Detecting Anatomical and Functional Connectivity Relations in Biomedical Literature via Language Representation Models

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Abstract

Understanding of nerve-organ interactions is crucial to facilitate the development of effective bioelectronic treatments. Towards the end of developing a systematized and computable wiring diagram of the autonomic nervous system (ANS), we introduce a curated ANS connectivity corpus together with several neural language representation model based connectivity relation extraction systems. We also show that active learning guided curation for labeled corpus expansion significantly outperforms randomly selecting connectivity relation candidates minimizing curation effort. Our final relation extraction system achieves $F_1 = 72.8\%$ on anatomical connectivity and $F_1 = 74.6\%$ on functional connectivity relation extraction.

1 Introduction

The NIH Common Fund’s Stimulating Peripheral Activity to Relieve Conditions (SPARC) program aims to transform our understanding of nerve-organ interactions to help spur the development of effective bioelectronic treatments. Bioelectronic medicine represents the convergence of molecular medicine, neuroscience, engineering and computing to develop devices to diagnose and treat diseases (Olofsson and Tracey, 2017). One of the projects within this large consortium is to create a systematized and computable wiring diagram of the autonomic nervous system, a part of the “wiring system” that travels throughout the body transmitting messages between the peripheral organs and the brain or spinal cord. While diagrams of nerves are currently available in medical texts (Standring and Gray, 2008), the SPARC program seeks to map these connections at higher levels of detail and with greater accuracy. Additionally, the diagrams in these medical texts are not generally queryable, nor are they sufficiently detailed to include the granular paths that these nerves travel. Such information would be needed, for example, to understand where reliable access points to a particular nerve might be so that stimulation only affects the most relevant nerve or to understand the mechanisms behind stimulation applied at particular locations. Many scientific studies contain information about individual nerves and at times the paths they traverse, but to our knowledge, no systematic approach has been attempted to bring these large quantities of information together into a computationally accessible format.

The SPARC project is building a cross-species connectivity knowledge base that contains detailed information about individual nerves, their pathways, cells of origin and synaptic targets. To date, this knowledge base has been populated through the development of detailed models of circuitry by experts funded through the SPARC project using the ApiNATOMY platform (Kokash and de Bono, 2021). ApiNATOMY provides a modeling language for representing the complexity of functional and anatomical circuitry in a standardized form. The circuitry contained in these models represent expert knowledge derived from the synthesis of the expert’s own work and the synthesis of, in some cases, hundreds of scientific publications. However, to ensure that information in the SPARC knowledge base is comprehensive and up to date, i.e., it represents the current state of knowledge about autonomic nervous system (ANS) connectivity, we sought to augment the expert-based model approach with experimental information derived from the primary scientific literature. As there are thousands of papers and additional sources like text books, we utilized natural language processing to identify sentences that contained information on neuronal connectivity in the ANS.
The task was approached by first gathering the relevant scientific literature by matching bodily structures at a variety of anatomical levels (i.e. gasserian ganglion, vagus nerve, brainstem, etc.) from a constructed set of vocabulary at sentence level. Then, annotators classified each structure to structure relationship using only the information provided within the sentence based on the connectivity types defined in our annotation guideline. This structured data were then used to train our connectivity relations models. Data from two curators was used to assess the inter-curator agreement to determine if the annotation guidelines are sufficient to “teach” the task to humans. We assessed connectivity statements into several types including, anatomical connectivity, functional connectivity, structural connectivity, topological connectivity and general connectivity as well as no connectivity. The general connectivity and no connectivity categories can be thought of as statements that are too vague to be of much direct use for our use case. The most important statements are anatomical connectivity, elucidating which parts are connected physically and functional connectivity, elucidating which parts are connected functionally. A definition and an example for each connectivity type used for annotation is shown in Table 1. Of course with single sentences, it is difficult to define a direct functional relationship, which typically rests on the latency with which a signal is detected between two elements (Bennett, 2001). However, statements about latency are very rare in the subset of the peripheral nervous system literature, whereas somewhat more general statements about functional relationships that, for example, describe damage to one area and altered functioning in another, are more abundant. We hypothesize that when such statements are reasonably abundant, a detection classifier will be easier to train.

In relation extraction, long-range relations are usually handled using dependency parse tree information. In traditional feature-based models, paths in the dependency parse tree between entities are used used as features (Kambhatla, 2004) which suffered from the sparsity of the feature patterns. More recently, neural models are increasingly employed for relation extraction instead of feature engineering using vectorized word embeddings. The dependency information is represented as computation graphs along the parse tree (Zhang et al., 2018). Sequence models, on the other hand, work at the surface level and represent long distance relationships via either convolutional or recurrent neural networks and an attention mechanism (Zhang et al., 2017).

In biomedical domain, relation extraction work is traditionally focused on protein-protein, gene-disease or protein-chemical interactions. Several labeled datasets, such as GAD (Bravo et al., 2015) (a gene-disease relation dataset) and CHEMPROT (Krallinger et al., 2017) (a protein-chemical multi-relation dataset) are publicly available. Neural sequence models have also been applied to protein-chemical relation extraction task (Lim and Kang, 2018).

Recently, sentence level transformer based language representation models such as BERT (Devlin et al., 2019) have shown superior downstream performance on many NLP tasks. A biomedical domain adapted version of BERT called BioBERT (Lee et al., 2019) has been shown state of the art performance on several biomedical relation extraction tasks.

While most of the transformer based language representation models are pretrained on sentences where a predefined percentage of the tokens are masked and the model learns to predict the masked tokens, a recently introduced language representation model, ELECTRA (Clark et al., 2020) learns to discriminate if a token in the original input is replaced by a language generator model or not. The generator model is a BERT like generative model that is co-trained with the discriminative model.

While there are efforts to extract brain connectivity information from neuroscience literature (Richardet et al., 2015), their focus is in the cognitive parts of the brain instead of ANS. In this paper, we introduce a labeled ANS connectivity corpus, together with four biomedical domain adapted ELECTRA models, that we have used to develop an anatomical and functional connectivity relation extraction system that outperforms BioBERT.

2 Methods

2.1 Vocabulary

In order to better structure information from papers, anatomical structure labels were drawn from a set of relevant ontologies, also approved for use by the SPARC project. These ontology terms include primarily FMA (RRID:SCR_003379), UBERON (RRID:SCR_010668), and NIFSTD (RRID:SCR_005414) terms, and they are listed
Relation | Definition | Example
--- | --- | ---
functional | a relationship was determined to exist between two structures using physiological techniques | The HB reflex is a reflex initiated by lung inflation, which excited the myelinated fibers of vagus nerve, pulmonary stretch receptors [11,19].
anatomical | a physical synaptic relationship was observed between two structures using anatomical techniques such as tract tracing | Only the most prominent nervous connections, such as the *penis nerve cord* (pnc, Fig. 8a), connecting the *ventral ganglion* to the penis ganglion can be detected.
structural | a relationship that reflected continuity between segments of nerves | The term vocal fold paralysis (VFP) refers to the reduced or absent function of the *vagus nerve* or its distal branch, the *recurrent laryngeal nerve* (RLN) [1-3].
topological | a relationship that reflected the course of a nerve | Oculomotor nerve (III) exited from the middle tectum nearby ventro-medial *midbrain* and was observed on 6-day-old fish.
genral | a statement that contained general information about connectivity but did not specify the technique used or otherwise failed to elucidate the exact type of connectivity discussed | Moreover, an interoceptive circuit connecting the *gut* to the *nucleus tractus solitarius* (NTS) via the vagus nerve has been demonstrated to convey the state of the gut to the limbic system (Figure 9; Maniscalco and Rinaman, 2018).

Table 1: Connectivity relation types

on the SPARC anatomy working group web pages, which include term lists. In order to provide a more targeted set of sentences for training, we selected a set of terms that was specifically associated with the ANS. These terms included sympathetic and parasympathetic nerves and ganglia from the FMA and UBERON. Terms were selected by the SPARC Anatomical Working Group, a group of anatomical experts who provide expertise to the SPARC knowledge engineers.

2.2 Corpus Generation

The sentences of interest for connectivity relation extraction were detected by longest phrase match from the target vocabulary of anatomical terms. We have used four million full-length PMC open access papers downloaded on November 2020 to search for sentences of interest. All the sentences mentioning at least two distinct anatomical structures from our vocabulary are selected. To focus our curation effort to a manageable portion of the vocabulary, a smaller vocabulary consisting of only ANS nerve and ganglion terms were selected to further filter the candidate sentence set where only sentences having at least one structure from the focused vocabulary set is selected. Since the resulting candidate set was still too large for curation, up to three examples from each unique focused vocabulary term encountered is randomly sampled to create our base corpus of 808 sentences to be curated.

2.2.1 Annotation of the Corpus and Inter-annotator Agreement

Three curators/domain experts were involved in the connectivity corpus labeling process. Our main curator (J.M.) is a full-time curator with several years of experience with biomedical named entity recognition, relation extraction and text classification curation tasks. Dr. A.B. is a neurophysiologist by training with expertise in microcircuitry. Dr. M.M. is a trained anatomist with expertise in microcircuitry and a professor of neuroscience. All curators annotated training sets using the relation annotator tool developed in-house. The tool allows the curators both to edit entities (if automatically detected anatomical structure boundaries are not correct) and to label automatically generated binary relation combinations arising from the two or more anatomical structures detected in the curated sentence.

To start, A.B. and J.M. completed 30 sentences together to gauge the difficulty of the task and train J.M. on the differences between anatomical and functional connectivity. Then, M.M. and J.M. independently annotated 102 relation labels. In this first iteration, connectivity between structures
could only be classified as anatomical connectivity, functional connectivity, or no relation. The initial inter-annotator agreement was 66.7%; Cohen’s kappa was 0.25. Even when simply comparing binary connectivity vs. no relation, there was only an inter-annotator agreement of 72%; Cohen’s kappa was 0.34. After discussing disagreements, additional connectivity types were added to the relation annotator tool. After expansion, connectivity could be classified as structural, topological, or general in addition to the previous versions labels: anatomical, functional, and no relation. This was done to make each connectivity type more explicit with less potential overlap, especially between our main connectivity types of interest (anatomical vs. functional).

In our second iteration, M.M. and J.M. independently annotated another 170 relation labels across 100 sentences. This time, the inter-annotator agreement was 73.5%; Cohen’s kappa was 0.25. Annotation differences though were primarily found to be between our general connectivity and no relation tags. If we consider general connectivity to be the same as no relation (collapsing them together), then our inter-annotator agreement jumps to 91.2%; likewise, Cohen’s kappa also increases to 0.55. Because our primary disagreements were between two tags of less interest (general connectivity vs. no relation), we believe our inter-annotator agreement is acceptable for this high difficulty task.

2.3 Models

2.3.1 ELECTRA based language representation models for Biomedical Domain

Domain specific language representation models result in performance improvements on downstream NLP tasks as demonstrated by BioBERT (Lee et al., 2019). Similarly, we have pretrained four ELECTRA (Clark et al., 2020) based models on biomedical corpus.

For pretraining corpus we have used both PubMed abstracts and PubMed Central (PMC) open access full-length papers. 21.2 million PubMed abstracts from the January 2021 baseline distribution are used to build our main pretraining corpus. Sentences extracted from the paper title and abstract text resulted in a corpus of 3.6 billion words. For the PMC open access papers, sentences extracted from all sections except the references section of the full-length papers are used to build a 12.3 billion words corpus. A domain specific word piece vocabulary is generated using SentencePiece byte-pair-encoding (BPE) model (Sennrich et al., 2016) from PubMed abstract texts. The models are pretrained for one million steps on the PubMed abstracts corpus followed by 200,000 steps training on the PMC open access papers corpus.

During training, ELECTRA uses a small transformers based encoder model using masked language objective like in BERT to generate possible replacements for the larger discriminative model which is also based on transformers architecture. Both models are trained jointly. During fine-tuning, only the discriminative model parameters are used. The discriminative model has essentially the same architecture as BERT but trained in a discriminative manner using a different objective. We have trained three different model sizes; a base model with embedding and hidden size of 768, 12 attention heads and 12 transformer layers; a mid sized model with embedding size of 384, hidden size of 512, 8 attention heads and 12 transformer layers; a mid sizes tall model having same parameters as the mid sized model but with 24 transformer layers. We have also trained another mid sized model with the combined PubMed abstract and PMC open access full paper corpus instead of the two corpus cascaded training approach used for the other three Bio-ELECTRA models. For all models, the maximum allowed input sequence length was set to 512. For all models besides the mid-tall model, the batch size was set to 256. The mid-tall model had a batch size of 128 because of the memory limitations of a single tensor processing unit (TPU). The model architectures, sizes and training times are summarized in Table 2. All the models are trained on a single 8 core version 3 TPU with 128 GB RAM.

While, we have used our Bio-ELECTRA models for connectivity relation extraction only, the models like BioBERT are applicable to many downstream biomedical NLP tasks.

3 Experiments

We conducted our experiments in two phases. In the first phase, all the binary connectivity relation candidates in the 805 sentences extracted from the open access subset of PubMed Central is annotated by a curator. The curated base set is then randomly split into 80/20% train/test set. Afterwards, ten randomly initialized models are trained. The reported results are average of 10 runs together with
| Model            | Params | Architecture       | Steps | Train Time/Hardware |
|------------------|--------|--------------------|-------|---------------------|
| Mid              | 50M    | hidden:512, layers:12 | 1.2M  | 6.5d on 8 TPUv3s    |
| Base             | 110M   | hidden:768, layers:12 | 1.2M  | 12.5d on 8 TPUv3s   |
| Mid-tall         | 88M    | hidden:512, layers:24, batch:128 | 1M    | 5.5d on 8 TPUv3s    |
| Mid Combined     | 50M    | hidden:512, layers:12 | 1.2M  | 6.5d on 8 TPUv3s    |

Table 2: ELECTRA Models for Biomedical Domain

standard deviation. In the second phase, the base set is enhanced via active learning.

As our baseline model, we have used a graph convolution over dependency parse tree neural model (Zhang et al., 2018) where the dependency graph structure is represented by an adjacency matrix over which convolution operations are performed. The model uses word embedding vectors for input encoding and stacked layers of graph convolution network (GCN) layers to encode relations. The input encodings can be further contextualized via a bi-directional long-short-term memory (LSTM) layer, which we have used in our experiments. For word embeddings we have used 300 dimensional Common Crawl (840B tokens) trained GloVe (Pennington et al., 2014) vectors. The dependency parse trees for the input sentences were generated via Stanford CoreNLP (Manning et al., 2014) package.

All the other models are fine-tuned from pre-trained transformers based language representation models. We have downloaded Bio-ELECTRA++ from Zenodo\(^1\). Besides our four biomedical corpus pretrained ELECTRA models, we have used BioBERT (Lee et al., 2019) version 1.1 and ELECTRA Base models. The binary anatomical structure entities are masked in candidate sentences as in (Zhang et al., 2018; Lee et al., 2019). Besides that, no further preprocessing is done. All the models are trained for three epochs, using the the default learning rate and maximum allowed batch size for our 8GB Nvidia RTX 2070 GPU.

The test performance of models tested are summarized in Table 3. Even after the benefit of dependency parses, contextualized graph convolution networks were at the bottom of the performance rank tying with the smallest language representation model. Two Bio-ELECTRA models, namely Bio-ELECTRA Base and Bio-ELECTRA Mid outperformed BioBERT. Given that the Bio-ELECTRA Mid has less than half the parameters of Bio-BERT, its performance is especially impressive. We chose the best performing Bio-ELECTRA Base model for the second stage.

3.1 Extending Curation Set via Active Learning

Since labeled data set generation is costly and time consuming, we have tried to leverage active learning to minimize curation effort while trying to maximize prediction performance. To this end, 250 randomly selected candidate sentences from the nerve-ganglia PMC data set, are interactively curated by our curator in ten iterations. Each iteration has consisted of 25 candidate sentences selected by the binary relation extraction classifier trained on all the curated sentences from the previous iterations plus the base training set. In the first iteration, the classifier is trained on the base training set only. For the control set, we have randomly selected 250 candidate sentences from the nerve-ganglia PMC data set, which are annotated separately by our curator. We have used uncertainty sampling as our oracle query strategy where the 25 unlabeled sentences that are closest to the decision boundary (probability estimate of 0.5) are selected for curation at each iteration. After each iteration, the extended training set is used to train ten randomly initialized models which are tested on the testing set. The precision and \(F_1\) performance scores over the active learning set is shown in Figure 1.

The testing performance of active learning based vs random selection based training set expansion is shown in Table 4. Active learning strategy was significantly better than random selection based on two-tailed t test.

3.2 Effect of Hyperparameter Optimization

The additional 500 curated sentences (250 from active learning, 250 from random control set) are combined with the base training set. To maximize relation extraction performance, we used hyperparameter tuning on the 80%/20% training/dev set split of the combined training set. Using hyper-

\(^1\)https://doi.org/10.5281/zenodo.3971235
| Model                  | Parameters | Precision | Recall | $F_1$ |
|------------------------|------------|-----------|--------|-------|
| Contextualized-GCN     |            | 71.05 (4.36) | 54.23 (4.20) | 61.36 (3.01) |
| ELECTRA Base           | 110M       | 69.35 (4.23) | 70.85 (5.43) | 70.03 (4.39) |
| BioBERT                | 110M       | 67.82 (4.71) | 72.34 (2.18) | 69.89 (2.40) |
| Bio-ELECTRA++          | 11M        | 54.41 (2.11) | 70.32 (3.38) | 61.26 (1.33) |
| Bio-ELECTRA Mid        | 50M        | 69.16 (3.53) | 73.83 (2.24) | 71.36 (2.16) |
| Bio-ELECTRA Base       | 110M       | 69.93 (2.91) | 74.26 (3.55) | 71.99 (2.76) |
| Bio-ELECTRA Mid Combined | 50M   | 67.66 (2.38) | 74.36 (5.80) | 70.70 (2.78) |
| Bio-ELECTRA Mid-tall   | 88M        | 63.89 (4.51) | 65.96 (3.81) | 64.78 (2.98) |

Table 3: Binary connectivity/no-connectivity relation extraction on base set

| Data Set                | Precision | Recall | $F_1$ |
|-------------------------|-----------|--------|-------|
| Random                  | 70.29 (1.69) | 74.04 (3.27) | 72.06 (1.68) |
| Active learning         | **75.88 (2.70)** | **75.11 (2.39)** | **75.47 (2.30)** |

Table 4: Test performance effect for active learning vs random selection based labeled set expansion

Figure 1: Average test performance over active learning iterations

To detect anatomical and functional connectivity relations among candidate structure binary relation sentences, we have introduced a three class classifier based on the same Bio-ELECTRA Base language representation model as the connectivity/no-connectivity classifier. Ten randomly initialized classifiers are trained using optimized hyperparameters. The test performance is shown in Table 6.

4 Discussion

Connectivity relations constituted only about 12% of the connectivity relationship candidates in our corpus. Taking this into account, the anatomical and functional connectivity detection performance of our final classifier is good enough to be used for ANS connectivity knowledge base construction with drastically reduced domain expert curation.

When looking at our model’s performance, we considered errors at the level of individual connectivity relations labels, meaning we could (and did) have some sentences with multiple errors. We defined errors as cases where relation labels tagged by the model and the annotator did not agree. We performed our error analysis in two phases. In phase 1, our analysis was performed using only binary connectivity data (i.e., did annotators mark a relationship using any type of connectivity or as no relation) from 40 connectivity errors: 19 false positives and 21 false negatives. False positives were defined as instances when the model predicted connectivity when there was actually no relationship (as defined by the annotator). Inversely, false negatives were cases when the model predicted structures to have no relationship when there was actually some type of connectivity. In phase 2, our
Table 5: Connectivity/No-Relation test performance on the extended training set

| Model                        | Precision | Recall | F1     |
|------------------------------|-----------|--------|--------|
| Bio-ELECTRA Base (default)   | 76.97 (2.72) | 74.68 (2.32) | 75.77 (1.99) |
| Bio-ELECTRA Base (opt)       | 77.32 (2.39) | 77.98 (1.65) | 77.62 (1.33) |

Table 6: Anatomical/functional connectivity test performance on the extended training set

| Relation                  | Precision | Recall | F1     |
|---------------------------|-----------|--------|--------|
| Anatomical connectivity   | 68.93 (2.94) | 77.12 (1.37) | 72.77 (1.99) |
| Functional connectivity   | 82.79 (2.39) | 68.00 (2.80) | 74.61 (2.35) |

analysis was performed on anatomical connectivity errors (18 false positives; 15 false negatives) and functional connectivity errors (5 false positives; 11 false negatives). With both phases, we noticed patterns emerging among the errors, although in most cases, these errors were present across all connectivity types. In other words, there was very little difference between the errors seen in phase 1 vs. the errors seen in phase 2.

The first identified error pattern was mislabeled data due to human error. We noticed 4 instances where the data was mislabeled. The second type of error occurred because a solid line of demarcation between connectivity types was difficult to establish due to ambiguities in our curation guidelines and the overall difficulty of the task. When we began annotating, connectivity between structures could only be defined as anatomical, functional or having no relation. After discussing the differences in our annotations though, it became apparent we needed to add additional connectivity types to clarify the lines of demarcation between each. As a result, we added structural, topological, and general connectivities, and while we did see improved classifier performance after adding these, it appears we weren’t entirely successful in our attempts to explicate our connectivity types. In the example,

“Chemoreceptors in the carotid body or aortic body in the walls of the internal carotid artery or the aorta sense the level of oxygen or carbon dioxide in the blood and convey these signals via the glossopharyngeal and vagus nerves to the nucleus of the tractus solitarius.”

Just from a cursory glance, it becomes obvious that this sentence is complicated; it contains multiple subject and verb phrases clouded by prepositional phrases. Unfortunately, the convoluted nature of the sentence hurts readability for both humans and machines. Because humans also tend to have issues understanding these highly complex sentences, we feel the best solution is for authors to limit the complexity of their sentences to reasonable levels when possible. If a sentence is too complex for a human to understand, it will most likely be too complex
for a computer. Additionally, we noticed persistent issues when subjects were not explicit. Unresolved pronouns (e.g. pronouns whose antecedents are unknown) and ambiguous body structures (i.e. fibers) tended to cause errors wherein the model would correctly identify that the sentence contained connectivity but would incorrectly identify which structures are connected. With regards to verb usage, our model seemed to perform better when the connectivity between structures was described in active voice rather than passive. One potential explanation is that sentences using active voice tend to be more clear and simple than sentences using passive voice. Lastly, our model seemed to perform worse the further apart the two connecting structures were within the sentence.

5 Conclusions

In this paper, we introduced a labeled corpus for ANS connectivity relations which is further expanded via active learning. The labeled ANS connectivity relation corpus is used to develop relation extraction systems mostly based on language representation neural models. We have introduced four biomedical domain pretrained ELECTRA (Clark et al., 2020) based discriminative language representation models, two of which have outperformed BioBERT (Lee et al., 2019) on the ANS connectivity relation extraction task. Using active learning guided curation, the labeled corpus is expanded minimizing the curation effort while significantly improving ANS connectivity relation extraction performance.

Based on the observed benefits of the active learning, we are planning to use our Bio-ELECTRA based relation extraction system in a web based tool for ANS connectivity knowledge base construction with active learning based continuous learning ability.

Software and Data Availability

All pretrained Bio-ELECTRA models are available on Zenodo (https://doi.org/10.5281/zenodo.4699034). The labeled connectivity corpus and codebase including the connectivity relation annotation tool are available on Github (https://github.com/SciCrunch/connectivity-re).

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Conflicts of interest

AB, MM, and IBO are co-founders of SciCrunch Inc, a company devoted to improving scholarly communication.

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