WNUT-2020 Task 1 Overview: Extracting Entities and Relations from Wet Lab Protocols

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Abstract

This paper presents the results of the wet lab information extraction task at WNUT 2020. This task consisted of two sub tasks: (1) a Named Entity Recognition (NER) task with 13 participants and (2) a Relation Extraction (RE) task with 2 participants. We outline the task, data annotation process, corpus statistics, and provide a high-level overview of the participating systems for each sub task.

1 Introduction

Wet Lab protocols consist of natural language instructions for carrying out chemistry or biology experiments (for an example, see Figure 1). While there have been efforts to develop domain-specific formal languages in order to support robotic automation\(^1\) of experimental procedures (Bates et al., 2017), the vast majority of knowledge about how to carry out biological experiments or chemical synthesis procedures is only documented in natural language texts, including in scientific papers, electronic lab notebooks, and so on.

Recent research has begun to apply human language technologies to extract structured representations of procedures from natural language protocols (Kuniyoshi et al., 2020; Vaucher et al., 2020; Kulkaeni et al., 2018; Soldatova et al., 2014; Vasilev et al., 2011; Ananthanarayanan and Thies, 2010). Extraction of named entities and relations from these protocols is an important first step towards machine reading systems that can interpret the meaning of these noisy human generated instructions.

However, performance of state-of-the-art tools for extracting named entity and relations from wet lab protocols still lags behind well edited text genres (Jiang et al., 2020). This motivates the need for continued research, in addition to new datasets and tools adapted to this noisy text genre.

1https://autoprotocol.org/

Figure 1: Examples of named entities and relations in a wet lab protocol

In this overview paper, we describe the development and findings of a shared task on named entity and relation extraction from the noisy wet lab protocols, which was held at the 6-th Workshop on Noisy User-generated Text (WNUT 2020) and attracted 15 participating teams.

In the following sections, we describe details of the task including training and development datasets in addition to the newly annotated test data. We briefly summarize the systems developed by selected teams, and conclude with results.

2 Wet Lab Protocols

Wet lab protocols consist of the guidelines from different lab procedures which involve chemicals, drugs, or other materials in liquid solutions or volatile phases. The protocols contain a sequence of steps that are followed to perform a desired task. These protocols also include general guidelines or warnings about the materials being used. The publicly available archive of protocol.io contains such guidelines of wet lab experiments, written by researchers and lab technicians around the world. This protocol archive covers a large spectrum of experimental procedures including neurology, epigenetics, metabolomics, stem cell biology, etc. Figure 1 shows a representative wet lab protocol.

The wet lab protocols, written by users from all over the worlds, contain domain specific jargon as well as numerous nonstandard spellings, abbreviations, unreliable capitalization. Such diverse and
noisy style of user created protocols imposed crucial challenges for the entity and relation extraction systems. Hence, off-the-shelf named entity recognition and relation extraction tools, tuned for well edited texts, suffer a severe performance degradation when applied to noisy protocol texts (Kulkarni et al., 2018).

To address these challenges, there has been an increasing body of work on adapting entity and relation extraction recognition tools for noisy wet lab texts (Jiang et al., 2020; Luan et al., 2019; Kulkarni et al., 2018). However, different research groups have used different evaluation setups (e.g., training / test splits) making it challenging to perform direct comparisons across systems. By organizing a shared evaluation, we hope to help establish a common evaluation methodology (for at least one dataset) and also promote research and development of NLP tools for user generated wet-lab text genres.

2.1 Annotated Corpus

Our annotated wet lab corpus includes 726 experimental protocols from the 8-year archive of ProtocolIO (April 2012 to March 2020). These protocols are manually annotated with 15 types of relations among the 18 entity types\(^2\). The fine-grained entities can be broadly classified into 5 categories: ACTION, CONSTITUENTS, QUANTIFIERS, SPECIFIERS, and MODIFIERS. The CONSTITUENTS category includes mentions of REAGENT, LOCATION, DEVICE, MENTION, and SEAL. The QUANTIFIERS category includes mentions of AMOUNT, CONCENTRATION, SIZE, TIME, TEMPERATURE, pH, SPEED, GENERIC-MEASURE and NUMERICAL. The SPECIFIERS category includes mentions of MODIFIER, MEASURE-TYPE and METHOD. The ACTION entity refers to the phrases denoting tasks that are performed to complete a step in the protocol. The mentions of these entities contain different types of relations, including– SITE, SETTING, CREATES, MEASURE-

\(^2\)Our annotated corpus is available at: https://github.com/jeniyat/WNUT_2020_NER.

Table 1: Statistics of the Wet Lab Protocol corpus.

|                | Train | Dev  | Test-18 | Test-20 | Total |
|----------------|-------|------|---------|---------|-------|
| #protocols     | 370   | 122  | 123     | 111     | 726   |
| #sentences     | 8444  | 2839 | 2813    | 3562    | 17658 |
| #tokens        | 107038| 36106| 36597   | 51688   | 231429|
| #entities      | 48197 | 15972| 16490   | 104654  | 185313|
| #relations     | 32158 | 10812| 11242   | 70591   | 124803|

|                | per Protocol | per Sentence |
|----------------|--------------|--------------|
| avg. #sentences| 24.32        | -            |
| avg. #tokens   | 318.77       | 13.11        |
| avg. #entities | 255.25       | 10.49        |
| avg. #relation | 171.90       | 7.07         |

TYPE-LINK, CO-REFERENCE-LINK, MOD-LINK, COUNT, MERONYM, USING, MEASURE, COMMANDS, OF-TYPE, OR, PRODUCT, and ACTS-ON.

2.1.1 Train and Development data

The training and development dataset for our task was taken from previous work on wet lab corpus (Kulkarni et al., 2018) that consists of from the 623 protocols. We excluded the eight duplicate protocols from this dataset and then re-annotated the 615 unique protocols in BRAT (Stenetorp et al., 2012). This re-annotation process aided us to add the previously missing 20,613 missing entities along with 10,824 previously missing relations and also to facilitate removing the inconsistent annotations. The updated corpus statics is provided in Table 1. This full dataset (Train, Dev, Test-18) was provided to the participants at the beginning of the task and they were allowed to use any of part of this dataset to train their final model.

2.1.2 Test Data

For this shared task we added 111 new protocols (Test-20) which were used to evaluate the submitted models. Test-20 dataset consists of 100 randomly sampled general protocols and 11 manually selected covid-related protocols from ProtocolIO (https://www.protocols.io/). This 111 protocols were double annotated by three annotators using a web-based annotation tool, BRAT (Stenetorp et al., 2012). Figure 1 presents a screenshot of our annotation interface. We also provided the annotators a set of guidelines containing the entity and relation type definitions. The annotation task was split in multiple iterations. In each iteration, an annotator was given a set of 10 protocols. An adjudicator then went through all the entity and relation annotations in these protocols and resolved the disagreements. Before adjudication, the inter-annotator agreement is 0.75, measured by Cohen’s Kappa (Cohen, 1960).

2.2 Baseline Model

We provided the participants baseline model for both of the subtasks. The baseline model for named
entity recognition task utilized a feature-based CRF tagger developed using the CRF-Suite\(^3\) with a standard set of contextual, lexical and gazetteer features. The baseline relation extraction system employed a feature-based logistic regression model developed using the Scikit-Learn\(^4\) with a standard set of contextual, lexical and gazetteer features.

2.3 NER Systems

Thirteen teams (Table 3) participated in the named entity recognition sub-task. A wide variety of approaches were taken to tackle this task. Table 2 summarizes the word representations, features and the machine learning approaches taken by each team. Majority of the teams (11 out of 13) utilized contextual word representations. Four teams combined the contextual word representations with global word vectors. Only two teams did not use any type of word representations and relied entirely on hand-engineered features and a CRF taggers. The best performing teams utilized a combination of contextual word representation with ensemble of learning. Below we provide a brief description of the approach taken by each team.

**B-NLP** (Lange et al., 2020) modeled the NER as a parsing task and uses a biaffine classifier. The second classifier of their system used the predictions from the first classifier and then updated the labels of the predicted entities. Both of the classifiers utilized word2vec (Mikolov et al., 2013) and SciBERT (Lee et al., 2019) word representations.

**BIO-BIO** (Kecheng et al., 2020) implemented a BiLSTM-CRF tagger that utilized BioBERT (Lee et al., 2020) word representation.

**BiTeM** (Knafou et al., 2020) developed a voting based ensemble classifier containing 14 transformer models, and utilized 7 different word representations including BERT (Devlin et al., 2019), ClinicalBERT (Huang et al., 2019), PubMedBERT\(_\text{base}\) (Gu et al., 2020), BioBERT (Lee et al., 2020), RoBERTa (Liu et al., 2019), Biomed-RoBERTa\(_\text{base}\) (Gururangan et al., 2020) and XLNet (Yang et al., 2019).

**DSC-IITISM** (Gupta et al., 2020) developed a BiLSTM-CRF model that utilized a concatenation of CamemBERT\(_\text{base}\) (Martin et al., 2020), Flair(PubMed) (Akbik et al., 2018), and GloVe(en) (Pennington et al., 2014) word representations.

**Fancy Man** (Zeng et al., 2020) fine-tuned the BERT\(_\text{base}\) (Devlin et al., 2019) model with an additional linear layer.

**IBS** (Sikdar et al., 2020) utilized an ensemble classifier with 4 feature based on CRF taggers.

**Kabir** (Khan, 2020) employed an RNN-CRF model that utilized concatenation of Flair(PubMed) (Akbik et al., 2018) and ELMo(PubMed) (Peters et al., 2018) word representations.

**Kaushik Acharya** (Acharya, 2020) employed a linear CRF with hand-crafted features.

**mahab** (Pour and Farinnia, 2020) fine-tuned the BERT\(_\text{base}\) (Devlin et al., 2019) sequence tagging model.

**mgsohrab** (Sohrab et al., 2020) fine-tuned the SciBERT (Beltyag et al., 2019) model.

**PublishInCovid19** (Singh and Wadhawan, 2020) employed a structured ensemble classifier (Nguyen and Guo, 2007) consisting of 11 BiLSTM-CRF taggers, that utilized the PubMedBERT (Gu et al., 2020) word representation.

**Sudeshna TCS** (Jana, 2020) fine-tuned XLNet (Yang et al., 2019) model.

**IITKGP** (Kaushal and Vaidhya, 2020) fine-tuned the Bio-BERT (Lee et al., 2020) model.

2.4 RE Systems

Two teams (Table 3) participated in the relation extraction sub-task. Both of the teams followed fine-tuning of contextual word representation and did not use any hand-crafted features. Table 5 summarizes the word representations and the machine learning approaches followed by each team. Below we provide a brief description of the model developed by taken by each team.

**Big Green** (Miller and Vosoughi, 2020) considered the protocols as a knowledge graph, in which relationships between entities are edges in the knowledge graph. They trained a BERT (Devlin et al., 2019) based system to classify edge presence and type between two entities, given entity text, label, and local context.

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\(^3\)http://www.chokkan.org/software/crfsuite/

\(^4\)https://scikit-learn.org/
Table 2: Summary of NER systems designed by each team.

| Team Name                | Affiliation                                                                 |
|--------------------------|-----------------------------------------------------------------------------|
| BiTeM                    | University of Applied Sciences and Arts of Western Switzerland, Swiss Institute of Bioinformatics, University of Geneva |
| PublishInCovid19         | Flipkart Private Limited                                                    |
| Fancy Man                | University of Manchester, Xian Jiaotong University, East China University of Science and Technology, Zhejiang University |
| mahab                    | Amirkabir University of Technology                                           |
| mgsohrab                 | National Institute of Advanced Industrial Science and Technology             |
| SadeshtaTCS              | Flipkart Private Limited                                                    |
| IITKGP                   | IIT, Kharagpur                                                              |
| BIO-BIO                  | IBS Software Pvt. Ltd. NTNU                                                 |
| DSC-IITISM               | IIT(ISM) Dhanbad                                                           |
| Kabir                    | IBS                                                                         |
| Baseline                 | IIT, Kharagpur                                                              |

Table 3: Team Name and affiliation of the participant.

| Team Name                | Affiliation                                                                 |
|--------------------------|-----------------------------------------------------------------------------|
| B-NLP                    | Bosch Center for Artificial Intelligence                                    |
| Big Green                | Dartmouth College                                                           |
| BIO-BIO                  | Harbin Institute of technology, Shenzhen                                     |
| BiTeM                    | University of Applied Sciences and Arts of Western Switzerland, Swiss Institute of Bioinformatics, University of Geneva |
| DSC-IITISM               | IIT(ISM) Dhanbad                                                           |
| Kabir                    | Microsoft                                                                   |
| KaushikAcharya           | Philips                                                                     |
| mgsohrab                 | National Institute of Advanced Industrial Science and Technology             |
| PublishInCovid19         | Flipkart Private Limited                                                    |
| SadeshtaTCS              | TCS Research & Innovation Lab                                               |

Table 4: Results on extraction of 18 Named Entity types from the Test-20 dataset.

| Team  | Exact Match  | P   | R   | F1  |
|-------|--------------|-----|-----|-----|
| BiTeM | 84.73        | 82.25 | 77.39 |
| PublishInCovid19 | 81.36 | 74.12 | 77.57 |
| Fancy Man | 76.21 | 71.76 | 73.92 |
| mahab | 50.19 | 52.96 | 51.54 |
| mgsohrab | 83.69 | 70.62 | 76.60 |
| SadeshtaTCS | 74.99 | 71.43 | 73.16 |
| IITKGP | 77.00 | 63.93 | 70.25 |
| BIO-BIO | 78.49 | 71.06 | 74.59 |
| DSC-IITISM | 64.20 | 57.07 | 60.42 |
| Kabir | 78.79 | 72.20 | 75.35 |
| IBS | 74.26 | 65.52 | 69.00 |
| KaushikAcharya | 73.68 | 63.98 | 68.48 |
| Baseline | 70.06 | 61.91 | 65.73 |

| Team  | Partial Match  | P   | R   | F1  |
|-------|----------------|-----|-----|-----|
| BiTeM | 88.72 | 75.66 | 81.67 |
| PublishInCovid19 | 85.74 | 78.11 | 81.75 |
| Fancy Man | 81.15 | 76.41 | 78.71 |
| mahab | 55.09 | 58.14 | 56.57 |
| mgsohrab | 87.95 | 74.22 | 80.50 |
| SadeshtaTCS | 70.73 | 75.95 | 77.80 |
| IITKGP | 81.76 | 77.43 | 79.54 |
| BIO-BIO | 84.85 | 69.39 | 76.46 |
| DSC-IITISM | 83.16 | 75.29 | 79.03 |
| Kabir | 79.52 | 60.90 | 64.49 |
| IBS | 83.73 | 76.73 | 79.08 |
| KaushikAcharya | 79.31 | 68.87 | 73.73 |
| Baseline | 75.66 | 66.85 | 70.98 |

mgsohrab (Sohrab et al., 2020) utilized PubMedBERT (Gu et al., 2020) as input to the relation extraction model that enumerates all possible pairs of arguments using deep exhaustive span representation approach.

3 Evaluation

In this section, we present the performance of each participating systems along with a description of the errors made by the model types.

3.1 NER Errors Analysis

Table 4 shows the comparison of precision (P), recall (R) and F1 score among different teams, evaluated on the Test-20 corpus. Here the exact match refers to the cases where a predicted entity is considered correct, only if the predicted type and boundary is exactly same as the gold entity. Whereas, in partial match, a predicted entity is considered correct if the predicted type is the same as the gold entity type and predicted entity boundary has some overlap with gold entity boundary.
Table 5: Summary of relation extraction systems designed by each team.

| Team            | Word Representation | Features                  | Approach                  |
|-----------------|---------------------|----------------------------|----------------------------|
| mgsohrab        | PubMedBERT          | -                          | PubMedBERT fine-tuning    |
| Big Green       | BERT                | -                          | BERT fine-tuning          |
| Baseline        | -                   | Gazetteers, Lexical, Contextual Logistic Regression | - |

We observe that ensemble models with contextual word representations outperforms all other approaches by achieving 77.99 F$_1$ score in exact match (Team:BiTeM) and 81.75 F$_1$ score in partial match (Team:PublishInCovid19). Fine tuning of contextual word representation systems demonstrated quite competent performance with SciBERT-fine tuning being the best (Team:mgsohrab).

In Figure 2, we present an error analysis. Among the best performing models, the ensemble of transformer (Team:BiTeM) had significantly lower amount of ‘over prediction’ error (i.e., tagging a non-entity token as entity), compared to the system with ensemble of BiLSTM-CRFs (Team:PublishInCovid19).

3.2 RE Errors Analysis

Table 6 shows the comparison of precision (P), recall (R) and F$_1$ score among the participant teams, evaluated on the Test-20 corpus. Both of the teams utilized the gold entities and then predict the relations among these entities by fine-tuning contextual word representations. We observed that fine-tuning of domain related PubMedBERT, provides significantly higher performance compared to the general domain BERT. While examining the relation predictions from both of these systems, we found that model with fine-tuned PubMedBERT (Team:mgsohrab) resulted in significantly less amount of errors in every category (Figure 3).

Table 6: Results on extraction of 15 relation types from the Test-20 dataset.

| Team            | P  | R  | F$_1$ |
|-----------------|----|----|-------|
| mgsohrab        | 80.86 | 80.07 | 80.46 |
| Big Green       | 45.42 | 86.54 | 59.57 |
| Baseline        | 80.10 | 66.21 | 72.50 |

4 Related Work

The task of information extraction from wet lab protocols is closely related to the event trigger extraction task. The event trigger task has been studied extensively, mostly using ACE data (Doddington et al., 2004) and the BioNLP data (Nédellec et al., 2013). Broadly, there are two ways to classify various event trigger detection models: (1) Rule-based methods using pattern matching and regular expression to identify triggers (Vlachos et al., 2009) and (2) Machine Learning based methods focusing on generation of high-end hand-crafted features to be used in classification models like SVMs or maxent classifiers (Pyysalo et al., 2012). Kernel based learning methods have also been utilized with embedded features from the syntactic and semantic contexts to identify and extract the biomed-
ical event entities (Zhou et al., 2014). In order to counteract highly sparse representations, different neural models were proposed. These neural models utilized the dependency based word embeddings with feed forward neural networks (Wang et al., 2016b), CNNs (Wang et al., 2016a) and Bidirectional RNNs (Rahul et al., 2017).

Previous work has experimented on datasets of well-edited biomedical publications with a small number of entity types. For example, the JNLPBA corpus (Kim et al., 2004) with 5 entity types (CELL LINE, CELL TYPE, DNA, RNA, and PROTEIN) and the BC2GM corpus (Hirschman et al., 2005) with a single entity class for genes/proteins. In contrast, our dataset addresses the challenges of recognizing 18 fine-grained named entities along with 15 types of relations from the user-created wet lab protocols.

5 Summary

In this paper, we presented a shared task for consisting of two sub-tasks: named entity recognition and relation extraction from the wet lab protocols. We described the task setup and datasets details, and also outlined the approach taken by the participating systems. The shared task included larger and improvised dataset compared to the prior literature (Kulkarni et al., 2018). This improvised dataset enables us to draw stronger conclusions about the true potential of different approaches. It also facilitates us in analyzing the results of the participating systems, which aids us in suggesting potential research directions for both future shared tasks and noisy text processing in user generated lab protocols.

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References

Kaushik Acharya. 2020. KaushikAcharya at WNUT 2020 Shared Task-1: Conditional Random Field(CRF) based Named Entity Recognition(NER) for Wet Lab Protocols. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Alan Akbik, Duncan Blythe, and Roland Vollgraf. 2018. Contextual String Embeddings for Sequence Labeling. In COLING 2018, 27th International Conference on Computational Linguistics.

Vaishnavi Ananthanarayanan and William Thies. 2010. Biocoder: A programming language for standardizing and automating biology protocols. Journal of biological engineering.

Maxwell Bates, Aaron J Berliner, Joe Lachoff, Paul R Jaschke, and Eli S Groban. 2017. Wet lab accelerator: a web-based application democratizing laboratory automation for synthetic biology. ACS synthetic biology.

Iz Beltagy, Arman Cohan, and Kyle Lo. 2019. SciBERT: Pretrained Contextualized Embeddings for Scientific Text. In Proceedings of the 2019 Conference on Empirical Methods in Natural Language Processing (EMNLP).

Jacob Cohen. 1960. A Coefficient of Agreement for Nominal Scales. Educational and Psychological Measurement.

Jacob Devlin, Ming-Wei Chang, Kenton Lee, and Kristina Toutanova. 2019. BERT: Pre-training of Deep Bidirectional Transformers for Language Understanding. In Proceedings of the 2019 Annual Conference of the North American Chapter of the Association for Computational Linguistics (NAACL).

George R Doddington, Alexis Mitchell, Mark A Przybocki, Lance A Ramshaw, Stephanie Strassel, and Ralph M Weischedel. 2004. The automatic content extraction (ace) program-tasks, data, and evaluation. In LREC.

Yu Gu, Robert Tinn, Hao Cheng, Michael Lucas, Naoto Usuyama, Xiaodong Liu, Tristan Naumann, Jianfeng Gao, and Hoi-fung Poon. 2020. Domain-Specific Language Model Pretraining for Biomedical Natural Language Processing. arXiv preprint arXiv:2007.15779.

Saket Gupta, Aman Sinha, and Rohit Agarwal. 2020. DSC-IITISM at WNUT 2020 Shared Task-1: Name Entity Extraction from Wet Lab Protocol. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Suchin Gururangan, Ana Marasović, Swabha Swayamdipta, Kyle Lo, Iz Beltagy, Doug Downey, and Noah A Smith. 2020. Don’t Stop Pretraining: Adapt Language Models to Domains and Tasks. arXiv preprint arXiv:2004.10964.

Lynette Hirschman, Alexander Yeh, Christian Blaschke, and Alfonso Valencia. 2005. Overview of biocreative: critical assessment of information extraction for biology.
Mohammad Mahdi Abdollah Pour and Parsa Farinnia. 2020. mahab at WNUT 2020 Shared Task-1: Name Entity Extraction from Wet Lab Protocol. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Sampo Pyysalo, Tomoko Ohta, Makoto Miwa, Hancheol Cho, Jun’ichi Tsujii, and Sophia Ananiadou. 2012. Event extraction across multiple levels of biological organization. Bioinformatics.

Patchigolla VSS Rahul, Sunil Kumar Sahu, and Ashish Anand. 2017. Biomedical event trigger identification using bidirectional recurrent neural network based models. arXiv preprint arXiv:1705.09516.

Utpal Kumar Sikdar, Bjorn Gamback, and M Krishna Kumar. 2020. IBS at WNUT 2020 Shared Task-1: Name Entity Extraction from Wet Lab Protocol. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Janvijay Singh and Anshul Wadhawan. 2020. PublishInCovid19 at WNUT 2020 Shared Task-1: Entity Recognition in Wet Lab Protocols using Structured Learning Ensemble and Contextualised Embeddings. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Mohammad Golam Sohrab, Khoa Duong, Makoto Miwa, and Hiroya Takamura. 2020. mgsohrab at WNUT 2020 Shared Task-1: Neural Exhaustive Approach for Entity and Relation Recognition Over Wet Lab Protocols. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Larisa N Soldatova, Daniel Nadis, Ross D King, Piyali S Basu, Emma Haddi, Véronique Baumlé, Nigel J Saunders, Wolfgang Marwan, and Brian B Rudkin. 2014. EXACT2: the semantics of biomedical protocols. BMC bioinformatics.

Pontus Stenetorp, Sampo Pyysalo, Goran Topić, Tomoko Ohta, Sophia Ananiadou, and Jun’ichi Tsujii. 2012. brat: a Web-based Tool for NLP-Assisted Text Annotation. In Proceedings of the Demonstrations at the 13th Conference of the European Chapter of the Association for Computational Linguistics (EACL).

Viktor Vasilev, Chenkai Liu, Traci Haddock, Swapnil Bhatia, Aaron Adler, Fusun Yaman, Jacob Beal, Jonathan Babb, Ron Weiss, Douglas Densmore, et al. 2011. A software stack for specification and robotic execution of protocols for synthetic biological engineering. In 3rd international workshop on bio-design automation.

Alain C Vaucher, Federico Zipoli, Joppe Geluykens, Vishnu H Nair, Philippe Schwaller, and Teodoro Laino. 2020. Automated extraction of chemical synthesis actions from experimental procedures.

Andreas Vlachos, Paula Buttery, Diarmuid O Séaghdha, and Ted Briscoe. 2009. Biomedical event extraction without training data. In Proceedings of the Workshop on Current Trends in Biomedical Natural Language Processing: Shared Task. Association for Computational Linguistics.

Jian Wang, Honglei Li, Yuan An, Hongfei Lin, and Zhihao Yang. 2016a. Biomedical event trigger detection based on convolutional neural network. International Journal of Data Mining and Bioinformatics.

Jian Wang, Jianhui Zhang, Yuan An, Hongfei Lin, Zhihao Yang, Yijia Zhang, and Yuanyuan Sun. 2016b. Biomedical event trigger detection by dependency-based word embedding. BMC medical genomics.

Zhilin Yang, Zihang Dai, Yiming Yang, Jaime Carbonell, Russ R Salakhutdinov, and Quoc V Le. 2019. Xlnet: Generalized autoregressive pretraining for language understanding. In Advances in neural information processing systems.

Qingcheng Zeng, Haoding Meng, Xiaoyang Fang, and Zhexin Liang. 2020. Fancy Man Launches Zippo at WNUT 2020 Shared Task-1: A Bert Case Model for Wet Lab Entity Extraction. In Proceedings of EMNLP 2020 Workshop on Noisy User-generated Text (WNUT).

Deyu Zhou, Dayou Zhong, and Yulan He. 2014. Event trigger identification for biomedical events extraction using domain knowledge. Bioinformatics.