LISN @ WMT 2021

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Abstract

This paper describes LISN’s submission to two shared tasks at WMT’21. For the biomedical translation task, we have developed resource-heavy systems for the English-French language pair, using both out-of-domain and in-domain corpora. The target genre for this task (scientific abstracts) corresponds to texts that often have a standardized structure. Our systems attempt to take this structure into account using a hierarchical system of sentence-level tags. Translation systems were also prepared for the News task for the French-German language pair. The challenge this year was to perform unsupervised adaptation to the target domain (financial news). For this, we explored the potential of using a retrieval-based strategy, with sentences that are similar to the test instances used to prime the decoder.

1 Introduction

This paper describes LISN’s submission to the translation shared tasks at WMT’21, where we took part in two shared tasks. For the biomedical translation tasks, we have developed resource-heavy systems for the English-French language pair, using a diversity of out-of-domain and in-domain corpora, thus continuing the efforts reported in (Abdul Rauf et al., 2020). Like for previous years shared task, the target genre (scientific abstract) corresponds to texts that often have a standardized structure comprising typical subsections of one to five lines. Standard subsections report the OBJECTIVE, the METHOD, or the RESULTs of the study. Our systems for this year attempt to take this structure into account using sentence-level tags, with the hope to capture some of the document structure and the phraseology of the domain into account. These systems are documented in Section 2.

2 MT for biomedical texts

In this section, we describe our participation to the biomedical task for WMT’21, in which we participated in both English to French and French to English directions. English-French is a reasonably resourced language pair with respect to biomedical parallel corpora, allowing us to train our Neural Machine Translation (NMT) systems (Vaswani et al., 2017) with in-domain corpora as well as large out-of-domain data that exists for this language pair. Like for last year (Abdul Rauf et al., 2020), our first goal is to make the best of all the available data, including supplementary in-domain monolingual data. Our corpora are described in Section 2.1.

For this year’s participation, we also attempt to take the internal structure of biomedical abstracts into account. Many of these abstracts follow what is often referred to as the “IMRAD format”, comprising the following subsections: INTRODUCTION,
METHODS, RESULTS, and DISCUSSION (Sollici and Pereira, 2004). This structure can be explicit in documents through dedicated headings or remain implicit. Our experiments aim to explore how to use this information in NMT and to measure the correlated impact. We notably expect that by informing the system with sub-document information, it will learn the typical style and phraseology of sentences occurring in each part.

For this purpose, we identified in our data all the abstracts that were conforming to this basic structure and worked to make this structure as explicit and standardized as possible. This notably implied to normalize the mains headings, as some variation was observed: for instance, ANALYSIS may be replaced with DISCUSSION, and additional subparts (OBJECTIVES, CONCLUSION) are also be observed. To incorporate the standard IMRaD format we mapped each subheading to the corresponding IMRaD subpart using a system of tags. Details regarding this process are given in Section 2.2.

All systems are based on the Transformer architecture of Vaswani et al. (2017). We were able to achieve appreciable gains both from back-translation and document structure processing. The results are discussed in Section 2.4.

2.1 Corpus and preprocessing

We trained our baseline systems on a collection of in domain biomedical texts as well as out-of-domain parallel corpus. Table 1 details the corpora used in training.

2.1.1 Parallel corpora

We gathered parallel and monolingual corpora available for English-French in the biomedical domain. The former included the biomedical texts provided by the WMT’20 organizers: Edp, Medline abstracts and titles (Jimeno Yepes et al., 2017), Scielo (Neves et al., 2016) and the Ufal Medical corpus consisting of Cesta, Ecdc, Emea (OpenSubtitles), PatTR Medical and Subtitles. In addition, we used the Cochrane bilingual parallel corpus (Ive et al., 2016), the Taus Corona Crisis corpus and the Mlia Covid corpus. We finally experimented with additional in-domain data selected using Information Retrieval (IR) techniques from general domain corpora including News-Commentary, Books and Wikipedia corpus obtained from the Open Parallel Corpus (OPUS) (Lison and Tiedemann, 2016). These were selected using the data selection scheme described in (Abdul-Rauf and Schwenk, 2009). Medline titles were used as queries to find relevant sentences. We used the 2-best sentences returned from the IR pipeline as additional corpus.

Our out-of-domain corpora include the parallel data provided by the WMT14 campaign for French-English: Gigafr-en, Common Crawl, Europarl, News Commentary and the UN corpora. For development purposes, we used Medline test sets of WMT’18 and 19, while Medline 20 was used as internal test data. This data was sentence-aligned with in-house

| Parallel Corpus | Wash (M) | Sents. |
|-----------------|----------|-------|
| English         | French   |       |
| Ufal            | 89.5     | 100.3 | 2.72 M |
| Edp             | 0.04     | 0.04  | 2.44 K |
| Medline titles  | 5.97     | 6.43  | 0.63 M |
| Medline abstracts | 1.23    | 1.44  | 0.06 M |
| Scielo          | 0.17     | 0.21  | 7.84 K |
| Cochrane-Reference | 2.23  | 2.74  | 0.12 M |
| Cochrane-PE     | 0.43     | 0.53  | 20.5 K |
| Cochrane-GooglePE | 0.63  | 0.77  | 30.3 K |
| Taus            | 20.1     | 23.2  | 0.88 M |
| Mlia            | 19.0     | 23.0  | 1.0M |
| IR Retrieved    | 13.2     | 14.7  | 3.6M |

| Development | Wash (K) | Sents. |
|-------------|----------|-------|
| Medline 18  | 5.7K     | 6.9K  | 265   |
| Medline 19  | 9.8K     | 12.4K | 537   |

| Test | Wash (K) | Sents. |
|------|----------|-------|
| Medline 20 | 12.7K | 16.2K | 699

| Monolingual Corpus | Wash (M) | Sent. |
|-------------------|----------|------|
| English Fr        | 8.79     | 7.70  | 0.33 M |
| Med Fr            | 16.3     | 16.2  | 0.06 M |
| IsTex Fr          | 6.92     | 7.84  | 0.42M |
| Med En            | 3.40     | 4.02  | 0.22M |

| Out Domain | Wash (M) | Sent. |
|------------|----------|------|
| Out-of-domain | 1139 | 1292 | 35M |

Table 1: Data sources for the biomedical task
2.1.2 Monolingual sources

The back-translation of monolingual sources has often been effectively used to cater for parallel corpus shortage in the Biomedical domain in (Stojanovski et al., 2019; Peng et al., 2019). We also adopt this approach here.

Supplementary French data from three monolingual sources were collected from public archives: abstracts of medical papers published by Elsevier from the Lissa portal; a collection of research articles collected from various sources henceforth referred to as Med_Fr (Maniez, 2009). These documents were automatically translated into English with an NMT system trained on biomedical corpora, with a BLEU score of 33.6 on Medline20 testset.

The English side of Medline German and Spanish corpora is used as supplementary English data for back translation. Duplicate documents were removed based on the document id. For these, the internal structure of documents is often available and has been tagged as for the parallel data. These texts were then split into sentences and translated into French using a NMT system trained on all Biomedical corpora with a BLEU score of 36.4 on Medline20 testset. All back-translated data is tagged using the proposal of Caswell et al. (2019).

Parallel and monolingual data are further processed using SentencePiece (Kudo and Richardson, 2018) tokenisation and detokenisation scheme to segment texts into subword units using a vocabulary of 32K subwords. These units were learned on all the in-domain corpora.

2.2 Sentence tagging: a three-level scheme

2.2.1 Tagging domains and corpora

As explained above, our training data is diverse, comprising in-domain parallel, out-of-domain parallel, and in-domain monolingual that is automatically back-translated. Some are made of lists of isolated sentences, while others retain the document information. Even within the in-domain data, some texts precisely match the genre of the testset (scientific abstracts) - this is the case for instance of Medline and to a lesser extent, Cochrane; while others are more remote (eg. the Ufal collection, or the Mlia corpus). In order to reflect this diversity, we designed a three-level sentence tagging scheme that is used for the experiments in Section 2.4.2. These tags appear as prefix of each source sentence.

The first level of tags distinguishes between out-of-domain data (<G>), and in-domain data (tagged <M>). The second level of tag aims to distinguish between data sources, hence the use of one dedicated tag for each corpus, except for the monolingual data, which is simply tagged with <BT>.

2.2.2 Tagging sections within documents

The third level of annotation is intended to enhance the translation context with information regarding the position of a sentence within the abstract. The structure of scientific abstracts in the medical domain often obey the IMRAD structure, and the third tag aims to include this structural information as an additional document-level context. Document level information is necessary to model long-range dependencies between words, phrases, or sentences, or document parts. For a translation system, the ability to model the context may notably improve certain translation decisions, e.g. a better or most consistent lexical choice (Kuang et al., 2018) or a better translation of anaphoric pronouns (Voita et al., 2018; Bawden et al., 2019). A recent review of these themes is in (Maruf et al., 2021).

For this purpose, we further pre-processed 6 corpora containing scientific abstracts. These corpora had different subheadings and structures as given below, which were mapped to a restricted set of section tags listed in Table 2:

1. Medline and Scielo: Abstracts and subheadings often without title. We identified a total of 189 subheadings including spelling variations. Examples include: Presenting Concerns of the Patient, Sources of Information, Novel finding, Study Selection etc.
2. Edp: Abstracts and subheadings mostly contain titles. 45 subheadings were found, such as: Case report, Observation, Subjects and Methods, Commentary, Pedagogical objectives etc.
3. Cochrane: only 10 different subheadings were found, including: Abs selection criteria, abs search strategy, abs data collection, summary title etc.

The identification and standardization of subheading information was a tedious process, involving a lot of rule-based processed to take the variability of sub-headings into account. In order to reconstruct fully parallel versions with subheadings, we also had to reinsert explicit headings in

tools and are shared at [https://github.com/fyvo/WMT-Biomed-Test](https://github.com/fyvo/WMT-Biomed-Test), [https://www.lissa.fr/dc/#env=lissa](https://www.lissa.fr/dc/#env=lissa), [https://www.istex.fr](https://www.istex.fr), [https://crtt.univ-lyon2.fr/les-corpus-medicaux-du-crtt-613310.kjsp](https://crtt.univ-lyon2.fr/les-corpus-medicaux-du-crtt-613310.kjsp), [https://pypi.org/project/sentence-splitter/](https://pypi.org/project/sentence-splitter/)
the source or the target files. Also note that this information was not available for all abstracts. After preprocessing files for which the full subheading information was available, we obtained the 6 fully-tagged corpora (see statistics in Table 3). A similar process was used for test sets (see Table 4).

| Corpus          | Lines | En words | Fr words |
|-----------------|-------|----------|----------|
| Medline         | 34836 | 742891   | 920811   |
| Edp             | 1682  | 34167    | 37508    |
| Scielo (wmt16)  | 7088  | 163275   | 199829   |
| Cochrane-Reference | 123598 | 2741426  | 3308485  |
| Cochrane-GooglePE | 30866  | 685490   | 828436   |
| Cochrane-PE     | 20693 | 468691   | 568262   |

| Testset       | en-fr | fr-en |
|---------------|-------|-------|
| medline20     | 735   | 580   |
| medline18     | 321   | 347   |
| medline19     | 493   | 469   |

Finally, we also introduced a third tag in all other documents as follows: sentences within an abstract where tagged as <ABS>, while all remaining sentences from other corpora where simply tagged as “unspecified subheading” (<US>).

### 2.3 Translation framework

Our translation systems mostly used the basic Transformer models, while a few contrastive systems used the large version (Vaswani et al., 2017). They all rely on Facebook’s seq-2-seq library (fairseq) (Ott et al., 2019) with parameters settings borrowed from transformer_wmt_de_en. The ReLU activation function was used in all encoder and decoder layers. We optimize with Adam (Kingma and Ba, 2015), set up with a maximum learning rate of 0.0005 and an inverse square root decay schedule, as well as 4000 warmup updates. We share the decoder input and output embedding matrices. Models are trained with mixed precision and a batch size of 4096 tokens on 4 V100 GPUs for 300k updates. Systems were trained until convergence based on the BLEU score on the development sets. Evaluation was performed using SacreBleu (Post, 2018). Scores are chosen based on the best score on the development set (Medline 18, 19) and the corresponding scores for that checkpoint are reported on Medline 20 test set.

For fine-tuned systems, the process starts with models trained to convergence, based on BLEU score on dev sets. Training then resumes using a selected portion of the training corpus using the same parameters and criterion as for the base systems. In our results corresponding systems are post-fixed with *-ft.

### 2.4 Results

We present our results for the two directions in two tables, Table 5 and 6, differentiating the normal versus the tag-based systems. Base systems are given on the left, (⇒) identifies the derived (fine-tuned) systems.

#### 2.4.1 Regular MT systems

Results for the untagged systems are reported in Table 5 and are denoted by X*, with E* and F* representing the English to French and French to English systems respectively.

We first built baseline systems. X0 denotes the systems built using only the in-domain data provided by the organizers. X1 are our baseline systems built using all in-domain parallel data. We see good improvement in both directions amounting to 4.2 and 4.8 BLEU points, which is obtained by adding around 1M sentences of additional Cochrane and Taus corpora to the already available 3.4M sentences from WMT’20. This hints at the relevance of the additional in-domain parallel corpora used.

We used the X1 systems as strong in-domain baselines to study the effect of adding back-translated in domain data. These appear as X2 and X3 in Table 5. Adding around 0.8M French to English and around 0.2M English to French back translated sentences did not help as much as we were expecting. We saw similar results last year and increased the amount of back translations this
Table 5: Results for systems using in-domain and out-of-domain corpora. Superscripts *n denote runs submitted.

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| X0  | WMT biomed data              | 3.4M      | 31.6       |
| X1  | All biomed                   | 4.5M      | 35.8       |

Back translations of monolingual data

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| X2  | X1 + BT                      | 5.3M      | 34.8       |
| X3  | X1 + BT-tag                  | 5.3M      | 36.6       |

Out of domain fine-tuned with in domain

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| X4  | outdomain⇒biomed             | 40.5M     | 32.3       |

Table 6: Results for systems with sentences tagged with our 3 level tagging scheme. Test sets are decoded 3 times, where the third tag is varied from the more specific (<SUBHEAD>) to the more generic (<US>). Superscripts *n denote the runs submitted.

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| TE1 | Out+In                       | 41.7M     | 36.2       |
| TE2 | TE1⇒fbibomedplusbt           | 47.2M     | 38.7       |
| TE3 | TE2⇒ftCocMed                 | 48.0M     | 38.2       |

Transformer Large

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| TE4 | Out+In                       | 41.7M     | 36.1       |
| TE5 | TE4⇒fbibomedplusbt           | 47.2M     | 38.4       |

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| TF1 | Out+In                       | 40.9M     | 32.1       |

Mixed baseline finetuned with in-domain

| ID  | Train                        | Sentences | Medline 20 |
|-----|------------------------------|-----------|------------|
| TF2 | TF1⇒fbibomedplusbt           | 46.4M     | 35.7       |
| TF3 | TF2⇒ftCocMed                 | 48.8M     | 35.3       |

year. X3 denote systems built using the tagging scheme proposed by Caswell et al. (2019), where back translations are prefixed with the <BT> tag on the source side.

Indicating that a training sentence is back-translated allows the model to separate the helpful and harmful signal. This proved particularly true for English into French where adding tag to back translations improved the BLEU score by 0.8 points; but it was not helpful in the reverse direction where the amount of back translated data was may be too small (0.2M lines). back-translations as compared to the baseline corpora.

Finally, systems were built by initialising the parameters from huge out-of-domain corpora and later fine tuned on in-domain corpora (X4), where in-domain sub words learned from all the Biomedi-cal data are used to segment the out-of-domain data. The initial systems were trained for 4 epochs on general domain WMT14 EN-FR corpora. The FR-EN system (F4) is the best system in this direction, reaching a BLEU score of 35.8.

2.4.2 Tagged Systems

As our 3-level tagging scheme, described in Section 2.2, is adding information about the domain of each sentence, we specifically focused on larger systems by using all the available in- and out-of-
domain corpora.

Results are summarized in Table 6 with TE* representing the Tagged English to French systems and TF* representing the French to English systems. TE1 is the baseline system for EN-FR built with all the available in-domain and out-domain data. TE4 is the corresponding baseline using a Large Transformer12. We then fine-tune these systems with all the in-domain data including the back translations, these are represented by TE2 and TE5 respectively. This gives an appreciable gain of 2.5 and 2.3 BLEU points for Transformer and Transformer large systems. As we saw no major difference in scores for Transformer versus Transformer large, so we continue the rest of experiments with the simple Transformer architecture. Fine-tuning further with just abstracts from Cochrane and Medline did not yield any further improvement.

French to English results display similar trends. The baseline (TF1) using all available (in domain + out-of-domain) data tagged with our 3 level scheme yielded a BLEU score of 32.1. Fine-tuning it further with all in-domain data (TF2) gives an improvement of 3.6 BLEU points which does not improve further when fine-tuning continues with just Cochrane and Medline abstracts (TF3).

To measure whether the model learned document domain and/or sentence origin information, we tested by tagging the test set with three different tags in the third position, using either the exact subheading, or abstract or UnSpecified for sentences for which the sub-section is unknown. Table 6 reports the scores for the three cases. Though the difference in scores for the three cases is minute, in-domain systems{TE2, TE3, TE5} and {TF2, TF3} achieve their best results when the test set is tagged with the subheading or the abstract tag, typical feature of the biomedical corpora. Conversely, for out-of-domain systems {TE1, TE4, TF1}, the best scores are always for the test set tagged with <US>. This strongly hints that the system is using the extra-information provided by the tag. These observations need to be confirmed using other metrics, as BLEU may not properly reflect these differences.

For English to French direction we got better scores with the tagged systems, with the best system (TE2 = 38.7) achieving 2.1 BLEU points more than the best un-tagged system (E3 = 36.6). This was however not the case for French-English where both tagged and un-tagged systems had more or less similar scores.

Systems in Tables 5 and 6 have different baselines, thus to establish a fair comparison we report numbers for comparable systems in Table 7. Systems {E2, F2} are built by adding back-translated data to the baseline. In systems {E3, E3}, the added back-translated data start with <BT> tag. Systems {TE, TF} use our 3-level tagging scheme for all sentences.

2.5 Conclusion

In this section, we have presented our work for the biomedical task. We notably have tried to incorporate document origin and structure information and improve strong baseline systems that were using a wealth of in-domain and out-of-domain data. Overall, our systems for this year are significantly better than last year’s, even though the benefits of adding document structures as tags need to be confirmed by more experiments and analyses.

3 News translation task: De ↔ Fr

In the 2021 News translation task, we focused on the German-French language pair in which the participants are asked to build MT systems for News in the financial domain. In this section, we discuss details of our approach and the rationale behind it.

| EN-FR      | FR-EN      |
|------------|------------|
| E2 base+bt | F2 base+bt |
| 34.8       | 33.5       |
| E3 base+bt-tag | F3 base+bt-tag |
| 36.6       | 32.4       |

Table 7: Comparison of our 3 level tagged systems with the corresponding untagged systems. Systems {E2, F2} are built by adding back-translated data to the baseline. In systems {E3, E3}, the added back-translated data start with <BT> tag. Systems {TE, TF} use our 3-level tagging scheme for all sentences.

12Hidden size of 1024 and a feed forward size of 4096. Rest of the parameters same as for other systems.
3.1 Unsupervised adaptation

As the training and development data do not contain domain information, the supervised domain adaptation paradigm is not suitable here. However, non-parametric adaptation (Bapna and Firat, 2019), example-based guided machine translation (Zhang et al., 2018), unsupervised domain adaptation (Farajian et al., 2017) or priming NMT (Xu et al., 2020; Pham et al., 2020) have showed promising results for this problem. These approaches retrieve translation examples that are similar to the input source sentence, and use them to guide the inference and to reproduce existing translations or to locally adapt the pre-trained NMT system to the input sentence.

Even though all of the approaches mentioned above have merits of their own, we decided to focus on computationally cheaper methods such as (Bulte and Tezcan, 2019; Xu et al., 2020) where the retrieved instances provide an extra conditioning context for the decoder. Pham et al. (2020) further improved these techniques by proposing to simultaneously prime the source and the target side of the retrieved examples (see Section 3.4.1) and has been our main source of inspiration.

3.2 Data and preprocessing

We use all available parallel data for De ↔ Fr, with the exception of the ParaCrawl data, for training. We also use monolingual data to improve translation quality. For both languages, we choose NewsCrawl 2020. We additionally use NewsCrawl 2018 and 2019 French data at inference time to explore the ability of our priming model to make use of extra data. Details are in Section 3.4.2. We use newstest2019 as development set and test our models on newstest2020.

We filter out sentence pairs with invalid language tag using fasttext language id model (Bojanowski et al., 2017). We use Moses tools to normalize punctuation, to remove non-printing characters and to tokenize into words. The final parallel data contains 5.6M sentences. We use a shared source-target vocabulary built with 40K Byte Pair Encoding (BPE) units using the subword-nmt implementation (Sennrich et al., 2016b).

3.3 Baseline systems

We build our Transformer-based (Vaswani et al., 2017) systems using fairseq (Ott et al., 2019). Our baseline system is a large Transformer with a hidden size of 1024 and a feedforward size of 4096. We optimize with Adam (Kingma and Ba, 2015), set up with a maximum learning rate of 0.0007 and an inverse square root decay schedule, as well as 4000 warmup updates. We tie the encoder and decoder input embedding matrices with the decoder output embedding matrix and we apply layer normalization before each block. Models are trained with mixed precision and a batch size of 4096 tokens on 4 V100 GPUs for 300k updates.

3.4 Submitted systems

3.4.1 Boosting NMT by similar translations

Our approach comprises 2 steps: similar translation retrieval and inference where the priming example is processed in forced-decoding mode.

The retrieval of relevant examples for a given source sentence is based on their distance in some high-dimensional numerical representation space. These representations are computed using the encoder of the baseline system (see Section 3.3) so as to keep our systems in the "constrained" track, as the use of large pre-trained models such as BERT (Devlin et al., 2019), XLM (Conneau and Lample, 2019), etc., was only allowed in unconstrained submissions. More precisely, for each sentence, we average the contextualized embeddings output at the last layer of the encoder. From the training dataset, we create a datastore of pairs (K, V) in which the key K is the sentence embedding of some source sentence f and the value is the sentence pair (f, e) whose source sentence is f. For each query, we retrieve k keys (k = 10 in all experiments).

The similarity between two sentences is the cosine similarity and the retrieval of the nearest neighbor(s) is performed using FAISS (Johnson et al., 2017). In order to search through a large datastore, we divide it into shards containing at most 500K data points; we conduct the k nearest neighbor search on each shard, gather all the retrieved keys from all shards into a list and reduce it to the k nearest keys. Given an input sentence and the list of its k nearest neighbours, we append m (m ≤ k) retrieved source sentences to the input sentence and initialize the target side by the concatenation of

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15https://github.com/rsennrich/subword-nmt

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16https://github.com/pytorch/fairseq
239

$m$ corresponding target sentences. We use a special
token to separate sentences.

During training, we train the NMT model with
two types of examples (with and without retrieval):
this means that each training sample will occur
twice, once with and once without priming. The
former examples have the following format:

\[
\text{f}_1 \ast \cdots \ast \text{f}_m | | f
\]

\[
\text{e}_1 \ast \cdots \ast \text{e}_m | | e
\]

while the latter is presented as the original data.

During inference, we use the same format as
for the source-side, while we initialize the decoder
with the prefix \( \text{e}_1 \ast \cdots \ast \text{e}_m | | \). We therefore call
this initialization "force-decoding". The special to-
kens, which serve as joiners between the retrieved
sentences and the source/target sentence, are care-
fully chosen so that they never occur in the real
text to avoid ambiguity. As discussed in Pham et al.
(2020), it is possible to concatenate several similar
sentences i.e. use \( m > 1 \); we however only report
results with \( m = 1 \), since our preliminary experi-
ments did not show superior results with \( m > 1 \).

3.4.2 Monolingual retrieval
Pham et al. (2020) suggested that monolingual texts
in the target language can also be helpful to in-
form the inference. To make use of monolingual
data, we create pseudo translation pairs with back-
translation to generate the missing source language
side. For this step, we leverage the baseline NMT
system in Section 3.3 for one direction to back-
translate the monolingual target text in the inverse
direction. We use Newscrawl 2020 as monolin-
guial resource for both directions. The monolingual
French data contains approximately 10M sentences
while the German data is much larger. We ran-
domly extract 10M sentences from the German
monolingual data as the pseudo corpus. The back-
translated corpora are added to the real parallel
corpora to create a larger datastore for retrieval.

3.5 Evaluation
3.5.1 Priming and Back-translation
We mainly evaluate our method on the De$\rightarrow$Fr di-
rection. Results on both Newstest2019 and 2020
are in Table 8. Our priming model is able to im-
prove for 0.4 BLEU on newstest2019. However,
the same improvement is not observed for new-
stest2020. As indicated in Pham et al. (2020),
monolingual back-translated data could be directly
applied during inference without any additional
training. We thus search similar sentences on both
original and synthetic data for the test sets. As
shown in Table 8 (+ bt inference), searching on
synthetic data directly improves our results by 0.6
BLEU point on newstest2019.

| Model   | newstest2019 | newstest2020 |
|---------|--------------|--------------|
| baseline| 35.7         | 32.8         |
| + bt    | 37.5         | 33.7         |
| + tag   | 37.5         | 34.3         |
| priming | 34.6         | 33.2         |
| + bt inference | 35.2 | 33.2 |
| priming + bt | 37.4 | 33.9 |
| + tag   | 36.9         | 34.1         |
| + min sim 0.85 | 37.5     | 34.3         |

Table 8: BLEU scores of models for De$\rightarrow$Fr direction.
Our best submitted system obtained a BLEU score of
28.1 on newstest2021.

Even though priming model could benefit from
back-translated data at inference time, training with
synthetic data has proven to be effective in many
previous works (Sennrich et al., 2016a; Edunov
et al., 2018; Ng et al., 2019). Therefore, we also ex-
periment by adding back-translated data to the orig-
inal data and retrain a translation model. Results
(+ bt) demonstrate that training with synthetic data
clearly improves the performance on both test sets.
Caswell et al. (2019) reports that using explicit tags
to distinguish original from back-translated data
provides further gains; however in our experiments,
taggind BT data was not very helpful.

Our model using priming with synthetic data
was not able to surpass the baseline model trained
with additional back-translated data. One possible
reason is that similar sentences retrieved with low
similarity scores may be too noisy, and therefore
decrease the overall performance. Filtering out
noisy similar sentences (with a threshold of 0.85) help
to further improve the performance and makes
it our best system (+ min sim 0.85). This setting
was used for our primary submission for both di-
rections.

We directly apply the best settings found for
De$\rightarrow$Fr to the reverse direction (Fr$\rightarrow$De) and report
the corresponding results in Table 9.

17 Thresholding the minimum similarity score is the result
of a trade-off: using a high threshold selects good sentences
for priming, at the risk of leaving many examples without
any priming data, while a low threshold retrieves more ex-
amples, many of which are of poor quality. Our preliminary
experiments showed that that 0.85 was a reasonable value.
### 3.5.2 Priming and domain adaptation

In this section, we try to assess the relationship between domain adaptation (DA) and priming, and question our initial assumption that priming performs some kind of unsupervised adaptation. Our test set for this part contains 1000 lines extracted from the European Central Bank (ECB) corpus, also available from OPUS website.

As an alternative to priming, we first consider a simple unsupervised domain adaptation technique, where we retrieve \( k = 10 \) most similar sentences for each test sample, yielding a corpus of \( 10 \times k \) sentences that we use to fine-tune for two epochs the baseline systems. Again, filtering based on a similarity scores helps to accumulate a smaller number of sentences that are closer to the test domain.

We then try to combine priming and fine-tuning in the following manner: for each test sentence, we use the \( k \) nearest examples \((f_1, e_1), \ldots (f_k, e_k)\) to derive \( k \) domain-adaptation examples with priming as follows: the first primes \( f_2 \) with \( f_1 \), the second \( f_3 \) with \( f_2 \), and so on, until finally \( f_1 \) is primed with \( f_k \) (the target part is built accordingly). This corpus is used for fine-tuning, and decoding proceeds as before (with \( f_1 \) as prime).

These approaches (priming, unsupervised DA, and priming+DA) are compared in Table 10. We first see that using back-translated data is detrimental to the BLEU score of the baseline system, an effect that might be due to the difference between News texts and ECB domain. We also see that unsupervised adaptation with highly similar sentences yields a small gain. Priming alone achieves the same result as the baseline, but can also benefit somewhat from unsupervised DA. Our best results are obtained when we mix the two strategies, only keeping highly similar sentences.

#### Table 9: BLEU scores of models for Fr→De. Our best submitted system obtained a BLEU score of 37.2 on newstest2021.

| Model               | newstest2019 | newstest2020 |
|---------------------|--------------|--------------|
| baseline            | 27.7         | 27.2         |
| + bt                | 32.4         | 32.9         |
| + tag               | 30.9         | 31.0         |
| priming + bt        | 29.8         | 29.3         |
| + tag               | 29.5         | 29.6         |
| + min sim 0.85      | 30.4         | 30.1         |

#### Table 10: BLEU scores for De→Fr on ECB.

| Model               | ECB           |
|---------------------|---------------|
| baseline            | 26.7          |
| baseline + bt + tag | 25.9          |
| + FT min sim 0.7    | 26.3          |
| + FT min sim 0.8    | 26.1          |
| priming + bt + tag  | 25.9          |
| + FT                | 26.6          |
| + FT min sim 0.7    | 26.3          |
| + FT min sim 0.8    | 26.0          |
| priming + bt + tag + min sim 0.7 | 26.3 |
| + FT min sim 0.7    | 26.5          |
| + FT min sim 0.8    | 26.3          |
| + FT min sim 0.8    | 26.3          |

### 3.6 Conclusion

In this section, we have reported our attempt to perform domain adaptation through priming, a technique which uses sentences that are similar to the test instances to provide additional context in training and decoding. In our experiments with the translation of News between French and German, we had little success with this technique, even when using massive amounts of back-translated data to search for relevant primes. This suggests that priming is not so useful for “open” domains such as News (Pham et al., 2020), and should better be used for standardized types of texts that occur in more specialized domains. We also tried to compare unsupervised DA and priming, showing that, in our context, the former was yielding better results than the latter and also proposed a promising way to combine these two complementary techniques.

## 4 Conclusion and outlook

In this paper, we have described the systems prepared for this year’s participation to WMT shared tasks. For the biomedical track, most of our efforts have been invested in the development of high resource systems, trying to take the structure of medical abstracts into account. In the News task, we have explored ways to perform unsupervised domain adaptation using retrieval based techniques and back-translated data.

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