

# Package ‘RcppCWB’

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**Type** Package

**Title** 'Rcpp' Bindings for the 'Corpus Workbench' ('CWB')

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**Description** 'Rcpp' Bindings for the C code of the 'Corpus Workbench' ('CWB'), an indexing and query engine to efficiently analyze large corpora (<<https://cwb.sourceforge.io>>). 'RcppCWB' is licensed

under the GNU GPL-3, in line with the GPL-

3 license of the 'CWB' (<<https://www.r-project.org/Licenses/GPL-3>>).

The 'CWB' relies on 'pcre2' (BSD license, see <<https://github.com/PCRE2Project/pcre2/blob/master/LICENCE.md>>)

and 'GLib' (LGPL license, see <<https://www.gnu.org/licenses/lgpl-3.0.en.html>>).

See the file LICENSE.note for further information. The package includes modified code of the 'rcqp' package (GPL-2, see <<https://cran.r-project.org/package=rcqp>>). The original work of the authors of the 'rcqp' package is acknowledged with great respect, and they are listed as authors of this package. To achieve cross-

platform portability (including Windows), using 'Rcpp' for wrapper code

is the approach used by 'RcppCWB'.

**License** GPL-3

**Encoding** UTF-8

**Copyright** For the copyrights for the 'Corpus Workbench' ('CWB') and acknowledgement of authorship, see file COPYRIGHTS.

**NeedsCompilation** yes

**SystemRequirements** GNU make, pcre2 (>= 10.00), GLib (>= 2.0.0). On Windows, no prior installations are necessary, as pre-built (i.e. cross-compiled) binaries of required libraries are downloaded from a GitHub repository (<<https://github.com/PolMine/libcl>>) during installation. On macOS, static libraries of Glib are downloaded (<<https://github.com/PolMine/libglib>>) if Glib is not present.

**Imports** Rcpp (>= 1.0.10), fs

**Suggests** knitr, testthat, rmarkdown

**LinkingTo** Rcpp

**Biarch** true

**VignetteBuilder** knitr

**URL** <https://github.com/PolMine/RcppCWB>

**BugReports** <https://github.com/PolMine/RcppCWB/issues>

**RoxygenNote** 7.3.2

**Collate** 'RcppCWB\_package.R' 'cl.R' 'cqp.R' 'cwb.R' 'checks.R'  
 'count.R' 'RcppExports.R' 'decode.R' 'cbow.R' 'region\_matrix.R'  
 'misc.R' 'zzz.R' 'xml.R'

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## Description

The RcppCWB package is a wrapper library to expose core functions of the Open Corpus Workbench (CWB). This includes the low-level functionality of the Corpus Library (CL) as well as capacities to use the query syntax of the Corpus Query Processor (CQP).

## The Idea Behind RcppCWB

The Open Corpus Workbench (CWB) is an indexing and querying engine popular in corpus-assisted research. Its core aim is to support working efficiently with large, structurally and linguistically annotated corpora. First of all, the CWB includes tools to index and compress corpora. Second, the Corpus Library (CL) offers low-level functionality to retrieve information from CWB indexed corpora. Third, the Corpus Query Processor (CQP) offers a syntax that allows to perform anything from simple to complex queries, using different annotation layers of corpora.

The CWB is a classical tool which has inspired a set of developments. A persisting advantage of the CWB is its mature, open source code base that is actively maintained by a community of developers. It is used as a robust and efficient backend for widely used tools such as TXM(<https://txm.gitpages.huma-num.fr/textometrie/>) or CQPweb (<https://cwb.sourceforge.io/cqpweb.php>). Its uncompromising C implementation guarantees speed and makes it well suited to be integrated with R at the same time.

The package RcppCWB is a follow-up on the `rcqp` package that has pioneered to expose CWB functionality from within R. Indeed, the `rcqp` package, published at CRAN in 2015, offers robust access

to CWB functionality. However, the "pure C" implementation of the `rcqp` package creates difficulties to make the package portable to Windows. The primary purpose of the `RcppCWB` package is to reimplement a wrapper library for the CWB using a design that makes it easier to achieve cross-platform portability.

Even though `RcppCWB` functions may be used directly, the package is designed to serve as an interface to CWB indexed corpora in packages with higher-level functionality. In this regard, `RcppCWB` is the backend of the `polmineR` package. It is deliberately open to be used in other contexts. The package may stimulate using linguistically annotated, indexed and compressed corpora on all platforms. The paradigm of working with text as linguistic data may benefit from `RcppCWB`.

## **Implementation**

When building the package, the first step is to compile the relevant parts of the CWB on Linux and macOS machines. On Windows, cross-compiled binaries are downloaded from a GitHub repository of the PolMine Project (<https://github.com/PolMine/libc1>). Second, Rcpp wrappers are compiled and make the relevant functions of the Corpus Library and CQP accessible. In addition to genuine CWB functions, `RcppCWB` offers a set of higher level functions implemented using Rcpp for common performance critical tasks.

## **Getting Started with RcppCWB**

To understand the data storage model of the CWB, in particular the notions of positional and structural attributes (s- and p-attributes), the vignette of the `rcqp` package is a very good starting point (see references).

The CWB 'Corpus Encoding Tutorial' explains how to create your own corpus, the 'CQP Query Language Tutorial' introduces the syntax of CQP (see references).

The `RcppCWB` package includes a sample corpus (REUTERS, the data also included in the `tm` package). The examples in the documentation of the functions may be a good starting point to understand how to use `RcppCWB`.

## **Digging Deeper**

The original paper of Christ (1994) explains the design choices of the CWB. The indexing and compression techniques of the CWB (Huffman coding) are explained in Witten et al. (1999).

## **Acknowledgements**

The work of the all developers of the CWB is gratefully acknowledged. There is a particular intellectual debt to Bernard Desgraupes and Sylvain Loiseau, and the `rcqp` package they developed as the original R wrapper to expose the functionality of the CWB.

## **Author(s)**

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## References

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- Open Corpus Workbench (<https://cwb.sourceforge.io>)
- Witten, I.H.; Moffat, A.; Bell, T.C. (1999). Managing Gigabytes. Morgan Kaufmann Publishing, San Francisco, 2nd edition.

## See Also

Useful links:

- <https://github.com/PolMine/RcppCWB>
- Report bugs at <https://github.com/PolMine/RcppCWB/issues>

## Examples

```
# functions of the corpus library (starting with cl) expose the low-level
# access to the CWB corpus library (CL)

ids <- cl_cpos2id("REUTERS", cpos = 1:20, p_attribute = "word", registry = get_tmp_registry())
tokens <- cl_id2str("REUTERS", id = ids, p_attribute = "word", registry = get_tmp_registry())
print(paste(tokens, collapse = " "))

# To use the corpus query processor (CQP) and its syntax, it is necessary first
# to initialize CQP (example: get concordances of 'oil')

cqp_query("REUTERS", query = '[]{5} "oil" []{5}')
cpos_matrix <- cqp_dump_subcorpus("REUTERS")
concordances_oil <- apply(
  cpos_matrix, 1,
  function(row){
    ids <- cl_cpos2id("REUTERS", p_attribute = "word", cpos = row[1]:row[2], get_tmp_registry())
    tokens <- cl_id2str("REUTERS", p_attribute = "word", id = ids, get_tmp_registry())
    paste(tokens, collapse = " ")
  }
)
```

**attribute\_size***Rcpp wrappers for CWB Corpus Library functions***Description**

Rcpp wrappers for CWB Corpus Library functions

**Usage**

```
attribute_size(corpus, attribute, attribute_type, registry)

cpos2str(corpus, p_attribute, registry, cpos)

cpos2id(corpus, p_attribute, registry, cpos)

struc2cpos(corpus, s_attribute, registry, struc)

id2str(corpus, p_attribute, registry, id)
```

**Arguments**

<code>corpus</code>	The ID of a CWB corpus.
<code>attribute</code>	Either a positional, or a structural attribute.
<code>attribute_type</code>	Either "p" (positional attribute) or "s" (structural attribute).
<code>registry</code>	Path to the corpus registry.
<code>p_attribute</code>	A positional attribute.
<code>cpos</code>	An integer vector of corpus positions.
<code>s_attribute</code>	A structural attribute.
<code>struc</code>	An integer value with struc.
<code>id</code>	An integer vector with token ids.

**check***Check Input to Rcpp Functions.***Description**

A set of functions to check whether the input values to the Rcpp wrappers for the C functions of the Corpus Workbench potentially causing crashes are valid. These auxiliary functions are called by the `cl_` and `cqp_` functions.

**Usage**

```
check_registry(registry)

check_corpus(corpus, registry, cl = TRUE, cqp = TRUE)

check_s_attribute(
  s_attribute,
  corpus,
  registry = Sys.getenv("CORPUS_REGISTRY")
)

check_p_attribute(
  p_attribute,
  corpus,
  registry = Sys.getenv("CORPUS_REGISTRY")
)

check_strucs(corpus, s_attribute, strucs, registry)

check_region_matrix(region_matrix)

check_query(query)

check_cpos(
  corpus,
  p_attribute = "word",
  cpos,
  registry = Sys.getenv("CORPUS_REGISTRY")
)

check_id(corpus, p_attribute, id, registry = Sys.getenv("CORPUS_REGISTRY"))
```

**Arguments**

registry	path to registry directory
corpus	name of a CWB corpus
cl	A logical value, whether CL availability of corpus is required for positive result.
cqp	A logical value, whether CQP availability of corpus is required for positive result.
s_attribute	a structural attribute
p_attribute	a positional attribute
strucs	strucs (indices of structural attributes)
region_matrix	a region matrix
query	a CQP query
cpos	vector of corpus positions

id	id (encoded p-attribute), integer value
----	-----------------------------------------

---

### check\_pkg\_registry\_files

*Check Paths in Registry Files*

---

#### Description

Check Paths in Registry Files

#### Usage

```
check_pkg_registry_files(pkg = system.file(package = "RcppCWB"), set = FALSE)
```

#### Arguments

pkg	Full path to package directory
set	Logical, whether

#### Value

Logical value, whether home directories are set correctly.

---

CL: p\_attributes

*Using Positional Attributes.*

---

#### Description

CWB indexed corpora store the text of a corpus as numbers: Every token in the token stream of the corpus is identified by a unique corpus position. The string value of every token is identified by a unique integer id. The corpus library (CL) offers a set of functions to make the transitions between corpus positions, token ids, and the character string of tokens.

#### Usage

```
cl_cpos2str(
  corpus,
  p_attribute,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  cpos
)
cl_cpos2id(corpus, p_attribute, registry = Sys.getenv("CORPUS_REGISTRY"), cpos)
cl_id2str(corpus, p_attribute, registry = Sys.getenv("CORPUS_REGISTRY"), id)
```

```

cl_regex2id(
  corpus,
  p_attribute,
  regex,
  registry = Sys.getenv("CORPUS_REGISTRY")
)

cl_str2id(corpus, p_attribute, str, registry = Sys.getenv("CORPUS_REGISTRY"))

cl_id2freq(corpus, p_attribute, id, registry = Sys.getenv("CORPUS_REGISTRY"))

cl_id2cpos(corpus, p_attribute, id, registry = Sys.getenv("CORPUS_REGISTRY"))

```

## Arguments

corpus	name of a CWB corpus (upper case)
p_attribute	a p-attribute (positional attribute)
registry	path to the registry directory, defaults to the value of the environment variable CORPUS_REGISTRY
cpos	corpus positions (integer vector)
id	id of a token
regex	a regular expression
str	a character string

## Examples

```

# registry directory and cpos_total will be needed in examples
cpos_total <- cl_attribute_size(
  corpus = "REUTERS", attribute = "word",
  attribute_type = "p", registry = get_tmp_registry()
)

# decode the token stream of the corpus (the quick way)
token_stream_str <- cl_cpos2str(
  corpus = "REUTERS", p_attribute = "word",
  cpos = seq.int(from = 0, to = cpos_total - 1),
  registry = get_tmp_registry()
)

# decode the token stream (cpos2id first, then id2str)
token_stream_ids <- cl_cpos2id(
  corpus = "REUTERS", p_attribute = "word",
  cpos = seq.int(from = 0, to = cpos_total - 1),
  registry = get_tmp_registry()
)
token_stream_str <- cl_id2str(
  corpus = "REUTERS", p_attribute = "word",
  id = token_stream_ids, registry = get_tmp_registry()
)

```

```

)
# get corpus positions of a token
token_to_get <- "oil"
id_oil <- cl_str2id(
  corpus = "REUTERS", p_attribute = "word",
  str = token_to_get, registry = get_tmp_registry()
)
cpos_oil <- cl_id2cpos <- cl_id2cpos(
  corpus = "REUTERS", p_attribute = "word",
  id = id_oil, registry = get_tmp_registry()
)
length(cpos_oil) # needs to be the same as oil_freq

# use regular expressions
ids <- cl_regex2id(
  corpus = "REUTERS", p_attribute = "word",
  regex = "M.*", registry = get_tmp_registry()
)
m_words <- cl_id2str(
  corpus = "REUTERS", p_attribute = "word",
  id = ids, registry = get_tmp_registry()
)

```

CL: s\_attributes      *Using Structural Attributes.*

## Description

Structural attributes store the metadata of texts in a CWB corpus and/or any kind of annotation of a region of text. The fundamental unit are so-called strucs, i.e. indices of regions identified by a left and a right corpus position. The corpus library (CL) offers a set of functions to make the translations between corpus positions (cpos) and strucs (struc).

## Usage

```

cl_cpos2struc(
  corpus,
  s_attribute,
  cpos,
  registry = Sys.getenv("CORPUS_REGISTRY")
)

```

```

cl_struct2cpos(
  corpus,
  s_attribute,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  struc
)
cl_struct2str(
  corpus,
  s_attribute,
  struc,
  registry = Sys.getenv("CORPUS_REGISTRY")
)
cl_cpos2lbound(corpus, s_attribute, cpos, registry = NULL)
cl_cpos2rbound(corpus, s_attribute, cpos, registry = NULL)

```

## Arguments

<code>corpus</code>	name of a CWB corpus (upper case)
<code>s_attribute</code>	name of structural attribute (character vector)
<code>cpos</code>	An integer vector with corpus positions.
<code>registry</code>	path to the registry directory, defaults to the value of the environment variable <code>CORPUS_REGISTRY</code>
<code>struc</code>	a struc identifying a region

## Details

`cl_cpos2rbound()` and `cl_cpos2lbound()` return NA for values of `cpos` that are outside a struc for the structural attribute given.

## Examples

```

# get metadata for matches of token
# scenario: id of the texts with occurrence of 'oil'
token_to_get <- "oil"
token_id <- cl_struct2id("REUTERS", p_attribute = "word", str = "oil", get_tmp_registry())
token_cpos <- cl_struct2cpos("REUTERS", p_attribute = "word", id = token_id, get_tmp_registry())
strucs <- cl_struct2struc("REUTERS", s_attribute = "id", cpos = token_cpos, get_tmp_registry())
strucs_unique <- unique(strucs)
text_ids <- cl_struct2str("REUTERS", s_attribute = "id", struc = strucs_unique, get_tmp_registry())

# get the full text of the first text with match for 'oil'
left_cpos <- cl_cpos2lbound(
  "REUTERS", s_attribute = "id",
  cpos = min(token_cpos),
  registry = get_tmp_registry()
)

```

```

right_cpos <- cl_cpos2rbound(
  "REUTERS",
  s_attribute = "id",
  cpos = min(token_cpos),
  registry = get_tmp_registry()
)
txt <- cl_cpos2str(
  "REUTERS", p_attribute = "word",
  cpos = left_cpos:right_cpos,
  registry = get_tmp_registry()
)
fulltext <- paste(txt, collapse = " ")

# alternativ approach to achieve same result
first_struc_match_oil <- cl_cpos2struc(
  "REUTERS", s_attribute = "id",
  cpos = min(token_cpos),
  registry = get_tmp_registry()
)
cpos_struc <- cl_struc2cpos(
  "REUTERS", s_attribute = "id",
  struc = first_struc_match_oil,
  registry = get_tmp_registry()
)
txt <- cl_cpos2str(
  "REUTERS",
  p_attribute = "word",
  cpos = cpos_struc[1]:cpos_struc[2],
  registry = get_tmp_registry()
)
fulltext <- paste(txt, collapse = " ")

```

**cl\_attribute\_size**      *Get Attribute Size (of Positional/Structural Attribute).*

## Description

Use `cl_attribute_size()` to get the total number of values of a positional attribute (param `attribute_type = "p"`), or structural attribute (param `attribute_type = "s"`). Note that indices are zero-based, i.e. the maximum position of a positional / structural attribute is attribute size minus 1 (see examples).

## Usage

```

cl_attribute_size(
  corpus,
  attribute,
  attribute_type,
  registry = Sys.getenv("CORPUS_REGISTRY")
)

```

## Arguments

corpus	name of a CWB corpus (upper case)
attribute	name of a p- or s-attribute
attribute_type	either "p" or "s", for structural/positional attribute
registry	path to the registry directory, defaults to the value of the environment variable CORPUS_REGISTRY

## Examples

```

token_no <- cl_attribute_size(
  "REUTERS",
  attribute = "word",
  attribute_type = "p",
  registry = get_tmp_registry()
)
corpus_positions <- seq.int(from = 0, to = token_no - 1)
cl_cpos2id(
  "REUTERS",
  "word",
  cpos = corpus_positions,
  registry = get_tmp_registry()
)

places_no <- cl_attribute_size(
  "REUTERS",
  attribute = "places",
  attribute_type = "s",
  registry = get_tmp_registry()
)
strucs <- seq.int(from = 0, to = places_no - 1)
cl_struc2str(
  "REUTERS",
  "places",
  struc = strucs,
  registry = get_tmp_registry()
)

```

cl\_charset\_name      *Get charset of a corpus.*

## Description

The encoding of a corpus is declared in the registry file (corpus property "charset"). Once a corpus is loaded, this information is available without parsing the registry file again and again. The cl\_charset\_name offers a quick access to this information.

## Usage

```
cl_charset_name(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))
```

### Arguments

<code>corpus</code>	Name of a CWB corpus (upper case).
<code>registry</code>	Path to the registry directory, defaults to the value of the environment variable <code>CORPUS_REGISTRY</code>

### Examples

```
cl_charset_name(
  corpus = "REUTERS",
  registry = system.file(package = "RcppCWB", "extdata", "cwb", "registry")
)
```

`cl_delete_corpus`      *Drop loaded corpus.*

### Description

Remove a corpus from the list of loaded corpora of the corpus library (CL).

### Usage

```
cl_delete_corpus(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))
```

### Arguments

<code>corpus</code>	name of a CWB corpus (upper case)
<code>registry</code>	path to the registry directory, defaults to the value of the environment variable <code>CORPUS_REGISTRY</code>

### Details

The corpus library (CL) internally maintains a list of corpora including information on positional and structural attributes so that the registry file needs not be parsed again and again. However, when an attribute has been added to the corpus, it will not yet be visible, because it is not part of the data that has been loaded. The `cl_delete_corpus` function exposes a CL function named identically, to force reloading the corpus (after it has been deleted), which will include parsing an updated registry file.

### Value

An integer value 1 is returned invisibly if a previously loaded corpus has been deleted, or 0 if the corpus has not been loaded and has not been deleted.

### Examples

```
cl_attribute_size("UNGA", attribute = "word", attribute_type = "p")
corpus_is_loaded("UNGA")
cl_delete_corpus("UNGA")
corpus_is_loaded("UNGA")
```

---

cl_find_corpus	<i>Load corpus.</i>
----------------	---------------------

---

### Description

Load corpus.

### Usage

```
cl_find_corpus(corpus, registry)
```

### Arguments

corpus	name of a CWB corpus (upper case)
registry	path to the registry directory, defaults to the value of the environment variable CORPUS_REGISTRY

### Value

A externalptr referencing the C representation of the corpus.

---

cl_lexicon_size	<i>Get Lexicon Size.</i>
-----------------	--------------------------

---

### Description

Get the total number of unique tokens/ids of a positional attribute. Note that token ids are zero-based, i.e. when iterating through tokens, start at 0, the maximum will be cl\_lexicon\_size() minus 1.

### Usage

```
cl_lexicon_size(corpus, p_attribute, registry = Sys.getenv("CORPUS_REGISTRY"))
```

### Arguments

corpus	name of a CWB corpus (upper case)
p_attribute	name of positional attribute
registry	path to the registry directory, defaults to the value of the environment variable CORPUS_REGISTRY

**Examples**

```
lexicon_size <- cl_lexicon_size(
  "REUTERS",
  p_attribute = "word",
  registry = get_tmp_registry()
)

token_ids <- seq.int(from = 0, to = lexicon_size - 1)
cl_id2str(
  "REUTERS",
  p_attribute = "word",
  id = token_ids,
  registry = get_tmp_registry()
)
```

**cl\_list\_corpora**      *Show CL corpora*

**Description**

Show CL corpora

**Usage**

```
cl_list_corpora()
```

**Value**

A character vector.

**Examples**

```
cl_list_corpora()
```

**cl\_load\_corpus**      *Load corpus*

**Description**

Load corpus

**Usage**

```
cl_load_corpus(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))
```

**Arguments**

- |          |                                                            |
|----------|------------------------------------------------------------|
| corpus   | A length-one character vector with the corpus ID.          |
| registry | A length-one character vector with the registry directory. |

**Value**

TRUE if corpus could be loaded and FALSE if not.

**Examples**

```
cl_load_corpus("REUTERS")
```

---

cl_rework	<i>Low-level CL access.</i>
-----------	-----------------------------

---

**Description**

Wrappers for CWB Corpus Library functions suited for writing performance code.

**Usage**

```
s_attr(corpus, s_attribute, registry)  
p_attr(corpus, p_attribute, registry)  
p_attr_size(p_attr)  
s_attr_size(s_attr)  
p_attr_lexicon_size(p_attr)  
cpos_to_struc(s_attr, cpos)  
cpos_to_str(p_attr, cpos)  
cpos_to_id(p_attr, cpos)  
struc_to_cpos(s_attr, struc)  
struc_to_str(s_attr, struc)  
regex_to_id(p_attr, regex)  
str_to_id(p_attr, str)  
id_to_freq(p_attr, id)
```

```

id_to_cpos(p_attr, id)

cpos_to_lbound(s_attr, cpos)

cpos_to_rbound(s_attr, cpos)

```

### Arguments

<code>corpus</code>	ID of a CWB corpus (length-one character vector).
<code>s_attribute</code>	A structural attribute (length-one character vector).
<code>registry</code>	Registry directory.
<code>p_attribute</code>	A positional attribute (length-one character vector).
<code>p_attr</code>	A <code>externalptr</code> referencing a p-attribute.
<code>s_attr</code>	A <code>externalptr</code> referencing a p-attribute.
<code>cpos</code>	An <code>integer</code> vector of corpus positions.
<code>struc</code>	A length-one <code>integer</code> vector with a struc.
<code>regex</code>	A regular expression.
<code>str</code>	A character vector.
<code>id</code>	An <code>integer</code> vector with token ids.

### Details

The default `cl_*` R wrappers for the functions of the CWB Corpus Library involve a lookup of a corpus and its p- or s-attributes (using the corpus ID, registry and attribute indicated by length-one character vectors) every time one of these functions is called. It is more efficient looking up an attribute only once. This set of functions passes "externalptr" classes to reference attributes that have been looked up. A relevant scenario is writing functions with a C++ implementation that are compiled and linked using `Rcpp::cppFunction()` or `Rcpp::sourceCpp()`

### Examples

```

library(Rcpp)

cppFunction(
  'Rcpp::StringVector get_str(
    SEXP corpus,
    SEXP p_attribute,
    SEXP registry,
    Rcpp::IntegerVector cpos
  ){
    SEXP attr;
    Rcpp::StringVector result;
    attr = RcppCWB::p_attr(corpus, p_attribute, registry);
    result = RcppCWB::cpos_to_str(attr, cpos);
    return(result);
  }',
  depends = "RcppCWB"
)

```

```
)
result <- get_str("REUTERS", "word", RcppCWB::get_tmp_registry(), 0:50)
```

**cl\_struct\_values***Check whether structural attribute has values***Description**

Structural attributes do not necessarily have values, structural attributes (such as annotations of sentences or paragraphs) may just define regions of corpus positions. Use this function to test whether an attribute has values.

**Usage**

```
cl_struct_values(corpus, s_attribute, registry = Sys.getenv("CORPUS_REGISTRY"))
```

**Arguments**

- |                          |                                                               |
|--------------------------|---------------------------------------------------------------|
| <code>corpus</code>      | Corpus ID, a length-one character vector.                     |
| <code>s_attribute</code> | Structural attribute to check, a length-one character vector. |
| <code>registry</code>    | The registry directory of the corpus.                         |

**Value**

TRUE if the attribute has values and FALSE if not. NA if the structural attribute is not available.

**Examples**

```
cl_struct_values("REUTERS", "places") # TRUE - attribute has values
cl_struct_values("REUTERS", "date") # NA - attribute does not exist
```

**corpus\_data\_dir***Get information from registry file***Description**

Extract information from the internal C representation of registry data.

**Usage**

```
corpus_data_dir(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

corpus_info_file(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

corpus_full_name(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

corpus_p_attributes(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

corpus_s_attributes(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

corpus_properties(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

corpus_property(corpus, registry = Sys.getenv("CORPUS_REGISTRY"), property)

corpus_registry_dir(corpus)
```

**Arguments**

- |                       |                                                            |
|-----------------------|------------------------------------------------------------|
| <code>corpus</code>   | A length-one character vector with the corpus ID.          |
| <code>registry</code> | A length-one character vector with the registry directory. |
| <code>property</code> | A corpus property defined in the registry file (.          |

**Details**

`corpus_data_dir()` will return the data directory (class `fs_path`) where the binary files of a corpus are kept (a directory also known as 'home' directory).

`corpus_info_file()` will return the path to the info file for a corpus (class `fs_path` object). If info file does not exist or INFO line is missing in the registry file, NA is returned.

`corpus_full_name()` will return the full name of the corpus defined in the registry file.

`corpus_p_attributes()` returns a character vector with the positional attributes of a corpus.

`corpus_s_attributes()` returns a character vector with the structural attributes of a corpus.

`corpus_properties()` returns a character vector with the corpus properties defined in the registry file. If the corpus cannot be located, NA is returned.

`corpus_property()` returns the value of a corpus property defined in the registry file, or NA if the corpus does not exist, is not loaded or if the property requested is undefined.

`corpus_get_registry()` will extract the registry directory with the registry file defining a corpus from the internal C representation of loaded corpora. The character vector that is returned may be > 1 if there are several corpora with the same id defined in registry files in different (registry) directories. If the corpus is not found, NA is returned.

**Examples**

```
corpus_data_dir("REUTERS", registry = get_tmp_registry())
corpus_info_file("REUTERS", registry = get_tmp_registry())
corpus_full_name("REUTERS", registry = get_tmp_registry())
```

```

corpus_p_attributes("REUTERS", registry = get_tmp_registry())
corpus_s_attributes("REUTERS", registry = get_tmp_registry())
corpus_properties("REUTERS", registry = get_tmp_registry())
corpus_property(
  "REUTERS",
  registry = get_tmp_registry(),
  property = "language"
)
corpus_registry_dir("REUTERS")
corpus_registry_dir("FOO") # NA returned

```

<code>corpus_is_loaded</code>	<i>Check whether corpus is loaded</i>
-------------------------------	---------------------------------------

### Description

Check whether corpus is loaded

### Usage

```
corpus_is_loaded(corpus, registry = Sys.getenv("CORPUS_REGISTRY"))
```

### Arguments

<code>corpus</code>	A length-one character vector with the corpus ID.
<code>registry</code>	A length-one character vector with the registry directory.

### Value

TRUE if corpus is loaded and FALSE if not.

<code>cqp_initialize</code>	<i>Initialize Corpus Query Processor (CQP).</i>
-----------------------------	-------------------------------------------------

### Description

CQP needs to know where to look for CWB indexed corpora. To initialize CQP, call `cqp_initialize`. To reset the registry, use the function `cqp_reset_registry`. To get the registry used by CQP, use `cqp_get_registry`. To get the initialization status, use `cqp_is_initialized`

**Usage**

```
cqp_initialize(registry = Sys.getenv("CORPUS_REGISTRY"))

cqp_is_initialized()

cqp_verbosity(silent, verbose)

cqp_get_registry()

cqp_reset_registry(registry = Sys.getenv("CORPUS_REGISTRY"))

cqp_load_corpus(corpus, registry)
```

**Arguments**

<code>registry</code>	the registry directory
<code>silent</code>	A single logical value, whether to be silent and suppress CQP messages (TRUE), or not (FALSE).
<code>verbose</code>	A single logical value, whether to show verbose parser output (TRUE) or not (FALSE).
<code>corpus</code>	ID of a CWB corpus (length-one character).

**Details**

`cqp_load_corpus` will return a logical value - TRUE if corpus has been loaded successfully, FALSE if not.

**Author(s)**

Andreas Blaette, Bernard Desgraupes, Sylvain Loiseau

**Examples**

```
cqp_is_initialized() # check initialization status
if (!cqp_is_initialized()) cqp_initialize()
cqp_is_initialized() # check initialization status (TRUE now?)
cqp_get_registry() # get registry dir used by CQP
cqp_list_corpora() # get list of corpora
```

`cqp_list_corpora`      *List Available CWB Corpora.*

**Description**

List the corpora described by the registry files in the registry directory that is currently set.

**Usage**

```
cqp_list_corpora()
```

**Author(s)**

Andreas Blaette, Bernard Desgraupes, Sylvain Loiseau

**Examples**

```
cqp_list_corpora()
```

---

cqp\_query

*Execute CQP Query and Retrieve Results.*

---

**Description**

Using CQP queries requires a two-step procedure: At first, you execute a query using `cqp_query`. Then, `cqp_dump_subcorpus` will return a matrix with the regions of the matches for the query.

**Usage**

```
cqp_query(corpus, query, subcorpus = "QUERY")  
cqp_dump_subcorpus(corpus, subcorpus = "QUERY")  
cqp_subcorpus_size(corpus, subcorpus = "QUERY")  
cqp_list_subcorpora(corpus)  
cqp_drop_subcorpus(corpus)
```

**Arguments**

corpus	a CWB corpus
query	a CQP query
subcorpus	subcorpus name

**Details**

The `cqp_query` function executes a CQP query. The `cqp_subcorpus_size` function returns the number of matches for the CQP query. The `cqp_dump_subcorpus` function will return a two-column matrix with the left and right corpus positions of the matches for the CQP query.

**Author(s)**

Andreas Blaette, Bernard Desgraupes, Sylvain Loiseau

## References

Evert, S. 2005. The CQP Query Language Tutorial. Available online at [https://cwb.sourceforge.io/files/CWB\\_Encoding\\_Tutorial.pdf](https://cwb.sourceforge.io/files/CWB_Encoding_Tutorial.pdf)

## Examples

```
cqp_query(corpus = "REUTERS", query = '"oil";')
cqp_subcorpus_size("REUTERS")
cqp_dump_subcorpus("REUTERS")

cqp_query(corpus = "REUTERS", query = '"crude" "oil";')
cqp_subcorpus_size("REUTERS", subcorpus = "QUERY")
cqp_dump_subcorpus("REUTERS")
```

cwb\_charsets

*Character sets supported by CWB*

## Description

The function returns a character vector with characters sets (charsets) supported by the Corpus Workbench (CWB). The vector is derived from the the CorpusCharset object defined in the header file of the corpus library (CL).

## Usage

```
cwb_charsets()
```

## Details

Early versions of the CWB were developed for "latin1", "utf8" support has been introduced with CWB v3.2. Note that RcppCWB is tested only for "latin1" and "utf8" and that R uses "UTF-8" rather than utf8" (CWB) by convention.

## Examples

```
cwb_charsets()
```

## Description

Wrappers for the CWB tools cwb-makeall, cwb-huffcode and cwb-compress-rdx. Unlike the 'original' command line tools, these wrappers will always perform a specific indexing/compression step on one positional attribute, and produce all components.

## Usage

```
cwb_encode(  
  corpus,  
  registry = Sys.getenv("CORPUS_REGISTRY"),  
  data_dir,  
  vrt_dir,  
  encoding = "utf8",  
  p_attributes = c("word", "pos", "lemma"),  
  s_attributes = list(),  
  skip_blank_lines = TRUE,  
  strip_whitespace = TRUE,  
  xml = TRUE,  
  quietly = FALSE,  
  verbose = FALSE  
)  
  
cwb_makeall(  
  corpus,  
  p_attribute,  
  registry = Sys.getenv("CORPUS_REGISTRY"),  
  quietly = FALSE,  
  logfile  
)  
  
cwb_huffcode(  
  corpus,  
  p_attribute,  
  registry = Sys.getenv("CORPUS_REGISTRY"),  
  quietly = FALSE,  
  logfile,  
  delete = TRUE  
)  
  
cwb_compress_rdx(  
  corpus,  
  p_attribute,  
  registry = Sys.getenv("CORPUS_REGISTRY"),
```

```

quietly = FALSE,
logfile,
delete = TRUE
)

```

## Arguments

corpus	Name of a CWB corpus (upper case).
registry	Path to the registry directory, defaults to the value of the environment variable CORPUS_REGISTRY.
data_dir	The data directory where cwb_encode will save the binary files of the indexed corpus. Tilde expansion is performed on data_dir using path.expand() to avoid a crash.
vrt_dir	Directory with input corpus files (verticalised format / file ending *.vrt). Tilde expansion is performed on vrt_dir using path.expand() to avoid a crash.
encoding	The encoding of the files to be encoded. Needs to be an encoding supported by CWB, see cwb_charsets(). "UTF-8" is taken as "utf8". Defaults to "utf8" (recommended charset).
p_attributes	Positional attributes (p-attributes) to be declared.
s_attributes	A list of named character vectors to declare structural attributes that shall be encoded. The names of the list are the XML elements present in the corpus. Character vectors making up the list declare the attributes that include the metadata of regions. To declare a structural attribute without annotations, provide a zero-length character vector using character() - see examples.
skip_blank_lines	A logical value, whether to skip blank lines in the input.
strip_whitespace	A logical value, whether to strip whitespace from tokens
xml	A logical value, whether input is XML.
quietly	A logical value, whether to turn off messages (including warnings).
verbose	A logical value, whether to show progress information (counter of tokens processed).
p_attribute	Name of p-attribute.
logfile	Redirect messages of cwb_makeall(), cwb_huffcode() or cwb_compress_rdx() to this file. Requires that quietly is TRUE.
delete	A logical value, whether to remove redundant file (p_attribute).corpus after compression.

## Details

Running cwb\_huffcode() and cwb\_compress\_rdx() is optional. Corpora can be fully used without compression. It is recommended when reducing the size of corpus data has relevant benefits, e.g. for sharing data. On Windows, compression is not stable and not recommended. A respective warning is issued when running cwb\_huffcode() and cwb\_compress\_rdx() on Windows.

## Examples

```

data_dir <- file.path(tempdir(), "bt_data_dir")
dir.create(data_dir)

cwb_encode(
  corpus = "BTMIN",
  registry = Sys.getenv("CORPUS_REGISTRY"),
  vrt_dir = system.file(package = "RcppCWB", "extdata", "vrt"),
  data_dir = data_dir,
  p_attributes = c("word", "pos", "lemma"),
  s_attributes = list(
    plenary_protocol = c(
      "lp", "protocol_no", "date", "year", "birthday", "version",
      "url", "filetype"
    ),
    speaker = c(
      "id", "type", "lp", "protocol_no", "date", "year", "ai_no", "ai_id",
      "ai_type", "who", "name", "parliamentary_group", "party", "role"
    ),
    p = character()
  )
)

unlink(data_dir)
unlink(file.path(Sys.getenv("CORPUS_REGISTRY"), "btmin"))
# The package includes an 'unfinished' corpus of debates in the UN General
# Assembly ("UNGA"), i.e. it does not yet include the reverse index, and it
# is not compressed.
#
# The first step in the following example is to copy the raw
# corpus to a temporary place.

home_dir <- system.file(
  package = "RcppCWB",
  "extdata", "cwb", "indexed_corpora", "unga"
)

tmp_data_dir <- file.path(tempdir(), "indexed_corpora")
tmp_unga_dir <- file.path(tmp_data_dir, "unga2")
if (!file.exists(tmp_data_dir)) dir.create(tmp_data_dir)
if (!file.exists(tmp_unga_dir)){
  dir.create(tmp_unga_dir)
} else {
  file.remove(list.files(tmp_unga_dir, full.names = TRUE))
}

regfile <- readLines(
  system.file(package = "RcppCWB", "extdata", "cwb", "registry", "unga")
)
regfile[grep("^HOME", regfile)] <- sprintf('HOME "%s"', tmp_unga_dir)
regfile[grep("^ID", regfile)] <- "ID unga2"
writeLines(text = regfile, con = file.path(get_tmp_registry(), "unga2"))

```

```

for (x in list.files(home_dir, full.names = TRUE)){
  file.copy(from = x, to = tmp_unga_dir)
}

# perform cwb_makeall (equivalent to cwb-makeall command line utility)
cwb_makeall(
  corpus = "UNGA2",
  p_attribute = "word",
  registry = get_tmp_registry()
)
cl_load_corpus("UNGA2", registry = get_tmp_registry())
cqp_load_corpus("UNGA2", registry = get_tmp_registry())

# see whether it works
ids_sentence_1 <- cl_cpos2id(
  corpus = "UNGA2", p_attribute = "word", registry = get_tmp_registry(),
  cpos = 0:83
)
tokens_sentence_1 <- cl_id2str(
  corpus = "UNGA2", p_attribute = "word",
  registry = get_tmp_registry(), id = ids_sentence_1
)
sentence <- gsub(
  "\\\\s+([\\\\.,])",
  "\\\\1",
  paste(tokens_sentence_1, collapse = " "))
)

# perform cwb_huffcode (equivalent to cwb-makeall command line utility)
cwb_huffcode(
  corpus = "UNGA2",
  p_attribute = "word",
  registry = get_tmp_registry()
)
cwb_compress_rdx(
  corpus = "UNGA2",
  p_attribute = "word",
  registry = get_tmp_registry()
)

```

**cwb\_version***Get CWB version*

## Description

Get the CWB version used and available when compiling the source code.

## Usage

```
cwb_version()
```

**Value**

A numeric\_version object.

**Examples**

```
cwb_version()
```

get_cbow_matrix	<i>Get CBOW Matrix.</i>
-----------------	-------------------------

**Description**

Get matrix with moving windows. Negative integer values indicate absence of a token at the respective position.

**Usage**

```
get_cbow_matrix(
  corpus,
  p_attribute,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  matrix,
  window
)
```

**Arguments**

corpus	a CWB corpus
p_attribute	a positional attribute
registry	the registry directory
matrix	a matrix
window	window size

**Examples**

```
m <- get_region_matrix(
  corpus = "REUTERS", s_attribute = "places",
  strucs = 0L:5L, registry = get_tmp_registry()
)
windowsize <- 3L
m2 <- get_cbow_matrix(
  corpus = "REUTERS", p_attribute = "word",
  registry = get_tmp_registry(), matrix = m, window = windowsize
)
colnames(m2) <- c(-windowsize:-1, "node", 1:windowsize)
```

`get_count_vector`      *Get Vector with Counts for Positional Attribute.*

## Description

The return value is an integer vector. The length of the vector is the number of unique tokens in the corpus / the number of unique ids. The order of the counts corresponds to the number of ids.

## Usage

```
get_count_vector(corpus, p_attribute, registry = Sys.getenv("CORPUS_REGISTRY"))
```

## Arguments

<code>corpus</code>	a CWB corpus
<code>p_attribute</code>	a positional attribute
<code>registry</code>	registry directory

## Value

an integer vector

## Examples

```
y <- get_count_vector(
  corpus = "REUTERS", p_attribute = "word",
  registry = get_tmp_registry()
)
df <- data.frame(token_id = 0:(length(y) - 1), count = y)
df[["token"]] <- cl_id2str(
  "REUTERS", p_attribute = "word",
  id = df[["token_id"]], registry = get_tmp_registry()
)
df <- df[,c("token", "token_id", "count")] # reorder columns
df <- df[order(df[["count"]]), decreasing = TRUE,]
head(df)
```

`get_pkg_registry`      *Get Registry Directory Within Package*

## Description

Get Registry Directory Within Package

## Usage

```
get_pkg_registry(pkgname = "RcppCWB")
```

**Arguments**

<code>pkgname</code>	Name of package (character vector)
----------------------	------------------------------------

<code>get_region_matrix</code>	<i>Get Matrix with Regions for Strucs.</i>
--------------------------------	--------------------------------------------

**Description**

The return value is an integer matrix with the left and right corpus positions of the strucs in columns one and two, respectively. For negative struc values in the input vector, the matrix reports NA values.

**Usage**

```
get_region_matrix(
  corpus,
  s_attribute,
  strucs,
  registry = Sys.getenv("CORPUS_REGISTRY")
)
```

**Arguments**

<code>corpus</code>	A CWB corpus (length-one character vector).
<code>s_attribute</code>	A structural attribute (length-one character vector).
<code>strucs</code>	Integer vector with strucs.
<code>registry</code>	Registry directory with registry file.

**Value**

A matrix with integer values indicating left and right corpus positions (columns 1 and 2, respectively).

**Examples**

```
y <- get_region_matrix(
  corpus = "REUTERS",
  s_attribute = "id",
  strucs = 0L:5L,
  registry = get_tmp_registry()
)
```

`ids_to_count_matrix`    *Perform Count for Vector of IDs.*

## Description

The return value is a two-column integer matrix. Column one represents the unique ids of the input vector, column two the respective number of occurrences / counts.

## Usage

```
ids_to_count_matrix(ids)
```

## Arguments

ids	a vector of ids (integer values)
-----	----------------------------------

## Examples

```
ids <- c(1L, 5L, 5L, 7L, 7L, 7L)
ids_to_count_matrix(ids)
table(ids) # alternative to get a similar result
```

`matrix_to_subcorpus`    *Create CWB subcorpus from matrix with regions.*

## Description

Create CWB subcorpus from matrix with regions.

## Usage

```
matrix_to_subcorpus(region_matrix, corpus, subcorpus)
```

## Arguments

region_matrix	A two-colum matrix with regions in rows: Start position of region in first column, end position in second column.
corpus	A externalptr referencing a corpus such as generated by <code>cl_find_corpus()</code> .
subcorpus	A length-one character vector providing the name for the subcorpus.

## Examples

```
## Not run:
# First we generate a subcorpus from a query result
oil_context <- cqp_query("REUTERS", subcorpus = "OIL", query = '[]{3}"oil" []{3}')
m <- subcorpus_get_ranges(oil_context)
reuters <- cl_find_corpus("REUTERS", registry = get_tmp_registry())
p <- matrix_to_subcorpus(subcorpus = "OIL2", corpus = reuters, region_matrix = m)
cqp_list_subcorpora("REUTERS")

x <- cqp_query("REUTERS:OIL2", query = '"crude";', subcorpus = "CRUDEOIL")
subcorpus_get_ranges(x)

# clean up
cqp_drop_subcorpus("REUTERS:OIL")
cqp_drop_subcorpus("REUTERS:OIL2")
cqp_drop_subcorpus("REUTERS:CRUDEOIL")

## End(Not run)
```

**p\_attr\_default**      *Get default p-attribute*

## Description

Usually the default p-attribute will be "word". Use this function to avoid a hard-coded solution. Extracts the default attribute defined in the CWB source code.

## Usage

```
p_attr_default()
```

## Value

A length-one character vector.

**region\_matrix\_ops**      *Get IDs and Counts for Region Matrices.*

## Description

Get IDs and Counts for Region Matrices.

**Usage**

```

region_matrix_to_ids(
  corpus,
  p_attribute,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  matrix
)

region_matrix_to_count_matrix(
  corpus,
  p_attribute,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  matrix
)

region_matrix_context(
  corpus,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  matrix,
  p_attribute,
  s_attribute,
  boundary,
  left,
  right
)

ranges_to_cpos(ranges)

```

**Arguments**

<code>corpus</code>	a CWB corpus
<code>p_attribute</code>	a positional attribute
<code>registry</code>	registry directory
<code>matrix</code>	a regions matrix
<code>s_attribute</code>	If not NULL, a structural attribute (length-one character vector), typically indicating a sentence ("s").
<code>boundary</code>	Structural attribute (length-one character vector) that serves as a boundary and that shall not be transgressed.
<code>left</code>	An integer value, number of strucs to move to the left.
<code>right</code>	An integer value, number of strucs to move to the right.
<code>ranges</code>	A two-column integer matrix of ranges (left and right corpus positions in first and second column, respectively).

**Details**

`ranges_to_cpos()` will turn a `matrix` of ranges into an `integer` vector with the individual corpus positions covered by the ranges.

## Examples

```
# Scenario 1: Get full text for a subcorpus defined by regions
m <- get_region_matrix(
  corpus = "REUTERS", s_attribute = "places",
  strucs = 4L:5L, registry = get_tmp_registry()
)
ids <- region_matrix_to_ids(
  corpus = "REUTERS", p_attribute = "word",
  registry = get_tmp_registry(), matrix = m
)
tokenstream <- cl_id2str(
  corpus = "REUTERS", p_attribute = "word",
  registry = get_tmp_registry(), id = ids
)
txt <- paste(tokenstream, collapse = " ")
txt

# Scenario 2: Get data.frame with counts for region matrix
y <- region_matrix_to_count_matrix(
  corpus = "REUTERS", p_attribute = "word",
  registry = get_tmp_registry(), matrix = m
)
df <- as.data.frame(y)
colnames(df) <- c("token_id", "count")
df[["token"]] <- cl_id2str(
  "REUTERS", p_attribute = "word",
  registry = get_tmp_registry(), id = df[["token_id"]]
)
df[order(df[["count"]]), decreasing = TRUE,]
head(df)
```

## region\_matrix\_to\_struct\_matrix

*Get min and max strucs of s-attribute present in region*

## Description

Look up the minimum and maximum struc of a s-attribute within a region, including scenario of nested s-attributes. If there are no regions of the s-attribute within the region, NA values are returned.

## Usage

```
region_matrix_to_struct_matrix(
  corpus,
  s_attribute,
  region_matrix,
  registry = NULL
)

region_to_strucs(corpus, s_attribute, region, registry = NULL)
```

**Arguments**

<code>corpus</code>	ID of a CWB corpus.
<code>s_attribute</code>	Name of structural attribute. The attribute may be nested.
<code>region_matrix</code>	A two-column matrix with regions, left corpus positions in column 1, right corpus positions in column 2.
<code>registry</code>	Path of the registry directory. If NULL (default), value of environment variable 'CORPUS_REGISTRY' will be used.
<code>region</code>	Vector with left and right corpus position of region.

**Value**

Depending whether input is a vector (argument `region`) or a matrix (argument `region_matrix`), a vector or a matrix.

`subcorpus_get_ranges`    *Get ranges of subcorpus*

**Description**

Get ranges of subcorpus

**Usage**

```
subcorpus_get_ranges(subcorpus_pointer)
```

**Arguments**

<code>subcorpus_pointer</code>	A pointer (class <code>externalptr</code> ) referencing a CWB subcorpus.
--------------------------------	--------------------------------------------------------------------------

`s_attribute_decode`    *Decode Structural Attribute.*

**Description**

Get `data.frame` with left and right corpus positions (cpos) for structural attributes and values.

**Usage**

```
s_attribute_decode(
  corpus,
  data_dir,
  s_attribute,
  encoding = NULL,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  method = c("R", "Rcpp")
)
```

## Arguments

corpus	A CWB corpus (ID in upper case).
data_dir	The data directory where the binary files of the corpus are stored.
s_attribute	A structural attribute (length 1 character vector).
encoding	Encoding of the values ("latin-1" or "utf-8")
registry	The CWB registry directory.
method	A length-one character vector, whether to use "R" or "Rcpp" implementation for decoding structural attribute.

## Details

Two approaches are implemented: A pure R solution will decode the files directly in the directory specified by `data_dir`. An implementation using Rcpp will use the registry file for `corpus` to find the data directory.

## Value

A `data.frame` with three columns, if the `s`-attribute has values, or two columns, if not. Column `cpos_left` are the start corpus positions of a structural annotation, `cpos_right` the end corpus positions. Column `value` is the value of the annotation.

## Examples

```
# pure R implementation (Rcpp implementation fails on Windows in vanilla mode)
b <- s_attribute_decode(
  corpus = "REUTERS",
  data_dir = system.file(package = "RcppCWB", "extdata", "cwb", "indexed_corpora", "reuters"),
  registry = get_tmp_registry(),
  s_attribute = "places", method = "R"
)

# Using Rcpp wrappers for CWB C code
b <- s_attribute_decode(
  corpus = "REUTERS",
  data_dir = system.file(package = "RcppCWB", "extdata", "cwb", "indexed_corpora", "reuters"),
  s_attribute = "places",
  method = "Rcpp",
  registry = get_tmp_registry()
)
```

---

*s\_attr\_is\_descendent   Explore XML structure of CWB corpus*

---

## Description

The data format of the Corpus Workbench (CWB) allows nested XML as import data. Auxiliary functions assist detecting whether two structural attributes are nested or at the same level (i.e. defining the same regions).

## Usage

```
s_attr_is_descendent(
  x,
  y,
  corpus,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  sample = NULL
)

s_attr_is_sibling(x, y, corpus, registry = Sys.getenv("CORPUS_REGISTRY"))

s_attr_relationship(x, y, corpus, registry = Sys.getenv("CORPUS_REGISTRY"))
```

## Arguments

x	A structural attribute, stated as length-one character vector.
y	Another structural attribute, stated as length-one character vector.
corpus	A corpus ID (length-one character vector).
registry	The directory with the registry file for the corpus.
sample	An integer vector with a sample number of strucs to evaluate. Evaluating only a sample may be an efficient choice for large corpora. If NULL (default), all strucs are evaluated.

## Details

`s_attr_is_descendent()` will evaluate whether s\_attribute x is a child of s\_attribute y. The return value is TRUE (a single logical value) if all regions defined by x are within the regions defined by y. If not, FALSE is returned. The return values is also FALSE if all regions of x and y are identical. Attributes will be siblings in this case, and not in an ancestor-sibling relationship.

`s_attr_is_sibling()` will test whether the regions defined for structural attribute x and structural attribute y are identical. If yes, TRUE is returned, assuming that both attributes are at the same level (siblings). If not, FALSE is returned.

`s_attr_relationship()` will return 0 if s-attributes x and y are siblings in the sense that they define identical regions. The return value is 0 if x is an ancestor of y and 1 if x is a descencdent of y.

## Examples

```
s_attr_is_descendent("id", "places", corpus = "REUTERS", registry = get_tmp_registry())
s_attr_is_sibling(x = "id", y = "places", corpus = "REUTERS", registry = get_tmp_registry())
s_attr_is_sibling(x = "id", y = "places", corpus = "REUTERS", registry = get_tmp_registry())
```

---

<code>s_attr_regions</code>	<i>Get regions defined by a structural attribute</i>
-----------------------------	------------------------------------------------------

---

## Description

Get all regions defined by a structural attribute. Unlike `get_region_matrix()` that returns a region matrix for a defined subset of strucs, all regions are returned. As it is the fastest option, the function reads the binary \*.rng file for the structural attribute directly. The corpus library (CL) is not used in this case.

## Usage

```
s_attr_regions(
  corpus,
  s_attr,
  registry = Sys.getenv("CORPUS_REGISTRY"),
  data_dir = corpus_data_dir(corpus = corpus, registry = registry)
)
```

## Arguments

<code>corpus</code>	A length-one character vector with a corpus ID.
<code>s_attr</code>	A length-one character vector stating a structural attribute.
<code>registry</code>	A length-one character vector stating the registry directory (defaults to CORPUS_REGISTRY environment variable).
<code>data_dir</code>	The data directory of the corpus.

## Value

A two-column matrix with the regions defined by the structural attribute: Column 1 defines left corpus positions and column 2 right corpus positions of regions.

## Examples

```
s_attr_regions("REUTERS", s_attr = "id", registry = get_tmp_registry())
```

---

<code>use_tmp_registry</code>	<i>Use Temporary Registry</i>
-------------------------------	-------------------------------

---

## Description

Use and get temporary registry directory to describe and access the corpora in a package.

**Usage**

```
use_tmp_registry(pkg = system.file(package = "RcppCWB"))

get_tmp_registry()
```

**Arguments**

pkg	Full path to a package.
-----	-------------------------

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