_subset(data_corpus_inaugural, Year > 1930 & President == "Roosevelt"))
```
char_trim(
  x,
  what = c("sentences", "paragraphs", "documents"),
  min_ntoken = 1,
  max_ntoken = NULL,
  exclude_pattern = NULL
)

Arguments

- **x**: corpus or character object whose sentences will be selected.
- **what**: units of trimming, "sentences" or "paragraphs", or "documents"
- **min_ntoken**, **max_ntoken**: minimum and maximum lengths in word tokens (excluding punctuation)
- **exclude_pattern**: a string regular expression whose match (at the sentence level) will be used to exclude sentences

Value

a corpus or character vector equal in length to the input. If the input was a corpus, then all docvars and metadata are preserved. For documents whose sentences have been removed entirely, a null string (""") will be returned.

Examples

txt <- c("PAGE 1. This is a single sentence. Short sentence. Three word sentence.",
  "PAGE 2. Very short! Shorter.",
  "Very long sentence, with multiple parts, separated by commas. PAGE 3.")
corp <- corpus(txt, docvars = data.frame(serial = 1:3))
corp

# exclude sentences shorter than 3 tokens
corp_trim(corp, min_ntoken = 3)
# exclude sentences that start with "PAGE <digit(s)>"
corp_trim(corp, exclude_pattern = "PAGE \d+")

# trimming character objects
char_trim(txt, "sentences", min_ntoken = 3)
char_trim(txt, "sentences", exclude_pattern = "sentence\.")

Description

This is a long paragraph (2,914 characters) of text taken from a debate on Joe Higgins, delivered December 8, 2011.
data_char_ukimmig2010

Usage

data_char_sampletext

Format

character vector with one element

Source


Examples

tokens(data_char_sampletext, remove_punct = TRUE)

data_char_ukimmig2010 Immigration-related sections of 2010 UK party manifestos

Description

Extracts from the election manifestos of 9 UK political parties from 2010, related to immigration or asylum-seekers.

Usage

data_char_ukimmig2010

Format

A named character vector of plain ASCII texts

Examples

data_corpus_ukimmig2010 <-
corpus(data_char_ukimmig2010,
   docvars = data.frame(party = names(data_char_ukimmig2010)))
summary(data_corpus_ukimmig2010, showmeta = TRUE)
Description

US presidential inaugural address texts, and metadata (for the corpus), from 1789 to present.

Usage

data_corpus_inaugural

Format

a corpus object with the following docvars:

- Year a four-digit integer year
- President character; President’s last name
- FirstName character; President’s first name (and possibly middle initial)
- Party factor; name of the President’s political party

Details

data_corpus_inaugural is the quanteda-package corpus object of US presidents’ inaugural addresses since 1789. Document variables contain the year of the address and the last name of the president.

Source


Examples

# some operations on the inaugural corpus
summary(data_corpus_inaugural)
head(docvars(data_corpus_inaugural), 10)
data_dfm_lbgexample

**dfm from data in Table 1 of Laver, Benoit, and Garry (2003)**

**Description**

Constructed example data to demonstrate the Wordscores algorithm, from Laver Benoit and Garry (2003), Table 1.

**Usage**

data_dfm_lbgexample

**Format**

A dfm object with 6 documents and 37 features.

**Details**

This is the example word count data from Laver, Benoit and Garry’s (2003) Table 1. Documents R1 to R5 are assumed to have known positions: -1.5, -0.75, 0, 0.75, 1.5. Document V1 is assumed unknown, and will have a raw text score of approximately -0.45 when computed as per LBG (2003).

**References**


data_dictionary_LSD2015

**Lexicoder Sentiment Dictionary (2015)**

**Description**

The 2015 Lexicoder Sentiment Dictionary in quanteda dictionary format.

**Usage**

data_dictionary_LSD2015
Format

A dictionary of four keys containing glob-style pattern matches.

negative 2,858 word patterns indicating negative sentiment
positive 1,709 word patterns indicating positive sentiment
neg_positive 1,721 word patterns indicating a positive word preceded by a negation (used to convey negative sentiment)
neg_negative 2,860 word patterns indicating a negative word preceded by a negation (used to convey positive sentiment)

Details

The dictionary consists of 2,858 "negative" sentiment words and 1,709 "positive" sentiment words. A further set of 2,860 and 1,721 negations of negative and positive words, respectively, is also included. While many users will find the non-negation sentiment forms of the LSD adequate for sentiment analysis, Young and Soroka (2012) did find a small, but non-negligible increase in performance when accounting for negations. Users wishing to test this or include the negations are encouraged to subtract negated positive words from the count of positive words, and subtract the negated negative words from the negative count.

Young and Soroka (2012) also suggest the use of a pre-processing script to remove specific cases of some words (i.e., "good bye", or "nobody better", which should not be counted as positive). Pre-processing scripts are available at http://www.snsoroka.com/data-lexicoder/.

License and Conditions

The LSD is available for non-commercial academic purposes only. By using data_dictionary_LSD2015, you accept these terms.

Please cite the references below when using the dictionary.

References


Examples

# simple example
txt <- "This aggressive policy will not win friends."

tokens_lookup(tokens(txt), dictionary = data_dictionary_LSD2015, exclusive = FALSE)
## tokens from 1 document.
## text1 :
## [1] "This" "NEGATIVE" "policy" "will" "NEG_POSITIVE" "POSITIVE" "POSITIVE" ","
# notice that double-counting of negated and non-negated terms is avoided
# when using nested_scope = "dictionary"
tokens_lookup(tokens(txt), dictionary = data_dictionary_LSD2015,
  exclusive = FALSE, nested_scope = "dictionary")
## tokens from 1 document.
## text1 :
## [1] "This" "NEGATIVE" "policy" "will" "NEG_POSITIVE" "POSITIVE."

# on larger examples - notice that few negations are used
dfm(data_char_ukimmig2010[1:5], dictionary = data_dictionary_LSD2015)

# compound neg_negative and neg_positive tokens before creating a dfm object
toks <- tokens_compound(tokens(txt), data_dictionary_LSD2015)
dfm_lookup(dfm(toks), data_dictionary_LSD2015)

---

**dfm**

*Create a document-feature matrix*

**Description**

Construct a sparse document-feature matrix, from a character, corpus, tokens, or even other dfm object.

**Usage**

```r
dfm(
  x,
  tolower = TRUE,
  remove_padding = FALSE,
  verbose = quanteda_options("verbose"),
  ...
)
```

**Arguments**

- **x** a tokens or dfm object
- **tolower** convert all features to lowercase
- **remove_padding** logical: if TRUE, remove the "pads" left as empty tokens after calling tokens() or tokens_remove() with padding = TRUE
- **verbose** display messages if TRUE
- **...** not used directly

**Value**

- a dfm object
Changes in version 3

In **quanteda** v3, many convenience functions formerly available in `dfm()` were deprecated. Formerly, `dfm()` could be called directly on a character or corpus object, but we now steer users to tokenise their inputs first using `tokens()`. Other convenience arguments to `dfm()` were also removed, such as `select`, `dictionary`, `thesaurus`, and `groups`. All of these functions are available elsewhere, e.g. through `dfm_group()`. See `news(Version >= "2.9",package = "quanteda")` for details.

See Also

`dfm_select()`, `dfm`

Examples

```r
## for a corpus
toks <- data_corpus_inaugural %>%
  corpus_subset(Year > 1980) %>%
  tokens()
dfm(toks)

# removal options
toks <- tokens(c("a b c", "A B C D")) %>%
  tokens_remove("b", padding = TRUE)
toks
dfm(toks)
dfm(toks, remove = "") # remove "pads"

# preserving case
dfm(toks, tolower = FALSE)
```

---

**dfm_compress**

Recombine a dfm or fcm by combining identical dimension elements

Description

"Compresses" or groups a dfm or fcm whose dimension names are the same, for either documents or features. This may happen, for instance, if features are made equivalent through application of a thesaurus. It could also be needed after a `cbind.dfm()` or `rbind.dfm()` operation. In most cases, you will not need to call `dfm_compress`, since it is called automatically by functions that change the dimensions of the dfm, e.g. `dfm_tolower()`.

Usage

```r
dfm_compress(x, margin = c("both", "documents", "features"))
```

```r
fcm_compress(x)
```
dfm_compress

Arguments

x  input object, a dfm or fcm

margin  character indicating on which margin to compress a dfm, either "documents", "features", or "both" (default). For fcm objects, "documents" has no effect.

Value

dfm_compress returns a dfm whose dimensions have been recombined by summing the cells across identical dimension names (docnames or featnames). The docvars will be preserved for combining by features but not when documents are combined.

fcm_compress returns an fcm whose features have been recombined by combining counts of identical features, summing their counts.

Note

fcm_compress works only when the fcm was created with a document context.

Examples

# dfm_compress examples
dfmt <- rbind(dfm(tokens(c("b A A", "C C a b B")), tolower = FALSE),
               dfm(tokens("A C C C C C"), tolower = FALSE))
colnames(dfmt) <- char_tolower(featnames(dfmt))
dfmt
dfm_compress(dfmt, margin = "documents")
dfm_compress(dfmt, margin = "features")
dfm_compress(dfmt)

# no effect if no compression needed
dfmatsubset <- dfm(tokens(data_corpus_inaugural[1:5]))
dim(dfmatsubset)
dim(dfm_compress(dfmatsubset))

# compress an fcm
fcmat1 <- fcm(tokens("A D a C E a d F e B A C E D"),
              context = "window", window = 3)
## this will produce an error:
# fcm_compress(fcmat1)
x <- c("The fox JUMPED over the dog."
       , "The dog jumped over the fox.")
toks <- tokens(x, remove_punct = TRUE)
fcmat2 <- fcm(toks, context = "document")
colnames(fcmat2) <- rownames(fcmat2) <- tolower(colnames(fcmat2))
colnames(fcmat2)[5] <- rownames(fcmat2)[5] <- "fox"
fcmat2
fcm_compress(fcmat2)
dfm_group

Combine documents in a dfm by a grouping variable

Description

Combine documents in a dfm by a grouping variable, by summing the cell frequencies within group and creating new "documents" with the group labels.

Usage

dfm_group(x, groups = docid(x), fill = FALSE, force = FALSE)

Arguments

x a dfm

groups grouping variable for sampling, equal in length to the number of documents. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for groups. See news(Version >= "3.0", package = "quanteda") for details.

fill logical; if TRUE and groups is a factor, then use all levels of the factor when forming the new documents of the grouped object. This will result in a new "document" with empty content for levels not observed, but for which an empty document may be needed. If groups is a factor of dates, for instance, then fill = TRUE ensures that the new object will consist of one new "document" by date, regardless of whether any documents previously existed with that date. Has no effect if the groups variable(s) are not factors.

force logical; if TRUE, group by summing existing counts, even if the dfm has been weighted. This can result in invalid sums, such as adding log counts (when a dfm has been weighted by "logcount" for instance using dfm_weight()). Not needed when the term weight schemes "count" and "prop".

Value
dfm_group returns a dfm whose documents are equal to the unique group combinations, and whose cell values are the sums of the previous values summed by group. Document-level variables that have no variation within groups are saved in docvars. Document-level variables that are lists are dropped from grouping, even when these exhibit no variation within groups.

Examples
corp <- corpus(c("a a b", "a b c c", "a c d d", "a c c d"),
        docvars = data.frame(grp = c("grp1", "grp1", "grp2", "grp2")))
dfm <- dfm(tokens(corp))
dfm_group(dfm, groups = grp)
dfm_group(dfm, groups = c(1, 1, 2, 2))

# with fill = TRUE
dfm_lookup

Apply a dictionary to a dfm

Description

Apply a dictionary to a dfm by looking up all dfm features for matches in a set of dictionary values, and replace those features with a count of the dictionary’s keys. If exclusive = FALSE then the behaviour is to apply a "thesaurus", where each value match is replaced by the dictionary key, converted to capitals if capkeys = TRUE (so that the replacements are easily distinguished from features that were terms found originally in the document).

Usage

dfm_lookup(
  x,
  dictionary,  
  levels = 1:5,
  exclusive = TRUE,
  valuetype = c("glob", "regex", "fixed"),
  case_insensitive = TRUE,
  capkeys = !exclusive,
  nomatch = NULL,
  verbose = quanteda_options("verbose")
)

Arguments

x the dfm to which the dictionary will be applied
dictionary a dictionary class object
levels levels of entries in a hierarchical dictionary that will be applied
exclusive if TRUE, remove all features not in dictionary, otherwise, replace values in dictionary with keys while leaving other features unaffected
valuetype the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetye for details.
case_insensitive logical; if TRUE, ignore case when matching a pattern or dictionary values
capkeys if TRUE, convert dictionary keys to uppercase to distinguish them from other features
nomatch an optional character naming a new feature that will contain the counts of features of x not matched to a dictionary key. If NULL (default), do not tabulate unmatched features.
verbose print status messages if TRUE
Note

If using dfm_lookup with dictionaries containing multi-word values, matches will only occur if the features themselves are multi-word or formed from ngrams. A better way to match dictionary values that include multi-word patterns is to apply tokens_lookup() to the tokens, and then construct the dfm.

See Also

dfm_replace

Examples

dict <- dictionary(list(christmas = c("Christmas", "Santa", "holiday"),
    opposition = c("Opposition", "reject", "notincorpus"),
    taxglob = "tax*",
    taxregex = "tax.+$",
    country = c("United_States", "Sweden")))
dfmat <- dfm(tokens(c("My Christmas was ruined by your opposition tax plan.",
                          "Does the United_States or Sweden have more progressive taxation?")),
            remove = stopwords("english"))

dfmat

# glob format
dfm_lookup(dfmat, dict, valuetype = "glob")
dfm_lookup(dfmat, dict, valuetype = "glob", case_insensitive = FALSE)

# regex v. glob format: note that "united_states" is a regex match for "tax*"
dfm_lookup(dfmat, dict, valuetype = "glob")
dfm_lookup(dfmat, dict, valuetype = "regex", case_insensitive = TRUE)

# fixed format: no pattern matching
dfm_lookup(dfmat, dict, valuetype = "fixed")
dfm_lookup(dfmat, dict, valuetype = "fixed", case_insensitive = FALSE)

# show unmatched tokens
dfm_lookup(dfmat, dict, nomatch = "_UNMATCHED")

---

**dfm_match**

*Match the feature set of a dfm to given feature names*

**Description**

Match the feature set of a dfm to a specified vector of feature names. For existing features in x for which there is an exact match for an element of features, these will be included. Any features in x not features will be discarded, and any feature names specified in features but not found in x will be added with all zero counts.
dfm_match

Usage

    dfm_match(x, features)

Arguments

x               a dfm
features        character; the feature names to be matched in the output dfm

Details

Selecting on another dfm’s featnames() is useful when you have trained a model on one dfm, and need to project this onto a test set whose features must be identical. It is also used in bootstrap_dfm().

Value

A dfm whose features are identical to those specified in features.

Note

Unlike dfm_select(), this function will add feature names not already present in x. It also provides only fixed, case-sensitive matches. For more flexible feature selection, see dfm_select().

See Also

dfm_select()

dfm_replace

Replace features in dfm

Description

Substitute features based on vectorized one-to-one matching for lemmatization or user-defined stemming.
Usage

dfm_replace(
  x,  
  pattern,  
  replacement,  
  case_insensitive = TRUE,  
  verbose = quanteda_options("verbose")
)

Arguments

  x  
  dfm whose features will be replaced  
  pattern  
  a character vector. See pattern for more details.  
  replacement  
  if pattern is a character vector, then replacement must be character vector of  
  equal length, for a 1:1 match.  
  case_insensitive  
  logical; if TRUE, ignore case when matching a pattern or dictionary values  
  verbose  
  print status messages if TRUE

Examples

dfmat1 <- dfm(data_corpus_inaugural)

# lemmatization
taxwords <- c("tax", "taxing", "taxed", "taxed", "taxation")
lemma <- rep("TAX", length(taxwords))
featnames(dfm_select(dfmat1, pattern = taxwords))
dfmat2 <- dfm_replace(dfmat1, pattern = taxwords, replacement = lemma)
featnames(dfm_select(dfmat2, pattern = taxwords))

# stemming
feat <- featnames(dfmat1)
featstem <- char_wordstem(feat, "porter")
dfmat3 <- dfm_replace(dfmat1, pattern = feat, replacement = featstem, case_insensitive = FALSE)
identical(dfmat3, dfm_wordstem(dfmat1, "porter"))

dfm_sample  
Randomly sample documents from a dfm

Description

  Take a random sample of documents of the specified size from a dfm, with or without replacement,  
  optionally by grouping variables or with probability weights.

Usage

dfm_sample(x, size = NULL, replace = FALSE, prob = NULL, by = NULL)
dfm_select

Select features from a dfm or fcm

**Description**

This function selects or removes features from a dfm or fcm, based on feature name matches with pattern. The most common usages are to eliminate features from a dfm already constructed, such as stopwords, or to select only terms of interest from a dictionary.

**Arguments**

- **x**
  - the dfm object whose documents will be sampled

- **size**
  - a positive number, the number of documents to select; when used with by, the number to select from each group or a vector equal in length to the number of groups defining the samples to be chosen in each category of by. By defining a size larger than the number of documents, it is possible to oversample when replace = TRUE.

- **replace**
  - if TRUE, sample with replacement

- **prob**
  - a vector of probability weights for obtaining the elements of the vector being sampled. May not be applied when by is used.

- **by**
  - optional grouping variable for sampling. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for by. See news(Version >= "2.9", package = "quanteda") for details.

**Value**

A dfm object (re)sampled on the documents, containing the document variables for the documents sampled.

**See Also**

- sample

**Examples**

```r
set.seed(10)
dfmat <- dfm(tokens(c("a b c c d", "a a c c d d d", "a b b c")))
dfmat
dfm_sample(dfmat)
dfm_sample(dfmat, replace = TRUE)

# by groups
dfmat <- dfm(tokens(data_corpus_inaugural[50:58]))
dfm_sample(dfmat, by = Party, size = 2)
```
dfm_select(x, pattern = NULL, selection = c("keep", "remove"),
valuetype = c("glob", "regex", "fixed"),
case_insensitive = TRUE,
min_nchar = NULL,
max_nchar = NULL,
verbose = quanteda_options("verbose")
)

dfm_remove(x, ...)

dfm_keep(x, ...)

fcm_select(x, pattern = NULL, selection = c("keep", "remove"),
valuetype = c("glob", "regex", "fixed"),
case_insensitive = TRUE,
verbose = quanteda_options("verbose"), ...
)

fcm_remove(x, ...)

fcm_keep(x, ...)

Arguments

x the dfm or fcm object whose features will be selected
pattern a character vector, list of character vectors, dictionary, or collocations object. See pattern for details.
selection whether to keep or remove the features
valuetype the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.
case_insensitive logical; if TRUE, ignore case when matching a pattern or dictionary values
min_nchar, max_nchar optional numerics specifying the minimum and maximum length in characters for tokens to be removed or kept; defaults are NULL for no limits. These are applied after (and hence, in addition to) any selection based on pattern matches.
verbose if TRUE print message about how many pattern were removed
dfm_select

... used only for passing arguments from dfm_remove or dfm_keep to dfm_select. Cannot include selection.

Details

dfm_remove and fcm_remove are simply a convenience wrappers to calling dfm_select and fcm_select with selection = "remove".

dfm_keep and fcm_keep are simply a convenience wrappers to calling dfm_select and fcm_select with selection = "keep".

Value

A dfm or fcm object, after the feature selection has been applied.

For compatibility with earlier versions, when pattern is a dfm object and selection = "keep", then this will be equivalent to calling dfm_match(). In this case, the following settings are always used: case_insensitive = FALSE, and valuetype = "fixed". This functionality is deprecated, however, and you should use dfm_match() instead.

Note

This function selects features based on their labels. To select features based on the values of the document-feature matrix, use dfm_trim().

See Also

dfm_match()

Examples

dfmat <- tokens(c("My Christmas was ruined by your opposition tax plan.",
                    "Does the United_States or Sweden have more progressive taxation?")) %>%
  dfm(tolower = FALSE)
dict <- dictionary(list(countries = c("United_States", "Sweden", "France"),
                         wordsEndingInY = c("by", "my"),
                         notintext = "blahblah"))
dfm_select(dfmat, pattern = dict)
dfm_select(dfmat, pattern = dict, case_insensitive = FALSE)
dfm_select(dfmat, pattern = c("s$", ".y"), selection = "keep", valuetype = "regex")
dfm_select(dfmat, pattern = c("s$", ".y"), selection = "remove", valuetype = "regex")
dfm_select(dfmat, pattern = stopwords("english"), selection = "keep", valuetype = "fixed")
dfm_select(dfmat, pattern = stopwords("english"), selection = "remove", valuetype = "fixed")

# select based on character length
dfm_select(dfmat, min_nchar = 5)

dfmat <- dfm(tokens(c("This is a document with lots of stopwords.",
                     "No if, and, or but about it: lots of stopwords.")))

dfmat
dfm_remove(dfmat, stopwords("english"))
toks <- tokens(c("this contains lots of stopwords",
                "no if, and, or but about it: lots"),
dfm_sort

Sort a dfm by frequency of one or more margins

Description

Sorts a dfm by descending frequency of total features, total features in documents, or both.

Usage

dfm_sort(x, decreasing = TRUE, margin = c("features", "documents", "both"))

Arguments

x
Document-feature matrix created by dfm()

decreasing
logical; if TRUE, the sort will be in descending order, otherwise sort in increasing order

margin
which margin to sort on features to sort by frequency of features, documents to sort by total feature counts in documents, and both to sort by both

Value

A sorted dfm matrix object

Author(s)

Ken Benoit

Examples

dfmat <- dfm(data_corpus_inaugural)
head(dfmat)
head(dfm_sort(dfmat))
head(dfm_sort(dfmat, decreasing = FALSE, "both"))
dfm_subset

Extract a subset of a dfm

Description

Returns document subsets of a dfm that meet certain conditions, including direct logical operations on docvars (document-level variables). dfm_subset functions identically to subset.data.frame(), using non-standard evaluation to evaluate conditions based on the docvars in the dfm.

Usage

dfm_subset(x, subset, drop_docid = TRUE, ...)

Arguments

- **x**: dfm object to be subsetted
- **subset**: logical expression indicating the documents to keep: missing values are taken as false
- **drop_docid**: if TRUE, docid for documents are removed as the result of subsetting.
- **...**: not used

Details

To select or subset features, see dfm_select() instead.

When select is a dfm, then the returned dfm will be equal in document dimension and order to the dfm used for selection. This is the document-level version of using dfm_select() where pattern is a dfm: that function matches features, while dfm_subset will match documents.

Value

dfm object, with a subset of documents (and docvars) selected according to arguments

See Also

subset.data.frame()

Examples

corp <- corpus(c(d1 = "a b c d", d2 = "a a b e", d3 = "b b c e", d4 = "e e f a b"),
docvars = data.frame(grp = c(1, 1, 2, 3)))
dfm <- dfm(tokens(corp))
# selecting on a docvars condition
dfm_subset(dfm, grp > 1)
# selecting on a supplied vector
dfm_subset(dfm, c(TRUE, FALSE, TRUE, FALSE))
**dfm_tfidf**

*Weight a dfm by tf-idf*

---

**Description**

Weight a dfm by term frequency-inverse document frequency (*tf-idf*), with full control over options. Uses fully sparse methods for efficiency.

**Usage**

```
dfm_tfidf(
  x,  
  scheme_tf = "count",  
  scheme_df = "inverse",  
  base = 10,  
  force = FALSE,  
  ...  
)
```

**Arguments**

- `x` object for which idf or tf-idf will be computed (a document-feature matrix)
- `scheme_tf` scheme for `dfm_weight()`; defaults to "count"
- `scheme_df` scheme for `docfreq()`; defaults to "inverse".
- `base` the base for the logarithms in the `dfm_weight()` and `docfreq()` calls; default is 10
- `force` logical; if TRUE, apply weighting scheme even if the dfm has been weighted before. This can result in invalid weights, such as as weighting by "prop" after applying "logcount", or after having grouped a dfm using `dfm_group()`.
- `...` additional arguments passed to `docfreq`.

**Details**

`dfm_tfidf` computes term frequency-inverse document frequency weighting. The default is to use counts instead of normalized term frequency (the relative term frequency within document), but this can be overridden using `scheme_tf = "prop"`.

**References**

dfm_tolower

Examples

dfmat1 <- as.dfm(data_dfm_lbgexample)
head(dfmat1[, 5:10])
head(dfm_tfidf(dfmat1)[, 5:10])
docfreq(dfmat1)[5:15]
head(dfm_weight(dfmat1)[, 5:10])

# replication of worked example from
# https://en.wikipedia.org/wiki/Tf-idf#Example_of_tf.80.93idf
dfmat2 <-
  matrix(c(1,1,2,1,0,0, 1,1,0,0,2,3),
        byrow = TRUE, nrow = 2,
        dimnames = list(docs = c("document1", "document2"),
                        features = c("this", "is", "a", "sample",
                                     "another", "example")))

as.dfm()
dfmat2
docfreq(dfmat2)
dfm_tfidf(dfmat2, scheme_tf = "prop")

## Not run:
# comparison with tm
if (requireNamespace("tm")) {
  convert(dfmat2, to = "tm")
  tm::weightTfIdf()
}
## End(Not run)

dfm_tolower

Convert the case of the features of a dfm and combine

Description

dfm_tolower() and dfm_toupper() convert the features of the dfm or fcm to lower and upper case, respectively, and then recombine the counts.

Usage

dfm_tolower(x, keep_acronyms = FALSE)
dfm_toupper(x)

fcm_tolower(x, keep_acronyms = FALSE)
fcm_toupper(x)
Arguments

x the input object whose character/tokens/feature elements will be case-converted

keep_acronyms logical; if TRUE, do not lowercase any all-uppercase words (applies only to *tolower() functions)

Details

fcm_tolower() and fcm_toupper() convert both dimensions of the fcm to lower and upper case, respectively, and then recombine the counts. This works only on fcm objects created with context = "document".

Examples

# for a document-feature matrix
dfm <- dfm(tokens(c("b A A", "C a b B")), tolower = FALSE)
dfm
dfm_tolower(dfm)
dfm_toupper(dfm)

# for a feature co-occurrence matrix
fcm <- fcm(tokens(c("b A A d", "C a b B e")),
            context = "document")
fcm
fcm_tolower(fcm)
fcm_toupper(fcm)

dfm_trim

Trim a dfm using frequency threshold-based feature selection

Description

Returns a document by feature matrix reduced in size based on document and term frequency, usually in terms of a minimum frequency, but may also be in terms of maximum frequencies. Setting a combination of minimum and maximum frequencies will select features based on a range. Feature selection is implemented by considering features across all documents, by summing them for term frequency, or counting the documents in which they occur for document frequency. Rank and quantile versions of these are also implemented, for taking the first n features in terms of descending order of overall global counts or document frequencies, or as a quantile of all frequencies.

Usage

dfm_trim(
  x,
  min_termfreq = NULL,
  max_termfreq = NULL,
  termfreq_type = c("count", "prop", "rank", "quantile"),
  min_docfreq = NULL,
max_docfreq = NULL,
docfreq_type = c("count", "prop", "rank", "quantile"),
sparsity = NULL,
verbose = quanteda_options("verbose"),
...)

Arguments

x a dfm object
min_termfreq, max_termfreq minimum/maximum values of feature frequencies across all documents, below/above which features will be removed
termfreq_type how min_termfreq and max_termfreq are interpreted. "count" sums the frequencies; "prop" divides the term frequencies by the total sum; "rank" is matched against the inverted ranking of features in terms of overall frequency, so that 1, 2, ... are the highest and second highest frequency features, and so on; "quantile" sets the cutoffs according to the quantiles (see quantile()) of term frequencies.
min_docfreq, max_docfreq minimum/maximum values of a feature’s document frequency, below/above which features will be removed
docfreq_type specify how min_docfreq and max_docfreq are interpreted. "count" is the same as [docfreq](x, scheme = "count"); "prop" divides the document frequencies by the total sum; "rank" is matched against the inverted ranking of document frequency, so that 1, 2, ... are the features with the highest and second highest document frequencies, and so on; "quantile" sets the cutoffs according to the quantiles (see quantile()) of document frequencies.
sparsity equivalent to 1 -min_docfreq, included for comparison with tm
verbose print messages
... not used

Value

A dfm reduced in features (with the same number of documents)

Note

Trimming a dfm object is an operation based on the values in the document-feature matrix. To select subsets of a dfm based on the features themselves (meaning the feature labels from featnames()) – such as those matching a regular expression, or removing features matching a stopword list, use dfm_select().

See Also

dfm_select(), dfm_sample()
dfm_weight

Weight the feature frequencies in a dfm

Usage

dfm_weight(
  x,
  scheme = c("count", "prop", "propmax", "logcount", "boolean", "augmented", "logave"),
  weights = NULL,
  base = 10,
  k = 0.5,
  smoothing = 0.5,
  force = FALSE
)
dfm_weight

dfm_smooth(x, smoothing = 1)

Arguments

x  
  document-feature matrix created by dfm

scheme  
  a label of the weight type:
  count  $tf_{ij}$, an integer feature count (default when a dfm is created)
  prop  the proportion of the feature counts of total feature counts (aka relative frequency), calculated as $tf_{ij}/\sum_j tf_{ij}$
  propmax  the proportion of the feature counts of the highest feature count in a document, $tf_{ij}/\max_j tf_{ij}$
  logcount  take the $1 + \log$ of each count, for the given base, or 0 if the count was zero: $1 + \log_{\text{base}}(tf_{ij})$ if $tf_{ij} > 0$, or 0 otherwise.
  boolean  recode all non-zero counts as 1
  augmented  equivalent to $k + (1 - k)\cdot dfm_weight(x, \text{"propmax"})$
  logave  $(1 + \log\text{ of the counts})/(1 + \log\text{ of the average count within document})$, or
  
  $\frac{1 + \log_{\text{base}}(tf_{ij})}{1 + \log_{\text{base}}(\sum_j tf_{ij}/N_i)}$
  logsmooth  log of the counts + smooth, or $tf_{ij} + s$
  weights  if scheme is unused, then weights can be a named numeric vector of weights to be applied to the dfm, where the names of the vector correspond to feature labels of the dfm, and the weights will be applied as multipliers to the existing feature counts for the corresponding named features. Any features not named will be assigned a weight of 1.0 (meaning they will be unchanged).

base  
  base for the logarithm when scheme is "logcount" or logave

k  
  the k for the augmentation when scheme = "augmented"

smoothing  
  constant added to the dfm cells for smoothing, default is 1 for dfm_smooth() and 0.5 for dfm_weight()

force  
  logical; if TRUE, apply weighting scheme even if the dfm has been weighted before. This can result in invalid weights, such as as weighting by "prop" after applying "logcount", or after having grouped a dfm using dfm_group().

Value

dfm_weight returns the dfm with weighted values. Note the because the default weighting scheme is "count", simply calling this function on an unweighted dfm will return the same object. Many users will want the normalized dfm consisting of the proportions of the feature counts within each document, which requires setting scheme = "prop".

dfm_smooth returns a dfm whose values have been smoothed by adding the smoothing amount. Note that this effectively converts a matrix from sparse to dense format, so may exceed memory requirements depending on the size of your input matrix.
dictionary

Create a dictionary

Description

Create a quanteda dictionary class object, either from a list or by importing from a foreign format. Currently supported input file formats are the WordStat, LIWC, Lexicoder v2 and v3, and Yoshikoder formats. The import using the LIWC format works with all currently available dictionary files supplied as part of the LIWC 2001, 2007, and 2015 software (see References).
dictionary

Usage

dictionary(
  x,
  file = NULL,
  format = NULL,
  separator = " ",
  tolower = TRUE,
  encoding = "utf-8"
)

Arguments

x a named list of character vector dictionary entries, including valuertype pattern matches, and including multi-word expressions separated by concatenator. See examples. This argument may be omitted if the dictionary is read from file.

file file identifier for a foreign dictionary

format character identifier for the format of the foreign dictionary. If not supplied, the format is guessed from the dictionary file’s extension. Available options are: "wordstat" format used by Provalis Research’s WordStat software "LIWC" format used by the Linguistic Inquiry and Word Count software "yoshikoder" format used by Yoshikoder software "lexicoder" format used by Lexicoder "YAML" the standard YAML format

separator the character in between multi-word dictionary values. This defaults to " ".

tolower if TRUE, convert all dictionary values to lowercase

encoding additional optional encoding value for reading in imported dictionaries. This uses the iconv labels for encoding. See the "Encoding" section of the help for file.

Details

Dictionaries can be subsetted using [ and [[, operating the same as the equivalent list operators.

Dictionaries can be coerced from lists using as.dictionary(), coerced to named lists of characters using as.list(), and checked using is.dictionary().

Value

A dictionary class object, essentially a specially classed named list of characters.

References


Yoshikoder page, from Will Lowe https://conjugateprior.org/software/yoshikoder/.

See Also
dfm, as.dictionary(), as.list(), is.dictionary()

Examples
corp <- corpus_subset(data_corpus_inaugural, Year>1900)
dict <- dictionary(list(christmas = c("Christmas", "Santa", "holiday"),
opposition = c("Opposition", "reject", "notincorpus"),
taxing = "taxing",
taxation = "taxation",
taxregex = "tax*",
country = "america"))
head(dfm(tokens(corp), dictionary = dict))

# subset a dictionary
dict[1:2]
dict[[c("christmas", "opposition")]]
dict[["opposition"]]

# combine dictionaries
c(dict["christmas"], dict["country"])

## Not run:
# import the Laver-Garry dictionary from Provalis Research
dictfile <- tempfile()
download.file("https://provalisresearch.com/Download/LaverGarry.zip",
dictfile, mode = "wb")
unzip(dictfile, exdir = (td <- tempdir()))
dictlg <- dictionary(file = paste(td, "LaverGarry.cat", sep = "/"))
head(dfm(data_corpus_inaugural, dictionary = dictlg))

## End(Not run)

docfreq

Compute the (weighted) document frequency of a feature

Description

For a dfm object, returns a (weighted) document frequency for each term. The default is a simple count of the number of documents in which a feature occurs more than a given frequency threshold. (The default threshold is zero, meaning that any feature occurring at least once in a document will be counted.)
**Usage**

docfreq(
  x,
  scheme = c("count", "inverse", "inversemax", "inverseprob", "unary"),
  base = 10,
  smoothing = 0,
  k = 0,
  threshold = 0
)

**Arguments**

- **x**: a dfm
- **scheme**: type of document frequency weighting, computed as follows, where \( N \) is defined as the number of documents in the dfm and \( s \) is the smoothing constant:
  - count: \( df_j \), the number of documents for which \( n_{ij} > \text{threshold} \)
  - inverse: \( \log_{\text{base}} \left( s + \frac{N}{k + df_j} \right) \)
  - inversemax: \( \log_{\text{base}} \left( s + \frac{\max(df_j)}{k + df_j} \right) \)
  - inverseprob: \( \log_{\text{base}} \left( \frac{N - df_j}{k + df_j} \right) \)
  - unary: 1 for each feature
- **base**: the base with respect to which logarithms in the inverse document frequency weightings are computed; default is 10 (see Manning, Raghavan, and Schütze 2008, p123).
- **smoothing**: added to the quotient before taking the logarithm
- **k**: added to the denominator in the "inverse" weighting types, to prevent a zero document count for a term
- **threshold**: numeric value of the threshold above which a feature will considered in the computation of document frequency. The default is 0, meaning that a feature’s document frequency will be the number of documents in which it occurs greater than zero times.

**Value**

a numeric vector of document frequencies for each feature
References


Examples

```r
dfmat1 <- dfm(data_corpus_inaugural[1:2])
docfreq(dfmat1[, 1:20])

# replication of worked example from
dfmat2 <-
  matrix(c(1,1,2,1,0,0, 1,1,0,0,2,3),
         byrow = TRUE, nrow = 2,
         dimnames = list(docs = c("document1", "document2"),
                         features = c("this", "is", "a", "sample",
                                     "another", "example")))

dfmat2

docfreq(dfmat2)
docfreq(dfmat2, scheme = "inverse")
docfreq(dfmat2, scheme = "inverse", k = 1, smoothing = 1)
docfreq(dfmat2, scheme = "unary")
docfreq(dfmat2, scheme = "inversemax")
docfreq(dfmat2, scheme = "inverseprob")
```

docnames

*Get or set document names*

**Description**

Get or set the document names of a corpus, tokens, or dfm object.

**Usage**

```r
docnames(x)
docnames(x) <- value
docid(x)
```

**Arguments**

- `x` the object with docnames
- `value` a character vector of the same length as `x`
Value

docnames <- assigns new values to the document names of an object. docnames can only be character, so any non-character value assigned to be a docname will be coerced to mode character.

docid returns an internal variable denoting the original "docname" from which a document came. Unless an object has been reshaped (e.g. \texttt{corpus\_reshape()}, \texttt{split}(e.g.\texttt{tokens\_split()}), or segmented (e.g. \texttt{corpus\_segment()}), \texttt{docid(x)} will return the docnames.

Note

docid is designed primarily for developers, not for end users. In most cases, you will want \texttt{docnames} instead. It is, however, the default for \texttt{groups}, so that documents that have been previously reshaped (e.g. \texttt{corpus\_reshape()}, \texttt{split}(e.g.\texttt{tokens\_split()}), or segmented (e.g. \texttt{corpus\_segment()} will be regrouped into their original \texttt{docnames} when \texttt{groups = docid(x)}.

See Also

\texttt{featnames()}

Examples

\begin{verbatim}
# get and set document names to a corpus
corp <- data_corpus_inaugural
docnames(corp) <- char_tolower(docnames(corp))

# get and set document names to a tokens
toks <- tokens(data_corpus_inaugural)
docnames(toks) <- char_tolower(docnames(toks))

# get and set document names to a dfm
dfmat <- dfm(data_corpus_inaugural[1:5])
docnames(dfmat) <- char_tolower(docnames(dfmat))

# reassign the document names of the inaugural speech corpus
docnames(data_corpus_inaugural) <- paste("Speech", 1:ndoc(data_corpus_inaugural), sep="")

# docid
corp <- corpus(c(textone = "This is a sentence. Another sentence. Yet another. ",
        texttwo = "Sentence 1. Sentence 2. " ))
corpsent <- corp %>%
    corpus_reshape(to = "sentences")
docnames(corpsent)  
docid(corpsent)  
docid(tokens(corpsent))
docid(dfm(tokens(corpsent)))
\end{verbatim}
docvars  Get or set document-level variables

Description

Get or set variables associated with a document in a corpus, tokens or dfm object.

Usage

docvars(x, field = NULL)
docvars(x, field = NULL) <- value

## S3 method for class 'corpus'
x$name

## S3 replacement method for class 'corpus'
x$name <- value

## S3 method for class 'tokens'
x$name

## S3 replacement method for class 'tokens'
x$name <- value

## S3 method for class 'dfm'
x$name

## S3 replacement method for class 'dfm'
x$name <- value

Arguments

x  corpus, tokens, or dfm object whose document-level variables will be read or set
field  string containing the document-level variable name
value  a vector of document variable values to be assigned to name
name  a literal character string specifying a single docvars name

Value

docvars returns a data.frame of the document-level variables, dropping the second dimension to form a vector if a single docvar is returned.
docvars<- assigns value to the named field

Accessing or assigning docvars using the $ operator

As of quanteda v2, it is possible to access and assign a docvar using the $ operator. See Examples.
Note

Reassigning document variables for a tokens or dfm object is allowed, but discouraged. A better, more reproducible workflow is to create your docvars as desired in the corpus, and let these continue to be attached "downstream" after tokenization and forming a document-feature matrix. Recognizing that in some cases, you may need to modify or add document variables to downstream objects, the assignment operator is defined for tokens or dfm objects as well. Use with caution.

Examples

# retrieving docvars from a corpus
head(docvars(data_corpus_inaugural))
tail(docvars(data_corpus_inaugural, "President"), 10)
head(data_corpus_inaugural$President)

# assigning document variables to a corpus
corp <- data_corpus_inaugural
docvars(corp, "President") <- paste("prez", 1:ndoc(corp), sep = "")
head(docvars(corp))
corp$fullname <- paste(data_corpus_inaugural$FirstName,
                       data_corpus_inaugural$President)
tail(corp$fullname)

# accessing or assigning docvars for a corpus using "$"
data_corpus_inaugural$Year
data_corpus_inaugural$century <- floor(data_corpus_inaugural$Year / 100)
data_corpus_inaugural$century

# accessing or assigning docvars for tokens using "$"
toks <- tokens(corpus_subset(data_corpus_inaugural, Year <= 1805))
toks$Year
toks$Year <- 1991:1995
toks$Year
toks$nonexistent <- TRUE
docvars(toks)

# accessing or assigning docvars for a dfm using "$"
dfm <- dfm(toks)
dfm$Year
dfm$Year <- 1991:1995
dfm$Year
dfm$nonexistent <- TRUE
docvars(dfm)

---

fcm Create a feature co-occurrence matrix
Description

Create a sparse feature co-occurrence matrix, measuring co-occurrences of features within a user-defined context. The context can be defined as a document or a window within a collection of documents, with an optional vector of weights applied to the co-occurrence counts.

Usage

fcm(
  x,
  context = c("document", "window"),
  count = c("frequency", "boolean", "weighted"),
  window = 5L,
  weights = NULL,
  ordered = FALSE,
  tri = TRUE,
  ...
)

Arguments

x  a tokens, or dfm object from which to generate the feature co-occurrence matrix
context  the context in which to consider term co-occurrence: "document" for co-occurrence counts within document; "window" for co-occurrence within a defined window of words, which requires a positive integer value for window. Note: if x is a dfm object, then context can only be "document".
count  how to count co-occurrences:
  "frequency" count the number of co-occurrences within the context
  "boolean" count only the co-occurrence or not within the context, irrespective of how many times it occurs.
  "weighted" count a weighted function of counts, typically as a function of distance from the target feature. Only makes sense for context = "window".
window  positive integer value for the size of a window on either side of the target feature, default is 5, meaning 5 words before and after the target feature
weights  a vector of weights applied to each distance from 1:window, strictly decreasing by default; can be a custom-defined vector of the same length as window
ordered  if TRUE, count only the forward co-occurrences for each target token for bigram models, so that the i, j cell of the fcm is the number of times that token j occurs before the target token i within the window. Only makes sense for context = "window", and when ordered = TRUE, the argument tri has no effect.
tri  if TRUE return only upper triangle (including diagonal). Ignored if ordered = TRUE.
...  not used here
Details

The function \texttt{fcm()} provides a very general implementation of a "context-feature" matrix, consisting of a count of feature co-occurrence within a defined context. This context, following Momtazi et. al. (2010), can be defined as the \texttt{document}, \texttt{sentences} within documents, \texttt{syntactic relationships} between features (nouns within a sentence, for instance), or according to a \texttt{window}. When the context is a window, a weighting function is typically applied that is a function of distance from the target word (see Jurafsky and Martin 2015, Ch. 16) and ordered co-occurrence of the two features is considered (see Church & Hanks 1990).

\texttt{fcm} provides all of this functionality, returning a $V \times V$ matrix (where $V$ is the vocabulary size, returned by \texttt{nfeat()}). The \texttt{tri = TRUE} option will only return the upper part of the matrix.

Unlike some implementations of co-occurrences, \texttt{fcm} counts feature co-occurrences with themselves, meaning that the diagonal will not be zero.

\texttt{fcm} also provides "boolean" counting within the context of "window", which differs from the counting within "document".

\texttt{is.fcm(x)} returns \texttt{TRUE} if and only if its \texttt{x} is an object of type \texttt{fcm}.

Author(s)

Kenneth Benoit (R), Haiyan Wang (R, C++), Kohei Watanabe (C++)

References


Examples

```
# see http://bit.ly/29b2zOA
toks1 <- tokens(c("A D A C E A D F E B A C E D"))
fcm(toks1, context = "window", window = 2)
fcm(toks1, context = "window", count = "weighted", window = 3)
fcm(toks1, context = "window", count = "weighted", window = 3,
weights = c(3, 2, 1), ordered = TRUE, tri = FALSE)

# with multiple documents
toks2 <- tokens(c("a a a b b c", "a a c e", "a c e f g"))
fcm(toks2, context = "document", count = "frequency")
fcm(toks2, context = "document", count = "boolean")
fcm(toks2, context = "window", window = 2)
```
fcm_sort

Sort an fcm in alphabetical order of the features

Description
Sorts an fcm in alphabetical order of the features.

Usage
fcm_sort(x)

Arguments
x
fcm object

Value
A fcm object whose features have been alphabetically sorted. Differs from fcm_sort() in that this function sorts the fcm by the feature labels, not the counts of the features.

Author(s)
Kenneth Benoit

Examples
# with tri = FALSE
fcm1 <- fcm(tokens(c("A X Y C B A", "X Y C A B B")), tri = FALSE)
rownames(fcm1)[3] <- colnames(fcm1)[3] <- "Z"
fcm1
fcm_sort(fcm1)

# with tri = TRUE
fcm2 <- fcm(tokens(c("A X Y C B A", "X Y C A B B")), tri = TRUE)
rownames(fcm2)[3] <- colnames(fcm2)[3] <- "Z"
fcm2
fcm_sort(fcm2)
featfreq

Compute the frequencies of features

Description
For a dfm object, returns a frequency for each feature, computed across all documents in the dfm. This is equivalent to colSums(x).

Usage
featfreq(x)

Arguments
x a dfm

Value
a (named) numeric vector of feature frequencies

See Also
dfm_tfidf(), dfm_weight()

Examples
dmat <- dfm(data_char_sampletext)
featfreq(dmat)

featnames
Get the feature labels from a dfm

Description
Get the features from a document-feature matrix, which are stored as the column names of the dfm object.

Usage
featnames(x)

Arguments
x the dfm whose features will be extracted
**Value**

character vector of the feature labels

**Examples**

```r
dfmat <- dfm(data_corpus_inaugural)

# first 50 features (in original text order)
head(featnames(dfmat), 50)

# first 50 features alphabetically
head(sort(featnames(dfmat)), 50)

# contrast with descending total frequency order from topfeatures()
names(topfeatures(dfmat, 50))
```

---

**index**  
*Locate a pattern in a tokens object*

**Description**

Locates a *pattern* within a tokens object, returning the index positions of the beginning and ending tokens in the pattern.

**Usage**

```r
index(
  x,  
  pattern,  
  valuetype = c("glob", "regex", "fixed"),  
  case_insensitive = TRUE
)
```

**Arguments**

- **x**  
  an input *tokens* object

- **pattern**  
  a character vector, list of character vectors, *dictionary*, or collocations object. See *pattern* for details.

- **valuetype**  
  the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See *value-type* for details.

- **case_insensitive**  
  logical; if TRUE, ignore case when matching a pattern or *dictionary* values
is.collocations

Value

a data.frame consisting of one row per pattern match, with columns for the document name, index positions from and to, and the pattern matched.

is.index returns TRUE if the object was created by index(); FALSE otherwise.

Examples

toks <- tokens(data_corpus_inaugural[1:8])
index(toks, pattern = "secure*")
index(toks, pattern = c("secure*", phrase("united states"))) %>% head()

is.collocations

Check if an object is collocations

Description

Function to check if an object is a collocations object, created by quanteda.textstats::textstat_collocations().

Usage

is.collocations(x)

Arguments

x object to be checked

Value

TRUE if the object is of class collocations, FALSE otherwise

kwic

Locate keywords-in-context

Description

For a text or a collection of texts (in a quanteda corpus object), return a list of a keyword supplied by the user in its immediate context, identifying the source text and the word index number within the source text. (Not the line number, since the text may or may not be segmented using end-of-line delimiters.)
Usage

kwic(
  x,
  pattern,
  window = 5,
  valuetype = c("glob", "regex", "fixed"),
  separator = " ",
  case_insensitive = TRUE,
  index = NULL,
  ...
)

is.kwic(x)

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

Arguments

x a character, corpus, or tokens object

pattern a character vector, list of character vectors, dictionary, or collocations object. See pattern for details.

window the number of context words to be displayed around the keyword

valuetype the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.

separator a character to separate words in the output

case_insensitive logical; if TRUE, ignore case when matching a pattern or dictionary values

index an index object to specify keywords

... unused

Value

A kwic classed data.frame, with the document name (docname) and the token index positions (from and to, which will be the same for single-word patterns, or a sequence equal in length to the number of elements for multi-word phrases).

Note

pattern will be a keyword pattern or phrase, possibly multiple patterns, that may include punctuation. If a pattern contains whitespace, it is best to wrap it in phrase() to make this explicit. However if pattern is a collocations or dictionary object, then the collocations or multi-word dictionary keys will automatically be considered phrases where each whitespace-separated element matches a token in sequence.
See Also

print-methods

Examples

# single token matching
toks <- tokens(data_corpus_inaugural[1:8])
kwic(toks, pattern = "secure*", valuetype = "glob", window = 3)
kwic(toks, pattern = "secur", valuetype = "regex", window = 3)
kwic(toks, pattern = "security", valuetype = "fixed", window = 3)

# phrase matching
kwic(toks, pattern = phrase("secur* against"), window = 2)
kwic(toks, pattern = phrase("war against"), valuetype = "regex", window = 2)

# use index
idx <- index(toks, phrase("secur* against"))
kwic(toks, index = idx, window = 2)
kw <- kwic(tokens(data_corpus_inaugural[1:20]), "provident*")
is.kwic(kw)
is.kwic("Not a kwic")
is.kwic(kw[, c("pre", "post")])
toks <- tokens(data_corpus_inaugural[1:8])
kw <- kwic(toks, pattern = "secure*", valuetype = "glob", window = 3)
as.data.frame(kw)

meta

Get or set object metadata

Description

Get or set the object metadata in a corpus, tokens, dfm, or dictionary object. With the exception of dictionaries, this will be corpus-level metadata.

Usage

meta(x, field = NULL, type = c("user", "object", "system", "all"))

meta(x, field = NULL) <- value

Arguments

x an object for which the metadata will be read or set
field metadata field name(s); if NULL (default), return all metadata names
type "user" for user-provided corpus-level metadata; "system" for metadata set automatically when the corpus is created; or "all" for all metadata.
value new value of the metadata field
Value

For `meta`, a named list of the metadata fields in the corpus.

For `meta <-`, the corpus with the updated user-level metadata. Only user-level metadata may be assigned.

Examples

```r
meta(data_corpus_inaugural)
meta(data_corpus_inaugural, "source")
meta(data_corpus_inaugural, "citation") <- "Presidential Speeches Online Project (2014)."
meta(data_corpus_inaugural, "citation")
```

ndoc  

Count the number of documents or features

Description

Get the number of documents or features in an object.

Usage

```r
ndoc(x)
nfeat(x)
```

Arguments

- `x` a `quanteda` object: a `corpus`, `dfm`, or `tokens` object, or a `readtext` object from the `readtext` package.

Details

`ndoc` returns the number of documents in an object whose texts are organized as "documents" (a `corpus`, `dfm`, or `tokens` object, a `readtext` object from the `readtext` package).

`nfeat` returns the number of features from a `dfm`; it is an alias for `ntype` when applied to `dfm` objects. This function is only defined for `dfm` objects because only these have "features". (To count tokens, see `ntoken()`.)

Value

an integer (count) of the number of documents or features

See Also

`ntoken()`
Examples

# number of documents
ndoc(data_corpus_inaugural)
ndoc(corpus_subset(data_corpus_inaugural, Year > 1980))
ndoc(tokens(data_corpus_inaugural))
ndoc(dfm(tokens(corpus_subset(data_corpus_inaugural, Year > 1980))))

# number of features
toks1 <- tokens(corpus_subset(data_corpus_inaugural, Year > 1980), remove_punct = FALSE)
toks2 <- tokens(corpus_subset(data_corpus_inaugural, Year > 1980), remove_punct = TRUE)
nfeat(dfm(toks1))
nfeat(dfm(toks2))

tsentence

nsentence(x)  Count the number of sentences

Description

Return the count of sentences in a corpus or character object.

Usage

nsentence(x)

Arguments

x a character or corpus whose sentences will be counted

Value

count(s) of the total sentences per text

Note

nsentence() relies on the boundaries definitions in the stringi package (see stri_opts_brkiter). It does not count sentences correctly if the text has been transformed to lower case, and for this reason nsentence() will issue a warning if it detects all lower-cased text.

Examples

# simple example
txt <- c(text1 = "This is a sentence: second part of first sentence.",
         text2 = "A word. Repeated repeated.",
         text3 = "Mr. Jones has a PhD from the LSE. Second sentence.")
nsentence(txt)
Description

Get the count of tokens (total features) or types (unique tokens).

Usage

```
ntoken(x,)
```

```
ntype(x,)
```

Arguments

```
x a quanteda object: a character, corpus, tokens, or dfm object
...
```

additional arguments passed to `tokens()`

Details

The precise definition of "tokens" for objects not yet tokenized (e.g. character or corpus objects) can be controlled through optional arguments passed to `tokens()` through ... .

For `dfm` objects, `ntype` will only return the count of features that occur more than zero times in the `dfm`.

Value

named integer vector of the counts of the total tokens or types

Note

Due to differences between raw text tokens and features that have been defined for a `dfm`, the counts may be different for dfm objects and the texts from which the dfm was generated. Because the method tokenizes the text in order to count the tokens, your results will depend on the options passed through to `tokens()`.

Examples

```
# simple example
txt <- c(text1 = "This is a sentence, this.", text2 = "A word. Repeated repeated.")
ntoken(txt)
nype(txt)
nntoken(char_tolower(txt))  # same
ntype(char_tolower(txt))  # fewer types
ntoken(char_tolower(txt), remove_punct = TRUE)
nype(char_tolower(txt), remove_punct = TRUE)
```

# with some real texts
phrase

Declare a pattern to be a sequence of separate patterns

Description

Declares that a character expression consists of multiple patterns, separated by an element such as whitespace. This is typically used as a wrapper around `pattern()` to make it explicit that the pattern elements are to be used for matches to multi-word sequences, rather than individual, unordered matches to single words.

Usage

```r
phrase(x, separator = " ")
```

```r
as.phrase(x)
```

```r
is.phrase(x)
```

Arguments

- **x** character, `dictionary`, list, collocations, or tokens object; the compound patterns to be treated as a sequence separated by `separator`. For list, collocations, or tokens objects, use `as.phrase()`.
- **separator** character; the character in between the patterns. This defaults to " ". For `phrase()` only.

Value

`phrase()` and `as.phrase()` return a specially classed list whose elements have been split into separate character (pattern) elements.

`is.phrase` returns `TRUE` if the object was created by `phrase()`: `FALSE` otherwise.

See Also

`as.phrase()`
Examples

# make phrases from characters
phrase(c("natural language processing"))
phrase(c("natural_language_processing", "text_analysis"), separator = "_")

# from a dictionary
phrase(dictionary(list(catone = c("a b"), cattwo = "c d e", catthree = "f")))

# from a list
as.phrase(list(c("natural", "language", "processing")))

# from tokens
as.phrase(tokens("natural language processing"))

print-methods
Print methods for quanteda core objects

Description

Print method for quanteda objects. In each max_n* option, 0 shows none, and -1 shows all.

Usage

## S3 method for class 'corpus'
print(
  x,
  max_ndoc = quanteda_options("print_corpus_max_ndoc"),
  max_nchar = quanteda_options("print_corpus_max_nchar"),
  show_summary = quanteda_options("print_corpus_summary"),
  ...
)

## S4 method for signature 'dfm'
print(
  x,
  max_ndoc = quanteda_options("print_dfm_max_ndoc"),
  max_nfeat = quanteda_options("print_dfm_max_nfeat"),
  show_summary = quanteda_options("print_dfm_summary"),
  ...
)

## S4 method for signature 'dictionary2'
print(
  x,
  max_nkey = quanteda_options("print_dictionary_max_nkey"),
  max_nval = quanteda_options("print_dictionary_max_nval"),
  show_summary = quanteda_options("print_dictionary_summary"),
  ...)
...)

## S4 method for signature 'fcm'
print(
  x,
  max_nfeat = quanteda_options("print_dfm_max_nfeat"),
  show_summary = TRUE,
  ...
)

## S3 method for class 'kwic'
print(
  x,
  max_nrow = quanteda_options("print_kwic_max_nrow"),
  show_summary = quanteda_options("print_kwic_summary"),
  ...
)

## S3 method for class 'tokens'
print(
  x,
  max_ndoc = quanteda_options("print_tokens_max_ndoc"),
  max_ntoken = quanteeda_options("print_tokens_max_ntoken"),
  show_summary = quanteda_options("print_tokens_summary"),
  ...
)

**Arguments**

- **x**
  - the object to be printed
- **max_ndoc**
  - max number of documents to print; default is from the print_*_max_ndoc setting of `quanteda_options()`
- **max_nchar**
  - max number of tokens to print; default is from the print_corpus_max_nchar setting of `quanteda_options()`
- **show_summary**
  - print a brief summary indicating the number of documents and other characteristics of the object, such as docvars or sparsity.
- **...**
  - not used
- **max_nfeat**
  - max number of features to print; default is from the print_dfm_max_nfeat setting of `quanteda_options()`
- **max_nkey**
  - max number of keys to print; default is from the print_dictionary_max_max_nkey setting of `quanteda_options()`
- **max_nval**
  - max number of values to print; default is from the print_dictionary_max_nval setting of `quanteda_options()`
- **max_nrow**
  - max number of documents to print; default is from the print_kwic_max_nrow setting of `quanteda_options()`
max_ntoken  max number of tokens to print; default is from the print_tokens_max_ntoken
setting of quanteda_options()

See Also

quanteda_options()

Examples

corp <- corpus(data_char_ukimmig2010)
print(corp, max_ndoc = 3, max_nchar = 40)

toks <- tokens(corp)
print(toks, max_ndoc = 3, max_ntoken = 6)

dfm <- dfm(toks)
print(dfm, max_ndoc = 3, max_nfeat = 10)

quanteda_options  Get or set package options for quanteda

Description

Get or set global options affecting functions across quanteda.

Usage

quanteda_options(..., reset = FALSE, initialize = FALSE)

Arguments

... options to be set, as key-value pair, same as options(). This may be a list of valid key-value pairs, useful for setting a group of options at once (see examples).
reset logical; if TRUE, reset all quanteda options to their default values
initialize logical; if TRUE, reset only the quanteda options that are not already defined. Used for setting initial values when some have been defined previously, such as in .Rprofile.

Details

Currently available options are:

verbose  logical; if TRUE then use this as the default for all functions with a verbose argument
threads  integer; specifies the number of threads to use in parallelized functions; defaults to RcppParallel::defaultNumThreads(); the number of threads can be changed only once in a session
print_dfm_max_ndoc  integer; specifies the number of documents to display when using the defaults for printing a dfm
quanteda_options

print_dfm_max_nfeat integer; specifies the number of features to display when using the defaults for printing a dfm

base_docname character; stem name for documents that are unnamed when a corpus, tokens, or dfm are created or when a dfm is converted from another object

base_featname character; stem name for features that are unnamed when they are added, for whatever reason, to a dfm through an operation that adds features

base_compname character; stem name for components that are created by matrix factorization

language_stemmer character; language option for char_wordstem(), tokens_wordstem(), and dfm_wordstem()

pattern_hashtag, pattern_username character; regex patterns for (social media) hashtags and usernames respectively, used to avoid segmenting these in the default internal "word" tokenizer

tokens_block_size integer; specifies the number of documents to be tokenized at a time in blocked tokenization. When the number is large, tokenization becomes faster but also memory-intensive.

tokens_locale character; specify locale in stringi boundary detection in tokenization and corpus reshaping. See stringi::stri_opts_brkiter().

Value

When called using a key = value pair (where key can be a label or quoted character name)), the option is set and TRUE is returned invisibly.

When called with no arguments, a named list of the package options is returned.

When called with reset = TRUE as an argument, all arguments are options are reset to their default values, and TRUE is returned invisibly.

Examples

(opt <- quanteda_options())
quanteda_options(verbos = TRUE)
quanteda_options("verbos" = FALSE)
quanteda_options("threads")
quanteda_options(print_dfm_max_ndoc = 50L)
# reset to defaults
quanteda_options(reset = TRUE)
# reset to saved options
quanteda_options(opt)
readtext-methods  Extensions for readtext objects

Description

These functions provide quanteda methods for readtext objects.

Arguments

- **x**: an object returned by spacy_parse, or (for spacy_parse) a corpus object
- **...**: not used for these functions

Details

- `texts(x)` returns the texts from a readtext object
- `docnames(x)` returns the document names from a readtext object
- `docvars(x, field = NULL)` returns a data.frame of the document variables from a readtext object
- `ndoc(x)` returns the number of documents from a readtext object

spacyr-methods  Extensions for and from spacy_parse objects

Description

These functions provide quanteda methods for spacyr objects, and also extend spacy_parse and spacy_tokenize to work directly with corpus objects.

Arguments

- **x**: an object returned by spacy_parse, or (for spacy_parse) a corpus object
- **...**: not used for these functions

Details

- `spacy_parse(x,...)` and `spacy_tokenize(x,...)` work directly on quanteda corpus objects.
- `docnames(x)` returns the document names
- `ndoc(x)` returns the number of documents
- `ntoken(x,...)` returns the number of tokens by document
- `ntype(x,...)` returns the number of types (unique tokens) by document
- `nsentence(x)` returns the number of sentences by document
### Examples

```r
## Not run:
library("spacyr")
spacy_initialize()

corp <- corpus(c(doc1 = "And now, now, now for something completely different.",
                 doc2 = "Jack and Jill are children."))
spacy_tokenize(corp)
(parsed <- spacy_parse(corp))

ntype(parsed)
ntoken(parsed)
ndoc(parsed)
docnames(parsed)

## End(Not run)
```

---

**sparsity**  
*Compute the sparsity of a document-feature matrix*

### Description

Return the proportion of sparseness of a document-feature matrix, equal to the proportion of cells that have zero counts.

### Usage

`sparsity(x)`

### Arguments

- `x`  
  the document-feature matrix

### Examples

```r
dfmat <- dfm(data_corpus_inaugural)
sparsity(dfmat)
sparsity(dfm_trim(dfmat, min_termfreq = 5))
```
textmodels  
*Models for scaling and classification of textual data*

Description
The `textmodel_*()` functions formerly in `quanteda` have now been moved to the `quanteda.textmodels` package.

See Also
`quanteda.textmodels::quanteda.textmodels-package`

---

textplots  
*Plots for textual data*

Description
The `textplot_*()` functions formerly in `quanteda` have now been moved to the `quanteda.textplots` package.

See Also
`quanteda.textplots::quanteda.textplots-package`

---

textstats  
*Statistics for textual data*

Description
The `textstat_*()` functions formerly in `quanteda` have now been moved to the `quanteda.textstats` package.

See Also
`quanteda.textstats::quanteda.textstats-package`
Construct a tokens object, either by importing a named list of characters from an external tokenizer, or by calling the internal `quanteda` tokenizer.

```r
tokens(x, what = "word", remove_punct = FALSE, remove_symbols = FALSE, remove_numbers = FALSE, remove_url = FALSE, remove_separators = TRUE, split_hyphens = FALSE, include_docvars = TRUE, padding = FALSE, verbose = quanteda_options("verbose"), ...)
```

**Arguments**

- **x**
  - the input object to the tokens constructor, one of: a (uniquely) named list of characters; a tokens object; or a corpus or character object that will be tokenized

- **what**
  - character; which tokenizer to use. The default `what = "word"` is the version 2 `quanteda` tokenizer. Legacy tokenizers (version < 2) are also supported, including the default `what = "word1"`. See the Details and quanteda Tokenizers below.

- **remove_punct**
  - logical; if TRUE remove all characters in the Unicode "Punctuation" [P] class, with exceptions for those used as prefixes for valid social media tags if `preserve_tags = TRUE`

- **remove_symbols**
  - logical; if TRUE remove all characters in the Unicode "Symbol" [S] class

- **remove_numbers**
  - logical; if TRUE remove tokens that consist only of numbers, but not words that start with digits, e.g. 2day

- **remove_url**
  - logical; if TRUE find and eliminate URLs beginning with http(s)

- **remove_separators**
  - logical; if TRUE remove separators and separator characters (Unicode "Separator" [Z] and "Control" [C] categories)

- **split_hyphens**
  - logical; if TRUE, split words that are connected by hyphenation and hyphenation-like characters in between words, e.g. "self-aware" becomes c("self","-","aware")
include_docvars
  if TRUE, pass docvars through to the tokens object. Does not apply when the
  input is a character data or a list of characters.

padding
  if TRUE, leave an empty string where the removed tokens previously existed.
  This is useful if a positional match is needed between the pre- and post-selected
  tokens, for instance if a window of adjacency needs to be computed.

verbose
  if TRUE, print timing messages to the console

... used to pass arguments among the functions

Details
tokens() works on tokens class objects, which means that the removal rules can be applied post-
tokenization, although it should be noted that it will not be possible to remove things that are not
present. For instance, if the tokens object has already had punctuation removed, then tokens(x, remove_punct
  = TRUE) will have no additional effect.

Value
  quanteda tokens class object, by default a serialized list of integers corresponding to a vector of
types.

Details
As of version 2, the choice of tokenizer is left more to the user, and tokens() is treated more as
a constructor (from a named list) than a tokenizer. This allows users to use any other tokenizer
that returns a named list, and to use this as an input to tokens(), with removal and splitting rules
applied after this has been constructed (passed as arguments). These removal and splitting rules are
conservative and will not remove or split anything, however, unless the user requests it.

Using external tokenizers is best done by piping the output from these other tokenizers into the
tokens() constructor, with additional removal and splitting options applied at the construction
stage. These will only have an effect, however, if the tokens exist for which removal is specified
at in the tokens() call. For instance, it is impossible to remove punctuation if the input list to
tokens() already had its punctuation tokens removed at the external tokenization stage.

To construct a tokens object from a list with no additional processing, call as.tokens() instead of
tokens().

Recommended tokenizers are those from the tokenizers package, which are generally faster than
the default (built-in) tokenizer but always splits infix hyphens, or spacyr.

quanteda Tokenizers
The default word tokenizer what = "word" splits tokens using stri_split_boundaries(x, type = "word")
but by default preserves infix hyphens (e.g. "self-funding"), URLs, and social media "tag" charac-
ters (#hashtags and @usernames), and email addresses. The rules defining a valid "tag" can
be found at https://www.hashtags.org/featured/what-characters-can-a-hashtag-include/ for hashtags

In versions < 2, the argument remove_twitter controlled whether social media tags were preserved
or removed, even when remove_punct = TRUE. This argument is not longer functional in versions
>= 2. If greater control over social media tags is desired, you should use an alternative tokenizer, including non-quanteda options.

For backward compatibility, the following older tokenizers are also supported through what:

"word1" (legacy) implements similar behavior to the version of what = "word" found in pre-
version 2. (It preserves social media tags and infix hyphens, but splits URLs.) "word1" is also 
slower than "word".

"fasterword" (legacy) splits on whitespace and control characters, using stringi::stri_split_charclass(x,"[\p{Z}\p{C}]+")

"fastestword" (legacy) splits on the space character, using stringi::stri_split_fixed(x," ")

"character" tokenization into individual characters

"sentence" sentence segmenter based on stri_split_boundaries, but with additional rules to avoid 
splits on words like "Mr." that would otherwise incorrectly be detected as sentence boundaries.
For better sentence tokenization, consider using spacyr.

See Also

tokens_ngrams(), tokens_skipgrams(), as.list.tokens(), as.tokens()

Examples

txt <- c(doc1 = "A sentence, showing how tokens() works.",
        doc2 = "@quantedainit and #textanalysis https://example.com?p=123.",
        doc3 = "Self-documenting code??",
        doc4 = "£1,000,000 for 50¢ is gr8 4ever \U0001f600")
tokens(txt)
tokens(txt, what = "word1")

# removing punctuation marks but keeping tags and URLs
tokens(txt[1:2], remove_punct = TRUE)

# splitting hyphenated words
tokens(txt[3])
tokens(txt[3], split_hyphens = TRUE)

# symbols and numbers
tokens(txt[4])
tokens(txt[4], remove_numbers = TRUE)
tokens(txt[4], remove_numbers = TRUE, remove_symbols = TRUE)

## Not run: # using other tokenizers
tokens(tokenizers::tokenize_words(txt[4]), remove_symbols = TRUE)
tokenizers::tokenize_words(txt, lowercase = FALSE, strip_punct = FALSE) %>%
tokens(remove_symbols = TRUE)
tokenizers::tokenize_characters(txt[3], strip_non_alphanum = FALSE) %>%
tokens(remove_punct = TRUE)
tokenizers::tokenize_sentences(
    "The quick brown fox. It jumped over the lazy dog.") %>%
tokens()
## End(Not run)

### tokens_chunk

*Segment tokens object by chunks of a given size*

**Description**

Segment tokens into new documents of equally sized token lengths, with the possibility of overlapping the chunks.

**Usage**

```
tokens_chunk(x, size, overlap = 0, use_docvars = TRUE)
```

**Arguments**

- **x**: `tokens` object whose token elements will be segmented into chunks
- **size**: integer; the token length of the chunks
- **overlap**: integer; the number of tokens in a chunk to be taken from the last `overlap` tokens from the preceding chunk
- **use_docvars**: if TRUE, repeat the docvar values for each chunk; if FALSE, drop the docvars in the chunked tokens

**Value**

A `tokens` object whose documents have been split into chunks of length `size`.

**See Also**

- `tokens_segment()`

**Examples**

```
txts <- c(doc1 = "Fellow citizens, I am again called upon by the voice of
  my country to execute the functions of its Chief Magistrate.",
  doc2 = "When the occasion proper for it shall arrive, I shall
  endeavor to express the high sense I entertain of this
distinguished honor.")
toks <- tokens(txts)
tokens_chunk(toks, size = 5)
tokens_chunk(toks, size = 5, overlap = 4)
```
tokens_compound

Convert token sequences into compound tokens

Description

Replace multi-token sequences with a multi-word, or "compound" token. The resulting compound tokens will represent a phrase or multi-word expression, concatenated with concatenator (by default, the ",_" character) to form a single "token". This ensures that the sequences will be processed subsequently as single tokens, for instance in constructing a dfm.

Usage

tokens_compound(
  x,
  pattern,
  valuetype = c("glob", "regex", "fixed"),
  concatenator = ",_",
  window = 0L,
  case_insensitive = TRUE,
  join = TRUE
)

Arguments

x an input tokens object

pattern a character vector, list of character vectors, dictionary, or collocations object. See pattern for details.

valuetype the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.

concatenator the concatenation character that will connect the words making up the multi-word sequences. The default _ is recommended since it will not be removed during normal cleaning and tokenization (while nearly all other punctuation characters, at least those in the Unicode punctuation class [P] will be removed).

window integer; a vector of length 1 or 2 that specifies size of the window of tokens adjacent to pattern that will be compounded with matches to pattern. The window can be asymmetric if two elements are specified, with the first giving the window size before pattern and the second the window size after. If paddings (empty "" tokens) are found, window will be shrunk to exclude them.

case_insensitive logical; if TRUE, ignore case when matching a pattern or dictionary values

join logical; if TRUE, join overlapping compounds into a single compound; otherwise, form these separately. See examples.
Value

A tokens object in which the token sequences matching pattern have been replaced by new compounded "tokens" joined by the concatenator.

Note

Patterns to be compounded (naturally) consist of multi-word sequences, and how these are expected in pattern is very specific. If the elements to be compounded are supplied as space-delimited elements of a character vector, wrap the vector in phrase(). If the elements to be compounded are separate elements of a character vector, supply it as a list where each list element is the sequence of character elements.

See the examples below.

Examples

txt <- "The United Kingdom is leaving the European Union."
toks <- tokens(txt, remove_punct = TRUE)

# character vector - not compounded
tokens_compound(toks, c("United", "Kingdom", "European", "Union"))

# elements separated by spaces - not compounded
tokens_compound(toks, c("United Kingdom", "European Union"))

# list of characters - is compounded
tokens_compound(toks, list(c("United", "Kingdom"), c("European", "Union")))

# elements separated by spaces, wrapped in phrase() - is compounded
tokens_compound(toks, phrase(c("United Kingdom", "European Union")))

# supplied as values in a dictionary (same as list) - is compounded
# (keys do not matter)
tokens_compound(toks, dictionary(list(key1 = "United Kingdom", key2 = "European Union")))

# pattern as dictionaries with glob matches
tokens_compound(toks, dictionary(list(key1 = c("U* K*")), valuetype = "glob"))

# note the differences caused by join = FALSE
compounds <- list(c("the", "European"), c("European", "Union"))
tokens_compound(toks, pattern = compounds, join = TRUE)
tokens_compound(toks, pattern = compounds, join = FALSE)

# use window to form ngrams
tokens_remove(toks, pattern = stopwords("en")) %>%
  tokens_compound(pattern = "leav*", join = FALSE, window = c(0, 3))
tokens_group

Combine documents in a tokens object by a grouping variable

Description

Combine documents in a tokens object by a grouping variable, by concatenating the tokens in the order of the documents within each grouping variable.

Usage

tokens_group(x, groups = docid(x), fill = FALSE)

Arguments

x  
tokens object

groups  
grouping variable for sampling, equal in length to the number of documents. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for groups. See news(Version >= "3.0", package = "quanteda") for details.

fill  
logical; if TRUE and groups is a factor, then use all levels of the factor when forming the new documents of the grouped object. This will result in a new "document" with empty content for levels not observed, but for which an empty document may be needed. If groups is a factor of dates, for instance, then fill = TRUE ensures that the new object will consist of one new "document" by date, regardless of whether any documents previously existed with that date. Has no effect if the groups variable(s) are not factors.

Value

a tokens object whose documents are equal to the unique group combinations, and whose tokens are the concatenations of the tokens by group. Document-level variables that have no variation within groups are saved in docvars. Document-level variables that are lists are dropped from grouping, even when these exhibit no variation within groups.

Examples

corp <- corpus(c("a a b", "a b c c", "a c d d", "a c c d"),
   docvars = data.frame(grp = c("grp1", "grp1", "grp2", "grp2")))
toks <- tokens(corp)
tokens_group(toks, groups = grp)
tokens_group(toks, groups = c(1, 1, 2, 2))

# with fill
tokens_group(toks, groups = factor(c(1, 1, 2, 2), levels = 1:3))
tokens_group(toks, groups = factor(c(1, 1, 2, 2), levels = 1:3), fill = TRUE)
tokens_lookup  

Apply a dictionary to a tokens object

Description

Convert tokens into equivalence classes defined by values of a dictionary object.

Usage

tokens_lookup(
  x,
  dictionary,
  levels = 1:5,
  valuetype = c("glob", "regex", "fixed"),
  case_insensitive = TRUE,
  capkeys = !exclusive,
  exclusive = TRUE,
  nomatch = NULL,
  nested_scope = c("key", "dictionary"),
  verbose = quanteda_options("verbose")
)

Arguments

x  
tokens object to which dictionary or thesaurus will be supplied
dictionary  
the dictionary-class object that will be applied to x
levels  
integers specifying the levels of entries in a hierarchical dictionary that will be applied. The top level is 1, and subsequent levels describe lower nesting levels. Values may be combined, even if these levels are not contiguous, e.g. levels = c(1:3) will collapse the second level into the first, but record the third level (if present) collapsed below the first (see examples).
valuetype  
the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.
case_insensitive  
logical; if TRUE, ignore case when matching a pattern or dictionary values
capkeys  
if TRUE, convert dictionary keys to uppercase to distinguish them from other features
exclusive  
if TRUE, remove all features not in dictionary, otherwise, replace values in dictionary with keys while leaving other features unaffected
nomatch  
an optional character naming a new key for tokens that do not matched to a dictionary values If NULL (default), do not record unmatched tokens.
nested_scope  
how to treat matches from different dictionary keys that are nested. When one value is nested within another, such as "a b" being nested within "a b c", the
tokens_lookup() will match the longer. When nested_scope = "key", this longer-match priority is applied only within the key, while "dictionary"'s applies it across keys, matching only the key with the longer pattern, not the matches nested within that longer pattern from other keys. See Details.

### Details

Dictionary values may consist of sequences, and there are different methods of counting key matches based on values that are nested or that overlap.

When two different keys in a dictionary are nested matches of one another, the nested_scope options provide the choice of matching each key's values independently (the "key") option, or just counting the longest match (the "dictionary" option). Values that are nested within the same key are always counted as a single match. See the last example below comparing the New York and New York Times for these two different behaviours.

Overlapping values, such as "a b" and "b a" are currently always considered as separate matches if they are in different keys, or as one match if the overlap is within the same key. Overlapped

### See Also

tokens_replace

### Examples

toks1 <- tokens(data_corpus_inaugural)
dict1 <- dictionary(list(country = "united states",
law=c("law*", "constitution"),
freedom=c("free*", "libert*")))
dfm(tokens_lookup(toks1, dict1, valuetype = "glob", verbose = TRUE))
dfm(tokens_lookup(toks1, dict1, valuetype = "glob", verbose = TRUE, nomatch = "NONE"))

dict2 <- dictionary(list(country = "united states",
law = c("law", "constitution"),
freedom = c("freedom", "liberty")))
# dfm(applyDictionary(toks1, dict2, valuetype = "fixed"))
dfm(tokens_lookup(toks1, dict2, valuetype = "fixed"))

# hierarchical dictionary example
txt <- c(d1 = "The United States has the Atlantic Ocean and the Pacific Ocean.",
d2 = "Britain and Ireland have the Irish Sea and the English Channel." )
toks2 <- tokens(txt)
dict3 <- dictionary(list(US = list(Countries = c("States"), oceans = c("Atlantic", "Pacific")),
Europe = list(Countries = c("Britain", "Ireland"), oceans = list(west = "Irish Sea", east = "English Channel"))))
tokens_lookup(toks2, dict3, levels = 1)
tokens_lookup(toks2, dict3, levels = 2)
tokens_lookup(toks2, dict3, levels = 1:2)
tokens_lookup(toks2, dict3, levels = 3)
tokens_lookup(toks2, dict3, levels = c(1,3))
tokens_lookup(toks2, dict3, levels = c(2,3))

# show unmatched tokens
tokens_lookup(toks2, dict3, nomatch = "_UNMATCHED")

# nested matching differences
dict4 <- dictionary(list(paper = "New York Times", city = "New York"))
toks4 <- tokens("The New York Times is a New York paper.")
tokens_lookup(toks4, dict4, nested_scope = "key", exclusive = FALSE)
tokens_lookup(toks4, dict4, nested_scope = "dictionary", exclusive = FALSE)

---

tokens_ngrams

Create ngrams and skipgrams from tokens

Description

Create a set of ngrams (tokens in sequence) from already tokenized text objects, with an optional skip argument to form skipgrams. Both the ngram length and the skip lengths take vectors of arguments to form multiple lengths or skips in one pass. Implemented in C++ for efficiency.

Usage

tokens_ngrams(x, n = 2L, skip = 0L, concatenator = "_")

char_ngrams(x, n = 2L, skip = 0L, concatenator = "_")

tokens_skipgrams(x, n, skip, concatenator = "_")

Arguments

x a tokens object, or a character vector, or a list of characters

n integer vector specifying the number of elements to be concatenated in each ngram. Each element of this vector will define a n in the n-gram(s) that are produced.

skip integer vector specifying the adjacency skip size for tokens forming the ngrams, default is 0 for only immediately neighbouring words. For skipgrams, skip can be a vector of integers, as the "classic" approach to forming skip-grams is to set skip = k where k is the distance for which k or fewer skips are used to construct the n-gram. Thus a "4-skip-n-gram" defined as skip = 0:4 produces results that include 4 skips, 3 skips, 2 skips, 1 skip, and 0 skips (where 0 skips are typical n-grams formed from adjacent words). See Guthrie et al (2006).

concatenator character for combining words, default is _ (underscore) character
Details

Normally, these functions will be called through \( \text{tokens}(x, \text{ngrams = ...}), \) but these functions are provided in case a user wants to perform lower-level ngram construction on tokenized texts.

\texttt{tokens\_skipgrams()} is a wrapper to \texttt{tokens\_ngrams()} that requires arguments to be supplied for both \( n \) and \( \text{skip} \). For \( k \)-skip skipgrams, set \( \text{skip} \) to \( 0:k \), in order to conform to the definition of skip-grams found in Guthrie et al (2006): A \( k \) skip-gram is an ngram which is a superset of all ngrams and each \( (k - i) \) skipgram until \( (k - i) = 0 \) (which includes 0 skip-grams).

Value

a tokens object consisting a list of character vectors of ngrams, one list element per text, or a character vector if called on a simple character vector

Note

\texttt{char\_ngrams} is a convenience wrapper for a (non-list) vector of characters, so named to be consistent with \texttt{quanteda}'s naming scheme.

References


Examples

\begin{verbatim}
# ngrams
tokens\_ngrams(tokens(c("a b c d e", "c d e f g")), n = 2:3)

toks <- tokens(c(text1 = "the quick brown fox jumped over the lazy dog"))
tokens\_ngrams(toks, n = 1:3)
tokens\_ngrams(toks, n = c(2,4), concatenator = " ")
tokens\_ngrams(toks, n = c(2,4), skip = 1, concatenator = " ")
# on character
char\_ngrams(letters[1:3], n = 1:3)

# skipgrams

toks <- tokens("insurgents killed in ongoing fighting")
tokens\_skipgrams(toks, n = 2, skip = 0:1, concatenator = " ")
tokens\_skipgrams(toks, n = 2, skip = 0:2, concatenator = " ")
tokens\_skipgrams(toks, n = 3, skip = 0:2, concatenator = " ")
\end{verbatim}
Description

Substitute token types based on vectorized one-to-one matching. Since this function is created for lemmatization or user-defined stemming. It supports substitution of multi-word features by multi-word features, but substitution is fastest when pattern and replacement are character vectors and valuetype = "fixed" as the function only substitute types of tokens. Please use tokens_lookup() with exclusive = FALSE to replace dictionary values.

Usage

tokens_replace(
  x,  
  pattern,  
  replacement,  
  valuetype = "glob",  
  case_insensitive = TRUE,  
  verbose = quanteda_options("verbose")
)

Arguments

x: tokens object whose token elements will be replaced
pattern: a character vector or list of character vectors. See pattern for more details.
replacement: a character vector or (if pattern is a list) list of character vectors of the same length as pattern
valuetype: the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.
case_insensitive: logical; if TRUE, ignore case when matching a pattern or dictionary values
verbose: print status messages if TRUE

See Also
tokens_lookup

Examples

toks1 <- tokens(data_corpus_inaugural, remove_punct = TRUE)

# lemmatization
taxwords <- c("tax", "taxing", "taxed", "taxed", "taxation")
lemma <- rep("TAX", length(taxwords))
toks2 <- tokens_replace(toks1, taxwords, lemma, valuetype = "fixed")
kwic(toks2, "TAX") %>%
  tail(10)

# stemming
type <- types(toks1)
stem <- char_wordstem(type, "porter")
toks3 <- tokens_replace(toks1, type, stem, valuetype = "fixed", caseInsensitive = FALSE)
identical(toks3, tokens_wordstem(toks1, "porter"))

# multi-multi substitution
toks4 <- tokens_replace(toks1, phrase(c("Supreme Court")), phrase(c("Supreme Court of the United States")))
kwic(toks4, phrase(c("Supreme Court of the United States")))

tokens_sample  Randomly sample documents from a tokens object

Description

Take a random sample of documents of the specified size from a corpus, with or without replacement, optionally by grouping variables or with probability weights.

Usage

tokens_sample(x, size = NULL, replace = FALSE, prob = NULL, by = NULL)

Arguments

x  a tokens object whose documents will be sampled
size  a positive number, the number of documents to select; when used with by, the number to select from each group or a vector equal in length to the number of groups defining the samples to be chosen in each category of by. By defining a size larger than the number of documents, it is possible to oversample when replace = TRUE.
replace  if TRUE, sample with replacement
prob  a vector of probability weights for obtaining the elements of the vector being sampled. May not be applied when by is used.
by  optional grouping variable for sampling. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for by. See news(Version >= "2.9", package = "quanteda") for details.

Value

a tokens object (re)sampled on the documents, containing the document variables for the documents sampled.

See Also

describe
Examples

```r
set.seed(123)
toks <- tokens(data_corpus_inaugural[1:6])
toks
tokens_sample(toks)
tokens_sample(toks, replace = TRUE) %>% docnames()
tokens_sample(toks, size = 3, replace = TRUE) %>% docnames()

# sampling using by
docvars(toks)
tokens_sample(toks, size = 2, replace = TRUE, by = Party) %>% docnames()
```

tokens_select

Select or remove tokens from a tokens object

Description

These function select or discard tokens from a tokens object. For convenience, the functions `tokens_remove` and `tokens_keep` are defined as shortcuts for `tokens_select(x, pattern, selection = "remove")` and `tokens_select(x, pattern, selection = "keep")`, respectively. The most common usage for `tokens_remove` will be to eliminate stop words from a text or text-based object, while the most common use of `tokens_select` will be to select tokens with only positive pattern matches from a list of regular expressions, including a dictionary. `startpos` and `endpos` determine the positions of tokens searched for `pattern` and areas affected are expanded by `window`.

Usage

```r
tokens_select(
  x,
  pattern,
  selection = c("keep", "remove"),
  valuetype = c("glob", "regex", "fixed"),
  case_insensitive = TRUE,
  padding = FALSE,
  window = 0,
  min_nchar = NULL,
  max_nchar = NULL,
  startpos = 1L,
  endpos = -1L,
  verbose = quanteda_options("verbose")
)
```

tokens_remove(x, ...)

tokens_keep(x, ...)
Arguments

- **x**
  - tokens object whose token elements will be removed or kept

- **pattern**
  - a character vector, list of character vectors, dictionary, or collocations object. See pattern for details.

- **selection**
  - whether to "keep" or "remove" the tokens matching pattern

- **valuetype**
  - the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.

- **case_insensitive**
  - logical; if TRUE, ignore case when matching a pattern or dictionary values

- **padding**
  - if TRUE, leave an empty string where the removed tokens previously existed. This is useful if a positional match is needed between the pre- and post-selected tokens, for instance if a window of adjacency needs to be computed.

- **window**
  - integer of length 1 or 2; the size of the window of tokens adjacent to pattern that will be selected. The window is symmetric unless a vector of two elements is supplied, in which case the first element will be the token length of the window before pattern, and the second will be the token length of the window after pattern. The default is 0, meaning that only the pattern matched token(s) are selected, with no adjacent terms. Terms from overlapping windows are never double-counted, but simply returned in the pattern match. This is because tokens_select never redefines the document units; for this, see kwic().

- **min_nchar, max_nchar**
  - optional numerics specifying the minimum and maximum length in characters for tokens to be removed or kept; defaults are NULL for no limits. These are applied after (and hence, in addition to) any selection based on pattern matches.

- **startpos, endpos**
  - integer; position of tokens in documents where pattern matching starts and ends, where 1 is the first token in a document. For negative indexes, counting starts at the ending token of the document, so that -1 denotes the last token in the document, -2 the second to last, etc. When the length of the vector is equal to ndoc, tokens in corresponding positions will be selected. Otherwise, only the first element in the vector is used.

- **verbose**
  - if TRUE print messages about how many tokens were selected or removed

- **...**
  - additional arguments passed by tokens_remove and tokens_keep to tokens_select. Cannot include selection.

Value

a tokens object with tokens selected or removed based on their match to pattern

Examples

```r
## tokens_select with simple examples
toks <- as.tokens(list(letters, LETTERS))
tokens_select(toks, c("b", "e", "f"), selection = "keep", padding = FALSE)
```
tokens_split(toks, c("b", "e", "f"), selection = "keep", padding = TRUE)
tokens_split(toks, c("b", "e", "f"), selection = "remove", padding = FALSE)
tokens_split(toks, c("b", "e", "f"), selection = "remove", padding = TRUE)

# how case_insensitive works
tokens_select(toks, c("b", "e", "f"), selection = "remove", case_insensitive = TRUE)
tokens_select(toks, c("b", "e", "f"), selection = "remove", case_insensitive = FALSE)

# use window
tokens_select(toks, c("b", "f"), selection = "keep", window = 1)
tokens_select(toks, c("b", "f"), selection = "remove", window = 1)
tokens_remove(toks, c("b", "f"), window = c(0, 1))
tokens_select(toks, tokens, pattern = c("e", "g"), window = c(1, 2))

# tokens_remove example: remove stopwords
txt <- c(wash1 <- "Fellow citizens, I am again called upon by the voice of my
country to execute the functions of its Chief Magistrate.",
        wash2 <- "When the occasion proper for it shall arrive, I shall
endeavor to express the high sense I entertain of this
distinguished honor.")
tokens_remove(tokens(txt, remove_punct = TRUE), stopwords("english"))

# token_keep example: keep two-letter words
tokens_keep(tokens(txt, remove_punct = TRUE), "??")

tokens_split

Split tokens by a separator pattern

Description

Replaces tokens by multiple replacements consisting of elements split by a separator pattern, with
the option of retaining the separator. This function effectively reverses the operation of tokens_compound().

Usage

tokens_split(
  x,
  separator = " ",
  valuetype = c("fixed", "regex"),
  remove_separator = TRUE
)

Arguments

  x  a tokens object

  separator  a single-character pattern match by which tokens are separated
valuetype the type of pattern matching: "glob" for "glob"-style wildcard expressions; "regex" for regular expressions; or "fixed" for exact matching. See valuetype for details.

remove_separator if TRUE, remove separator from new tokens

Examples

# undo tokens_compound()
toks1 <- tokens("pork barrel is an idiomatic multi-word expression")
tokens_compound(toks1, phrase("pork barrel"))
tokens_compound(toks1, phrase("pork barrel")) %>%
tokens_split(separator = ".")

# similar to tokens(x, remove_hyphen = TRUE) but post-tokenization
toks2 <- tokens("UK-EU negotiation is not going anywhere as of 2018-12-24.")
tokens_split(toks2, separator = "-", remove_separator = FALSE)

tokens_subset Extract a subset of a tokens

description

Returns document subsets of a tokens that meet certain conditions, including direct logical operations on docvars (document-level variables). tokens_subset() functions identically to subset.data.frame(), using non-standard evaluation to evaluate conditions based on the docvars in the tokens.

Usage

tokens_subset(x, subset, drop_docid = TRUE, ...)

Arguments

x tokens object to be subsetted
subset logical expression indicating the documents to keep: missing values are taken as false
drop_docid if TRUE, docid for documents are removed as the result of subsetting.
... not used

Value

tokens object, with a subset of documents (and docvars) selected according to arguments

See Also

subset.data.frame()
Examples

corp <- corpus(c(d1 = "a b c d", d2 = "a a b e",
    d3 = "b b c e", d4 = "e e f a b"),
    docvars = data.frame(grp = c(1, 1, 2, 3)))
toks <- tokens(corp)
# selecting on a docvars condition
tokens_subset(toks, grp > 1)
# selecting on a supplied vector
tokens_subset(toks, c(TRUE, FALSE, TRUE, FALSE))

tokens_tolower  Convert the case of tokens

Description
tokens_tolower() and tokens_toupper() convert the features of a tokens object and re-index the types.

Usage
tokens_tolower(x, keep_acronyms = FALSE)
tokens_toupper(x)

Arguments
x the input object whose character/tokens/feature elements will be case-converted
keep_acronyms logical; if TRUE, do not lowercase any all-uppercase words (applies only to _tolower() functions)

Examples
# for a document-feature matrix
toks <- tokens(c(txt1 = "b A A", txt2 = "C C a b B"))
tokens_tolower(toks)
tokens_toupper(toks)
**tokens_wordstem**

*Stem the terms in an object*

### Description

Apply a stemmer to words. This is a wrapper to `wordStem` designed to allow this function to be called without loading the entire SnowballC package. `wordStem` uses Martin Porter's stemming algorithm and the C libstemmer library generated by Snowball.

### Usage

```r
tokens_wordstem(x, language = quanteda_options("language_stemmer"))
char_wordstem(x, language = quanteda_options("language_stemmer"))
dfm_wordstem(x, language = quanteda_options("language_stemmer"))
```

### Arguments

- **x**
  - a character, tokens, or dfm object whose word stems are to be removed. If tokenized texts, the tokenization must be word-based.

- **language**
  - the name of a recognized language, as returned by `getStemLanguages`, or a two- or three-letter ISO-639 code corresponding to one of these languages (see references for the list of codes).

### Value

- `tokens_wordstem` returns a `tokens` object whose word types have been stemmed.
- `char_wordstem` returns a `character` object whose word types have been stemmed.
- `dfm_wordstem` returns a `dfm` object whose word types (features) have been stemmed, and recombined to consolidate features made equivalent because of stemming.

### References

- [http://snowball.tartarus.org/](http://snowball.tartarus.org/)
- [http://www.iso.org/iso/home/standards/language_codes.htm](http://www.iso.org/iso/home/standards/language_codes.htm) for the ISO-639 language codes

### See Also

- `wordStem`
Examples

# example applied to tokens
txt <- c(one = "eating eater eaters eats ate",
         two = "taxing taxes taxed my tax return")
th <- tokens(txt)
tokens_wordstem(th)

# simple example
char_wordstem(c("win", "winning", "wins", "won", "winner"))

# example applied to a dfm
(origdfm <- dfm(tokens(txt)))
dfm_wordstem(origdfm)

topfeatures

Identify the most frequent features in a dfm

Description

List the most (or least) frequently occurring features in a dfm, either as a whole or separated by document.

Usage

```r
topfeatures(
  x,
  n = 10,
  decreasing = TRUE,
  scheme = c("count", "docfreq"),
  groups = NULL
)
```

Arguments

- `x` the object whose features will be returned
- `n` how many top features should be returned
- `decreasing` If TRUE, return the `n` most frequent features; otherwise return the `n` least frequent features
- `scheme` one of count for total feature frequency (within group if applicable), or docfreq for the document frequencies of features
- `groups` grouping variable for sampling, equal in length to the number of documents. This will be evaluated in the docvars data.frame, so that docvars may be referred to by name without quoting. This also changes previous behaviours for groups. See `news(Version >= "3.0", package = "quanteda")` for details.
Value

A named numeric vector of feature counts, where the names are the feature labels, or a list of these if groups is given.

Examples

dfmat1 <- corpus_subset(data_corpus_inaugural, Year > 1980) %>%
tokens(remove_punct = TRUE) %>%
dfm()
dfmat2 <- dfm_remove(dfmat1, stopwords("en"))

# most frequent features
topfeatures(dfmat1)
topfeatures(dfmat2)

# least frequent features
topfeatures(dfmat2, decreasing = FALSE)

# top features of individual documents
topfeatures(dfmat2, n = 5, groups = docnames(dfmat2))

# grouping by president last name
topfeatures(dfmat2, n = 5, groups = President)

# features by document frequencies
tail(topfeatures(dfmat1, scheme = "docfreq", n = 200))

---

types

Get word types from a tokens object

Description

Get unique types of tokens from a tokens object.

Usage

types(x)

Arguments

x a tokens object

See Also

featnames

Examples

toks <- tokens(data_corpus_inaugural)
types(toks)
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