Abstract

Interpreting patient case descriptions has emerged as a challenging problem for biomedical NLP, where the aim is typically to predict diagnoses, to recommend treatments, or to answer questions about cases more generally. Previous work has found that biomedical language models often lack the knowledge that is needed for such tasks. In this paper, we aim to improve their performance through a self-supervised intermediate fine-tuning strategy based on PubMed abstracts. Our solution builds on the observation that many of these abstracts are case reports, and thus essentially patient case descriptions. As a general strategy, we propose to fine-tune biomedical language models on the task of predicting masked medical concepts from such abstracts. We find that the success of this strategy crucially depends on the selection of the medical concepts to be masked. By ensuring that these concepts are sufficiently salient, we can substantially boost the performance of biomedical language models, achieving state-of-the-art results on two benchmarks.

1 Introduction

Natural Language Processing (NLP) in the biomedical domain poses a number of particular challenges. For this reason, several Language Models (LMs) that are specialised towards the biomedical domain have been proposed, including BioBERT (Lee et al., 2020), ClinicalBERT (Alsentzer et al., 2019), SciBERT (Beltagy et al., 2019a), and PubMedBERT (Gu et al., 2021). Recent work has focused on analysing the capabilities of such models (Jin et al., 2019; Alghanmi et al., 2021; Sung et al., 2021) and enhancing them further (He et al., 2020b; Yuan et al., 2021; Zhang et al., 2021; Fei et al., 2021). Broadly speaking, biomedical LMs have proven successful in capturing the meaning of specialised terminology, but they have been far less successful in enabling medical reasoning, e.g. for predicting a likely diagnosis from a given patient case description. Currently, the main strategies for alleviating this latter issue have centered on incorporating structured knowledge, especially in the form of knowledge graphs. For example, Meng et al. (2021) proposed a method to integrate a large biomedical knowledge graph into a language model through the use of adapters (Pfeiffer et al., 2020), while Zhang et al. (2022) used graph neural networks to jointly reason about language model outputs and knowledge graphs.

We focus on the task of interpreting patient case descriptions. To illustrate this task, Table 1 shows a question from the MedQA benchmark (Jin et al., 2021). In this context, the aim is typically to infer a diagnosis or to recommend a treatment. This is highly challenging, even for biomedical language models, because many pieces of information may need to be combined to find the right answer, and often some degree of clinical judgment is needed. Accordingly, the performance of state-of-the-art biomedical language models remains rather low for benchmarks such as MedQA. We argue that this can, to some extent, be explained by the fact that

| Question: A 38-year-old woman comes to the emergency department because of progressive headache, blurry vision, and nausea for 1 day. Four days ago, she was diagnosed with a right middle ear infection. She appears lethargic. Her temperature is 39.1°C (102.3°F), and blood pressure is 148/95 mm Hg. Ophthalmologic examination shows bilateral swelling of the optic disc. The corneal reflex in the right eye is absent. Sensation to touch is reduced on the upper right side of the face. Serum studies show increased concentrations of fibrin degradation products. Which of the following is the most likely diagnosis?

(A) Cerebral venous thrombosis
(B) Hypertensive emergency
(C) Subarachnoid hemorrhage
(D) Viral meningitis |

Table 1: Example of a question from MedQA, along with the answer candidates.
interpreting patient case descriptions is a paragraph-level task, whereas the standard masked language modelling objective encourages the model to primarily focus on sentence-level context.

Ideally, biomedical language models for interpreting patient case descriptions would be pre-trained on a task that involves predicting diagnoses, or other salient aspects of these patient cases. Unfortunately, beyond the training fragment of benchmarks such as MedQA, such labelled data is not readily available. As an alternative, we propose to generate a pseudo-labelled dataset, based on the heuristic that whenever a case description mentions a disease, it is likely (although by no means guaranteed) that this disease is a valid diagnosis, and similar for other medical concepts such as treatments. To get access to a large set of case descriptions, we rely on abstracts of published case reports. In particular, starting from a collection of PubMed abstracts, we first use a simple heuristic to identify those that are likely to correspond to case reports. Given a case report that mentions some disease, we then fine-tune the language model on the task of predicting that disease. Note that the target disease is masked, as the task would otherwise be trivial. The pre-training task is formulated as a binary classification problem, i.e. given a patient description and a disease, is that disease the correct diagnosis (or more precisely, is it the disease that was masked). This formulation has the advantage that the input format is similar to that of multiple-choice question answering (QA) and natural language inference (NLI). Beyond diseases, we also experiment with predicting masked treatments. Similar to the usual masked language modelling objective, our pre-training task involves making predictions about masked text spans. However, due to the fact that we specifically mask diseases and treatments, we hypothesize that this will improve the model’s ability to take the whole case description into account when making predictions. Finally, note that we consider this to be an intermediate fine-tuning step. In other words, we start from a state-of-the-art biomedical language model, which is then fine-tuned on the proposed task, before finally being fine-tuned on a downstream task.

We find that this intermediate fine-tuning leads to substantial improvements in downstream tasks, even when using a biomedical LM that was already pre-trained on PubMed. To some extent, this comes from the fact that we specifically fine-tune the model on case reports. However, this in itself is not sufficient. To achieve good results, we find that a careful selection of the target concepts is needed. For instance, strong results are obtained when only masking medical treatments. When masking diseases, the improvements over the baseline are sometimes smaller. This is surprising, given that most questions in the considered benchmarks are about diagnosing diseases. Upon closer inspection, the under-performance of strategies that rely on masking diseases appears to be related to the fact that diseases can be mentioned for two common reasons: (i) because the patient has been diagnosed with that disease, which is the case that underpins the intuition behind our proposed approach, or (ii) because the disease is relevant to the medical history of the patient. In the latter case, only a small part of the abstract may be relevant to the disease, which hampers the extent to which the model learns to focus on the case description as a whole. To address this issue, we propose to split abstracts in which multiple diseases are mentioned. Despite the simplicity of the overall approach, our fine-tuning strategies enable significant improvements over the current state-of-the-art in two benchmarks that are focused on patient case descriptions: MedQA (Jin et al., 2021) and DisKnE (Alghanmi et al., 2021).

2 Related Work

The standard paradigm in NLP at the moment is to fine-tune a pre-trained LM, such as BERT (Devlin et al., 2018), on task-specific training data. However, it has been observed that adding an intermediate step, where the LM is first fine-tuned on a different task, for which training data is more abundant, can be highly beneficial (Phang et al., 2018, 2020; Oğuz et al., 2021; Park and Caragea, 2020; Poth et al., 2021). Several works have investigated the role of intermediate tasks, in particular with the aim of analyzing when and why results improve (Pruksachatkun et al., 2020; Chang and Lu, 2021).

For the biomedical domain, one strategy has been to rely on transfer learning from general-domain tasks. For instance, Soni and Roberts (2020) use general-domain question answering for intermediate training, to improve a clinical question answering system. Another strategy has been to rely on different, but related tasks, such as pre-

\[ \text{Intermediate-FT-Biomedical-LMs} \]
training on natural language inference to develop a question answering system (Jeong et al., 2020). Furthermore, several authors have proposed techniques for infusing the knowledge from biomedical knowledge graphs into LMs (He et al., 2020a; Meng et al., 2021; Jha and Zhang, 2022). More closely related to our approach, He et al. (2020c) propose a strategy which relies on the structure of Wikipedia to infuse knowledge about diseases. For instance, to teach the model about how diseases are treated, they rely on the fact that disease-centric Wikipedia articles tend to have a section called Treatment. They then combine the content of that section with a generated question-style sentence mentioning the aspect considered (i.e. treatment in this case) and a masked disease. However, rather than infusing encyclopedic knowledge, our aim is to teach LMs to interpret patient case descriptions. Another related approach was introduced by Pergola et al. (2021), who propose to fine-tune a biomedical language model by using a masked language modelling objective which is modified such that only biomedical concepts are masked. This approach has some similarities with our work, e.g. the idea of masking biomedical concepts as an intermediate fine-tuning task, but there are also some clear differences. First, we formulate our task as a binary classification problem, rather than masked language modelling. Moreover, we specifically target diseases and treatments, and we only mask one concept at a time (although all occurrences of that concept are masked). Finally, since we focus on paragraph-level understanding, we pay particular attention to how these input paragraphs can be selected. As we will see, each of these differences has a clear impact on the empirical results.

3 Proposed Method

We consider the problem of making predictions from patient case descriptions. For instance, given a description that lists symptoms and other information about the patient (e.g. gender, age, and medical history), we would like to infer the corresponding diagnosis or to recommend suitable treatments. We are specifically interested in the potential of using freely available case reports from the medical literature to improve the ability of standard biomedical LMs to make such predictions. In Section 3.1, we first explain our overall strategy. Subsequently, in Section 3.2 we describe the specific variants that we included in our analysis.

3.1 Overall Strategy

Our aim is to design an intermediate fine-tuning task for specialising biomedical LMs towards the task of interpreting patient case descriptions. This fine-tuning task relies on passages from PubMedDS (Vashishth et al., 2021), a corpus which primarily consists of abstracts from PubMed. First, we split the abstracts into passages of up to 250 words, to address the limitations on input length of BERT-based LMs. Next, we aim to identify those passages that contain a case report, describing a specific patient rather than more general findings. To this end, we rely on the simple but effective heuristic that case reports often mention the age of the patient. In particular, we select those passages that contain at least one keyword from the following list: year-old male, year-old female, year-old boy, year-old girl, year-old woman, year-old man. Let us write $D$ for the resulting corpus, i.e. the set of passages that contain at least one of the aforementioned keywords. Subsequently, we determine which medical concepts are mentioned in the passages from $D$. To this end, we use QuickUMLS (Soldaini and Goharian, 2016) with UMLS-2020AA to identify both the spans and the semantic types (e.g. diseases, treatments) of the mentioned concepts. Finally, we create positive training examples of the form $(P, C)$, where $C$ is a medical concept, and $P$ is a passage from $D$ in which all mentions of $C$ have been replaced by a single <mask> token. To generate negative training examples, we simply replace the medical concept $C$ by another concept, as explained below. A given example (passage,concept) is encoded as follows: 

```
<cls> passage <sep> concept
```

mimicking the input format that is typically used for question answering and natural language inference models. The LM is fine-tuned on these examples using a standard cross-entropy loss.

3.2 Training Strategies

We now describe the different variants that we considered. These variants primarily differ in the kinds of medical concepts that are selected as target concepts. Across all variants, we never mask the concept disorder, as constructing training examples from such mentions was found to be highly detrimental, given its prevalence and generic meaning. For all variants, we attempt to balance the number of positive and negative examples. Table 2 provides an overview of the total number of training exam-
AnyType 1,011,482
SpecificType – diseases 160,534 – treatments 2,460
SplitDis 100,225
OneDis 3,310

Table 2: The total number of training examples for each of the intermediate fine-tuning tasks (#).

samples arising from each of the following strategies.

**AnyType** We create a positive example for every medical concept that is found (with the exception of disorder). Note that passages typically mention several concepts, hence this strategy allows us to derive multiple positive examples from the same passage, each time masking a different concept. To construct negative examples, we corrupt positive examples by randomly selecting a concept from those that have been identified in the corpus, regardless of the semantic type.

**SpecificType** In this variant, we only construct training examples from medical concepts of particular types. Specifically, we have experimented with diseases and treatments. Negative examples are constructed by replacing the target concept with another concept of the same semantic type, i.e. diseases are replaced by diseases, and treatments are replaced by treatments.

**SplitDis** Many passages contain more than one disease, which may confuse the model. For instance, diseases which are mentioned as part of the patient history may only be loosely related to the rest of the case report. Since our aim is to train the model to make predictions based on the whole case description, in this variant, passages containing more than one disease are split into sub-passages. In particular, when constructing a positive example for a target disease $d$, we select the sub-passage which begins with the first sentence in which $d$ is mentioned, and includes all the subsequent sentences, until we reach a sentence that mentions another disease (where this final sentence is excluded from the selected sub-passage). If the target disease is mentioned in a sentence that also contains another disease, it is excluded altogether. For illustration, training examples that were obtained with the SplitDis strategy are presented in Table 3.

**OneDis** Instead of splitting passages mentioning more than one disease into sub-passages, as with SplitDis, here we simply discard such passages. This results in a much smaller number of positive examples, but with stronger guarantees that the disease being masked is salient. In both this and the SplitDis method, negative examples are obtained by using randomly selected diseases.

### 4 Experiments

In this section, we empirically analyse the different variants of the intermediate fine-tuning strategy.

**Evaluation Datasets** We mainly focus on two benchmarks that are specifically focused on interpreting and reasoning about patient cases. First, we use MedQA (Jin et al., 2021), which is a multiple-choice question answering benchmark. The questions are taken from medical exams and are specifically asking about what can be inferred from a given patient case description. We use the English version of this dataset (USMLE). Results for this benchmark are reported in terms of accuracy (Acc). Second, we use DisKnE (Alghanmi et al., 2021), which has been derived from MedNLI (Romanov and Shivade, 2018). Therefore, to use DisKnE, a license and access to MedNLI is required. Instances of this benchmark consist of a patient case description and a disease, and the aim is to predict whether that disease can be inferred as diagnosis. This repurposing from the original MedNLI is of particular relevance to our experiments, given that many instances in MedNLI can be solved simply with linguistic knowledge. DisKnE contains a separate training-test split for each disease, and for each split, we consider the task of ranking all test cases, according to our confidence that the given target disease is a valid diagnosis. The results are averaged across all diseases and are reported in terms of Mean Average Precision (MAP). We use the medical-similar variant of the benchmark.

In addition, we also consider the English version of HeadQA (Vilares and Gómez-Rodríguez, 2019), as a more general healthcare-oriented QA dataset. This dataset contains a broad variety of healthcare questions, most of which do not involve patient descriptions. However, the questions are designed to require complex medical reasoning. As such, we use this benchmark to analyse whether our proposed approach may also benefit such settings.
As a baseline, we directly fine-tune the models on the training data from the downstream task. For the other configurations, we first fine-tune the models on the proposed intermediate task.

We use the official training, validation, and test splits for each dataset, with the exception that we excluded questions with images for HeadQA.

Some questions in this dataset require interpreting images. As this is beyond the scope of the paper, we discard all questions involving images for our experiments. This resulted in a total number of 2589 questions for training, 1336 for validation, and 2675 for testing.

**Setup** We use four pre-trained LMs for the baselines and main experiments:

- the cased version of the standard BERT<sub>base</sub> (Devlin et al., 2019);
- the cased version of SciBERT (Beltagy et al., 2019b);
- the cased version of ClinicalBERT (Alsentzer et al., 2019) that was trained on MIMIC-III while being initialized from BioBERT (Lee et al., 2020);
- the PubMedBERT model (Gu et al., 2021) that was trained from scratch on full-length PubMed articles as well as abstracts.

As a baseline, we directly fine-tune the models on the training data from the downstream task. For the other configurations, we first fine-tune the models on the proposed intermediate task.

We use the official training, validation, and test splits for each dataset, with the exception that we excluded questions with images for HeadQA.

**Training Details** We use the same settings and hyper-parameters for all datasets. For fine-tuning the models on the target task, we set the batch size to 8, the number of epochs to 4 and the learning rate to 2e-5. For the intermediate fine-tuning step, we again set the batch size to 8 and the learning rate to 2e-5. Regarding the number of epochs for intermediate fine-tuning, we note that the number of training examples varies greatly across the different variants. For this reason, and to mitigate the potential for catastrophic forgetting, we tuned the number of epochs, choosing from {2, 3, 4}, based on the development split of the downstream task.

**Limitations** Our method relies on an automated extraction tool for identifying the target medical concepts, which will inevitably lead to some noisy training examples. For example, SplitDis and OneDis rely on the assumption that we can detect all mentions of diseases in the text. More generally, regardless of performance, the predictions of a biomedical LM can clearly not be relied upon for diagnosing patients or recommending treatments in a clinical setting. Our purpose in studying these models is rather because a deeper understanding of patient records would make it possible to improve retrieval systems (e.g. suggesting relevant case reports to a clinician handling an unusual patient) or to identify hypotheses for medical research (e.g. by inducing patterns from large sets of case reports).
4.1 Results

Tables 4, 5 and 6 summarize our results. As can be seen, PubMedBERT clearly outperforms the other language models. In general, most variants of the intermediate fine-tuning tasks lead to clear improvements over the baselines. A clear and remarkable conclusion that can be observed for all benchmarks is that the type of intermediate fine-tuning data appears to be much more important than the number of training examples. For instance, the version of SpecificType which only uses treatments achieves the best overall results, outperforming the previous state-of-the-art for MedQA and achieving among the strongest results for both DisKnE and HeadQA. This is surprising, both because of the small number of training examples we can generate for this variant and because of the focus on diseases in DisKnE and many of the MedQA and HeadQA questions.

For MedQA, SpecificType with treatments outperforms the previous state-of-the-art (Zhang et al., 2022) by 1.9 percentage points, despite not relying on any structured knowledge graphs. Note that DisKnE is a recent benchmark, for which the only reported results thus far were obtained from simply fine-tuning biomedical LMs. These existing results were reported prior to the introduction of PubMedBERT, which outperforms these published results. The OneDis variant performs well for DisKnE, despite the low number of corresponding training examples. For MedQA, SplitDis outperforms SpecificType with diseases (with the exception of BERT), which supports the idea that simply masking diseases can lead to training examples that are too noisy. While HeadQA is not particularly focused on patients case descriptions, we still see consistent improvements over the baselines with SpecificType, SplitDis and OneDis, although the improvements are somewhat smaller than those for MedQA and DisKnE.

We can see that our proposed strategy outperforms the baselines for each of the different language models, with the exception of SciBERT with DisKnE. However, there are some differences between the language models in terms of which variant of our method performs best. For MedQA, for instance, we can see that SpecificType with diseases is highly competitive for BERT and ClinicalBERT (compared to the other variants for these language models). As these are the language models that are least adapted to the considered task, we can indeed expect that more pre-training data might be needed for these models. This can explain the relative success of SpecificType with diseases and SplitDis, given that these are associated with a larger number of training examples.

| Baseline | BERT | ClinicalBERT | SciBERT | PubMedBERT |
|----------|------|--------------|---------|-------------|
| 27.8     | 29.1 | 29.2         | 35.5    |             |
| AnyType  |      |              |         |             |
| SpecificType |      |              |         |             |
| – diseases | 28.2 | 31.5         | 30.4    | 38.0        |
| – treatments | 27.8 | 31.0         | 34.5    | 40.4        |
| SplitDis | 27.7 | 31.8         | 33.4    | 38.7        |
| OneDis   | 27.0 | 29.6         | 33.3    | 35.6        |

Table 4: Results for MedQA in terms of Accuracy.

| Baseline | BERT | ClinicalBERT | SciBERT | PubMedBERT |
|----------|------|--------------|---------|-------------|
| 57.0     | 67.5 | 69.2         | 69.7    |             |
| AnyType  |      |              |         |             |
| SpecificType |      |              |         |             |
| – diseases | 60.2 | 70.0         | 67.0    | 72.9        |
| – treatments | 57.5 | 67.5         | 68.3    | 73.6        |
| SplitDis | 58.3 | 74.1         | 68.1    | 72.2        |
| OneDis   | 64.0 | 68.2         | 66.2    | 74.4        |

Table 5: Results for DisKnE in terms of Mean Average Precision (MAP).

4.2 Analysis

Table 7 shows the results of some variants of the SpecificType with diseases and SplitDis strategies, as explained next. We use PubMedBERT for these experiments, as this model achieved the best results in the main experiments. We focus on the MedQA benchmark as this is the most representative benchmark for our problem setting.

**Frequent vs Rare** We analyze whether there is any advantage in focusing specifically on common diseases, or conversely, in focusing on rare diseases. Table 7 shows the results of two variants of Speci-
Table 6: Results for HeadQA in terms of Accuracy.

|                      | BERT | ClinicalBERT | SciBERT | PubMedBERT |
|----------------------|------|--------------|---------|------------|
| Baseline             | 28.8 | 29.3         | 32.8    | 39.5       |
| AnyType              | 29.3 | 30.0         | 31.7    | 39.1       |
| SpecificType         |      |              |         |            |
| – diseases           | 29.8 | 30.1         | 34.5    | **41.8**   |
| – treatments         | **30.3** | **31.1** | **35.7** | 41.0       |
| SplitDis             | 29.8 | 29.6         | 32.6    | 40.7       |
| OneDis               | 29.7 | 29.8         | 34.0    | 40.8       |

Table 7: Analysis results for MedQA (Accuracy). We also report the total number of training examples for each of the intermediate fine-tuning tasks (#). Results were obtained using PubMedBERT.

| SpecificType         | #   | MedQA (Acc) |
|----------------------|-----|-------------|
| Most-Frequent        | 49,816 | 36.8   |
| Least-Frequent       | 8,466   | 38.0   |
| Most-General         | 7,229   | 36.6   |
| Most-Specific        | 8,778   | 36.9   |
| Treatment-Case-Dis   | 6,934   | 38.2   |
| SplitDis             |      |            |
| Most-Similar         | 1,858   | 37.7   |
| Least-Similar        | 1,870   | 36.7   |
| SplitDis+Def         | 105,952 | 37.7   |
| Treatment-Case-Dis   | 2,430   | 38.4   |

for the Most-General variant, we only consider diseases with fewer than 5 ancestors in SNOMED CT. For the Most-Specific variant, we only consider diseases with at least 30 ancestors. We find that both variants of SpecificType perform similarly.

Similar vs Different We explore a setting in which only case reports about diseases similar to “heart disease” are provided during training. Specifically, we use cui2vec (Beam et al., 2020) to identify the 50 most similar diseases that occur at least once in our corpus of case reports. We then consider a variant of SplitDis in which only passages with heart disease, or any of the 50 similar diseases, occurs as the target disease. Our aim in this experiment is to see whether training on one type of diseases is sufficient to obtain good results. Furthermore, we may also assume that because the resulting corpus only involves similar diseases, the model is forced to focus on more subtle details in the paragraphs, and might thus improve as a result. To test this hypothesis, we also consider the variant Least-Similar, where we instead use the diseases that are least similar to heart disease. Rather than fixing the number of diseases at 50, in this case we chose the number to ensure a similar number of training examples as for Most-Similar. The results for both variants are below those of the standard SplitDis variant. However, we can see that Most-Similar clearly outperforms Least-Similar.

Adding Definitions We analyse the usefulness of UMLS definitions. Specifically, we augment the SplitDis training examples with examples of the form (def, dis), where def is the UMLS definition of a disease, and dis is the corresponding disease. Negative examples are again created by replacing the target disease with a randomly chosen other disease. The results in Table 7 show that adding definitions does not improve the results.

Diseases in Treatment Cases The good performance of the SpecificType variant with treatments, despite the small number of training examples we have for that setting, is one of the most surprising findings from the main experiments. Here we analyse whether this might be related to the quality of the case reports that were selected in that setting, i.e. the case reports that mention a treatment. To this end, we consider all such case reports, but instead of using the treatments as the target concepts, we instead focus on diseases. In other words, we use the SpecificType setting for diseases, but applied to
Table 8: Ablation results for MedQA in terms of Accuracy. Results were obtained using PubMedBERT.

| Method                        | Acc  |
|-------------------------------|------|
| MLM-RandomMask                |      |
| – SplitDis                    | 36.4 |
| – SpecificType: treatments    | 35.2 |
| MLM-SpecificMask              |      |
| – SplitDis                    | 37.6 |
| – SpecificType: treatments    | 38.5 |
| Random-Abstracts               |      |
| – SplitDis                    | 38.2 |
| – SpecificType: treatments    | 37.6 |
| No Mask                       |      |
| – SplitDis                    | 36.7 |
| – SpecificType: treatments    | 37.8 |
| Remove-Sent (treatments)      | 38.9 |

the case reports that mention treatments. We also consider a variant in which the SplitDis setting is applied to these case reports. The results in Table 8, shown as Treatment-Case-Dis, reveal that this variant still underperforms the SpecificCase variant with treatments.

4.3 Ablation Experiments

In this section, we analyse the importance of a number of our design choices. We again focus on PubMedBERT and MedQA. We specifically consider the SplitDis and SpecificType with treatments, as these yielded the best results in the main experiments. The results are summarized in Table 8.

Masked Language Modelling  We experimented with two variants of the masked language modelling (MLM) objective for the intermediate fine-tuning task. For the MLM-RandomMask variant, we randomly mask tokens, following the standard approach that is used for LM pre-training. For the MLM-SpecificMask variant, we specifically mask the tokens corresponding to diseases (for the SplitDis setting) or treatments (for the SpecificType setting). The results show that our approach outperforms both MLM strategies, while MLM-SpecificMask outperforms MLM-RandomMask.

Random Abstracts vs Case Reports  We analyse the importance of specifically focusing on case reports. In the Random-Abstracts variant, rather than targeting abstracts which are likely to correspond to case reports, we consider a set of 60,000 randomly sampled abstracts from PubMedDS. We then use our SplitDis and SpecificType settings to construct the examples. The results in Table 8 show that using randomly chosen abstracts leads to worse results, compared to our standard setting.

Masking vs not Masking  We consider a variant of the method in which the original passage is used, i.e., where we do not replace occurrences of the target disease with a <mask> token. The results in Table 8 clearly show that masking is essential to achieve the best results. Nonetheless, even without masking we obtain results that are clearly better than those of the baseline (i.e., PubMedBERT without intermediate fine-tuning).

Masking vs Removing Sentences  Instead of replacing the target concept with a <mask> token, here we remove the entire sentence in which this concept is mentioned. For this variant, called Remove-Sent, we only consider the SpecificType setting (with treatments), as using SplitDis would result in too few examples, given that several SplitDis examples consist of a single sentence. The results show that removing the sentence underperforms masking the concept.

5 Conclusions

We have proposed a strategy for intermediate fine-tuning of biomedical language models, to improve their ability to interpret patient case descriptions. The core of our strategy is to exploit abstracts of case reports found in the literature, as a surrogate of patient case descriptions, and to rely on the heuristic that diseases and treatments that are mentioned in such abstracts are likely to correspond to diagnoses and recommendations, respectively. Despite its conceptual simplicity and without the cost of manual annotation, this approach was found to lead to clear performance gains, setting a new state-of-the-art in MedQA and DisKnE, while also benefiting more diverse datasets such as HeadQA.

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