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Natural Language Explanatory Arguments for Correct and Incorrect Diagnoses of Clinical Cases

Santiago Marro[®], Benjamin Molinet[®], Elena Cabrio[®] and Serena Villata[®] Université Côte d'Azur, Inria, CNRS, I3S, France {santiago.marro, benjamin.molinet, elena.cabrio, serena.villata}@univ-cotedazur.fr

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Abstract:

The automatic generation of explanations to improve the transparency of machine predictions is a major challenge in Artificial Intelligence. Such explanations may also be effectively applied to other decision making processes where it is crucial to improve critical thinking in human beings. An example of that consists in the clinical cases proposed to medical residents together with a set of possible diseases to be diagnosed, where only one correct answer exists. The main goal is not to identify the correct answer, but to be able to explain why one is the correct answer and the others are not. In this paper, we propose a novel approach to generate argument-based natural language explanations for the correct and incorrect answers of standardized medical exams. By combining information extraction methods from heterogeneous medical knowledge bases, we propose an automatic approach where the symptoms relevant to the correct diagnosis are automatically extracted from the case, to build a natural language explanation. To do so, we annotated a new resource of 314 clinical cases, where 1843 different symptoms are identified. Results in retrieving and matching the relevant symptoms for the clinical cases to support the correct diagnosis and contrast incorrect ones outperform standard baselines.

1 INTRODUCTION

Explanatory Artificial Intelligence (XAI) is a main topic in AI research nowadays, given, on the one side, the predominance of black box methods, and on the other side, the application of these methods to sensitive scenarios like medicine. Among the huge set of contributions in this area (Tjoa and Guan, 2019; Saeed and Omlin, 2021), some approaches highlight the need to build explanations which are clearly interpretable and possibly convincing, leading to the investigation of the generation of argument-based explanations (Cyras et al., 2021). These explanations are intended to be not only rational, but "manifestly" rational (Johnson, 2000), such that arguers can see for themselves the rationale behind inferential steps taken. This task becomes even more challenging if we target the generation of natural language argumentbased explanations (Cyras et al., 2021; Vassiliades et al., 2021).

^a https://orcid.org/0000-0001-6220-0559

^b https://orcid.org/0000-0002-8208-2139

clo https://orcid.org/0000-0002-8208-2139

d https://orcid.org/0000-0003-3495-493X

In this paper, we tackle this challenging task, focusing on a specific application scenario, i.e., the generation of explanatory natural language arguments in medicine. More precisely, our goal is to automatically generate natural language argument-based explanations to be used for educational purposes to train medical residents. These students are trained through tests where first there is the description of a clinical case (i.e., symptoms experienced by the patient, results of clinical exams and analysis, and some further information concerning the patient herself like age, gender, or population group), and they need to answer the following question: "Which of the following is the most likely diagnosis?". The test is composed of a number of possible answers to this question, i.e., potential diagnoses, among which, one of them is the correct diagnosis, and the others are incorrect. The solution consists in selecting the correct answer. In addition, medical residents are asked to justify their answer through an explanation. In order to automatize this training phase, we address the task of automatically generating explanations of the kind: "The patient is affected by $[diagnosis_x]$ because the following relevant symptoms have been identified: [correct diagnosis symptoms]. The [diagnosis_y] is incorrect because the patient is not showing the symptoms [incorrect diagnosis symptoms]".

To address this task, a full pipeline needs to be designed in order to (i) detect the symptoms in the clinical case description, (ii) match them with the symptoms, in a medical knowledge base, to identify to which diseases they are associated with, and what is their frequency, and finally, (iii) generate pattern-based natural language explanations employing the elements identified in the two previous steps. To do so, we first annotate a new resource of 314 unique clinical cases in English, with the symptoms which are relevant to derive the correct and incorrect diagnoses. These symptoms are extracted from the Human Phenotype Ontology (HPO) knowledge base (Köhler et al., 2021), where each disease is associated with the list of symptoms that can be manifested in this disease.

Relying on contextual embedding search, our contribution is threefold: (i) we detect in the clinical case description, the symptoms from a newly annotated resource; (ii) we automatically match the symptoms with those available on HPO, with the aim to associate them to the correct and incorrect diagnoses, and (iii) natural language explanatory arguments are automatically generated. We address an extensive evaluation of this new full pipeline to generate natural language explanations for clinical cases, obtaining very promising results. The work we present in this paper is motivated by the lack of existing medical textual resources annotated with symptoms associated with diagnoses and the need for effective methods to address natural language explanations in medicine. To the best of our knowledge, this is the first approach to generate such a kind of natural language explanations in the medical domain for educational purposes, i.e., to train medical residents to generate effective natural language explanations about the correct and incorrect diagnosis of a clinical case.

2 RELATED WORK

Since the introduction of BERT (Devlin et al., 2019), transformer-based models have recently had a major impact on most NLP tasks. Multiple models evolved from it with different design choices, like RoBERTa (Liu et al., 2019), ELECTRA (Clark et al., 2020) and ALBERT (Lan et al., 2019). These models are trained on a large amount of data from multiple sources and domains, which means that they are not necessarily prepared for the biomedical domain.

In recent years, a great number of resources

and NLP tools have been developed specifically for the biomedical domain. For entity extraction, the most popular datasets are BC4CHEMD (Krallinger et al., 2015), B5CDR-Chem (Li et al., 2016), NCBI-Disease (Doğan et al., 2014), BC2GM (Smith et al., 2008), JNLPBA (Kim et al., 2004), where the annotations range from drug-disease interactions to the identification of diseases, genes, and molecular entities such as protein, DNA, RNA. Symptom detection, i.e., the task we address in this paper, can be seen as a sub-task of the broader task of medical entity extraction.

Off-the-shelf NLP tool-kits such as Spacy (Honnibal and Montani, 2017), MedSpacy (Eyre et al., 2021) and CLAMP (Soysal et al., 2018) provide multiple modules for text processing. In particular, MedSpacy is built on top of Spacy specifically for clinical natural language processing, while CLAMP offers a method for named entity recognition (NER) as well as a visual interface for annotating and labeling clinical text.

Most of the recent approaches treat NER as a sequence labeling task where specialized transformerbased models hold the best results. ple, (Naseem et al., 2021) showed that pre-training the ALBERT model on a huge biomedical corpus ensured that the model captured better biomedical contextdependent NER. Results outperform non-specialized models obtaining SOTA results in a lot of datasets. Similar results can be seen in (raj Kanakarajan et al., 2021), where the authors pre-train a biomedical language model using biomedical text and vocabulary with the technique proposed by ELECTRA. Other specialized models based on BERT have been proposed by (Beltagy et al., 2019), (Lee et al., 2020) and (Gu et al., 2020) and BioMed-RoBERTa (Gururangan et al., 2020) based on RoBERTa.

(Michalopoulos et al., 2020) propose UmlsBERT, a contextual embedding model that integrates domain knowledge from the Unified Medical Language System (UMLS) (Bodenreider, 2004), taking into consideration structured expert domain knowledge. They show that UmlsBERT can associate different clinical terms with similar meanings in the UMLS knowledge base and create meaningful input embeddings by leveraging information from the semantic type of each word. In our work, we compare the representation of the symptoms found in the clinical case with different contextual embeddings with the goal to find a representation which matches the one provided in the Human Phenotype Ontology (HPO).

Ngai et al. (Ngai and Rudzicz, 2022) also tackle the problem of finding relevant clinical information, where among the entities they also identify symptoms. In contrast to our work, they only focus on 6 specific diagnoses. Furthermore, their goal is to predict the correct diagnosis and explain these predictions using feature attribution methods, whilst ours is to generate high-quality explanations in natural language for educational purposes, i.e., to improve medical residents' skills in explaining their answer to the test.

Besides detecting symptoms from clinical cases, in our work, we also aim to accurately map them to medical ontologies, such as the Human Phenotype Ontology (HPO), to identify the relationship between the symptoms (originally described in layperson terms) and diseases. Recent work by (Manzini et al., 2022) proposes a tool for automatically translating between layperson terminology and HPO, using a vector space and a neural network to create vector representations of medical terms and compare them to layperson versions. However, this approach has a limitation in that it translates layperson terms without considering their context, potentially missing relevant information that may change the semantics of the term. In our work, we propose a method that takes into account the context in which the layperson term is introduced, leading therefore to an accurate mapping to an HPO term.

Natural language explanation generation has received a lot of attention in recent years, grounding on the progress of generative models to train specific models for explanations. (Camburu et al., 2018) generate explanations by justifying a relation (entailment, contradiction or neutral) for a premise-hypothesis pair by training a Bi-LSTM on their e-SNLI dataset, i.e., the Stanford Natural Language Inference (Bowman et al., 2015) dataset augmented with an explanation layer which explains the SLNI relations. (Kumar and Talukdar, 2020) propose to generate short explanations with GPT-2 (Radford et al., 2019), learned together with the input by a classifier to improve the final label prediction, using e-SNLI (Camburu et al., 2018). These solutions are not applicable to our use case given that explaining a medical diagnosis is a more challenging task than restraining the explanations to the three basic relations considered by (Camburu et al., 2018) and (Kumar and Talukdar, 2020). (Narang et al., 2020) propose an approach based on the T5 model (Raffel et al., 2019) to generate an explanation after prediction. Again, this solution is not applicable to the specific medical scenario we target, where explanations require to be structured following precise argumentative structures (Josephson and Josephson, 1994; Campos, 2011; Dragulinescu, 2016) and to ground on medical knowledge, like the one we inject through the HPO.

Other approaches use explanations via tem-

plates (Reiter and Dale, 1997), e.g., (Abujabal et al., 2017) uses templates and inject the reasoning steps and query of their Q&A system. To the best of our knowledge, no related work generates natural language post-hoc explanations for the medical domain.

3 DATASET

To train and evaluate the proposed approach to build natural language explanatory arguments, we rely on the MEDQA dataset (Jin et al., 2021), which contains a set of clinical case descriptions together with a set of possible questions and answers on the correct diagnosis. The questions and their associated answers were collected from the National Medical Board Examination in the USA (USMLE), Mainland China (MCMLE), and Taiwan (TWMLE). In this work, we only focus on the clinical cases and the questions in English (i.e., USMLE). In total, the MEDQA-USMLE dataset consists of 12,723 unique questions on different topics, ranging from questions like "Which of the following symptoms belongs to schizophrenia?" to questions about the most probable diagnosis, treatment or outcomes for a certain clinical case which is described (Jin et al., 2021). To reach our goal, we extract the clinical cases belonging to the latter group, which are intended to test medical residents to make the correct diagnosis. We end up with 314 unique clinical cases associated with the list of possible diagnoses.

Annotation of the MEDQA-USMLE Clinical Cases. To annotate the clinical cases from the MEDQA-USMLE dataset, we rely on the labels from the Unified Medical Language System (UMLS) (Bodenreider, 2004) Semantic Types, making it consistent with standard textual annotations in the medical domain (Campillos-Llanos et al., 2021; Albright et al., 2013; Mohan and Li, 2019). In particular, we annotate the following elements in the clinical case descriptions: Sign or Symptom, Finding, No Symptom Occurrence, Population Group, Age Group, Location and Temporal Concept. In this paper, we use only the symptoms, but we addressed a complete annotation to employ these data for future work. Quantifiers defining a symptom have not been annotated (e.g., we can find "moderate pain", where we only annotate "pain"). The labels Sign or Symptom and No Symptom Occurrence are associated only to the text snippet defining the symptom in a sentence. Findings consist of such information discovered by direct observation or measurement of an organism's attribute or condition. For instance, components in "Her temperature is 39.3°C (102.8°F), pulse is 104/min, respirations are 24/min, and blood pressure is 135/88 mm Hg". Location refers to the location of a symptom in the human body, and Temporal Concept is used to tag time-related information, including duration and time intervals. Population Group and Age Group highlight information on the age and gender of the patient.

To address the annotation process of the MEDQA-USMLE dataset, we first carried out a semi-automatic annotation relying on the UMLS database. We processed each clinical case through the UMLS database and obtained all the entities detected along their Concept Unique Identifiers (CUI) and their semantic type. The semantic type is then used to disambiguate the entities and generate the pre-annotated files. After the definition of the detailed annotation guidelines (summarized above) in collaboration with clinical doctors, three annotators with a background in computational linguistics carried out the annotation of the 314 clinical cases. To ensure the reliability of the annotation task, the inter-annotator agreement (IAA) has been calculated on an unseen shared subset of 10 clinical cases annotated by four annotators, obtaining a Fleiss' kappa (Fleiss, 1971) of 0.70 for all of the annotated labels, 0.61 for Sign or Symptom, 0.94 for Location, 0.71 for Population Group, 0.66 for Finding, 0.96 for Age Group and 0.96 for No Symptoms Occurrence. We can see a substantial agreement for Sign or Symptom, Finding and Population Group, and an almost perfect agreement for Location, Age Group and No Symptoms Occurrence.

Table 1 reports on the statistics of the final dataset, named MEDQA-USMLE-Symp.1 The accuracy of the annotations provided by the three annotators has been validated from a medical perspective with a clinical doctor. Of the seven entity labels, only three contain medical vocabulary (Sign or Symptom, Finding, and No Symptom Occurrence) and they have been evaluated by this expert. More specifically, we randomly sampled 10% of the data (i.e., 30 cases) and we asked the clinician to verify whether the entity was correctly labeled and whether there were any missing or extra words. The results of the validation showed that 98% of the data was labeled correctly. Less than 2% of the instances were evaluated as incorrectly labeled (e.g., a Finding that was labeled as a Sign or Symptom or vice versa).

Knowledge Base of Diseases and Relevant Symptoms. To collect the medical knowledge needed to define whether a detected symptom is relevant with

Table 1: Statistics of the MEDQA-USMLE-Symp dataset.

| Label | # Entities |
|-----------------------|------------|
| Sign or Symptom | 1579 |
| Finding | 1169 |
| Temporal Concept | 567 |
| Location | 498 |
| Population Group | 364 |
| Age Group | 304 |
| No Symptom Occurrence | 264 |

respect to a given disease, we employ the HPO knowledge base to retrieve (i) the relevant information of each diagnosis which is proposed as an option to answer the question "Which of the following is the most likely diagnosis?", and (ii) the symptoms (named terms in HPO) associated to each diagnosis. This knowledge base also includes information on the frequency² of the occurrence of symptoms, defined in collaboration with ORPHA³ as follows: Excluded (0%); Very rare (1-4%); Occasional (5-29%); Frequent (30-79%); Very frequent (99-80%). Obligate (100%); HPO integrates different sources of symptoms, including ORPHA and OMIM⁴. This knowledge base is quite rich and contains also links and hierarchical links between symptoms (Symptom S2 subclass of Symptom S1), genes or definitions.

4 PROPOSED FRAMEWORK

An overview of the framework we propose to address automatic symptom relevancy assessment and matching to build our natural language explanations is visualized in Figure 1. Starting from the clinical cases in which the correct and incorrect diagnosis are already identified, the goal is to assess the relevant symptoms present in the case such that these symptoms can be used to *explain why* a certain diagnosis is the correct one and *why* the incorrect ones have to be discarded.

In order to accurately diagnose a patient's condition, it is important to identify the symptoms that are most relevant to the possible diagnoses. This means looking at all of the symptoms that have been detected and determining which ones are most likely to be related to the underlying cause of the patient's condition. This can be done by considering the individual symptoms and their potential connections to the possible diagnoses. It is also important to consider any additional information that may be available, such as

¹https://github.com/Wimmics/ MEDQA-USMLE-Symp

²https://hpo.jax.org/app/browse/term/HP:0040279

³https://www.orpha.net/consor/cgi-bin/index.php?lng=

FR

⁴https://www.omim.org/

the patient's medical history and other relevant factors, in order to be able to fully explain the diagnosis. Our work focuses on identifying relevant symptoms in order to accurately diagnose a patient's condition.

The relevancy assessment model associates, when possible, the pertinent symptoms mentioned in the clinical case description with a symptom of a diagnosis found in the HPO knowledge base. The proposed framework consists of two different steps, where: (i) we retrieve from HPO the required diagnosis information (i.e., the symptoms and how common they are), then the symptoms in the case are detected and extracted using an attention-based neural architecture which relies on the clinical case text only; (ii) the relevancy of each symptom is assessed by matching the detected symptoms with the ones retrieved from HPO. The matched symptoms are then used to generate natural language argument-based explanations for correct and incorrect diagnoses. In the following, we explain in detail each sub-task in the pipeline:

Symptoms Detection, consisting in detecting the different symptoms described in the clinical case (medical terms or symptoms described by the patient's own words). In order to detect these entities, we propose a neural approach based on pre-trained Transformer Language Models.

Symptoms Alignment, to align a symptom detected in the clinical case with an identical term in HPO. We first compute an embedding vector for each found symptom and then calculate the cosine distance with each term in HPO. We then assign the closest concept to that symptom. We evaluated both static and contextual embedding methods.

Explanation Generation We propose template-based explanations based solely on the symptoms that are relevant to explain the diagnosis. To do this we propose several templates that tackle different kinds of explanations, going from explaining why a patient was given a certain diagnosis to explaining why the alternatives cannot be considered viable options. We support our explanations with statistical information obtained from HPO such as the frequency of each symptom incidence, and we propose to look for possible symptoms that were not detected by the system but are frequent for a certain disease.

5 EXPERIMENTS

In this section, we report on the experimental setup, the obtained results and the error analysis for the symptom detection and symptom alignment methods. **Setup.** For the symptom detection task, we experimented with different transformer-based Language Models (LMs) such as BERT (Devlin et al., 2019), SciBERT (Beltagy et al., 2019), BioBERT (Lee et al., 2020), PubMedBERT (Gu et al., 2020) and Umls-BERT (Michalopoulos et al., 2020) initialized with their respective pre-trained weights. All the models we employ are specialized in the biomedical domain, with the exception of BERT which will serve us as a baseline. We cast the symptom detection problem as a sequence tagging task. Following the BIOtagging scheme, each token is labeled as either being at the Beginning, Inside or Outside of a component. This translates into a sequence tagging problem with three labels, i.e., B-Sign-or-Symptom, I-Sign-or-Symptom and Outside. To fine-tune the LMs, we use the PyTorch implementation of huggingface (Wolf et al., 2020) (v4.18). For BERT, we use the uncased base model with 12 transformer blocks, a hidden size of 768, 12 attention heads, and a learning rate of 2.5e-5 with Adam optimizer for 3 epochs. The same configuration was used to fine-tune SciBERT BioBERT, PubMedBERT and UmlsBERT. For SciB-ERT, we use both the cased and uncased versions, and for BioBERT we use version 1.2. Batch size was 8 with a maximum sequence length of 128 subword tokens per input example.

Regarding the matching module, we experimented with two different methods to align our detected symptoms with terms in HPO by (i) directly comparing the computed embeddings of the detected symptoms with the embeddings of the terms in HPO, and (ii) by taking into account the context in which the symptoms are detected and applying the same context to every term in HPO.

To align our detected symptoms (in the clinical case) with the equivalent HPO terms, we calculate the cosine distance of each embedding of the HPO terms with respect to the embedding of the detected symptom. In the experimental setting of (i) and (ii), we use the static pre-trained embeddings GloVe 6B as well as BERT, SciBERT, BioBERT and UmlsBERT in the same configurations as in the symptom detection task. For (ii), it is necessary to calculate the context embeddings "on the fly" because each context is unique and depends on the clinical case where it was detected. It is not reasonable to recalculate all HPO term embeddings on the fly for each new context since the ontology contains 10,319 unique terms, so we propose to generate all the HPO terms embedding at once and save them. Therefore, this module takes as input the symptoms detected by the previous module and finds the context⁵ of these symptoms in the clinical case.

⁵The context consists of the sentence(s) containing the

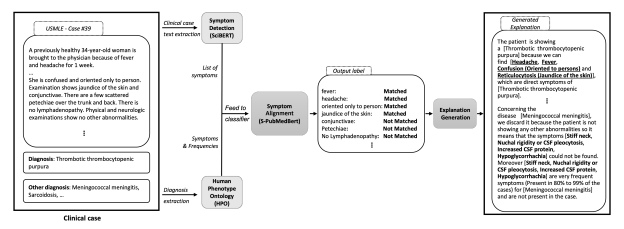


Figure 1: Overview of our full pipeline for symptom prediction and alignment, and NL explanation generation module.

The context C is embedded using sentence embedding methods and saved separately from the symptom S, and the two embeddings are added together (C+S) to form the reference R. This same context embedding C is added in the same way to each HPO term embedding T_1, T_2, \ldots, T_i to form the candidates C_1, C_2, \ldots, C_i .

We compute and retrieve the five best cosine distances between C and R to address a fair comparison with other systems.

We defined a test set of 23 cases where (i) we retrieved from HPO the symptoms related to the diseases for each case, and (ii) we manually aligned the annotated symptoms in the case to the concepts from HPO. This resulted in 162 symptoms aligned to a specific term in HPO that serve us as a testing set for our matching module.

As mentioned in Section 2, the system proposed by (Manzini et al., 2022) offers a similar approach to translating layperson terms to medical terms in HPO. However, their work does not take into account the context in which a symptom is found. To the best of our knowledge, this system constitutes the stateof-the-art when translating layperson terms to HPO terms so we decided to compare our proposal with theirs. However, due to the unavailability of their model, we rely on their online demo, which outputs only the top 5 ranking of the HPO terms that are closest to the input symptom. To perform a comparison with our pipeline, we first compute the accuracy of the aligned symptoms using our symptoms alignment module and then replaced it with (Manzini et al., 2022) proposed system (DASH). Results are shown in Table 4.

Since a symptom can be composed of several words (e.g., "shortness of breath"), we split the

To tackle both tasks we make use of our annotated dataset (Section 3). The annotations are converted into two datasets, one for each part of the pipeline. The first dataset is used for the symptom detection task, and it is in the CoNLL format for token-wise labels. The second dataset, for the symptom alignment task, is converted into a csv format, where each symptom in the clinical case description and available related knowledge (i.e., the list of symptoms and their frequencies for each possible diagnosis associated with the case) extracted from HPO are paired.

Results. Results for the symptom detection task are shown in Table 2 in macro multi-class precision, recall, and F1 score. We can observe that all models

symptom into words that we encode by either using each word as an input on Glove (Pennington et al., 2014), or extracting directly from the contextualized models the representation of the symptom by summarizing the hidden states of the last four layers in the model. We then sum the vectors of each word to get an n-gram representation of the symptom. We also explore sentence embeddings, by making use of Sentence-BERT (Reimers and Gurevych, 2019), a new model that derives semantically meaningful sentence embeddings (i.e., semantically similar sentences are close in vector space) that can be compared using cosine similarity. Sentence-BERT can be used with different pretrained models, in this work we focus on the models BERT (Devlin et al., 2019), SciBERT (Beltagy et al., 2019), UMLSBERT (Michalopoulos et al., 2020) and S-PubMedBert by (Deka et al., 2022). The first represents a competitive baseline in our experiments since it is the SOTA model for comparing sentences crossdomain, while the three latter models are pre-trained on scientific or medical data or both.

symptom and the entire clinical case.

Table 2: Results for entity recognition in macro multi-class precision, recall, and F1-score.

| Model | P | R | F1 |
|-----------------|------|------|------|
| BERT | 0.85 | 0.84 | 0.84 |
| BioBERT v1.2 | 0.84 | 0.85 | 0.84 |
| UmlsBERT | 0.85 | 0.85 | 0.85 |
| PubMedBERTbase | 0.83 | 0.84 | 0.83 |
| SciBERT cased | 0.85 | 0.85 | 0.85 |
| SciBERT uncased | 0.85 | 0.86 | 0.86 |

perform similarly, with the best results from the specialized SciBERT (Beltagy et al., 2019) model. The biggest difference in performance is given by comparing SciBERT uncased with PubMedBERT, with the SciBERT model performing better. Interestingly, BERT performs closely to the specialized models, and, in some cases, it outperforms them. This may be due to the fact that the clinical cases from our dataset are written for medical exams at the med school. They contain some technical specialized words, but overall the symptoms are described in layperson terms.

It is worth noticing that the majority of our labels do not pertain to medical terminology (e.g. Age and Population Group, Location and Temporal Concept). Sign or Symptom and Finding are the only labels that require specialized vocabulary.

Overall, SciBERT uncased is the best-performing model (in bold) with a macro F1-score of 0.86, outperforming the other approaches for each of the categories. In Table 3 we report the performances for each entity with the best-performing model. The *Sign or Symptom* detection task obtains a 0.82 F1 score. In the work of (Ngai and Rudzicz, 2022), the authors also detect symptoms obtaining an F1 score of 0.61. However, these results can not be directly compared since the datasets on which both models were fine-tuned are different: we train on clinical cases, while they use dialogues between doctors and patients. Moreover, given that the dataset they use is not released, we can not evaluate our approach to their data.

The results of the symptoms alignment module experiments are summarised in Table 4. As baseline models, we propose to use the same methods but without the context of the symptom, similarly to (Manzini et al., 2022) *DASH*. In Table 4 we show only the best-performing baseline *PubMedBERT no context* obtaining similar results to *DASH* (0.41 and 0.37, respectively). Adding contextual representation to the embeddings results in a significant improvement (up to 0.53 in accuracy) supporting the hypothesis that context plays an important role when translating layperson terms to formal medical terms.

Table 3: Results for entity recognition using our best performing model (SciBERT uncased) in P, R, and F1-score.

| Entity | P | R | F1 |
|-----------------------|------|------|------|
| Other | 0.93 | 0.91 | 0.92 |
| Age Group | 1.00 | 0.97 | 0.98 |
| Finding | 0.85 | 0.88 | 0.86 |
| Location | 0.74 | 0.80 | 0.77 |
| No Symptom Occurrence | 0.79 | 0.72 | 0.75 |
| Population Group | 0.88 | 0.95 | 0.91 |
| Sign or Symptom | 0.83 | 0.82 | 0.82 |
| Temporal Concept | 0.78 | 0.87 | 0.82 |
| Weighted avg | 0.89 | 0.89 | 0.89 |
| Macro avg | 0.85 | 0.86 | 0.86 |

Table 4: Results for DASH and our symptom alignment method using different embeddings with and without context (accuracy score).

| Model | Accuracy |
|------------------------|----------|
| DASH | 0.37 |
| PubMedBERT no context | 0.41 |
| BERT + context | 0.38 |
| SciBERT + context | 0.39 |
| UMLSBERT + context | 0.44 |
| S-PubMedBERT + context | 0.53 |

Error Analysis. HPO has limitations with respect to the number of symptoms associated with each diagnosis. For some diagnoses, we have multiple symptoms, while for others we can have only one or none. We notice that in those cases where the diagnosis is a mental disease, the model tends to make more mistakes. Inspecting HPO for this kind of diagnoses, we find that either the diagnosis does not appear in the HPO ontology or the symptoms tend to be more general, including a lot of common symptoms like changes in appetite or low energy, that alone may not be relevant but all together indicate a precise diagnosis. Moreover, some relevant symptoms may not be described explicitly but encoded in the clinical cases as Findings. These findings often refer to a relevant symptom that is not explicitly mentioned in the case, like in the example introduced in Section 3 about findings, where we have "respirations are 24/min" that, combined with the fact that the patient is a 34-yearold woman, means that she has dyspnea. Automatically deriving this implicit knowledge remains an open challenging issue. Given that we rely on HPO only, some diseases or diagnoses are not present in the knowledge base, preventing us to generate the associated explanations. Combining HPO with more specialized medical knowledge bases is a future direction for this work, both to complete the information we have, and also to integrate new diagnoses.

6 NATURAL LANGUAGE EXPLANATION GENERATION

In the previous section, we described the first steps of our pipeline for automatically identifying the relevant symptoms which occur in the clinical case description and then matching them with the symptoms associated with the diseases in the medical knowledge base HPO. We move now to the last step of the pipeline, i.e., the generation of natural language explanatory arguments, according to the identified relevant symptoms for the correct and incorrect diagnoses. Given the specificity of the clinical data we are dealing with, we decide to address this task by generating explanations through the definition of explanatory patterns (Josephson and Josephson, 1994; Campos, 2011; Dragulinescu, 2016). We have therefore defined different patterns which take into account the different requirements of our use case scenario, where we aim at (i) explaining the correct answer by the detected symptoms and their frequency, (ii) explaining why the incorrect options cannot hold, and (iii) highlighting the relevant symptoms not explicitly mentioned in the clinical case. Let us consider the following clinical case, where in bold we highlight the symptoms and we underline the relevant symptoms supporting the correct answer.

Clinical Case. A previously healthy 34-year-old woman is brought to the physician because of fever and **headache** for 1 week. She has not been exposed to any She takes no medications. Her temperature is 39.3°C (102.8°F), pulse is 104/min, respirations are 24/min, and blood pressure is 135/88 mm Hg. She is confused and oriented only to person. shows **jaundice** of the skin and conjunctivae. There are a few scattered petechiae over the trunk and back. There is no lymphadenopathy. Physical and neurologic examinations show no other abnormalities. Test of the stool for occult blood is positive. Laboratory studies show: Hematocrit 32% with fragmented and nucleated erythrocytes Leukocyte count 12,500/mm3 Platelet count 20,000/mm3 Prothrombin time 10 sec Partial thromboplastin time 30 sec Fibrin split products negative Serum Urea nitrogen 35 mg/dL Creatinine 3.0 mg/dL Bilirubin Total 3.0 mg/dL Direct 0.5 mg/dL Lactate dehydrogenase 1000 U/L Blood and urine cultures are negative. A CT scan of the head shows no abnormalities. Which of the following is the most likely diagnosis?

The correct diagnosis is Thrombotic thrombocytopenic purpura, whilst the other (incorrect) options are Disseminated intravascular coagulation, Immune thrombocytopenic purpura, Meningococcal meningitis, Sarcoidosis and Systemic lupus erythematosus.

Why Pattern. We focus here on the correct diagnosis explanation pattern, which allows explaining why this is the correct diagnosis. We define the following template to generate our natural language explanations:

Template 1. (Why for correct diagnosis) The patient is showing a [CORRECT DIAGNOSIS] as these following symptoms [PERFECT MATCHED SYMPTOMS, MATCHED SYMPTOMS] are direct symptoms of [CORRECT DIAGNOSIS].

Moreover, [OBLIGATORY SYMPTOMS] are obligatory symptoms (always present, i.e., in 100% of the cases) and [VERY FREQUENT SYMPTOMS] are very frequent symptoms (holding on 80% to 99% of the cases) for [CORRECT DIAGNOSIS] and are present in the case description.⁶

In Template 1, the [CORRECT DIAGNOSIS] represents the correct answer to the question "Which of the following is the most likely diagnosis?" and therefore the correct diagnosis of the described disease. The [SYMPTOMS] in bold represent the symptoms automatically detected through the first module of our pipeline, and they are also underlined when they are considered as relevant by our matching module, i.e., they are listed among the symptoms for the disease in the HPO knowledge base. Both [PERFECT MATCHED SYMPTOMS] and [MATCHED SYMPTOMS] in Template 1 are considered relevant but they differ in the confidence level the system assigns to the matched symptoms. This allows us to integrate a notion of granularity in our explanations and to rely on the symptoms detected in the clinical case that strongly match with a symptom in HPO. If the system does not detect any relevant symptom, no explanation is generated for the correct answer. Furthermore, we employ the information about the symptom frequencies (retrieved through HPO) in the [OBLIGATORY SYMPTOMS] and [VERY FREQUENT SYMPTOMS] to generate stronger evidence to support our natural language argumentative explanations. Sometimes the frequencies are not available in the HPO, in which case we do not display them in our final explanation.

We present now some examples of explanatory arguments automatically generated by our system.

Example 1. The patient is showing a [Thrombotic thrombocytopenic purpura] as these following symptoms [Headache, Fever, Confusion (Oriented to persons) and Reticulocytosis (Jaundice of the skin)] are direct symptoms of [Thrombotic thrombocytopenic purpura].

Moreover [Reticulocytosis (Jaundice of the skin)] are very frequent symptoms (holding on 80% to 99% of the cases) for [Thrombotic thrombocytopenic purpura] and are present in the case description.

⁶Sources from HPO: https://hpo.jax.org/app/browse/term/HP:0040279

When filling the [SYMPTOMS] span in Template 1, we inject only the symptoms matched in the **HPO** the [PERFECT MATCHED SYMPTOMS], and we combine the HPO symptoms with the symptoms detected in the case description for the [MATCHED SYMPTOMS] in this form: [matched symptom in HPO (detected symptom in the clinical case)] (e.g., in Exam-**Confusion (Oriented to persons)** Reticulocytosis (Jaundice of the skin))

Why not Template. Explaining why a diagnosis is the correct one is important, but it is also necessary to be able to say why the other options are not correct as possible diagnoses for the clinical case under investigation (Miller, 2019). We, therefore, propose to provide explanations based on the relevant symptoms for the incorrect options by contrasting them with the clinical case at hand.

Template 2. (Why not for incorrect diagnosis) Concerning the [INCORRECT DIAGNOSIS] diagnosis, it has to be discarded because the patient in the case description is not showing [INCORRECT DIAGNOSIS SYMPTOMS FROM HPO (MINUS DETECTED SYMPTOMS IN CASE)] symptoms.

Despite [SHARED CORRECT SYMPTOMS] symptoms shared with the [CORRECT DIAGNOSIS] correct diagnosis, the [INCORRECT DIAGNOSIS] diagnosis is based on [INCORRECT DIAGNOSIS SYMPTOMS].

Moreover, [OBLIGATORY SYMPTOMS] are obligatory symptoms (always present, i.e., in 100% of the cases) and [VERY FREQUENT SYMPTOMS] are very frequent symptoms (holding on 80% to 99% of the cases) for [IN-CORRECT DIAGNOSIS], and they are not present in the case description.

Template 2 can be applied to each incorrect possible answer of the case, individually. The incorrect answer corresponds to the [INCORRECT DIAG-NOSIS] and [INCORRECT DIAGNOSIS SYMP-**TOMS**] are all relevant symptoms associated with this disease in the HPO knowledge base, without the symptoms in common with the correct answer. Again, in the template, we use the frequencies provided by HPO to provide further evidence to make our explanatory arguments more effective. The template includes therefore with [OBLIGATORY SYMPTOMS] and [VERY FREQUENT SYMPTOMS] the mandatory and very frequent symptoms of the incorrect diagnosis, which are missing in the clinical case description. The following explanations are automatically generated for (one of) the incorrect diagnoses of the clinical case we introduced at the beginning of this section.

Example 2. Concerning the [Meningococcal meningitis] diagnostic, it has to be discarded because the patient in the case description is not showing [Stiff neck, Nuchal rigidity

or CSF pleocytosis, Increased CSF protein, Hypoglycorrhachia] symptoms.

Despite [Petechiae, Fever, Headache] symptoms shared with the [Thrombotic thrombocytopenic purpura] correct diagnosis, the [Meningococcal meningitis] diagnosis is based on [Stiff neck, Nuchal rigidity or CSF pleocytosis, Increased CSF protein and Hypoglycorrhachia].

Moreover, [Stiff neck, Nuchal rigidity, CSF pleocytosis, Increased CSF protein or Hypoglycorrhachia] are very frequent symptoms (holding on 80% to 99% of the cases) for [Meningococcal meningitis] and are not present in the case description.

Example 2 shows the NL explanation of why the possible answer [Meningococcal meningitis] is not the correct diagnosis given the symptoms discussed in the clinical case description. In case the disease is not found in HPO, we do not generate the associated explanation.

Additional Explanatory Arguments. In order to enrich our explanations with additional explanatory arguments to improve critical thinking in the medical residents, we also generate another template. Indeed, in some clinical cases, it is possible that the symptoms are not sufficient to explain the diagnosis or sometimes the symptom has to be combined with vital signs or other characteristics of the patient to be correctly interpreted. Some of these signs represent potentially important symptoms for the diagnosis, as in the previous example, where the sentence respirations are 24/min could be associated with the symptom of *Dyspnea* in HPO. Template 3 aims at drawing the medical residents' attention to (statistically) important symptoms that are missing or not explicitly mentioned in the clinical case description:

Template 3. Furthermore, [CORRECT DIAGNOSIS VERY FREQUENT SYMPTOMS (MINUS MATCHED SYMPTOMS)] are also frequent symptoms for [CORRECT DIAGNOSIS] and could be found in the findings of the clinical case.

Example 3 is generated by our system and brings attention to Dyspnea. This additional explanatory argument complements the explanation we generate for the correct and incorrect diagnoses in the case presented at the beginning of this section.

Example 3. Furthermore, [Dyspnea, Thrombocytopenia, Generalized muscle weakness, Reticulocytosis, and Microangiopathic hemolytic anemia] are also frequent symptoms for [Thrombotic thrombocytopenic purpura] and could be found in the findings of the clinical case.

Limitations. Our work aims to generate templatebased natural language explanations to explain from a symptomatic point of view why a diagnosis is correct and why the remaining ones are incorrect. Templatebased explanations are limited in several ways. First, they are design-dependent, which means that if the templates are not well-designed, they are not helping the user in getting a better understanding of the reason behind a correct/incorrect diagnosis. This can reduce the user's overall satisfaction with the program and make it less effective at achieving its intended goals, i.e. supporting medical residents' training. In our case, we tried to build our template in collaboration with doctors to have a result compliant with their expectations and requirements. Templates are also inflexible and are fixed in advance, they may not be able to adapt to changing circumstances or to new information. This can make them less effective in dynamic or rapidly-changing environments. Again, this is not a serious issue in our case because we are using only the symptoms as the source of data for the moment, which are not evolving.

7 CONCLUSION

In this paper, we present a full pipeline to generate natural language explanatory arguments for correct and incorrect diagnoses in clinical cases. More precisely, based on a novel annotated linguistic resource, our pipeline first automatically identifies in a clinical case description the relevant symptoms and matches them to the HPO medical knowledge base terms to associate symptoms to the correct and incorrect diagnoses proposed as potential answers to the test, and second, automatically generates a natural language explanatory argument which highlights why a certain answer is the correct diagnoses and why the others are not. Extensive experiments on a dataset of 314 clinical cases in English on various diseases show good results (0.86 F1-Score on symptom detection and 0.53 Accuracy on relevant symptom alignment for Top 5 matches), outperforming competitive baselines and SOTA approaches.

Several future work lines arise from this work. First, we plan to address a user evaluation with medical residents. Even though clinical doctors have been involved in the definition of the annotation guidelines we defined, a user evaluation with medical residents is required to get their feedback on our explanatory arguments. Second, we plan to make these explanations interactive to address a rule-based dialogue with the student to focus on precise aspects of the clinical case and go into more precise or generic explanations if required by the student.

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