Parallel Corpora for the Biomedical Domain

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Abstract

A vast amount of biomedical information is available in the form of scientific literature and government-authored patient information documents. While English is the most widely used language in many of these sources, there is a need to provide access to health information in languages other than English. Parallel corpora can be leveraged to implement cross-lingual information retrieval or machine translation tools. Herein, we review the extent of parallel corpus coverage in the biomedical domain. Specifically, we perform a scoping review of existing resources and we describe the recent development of new datasets for scientific literature (the EDP dataset and an extension of the Scielo corpus) and clinical trials (the ReBEC corpus). These corpora are currently being used in the biomedical task in the Conference on Machine Translation (WMT’16 and WMT’17), which illustrates their potential for improving and evaluating biomedical machine translation systems. Furthermore, we suggest additional applications for multilingual natural language processing using these resources, and plan to extend resource coverage to additional text genres and language pairs.

Keywords: Parallel corpus, biomedical domain, multilingual applications

1. Introduction

Machine translation (MT) is currently being used for a variety of tasks and domains. It is known to play an important role in supporting readers’ access to textual documents in a language other than their native language or for communicating in real time. The accuracy of MT systems has improved in recent years thanks to the availability of large collections of parallel and/or comparable corpora. In turn, these resources could be leveraged by deep learning methods, which created a paradigm shift for MT.

MT plays an important role in the health domain. For instance, it has the potential to enable patients to read documents written in a language in which they are not fluent and to hold a conversation with foreign health professionals in case of accidents or health issues in a foreign country. Further, it allows patients to access health information which is only available in a foreign language, for instance, in the case of disease outbreak with origin in other countries (e.g., Zika virus outbreak in Brazil (Bueno, 2017)).

MT can also support researchers to access scientific literature only available in a foreign language, for instance, when working on tropical diseases specific of a region or even when moving to another country for research purposes (Walker, 2016). Finally, MT can also support the biomedical natural language processing (BioNLP) domain when processing documents in languages other than English for which no specific NLP tools are available. This is often the case for clinical discharge reports that are usually only available in the local language. In such cases, researchers could translate the original document into English and rely on state-of-the-art BioNLP tools that are available for English (?). The biomedical and health domain is well known for its complex nomenclature, for which specific language resources and tools have been developed, e.g., lemmatizers (Liu et al., 2012). Therefore, specific training and test datasets are also necessary to precisely translate biomedical document across languages. However, despite its importance for the general population and researchers, there are very few parallel and comparable corpora specific for this domain.

In this paper, we present an overview of the state-of-the-art on parallel and comparable corpora for the biomedical domain. In a scoping review of existing resources, we characterize the resources available by language pairs and document type and provide pointers to more in-depth descriptions of the resources. Additionally, we present the parallel corpora that we assembled and built, such as EDP (French/English), ReBEC (Neves, 2017) (Portuguese/English) and Scielo (Neves et al., 2016) (French/English, Portuguese/English and Spanish/English). For the latter, we provide details on the corpus construction, insights on the data and their utilization for the biomedical task (Bojar et al., 2016; Jimeno Yepes et al., 2017) of the Conference for Machine Translation (WMT). All corpora are available in our repository in GitHub.

2. Related Work

One of the first efforts that involved the development of large-scale shareable parallel corpora for the biomedical domain was the OPUS collection that contained medical documents from the European Medicines Agency (EMEA) (Tiedemann, 2012). A number of biomedical parallel (Widdows et al., 2002; Ozdowska et al., 2005; Deleger et al., 2009) and comparable corpora (Chiao and Zweigenbaum, 2004) have been used for terminology translation only. Similarly, the Mantra project (Kors et al., 2013)
For all corpora we produced, we carried out the following procedure: (a) document retrieval or download; (b) document parsing and processing; (c) document (sentence) alignment; and (d) quality checking.

**Document retrieval and download.** Document retrieval varies depending on the document collection, some are readily available for download while others need to be crawled from the corresponding Web site.

**Document (sentence) alignment.** The corpora described in this section do not result from organized professional translation. For this reason, the texts were not translated sentence by sentence as is often the case for professional translation of technical documents. Empirical inspection of the corpora suggests that while some of the documents reflect sentence by sentence translation, others were created more freely and the content in one language could be structured differently in the other language. We made the hypothesis that documents could nonetheless be aligned at the sentence level and we relied on automatic tools for performing the alignment. We identified alignment tools based on an evaluation of alignment for literary texts which is a genre that also features fuzzy alignment.

**Quality checking.** After automatically aligning the sentences of the documents, we manually checked a sample of our corpora. This was carried out using the Appraise tool, and we evaluated whether the aligned sentences were translatable and whether the content available in one language could be structured differently in the other language. Native speakers of each foreign language were responsible for this task.

### 4. Application to Three Biomedical Corpora

Here we describe the three corpora that we developed and highlight the differences regarding the particular tools that we used for the various steps above.

**EDP** We identified five open access CC-BY journals, referenced EDP Science as having content in French and in English: the articles were originally written in French but the journals also publish the titles and abstracts in English, using a translation provided by the authors. Three journals are listed by the publisher under *Health*: “Actualités Odonto-Stomatologiques” and “Médecine Buccale Chirurgie Buccale”, which are journals addressing dentistry and “Les Cahiers de Myologie”, a journal addressing musculature. Two journals are listed under *Life & Environmental Sciences*: “Cahiers Agriculture” and “Oilsseeds and Fats, Crops and Lipids”. A list of the journal URLs was ob-

```plaintext
http://www.translatecochrane.fr/corpus/
http://www.edpsciences.org
```
Table 1: Overview of biomedical parallel corpus. We use ISO 639-1 two-letter language codes. A star indicates resources that include previously developed corpora as well as new data.

| Corpus         | genre                      | languages (other than English) | reference                  |
|----------------|----------------------------|--------------------------------|----------------------------|
| Cochrane       | Systematic Review (SR) abstracts | fr                             | (Ive et al., 2016)         |
| COPPA, PaTr    | Patents                    | de,fr                          | (Bojar et al., 2014)       |
| EDP            | Article titles and abstracts | fr                             | iibid.                     |
| EMEA           | Medication description     | cs,da,de,el,es,et,fi,fr,hu,it   | (Tiedemann, 2012)          |
| Himl*          | Patient information and SR abstracts | lt,lv,mt,nl,pl,pt,ro,sk,sl,sv | (Jimeno Yepes et al., 2017) |
| Khresmoi       | Short medical search queries | cs,de,fr                       | (Pecina et al., 2013)      |
| MEDLINE        | Article titles             | cs,de,fr                       | (Wu et al., 2011)          |
| MuchMore Springer |                        | de,es,fr,hu,pl,tu              | (Pecina et al., 2013)      |
| ReBEC          | Clinical Trial summaries   | pt                             | (Widdows et al., 2002)     |
| Santé Canada   | Patient information        | fr                             | (Neves, 2017)              |
| Scielo         | Article titles and abstracts | es,fr,pt                       | (Deleger et al., 2009)     |
| UFA/L*         | Medical web crawl          | cs,da,de,el,es.eu, et,fi,fr,he,hu,hr it,ja,ko,lt,lv,nl,no,pl,pt,sv,tr,zh | (Neves et al., 2016)      |
| UMLS           | Metathesaurus               |                                | (Jimeno Yepes et al., 2017) |

Table 2: Content of open biomedical parallel corpus.

| Corpus       | Tokens | Count method                          |
|--------------|--------|---------------------------------------|
| EDP          | EN 56,684 | wc -w on txt files                   |
|              | FR 62,333 | wc -w on txt files                   |
| ReBEC        | EN 625,881 | reported by (Neves, 2017)            |
|              | PO 665,325 | reported by (Neves, 2017)            |
| Scielo       | EN 20,337,385 | script BioC2txtWithCounts.py       |
|              | ES 21,651,629 | available on GitHub                 |
|              | EN 525,866 | reported by (Neves et al., 2016)     |
|              | FR 735,486 | reported by (Neves et al., 2016)     |
|              | EN 18,573,561 | script BioC2txtWithCounts.py       |
|              | PT 18,573,561 | available on GitHub                 |


tained and crawled on March 15, 2017. The html pages were parsed to extract the titles and abstracts in French and English as well as the author names. Any articles lacking some of this information were discarded. The dataset was pre-processed for sentence segmentation using the Stanford CoreNLP toolkit for use in the WMT17 biomedical task. A manual reference for sentence segmentation was then created independently by revising baseline segmentation after the punctuation marks: full stop, interrogation point, exclamation point and colon. Based on the manually validated sentence segmentation, the dataset was aligned automatically at the sentence level using YASA. Manual evaluation conducted on a sample set suggests that 94% of the sentences are correctly aligned, with about 20% of the sentence pairs exhibiting additional content in one of the languages.

**MEDLINE vernacular titles**  MEDLINE indexes journals in languages other than English that publish a title and abstract in English. In this case, MEDLINE citations include the title of the article in the vernacular language in addition to English. This has been used to develop parallel corpora to train machine translation methods. We have retrieved the MEDLINE citations for articles in French, Spanish and Portuguese available before the first WMT biomedical task. We collected titles in English and vernacular (Spanish, French and Portuguese). Titles are already aligned since they typically can be considered as one sentence. It can be noted that while our work was limited to the languages of interest in the WMT biomedical track, parallel titles and/or abstracts may also be retrieved for other languages. For instance, the query chinese [la] returns 286,151 results on September 29, 2017, and parsing the MEDLINE xml result file could yield several thousand aligned titles and abstract sentences for the relevant citations.

**ReBEC**  As already described in (Neves, 2017), the construction of the ReBEC corpus followed the workflow described in the previous section. The Website site of the Brazilian Clinical Trials Registry provides ways to easily download the trials in XML format, which was further

5Using [http://www.xsitemap.com/](http://www.xsitemap.com/)
6Using the perl utility wget [https://stanfordnlp.github.io/CoreNLP/](https://stanfordnlp.github.io/CoreNLP/)
parsed. However, given the various elements (sub-sections) in a trial, e.g., inclusion criteria, exclusion criteria, and given that some of these appear multiple times in the document, the automatic alignment of parallel documents is not straightforward.

Scielo Scielo (Scientific Electronic Library Online)\(^{16}\) is a database of open access scientific publications with a focus on developing and emerging countries, and especially on Latin America. All publications in Scielo are available under either the Creative Commons Attribution-Noncommercial 3.0 Unported (cc-by-nc) or Attribution 3.0 Unported (cc-by) licenses, which makes all documents suitable for redistribution and research purpose.

We developed a corpus based on Scielo (Scielo corpus \citep{Neves:2016}) using the following procedure. We crawled the Scielo site and retrieved articles periodically from Scielo. Our crawling has its starting point in the pages that list all journals from the "Biological Sciences" and "Health Sciences" subjects. These categories are used to compose the two datasets, with the corresponding names, of our corpus. Despite being distinct categories in Scielo, these are overlapping categories, as there are many journals that belong to both of them. From the list of journals, it is possible to retrieve a list of all issues of a particular journal, which is available in the regional web sites of Scielo in distinct countries, such as Brazil, Chile or Colombia. The HTML page of the journal’s list of issues was further parsed to retrieve the page containing the list of articles of a given issue.

Finally, we downloaded the page of a particular article and parsed the HTML code in order to extract the title and the abstract of each publication. Titles and abstracts were subsequently stored and indexed in the SAP HANA database. All translations of the abstracts in Scielo are the original texts provided by the authors of the publications, who are presumably not professional translators, and who may not have native proficiency in both languages. After the initial version of the corpus produced in 2016, we are using the same procedure to update the corpus on a yearly basis for the ENES and ENPT language pairs. For ENFR, there were no new documents added in 2017.

5. Results

5.1. Datasets Descriptive Statistics

Table \[\text{2}\] presents detailed statistics of the contents of the biomedical parallel corpora that we developed. Table \[\text{3}\] presents an overview of the corpora with the training and test set splits that were offered throughout the WMT campaigns.

5.2. Quality Assessment

We also provide a summary of the correct alignment rate for the various corpora, as shown in Table \[\text{4}\]. The alignment was automatically carried out using the respective tools as previously described and a sample was manually checked using the Appraise tools for ReBEC and Scielo, and manual inspection of text files for EDP.

For EDP the manual reference for sentence segmentation provides an evaluation of Stanford sentence segmentation, which comes to 0.77 F-measure on the French portion and 0.81 F-measure on the English portion. Overall, error analysis reveals that the segmentation errors produced by the Stanford tool mainly result from segmentation of the section titles in structured abstracts (Introduction, Material and Method, Results...) which were considered as separate segments by the manual reference but not by the tools. Other errors occur due to organism names (e.g. \textit{E. coli}, which may cause a sentence boundary to be set after "E.").

5.3. Data format

All corpora presented in the previous section are available from GitHub in the BioC format \citep{Comeau:2013}, a standard XML format in the BioNLP community.

6. Discussion

In this section we present a short discussion on some interesting topics that raised during both the corpus construction and its use in our shared tasks.

6.1. Lessons learned during corpus construction.

The challenges of building parallel corpora for the biomedical include the identification of high quality relevant data that can be shared with the community. Technical issues then lie with the identification of adequate tools for sentence segmentation and alignment.

\textbf{Sentence segmentation:} we relied on tools which are non-specific for the biomedical domains, such as Stanford CoreNLP, OpenNLP and SAP HANA. However, we did observe issues. A specific discussion of sentence segmentation errors is reported in \citep{Neves:2017} for ReBEC. For the EDP corpus, we used initially used Stanford CoreNLP for sentence segmentation (in the version of the corpus distributed at WMT17). Then, we manually validated sentence segmentation in both languages in order to create a reference corpus that may be used to train and evaluate sentence segmentation tools. Therefore, Updated versions of the corpus reflect the manual sentence segmentation.

\textbf{Sentence Alignment:} GMA was used for Scielo and ReBEC. Due to difficulties to install GMA, \citep{Yasa:2013} was used for EDP; however, Yasa may be limited to the language pair en/fr. We can refer readers to \citep{Xu:2015} for a discussion and evaluation of alignment tools for a specialized domain (literary texts). Nevertheless, both tools provided good automatic alignments (?). Additionally, GMA was used for two languages (es and pt) and two document types (scientific publications and clinical trials).

6.2. Differences across the corpora.

Despite the corpora presented in this work have been converted into a similar layout, we observed some differences across the results we obtained. These differences are mostly related to particularities of the corpora, such as its format. One such example is the lower rate of correct alignments for the ReBEC corpus that was due to problems when parsing the document format rather that the alignment tool itself, though some few errors could have come from GMA.

\footnote{\url{http://www.scielo.org}}
multilingual parallel corpora in the biomedical domain. To our knowledge, this is the first survey of biomedical parallel corpora. In addition we detailed the development of corpora that we recently provided for training and evaluating biomedical machine translation systems. The collections cover a total of four languages (including English) and various types of documents, such as scientific publications and clinical trials, from various sources and databases. Further, these corpora have been evaluated on the scope of two shared tasks and are freely available for the scientific community either for MT or other NLP tasks. Finally, future work will contribute towards the inclusion of additional languages, e.g., German, as well as other documents types, e.g., health-related news and clinical reports.

### 6.3. Gaps to be addressed.

We can see from table [1] that there is no clinical corpus or datasets from the social media. Also, some languages benefit from better coverage than others depending on the corpora: DE, ES, FR, PT. Finally, we do not yet cover Asian languages, which we plan to address in the future through collaboration with members of the BioNLP community. Typically, we intend to create an abstract collection from MEDLINE as indicated in section 4.

### 7. Conclusion

We presented a scoping review of the various parallel corpora that are available for the biomedical domain. This work was supported in part by the Agence Nationale pour la Recherche (French National Research Agency) under grant number ANR-13-JCJC-SIMI2-CABeReNeT. The authors would like to thank Arthur Boyer for his contribution to the EDP corpus.

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