Comparing Encoder-Only and Encoder-Decoder Transformers for Relation Extraction from Biomedical Texts: An Empirical Study on Ten Benchmark Datasets

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Abstract

Biomedical relation extraction, aiming to automatically discover high-quality and semantic relations between the entities from free text, is becoming a vital step for automated knowledge discovery. Pretrained language models have achieved impressive performance on various natural language processing tasks, including relation extraction. In this paper, we perform extensive empirical comparisons of encoder-only transformers with the encoder-decoder transformer, specifically T5, on ten public biomedicalextraction datasets. We study the relation extraction task from four major biomedical tasks, namely chemical-protein relation extraction, disease-protein relation extraction, drug-drug interaction, and protein-protein interaction. We also explore the use of multi-task fine-tuning to investigate the correlation among major biomedical relation extraction tasks. We report performance (micro F-score) using T5, BioBERT and PubMedBERT, demonstrating that T5 and multi-task learning can improve the performance of the biomedical relation extraction task.

1 Introduction

The scientific literature provides a rich source of biomedical knowledge (e.g., drug-drug interactions), and due to its rapid growth, it becomes increasingly difficult for scientists to keep up-to-date with the most recent discoveries hidden in literature (Zhang and Lu, 2019; Yadav et al., 2020). Moreover, manual curation of information from biomedical literature is time-consuming, costly, and insufficient to keep up with the rapid growth of the literature (Herrero-Zazo et al., 2013). Hence, there has been growing interest in using natural language processing (NLP) techniques for automatic relation extraction (RE) between biomedical entities from texts.

Recently, a variety of approaches based on pretrained language models such as BERT (Devlin et al., 2019) and other variants have shown promising results in various NLP tasks such as relation extraction (drissiya El-allaly et al., 2021b,a), question answering (Sarrouti et al., 2021c,a), text summarization (Goodwin et al., 2020; Yadav et al., 2021), and misinformation detection (Sarrouti et al., 2021b). In particular, RE with classification-based encoder-only pretrained transformers (BERT and variants) has been extensively studied (Lee et al., 2019; Peng et al., 2019a; Gu et al., 2022). In contrast, RE with pretrained language models based on encoder–decoder architecture, specifically Text-to-Text Transfer Transformer (T5) (Raffel et al., 2020), has not been well-studied. Unlike encoder-only transformers, which are designed to predict a single prediction for an input sequence, T5 generates target tokens based on an encoder-decoder architecture.

In this paper, our goal is to compare pretrained sequence-to-sequence transformers with the encoder-only transformers for RE from biomedical texts. In order to satisfy this aim, we compare T5 with in-domain BERT-based models such as BioBERT and PubMedBERT on ten biomedical RE benchmark datasets. We also explore the use of multi-task fine-tuning (MTFT) on ten biomedical RE datasets (each with different entities and relation types) to investigate the correlation among four major biomedical RE tasks, namely chemical-protein relation extraction, disease-protein relation extraction, drug-drug interaction, and protein-protein interaction. Our experiments show that T5 performs better than in domain BERT-based models (encoder-only) such as BioBERT and PubMedBERT. The results also show that fine-tuning T5 with multi-task learning substantially improves the performance compared to single task fine-tuning.

2 Related Work

There has been a recent surge in interest from the NLP community to automatically extract re-
lations between biomedical entities (proteins, gene, diseases, etc.) from the biomedical literature (Krallinger et al., 2008; Segura-Bedmar et al., 2013; Krallinger et al., 2017; Miranda et al., 2021). Recently, with the success of pretrained language models, several techniques based on transformers are widely utilized for extracting the relationships between entities from biomedical literature (Thillaisundaram and Togia, 2019; Wei et al., 2019; Hebbar and Xie, 2021; Hiai et al., 2021; Liu et al., 2021; Zhou et al., 2021; Su et al., 2021; Chang et al., 2021; Weber et al., 2021). The success of these systems has primarily been a result of encoder-only transformers such as BERT (Devlin et al., 2019) and its variants like SciBERT (Beltagy et al., 2019), BioBERT (Lee et al., 2019), and PubMedBERT (Gu et al., 2022). Unlike RE with classification-based encoder-only transformers which have been widely studied, RE with encoder-decoder transformers has not been well-explored. Encoder–decoder-based transformer, specifically T5, (Raffel et al., 2020) has shown strong performance in various NLP tasks such as question answering and text summarization, etc.

In this work, we perform comprehensive comparisons of encoder-only transformers with the encoder-decoder transformer, specifically T5, on ten public biomedical relation extraction datasets. We also explore the use of multi-task learning to learn the shared complementary features across multiple biomedical relation extraction datasets.

3 Experiments

3.1 Problem statement

Given an input sentence \( S \) consisting of \( n \) tokens, i.e., \( S = \{ w_1, w_2, \ldots, w_n \} \) and a pair of entities \( (e_1, e_2) \) where \( e_1 \in S \) and \( e_2 \in S \), RE models are tasked with predicting the maximum probable label \( \hat{y} \) from the set of labels in annotated data, \( y \).

3.2 Datasets and processing

We explore ten benchmark datasets of RE between various entity types such as protein-protein, drug-drug, chemical-protein and disease-protein. Since the vast majority of relation instances are within single sentences in datasets of the aforementioned relation types, we model the RE task as sentence-level relation classification. The statistics of biomedical RE datasets are listed in Table 1.

**Protein-protein interactions.** We use five benchmark datasets, namely BioInfer, AIMed, IEPA, HPRD50, and LLL. These datasets are converted to a unified format by Puysalo et al. (2008). Sentences that contain a pair of proteins are selected to generate positive and negative instances. All protein-protein pairs that occur in a sentence and do not have an explicit label in aforementioned datasets are considered as negative instances. Following previous work, we anonymized target named entities in a sentence using the pre-defined tag @PROTEINS. For instance, a sentence with two protein names is represented as “The POU domains of the @PROTEINS and Oct2 transcription factors mediate specific interaction with @PROTEINS.”.

**Drug-drug interactions.** We use an existing preprocessed version of the Drug-Drug Interaction (DDI) 2013 corpus (Herrero-Zazo et al., 2013) and its corresponding train/dev/test split created by Peng et al. (2019b). Drug names were anonymized using the tag @DRUGS. For instance, a sentence with a pair of drug names is represented as “Ketoconazole: @DRUGS may inhibit both synthetic and catabolic enzymes of @DRUGS”. We evaluate four types of DDI relationships: “mechanism”, “effect”, “advice”, and “Int”. The “mechanism” class defines the DDIs that are described by their pharmacokinetic mechanism. The “effect” type describes an effect or a pharmacodynamic mechanism in DDIs. The “advice” class describes DDIs that mention a recommendation or advice regarding a drug interaction. The “int” class is used when the text describes an interaction between drugs but without providing any additional information.

**Disease-protein relationships.** We use the existing preprocessed versions of the Genetic Association Database corpus (GAD) (Bravo et al., 2015) and EU-ADR datasets (van Mulligen et al., 2012). For both datasets, we use their corresponding train/dev/test splits created by Lee et al. (2019). Targeted entities were anonymized using the tags @DISEASES and @GENES. For instance, a sentence with a pair of two entities (gene and disease in this case) is represented as “In conclusion, @GENE 8092C > A polymorphism may modify the associations between cumulative cigarette smoking and @DISEASES risk.”.

**Chemical-protein relationships.** We use ChemProt (Krallinger et al., 2017) and DrugProt (Miranda et al., 2021) datasets that contain gene–chemical relations. For ChemProt, we use an existing preprocessed version and their corresponding train/dev/test split created by Peng et al.
### Table 1: Statistics of the biomedical relation extraction datasets. For DrugProt, we use the dev set as a test set.

| Dataset     | Train | Dev  | Test  | Metrics   |
|-------------|-------|------|-------|-----------|
| AIMed       | 4938  | -    | 549   | micro F1  |
| BioInfer    | 8544  | -    | 950   | micro F1  |
| HPRD50      | 389   | -    | 44    | micro F1  |
| IEPA        | 734   | -    | 82    | micro F1  |
| LLL         | 300   | -    | 34    | micro F1  |
| DD1         | 2937  | 1004 | 979   | micro F1  |
| ChemProt    | 4154  | 2416 | 3458  | micro F1  |
| DrugProt    | 17277 | 3765 | -     | micro F1  |
| GAD         | 4796  | -    | 3534  | micro F1  |
| EU-ADR      | 318   | -    | 37    | micro F1  |

We evaluate the same five classes: CPR:3, CPR:4, CPR:5, CPR:6, CPR:9. The CPR:3 class describes upregulator, activator, and indirect up-regulator. The CPR:4 class describes downregulator, inhibitor and indirect downregulator relation types. The CPR:5 category describes agonist, agonist activator and agonist inhibitor relation types. The CPR:6 type describes the antagonist relation. The CPR:9 class describes the following relation types: substrate, product of, and substrate product of. For DrugProt, we use the standard training and development sets in the DrugProt shared task and evaluate the same 13 classes: Activator, Agonist, Agonist-Inhibitor, Antagonist, Direct-Regulator, Indirect-Downregulator, Indirect-Upregulator, Inhibitor, Part-Of, Product-Of, Substrate, Substrate_Product-Of, Agonist-Activator.

We first split abstracts into sentences using NLTK and then anonymized target entities in a sentence using the tags @CHEMICAL$ and @GENE$. For instance, a sentence with a pair of two entities (chemical and gene in this case) is represented as “During differentiation, @CHEMICAL$ promoted early expression of osteoblast transcription factors, @GENE$ and osterix.”

### 3.3 Models and setups

We compare in-domain BERT-based language models such as BioBERT (Lee et al., 2019) and PubMedBERT (Gu et al., 2022) with T5 (Raffel et al., 2020) and its variant SciFive (Phan et al., 2021), which is trained on biomedical texts (PubMed abstracts). For BERT-based models, we use a [CLS] token for the classification of relations. The [CLS] representation is fed into a softmax layer for a multi-way classification. For the T5-based models, the input sequence for the relation extraction task is “Processed sentence: [s] Relation: [r]”. We fine-tuned T5 to generate tokens of relation types which are the ground truth labels in training datasets.

We also explore the use of MTFT on ten biomedical RE datasets. Figure 1 illustrates MTFT for RE tasks. We used the proportional and temperature-scaled task mixing as in (Raffel et al., 2020) for data mixture. During fine-tuning, a task-specific token (in our case, name of the dataset) is prepended to the input sequence.

In our experiments, we used the BioBERT (v1.1-base-PubMed), PubMedBERT, T5-base, and SciFive (SciFive-base-Pubmed) implementations provided in HuggingFace’s Transformers package version 4.16.2 (Wolf et al., 2020). All models were trained with a batch size of 16 and maximum sequence length of 300 tokens for 10 epochs using single GPU (16 GB VRAM) on Amazon SageMaker. Adam optimiser with a learning rate of 1e-5 was used.

### 3.4 Results

In Table 2, we show the results of T5-based models compared to the in-domain and SOTA BERT-based models (pretrained on biomedical text) on ten benchmarking biomedical RE datasets, listed in Table 1. We compare the micro F1 scores obtained by T5 and its variant SciFive (pretrained on PubMed abstracts) to the BioBERT and PubMedBERT. On average (micro), T5 which was only pre-trained on the general domain corpus, obtained a higher F1 score than BioBERT and PubMedBERT. T5 achieved the highest F1 scores on 5 out of 10 biomedical RE datasets. Models using biomedical text in pre-training generally perform better than models which pre-trained on general domain corpus. However, we observe that T5-scifive which was pre-trained on biomedical text (PubMed abstracts) did not perform well compared to T5.

We also explored the impact of MTFT on four
| Relation                | Datasets | BioBERT | PubMedBERT | T5    | T5-SciFive | T5-MTFT |
|------------------------|----------|---------|------------|-------|------------|---------|
| Protein-protein        | AIMed    | 92.36   | 93.31      | 94.35 | 94.17      | 93.62   |
|                        | BioInfer | 95.97   | 94.59      | 95.36 | 95.89      | 95.16   |
|                        | HPRD50   | 85.45   | 90.56      | 84.09 | 90.90      | 95.95   |
|                        | IEPA     | 86.58   | 86.46      | 87.80 | 87.80      | 90.24   |
|                        | LLL      | 88.24   |            | 97.05 | 94.11      | 97.05   |
| Drug-drug              | DDI      | 89.67   | 90.69      | 91.01 | 90.60      | 91.83   |
| Chemical-protein       | ChemProt | 90.11   | 91.64      | 90.45 | 92.39      | 96.56   |
|                        | DrugProt | 88.69   | 89.40      | 88.71 | 89.56      | 89.37   |
| Disease-protein        | GAD      | 79.91   | 80.87      | 81.46 | 81.27      | 80.71   |
|                        | EU-ADR   | 57.42   | 62.53      | 78.38 | 75.67      | 83.78   |
| Average score          |          | 85.44   | 88.22      | 89.47 | 89.23      | 91.42   |

Table 2: Biomedical relation extraction test results. In T5-MTFT, we fine-tuned T5 with multi-task learning on ten datasets and then evaluate on the test set for each dataset.

benchmark biomedical RE tasks, i.e., drug-drug interaction, protein-protein interaction, chemical-protein relation extraction, and disease-protein relation extraction. On average, the results clearly show that the performance improves when using MTFT (an improvement of 1.95 F-score over the best single performing model). For instance, on the ChemProt dataset, T5-MTFT was able to achieve significant performance improvement of 6.11 and 6.45 F-score points over T5 and BioBERT respectively. While overall results indicate that MTFT provides improved RE performance on the four biomedical RE tasks (tasks with clear knowledge transfer), we observe a slight drop in the performance on some datasets such as AIMed, BioInfer, and GAD. In MTFT, we believe that in addition to the sample size of each task, the difficulty of the task/dataset can have an impact on the overall performance (the model underfits or overfits a dataset). More efforts and ablation studies are needed to study the impact of different biomedical RE tasks/datasets on downstream performance.

### 3.5 Error analysis

We performed a manual analysis of the test sets where the best performing model (T5-MTFT) predicted an incorrect label. Table 3 presents some examples.

**Protein-protein interaction.** The error analysis has shown that sentences are mostly classified incorrectly when they contain repetitive protein mentions (examples #1 and #3). Multiple protein mentions tend to add noise, which can prevent the model to extract the relevant contextual information. In addition, numerical or statistical findings might be a cause of error (example #1). We also observed that when the protein interacting words (e.g., bind, interact, localization) are mentioned in a sentence, the model predicts the class label “true” (i.e, interacting) (examples #2, #3 and #4).

**Drug-drug interaction.** The model tends to classify “Int” class as “Effect” type (examples #5 and #6). “Int” type is used whenever there exists an interaction between two drugs (i.e., a coarse-grained relation type). Having coarse-grained and fine-grained categories can be a cause of error. We also observed that when the input sentence contains some class-specific words (e.g., effect, interact, interaction, advise) that are not associated with the target entities, the model fails to predict the correct label (examples #7 and #8).

**Chemical-protein relation extraction.** Being a common source of mis-classification, the CPR:3 type was often predicted as CPR:4 and vice versa (examples #9 and #10). The CPR:3 class usually describes up-regulation, and its instances usually include up-regulation words such as “promote”, “increase”, and “activate”. The CPR:4 class is usually related to down-regulation and contains down-regulation words such as “decrease”, “inhibitor”, and “deposition”. Having both up-regulation and down-regulation words in the same sentence creates confusion, which can lead to mis-classification. The model also misclassified some instances due to the presence of multiple entities in a single sentence (example #11). Multiple entities can also create noise and make it difficult for the model to identify if there is a relation between the two target entities.

**Disease-protein relation extraction.** We found that our model fails to predict the correct label for instances (examples #12, #13, #14 and #15) that contain association words (e.g., associated)
Table 3: Examples of sentences that were incorrectly classified by the MTFT model.

| Example | AIMed_sentence: Chemokines that could compete with high affinity for MIP-1beta binding could also compete for monomeric gp120 binding, although with variable potencies; maximal @PROTEINS binding inhibition was 80% for MCP-2, but only 30% for @PROTEINS. Gold label: TRUE Predicted label: FALSE |
| AIMed_sentence: We investigated whether @PROTEINS, which binds to tyrosine-phosphorylated ITAM, interacts with @PROTEINS following T cell activation. Gold label: FALSE Predicted label: TRUE |
| AIMed_sentence: We further demonstrated that @PROTEINS and E3 but not @PROTEINS can decrease the fusogenic activity of Abeta(29-42) via a direct interaction. Gold label: FALSE Predicted label: TRUE |
| Biolnder_sentence: In localization studies with mammalian cells, all fusion proteins showed the localization expected for @GENE. Gold label: TRUE Predicted label: TRUE |
| DDI_sentence: Other drugs which may enhance the neuromuscular blocking action of @DRUGS such as MIVACRON include certain antibiotics (e.g., aminoglycosides, tetracyclines, bacitracin, @DRUGS, lincomycin, clindamycin, colistin, and sodium colistimethate), magnesium salts, lithium, local anesthetics, procainamide, and quinidine. Gold label: INT Predicted label: EFFECT |
| DDI_sentence: @DRUGS may decrease the effectiveness of oral contraceptives, certain antibiotics, @DRUGS, theophylline, corticosteroids, anticoagulants, and beta blockers. Gold label: INT Predicted label: EFFECT |
| DDI_sentence: Drugs Eliminated by Active Tubular Secrecion: Although studies to assess drug-drug interactions with Sanhueta have not been conducted, @DRUGS has the potential for pharmacokinetic interactions with other drugs that are eliminated by active tubular secretion (e.g. digoxin, procainamide, panceuronium, morphine, @DRUGS, metformin and tenofovir). Gold label: MECHANISM Predicted label: INT |
| DDI_sentence: Since Celontin (@DRUGS) may interact with concurrently administered @DRUGS, periodic serum level determinations of these drugs may be necessary (eg methoxuximide may increase the plasma concentrations of phenytoin and phenobarbital). Gold label: ADVISE Predicted label: INT |
| ChemProt_sentence: Etvn-50 possessed a broad spectrum of in vitro anticancer activity for those tested cancer cells, especially sensitive to MDA-MB-435, SKOV-3, BXPC-3, SMMC-7721, MCF-7, HO-8910, SGC-7901, BEL-7402, HCT-116, and 786-O, with the respective IC50 below 10mg/ml. Treatment with @CHEMICALS or VB1 resulted in arresting the MDA-MB-435 and SMMC-7721 cells at G2/M phase, which was further supported by observations of increased phosphorylation of Histone 3 at Ser10, phosphorylation of @GENES at Tyr15, expression of cyclin B1, and decreased expression of Cdc25c. Gold label: CPR:3 Predicted label: CPR:4 |
| ChemProt_sentence: @CHEMICALS also increases Amyloid b (@GENES) deposition and tau pathology. Gold label: CPR-4 Predicted label: CPR-3 |
| ChemProt_sentence: Agonist and antagonist actions of yohimbine as compared to @CHEMICALS at alpha(2)-adrenergic receptors @GENES, serotonin (5-HT)(1A), 5-HT(1B), 5-HT(1D). Gold label: CPR-5 Predicted label: CPR-6 |
| GAD_sentence: Our results possibly indicate an association of @DISEASES with @GENES homozygosity (P=0.056). Gold label: FALSE Predicted label: TRUE |
| GAD_sentence: Our results suggest that the @GENES 108Hs variant is associated with reduced susceptibility to @DISEASES. Gold label: FALSE Predicted label: TRUE |
| GAD_sentence: Our results indicate that the intron 2 CYP46 @GENES genotype may predispose to @DISEASES, and this association is independent of the apolipoprotein E genotype. Gold label: FALSE Predicted label: TRUE |
| GAD_sentence: Although there remains a possibility that the @GENES TaqI A polymorphism plays some role in modifying the phenotype of the @DISEASES, these results suggest that neither the A1 allele nor the homozygous A1 genotype is associated with alcoholism. Gold label: FALSE Predicted label: TRUE |

4 Conclusion

In this paper, we present a comprehensive evaluation of encoder-only and encoder-decoder transformers on four benchmark biomedical RE tasks. We also explored the use of MTFT to investigate the correlation among these biomedical RE tasks. For that, we used ten popular datasets, namely AIMed, BioInfer, HPRD50, IEPA, LLL, DDI, ChemProt, DrugProt, GAD, and EU-ADR. The experiments showed that T5 and MTFT achieved better performance than BERT-based models (BioBERT and PubMedBERT) in extracting relations between bio-entities from texts. In the future, we plan to study the impact of each RE task/dataset on the downstream performance.

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