NLP-CIC-WFU at SocialDisNER: Disease Mention Extraction in Spanish Tweets Using Transfer Learning and Search by Propagation

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Abstract

Named entity recognition (e.g., disease mention extraction) is one of the most relevant tasks for data mining in the medical field. Although it is a well-known challenge, the bulk of the efforts to tackle this task have been made using clinical texts commonly written in English. In this work, we present our contribution to the SocialDisNER competition, which consists of a transfer learning approach to extracting disease mentions in a corpus from Twitter written in Spanish. We fine-tuned a model based on mBERT and applied post-processing using regular expressions to propagate the entities identified by the model and enhance disease mention extraction. Our system achieved a competitive strict F1 of 0.851 on the testing data set.

1 Motivation

Although there are several works for disease mention extraction, the bulk of them has been carried out for clinical texts written in English (Eftimov et al., 2017; Patra and Saha, 2013; Peng et al., 2019; Sachan et al., 2018; Lee et al., 2020; Akhtyamova, 2020; Alsentzer et al., 2019; Yasunaga et al., 2022; Gu et al., 2021). In this work, we present our contribution to the SocialDisNER competition (Gasco et al., 2022) at the SMM4H workshop, task 10 (Weissenbacher et al., 2022). Our system is focused on disease mention extraction from Twitter messages in Spanish. The nature of the texts written in this social network presents new challenges to the disease extraction task because misspellings are frequent (Magumba et al., 2018; Magge et al., 2021). Additionally, many disease mentions can be subsumed in hashtags, urls, or user names (Magumba et al., 2018), which, together with the above, makes it more difficult to identify entities than in common clinical texts as shown in Xiong et al. (2020); García-Pablos et al. (2020); Wang et al. (2019).

2 System description

In this work, we present a transfer learning approach using the model proposed by Tamayo et al. (2022), which is a version of multilingual BERT (Devlin et al., 2019) fine-tuned for disease mention extraction from clinical texts and we apply post-processing rules to extract diseases mentioned in a corpus of tweets in Spanish. Our system tackles the problem in three steps, namely, pre-processing, transfer learning, and post-processing. Below we describe each of them.

2.1 Pre-processing

To implement the fine-tuning process, the BIO scheme (Begin, Inside, Outside) (Ramshaw and Marcus, 1995) was used. Since the dataset provided by SocialDisNER is formatted in a different way, pre-processing was needed to take it to the BIO scheme. We used the disease mentions in the provided structured dataset as a reference to annotate disease mentions in each tweet with their corresponding labels in the BIO scheme. Tokenization was carried out using SpaCy (Honnibal and Montani, 2017) instead of a NER dedicated library such as SciSpacy (Neumann et al., 2019) because the former works for Spanish.

2.2 Transfer learning

We tackled disease mention extraction as a sequence labeling problem using the whole tweet as input, and the labels mentioned above as output. We randomly split partitions of the training dataset into training (75%) and validation (25%) sets. This partition was done iteratively five times with random seeds. Additionally, we carried out a hyperparameter tuning searching for the best model’s configuration using a grid search for the epochs (3, 5, 7) and the learning rate (5e-03, 5e-05, 5e-07). 7 epochs and a learning rate of 5e-05 yielded the best results. With regard to the rest of hyperparameters, default values were kept. For this process,
we used a transformer library and the model available at Hugging Face\(^1\). Google Colab Pro with a GPU Tesla P100 with 27.3 gigabytes of available RAM was used to run all the experiments. The data we used for our training process together with the source code to replicate this work are available at a GitHub repository\(^2\).

2.3 Post-processing plus search by propagation

Post-processing was carried out through a custom Python script to clean up and format the output as follows: 1) Because mBERT works with a subword tokenization system, we decoded the output that contained subwords. 2) We concatenated all the named entities detected by the model one after the other. This means that if the model detected a named entity whose final character position (or final character position plus one) concurred with the first position of the next named entity detected, our system considered that these two entities were part of one single entity. This was necessary because the model extracts parts of some entities separately. 3) We also applied simple but effective post-processing based on some orthographic and grammatical rules which are detailed in Table 1. 4) Under the assumption that SocialDisNER participants were required to extract all the mentions of a disease mention occurring in a tweet, we used the entities extracted by the model to identify and extract any repetitions of said entities in the same document. In order to retrieve misspelled mentions or mentions subsumed by hashtags, urls, or user names, we carried out a search by propagation applying the following steps: a) lowercase both the entity identified by the model and the tweet, b) concatenate multi-word entities, c) delete accents, and d) search entity occurrences throughout the tweet. Lastly, since we work with the BIO scheme, the last post-processing step consisted of decoding the predictions to put them in the data format required by SocialDisNER.

| If the disease mention detected … | … then apply this rule |
|-----------------------------------|------------------------|
| 1. Starts with punctuation mark   | 1. Delete the match and adjust the entity’s beginning index |
| 2. Contains a mark of new line    | 2. Replace the match with a space |
| 3. Contains a space before and/or after a hyphen or a parenthesis | 3. Delete the space(s) and adjust the entity’s ending index |
| 4. Ends with non-content words or punctuation marks | 4. Delete the match and adjust the entity’s ending index |
| 5. Concurs with non-content words or punctuation/hastag marks | 5. Leave out of the entities detected |

Table 1: Post-processing rules

| Model   | P   | R   | F1  |
|---------|-----|-----|-----|
| mBERT   | 0.861 | 0.876 | 0.868 |
| post-processing | | | |

Table 2: Results (5-iteration mean) on the development dataset

our system extracts false positives. They are meaningful entities, but they are not in the gold standard (e.g., EFyC, formación diabetológica). Second, the model truncates some entities (e.g., problemas de aprendizaje instead of problemas de aprendizaje, enfermedades respiratorias crónicas instead of Enfermedades respiratorias crónicas). The first example of the latter type of error is caused by the nature of the mBERT model which works with subwords tokenization. Finally, we consider that there are some errors resulting from an incorrect tagging of the dataset (e.g., our model extracts the entity cáncer but in the gold standard appears niñas y niños que hay hoy con cáncer).

4 Conclusions

In this work, we presented a system based on mBERT following a fine-tuning approach plus simple post-processing and search by propagation to extract disease mentions from Tweets in Spanish. We achieved competitive results with a strict F1 of 0.851, Precision of 0.842, and Recall of 0.860 on the test dataset of the SocialDisNER competition.
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