

Using the `tm.plugin.koRpus` Package for Corpus Analysis

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February 18, 2018

The R package `tm.plugin.koRpus` is an extension to the `koRpus` package, enhancing its usability for actual corpus analysis. It adds new classes and methods to `koRpus`, which are designed to work with complete text corpora in both `koRpus` and `tm` formats. This vignette gives you a quick overview.

1 What is `tm.plugin.koRpus`?

While the `koRpus` package focusses mostly on analysis steps of individual texts, `tm.plugin.koRpus` adds several new object classes and respective methods, which can be used to analyse complete text corpora in a single step. These classes are also a first step to combine object classes of both, the `koRpus` and `tm` packages.

There are three basic classes, which are hierarchically nested:

- class `kRp.topicCorpus` holds a list (named by topics) of objects of
 - class `kRp.sourcesCorpus`, which in its `sources` slot holds a list of objects of
 - * class `kRp.corpus`, which in turn contains objects of both `koRpus` and `tm` classes.

The idea behind this is to be able to categorize corpora on at least two levels. The default assumes that these levels are different *sources* and different *topics*, but apart from this naming (which is coded into the classes) you can actually use this for whatever levels you like.

If you don't need these levels, you can just use the function `simpleCorpus()` to create objects of class `kRp.corpus`. It represents a flat corpus of texts. To distinguish texts which came from different sources, use the function `sourcesCorpus()`, which will generate sub-corpora for each source given. And one level higher up, use the function `topicCorpus()`, to sort `kRp.sourcesCorpus` objects by different topics. Objects of this class will only be valid if there are texts of each topic from each source.

2 Tokenizing corpora

As with `koRpus`, the first step for text analysis is tokenizing and possibly POS tagging. This step is performed by the functions mentioned above, `simpleCorpus()`, `sourcesCorpus()`, or `topicCorpus()`, respectively. The package includes four sample texts taken from Wikipedia¹ in its `tests` directory we can use for an elaborate demonstration:

```
> library(tm.plugin.koRpus)
> # set the root path to the sample files
> sampleRoot <- file.path(path.package("tm.plugin.koRpus"), "tests",
+   "testthat", "samples")
> # now we can define the topics (names of the vector elements)
> # and their main path
> samplePaths <- c(
>   C3S=file.path(sampleRoot, "C3S"),
>   GEMA=file.path(sampleRoot, "GEMA")
> )
> # we also define the sources
> sampleSources <- c(
>   wpa="Wikipedia_alt",
>   wpn="Wikipedia_neu"
> )
> # and finally, we can tokenize all texts
> sampleTexts <- topicCorpus(paths=samplePaths, sources=sampleSources,
+   tagger="tokenize", lang="de")

processing topic "C3S", source "Wikipedia_alt", 1 texts...
processing topic "C3S", source "Wikipedia_neu", 1 texts...
processing topic "GEMA", source "Wikipedia_alt", 1 texts...
processing topic "GEMA", source "Wikipedia_neu", 1 texts...
```

Should you need to get hold of the nested objects inside `kRp.sourcesCorpus` or `kRp.topicCorpus` class objects, or replace them with updated ones, you can do so by using the methods `corpusTagged()`, `corpusSources()`, or `corpusTopics()`:

```
> allC3SSources <- corpusSources(corpusTopics(sampleTexts, "C3S"))
> names(allC3SSources)

[1] "wpa" "wpn"
```

3 Analysing corpora

After the initial tokenizing, we can analyse the corpus by calling the provided methods, for instance lexical diversity:

¹see the file `tests/testthat/samples/License_of_sample_texts.txt` for details

```
> sampleTexts <- lex.div(sampleTexts, char=FALSE, quiet=TRUE)
> corpusSummary(sampleTexts)
```

	text	topic	source	CTTR	HD-D	(vocd-D)	Herdan's C	Maas a	Maas	lgV0	
wpaC3S01	wpaC3S01	C3S	wpa	6.13		38.14	0.95	0.16		6.21	
wpnC3S01	wpnC3S01	C3S	wpn	6.82		38.05	0.94	0.17		6.10	
wpaGEMA01	wpaGEMA01	GEMA	wpa	7.07		37.61	0.94	0.17		6.11	
wpnGEMA01	wpnGEMA01	GEMA	wpn	7.13		37.87	0.94	0.16		6.24	
	MATTR	MSTTR	MTLD	MTLD-MA	Root	TTR	Summer	TTR	Uber	index	Yule's K
wpaC3S01	0.81	0.79	100.16	NA	8.68	0.93	0.78		39.92		49.92
wpnC3S01	0.82	0.76	123.01	NA	9.65	0.92	0.73		36.46		54.88
wpaGEMA01	0.80	0.78	106.94	192	10.00	0.92	0.71		35.96		65.08
wpnGEMA01	0.81	0.79	111.64	NA	10.08	0.92	0.73		37.47		60.14

As you can see, `corpusSummary()` fetches a data frame from the object which contains the summarised results of all texts below the given object level. That is, if you are only interested in the results for texts of the first topic, simply apply `corpusSummary()` on the result of `corpusTopics()`:

```
> corpusSummary(corpusTopics(sampleTexts, "C3S"))
```

	text	topic	source	CTTR	HD-D	(vocd-D)	Herdan's C	Maas a	Maas	lgV0	
wpaC3S01	wpaC3S01	C3S	wpa	6.13		38.14	0.95	0.16		6.21	
wpnC3S01	wpnC3S01	C3S	wpn	6.82		38.05	0.94	0.17		6.10	
	MATTR	MSTTR	MTLD	MTLD-MA	Root	TTR	Summer	TTR	Uber	index	Yule's K
wpaC3S01	0.81	0.79	100.16	NA	8.68	0.93	0.78		39.92		49.92
wpnC3S01	0.82	0.76	123.01	NA	9.65	0.92	0.73		36.46		54.88

There are quite a number of `corpus*()` getter/setter methods for slots of these objects, e.g., `corpusReadability()` to get the `readability()` results from objects of class `kRp.corpus`.

Using the summary data.frame, you could now perform ANOVAs or plot interactions:

```
> library(sciplot)
> lineplot.CI(
+   x.factor=corpusSummary(sampleTexts)[["source"]],
+   response=corpusSummary(sampleTexts)[["MTLD"]],
+   group=corpusSummary(sampleTexts)[["topic"]],
+   type="l",
+   main="MTLD",
+   xlab="Media source",
+   ylab="Lexical diversity score",
+   col=c("grey", "black"),
+   lwd=2
+ )
```

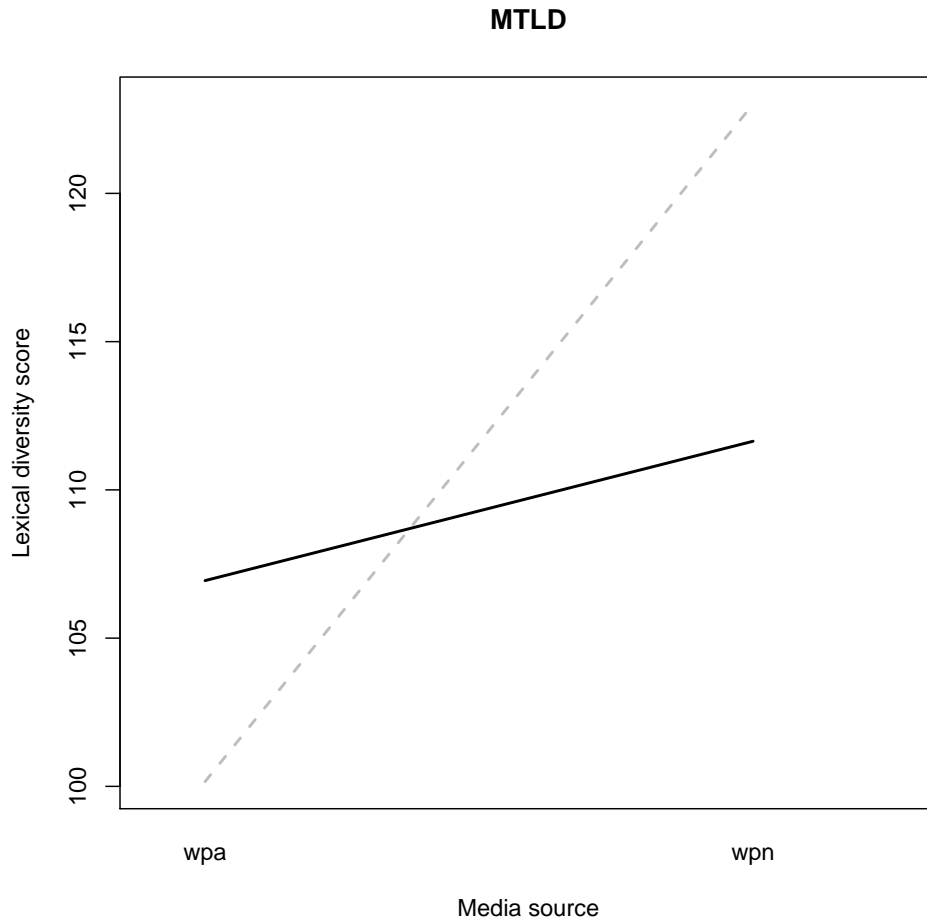


Figure 1: Well, the example texts aren't so impressive here, as there's not much variance in one text per source and topic.

3.1 Frequency analysis

The object classes make it quite comfortable to analyse type frequencies of corpora. There is a method `read.corp.custom()` for these classes, that will do this analysis recursively on all levels:

```
> sampleTexts <- read.corp.custom(sampleTexts, caseSens=FALSE)
> sampleTextsWordFreq <- query(corpusFreq(sampleTexts), var="wclass",
+   query=kRp.POS.tags(lang="de", list.classes=TRUE, tags="words"))
```

```
> head(sampleTextsWordFreq, 10)
```

	num	word	lemma	tag	wclass	lttr	freq	pct	pmio	log10
3	3	die		word.kRp	word	3	30	0.037220844	37220	4.570776
4	4	der		word.kRp	word	3	21	0.026054591	26054	4.415874
5	5	gema		word.kRp	word	4	17	0.021091811	21091	4.324097
6	6	und		word.kRp	word	3	17	0.021091811	21091	4.324097
7	7	einer		word.kRp	word	5	12	0.014888337	14888	4.172836
8	8	von		word.kRp	word	3	12	0.014888337	14888	4.172836
11	11	ist		word.kRp	word	3	10	0.012406948	12406	4.093632
12	12	bei		word.kRp	word	3	9	0.011166253	11166	4.047898
13	13	das		word.kRp	word	3	8	0.009925558	9925	3.996731
14	14	urheber		word.kRp	word	7	8	0.009925558	9925	3.996731
	rank.avg	rank.min	rank.rel.avg	rank.rel.min	inDocs	idf				
3	263.0	263	99.24528	99.24528	4	0				
4	262.0	262	98.86792	98.86792	4	0				
5	260.5	260	98.30189	98.11321	4	0				
6	260.5	260	98.30189	98.11321	4	0				
7	258.5	258	97.54717	97.35849	4	0				
8	258.5	258	97.54717	97.35849	4	0				
11	256.0	255	96.60377	96.22642	4	0				
12	254.0	254	95.84906	95.84906	4	0				
13	252.5	252	95.28302	95.09434	4	0				
14	252.5	252	95.28302	95.09434	2	0.301029995663981				

In combination with the `wordcloud` package, this can directly be used to plot word clouds:

```
> require(wordcloud)
> colors <- brewer.pal(8, "RdGy")
> wordcloud(
+   head(sampleTextsWordFreq[["word"]], 200),
+   head(sampleTextsWordFreq[["freq"]], 200),
+   random.color=TRUE,
+   colors=colors
+ )
```

