"Heavy-Light Chain Pair Identification in Antibodies using BERT (work title)"

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Abstract

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Glossary

Word Embeddings Mapping of words into vectors with real numbers.

1. Introduction

- 1.1. Background
- 1.2. Objectives
- 1.3. Scope of the Topic
- 1.4. Research Question
- 1.5. Nature of the Thesis
- 1.6. Relevance of the Thesis

2. Sequential Transfer Learning in NLP for Antibody Research

The following section explains the theoretical background necessary for the thesis with regard to the biology of antibodies and the functionality of the NLP models used. A basic understanding of machine learning and protein structures is assumed.

2.1. Introduction to Antibodies

Antibodies are Y-shaped proteins that consist of two identical light chains (LCs) and two identical heavy chains (HCs) (Chiu, Goulet, Teplyakov, & Gilliland, 2019).

2.2. Antibody Engineering and Therapeutic Applications

2.3. Deep Learning Methods for Antibody Research

Deep learning is a branch of machine learning that focuses on algorithms capable of identifying complex patterns in data by transforming low-level inputs (like pixels in an image) into high-level features (such as object shapes). It utilizes artificial neural networks (ANNs) with multiple layers between the input and output, making them "deep". These networks consist of nodes, or neurons, that process inputs and pass the outputs to subsequent layers, gradually extracting more abstract features. In the context of biochemistry, deep learning can start from basic data, like amino acid seguences, and learn to recognize complex biological structures or functions (Graves et al., 2020). NLP models can be effectively used for analyzing amino acid sequences due to the conceptual similarities between proteins and language. Proteins can be represented as strings of 20 amino acid letters, making them a natural fit for many NLP methods. This similarity in representation allows for the application of NLP algorithms to the study of proteins, leveraging the success and promise of NLP methods in other domains (Ofer, Brandes, & Linial, 2021). NLP methods have been successfully applied to protein sequences for tasks such as predicting protein families or properties. Word embedding models in NLP have been used to extract features of protein sequences and have demonstrated successful applications in

protein family classification (Xu et al., 2020). This can be explained by the following similarities between natural language and protein sequences: Like natural language, natural proteins generally consist of reused modular elements that exhibit slight variations and can be rearranged and reassembled in a hierarchical fashion. In this analogy, common protein motifs and domains, which are the basic functional building blocks of proteins, are comparable to words, phrases and sentences in human language. Another similarity between proteins and human language is the completeness of the information. A protein is more than just a sequence of amino acids, it is also a three-dimensional machine with a specific structure and associated function, which is largely determined by the amino acid sequence. From an information-theory perspective, this means that the information of the protein is contained in the protein sequence. However, the analogies between proteins and human language only go so far. We can read and understand human language, but not in the same way as the protein sequence. In addition, most human languages have uniform punctuation and stop words that clearly delineate structures such as words or sentences. There is no clear analogy between the building blocks of languages and those of proteins (Ofer et al., 2021).

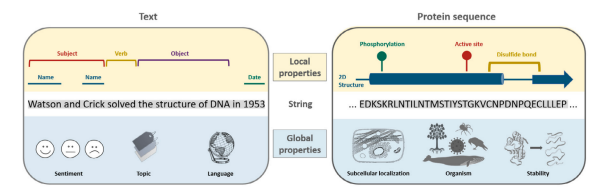


Figure 2.1.: (Ofer et al., 2021)

2.4. BERT and Transformers in Bioinformatics

BERT from Devlin, Chang, Lee, and Toutanova (2019) stands for "Bidirectional Encoder Representations from Transformers" and is based on a bidirectional language model. BERT uses the transformer according to Vaswani et al. (2017) as its architecture. At the time of publication, BERT was able to establish the state of the art in 11 natural language processing tasks (Devlin et al., 2019). The first forms of language modelling in connection

with machine learning can be found in Mikolov, Chen, Corrado, and Dean (2013) in the form of the "skip-gram" model. In the skip-gram model, text or unlabelled data is used to train the probability distribution of the next word based on the previous words in the sentence. This process can then be used to calculate static word vectors, which serve as a starting point for other NLP tasks (Mikolov, Sutskever, Chen, Corrado, & Dean, 2013). The idea of this unidirectional language model was subsequently used by various other publications and transferred to other architectures such as the Transformer according to Vaswani et al. (2017) (Radford, Narasimhan, Salimans, & Sutskever, 2018). In contrast to Radford et al. (2018), however, BERT uses a bidirectional language model. Figure 2.2 shows the structure of BERT with pretraining and fine-tuning in graphical form. The language model, or the step known as "pre-training", is trained using two tasks:

Masked language modelling: Since the words of the sentence are processed in parallel in the transformer architecture Vaswani et al. (2017), individual words must be masked in bidirectional prediction. In the case of BERT, these are replaced with the token "[MASK]". The model is then trained to correctly predict these masked words.

Next Sentence Prediction: In order to encode connections between whole sentences in the language model, "Next Sentence Prediction" is used in addition to "Masked Language Modelling". For two sentences A and B, in 50% of the sentences, a sentence B is actually used, which occurs in the text as a directly following sentence after A, and in 50% of the cases a random other sentence is used, which is taken from the corpus. The model must then predict whether these sentences belong together or not.

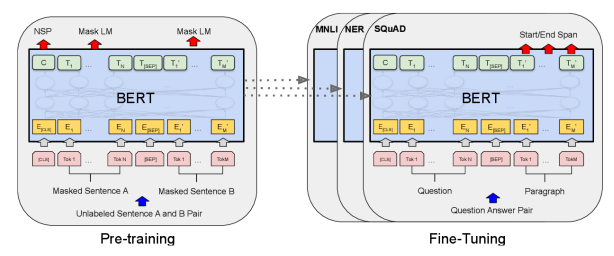


Figure 2.2.: BERT Overview (Devlin et al., 2019).

2.5. Heavy-Light Chain Pair Identification

- 2.5.1. Gap in the Literature
- 2.5.2. Conclusion

3. Materials & Methods

4. Results

5. Discussion

6. Conclusion

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A. Appendix

A.1. Link to the Code

The link to the entire code of this thesis can be found at: https://github.com/ibmm-unibe-ch/OAS_paired_sequences_cls.git

A.2. Declaration of Independence