Aims and scope

Genomics & Informatics is the official journal of the Korea Genome Organization (http://kogo.or.kr). Its abbreviated title is Genomics Inform. It was launched in 2003 by the Korea Genome Organization. It aims at making a substantial contribution to the understanding of any areas of genomics or informatics. Its scope includes novel data on the topics of gene discovery, comparative genome analyses, molecular and human evolution, informatics, genome structure and function, technological innovations and applications, statistical and mathematical methods, cutting-edge genetic and physical mapping, next generation sequencing and de novo assembly, and other topics that present data where sequence information is used to address biological concerns. Especially, Clinical genomics section is for a short report of all kinds of genome analysis data from clinical field such as cancer, diverse complex diseases and genetic diseases. It encourages submission of the cancer panel analysis data for a single cancer patient or a group of patients. It also encourages deposition of the genome data into designated database. Genome archives section is for a short manuscript announcing the genetic information of recently sequenced prokaryotic and eukaryotic genomes. These genome archives data can make the rationale for sequencing a specific organism.

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Contact information
Park, Taesung, Editor-in-Chief

Editorial office of Genomics & Informatics
Room No. 806, 193 Mallijae-ro, Jung-gu, Seoul 04501, Korea
Tel: +82-2-558-9394, Fax: +82-2-558-9434, email: kogo3@kogo.or.kr, URL address: https://genominfo.org

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Editor’s introduction to the special issue of the 6th Biomedical Linked Annotation Hackathon (BLAH6)

Jin-Dong Kim¹*, Kevin Bretonnel Cohen², Fabio Rinaldi³, Zhiyong Lu⁴, Nigel Collier⁵, Hyun-Seok Park⁶

¹Database Center for Life Science (DBCLS), Research Organization of Information and Systems (ROIS), Kashiwa, Chiba 277-0871, Japan
²School of Medicine, University of Colorado, Aurora, CO 80045, USA
³Dalle Molle Institute for Artificial Intelligence Research (IDSIA), 6928 Manno, Switzerland
⁴National Center for Biotechnology Information (NCBI), National Institutes of Health (NIH), Bethesda, MD 20894, USA
⁵Faculty of Modern & Medieval Languages, University of Cambridge, Cambridge CB3 9DP, UK
⁶Center for Convergence Research of Advanced Technologies, Ewha Womans University, Seoul 03760, Korea

As data science gains in importance and popularity, the need for accessing data in scientific literature is rapidly increasing. While structured databases are supposed to supply readily machine-readable data, unstructured contents, particularly scientific literature, are recognized as a biggest source of data with comprehensive details, e.g., experimental environments and actual observations.

Since the importance of scientific literature for data science has been widely recognized, several groups have invested to develop various text mining resources. While many of them are publicly available, interoperability of them remains a critical issue, hindering efficient use or reuse of them, particularly in mix with others.

The Biomedical Linked Annotation Hackathon (BLAH) series is annually organized to join forces of biomedical text mining for the goal to promote interoperability among text mining resources. The sixth edition of it was held in Tokyo, February 4–7, 2020, with 52 participants from 9 countries. The first day was held as a symposium to exchange and publicise the activities and ideas of the participants, and the following three days was held as a hackathon: the participants worked on implementing their ideas with collaboration with other participants.

While the main theme of the event was improving interoperability of biomedical literature mining, which include annotation datasets, tools, platforms, terminology resources, and so on, this year, "social media mining" was also explored as a special theme. Social media is recognized as a good source of raw signals on how people are thinking about what is going on in the world, which are largely missing in scientific literature. Therefore, social media mining is expected to complement literature mining.

This special issue is a collection of the reports on achievements from the hackathon, which address various issues of biomedical literature and social media mining, including document collection, automatic annotation, manual annotation, annotation platform, translation, terminology, ontology, and so on. Note that, except a few, many of the works began just before or even during the hackathon, and due to the limited time for work,
they are often small-sized works, which are expected to benefit from collaboration with other participants. Readers will find that many of the articles have co-authorship with, or acknowledgment of other participants, which is a typical nature of hackathon-oriented publications.

We hope that this will be an opportunity for the readers of the journal Genomics & Informatics to get aware of the state-of-the-art activities regarding interoperability of biomedical text mining, and at the same time to observe activities of hackathons like BLAH.

ORCID

Jin-Dong Kim: https://orcid.org/0000-0002-8877-3248  
Kevin Bretonnel Cohen: https://orcid.org/0000-0003-1749-8290  
Fabio Rinaldi: https://orcid.org/0000-0001-5718-5462  
Zhiyong Lu: https://orcid.org/0000-0002-8301-9553  
Nigel Collier: https://orcid.org/0000-0002-7230-4164  
Hyun-Seok Park: https://orcid.org/0000-0002-6617-2740

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Using the PubAnnotation ecosystem to perform agile text mining on *Genomics & Informatics*: a tutorial review

Hee-Jo Nam¹, Ryota Yamada², Hyun-Seok Park¹,³*

¹Bioinformatics Laboratory, ELTEC College of Engineering, Ewha Womans University, Seoul 03760, Korea
²Fuku Corporation, Tokyo 113-0033, Japan
³Center for Convergence Research of Advanced Technologies, Ewha Womans University, Seoul 03760, Korea

The prototype version of the full-text corpus of *Genomics & Informatics* has recently been archived in a GitHub repository. The full-text publications of volumes 10 through 17 are also directly downloadable from PubMed Central (PMC) as XML files. During the Biomedical Linked Annotation Hackathon 6 (BLAH6), we experimented with converting, annotating, and updating 301 PMC full-text articles of *Genomics & Informatics* using PubAnnotation, a system that provides a convenient way to add PMC publications based on PMCID. Thus, this review aims to provide a tutorial overview of practicing the iterative task of named entity recognition with the PubAnnotation/PubDictionaries/TextAE ecosystem. We also describe developing a conversion tool between the Genia tagger output and the JSON format of PubAnnotation during the hackathon.

**Keywords**: named entity recognition, natural language processing, text mining

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**Introduction**

*Genomics & Informatics* is the official journal of the Korea Genome Organization. The prototype version of the full-text corpus of *Genomics & Informatics* (GNI version 1.0) has recently been archived in a GitHub repository [1,2]. Further preprocessing and semi-automatic editing are underway to prepare the next version of GNI. As the volume numbers of *Genomics & Informatics* are growing, we needed a persistent and sharable repository to annotate and to upload the PMC articles of *Genomics & Informatics*.

During the Biomedical Linked Annotation Hackathon 6 (BLAH6), we experimented with annotating the PMC articles of *Genomics & Informatics*, making a custom dictionary using PubDictionaries, and uploading the annotation results into PubAnnotation. PubDictionaries is a public repository of dictionaries and PubAnnotation is a public repository of text annotations; these resources are primarily developed and maintained by the Database Center for Life Science (DBCLS), Japan [3,4]. PubAnnotation and PubDictionaries adopt a dictionary-based agile text mining approach, wherein iterative development cycles can be carried out by modifying a dictionary, manually reannotating, and automatically reannotating [5].

Thus, the purpose of this interdisciplinary tutorial review is to share our experiences of using the PubAnnotation ecosystem and writing a conversion script to apply to the *Genomics & Informatics* corpus [6,7]. We provide an introductory overview to briefly intro-
duce basic information extraction tasks and dictionary-based named entity recognition (NER) for non-experts in the field, and to provide some helpful pointers to start a deeper investigation into agile text mining and corpus annotation techniques in general.


Creating a PubAnnotation Pilot Project

PubAnnotation supports an agile approach to text mining by instantiating software components that allow for decomposed parallel development, while also facilitating continuous integration [5].

The PubAnnotation ecosystem is designed to be an open, API-driven system, and to harness changes for the user’s advantage. Annotations can be obtained from an external web service, which is called an annotation server. Through this principle, potential users are able to use the system to fine-tune and adjust their existing project [8].

During BLAH6, we created a PubAnnotation pilot project, called BLAH6-GNI-Corpus (http://pubannotation.org/projects/BLAH6-GNI-Corpus), initially to upload the Genomics & Informatics corpus. PubAnnotation provides a convenient way to add, annotate, and edit PMC publications based on PMCID. We specified the PMCID and uploaded the text files of Genomics & Informatics. In total, 301 documents were imported into the project.

Three components were used to implement the iterations of agile development, as shown in Fig. 1: PubAnnotation, a storage component for regression testing; TextAE, a manual annotation tool; and PubDictionaries, a dictionary-based annotator. These three components of the PubAnnotation ecosystem provide many ways to proceed with NER projects. The following sections provide one scenario, in which we conducted agile text mining with these components by adding the PMC publications of Genomics & Informatics to a PubAnnotation project, writing a script to upload the existing tagged documents, creating a PubDictionaries project to obtain annotations, and editing the annotations manually with TextAE [5].

A Tutorial Example

We initially used the GENIA tagger to annotate biological terms when developing the GNI corpus 1.0 [9,10]. It is easiest to under-

![Fig. 1. An Agile approach to text mining with PubAnnotation, PubDictionaries, and TextAE.](https://doi.org/10.5808/GI.2020.18.2.e19)
stand how PubAnnotation and PubDictionaries might be used to integrate the GNI corpus 1.0 on the basis of an example.

**An exemplary output format of the GENIA tagger**

The annotation result of PMCID 6440663—"We discovered inactivating mutations of tumor suppressor genes, including APC, TP53, and ARID1A, in three patients."—as shown in Fig. 2, is used as an example sentence.

The GENIA tagger outputs the base forms, part-of-speech (POS) tags, chunk tags, and named entity tags. The tagger is specifically tuned for biomedical texts such as MEDLINE abstracts. Fig. 2A is a direct output from the GENIA tagger, and 2B is a visualization of NER generated by TextAE [10], the default viewer and editor of PubAnnotation. Four different levels of tags are attached for each word in the example sentence: base forms, POS tags, chunk tags, and named-entity tags. For example, “TP53”, “TP53”, “NN”, “B-NP”, and “B-protein” indicate that the part of speech of the word “TP53” is a noun (“NN”), that the word begins a noun phrase (“B-NP”), and that it begins a phrase of a protein name (“B-protein”).

The last tag is a semantic-level tag to classify named entities in the text into pre-defined categories such as proteins, DNAs, RNAs, cell lines, and cell types. For named-entity tags, B/I/O notation was used, wherein the B/I/O terminology refers to the beginning of the phrase (B), internal to the phrase (I), and outside of the phrase (O).

Fig. 2 shows that APC was wrongly classified, because “APC” could refer to the adenomatous polyposis coli gene or to an antigen-presenting cell. Generally, biomedical NER faces difficulties for many reasons, prominent among which are the often-ambiguous abbreviations that are frequently used in the biomedical field.

**Wring a Python script to convert GENIA tagging results into a PubAnnotation format**

The desired result of a dictionary-based text annotation task would be an index of the dictionary entities corresponding to the referenced target texts. PubAnnotation’s text sequencer turns a document into a sequence of characters, so that positions in the document can be specified unambiguously by character offsets. For this reason, a conversion tool between the Genia tagger output and the JSON import/export format of PubAnnotation was written in Python during BLAH6. As shown in Fig. 3, we extracted the text field from a JSON file, and tokenized it by sentence, using the Natural Language Toolkit (NLTK) package, as in lines 67–73 [11]. This tokenizer divides a text into a list of sentences by using an unsupervised algorithm to build a model for abbreviation words, collocations, and named entities.
Fig. 3. A conversion tool written in Python.

```python
1 def _tagjson(sents, fp_json, tagger):
2     fp, _, pcid, divid, sec = fp_json, stem.split('-', maxsplit=1)
3     data = {
4         'text': None,
5         'sourcedb': 'PMC',
6         'sourceid': pcid,
7         'divid': divid,
8         'denotations': []
9     }
10
11 # tagging by GENIA Tagger
12 ans = []
13 begin, end = 0, 0
14 for sent in sents:
15     for word, word2, _, _, tag in tagger.parse(sent):
16         end = begin + len(word)
17         ann = {
18             'begin': begin,
19             'end': end,
20             'tag': tag,
21             'word': word,
22         }
23         begin = end + 1
24         ans.append(ann)
25         begin = end + 1
26         text = ' '.join([w for _, _, w in ann])
27         data['text'] = text
28
29 # combine B-I terms
30 reversed_anns = []
31 is_continue = False
32 _end = None
33 word_list = []
34 for begin, end, tag, word in reversed(anns):
35     if tag.startswith('O'):
36         continue
37         word_list.append(word)
38     if tag.startswith('I') and not is_continue:
39         is_continue = True
40         _end = end
41     elif tag.startswith('B'):
42         if _end is None:
43             _end = end
44         ner_tag = tag.split('-', maxsplit=1)[-1]
45         word = ' '.join(reversed(word_list))
46         reversed_anns.append(['begin', _end, ner_tag, word])
47         is_continue = False
48         _end = None
49         word_list = []
```

Fig. 4. PubAnnotation JSON format with its tag and span indexing information.

```json
{"target":"http://pubannotation.org/docs/sourcedb/PMC/sourceid/6440663/divs/0",
"sourcedb":"PMC","sourceid":"6440663",
"text":"We discovered inactivating mutations of tumor suppressor genes, including APC, TP53, and ARID1A, in three patients. ",
"divid":0,"project":"BLAH6-GNI-CORPUS",
"denotations": [{
  "id":"T1","span":{"begin":40,"end":62},"obj":"DNA"},
  {"id":"T2","span":{"begin":74,"end":77},"obj":"cell_type"},
  {"id":"T3","span":{"begin":79,"end":83},"obj":"protein"},
  {"id":"T4","span":{"begin":89,"end":95},"obj":"protein"}
]}
```
tions, and words that start sentences. In line 12–27, indexes for each word in the Genia output are calculated by adding up the white spaces and character lengths. In lines 29–60, a new list is created, containing the begin index, end index, named entity tag, and word; B-tags and I-tags are combined, and the indexes are recalculated. Finally, the list to the denotation field of the dictionary is appended and converted into JSON.

Manual editing using TextAE

A facility of visualization and manual editing is one of the primary aspects of making the PubAnnotation ecosystem adoptable by end-users. A user can easily add a new entry or delete an entry, in a try-and-revise manner.

In Fig. 4, there are four denotations for our example sentence, T1 through T4, with its tag and span information. The first one connects span 40–62 (the text spanning from the 40th to 62nd characters) to DNA, while the fourth connects span 89–95 to Protein. The default interpretation of T4 is as follows: the text span between “span”:“begin”:89, “end”:95” denotes an entity T1 “id”:“T1” of which the type is Protein.

Once an annotation file is prepared, TextAE can be used for manual editing of NER [12], as in Fig. 5. TextAE is a web-based graphical annotation editor, which was developed as an open-source project. APC is now tagged as a “gene,” as shown in Fig. 5, after manual editing. In addition to NER tagging, the example also presents the ease of using TextAE for manual editing of relation annotations, showing that the two entities, T1 and T2, that are introduced by the two denotations, are related to each other by the predicate “contains”, specified by the two different keys, so the relationship is directional.

Summary

In this tutorial review, we presented our experiences of conducting and agile text mining. During BLAH6, we created two separate PubAnnotation projects (BLAH6-GNI-Corpus and BLAH6-GNI-Corpus2), a dictionary (BLAH6-GNI-Dictionary), and an annotator (BLAH6-GNI-Annotator). A total of 12,908 labels were registered in the PubAnnotation ecosystem (https://pubannotation.org/annotators/BLAH6-GNI-ANNOTATOR).

While developing a conversion tool during BLAH6, indexing and calculating spans was a non-trivial task, as PubAnnotation utili-
lizes character-based indexing; enforcement of a fixed tokenization of the text is technically expensive.

Some minor suggestions relate to the user interface. In some menus, it was not fully obvious for a first-time user of the system what was clickable. We also had to create two separate projects, simply to utilize PubAnnotation’s text sequencer.

We assume that there are many categories of users with different levels of experience and familiarity with PubAnnotation, ranging from pure natural language processing specialists to biomedical research end users. We hope that some additional features will be added to the PubAnnotation ecosystem, to provide diversified access to different groups of users, who have different needs regarding workflow and information density.

**ORCID**

Hee-Jo Nam: https://orcid.org/0000-0001-6184-6737
Ryota Yamada: https://orcid.org/0000-0003-2237-5025
Hyun-Seok Park: https://orcid.org/0000-0002-6617-2740

**Authors’ Contribution**

Conceptualization: HSP. Data curation: HJN. Methodology: HJN, RY. Writing – original draft: HSP.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.

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**References**


WTO, an ontology for wheat traits and phenotypes in scientific publications

Claire Nédellec1*, Liliana Ibanescu2, Robert Bossy1, Pierre Sourdille3

1Paris-Saclay University, INRAE, MaIAGE, F-78350 Jouy-en-Josas, France
2Paris-Saclay University, INRAE, UMR MIA-Paris, AgroParisTech, F-75005, Paris, France
3University Clermont-Auvergne, INRAE, UMR 1095 GDEC, F-63000 Clermont-Ferrand, France

Phenotyping is a major issue for wheat agriculture to meet the challenges of adaptation of wheat varieties to climate change and chemical input reduction in crop. The need to improve the reuse of observations and experimental data has led to the creation of reference ontologies to standardize descriptions of phenotypes and to facilitate their comparison. The scientific literature is largely under-exploited, although extremely rich in phenotype descriptions associated with cultivars and genetic information. In this paper we propose the Wheat Trait Ontology (WTO) that is suitable for the extraction and management of scientific information from scientific papers, and its combination with data from genomic and experimental databases. We describe the principles of WTO construction and show examples of WTO use for the extraction and management of phenotype descriptions obtained from scientific documents.

Keywords: ontology, text mining, wheat trait and phenotype

Availability: WTO ontology (https://doi.org/10.15454/1.4382637738008071E12) is available on AgroPortal: http://agroportal.lirmm.fr/ontologies/WHEATPHENOTYPE under the license Creative Commons Attribution International 4.0 International (CC BY 4.0); The wheat trait bibliographic search engine SAMBlé AlvisIR is available at: http://bibliome.jouy.inra.fr/demo/wheat/alvisir/webapi/search.

Introduction

Improvement of most animal and plant species of agronomical interest has become an international stake because of the increasing demand for feeding a growing world population. The new environmental constraints such as the reduction of inputs (water, fertilizers, and pesticides) and the reduction of acreages involve the development of new breeding schemes that must be shorter and more powerful. This requires a significant improvement of the agronomical potential of the species through breeding. This is especially true for bread wheat (Triticum aestivum L.) which is the most widely grown crop worldwide.

The recent advent of genomic tools contributed to a better understanding of the biological mechanisms underlying the expression of phenotypes of agronomical interest. The availability of genetic information linked to genotyping and phenotyping experimental data obtained from fields and controlled environments has never been greater for understanding biological mechanisms and hypothesizing new models of plant biology [1].

As a consequence, reusing data from different platforms that are obtained through different methods, sensors and protocols, has become a major challenge. The standardization of the information for semantic interoperability of heterogeneous datasets is a key
milestone [2]. An ontologie, as defined in [3], is designed to represent the knowledge from one domain by concepts (or classes), relationships among these concepts and instances of these concepts. Therefore ontologies have long been identified as a critical tool for managing information systems in the fields of integrative plant biology, genetics and phenomics [4]: among others Gene Ontology [5,6] defines gene functions, biological processes and cellular components; the Plant Ontology (PO) Database [7,8] developed by the Planteome Project is a community resource for plant structure and developmental stages controlled vocabulary and annotations [9]. PO links plant anatomy, morphology and growth and development to plant genomics data.

Dedicated ontologies focus on controlled vocabulary for the description of the phenotypic information. The Plant Trait Ontology (TO) [10] of the Planteome project [11] defines general phenotypic traits in plants. Each trait is a distinguishable feature, characteristic, quality, or phenotypic feature of a developing or mature plant independently of the species. The Crop Ontology (CO) [12,13] is developed by several centers of the Consultative Group on International Agricultural Research (CGIAR) Biodiversity and their partners (Elixir, INRAE, iBET). This ontology focuses on the documentation of phenotype observations as variables that are grouped in nine high-level trait classes. The variables are triplets of observation methods, units of measurement, and traits that encompass the observed entity (e.g., grain, plant). CO distinguishes specific traits for 31 economically important plant species. Their vocabularies have reached different stages of development, ranging from pearl millet (52 variables) to wheat, the richest, with 498 variables.

Beside observation and experimental data, scientific literature is a significant source of genetic and phenotypic information on plants [14,15]. Automatic information retrieval and information extraction have been acknowledged as major challenges in Life Science for assisting manual biocuration, either to assess experimental or inferred data quality or to fill databases with complementary information [16,17]. However, most work focuses on molecular biology, functional and comparative genomics resource development, and phenotypic-related human health, as the BioCreative Track III interactive text mining task in 2012 [18].

In the plant biology domain, information extraction from text has attracted less attention [19], even though the quality and the abstraction of the textual information confer it a significant value for breeding. General properties of plant cultivars as described in the literature are of great interest for many research and innovation studies that are complementary to the detailed and partially unrelated phenotypic observations. Scientific literature summarizes, synthesize, abstracts and explains experimental results, filtering out spurious observations and highlighting important outcomes. As such it constitutes a valuable source of knowledge for the interpretation of phenotyping experimental results, as well as for the design of plant system biology models able to explain, predict, or simulate genotypic-phenotypic relationships.

Information extraction from text requires the establishment of dedicated ontologies and of text mining pipelines as largely recognized in the biomedical domain [20]. Ontologies improve text mining performances and conversely the information extracted is more reusable when linked to a reference resource such as an ontology through the normalization process. Normalization consists in assigning a class or a category from a controlled vocabulary to text mentions. It is a key step for the semantic interoperability of textual information and other sources of data and a major text mining challenge [21]. Plant traits and phenotypes expressed in textual sources are characterized by a great variability of the lexicon [15]. The text carries information at various levels of generality with different assessment status, ranging from experimental fine-grained data to general expert knowledge, through intermediate levels of synthesis and abstraction. The examples in Fig. 1 illustrate the variability of trait expressions in scientific documents in descending order of generality. In example (1) the trait “resistance to fungal and viral diseases” is a general trait. Example (2) mentions “FHB resistance” (i.e., Fusarium head blight resistance) which reflects resistance to a specific fungal disease FHB, and its effect on the related observations of six specific traits (e.g., plant height). Example (3) is the most specific: it is about the severity score of the trait “Russian wheat aphid resistance” observed for a given cultivar (i.e., Hatcher) whose value is 1.9.

This varying scope of phenotypic information in scientific papers answers to different needs and usages. It ranges from detailed documentation of experiments and inferred data, to review of shared and well-acknowledged bodies of knowledge supported by large sets of experimental and scientific results.

We have been developing the Wheat Trait Ontology (WTO) since 2010 to answer breeders and scientists’ needs for wheat trait and phenotype information management and retrieval at varying abstraction scales. WTO supports two objectives: (1) building a formal shared representation of wheat trait whose knowledge organization closely reflects the expert knowledge model and (2) making phenotypic information extraction from text easier. To achieve both objectives, the sources for building WTO include expert knowledge and textual documents: expert interviews, terminology analysis from the literature and gene catalogs. The richness of the WTO vocabulary, its similarity with scientific literature lexicon and its deep hierarchies make it a useful resource for both text mining and information management.
The paper is organized as follows. Section 2 describes the WTO. Section 3 presents the motivation and method for building WTO. Section 4 illustrates WTO usage through an application. Section 5 discusses WTO characteristics compared to other semantic resources and presents future work.

WTO Description

The WTO covers a wide range of bread wheat traits (e.g., observable physical plant properties), phenotypes (e.g., trait values) and their related environmental conditions (e.g., disease, extreme temperature) organized in three trees. The current version contains 596 classes. The population of the main classes and their subclasses is given in Table 1.

The maximum depth of WTO is 9 and the average number of children per class is 3. We chose a deep and balanced structure because the breeder’s needs to manage data at different levels of aggregation. Classes at intermediate levels support synthetic queries for searching high-level correlations between genetic, phenotypic, and physiological phenomena.

The classes of the ‘Trait’ subtree are linked to the corresponding phenotypes by the “Trait_has_value” relationship. For instance, ‘ear emergence time’ trait class is linked to the ‘late heading’ phenotype class.

The ‘Environmental condition’ subtree mainly represents abiotic conditions and biotic conditions that are linked to the corresponding responses of the plant to abiotic stresses and biotic stresses. The main root classes of ‘Response to environmental conditions’ range from response to chemical, radiation, temperature, to a large range of responses to biotic stresses as shown in Fig. 2.

Response to biotic stress is indeed a major concern for wheat breeding. Wheat is affected by several microbial, bacterial, viral and mainly fungal diseases that cause major crop loss [22]. WTO accounts for this situation with two large subtrees ‘Disease’ (58 classes) and ‘Pest’ (103 classes) of ‘Environmental condition’ (Table 1). The relation ‘Causes’ between the ‘Pathogen’ classes and the infectious “Disease” classes represents the causative link between the agent and the disease. WTO distinguishes between disease of bacterial, viral and fungal causal agents. A total of 55 different fungal species causing 44 diseases is described in WTO.

In a similar way, the ‘Response to biotic stress’ subtree finely distinguishes between the causal stress factors as shown in Figs. 3 and 4.

The responses of the plant to biotic stresses are expressed in two ways: either by the disease name or by the causative agent names. A given disease name may have synonyms and a disease may be caused by more than an agent (Fig. 5).

Moreover, fungi naming in scientific paper do not always strictly follows the nomenclature standard imposed by the mycologist community. For instance, names corresponding to different life stages can be found. For each resistance trait, the causative agents are given with their standard names and the other names as in Fig. 6.

WTO lexical synonymy relations and conceptual relations are complementary with respect to the intended uses to reflect expert knowledge model and make phenotypic information extraction from text easier.

As summarized by Fig. 7 WTO structure is mainly hierarchical with two transversal relations: a domain-specific causal one and a variable-value relation.

Table 1. Main classes of WTO with the number of some subclasses

<table>
<thead>
<tr>
<th>No.</th>
<th>Environmental condition</th>
<th>Abiotic condition (e.g., chemical, nutrient, water, wind)</th>
<th>Biotic condition</th>
<th>Biotic stress</th>
<th>Disease</th>
<th>Bacterial disease</th>
<th>Fungal disease</th>
<th>Viral disease</th>
<th>Diseased caused by nematode</th>
<th>Pest</th>
<th>Insects</th>
<th>Plant property</th>
<th>Phenotype</th>
<th>Trait</th>
<th>Development (plant habit, precocity, vernalization)</th>
<th>Growth (crop yield, nutrient use efficiency, density)</th>
<th>Morphology (of awn, glume, grain, spike)</th>
<th>Quality</th>
<th>Food property</th>
<th>Grain composition</th>
<th>Grain quality</th>
<th>Milling quality</th>
<th>Reproduction</th>
<th>Response to environmental conditions</th>
<th>Response to abiotic stress</th>
<th>Response to biotic stress</th>
</tr>
</thead>
<tbody>
<tr>
<td>221</td>
<td>Abiotic condition</td>
<td>51</td>
<td>171</td>
<td>170</td>
<td>58</td>
<td>6</td>
<td>44</td>
<td>6</td>
<td>2</td>
<td>103</td>
<td>21</td>
<td>374</td>
<td>45</td>
<td>326</td>
<td>19</td>
<td>41</td>
<td>23</td>
<td>58</td>
<td>30</td>
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<td>13</td>
<td>4</td>
<td>5,173</td>
<td>64</td>
<td>104</td>
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</tr>
<tr>
<td></td>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>WTO, Wheat Trait Ontology.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 1. Examples of phenotype mentions from scientific papers. Traits and phenotypes are in bold.
Wheat Trait Ontology Building

WTO was built using the NeOn Methodology [23], a scenario-based methodology that supports the collaborative aspects of ontology development and reuse. The WTO development process followed successively Scenario 1, From specification to implementation, and Scenario 2, Reusing non-ontological resources of the NeOn methodology. The first step is to specify the ontology requirements, provided in the next subsection. Then we present insights and rationales for design choices in the following subsection.

Ontology Requirements Specification

The needs for the development of shorter and more powerful breeding schemes is a strong motivation for sharing phenotypic information linked to genes of interest and traits. Building an open and shared database for marker-based assisted selection (MAS) in bread wheat was the SAMblé project objective (2010–2014) [24]. The SAMBlé database should support both short-term MAS-related goals of breeders, the intended users, and long-term research goals of researchers on underlying biological mechanisms of phenotypes. The information considered for the database was the existence of links between one or more markers and genes of agronomic interest in bread wheat. The information sources were the scientific literature, gene catalogs and in-field and high-throughput phenotyping experiments.

In scientific papers, phenotypic information is frequently linked to varieties, genes or markers and traits as in Fig. 8, which makes it extremely relevant for breeding [15].

This information was first automatically extracted from the literature, then assessed against reference material and elite material (335 varieties) under field conditions for different traits of interest. Finally, the markers that gave the best results and could be used in breeding selection schemes were recorded in the database to be queried by the partner breeders [24]. The traits considered in the SAMBlé project were related to four main large topics, namely, disease resistance, resistance to abiotic stress, plant development, and baking quality.

Representative queries of the breeders were, “which alleles and markers are involved in resistance to rust (e.g., leaf rust, stripe rust, stem rust)” and “what are the varieties tested.” Same question arises for “bread making quality (e.g., flour quality, color, composition, mechanical property, crumb firmness)?”.

A general objective of SAMBlé was to develop a shared database with the information collected by the project that would be easily searchable. The WTO was designed to support this goal. The ontology should support queries on traits and phenotypes at various levels of aggregation combined with other criteria on markers, genes, and varieties.

To this purpose we created WTO as deep non-strict hierarchies with the information collected by the project that would be easily searchable. The WTO was designed to support this goal. The ontology should support queries on traits and phenotypes at various levels of aggregation combined with other criteria on markers, genes, and varieties.

Fig. 2. Wheat Trait Ontology (WTO) subclasses for ‘response to environmental condition’.

Design and implementation

The design of WTO followed a top-down approach where the core model was first established based on project partner expertise on wheat phenotyping: the SAMblé project gathered breeders from French breeding companies, the French union of breeders (UFS) and Arvalis, it was led by the research unit GDEC-INRAE (Genetics, Diversity and Ecophysiology of Cereals). Text mining and plant information management were provided by Mathematics, Informatics and Genomics Laboratory, French National Research Institute for Agriculture, Food and the Environment (MIG-INRA) and Unité de Recherche Génomique Info, French National Research Institute for Agriculture, Food and the Environment (URGI-INRAE). The main classes of the WTO core model were similar as presented in Table 1 of Section 2. The core model was then extended by reusing information from three external sources: scientific literature, the Catalog of Gene Symbols for Wheat [25] and GrainGene database [26]. The biotic stress response, diseases, and pathogen WTO subtrees (see Section 2) were then significantly restructured by wheat disease experts. We adopted the Obo-Edit tool as ontology editor, to make it easier for biologists and breeders to revise and enrich WTO, compared to more powerful but less user friendly tools.
**Fig. 3.** An excerpt of the different ‘resistance to a fungal pathogen’ in Wheat Trait Ontology (WTO).

**Fig. 4.** Example of the rust disease family in Wheat Trait Ontology (WTO).

**Fig. 5.** Synonyms of the ‘Resistance to Leaf Rust’ label in Wheat Trait Ontology (WTO). Leaf Rust disease is caused by different fungi, namely ‘Puccinia recondita’ and ‘Puccinia triticina’.

**Fig. 6.** Example of various names of *Parastagonospora nodorum* fungus in Wheat Trait Ontology (WTO).
Scientific literature as a source of concepts
An Ontology Acquisition approach was first used to extract and conceptualize WTO concepts and relationships from scientific text expressed in natural language, following the same methodology as described in Nedellec et al. study [27].

We applied the term extractor BioYateA [28] to a scientific corpus to automatically extract relevant domain-specific terms. BioYateA's strength over other term extractors is the ability to extract prepositional phrases that are frequent in wheat trait terms, e.g., response to vernalization, florets without grain [29]. The scientific corpus was composed of the abstracts and titles of articles. They were obtained from the Web of Science (WoS) bibliographic search engine with the keywords 'wheat or Triticum aestivum and marker and gene'. It yielded 3,170 references (see Nedellec et al. study [15] for more details).

The candidate terms extracted by BioYateA were then used to derive concepts using the Terminology Design Interface (TyDI) tool. TyDI supports term collaborative assessment and structuring [27]. First, relevant terms were selected among candidate terms by manual screening. Validated terms were grouped in semantic classes of preferred terms, synonyms and typographic and acronym variations. They were structured in hypernym hierarchies consistent with the core model. Concepts and concept hierarchies were then derived from these semantic classes and hypernym trees to populate the core model. The preferred terms were kept as concept labels. This literature term analysis approach sped-up the discovery process of a very large set of trait, phenotype, disease, and pathogen related concepts and subsumption relationships.

Other external sources of wheat trait terms
To identify complementary relevant trait terms, we also used the Catalog of Gene Symbols for Wheat (WGC) [30] available online at the Wheat Genetics Resources Database of Japan as a PDF file at the date of WTO building in 2011. The main contribution to WTO from the catalog was related to plant morphology (e.g., plant height) and physiology (e.g., response to photoperiod).

The GrainGenes [26] database was also used for the study of biotic stress response. GrainGenes is a comprehensive resource for molecular and phenotypic information for wheat maintained by U.S. Department of Agriculture and mirrored by MaLAGE. GrainGenes web pages listed general traits and specific traits for wheat, barley, and oat species from which we identified some wheat disease names and their pathogen agents. INRAE experts of wheat diseases then controlled the naming because for some diseases American vernacular naming was not consistent with European naming.

WTO evolution
Fig. 9 displays the evolution of the WTO (formerly named Wheat Phenotype Ontology) between 2010 and 2020. The first public
version of WTO was released in August 2011. It contained 460 classes and 260 synonyms of labels. The 2011 version of WTO was revised in 2014. Confusions between pathogen names and synonyms were corrected which resulted in an increased number of synonyms and decreased number of classes. In 2015, the classes ‘Fiber quality’, ‘Food property’, ‘Milling quality’, ‘Grain composition’ and ‘Grain quality’ were grouped in a new ‘Quality’ class in order to reduce the number of root classes and to increase WTO readability. Conversely the ‘Development’ class that mixed phenological phenotypes, morphology (e.g., color, length) and growth (related to yield) was split into three distinct classes: ‘Development’, ‘Morphology’ and ‘Growth’.

In 2017, with the purpose of using WTO for managing other phenotypic databases than the SAMBlé one, we evaluated WTO scope with respect to two external resources. The WIPO (the Wheat INRA Phenotype Ontology formerly named the INRA Wheat Ontology) [31] developed by URGI-INRAE and the “list of wheat descriptors for Characterization and Evaluation” of the NARO GeneBank project. A few more morphology terms such as ‘presence of awn’, ‘glume pubescence’, ‘glume color’ were then added to WTO.

**WTO for marker-assisted selection**

To be used as the conceptual formalization of the SAMBlé database schema, WTO was integrated into the MAS (Marker Assisted Selection) knowledge model detailed in Nedellec et al.’s study [15]. The MAS model was designed to manage the entities and relations of the SAMBlé database. It contains 8 entity types and 14 n-ary relationships for the representation of the genotypic and phenotypic information and relationships collected from the literature and experiments of the SAMBlé project. The main MAS model entities are ‘Marker’, ‘Type’, ‘Allele’ and ‘Gene’, ‘Trait’ and ‘Phenotype’ and ‘Variety’. ‘Type’ represents the type of method used to identify the marker, e.g., amplified fragment length polymorphism, microsatellite. The main relationships are ‘Marker tags Gene in Variety’ between markers, genes and varieties, ‘Trait has Phenotype in Variety’ between traits, phenotypes and varieties and ‘Gene expresses Phenotype in Variety’ between genes, phenotypes, and varieties.

The connection of the MAS model to WTO is achieved through the straight forward alignment of two pairs of MAS and WTO classes: (1) the MAS ‘Trait’ class is aligned with WTO ‘Plant property’ class, the root of the trait subtree (2) the MAS ‘Phenotype’ class is aligned with WTO ‘Phenotype’ class, the root of WTO phenotype subtree. The other MAS classes (e.g., Gene, Marker, Variety) are also connected to nomenclatures and catalogs (e.g., Genes nomenclatures, Markers lists, and Variety catalogs) for data standardization. The integrated MAS and WTO model was successfully used for the management of SAMBlé database information and for information extraction from text.

### Wheat Trait Ontology Usage

WTO has been validated through the use by breeders and researchers involved in the SAMBlé project of two end-user applications, the SAMBlé database interface [24] and the Wheat literature semantic search engine AlvisIR. AlvisIR supports queries on genes, varieties, markers, phenotypes and traits extracted from PubMed references. Phenotype and trait expressions in text are normalized by WTO concepts.

Fig. 10 gives an example of a semantic search for phenotypic information. The example query asks for documents where the gene ‘Lr34’ is mentioned in relation to the trait ‘resistance to rust’ in ‘wheat’ by combining the three keywords, ‘Lr34’, ‘resistance to rust’, and ‘wheat’. The first hit displays a document extract where ‘adult plant stripe rust resistance’ (underlined in green) is tagged by the query ‘resistance to rust’ keyword.

‘Resistance to rust’ in the user query has been interpreted by three complementary mechanisms. A text mining workflow run in batch mode has first automatically extracted all terms from the documents, among which the term ‘adult plant stripe rust resistance’, and automatically mapped it to the relevant WTO class ‘Resistance to stripe rust’. The query interpreter executed on the fly has segmented the user query and mapped the query term ‘resistance to rust’ to the corresponding WTO class. The subsumption relation between the query class and the document class has then been verified. The document term is therefore validated as an instance of the query term and the document is displayed as a hit.

Fig. 11 shows the corresponding subpart of WTO with the two mapped classes. A navigation tool (Fig. 11) supports the expression of the query by the user by the combination of selected classes. The users from the SAMBlé project are satisfied with the balanced and deep tree structure of WTO that makes ontology browsing and class selection much easier than a flat and large list of classes would.

High-level queries as exemplified here are powerful for combining criteria on phenotypes with other genetic or environmental information as requested by the SAMBlé project.

The online version of AlvisIR indexes PubMed abstracts. PubMed has been preferred over WoS for its Open Access license to references. Current work includes the extension of the corpus to full papers of main scientific journals. Eighteen thousand papers have been identified among which half are available through Open Access and 1,361 journals targeted. The text mining workflow
named WheatLiterature used to fill in the database from the scientific literature is based on the AlvisNLP technology (AlvisNLP on Github). It is distributed as a component of the European text mining OpenMinTeD platform [32].

**Discussion**

Beyond semantic search, ontology-based fine-grained information extraction is a key component of the integration of textual information with experimental and genetic data. However, the reference knowledge models often differ with the sources and the nature of the information. Their alignment and user query rewriting are a major challenge for data integration [33].

Significant work has been done on Wheat Data Interoperability Guidelines [34] that focuses on Minimum Information About a Plant Phenotyping Experiment (MIAPPE) [2,35]. For experimental data, observation variables including traits but also observation protocol, unit of measure and development stage are critical for properly documenting the observations and determining if observations are comparable or not. This leads to building trait ontologies as WIPO [31] or Crop Ontology [13] where the trait leaves are database variable traits (e.g., ‘Susceptibility to leaf rust’ in WIPO, ‘Leaf rust severity’ in Crop Ontology). Phenotypes, the values of the traits, (e.g., ‘Susceptible to leaf rust’) are not conceptualized as classes but represented by the database numerical data as values of the trait variable. For instance, in WIPO the trait Disease

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**Fig. 9.** Statistics of the Wheat Trait Ontology (WTO) between 2010 and 2020.

**Fig. 10.** Screenshot of AlvisIR semantic search engine query web page.
intensity score takes values on a 1 to 9 increasing scale (1, no disease; 9, very severe). Such ontologies are suitable for accurately documenting observations and for the computation of correlations by statistical tools.

Conversely, WTO aimed at managing both traits and phenotype values represented by expressions, as they occur in the scientific literature. For instance, in the *the leaf rust susceptible cultivar 'GA 100'* phrase, the phenotype value is *'leaf rust susceptible'* and the variety is *'GA 100'*. In this way, WTO representation then supports SAMblé data discovery by direct queries on traits and phenotype values (e.g., *'Leaf rust susceptibility'*) at various levels of generality (e.g., *'Rust susceptibility', 'Fungal disease susceptibility'*) and their relation to other information (e.g., cultivars).

Similar queries on observation databases that follow MIAPPE recommendations would require the translation of numerical values by using value domains or thresholds, i.e., discretization and hierarchization of the phenotypes. Moreover, the lack of depth of ontologies such as WIPO or Crop Ontology with a comb-like structure does not allow high-level queries. An example in WIPO, is the trait *'Susceptibility to leaf rust'*, which is a direct subclass of the high level *'biotic stress trait'* without intermediate levels. Similarly, in the Crop Ontology the trait *'Fusarium head blight AUDPC'* is a direct subclass of *'biotic stress trait'*. Another representative example is *'Nitrogen harvest index'*. In WIPO and Crop Ontology, it has only one direct ancestor, which is *'Quality trait'*, with 51 other sibling traits in Crop Ontology. In WTO, *'Nitrogen harvest index'* has five successive ancestors: *'Nitrogen use efficiency', 'Macronutrient use efficiency', 'Nutrient use efficiency', 'Growth'*, by increasing order of generality.

The integration of the two sources of data, observations, and synthetic information from text in a same data management system should preserve the best of the two approaches. It would require the alignment of the ontology classes and the rewriting of the phenotype variable values to map them to qualitative descriptors.

In the SAMblé project and for development of the OpenMintTeD Wheat use case, we experienced this situation with the two ontologies: WIPO, which indexes experimental phenotype data, and WTO, which indexes PubMed phenotypic information. Their classes are not mappable in a straight-forward one-to-one way. It is noteworthy that the types of alignments and rewriting identified during these projects are not specific to wheat or even to plants, but are general to any phenotype observation data. Further investigation of this question is a future challenge for the integration of phenotype data from different sources allowing a better exploitation of textual data.

**Conclusion**

We proposed WTO, a reusable ontology of bread wheat traits and phenotypes and related environmental factors. The design of the model relies both on domain expert knowledge and the analysis of evidence published in the scientific literature. The WTO model is
deeply structured, well reflecting the domain knowledge. It facilitates navigation and reuse for data and knowledge discovery. The model was designed to support the extraction and the management of marker-assisted selection information. WTO is also a contribution to the description of the link between genetic and phenotypic information. Concept synonyms were directly extracted from the literature, which turns WTO a suitable resource for Information Extraction and Information Retrieval. WTO has been assessed for its consistency through its use. WTO is complementary to other ontologies dedicated to the documentation of phenotypic observations. We believe that future work on their alignment and mapping will favor data semantic interoperability from the literature and experimental sources.

ORCID

Claire Nédellec: https://orcid.org/0000-0002-0577-0595
Liliana Ibanescu: https://orcid.org/0000-0003-3373-437X
Robert Bossy: https://orcid.org/0000-0001-6652-9319
Pierre Sourdille: https://orcid.org/0000-0002-1027-2224

Authors’ Contribution

Conceptualization: CN, RB, PS. Funding acquisition: CN, PS. Methodology: CN, RB, LI. Writing – original draft: CN. Writing – review & editing: LI.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

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Extending TextAE for annotation of non-contiguous entities

Jake Lever1*, Russ Altman1, Jin-Dong Kim2

1Department of Bioengineering, Stanford University, Stanford, CA 94305, USA
2Database Center for Life Science, Research Organization of Information and Systems, Kashiwa 277-0871, Japan

Named entity recognition tools are used to identify mentions of biomedical entities in free text and are essential components of high-quality information retrieval and extraction systems. Without good entity recognition, methods will mislabel searched text and will miss important information or identify spurious text that will frustrate users. Most tools do not capture non-contiguous entities which are separate spans of text that together refer to an entity, e.g., the entity "type 1 diabetes" in the phrase "type 1 and type 2 diabetes." This type is commonly found in biomedical texts, especially in lists, where multiple biomedical entities are named in shortened form to avoid repeating words. Most text annotation systems, that enable users to view and edit entity annotations, do not support non-contiguous entities. Therefore, experts cannot even visualize non-contiguous entities, let alone annotate them to build valuable datasets for machine learning methods. To combat this problem and as part of the BLAH6 hackathon, we extended the TextAE platform to allow visualization and annotation of non-contiguous entities. This enables users to add new subspans to existing entities by selecting additional text. We integrate this new functionality with TextAE’s existing editing functionality to allow easy changes to entity annotation and editing of relation annotations involving non-contiguous entities. Finally, we roughly quantify the problem across the entire accessible biomedical literature to highlight that there are a substantial number of non-contiguous entities that appear in lists that would be missed by most text mining systems.

Keywords: editor, text annotation, text mining, visualization

Introduction

Information extraction and retrieval methods are essential tools to enable scientists to find and read the appropriate papers to enable discoveries. Many of these methods require identifying mentions of specific biomedical entities in the text and make use of named entity recognition (NER) tools for this task. Most entities are represented by a single span of text, e.g., the name of a drug. However, some entities are represented by multiple spans of text that are separated by other words and together identify the entity, for example, the separate words "skin" and "cancer" in "skin and lung cancer." These are known as non-contiguous, or discontiguous entities. Table 1 illustrates several more examples from public text mining resources. It should be noted that non-contiguous entities are different from anaphora or coreference resolution, in which multiple spans refer to the same entity separately and do not work together to identify the entity.

Robust annotation tools that are capable of annotating non-contiguous entities are important so that valuable entity information is not missed. These tools are needed to create
corpora with non-contiguous entities that can be used as training
data for machine learning-based NER methods and also evaluate
all NER methods fairly. The leading NER methods frequently use
machine-learning methods such as conditional random fields
(CRF) that are incapable of capturing non-contiguous entities
without additional postprocessing. Popular tools such as BAN-
NER [1], tmChem [2], and DNORM [3] do not support non-con-
tiguous entities.

Many annotation tools have been developed for manual tagging
of entities within a document for the biological domain and other
domains. A detailed recent review of the strengths and weaknesses
of different methods can be found in Neves and Seva’s study [4].
To gauge the support for non-contiguous entities, we manually
tested the 15 tools selected in that review with an overview shown
in Table 2. We were able to run all but one, PDFAnno which dis-
played an error message that others have reported on Github. We
found that only 2 support non-contiguous entities, BRAT [5], and
Catma. Furthermore the AlvisAE [6] tool that was not included in
the review also supports non-contiguous entity annotation. We
suggest that more tools need to provide support for non-contigu-
ous entities.

To that goal, we describe the addition of non-contiguous entity
support to TextAE. TextAE is an annotation platform that forms
part of the PubAnnotation system for storing and editing text an-
nnotations [7]. The PubAnnotation format currently has support for non-contiguous entities but are
converted to an alternative representation when edited using the
current release of TextAE, known as the chaining representation.
This representation converts an entity that contains multiple spans
to multiple entities and links them with a relation with type “_lexi-
ticallyChainedTo.” This representation is time-consuming to edit
and visualizes poorly. Fig. 1 illustrates the current representation of
domain with non-contiguous entities that can be used as training
data for machine learning-based NER methods and also evaluate
all NER methods fairly. The leading NER methods frequently use
machine-learning methods such as conditional random fields
(CRF) that are incapable of capturing non-contiguous entities
without additional postprocessing. Popular tools such as BAN-
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Catma. Furthermore the AlvisAE [6] tool that was not included in
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current release of TextAE, known as the chaining representation.
This representation converts an entity that contains multiple spans
to multiple entities and links them with a relation with type “_lexi-
ticallyChainedTo.” This representation is time-consuming to edit
and visualizes poorly. Fig. 1 illustrates the current representation of
tables from different public text mining datasets

<table>
<thead>
<tr>
<th>Source</th>
<th>PubMed ID</th>
<th>Snippet</th>
<th>Non-contiguous entity</th>
</tr>
</thead>
<tbody>
<tr>
<td>BioNLP 2019 Bacteria Biotope Task</td>
<td>23224222</td>
<td>Both French and German cheeses have previously been reported to contain M. psychrotolerans</td>
<td>French cheeses</td>
</tr>
<tr>
<td>CancerMine</td>
<td>19622846</td>
<td>...and used API tests to identify S. aureus and E-tests to determine methicil-</td>
<td>Methicillin resistance</td>
</tr>
<tr>
<td>CancerMine</td>
<td>19855840</td>
<td>It is suggested that DCL1 is a candidate tumour suppressor gene for human liver cancer, as well as for prostate, lung, colorectal and breast cancers</td>
<td>Prostate cancers</td>
</tr>
<tr>
<td>CancerMine</td>
<td>19734946</td>
<td>LARG at chromosome 11q23 has functional characteristics of a tumor sup-</td>
<td>Breast cancer</td>
</tr>
<tr>
<td>PGxMine</td>
<td>23385314</td>
<td>In vitro analysis and quantitative prediction of efavirenz inhibition of eight cytochrome P450 (CYP) enzymes: major effects on CYPs 2B6, 2C8, 2C9 and 2C19</td>
<td>CYP 2C19</td>
</tr>
</tbody>
</table>

The examples from CancerMine [8] and PGxMine [9] are not currently captured by the corresponding method and are false negatives.

Table 2. An analysis of the annotation tools reviewed in Neves and Seva’s study [4] for their capabilities to annotate non-contiguous entities

<table>
<thead>
<tr>
<th>Tool</th>
<th>URL</th>
<th>Can run?</th>
<th>Supports entity annotation?</th>
<th>Support non-contiguous entities?</th>
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<td>Y</td>
<td>N</td>
</tr>
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<td>brat</td>
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<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>Catma</td>
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<td>Y</td>
</tr>
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<td>Y</td>
<td>Y</td>
<td>N</td>
</tr>
<tr>
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<td>N</td>
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<tr>
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<td>Y</td>
<td>N</td>
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<td>Y</td>
<td>N</td>
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<td>PDFAnno</td>
<td><a href="https://github.com/paperaipdfanno">https://github.com/paperaipdfanno</a></td>
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<td>-</td>
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<td><a href="https://prodi.gy/">https://prodi.gy/</a></td>
<td>Y</td>
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<td>tagtog</td>
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<td>Y</td>
<td>Y</td>
<td>N</td>
</tr>
</tbody>
</table>

https://doi.org/10.5808/GI.2020.18.2.e19
method. With the current TextAE interface, it is time-consuming to annotate each entity. Assuming TextAE has been set up with appropriate entity and relation types, for each non-contiguous entity, it requires creating two entities (2 mouse clicks), designating one entity with the type “_FRAGMENT” (2 clicks), switching to the relation mode (1 click), creating a relation between the two entities (2 clicks), changing its type to “_lexicallyChainedTo” (2 clicks) and switching back to Term Mode to continue entity annotation (1 click). Even for a TextAE power user, ten clicks for each non-contiguous entity is time-consuming for a large-scale annotation and produces an unwieldy result which is not visually clear.

In this paper, we describe our solution of an extension to the existing TextAE annotation platform to provide seamless support for annotating non-contiguous entities. Finally, we provide evidence of the widespread nature of non-contiguous entities in the biomedical literature using a rule-based extraction system to roughly quantify the scale of non-contiguous entities across all PubMed abstracts and accessible PubMed Central full-text papers.

**Methods**

To develop improved methods to capture non-contiguous entities, well-annotated data needs to be prepared and examined that contain non-contiguous entities. We extend the TextAE annotation platform that is part of the PubAnnotation system [7]. This enables annotation of entities with multiple spans as shown in Fig. 2 with a new subspan mode. The user can select new spans of text that will be added to an existing entity and displayed clearly.

The first task for implementation was changing the underlying span model in TextAE so that all spans are represented as a list of subspans. We dynamically check the input annotation data (in PubAnnotation format) to check if an entity has a single span, or a list of spans, and convert all entities to contain lists of spans, even for single spans. Previously, spans were rendered using a single...
HTML span tag around the section with appropriate CSS styling to identify the span as an entity. To visualize the new subspans, we removed the styling from the span class, and create subspans for each part of the span and transferred the stylings to the subspans. TextAE implements an Undo/Redo system so changes were required across the codebase to enable the existing functionality to work with the new underlying data structure and allow entities to be manipulated as before.

A new toggleable button (Add subspan) was added to the toolbar. When this button is toggled, any new spans that are selected by the user are added to the previously selected entity. This requires checking that new subspans were compatible with the structure that is enforced by the HTML page. This means that spans and subspans cannot intersect unless one is contained within the other entirely. This means that in the snippet: “breast cancer gene”, it would not be possible for “breast cancer” and “cancer gene” to be tagged as entities. However “breast cancer” and “cancer” could be tagged as “cancer” is fully contained within “breast cancer.” We have not come across use-cases where this functionality is currently needed but cannot discount the potential of this limitation. Fig. 2 illustrates the user interface with an example of non-contiguous entities.

TextAE has several user interface shortcuts to enable fast annotation and correction of entities. Users can extend an entity annotation by highlighting text that begins within an entity annotation and goes beyond the entity. Inversely, users can also shorten entity annotations by highlighting text that begins outside an entity span and finishes within an entity, thereby removing the selected text from the entity annotation. We extended this functionality to work for the new subspan system so that it would extend the appropriate first or last subspan in an entity outwards, or would shorten or even remove subspans that are highlighted. We further added user interface tweaks so that when a user selected a subspan, it would select all the subspans for the entity. Finally, we implemented export functionality so that the new subspans would be correctly stored in the PubAnnotation format with a list of spans for those entities with multiple subspans.

The code for this paper is available at https://github.com/jakelever/textae.

Results

We first tested to check that all existing functionality of TextAE remained operational. We confirmed that the new subspan model was able to load data containing non-contiguous entities and annotations with non-contiguous entities could be saved correctly to the PubAnnotation format. Furthermore, we tested that all existing functionality, including relation annotation, worked correctly with non-contiguous entity annotations.

We quantified the user interactions required to annotate non-contiguous entities. With this new interface, the user needs to annotate a single span (1 click), enable the Add subspan mode (1 click), add a new subspan (1 click), and disable the Add subspan mode (1 click). With only four clicks, we have drastically reduced the user effort, compared to the 10 clicks required previously, and no longer require the user to switch annotation modes within TextAE. Furthermore, the output is visually clearer. This performance is similar to the Catma tool, which requires four clicks to annotate a non-contiguous entity (1 to activate the discontinuous mode, 2 to select the two spans and 1 to select the entity type). And it is marginally easier than BRAT which takes five clicks (2 to annotate the first entity, 1 to edit the entity, 1 to select Add Frag and 1 to select the new span).

Discussion

While non-contiguous entities initially seem like a limited problem for text annotation, we note that two other BLAH 6 hackathon projects requested this functionality during the event: a project working on annotations from the recent BioNLP Shared Task [10] and a project focused on Medical Device Indication annotation. To understand the scale of this problem, we quantified the number of non-contiguous entities that appear in lists, as shown in the CancerMine examples in Table 1. We focussed on this format as these can be extracted using a modified dictionary matching method.

We used the PubTator Central resource [11] as it provides text-level entity annotations of a very large set of biomedical publications and also a rough set of synonyms for different entity types. The annotations provide locations of biomedical entities that may be the final element in a list. For example, the phrase “prostate, skin and lung cancer” would only likely be tagged for “lung cancer” in PubTator. We aimed to retrieve other entities from these lists using the set of synonyms from PubTator Central, so that “prostate cancer” and “skin cancer” would be extracted from the example phrase. We used a simple rule-based system that identified candidate lists by searching for tagged biomedical entities that follow the word “and.” We then searched the preceding words in the candidate list and attempted word substitutions with the final term to find terms that were in the lexicon.

Across the 30,044,935 abstracts and 2,485,641 full-text papers that were minable, we find 3,269,632 potential mentions of non-contiguous entities in the example list format.
reviewed 100 of them to understand the error profile and found that 42% were true positives. The main errors were caused by spurious mistakes in the lexicon and a more conservative lexicon would likely improve precision but may affect overall recall. Nevertheless, this initial result suggests that many biomedical entities are described in the list form that would be missed with most current methods. While there are considerable false-positive dues to the dictionary matching method, we would argue that this will only be a fraction of non-contiguous entities across the biomedical literature as we examine only one type of linguistic structure that could contain non-contiguous entities.

Fig. 3 shows an overview of the results from the literature analysis. Lists appear more in full-text papers than in abstracts even when taking account of the substantially larger number of abstracts than full-text articles in the corpus. They can even appear in the article title. Furthermore, disease has substantially more non-contiguous entities, likely due to the larger number of multiple word terms in that lexicon (837,390 compared to 103,427 for genes for example).

This analysis strongly suggests that non-contiguous are a substantial problem in biomedical text mining and that methods that ignore them will be missing large amounts of potential extracted information. We hope our contribution to an annotation tool that could help visualize and annotate these problematic entities may take a step towards new methods to identify them.

**ORCID**

Jake Lever: https://orcid.org/0000-0001-8198-2939

Russ Altman: https://orcid.org/0000-0003-3859-2905

Jin-Dong Kim: https://orcid.org/0000-0002-8877-3248

**Authors’ Contribution**

Conceptualization: JL, JDK. Data curation: JL. Formal analysis: JL. Funding acquisition: RA, JDK. Methodology: JL. Writing – original draft: JL, RA, JDK. Writing – review & editing: JL, RA, JDK.

**Conflicts of Interest**

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**References**


There has been a dramatic increase in the popularity of utilizing social media data for research purposes within the biomedical community. In PubMed alone, there have been nearly 2,500 publication entries since 2014 that deal with analyzing social media data from Twitter and Reddit. However, the vast majority of those works do not share their code or data for replicating their studies. With minimal exceptions, the few that do, place the burden on the researcher to figure out how to fetch the data, how to best format their data, and how to create automatic and manual annotations on the acquired data. In order to address this pressing issue, we introduce the Social Media Mining Toolkit (SMMT), a suite of tools aimed to encapsulate the cumbersome details of acquiring, preprocessing, annotating and standardizing social media data. The purpose of our toolkit is for researchers to focus on answering research questions, and not the technical aspects of using social media data. By using a standard toolkit, researchers will be able to acquire, use, and release data in a consistent way that is transparent for everybody using the toolkit, hence, simplifying research reproducibility and accessibility in the social media domain.

Keywords: data mining, information storage and retrieval, machine learning, social media

Availability: All code described in this paper is fully available at: https://github.com/thepanaealab/SMMT.

Introduction

Only in the last six years, there has been a great influx of research works that describe different types of research works using Twitter and Reddit data, nearly 2,500 papers are found in PubMed [1]. These works encompass countless applications, such as the usage of opioids [2], the flu [3], eating disorder [4] networks analyses, depression symptoms detection [5], and diabetes interventions [6], etc. While all the listed studies use data from Twitter and Reddit, we can only find code available for one of them. Additionally, the data acquisition methodology is different on each study and seldomly reported, a crucial step towards reproducibility of any of their analyses. When it comes to using Twitter data for drug identification and pharmacovigilance tasks, authors of works like [7-9] have been consistently releasing publicly available datasets, software tools, and complete Natural Language Processing (NLP) systems with their works. In an attempt to shift the biomedical community into better practices for research transparency and reproducibility, we introduce the Social Media Mining Toolkit (SMMT), a suite of tools aimed to encapsulate the cumbersome details of acquiring, preprocessing, annotating and standardizing social media data. The need for a toolkit like SMMT arose from our work using Twitter data for the characterization of disease transmission during natural disasters [10] and mining large-scale repositories for drug usage related tweets for pharmacovigilance purposes [11]. We originally wanted to use other researcher’s tools and surprisingly we found very little code available with the majority outdated and non-functioning. Going one step
back, we did find rudimentary Python libraries to interact with the Twitter API, but some of their learning curves are steep and not overly documented. We then decided to clean and integrate our code into a toolkit that would help us provide a comprehensive resource for other researchers/users to replicate our work and to use in their own analyses, hence SMMT was born. Parallel to tools like SMMT, there are other research groups that are outlining frameworks to streamline the mining of social media like Sarker et al. [12], which are complementary to the use and need of this tool.

Methods

Programmed using Python version 3 and the latest Twitter API interfaces, the functionality of SMMT is divided into three separate sets of tools: data acquisition tools, data preprocessing tools, and data annotation and standardization tools. The particular versions of the additional Python libraries used by SMMT are available at the github documentation [13] since they are constantly updated and refreshed. Besides extensive usage documentation, the tool also provides two end-to-end usage examples, as well as additional Google Colaboratory [14] interactive Python notebooks with data usage examples. Note than in order to use most of the functionality of SMMT, users need to sign-up to acquire access to the Twitter Application Program Interface API. Once approved, users will be provided a set of API credential keys, more information can be found in [15]. Fig. 1 shows all current components of SMMT. In the following sections we provide additional details of each category of available tools.

Data acquisition tools

The tools in this category are used to gather data from social media sites, namely Twitter for this initial release of SMMT. The most common way of acquiring Tweets is to use the Twitter streaming API [16]. Our toolkit provides two separate utilities to capture streaming data, one will gather all available tweets and will continue running until terminated (streaming.py), and the other will take a list of search keywords and number of desired tweets and will pull those from the current tweet stream (search_generic.py). Details on how to use these utilities can be found on the README file.

The most common and permitted way of sharing Twitter data publicly is by only sharing the tweet id number. This number then needs to be ‘hydrated’, which means that the Twitter API needs to be used to fetch all the complete tweet and additional meta-data fields. This is a vital step for most users trying to replicate other studies or analyses. We provide a utility called get_metadata.py which reads a list of tweet ids and hydrates them automatically.

One of the major drawbacks of the Twitter API is the fact that unless having paid access to it, researchers cannot extract all historical tweets for any given Twitter user. Also, extracting all tweets from a given time range is not always easily and efficiently possible with the API. For these purposes we provide a utility called scrape.py which, once given a list of Twitter handles and corresponding date ranges, will automatically scrape the Twitter page and pull the tweet ids of the desired user and date range. These tweet ids then need to be ‘hydrated’ to be able to fully use them.

Data preprocessing tools

After having acquired enough data for research purposes from the Twitter stream, or identified and ‘hydrated’ a publicly available dataset, there is a need to subset the tweets and process the tweets JSON files to extract the fields of interest. While seemingly trivial, most biomedical researchers do not want to work with JSON objects, and since around 70% of the JSON fields are not populated, precise preprocessing steps need to be carried out to clean the data and render it useful in friendlier formats.

SMMT contains the parse_json_fine.py tool which takes a relatively small file (less than 1 Gigabyte in size) of Twitter JSON objects and separates these objects into a tab delimited file with each JSON field converted to a column and each tweet into a data row. With over 170 fields of meta-data, researchers are usually not interested in the vast majority of them. This tool can be configured to select which fields are of interest to be parsed and only process those into the tab delimited format. If the size of the tweets JSON objects file is larger than 1 Gigabyte, we provide an additional tool, parse_json_heavy.py, which can handle Terabyte sized files sequen-
tially rather than reading them all in memory for speed.

Once all the tweets are processed into the cleaner tab delimited format, which can even be read in Excel, there might be a need to further subset the tweets based on a given list of terms, or dictionary. For this purpose, we have included the `separate_tweet.tsv.py` file, which takes a term list in a format specified in the README file of SMMT and will return only the tweets that contain the provided terms.

**Data annotation and standardization tools**

After preprocessing the acquired social media data, researchers have the capabilities of standardizing their tweets’ text with our set of tools. Taking advantage of OntoGene Bio Term Hub [17] and their harmonization of biomedical terminologies into a unified format, we provide a tool, `create_dictionary.py`, that converts their downloads into SMMT-compatible dictionaries. To avoid creating a complicated and cumbersome format for our tool, we opted for simplicity and only rely on having a tab delimited file with an identifier column and a term name column. Other dictionaries that we have made available will standardize any annotations using the Observational Health Data Sciences and Informatics (OHDSI) vocabulary [18]. We are testing functionality to also convert our dictionaries to the PubDictionaries [19] format for the next release, allowing researchers to use their functionality and online REST services.

One of the most important tools of SMMT is the Spacy [20] NER annotator, `SMMT_NER_basic.py`, this tool will take the tab delimited tweets, a dictionary file, and the name of the output file for the annotations. In order to extend the usability of our tool, we provide the resulting annotations in a traditional format: document, span, term format; as well as pre-formatted outputs compatible with the brat annotation tool [21] and PubAnnotation and its viewer TextAE [22] as shown in Fig. 2.

**Discussion**

While all the tools have their own documentation, in order to ease the adoption of the tools available in SMMT, we have included an end-to-end example in the examples folder that performs the following tasks:

1. Download 300 tweets from the Twitter API stream for each of the following keywords: donald trump, coronavirus, cricket
2. We then preprocess those tweets to extract Tweet Id and their text into tab delimited files.
3. Using a Google Colab Notebook, we use these preprocessed files and then use the TF-IDF vectorizer on the text of the tweets to create a test and train set and build a Multi Nomial Naive Bayes Classifier to separate tweets based on their label. All details and steps of this process are outlined in the Colab Notebook.
4. We then test our trained model on the test set and generate a confusion matrix heat-map (Fig. 3) of the classification task, and show the model performance metrics.

The whole process of this example takes less than 30 minutes to complete and is heavily documented for SMMT users to overcome the learning curve of acquiring and preprocessing tweets. While our example is simple in nature, users can build upon it and modify it to better suit their needs.

The tools part of SMMT allow users to simplify their research workflows and to focus on determining which data they want to use and the analyses they want to perform, rather than deciphering how to acquire the data. While most cutting-edge and near real-time research will be done pulling tweets from the Twitter API stream, there are countless datasets available for historical research, from large general purpose databases like the Internet Archive’s Twitter Stream Grab dataset [23], which consists of data from 2014 to 2019, to more specialized and pre-curated datasets for.

![Sample SMMT_NER_basic annotation using an RxNorm–based dictionary and displayed on TextAE.](https://doi.org/10.5808/GI.2020.18.2.e16)
uses like Pharmacovigilance [11] among others. This initial version release of SMMT will continue growing with additional tools being developed for platforms like Reddit, Dark Web forums, and other social media data sources.

ORCID

Ramya Tekumalla: https://orcid.org/0000-0002-1606-4856
Juan M. Banda: https://orcid.org/0000-0001-8499-824X

Authors' Contribution

Conceptualization: RT, JMB. Formal analysis: RT, JMB. Methodology: RT, JMB. Writing – original draft: RT, JMB. Writing – review & editing: RT, JMB.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

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References


A proof-of-concept study of extracting patient histories for rare/intractable diseases from social media

Atsuko Yamaguchi¹, Núria Queralt-Rosinach²

¹Tokyo City University, Setagaya, Tokyo 157-0087, Japan
²Leiden University Medical Center, Leiden, 2333 ZA, The Netherlands

The amount of content on social media platforms such as Twitter is expanding rapidly. Simultaneously, the lack of patient information seriously hinders the diagnosis and treatment of rare/intractable diseases. However, these patient communities are especially active on social media. Data from social media could serve as a source of patient-centric knowledge for these diseases complementary to the information collected in clinical settings and patient registries, and may also have potential for research use. To explore this question, we attempted to extract patient-centric knowledge from social media as a task for the 3-day Biomedical Linked Annotation Hackathon 6 (BLAH6). We selected amyotrophic lateral sclerosis and multiple sclerosis as use cases of rare and intractable diseases, respectively, and we extracted patient histories related to these health conditions from Twitter. Four diagnosed patients for each disease were selected. From the user timelines of these eight patients, we extracted tweets that might be related to health conditions. Based on our experiment, we show that our approach has considerable potential, although we identified problems that should be addressed in future attempts to mine information about rare/intractable diseases from Twitter.

Keywords: intractable diseases, rare diseases, social media mining

Availability: In this paper, we used Twitter timelines and the Human Phenotype Ontology. We obtained user timelines from Twitter (https://twitter.com) using Python code (https://github.com/acopom/smm4rd) with Tweepy (https://www.tweepy.org/), which is a Python library for accessing the Twitter API (https://developer.twitter.com/). The Human Phenotype Ontology is available at https://hpo.jax.org/app/download/ontology.

Introduction

Social media has become a data source that is making a major contribution to big data. Recent scientific research has started to use and evaluate social media in the context of healthcare [1-4]. Svenstrup et al. [5] highlighted the potential of social media platforms dedicated to healthcare specialists as a means of knowledge-sharing for rare disease (RD) diagnoses. Schumacher et al. [6] introduced a case of online research and analysis of respondents using social media for the study of RDS. The role of social media was as a “participation caption” for recruiting a patient cohort and collecting clinical information. The authors concluded that the methodology and response patterns can be used for RD research. However, in those studies, social media platforms were used only from the viewpoint of healthcare specialists (e.g., medical doctors), even though a much broader range of people, including patients, are contributing to social media data. In particular, commu-
nities of patients suffering from RDs are very active on social media platforms. By definition, RDs affect small percentages of the population (https://ec.europa.eu/info/research-and-innovation/research-area/health-research-and-innovation/rare-diseases_en). These RD patient communities are small and patients are geographically scattered. Even though there are more than 8,000 RDs, only 5% have treatment. The lack of patient information available for research seriously hinders the diagnosis and treatment of rare and intractable diseases [7]. In general, RD patients suffer from very severe and heterogeneous symptoms and remain undiagnosed for several years [8]. Consequently, these disease communities use social media platforms to try to find other patients with similar health problems or expertise about their rare condition, sharing manifold types of information—including symptoms, treatments, side effects, and other diseases and activities—that go beyond what is normally captured in a clinical setting or patient registry [9]. Recently, Klein et al. [10] mined Twitter to collect data on rare health-related events reported by patients, and showed that this social media platform was useful for gathering patient-centric information that could be used for future epidemiological analyses. Our hypothesis was that data from RD patient histories posted on social media would capture patients’ perspectives of their health status, which may be valuable for research into ways of helping undiagnosed patients by accelerating the timeline to diagnosis and treatment.

The special theme of the Biomedical Linked Annotation Hackathon 6 (BLAH6) was “social media mining.” Therefore, we attempted to extract patient-centric knowledge from social media as a task for the 3-day hackathon. In this paper, we present our work that we conceived, designed, and developed during BLAH6 to explore the potential of social media data as a source of patient-centric knowledge. For this project, we focused on rare and intractable diseases and selected Twitter to obtain patients’ timelines, as this platform may contain descriptions of the history of their health conditions. By focusing on the date of diagnosis, we intended to obtain histories of their health conditions before and after diagnosis.

Methods

Due to the time constraints of the hackathon, we selected one RD and one intractable disease. Then, we searched for patients with the two diseases and obtained their timelines. We also tried to extract tweets related to the disease and symptoms from each timeline.

First, we selected a RD that is adult-onset and not too rare to facilitate the extraction of a proper amount of data for analysis. To do so, we used information on the number of patients diagnosed with rare and intractable diseases in Japan, provided by Japan Intractable Disease Information Center (https://www.nanbyou.or.jp/). Based on this information, we selected amyotrophic lateral sclerosis (ALS) as an RD, and for similar reasons, we selected multiple sclerosis (MS) as an intractable disease. Second, we obtained a list of Twitter users who were diagnosed with ALS or MS using the search terms “I was diagnosed” and the disease name. Then, we selected users diagnosed during the last 5 years who had more than 100 tweets, excluding retweets and replies. This resulted in four users for each disease. By using Tweepy with a Python script (https://github.com/acopom/smm4rd), we obtained the timestamp and the text of the Twitter timelines, including 6088 tweets without retweets and replies for the eight users.

To extract tweets dealing with a user’s health conditions, we used all terms in the Human Phenotype Ontology (HPO) [11] except for three (“all,” “left,” and “right”). All tweets that included HPO terms in the text were extracted. We then removed some tweets by manual search inspection because they described the health condition of someone else, such as the user’s child. Through this process, we obtained a set of tweets that were related to the user’s health condition. We called this set of tweets “tweets by HPO” for a user u and denoted it as $H(u)$.

Additionally, we extracted tweets dealing with health conditions using common words, such as “cold.” However, many tweets extracted in this way were not related to health conditions, for example, “It’s cold today.” Consequently, we manually removed many tweets from this extracted tweet set. We called this set of tweets “tweets by manual” for a user u and denoted it as $M(u)$.

We called $H(u) \cup M(u)$ “tweets about the disease” and denoted this set as $D(u)$. As each tweet in $D(u)$ may contain sensitive information from the viewpoint of user protection, a short summary of each tweet to conceal details was made manually.

Results and Discussion

To conceal the identity of the users with ALS and MS, we used ALS1, ALS2, ALS3, and ALS4 to refer to the ALS patients and MS1, MS2, MS3, and MS4 to refer to the MS patients instead of their Twitter user names. Table 1 shows the numbers of tweets, the number of tweets in $H(u)$, and the number of tweets in $M(u)$ for each user $u$. Of note, all tweets about ALS were posted after the users were diagnosed, whereas all tweets about MS, except for one, were posted before the diagnosis.

We next constructed a patient history for each user $u$ using tweets in $D(u)$. For example, ALS1 had two tweets in $H(\text{ALS1})$ extracted by the HPO term “pain” (HP:0012531). $M(\text{ALS1})$ in-
Table 1. Summary of the eight users analyzed in this experiment

<table>
<thead>
<tr>
<th>User</th>
<th>#Tweets</th>
<th>#H</th>
<th>#M</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALS1</td>
<td>2135</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>ALS2</td>
<td>1295</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>ALS3</td>
<td>213</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>ALS4</td>
<td>182</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>MS1</td>
<td>777</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>MS2</td>
<td>348</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>MS3</td>
<td>572</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>MS4</td>
<td>566</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>6088</td>
<td>16</td>
<td>13</td>
</tr>
</tbody>
</table>

#Tweets, #H, and #M show the total numbers of tweets, the number of tweets in H, and the number of tweets in M, respectively.

from these five tweets, we obtained four events related to health conditions because two of the tweets in $H(\text{ALS1})$ indicated one event. Fig. 1 shows the patient history of ALS1, who had four events after diagnosis. We set the date of diagnosis as a reference point. We presented short summaries such as “can talk” instead of showing real tweets because the extracted tweets may contain sensitive information from the viewpoint of user protection. At 270 days after the date of diagnosis, we can see that ALS1 could work, walk, and talk. However, ALS1 could no longer walk 644 days after the date of diagnosis.

Similarly, Fig. 2 shows the patient history of MS1, who had three events constructed by four tweets in $D(\text{MS1})$. MS, multiple sclerosis.

Fig. 1. Patient history with four events constructed by five tweets in $D(\text{ALS1})$. ALS1, amyotrophic lateral sclerosis 1.

Fig. 2. Patient history with three events constructed by four tweets in $D(\text{MS1})$. MS, multiple sclerosis.
events as constructed by four tweets in D(MS1). MS1 had an asthma attack 2,102 days before the diagnosis, and experienced anxiety and received a drug for it 375 days before the diagnosis.

This experiment showed the potential of Twitter data as a source of patient-centric knowledge, by extracting tweets related to health conditions and constructing a patient history from each user’s timeline. However, we found that the typical method of scientific data extraction did not work well for mining tweets. As shown in Table 1, we obtained a very small number of tweets related to health conditions. To address this limitation, the development of a dictionary for the healthcare domain specialized for social media data is vitally necessary to leverage and better understand the scientific value of data from social media for rare and intractable diseases.

ORCID

Atsuko Yamaguchi: https://orcid.org/0000-0001-7538-5337
Núria Queralt Rosinach: https://orcid.org/0000-0003-0169-8159

Authors’ Contribution

Conceptualization: AY, NQR. Data curation: AY. Formal analysis: AY. Funding acquisition: AY, NQR. Methodology: AY, NQR. Writing – original draft: AY. Writing – review & editing: AY, NQR.

Conflicts of Interest

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Introduction

According to the Directive 2010/63/EU (https://eur-lex.europa.eu/legal-content/EN/ALL/?uri=CELEX%3A32010L0063), researchers who plan to carry out animal experimentation are required to examine whether alternative methods not entailing the use of a live animal are already available for the planned research purpose (replacement). In addition, the chosen method should ensure that the animal number is reduced to a minimum without comprising the objective (reduction), and to reduce the possible pain, distress, and suffering (refinement). These measures are known as the 3R principles.

When searching for alternative methods to animal experiments, researchers have to carry out various queries to bibliographic databases, e.g., PubMed (https://www.ncbi.nlm.nih.gov/pubmed/), and carefully analyze the potential candidate articles. For each of these potential articles, the researcher should check whether it addresses two important issues: (1) a method for replacement, and (2) the planned research question. To assist researchers in their search for alternative methods, we are currently developing a Web application that addresses these two aspects. We rank the potential candidate articles based on the similarity of the research question (with regard to an input article) and iden-
tify the proposed methods in each of the articles.

For the implementation of the Web application, we rely on various tools, such as document classification, named-entity recognition, and annotation storage, among others. In the scope of the BLAH6 Hackathon (https://blah6.linkedannotation.org/), we integrated the PubAnnotation ecosystem [1] into the backend of our application. PubAnnotation contains three main components that can support our application for some of those tasks: the PubAnnotation repository, PubDictionaries, and the TextAE annotation tool.

Here we describe the integration of these tools into our application. We start by introducing our application, followed by how the components are being integrated into it. The Web application is still under development and not yet available for the final user. However, the resources that we created in the PubAnnotation platform are already available to the research community.

The Web Application

Fig. 1 shows an overview of the real-time interaction of the Web application with PubMed and PubAnnotation. Given a reference article as input, our application retrieves the so-called similar articles from PubMed (https://www.ncbi.nlm.nih.gov/pubmed), i.e., the ones that were pre-compiled by PubMed [2]. The tool performs two processing tasks based on the title and abstracts of these similar articles: (1) classification of the proposed methods; and (2) ranking of the retrieved similar articles according to the similarity of their research questions to the one in the reference article.

The classification of the proposed methods will utilize machine learning algorithms to be trained based on manually annotated articles (abstracts), which are currently being manually labeled. These labels cover the various types of methods that are relevant for our domain, such as whether the experiments have been carried out in vivo (e.g., vertebrates, invertebrates) or in vitro (e.g., cell lines or organs). Further, we are also experimenting with named-entity recognition tools to support this task. We focus on entities which are still not well supported by the existing tools, e.g., cell lines, and on a dictionary-based approach that relies on the comprehensive Cellosaurus resource [3].

For the ranking task, the application calculates the text similarity between the reference article and each of the PubMed similar articles, and the resulting scores are used to rank these articles. For the text similarity, we utilize the TextFlow tool [4]. However, instead of relying on the whole abstract of the articles, we utilize only the most relevant discourse categories (or zones), such as “introduction” and “results”. This is due to the fact that only some parts of the abstract potentially describe the research question.

We recently published a study in which we compared four tools for the extraction of the zones and evaluated them on seven use studies (https://github.com/mariananeves/scientific-elements-text-similarity) [5]. Our study also demonstrated that using pre-selected zones, instead of the whole abstracts, yields better performance in the ranking task.

The zones can be manually annotated or automatically detected. The manually annotated zones are the original ones included in the structured abstracts in Pubmed (https://www.nlm.nih.gov/bsd/policy/structured_abstracts.html). However, given that not all abstracts in PubMed are structured, we automatically extract the zones for the remaining articles using the ArguminSci tool [6]. This was the best performing tool according to our study [5].

Finally, our Web application will contain visual components to display the abstract of the articles involved in the search. We currently consider two scenarios. The first is the visualization of the reference article in order to obtain feedback from the user, e.g., the research question in mind, by asking the user to highlight this in-
formation on the text. The second scenario is a side-by-side display of the reference article and each one of the retrieved similar articles in order to compare two articles and further gather user feedback.

Integration with the PubAnnotation Ecosystem

We are currently integrating the PubAnnotation ecosystem in various components of our Web application in order to support various tasks, namely, storage, alignment, named-entity recognition, and visualization of annotations. Here we describe how each of the tools is being integrated in our application.

PubAnnotation database

We utilize the PubAnnotation database to allow easy storage and retrieval of PubMed titles, abstracts, and their annotations. We store the zones coming from the reference article and its similar articles into one of the two repositories that we created in PubAnnotation, depending on the origin of these zones: (1) the PubMed_Structured_Abstracts repository (http://pubannotation.org/projects/PubMed_Structured_Abstracts) for the original zones available in the structured abstracts in PubMed; and (2) the PubMed_ArguminSci repository (http://pubannotation.org/projects/PubMed_ArguminSci) for the zones automatically extracted by the ArguminSci tool [6]. Both repositories are public and the annotations (zones) can be retrieved using the PubAnnotation API (http://www.pubannotation.org/docs/intro/).

For each article processed by our application, both reference articles or similar articles, we first check whether the article is already included in the PubAnnotation (cf. “fetch article” in Fig. 2), i.e., in any of its repositories. The output is either a message that the article is inexistent or a JSON object that includes the article’s title, abstract, and its annotations, which may come from various repositories in PubAnnotation. We check whether annotations already exist from any of our two repositories described above (cf. “get zones” in Fig. 2). If any zones could be found, these are returned to be further processed by the Web application.

In case that no zones have been stored for the article in none of our two repositories, we first check whether the article contains a structured abstract. This information is contained in the data retrieved from PubMed. If the article contains a structured abstract, its zones are simply stored into PubAnnotation (cf. “store zones” in Fig. 2) and will be available for future queries. Otherwise, we extract the zones using the ArguminSci tool, followed by their stor-
age into PubAnnotation. For any of the two situations, we store the zones in PubAnnotation in a two-steps procedure: (1) we add the article into the corresponding repository (either PubMed_Structured_Abstracts or PubMed_ArguminSci), and (2) we add the annotations into the same repository. It is not possible to perform the second step if the article was not previously included in the repository.

As of May 2020, the PubMed_Structured_Abstracts repository contains more than 31k documents while the PubMed_ArguminSci repository holds almost 80k documents. Therefore, we can state that less than 30% of the documents processed by our application included a structured abstract, while we had to perform predictions for zones for more than 70% of them. These documents consist of reference articles and the corresponding similar articles derived from the various queries that we made to our application in the last months, but also from the machine learning experiments that we carried out for the document classification step. Currently, we do not plan to include zoning annotations for all articles in PubMed, but just for those that happen to be processed by our application during our various experiments, and later, from the queries made by the users. Therefore, the repositories should incrementally grow with the time.

Another interesting feature in PubAnnotation is the annotation alignment. We deal with annotations retrieved from two sources, i.e., ArguminSci and PubMed Structured Abstracts, whose annotations might have been derived from a slightly different version of the article’s abstract, or the corresponding text somehow altered by the tool during processing. The annotation alignment function in PubAnnotation automatically converts the offsets of these annotations to the article’s abstract that is stored in PubAnnotation. Therefore, this function relieves us from writing customized scripts for dealing with the annotations returned by the various resources or tools.

Currently, no storage in PubAnnotation is being carried out for annotations coming from the classification task. This is due to a couple of reasons. First, the performance of our algorithms is not yet satisfactory. Further, document-level annotations are currently not supported by the JSON format of PubAnnotation. However, we plan to store them in PubAnnotation in the near future.

TextAE (Text Annotation Editor)

Besides using the PubAnnotation ecosystem for annotation storage, we also plan to rely on other tools of the platform in our Web application. For instance, we are currently experimenting with the TextAE tool (http://textae.pubannotation.org/) for displaying articles and annotations to the user. TextAE can be embedded into a HTML page to display the text and annotations that are passed in the JSON format. TextAE can be used in both of our visualization scenarios, i.e., either for displaying single articles or a side-by-side comparison. For the first scenario, an editable version of TextAE can potentially be used for collecting user feedback on the reference article, i.e., through text highlighting. For the second scenario (side-by-side comparison), we currently display annotations for species, disease, and chemicals from PubTator Central [7] using its annotator (http://pubannotation.org/annotators/PubTator) that is currently available in PubAnnotation. We also envisage relying on a side-by-side comparison to gather feedback from the user about the similarity of both research questions.

PubDictionaries

We are also experimenting with PubDictionaries in the PubAnnotation ecosystem. Given a dictionary composed of terms (i.e., sets of identifiers and names), it is possible to perform a dictionary-based named-entity recognition by matching the terms in the dictionary to the title and abstract of articles in PubAnnotation. We are evaluating this functionality for the task of identifying cell lines, which might support our classification task, in addition to the machine learning approach.

For this purpose, we created the Cellosaurus_v33 dictionary (http://pubdictionaries.org/dictionaries/Cellosaurus_v33) that includes cell lines released in version 33 of Cellosaurus [3]. Based on this dictionary, we created a corresponding annotator in PubAnnotation (http://pubannotation.org/annotators/Cellosaurus_v33), which is a Web service that can be applied to any article in PubAnnotation for real-time annotation. We are currently developing a pre-processing script to filter out cell line names that match to a list of stopwords. Further, a post-processing script will also be applied to filter out mentions that match entities returned by PubTator Central, which are potential false positives.

Conclusion

We presented the integration of the PubAnnotation ecosystem in our planned Web application which aims to mine alternative methods to animal experiments. We use all main functionalities of the ecosystem, namely, the PubAnnotation repository for the storage and alignment of annotations, PubDictionaries for dictionary matching of cell lines, and the TextAE annotation tool for the visualization of articles and annotations. Two repositories for annotations of discourse elements were created and are available to the research community. Further, these two repositories are being frequently and automatically updated by our Web application. Finally, we also released a cell line dictionary and its corresponding annotator.
ORCID

Mariana Neves: https://orcid.org/0000-0002-6488-2394

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

References

Enabling a fast annotation process with the Table2Annotation tool

Pierre Larmande¹,²*, Kazim Muhammed Jibril²

¹DIADE, Univ. Montpellier, IRD, Montpellier 34398, France
²ICTLab, USTH, Hanoi 10000, Vietnam

In semantic annotation, semantic concepts are linked to natural language. Semantic annotation helps in boosting the ability to search and access resources and can be used in information retrieval systems to augment the queries from the user. In the research described in this paper, we aimed to identify ontological concepts in scientific text contained in spreadsheets. We developed a tool that can handle various types of spreadsheets. Furthermore, we used the NCBO Annotator API provided by BioPortal to enhance the semantic annotation functionality to cover spreadsheet data. Table2Annotation has strengths in certain criteria such as speed, error handling, and complex concept matching.

Keywords: bioinformatics, ontologies, semantic annotation
Availability: GitHub: https://github.com/pierrelarmande/Table2Annotation.

Introduction

Semantic annotation has been defined in various ways by various authors, but these definitions are all similar and reflect a single clear purpose. For instance, Oliveira and Rocha [1] defined semantic annotation as the process in which semantic concepts are linked to natural language. Liao et al. [2] defined semantic annotation as methods of describing resources (texts, images ...) with metadata where the meaning has been specified in an ontology. According to Oliveira and Rocha [1] semantic annotation can be seen as a methodology of adding metadata—comprising classes, properties, relations, and instances (i.e., the concepts of an ontology)—to web resources to be able to give or allocate semantics. Summarizing all of these definitions, we can simply state that semantic annotation is a way of matching resources to ontologies.

To make this point clearer, take this example of the text “...days to flowering...” With the help of semantic annotation, we would be able to match this text to the ontology concept “days to flowering trait” from the Trait Ontology [3], which has the concept ID of TO:0000344.

Semantic annotation helps to boost the ability to search and access resources. It is also a step towards data FAIRification [4]. According to Jovanovic and Bagheri [5], semantic annotation can be used in information retrieval to expand the queries from the user with some ontology terms and also to provide a grouping of documents retrieved based on specific content. Biomedical resources contain numerous abbreviations in the texts, sometimes with different meanings, which makes it hard to perform comprehensive searches. Semantic annotation helps to disambiguate these abbreviated terms based on the way they appear in a certain context.

In this paper, we sought to identify ontological concepts in scientific texts. This could
be seen as an ontology matching process, in which natural language texts are matched with concepts. There are already some existing web services and tools that use semantic annotation for ontology matching, as have been evaluated by Oliveira and Rocha [1]. However, few of these tools handle spreadsheet data as text input. We developed the Table2Annotation tool with that purpose because there is a need for such a tool in the life sciences community, which produces extensive experimental data in spreadsheets. Semantic annotations will facilitate more complex analyses across several datasets.

The paper is organized as follows. Section 2 defines the challenges of semantic annotation. Section 3 presents an overview of Table2Annotation. Section 4 analyzes the results of semantic annotation through some examples. Section 5 concludes the manuscript.

**Semantic Annotation Challenges**

Semantic annotation has some benefits, but some challenges are also faced during the annotation of biological texts or other resources. Some of these challenges, as also described in previous research [6-8], are follows:

- **Word sense disambiguation**: It is necessary to determine the correct meaning of a word as used in a sentence when a word has multiple meanings.
- **Spelling/grammatical error identification**: Correcting spelling or grammar in biomedical texts is very important. Spelling and grammar errors cause ambiguity in already sparse text.
- **Discontinuous entities**: Entities can be composed of multiple words in a discontinuous span. For example, “drought and salinity tolerance” means “drought tolerance and salinity tolerance,” but in this case we might only have matching for “salinity tolerance.”
- **Gene/protein disambiguation**: In the biomedical context, all proteins have associated genes, often with the same name, making it difficult to annotate texts dealing with genes and proteins.
- **Detection of name variants**: Variations of entity naming can take many forms, thereby complicating annotations. For example, abbreviations and shorthand texts are difficult to normalize with ontological concepts.

The challenges faced in annotation can be tackled by two approaches, which can be also combined. The first one is the term-to-concept matching method, which involve matching some parts of the provided text to structured knowledge databases, dictionaries, or vocabularies. However, it is difficult to maintain comprehensive lexicons to be used for annotation. The second approach is machine learning, which involves creating annotators for specific purposes and usage instead of more general ones [5].

Of particular note, the third challenge (discontinuous entities) can be tackled by creating algorithms that can transform texts with conjunctions like “and” or “or.” Thus, in the example of “drought and salinity tolerance,” an algorithm could transform this phrase to “drought tolerance and salinity tolerance” before the annotation process.

Although these are good solutions to tackle some challenges, some drawbacks remain. For instance, a drawback of term-to-concept matching is its inability to disambiguate terms, so annotators that inherit this method usually match terms with several possibilities. This drawback is encountered in the use of the NCBO annotator [9], and one way to solve this problem is to have several algorithms that use knowledge-based dictionaries to transform ambiguous terms into meanings that are clear for the annotator. These algorithms should also be able to correct incorrect grammar usage and wrong spellings by matching dictionary terms with similar spellings or phrases.

**Challenges of semantic annotation tools**

Diverse tools are used in semantic annotation [1]. These tools also encounter some challenges, a few of which are listed below:

- **Speed**: This is one of the most common challenges. Annotations performed on huge datasets can take a lot of time to process.
- **Language specificity**: Most annotators are in English, which makes it difficult to apply semantic annotation in other languages.
- **Document genre genericity**: Annotators that support document input can face the problem of having to annotate different document formats, and not supporting a particular format could be a challenge.
- **Text variation**: According to Jovanovic and Bagheri [5], challenges are faced also due to the fact that there are different kinds of biomedical texts and variations in texts, for example between biomedical and clinical texts.
- **Entity disambiguation**: Entities mentioned in biomedical texts sometimes do not have enough context to disambiguate them.

These challenges and others are been studied, and many experts have tried to figure out ways to tackle them in newly developed systems. It may not be possible to fully resolve these challenges, but they can be reduced, and the following section shows how we tackled some of these challenges in the system developed for this project.

**Overview of the Table2Annotation Tool**

In this section, we describe the proposed solution to build an ontology matching system. Our solution uses the NCBO annotator...
web service API for primary information retrieval.

The NCBO annotator annotates data with the MGrep term-to-concept matching tool and retrieves sets of annotations that are later expanded using various methods of semantic matching, meaning that this annotator goes through two stages. This annotator is unique because of the method it uses to associate concepts, instead of looking for the concept that best matches the provided context. This annotator uses BioPortal [10] and although it does not support disambiguation of terms, it is suitable for real-time processing. This annotator is available for free and is implemented through web services. This annotator is currently used in AgroPortal [11] and BioPortal.

The flow of Table2Annotation is quite simple and understandable. The system starts by taking an input dataset (CSV, Excel, etc.) and then processes the file by reading the data and fetching the necessary data to be annotated. It takes the necessary data and calls an external API provided by AgroPortal to annotate the data. The results returned from this process are processed by taking the Uniform Resource Identifier (URI), concept ID, and the matched words. Finally, the annotated terms are saved and written to an output file for the user to access.

The operation of the matching system is described diagrammatically in Supplementary Fig. 1. In building this Table2Annotation tool we decided to use the NCBO annotator (AgroPortal API) to support the annotation of terms.

Important algorithms

As discussed in the challenges section, there are several problems that must be dealt with. Thus, we developed specific algorithms to handle some of them.

Threading

First of all, the system was created in a functional independent approach where the major functions are independent. For example, obtaining inputs and annotation are independent. This allows us to better handle the slower part of the system. The function that slows down the system is the one that deals with iterating through the cells, taking the cell data, and then annotating this data. To reduce the problem of speed, we decided to create an algorithm to speed up the process. The algorithm uses the concept of multi-threading, allowing the function to be run by several processors (threads) concurrently.

Permutation

As discussed above regarding the problem of discontinuous entities, although this issue has not been fully resolved and future enhancements remain to be made, the problem of conjunctions can be reduced by creating an algorithm to handle this case.

Multiple dataset formats

The problem of document genre genericity was reduced by creating an algorithm to detect the format of the file being input by the user and then handling the process depending on the file format.

Running Table2Annotation

Table2Annotation is a Java-based program that is currently executed through the command line interface. The user must have a dataset that he or she wants to annotate first. Table2Annotation is compiled after the code and all the functions explained in the previous section have been fully implemented. The compilation of Table2Annotation is done with all the necessary libraries included in the Java project. To run the system the user needs to input the following parameters: input file (mandatory), column (mandatory), suggestions (optional), slice (optional), separator (optional), and sheet (optional).

First, the user provides the path to the input file (dataset) and then provides the name of the column to be annotated. These two parameters are mandatory and the others are optional.

The other functions that can be passed as parameters are as follows: (1) suggestions (recommendations) of ontologies, allowing the user to specify which AgroPortal (or BioPortal) ontologies to use for the annotation process; (2) the slice (grouping), allowing the user to define which slice to use for the annotation process (slices can be compared to an instance of AgroPortal or BioPortal for a defined subset of ontologies); (3) the separator, if the file is a separated file type, allowing the user to define the type of separator used to split the cells; and (4) the sheet number if the file is an Excel file with multiple sheets.

After the command is executed, the system starts processing and stops when the process is completed. The results of this operation are output to a file in same format as the input file and given to the user.

Results

In this section, we describe the results obtained from the Table2Annotation tool. We also describe the context of obtaining the results and an evaluation of the system.

First of all, we needed a dataset to test the tool, as shown in Fig. 1. The dataset that we used was quite small, as using a small dataset better demonstrates how the results are obtained, but the same principles are applied when using a large dataset. The dataset contains a “PROPERTY” column, which contains the terms to be annotated.
Test without recommendations or slices
In this test, we ran the system without giving recommendations or slice options (i.e., an ontology list to map on provided by AgroPortfolio), and the results are shown in Fig. 1. In the results obtained by processing, we can see that there are three new columns: "PROPERTY_id", "PROPERTY_id_uri," and "PROPERTY_id_match." The first added column contains the concept IDs obtained from the annotation, the second added column contains the URIs of the concepts, and the third added column contains the matching of the terms with the concept.

Test with a slice
In this test, we test-ran the system by giving it a slice called "agrold", which contains ontology groups for agronomy. The results of the test are shown in Fig. 2. In the results, we can see three terms (highlighted in yellow) that do not match with any concept, because they do not have ontologies belonging to the "agrold" group.

Test with recommendations
In this test, we ran the system with three suggestion parameters: "PO (Plant Ontology)", "TO (Plant Trait Ontology)", and "PATO (Phenotypic Quality Ontology)." The results of the test are shown in Fig. 3. In the results, we can see that six terms (highlighted in pink) had no matching concepts, because we filtered the annotation to the three ontologies given in the suggestions.

Test with a permutation algorithm
In this section, we tried to show the effect of having an algorithm to solve the problem of conjunctions in terms, which was mentioned earlier. We annotated the term "drought and salinity tolerance" and Fig. 4 shows the results. Fig. 4 (A, dataset result without algorithm) shows the results from the operation without the algorithm, and we can see that there are only matches for "drought" and "salinity tolerance." Fig. 4 (B, dataset result with algorithm) shows the results from the operation with the algorithm, which yields
Conclusion

In conclusion, Table2Annotation has strengths in certain criteria such as speed, error handling, and concept matching. First, we use a multi-threading algorithm that runs the process very effectively and efficiently. Second, it handles errors and exceptions by ignoring them whenever they occur. If there is an error while matching one term, it skips the term with an error and continues to the next one. If there is a general error, it still completes the matching process, but returns empty results. This method of error handling allows the user to run the process while multitasking and return to obtain the results without having to worry about system process terminations. Last, the matching results are good, and we see that cases of conjunctions are handled appropriately, so that the results contain more matches. The filters (slice and suggestions) also help to tailor the results to match the user’s expectations.

The system has strengths, but also has some weaknesses, such as relying on an internet connection and being dependent on the API. The system uses an external API, which can cause problems. Firstly, the system cannot work offline as it needs internet access to call the external API, which could be seen as a weakness. Secondly, if the external API is down for some reason, the system cannot be used. These weaknesses can be solved by building a full annotation system that does not depend on the availability of any external annotation API.

In the future, we think that we can improve algorithms to handle grammar problems and disambiguation. These algorithms should use language dictionaries to be able to transform terms without meaning (short forms) to something understandable to improve the process of matching to concepts. For example, when an abbreviated term is encountered, there should be a dictionary to look up matches for three terms: “drought”, “salinity tolerance”, and “drought tolerance.”
the term and return the full meaning. This will further help to reduce the problems of spelling and grammar mentioned earlier.

**ORCID**

Pierre Larmande: https://orcid.org/0000-0002-2923-9790
Kazim Muhammed Jibril: https://orcid.org/0000-0002-0493-4973

**Authors’ Contribution**

Conceptualization: PL, KMJ. Data curation: KMJ. Formal analysis: PL, KMJ. Funding acquisition: PL. Methodology: PL, KMJ. Writing – original draft: PL, KMJ. Writing – review & editing: PL.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.

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**Supplementary Materials**

Supplementary data can be found with this article online at http://www.genominfo.org.

**References**

Improving accessibility and distinction between negative results in biomedical relation extraction

Diana Sousa*, Andre Lamurias, Francisco M. Couto
LASIGE, Departamento de Informática, Faculdade de Ciências, Universidade de Lisboa, 1749-016 Lisboa, Portugal

Accessible negative results are relevant for researchers and clinicians not only to limit their search space but also to prevent the costly re-exploration of research hypotheses. However, most biomedical relation extraction datasets do not seek to distinguish between a false and a negative relation among two biomedical entities. Furthermore, datasets created using distant supervision techniques also have some false negative relations that constitute undocumented/unknown relations (missing from a knowledge base). We propose to improve the distinction between these concepts, by revising a subset of the relations marked as false on the phenotype-gene relations corpus and give the first steps to automatically distinguish between the false (F), negative (N), and unknown (U) results. Our work resulted in a sample of 127 manually annotated FNU relations and a weighted-F1 of 0.5609 for their automatic distinction. This work was developed during the 6th Biomedical Linked Annotation Hackathon (BLAH6).

Keywords: biomedical research, knowledge base, negative results, relation extraction

Introduction

Researchers and clinicians need to have access not only to known relations between biomedical entities but also to relations that were already disproven. Accessible negative results limit their search space and prevent the costly re-exploration of research hypotheses. However, most biomedical relation extraction datasets do not seek to distinguish between a false and a negative relation among two biomedical entities, and few knowledge bases hold negative examples. Some domain-specific exceptions are worth noticing, such as the Negatome database [1] for protein-protein interactions, and the phenotype-disease relations annotation file made available by the Human Phenotype Ontology (HPO) organization [2] that contains both positive and negative relations.

A false relation should express a context where the entities are not related. In contrast, a negative relation should express a context where there is an affirmation of no association between the two entities. Furthermore, datasets created using distant supervision techniques also have some false negative relations that constitute undocumented/unknown relations [3]. These relations are not marked true because they are not described in a knowledge base at the moment of the dataset creation, even though upon reading the context of these relations within their respective sentences one can support a true relation. Unknown relations are good examples of hypotheses to be further explored by re-
searchers and clinicians and can be of use to effectively populate the biomedical relations knowledge bases.

We propose to improve the distinction between false, negative, and unknown (FNU) relations, by:
- Revising a subset of the relations marked as false on the phenotype-gene relations (PGR) corpus [4] to create a sample dataset of FNU relations (made available on PubAnnotation platform (http://pubAnnotation.org/collections/Annotation%20of%20Human%20Phenotype-Gene%20Relations%20-%20Identification%20of%20False%20and%20Unknown%20Relations) [5])
- Implementing the first steps (using regular expressions and a neural network) to automatically distinguish between the FNU relations, using the previous sample FNU dataset as a test set.

**Methodology**

The PGR corpus consists of 1,712 abstracts, 5,676 human phenotype annotations, and 4,283 relations [4]. This automatically annotated corpus distinguishes between false and true relations but fails to identify different types of FNU relations. Fig. 1 illustrates the levels that we considered to represent true PGR relations (true, positive, and known), and false PGR relations (false, negative, and unknown).

Previously, our team had an expert curating a subset of the PGR corpus (around 30%). These annotations were initially divided into true and false, for a different scope out of the reach of this work. Nonetheless, for this work, we used the 127 false annotations curated by our domain expert in that subset to make the distinction between false (F), negative (N), and unknown (U) relations. The distribution of each type of relation is displayed in Table 1.

Some concrete examples of what sentences constitute each type of relation are presented in Fig. 2.

The manual annotations allowed for the assessment of common patterns for the false and negative types of relations:
- False relations are often enumerations or an explanation of protocol that does not imply any type of relation.
- Negative relations are more regular, with words that imply the negation of association, such as non, no, dissociation, and not, frequently combined with associated, and involved.

Contrarily, unknown relations follow intractable patterns and are the most heterogeneous.

The first approach towards catching false and negative examples that follow the specified patterns was using regular expressions by:
- Analyzing the list of detected negative expressions and of detected false expressions and possible equivalences (for instance, for the negative expressions list, not associated).
- Introducing patterns that use those expressions, such as ‘(+gene_entity+)+phenotype_entity+’ (.*’)+negative_expression+(.*’)+gene_entity+’ that translates to gene or phenotype followed by negative expression followed by gene or phenotype (for negative examples).
- Evaluating using the manually curated dataset of 127 FNU relations (gold standard dataset) if those patterns are able to correctly classify the FNU relations.

Using regular expressions based on the annotation process can and probably will introduce a bias towards the relations that we annotated. Further applications of these regular expressions should be explored for the approach to be fully validated. Nevertheless, the creation of the regular expressions was done posterior-

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**Table 1.** Distribution of each type of FNU relation: false, negative, unknown, and the total number of relations

<table>
<thead>
<tr>
<th></th>
<th>False</th>
<th>Negative</th>
<th>Unknown</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>No.</td>
<td>73</td>
<td>11</td>
<td>43</td>
<td>127</td>
</tr>
</tbody>
</table>

**Fig. 1.** Illustration of the levels that correspond to the true phenotype-gene relations (PGR) relations (true, positive, and known), and false PGR relations (false, negative, and unknown). Also, some generic sentences that elucidate the distinction between false and negative relations, and the distinction between known and unknown relations, according to the authors.
ly to the annotation process, solely based on the patterns described above, with the goal of generalizing as much as possible to avoid overfitting.

As a second approach, we briefly tried to apply a neural network using the Keras library (without any tuning, due to time constraints). For this purpose, we divided the FNU dataset into a training set (70%, 89 FNU relations) and a test set (30%, 38 FNU relations).

Results and Discussion

The application of a small subset of regular expressions to catch false and negative examples that follow the previously mentioned patterns had some promising results. We opted for the unknown relation as our default label since this type of sentences are more heterogeneous with irregular patterns that are difficult to capture by the use of regular expressions. Testing against the gold standard dataset shows a weighted-F1 of 0.5609. Other relevant metrics are displayed in Table 2.

The use of the neural network produced poor results (0.2308 accuracy) mainly due to the lack of tuning and the small size of our FNU dataset.

These preliminary results show that it is possible to capture common patterns of false and negative relations with high precision, but also shows the need for more work and possible exploration of machine learning techniques in order to capture more instances of those types of relations. More manual work, building regular expressions, should boost these preliminary results. Using syntax and dependency parsing to capture complex enumerations can also boost performance (e.g., enumerations where a group of genes is associated with a phenotype A and another group of genes is related to phenotype B).

Conclusions and Future Work

This work demonstrated that regular expressions are a feasible way of capturing differences between FNU relations, at least at a preliminary stage. The false and negative types of relations follow distinctive patterns that should be further explored to boost the weighted-F1 of 0.5609. Preliminary work with neural networks showed poor results (due to time constraints), but tuning the training and a larger dataset should boost these early results.

Future work could be revising all the false relations within the PGR corpus, and also of other datasets. Negative relations in manually annotated datasets should be easier to detect since the unknown relations would not be present. All of this will allow us to further explore machine learning approaches to tackle this problem more effectively.

Table 2. The evaluation metrics (precision, recall, and f-measure) for the false, negative, and unknown relations, and the weighted-F1 for all classes

<table>
<thead>
<tr>
<th>Type of relation</th>
<th>Precision</th>
<th>Recall</th>
<th>F-measure</th>
<th>Weighted-F1</th>
</tr>
</thead>
<tbody>
<tr>
<td>False</td>
<td>0.8438</td>
<td>0.3699</td>
<td>0.5143</td>
<td>0.5609</td>
</tr>
<tr>
<td>Negative</td>
<td>0.8333</td>
<td>0.4545</td>
<td>0.5882</td>
<td></td>
</tr>
<tr>
<td>Unknown</td>
<td>0.427</td>
<td>0.8837</td>
<td>0.5758</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 2. Example sentences for each type of false, negative, and unknown (FNU) relation: false (PMID:25343988), negative (PMID:16960806), and unknown (PMID:28698647). Also, the identified entities for each sentence, and their identifiers in the National Center for Biotechnology Information (NCBI) (for genes) and HPO (for human phenotypes).
Authors’ Contribution

Conceptualization: DS, FMC. Data curation: DS, AL. Formal analysis: DS. Funding acquisition: FMC. Methodology: DS, AL, FMC. Writing – original draft: DS. Writing – review & editing: DS, AL, FMC.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

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References

SciBabel: a system for crowd-sourced validation of automatic translations of scientific texts

Felipe Soares¹, Rozane Rebechi², Mark Stevenson¹

¹Computer Science Department, University of Sheffield, Sheffield S38RA, UK
²Instituto de Letras, Universidade Federal do Rio Grande do Sul, Porto Alegre 91540-000, Brazil

Scientific research is mostly published in English, regardless of the researcher’s nationality. However, this growing practice impairs or hinders the comprehension of professionals who depend on the results of these studies to provide adequate care for their patients. We suggest that machine translation (MT) can be used as a way of providing useful translation for biomedical articles, even though the translation itself may not be fluent. To tackle possible mistranslation that can harm a patient, we resort to crowd-sourced validation of translations. We developed a prototype of MT validation and edition, where users can vote for that translation as valid, or suggest modifications (i.e., post-editing the MT). A glossary match system is also included, aiming at terminology consistency.

Keywords: crowdsourcing, linguistics, machine translation, medical informatics applications, PubMed

Availability: Available online under the MIT license at https://github.com/soares-f/scibabel.

Introduction

Research in the biomedical domain, particularly about treatments and procedures for humans, can help improve the patient care offered by physicians. Evidence-based medicine is based on the premise that physicians give the best care possible when they base their treatments on reliable scientific evidence. But, although in practice this access is possible, there is a limitation that makes evidence-based medicine out of the reach for many physicians: almost all of its contents are written in English.

During the first half of the 20th century, scientific research was published in a variety of languages. But, as Gordin [1] described in detail, a complex set of factors led to English becoming the language of most scientific publications following the Second World War. Researchers tend to publish in English regardless of their native language. But, while academic researchers are often proficient in English, this may not be true for physicians in non-English speaking countries.

Translation of documents into the languages with which physicians are familiar seems like an obvious way to make the world’s scientific production accessible to them. But new research is produced so quickly and its results are published so rapidly that translating the information manually would be impractical. For example, in 2019 alone, more than 10,000 new articles were published in PubMed (PubMed Query: ((“2019”[Date - Publication] : “3000”[Date - Publication])) AND (treatment[Title/Abstract]) AND (procedure[Title/Abstract]) containing the keywords “treatment” and “procedure”—exactly
the kind of articles that would be of interest to physicians. However, there is a technology that could potentially do this translation automatically: machine translation (MT).

MT is a technology to render texts written in one language to another language. Modern MT research began just after the Second World War with the automatic translation of Russian scientific texts to English [2] as part of the scientific response to the Cold War (e.g., see Hutchins [3]). Machine translation research fell into decline soon thereafter due to considerable skepticism about whether practical MT systems were possible within the research community [4], but MT resurfaced in the 1990s with the advent of more powerful computers and alternative approaches. The field of MT experienced explosive growth after the September 2001 terrorist attacks and is an active area of scientific research [5-8]. This effort has led to a substantial improvement in the quality of translations produced by MT systems [9].

The earliest work on MT for scientific content concentrated on the physical sciences, however the focus of current research is shifting towards biomedical texts, especially due to shared tasks. This difference is important because, while users of translations in other scientific fields can tolerate some amount of error, as they do not have such a strict vocabulary and are not dealing directly with human beings, even a small mistranslation in this domain (e.g., a drug name being incorrectly translated, or a negation being ignored) could lead to disastrous consequences to patients. For example, consider Supplementary Table 1 which shows examples of a simple medical instruction (i.e., “Take two pills orally every day unless you feel dizzy or lightheaded”) usually found in drugs prescriptions translated into Finnish, Korean, Portuguese, Italian, Spanish, Japanese, French, German, Russian, Chinese (simplified) and Ukrainian by Google Translate. The third column contains their translations back into English by an educated native speaker (a common method of evaluating MT, similar to an approach known as back-translation) [10]. Contraindications that have been incorrectly translated are highlighted in bold font and it can be seen that these occur in six of the 11 translations. This demonstrates the need for automatic translations to be manually checked for critical mistranslations. However, this process is time-consuming and unlikely to scale well. Therefore, we propose a crowdsourced approach to validate automatic translations of biomedical articles and develop a prototype to facilitate such task.

In the proposed system volunteers who are able to read biomedical articles in English and also in another language would check MT output for critical mistranslations and vocabulary adequacy. The purpose of this system is to guarantee that the message in the source text is correctly conveyed in the translation, even though the translated text may lack fluency. Volunteers would accept the proposed translations if they are correct and be able to make editions when appropriate (e.g., incorrect terminology). We expect that our system, named SciBabel, would allow physicians and medical staff not proficient in English to access the most recent advances in medicine, enabling them to provide their patients with better treatment. The source code is available at https://github.com/soares-f/scibabel.

**Background**

An illustration of the recent improvements in MT can be seen from the performance of systems reported in the biomedical track of the Conference on Machine Translation (WMT), which focuses on the translation of PubMed abstracts. Translation quality increased by around 51% (or 16 percentage points) from 2016 to 2019 for English to Spanish. In Table 1 [11-16], we show the MT performance for some language pairs for biomedical texts with dates ranging from 2013 to 2019 for selected language pairs. Note that translation quality is measured automatically using the BLEU

<table>
<thead>
<tr>
<th>Reference</th>
<th>Language pair</th>
<th>Score (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>English → Spanish</td>
<td>BLEU: 31.11</td>
</tr>
<tr>
<td></td>
<td>English → Spanish</td>
<td>BLEU: 37.93</td>
</tr>
<tr>
<td></td>
<td>English → Spanish</td>
<td>BLEU: 47.01</td>
</tr>
<tr>
<td>Peng et al. (2019) [16]</td>
<td>English → German</td>
<td>BLEU: 35.26</td>
</tr>
<tr>
<td></td>
<td>English → French</td>
<td>BLEU: 38.29</td>
</tr>
<tr>
<td></td>
<td>English → Chinese</td>
<td>BLEU: 37.09</td>
</tr>
</tbody>
</table>

For years 2018 and 2019, metrics refer to the WMT challenge of the respective years.
score, a common MT metric that relies on the overlapping portions of the generated translations and the manually translated text [17].

In the two most recent WMT conferences (2018 and 2019) interesting results were reported for the English/Portuguese and English/Spanish language pairs. For instance, for the English to Spanish, the number of MT-generated sentences judged by humans as better than human translations was larger than the number of human sentences judged better than MT ones.

When combining the number of times that the best MT was equally good or better than human translation for WMT19, we get an average of 73% of correct translations according to human judgment, with surprising 90% for EN/ES and 82.09% for ZH/EN. This strengthens our point that MT can indeed be used to aid dissemination of biomedical scientific content.

However, as shown in Supplementary Table 1, MT systems can make critical mistakes when considering the usage of a medicine, for instance. It has been shown in literature that even human translation is prone to errors [18]. That is why the translation and localization industry usually has a two-step (or even more) process for translation. That is, at least one additional human is involved in checking the translation already carried out (also called proof-reading) [19].

Crowdsourcing of intensive tasks is not new in science. One example can be the Folding@Home initiative [20], which was popular in the first decade of the years 2000. This initiative consisted of crowdsourcing computational power from regular end-users (that signed to the initiative) to simulate protein folding, drug design, and molecular dynamics. Similarly, Seti@Home [21] tried to follow the same path to search for extraterrestrial life.

The crowdsourcing of manual annotation (or evaluation) was already explored by different authors [22,23]. For instance, the information retrieval (IR) shared tasks can be seen as the pioneers of human distributed annotation. Participants of IR shared tasks would blindly evaluate the participants’ automatic predictions. Another example of distributed annotation is the Amazon Mechanical Turk, which pays users to manually annotate tasks. Some authors developed games [24-26] or mobile apps [27] to gather human annotation.

Regarding crowdsourcing of translations, Zaidan and Callison-Burch [28] state that collecting translations by crowdsourcing using non-professionals may lead to low-quality results. They propose the use of distance among translations and LM perplexity to score collected translations to discriminate between “good” and “bad” translations.

Ambati et al. [29] explored the challenges involved in crowdsourcing translation based on their experiments with Amazon Mechanical Turk. Their main findings regarding challenges are related to the large label space, that is, even though there is a finite number of possible translations for a single translation, there is a much larger space of acceptable sentences in the target space, but that may not be adequate or not style compliant. The second one is the small number of bilingual speakers for low-resourced languages. The third one is low quality, as most of the crowd-sourced translators are not professional linguists. Given this scenario, they proposed a framework based on phases to enhance the final quality of crowd-sourced translations. The first step of the translation is done by weak bilingual translators, translations which are revised by bilingual translators and the final step is done by monolinguals of the target language or bilinguals whose mother tongue is the target one. Considering the potential of crowd-sourced annotation, we aimed at developing a prototype of a system to enable the manual evaluation of automatic translations tailored to biomedical texts and post-edition. Our goal was to produce a simple yet usable interface to annotate translations as valid in the target language, while enabling users to make adjustments in the translation to correct possible mistakes.

**Design**

When idealizing such a tool, we envisioned not to provide perfect and fluent translations, since that would require a considerable effort from users. We are rather interested in finding gross and dangerous MT mistakes, the ones that could completely hinder the interpretation of the article. That is, we are interested in assuring that the translated text conveys the same original message, even though it may not sound completely fluent for a native speaker.

We can see as an example the sentence “Nehmen Sie jeden Tag zwei Tabletten ein, es sei denn, Ihnen ist schwindelig oder benommen” in German. The direct translation, as seen in Supplementary Table 1, is “Take two pills every day by mouth unless you feel dizzy or lightheaded.” This may not sound natural, but it conveys the message that the dosage is two pills with a daily frequency and the contra-indication is if the person feels dizzy or lightheaded.

**Functionalities**

The following functionalities were implemented:

- Parallel visualization of the original text and the machine translated version.
- A “voting” system that allows users to flag a particular translation as correct (similar to a “like” in social media).
- An option to edit a suggested translation, allowing users to correct possible mistranslations.
- Only the last translation is available, since this is deemed to be
the one with best quality.

- When editing a translation, a terminology lookup is available. That is, for each matched string in the source text, the suggested translation is shown.

**Technical details**

In our prototype we aim at providing a simple and easily upgradable interface for document validation and modification. The prototype is coded in Python 3 using the Flask microframework. Our choice of Flask is due to its simplicity regarding back-end and front-end, while being able to scale if required.

For the interface, we opted for the Bootstrap library (https://getbootstrap.com/), since it provides responsive mobile-ready frontend components. The functionalities were expanded using jQuery and Javascript.

As for the backend, we took advantage of the SQLAlchemy toolkit (https://www.sqlalchemy.org/), which is an ORM (Object Relational Mapper) that abstracts database operations. By using SQLAlchemy, we were able to make the app database agnostic. That is, the user can easily switch among the RDBMS supported by the package without needing to change several parts in the code.

Regarding the translation system behind the prototype, we used an in-house model developed with OpenNMT (https://opennmt.net/) which is decoupled from the interface. We do not think that at this point it is extremely relevant to have an online translation system, since new articles can be batch translated overnight, for instance.

For the dictionary, we encourage the usage of UMLS, since it is a very comprehensive asset, already standardized and is available in many languages. Users can also make use of SNOMED CT available in more than one language, when compatible with licensing.

**Results**

We implemented our prototype following the design specified in Section 3. For such, we first created a simple interface to visualize the translated content in the source language (e.g., English in this case) and target language (e.g., French). In this first screen, bilingual users can check the translation, which is shown in column format. We also introduced a feature that allows users to hover over the source or target sentence and check which sentence it refers to on the other column of the parallel text. After checking the translation, bilingual users can flag (i.e., Like) the translation as good, or perform modifications (editing).

In Fig. 1, we show a screenshot of the article validation step. We have already included placeholders in the top bar to allow inclusion of alternative MT models as well as access to an Administrator backend which is under development.

In Fig. 2, we included a screenshot of the edition mode for the translated contents. In this view, the text is shown by sentences, with translations displayed as text boxes, such that users can perform post-edition on the suggested text. In addition, we included a glossary functionality, which can help users to guarantee terminology consistency. For this, a dictionary has to be supplied beforehand, and then a simple string matching is used to show the suggested translation. For instance, for the term "estrogen receptors", the suggested translation in French is "Récepteur des œstrogènes", while the automatic translation is "récepteurs aux œstrogènes". Although the automatic translation is not wrong, the suggested term "Récepteur des œstrogènes" is flagged in UMLS (https://www.nlm.nih.gov/research/umls/index.html, Unified Medical Language System) as preferred.

**Conclusion and Further Steps**

In this article, we pointed out the importance of making biomedical literature accessible to all healthcare professionals, despite the language they speak. As scientific publication, especially in biomedical sciences, has been fastly growing, manual translation of articles is an untractable approach to make such information multilingual. Thus, we argue that MT can be an alternative to alleviate such bottleneck.

However, despite the increasing performance of MT systems, some critical errors may occur when texts are translated, which can ultimately hinder patient safety. Thus, manual validation/evaluation of translations should be performed to mitigate potential risks. To enable validation to scale to several languages, we point out that crowdsourcing the effort may be a solution. Therefore, we developed a prototype of a system that can allow an easy translation validation and possible edition.

The prototype was developed using Python 3 and Flask (https://flask.palletsprojects.com/en/1.1.x/), with Bootstrap for the visual interface. A visualization and edition interface was created, and an Administrator interface is currently under development. We included visual features to help users when doing the validation or editing the text.

As future steps, we envision some important upgrades:

- Ability to export translations into TMX and TXML formats, since they are standard in the localization industry;
- Ability to flag different unit of measurements in translation (e.g., pounds to kilograms), since the numbers need to be converted accordingly.
Fig. 1. Interface for translation evaluation. Users can flag the translation as adequate (i.e., Like) or edit the proposed translation using the links in the Actions column.

Fig. 2. Interface for translation editions. Users can edit the proposed translation to make corrections on mistranslations or terminology adequacy. The prototype also shows suggested translations from terms matched in a dictionary, aiming at providing terminology consistency.
Include a voting scheme for rollback of manual edits and a “annotation” weight according to the mother tongue of the annotator. In addition, a similar approach for quality assurance as proposed by [29] could be used, by establishing a score for annotators as well as for annotations;

Develop an additional view to allow annotation transfer between source and target languages.

The last upgrade, related to annotation transfer, can be extremely helpful to create multilingual annotated datasets by leveraging existing annotations in one language. For instance, one could use annotations already made in a document in English to transfer those annotations to a translated text, making annotation quicker and less expensive.

ORCID

Felipe Soares: https://orcid.org/0000-0002-2837-1853
Rozane Rebechi: https://orcid.org/0000-0002-1878-7548
Mark Stevenson: https://orcid.org/0000-0002-9483-6006

Authors’ Contribution

Conceptualization: FS, RR. Data curation: RR. Formal analysis: FS. Funding acquisition: MS. Methodology: FS. Writing – original draft: FS, RR, MS. Writing – review & editing: MS.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

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Supplementary Materials

Supplementary data including one table can be found with this article online at http://www.genominfo.org.

References


open-japanese-mesh: assigning MeSH UIDs to Japanese medical terms via open Japanese–English glossaries

Ryota Yamada¹, Yuka Tateisi²

¹Fuku Inc., Tokyo 113-0033, Japan
²National Bioscience Database Center, Japan Science and Technology Agency, Tokyo 102-8666, Japan

The Medical Subject Headings (MeSH)thesaurus is a controlled vocabulary for indexing biomedical documents that is used for document retrieval and other natural language processing purposes. However, although the original English MeSH is freely available, its Japanese translation has a restricted license. We attempted to create an open alternative, and for this purpose we made a script for assigning MeSH UIDs to Japanese medical terms using Japanese–English glossaries. From the MeSpEn glossary and MEDUTX dictionary, we generated a 12,457-word Japanese–MeSH dictionary.

Keywords: dictionaries, Medical Subject Headings, natural language processing
Availability: The script is available from before the URL https://github.com/roy29fuku/open-japanese-mesh.

Introduction

The Medical Subject Headings (MeSH) [1] thesaurus is a controlled vocabulary developed and maintained by the United National Library of Medicine (NLM) that is used for indexing biomedical articles in PubMed.

MeSH is primarily used for indexing and searching the PubMed database, but it can also be used as a reliable dictionary of technical terms in the biomedical domain, as its headings and entry terms are representations of biomedical concepts approved by the NLM. Thus, MeSH is a valuable resource for natural language processing (NLP) applications. The metathesaurus in the Unified Medical Language Systems (UMLS) [2] includes translations of MeSH to several languages including Japanese.

However, although the original MeSH in English can be freely downloaded and used, the translations of MeSH in the UMLS are provided with “category 3” restrictions, which means that they cannot be incorporated into applications available outside the institution of the licensee. According to a mini-survey conducted in the 5th Biomedical Linked Annotation Hackathon (BLAH5) [3], although there are web-based dictionaries/thesauri that are freely consulted for finding MeSH UIDs or tree numbers by human readers, no dictionaries that are completely free for NLP applications are available.

Methods

MeSH consists of three types of records: descriptors (main headings), qualifiers, and supplementary concept records.
Descriptors are terms that characterize the subject matter. They are organized in a hierarchical structure based on broader/narrower relations of concepts. Qualifiers are used with descriptors and describe an aspect of a subject denoted by the descriptor. Supplementary concept records are names of chemicals, drugs, and new concepts. Supplementary concept records are not hierarchically ordered. Instead, each supplementary concept is linked to one or more descriptors. Descriptors and supplementary concepts have a heading (representative term) and entry terms (synonyms). Each record in MeSH is accompanied by an identifier (UID).

In order to link Japanese medical terms with medical concepts in MeSH, we used two open Japanese-English bilingual glossaries. The MeSpEn English-Japanese glossary is part of the MeSpEn multilingual medical glossary developed by the Text Mining Unit (TEMU) of the Barcelona Computing Center and available under a Creative Commons Attribution 4.0 International License [4]. The MEDUTX dictionary was developed by Kitasato University and is licensed under a Creative Commons Attribution 3.0 International License [5]. The MeSpEn English-Japanese glossary has 16,756 unique Japanese terms and 10,738 unique English terms (27,668 unique pairs). The MEDUTX dictionary has 21,821 unique Japanese terms and 22,276 unique English terms (27,122 unique pairs). Merging the two dictionaries yielded a resource with 35,903 unique Japanese terms and 30,853 unique English terms (54,790 unique pairs).

We used the 2020 MeSH ASCII files for descriptors (d2020.bin) and supplementary concepts (c2020.bin) downloaded from the FTP site of the NLM on February 5, 2020. The descriptors file contained 242,205 terms (headings and entry terms) that were mapped to 29,640 concepts (UIDs) and the supplementary concepts file had 649,322 terms that were mapped onto 268,825 UIDs.

Since the Japanese-English dictionaries we used were much smaller than the MeSH vocabulary, we developed a Python script that can be applied to any Japanese-English glossary (in the form of tab-separated list of Japanese terms and corresponding English terms) and assigned the UIDs to Japanese terms where applicable, in order to be able to expand the output dictionary when more Japanese-English resources are available.

We mapped Japanese medical terms to UIDs in the process illustrated in Fig. 1. First, a Japanese term was mapped to English term(s) with Japanese-English dictionary. The English terms were normalized as follows: they were placed in lowercase, zenkaku (full-width, non-ASCII) characters were converted to their hanka-ku (half-width, ASCII) counterparts, Greek characters were spelled out, and Roman numerals were converted into Arabic numerals. The jaconv library [6] was used for zenkaku-to-hankaku normalization. The MeSH terms were also normalized, and the normalized English terms from the dictionary were matched against the normalized MeSH terms.

The Python class for Japanese-English dictionaries, MeSH data, and normalization rules were defined in order to easily incorporate new dictionaries and new normalization rules. We also investigated the effect of each type of normalization.

### Results and Discussion

Without normalization of English terms, 2,838 out of 35,903 Japanese terms were mapped onto MeSH concepts (UIDs). With normalization of English terms, 12,457 Japanese terms out of 35,903 (about 34.7%) were mapped to UIDs. The contributions of each type of normalization are summarized in Table 1. The results show that case matching of the alphabet was the most effective normalization step, and the contributions of other types of normalization were small.

At least one Japanese term was assigned to 7,346 out of 298,465

### Table 1. Number of terms successfully assigned MeSH UIDs according to normalization

<table>
<thead>
<tr>
<th>Normalization</th>
<th>Example</th>
<th>Mapped Japanese terms</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>A → a</td>
<td>2,838</td>
</tr>
<tr>
<td>Lowercasing</td>
<td>A → a</td>
<td>12,406</td>
</tr>
<tr>
<td>Zenkaku-to-hankaku</td>
<td>A (uFF21) → A (u0041)</td>
<td>2,839</td>
</tr>
<tr>
<td>Greek-to-English</td>
<td>a → alpha</td>
<td>2,857</td>
</tr>
<tr>
<td>Roman numerals-to-Arabic</td>
<td>VIII → 8</td>
<td>2,838</td>
</tr>
<tr>
<td>All</td>
<td></td>
<td>12,457</td>
</tr>
</tbody>
</table>

![Fig. 1. The UID assignment process.](https://doi.org/10.5808/GI.2020.18.2.e22)
MeSH concepts (UIDs), of which 6,185 were descriptors and 1,161 were supplementary concepts. This means that Japanese terms were assigned to about 20.9% (6,185/29,640) of descriptors and 0.4% (1161/268,825) of supplementary concepts. Considering the size of the Japanese-English dictionary (about 3% of the MeSH vocabulary) this result seems reasonable. For improving its coverage, a list of translations of names of chemicals, drugs, and other named entities regarded as supplementary concepts in MeSH should be obtained.

**Conclusion**

We made a script for assigning MeSH UIDs to Japanese medical terms using Japanese-English glossaries. From the MeSpEn glossary and MEDUTX dictionary, we obtained a 12,457-word Japanese-MeSH dictionary. This dictionary could be enhanced by using additional Japanese-English dictionaries. The script is available from https://github.com/roy29fuku/open-japanese-mesh under the Creative Commons Attribution 4.0 International License. Our future work includes a comparison with the Japanese translations in the UMLS metathesaurus.

**ORCID**

Ryota Yamada: https://orcid.org/0000-0003-2237-5025  
Yuka Tateisi: https://orcid.org/0000-0002-3813-5782

**Authors' Contribution**

Conceptualization: YT. Formal analysis: YT, RY. Methodology: RY. Writing – original draft: YT. Writing – review & editing: RY, YT.

**Conflicts of Interest**

No potential conflict of interest relevant to this article was reported.

**References**

Choosing preferable labels for the Japanese translation of the Human Phenotype Ontology

Kota Ninomiya¹,²*, Terue Takatsuki³, Tatsuya Kushida⁴,⁵, Yasunori Yamamoto³, Soichi Ogishima⁶

¹National Institute of Public Health, Wako 351-0197, Japan
²Social Cooperation Program of IT Healthcare, Graduate School of Pharmaceutical Sciences, The University of Tokyo, Tokyo 113-0033, Japan
³Database Center for Life Science, Research Organization of Information and Systems, Kashiwa 277-0871, Japan
⁴BioResource Research Center, RIKEN, Tsukuba 305-0074, Japan
⁵National Bioscience Database Center, Japan Science and Technology Agency, Tokyo 102-8666, Japan
⁶Advanced Research Center for Innovations in Next-Generation Medicine, Tohoku University, Sendai 980-8573, Japan

Introduction

The Human Phenotype Ontology (HPO) [1] is the de facto standard ontology to describe human phenotypes in detail, and it is actively used, particularly in the field of rare disease diagnoses. For clinicians who are not fluent in English, the HPO has been translated into many languages, and there have been four initiatives to develop Japanese translations. At the Biomedical Linked Annotation Hackathon 6 (BLAH6), a rule–based approach was attempted to determine the preferable Japanese translation for each HPO term among the candidates developed by the four approaches. The relationship between the HPO and Mammalian Phenotype translations was also investigated, with the eventual goal of harmonizing the two translations to facilitate phenotype–based comparisons of species in Japanese through cross-species phenotype matching. In order to deal with the increase in the number of HPO terms and the need for manual curation, it would be useful to have a dictionary containing word–by-word correspondences and fixed translation phrases for English word order. These considerations seem applicable to HPO localization into other languages.

Keywords: biological ontologies, natural language processing, phenotype, rare diseases, translations

Availability: As these new translations are still in progress, only an old version of Japanese translations can be obtained from https://github.com/ogishima/HPO-japanese.
HPO term. At the Biomedical Linked Annotation Hackathon 6 (BLAH6), an attempt was made to select preferable unique Japanese terms.

The HPO has mainly been used in the field of rare diseases as the most comprehensive resource for deep phenotyping, which is defined as "the precise and comprehensive analysis of phenotypic abnormalities in which the individual components of the phenotype are observed and described" [4]. As approximately 80% of rare diseases, the number of which is estimated to be between 5,000 and 8,000, are thought to be genetic [5,6], they may occur anywhere.

For individuals with rare diseases, delays in diagnoses and frequent misdiagnoses lead to irreversible disease progression, and mistreatment based on a misdiagnosis can even harm patients in some circumstances. This problematic journey faced by patients with rare diseases is sometimes called the "diagnostic odyssey." It has been reported that it takes 5–7 years on average for patients with rare diseases to be diagnosed correctly in the UK and USA, and that patients received incorrect diagnoses two or three times [7]. In Japan, the average time to be diagnosed correctly with Fabry disease was found to be about 20 years [8].

Therefore, HPO localization is expected to help clinicians who are not fluent in English make early diagnoses based on medical records containing standardized and detailed phenotypic information. HPO terms are being translated into Japanese, French, German, Russian, Turkish, Spanish, Italian, Dutch, Portuguese, and Chinese.

In order to understand the pathology of a specific disease, researchers often use model animals that present the same symptoms or have the same genetic abnormalities. When they choose the appropriate model animals, standardized phenotyping can be a critical clue. In Exomiser [9], phenotypic data from several species, such as mice and zebrafish, are also used for functional annotation of genetic variants from human whole-genome sequencing data. The standardized description of phenotypes by the HPO and other phenotype ontologies has enabled a phenotype-based comparison of species through cross-species phenotype matching. Harmonization of translations is also expected to make it possible for researchers to search for bio-resources for human beings or other species only using the same terms in Japanese.

At BH15, which was held in 2015, HPO terms started to be translated into Japanese. As a result of the hackathon and subsequent efforts, each HPO term had four Japanese equivalent terms, which were translated using different English-Japanese dictionaries, and the translations have been made available to the public [10].

One of the four translations is based on the Life Science Dictionary (LSD) [11], which is an English-Japanese dictionary for the life sciences; this translation is updated by researchers at Kyoto University. The second translation is based on the Japanese translation of the Mammalian Phenotype (MP) ontology [12], and was created by Riken BioResource Research Center. The third translation was created by Kenji Naritomi, a medical expert who has translated many materials about genetic diseases into Japanese. He translated the HPO terms to the extent that he could. The last translation is an automatic translation using Google Translate.

At BLAH6, a unique Japanese translation for each English term in the four translations was selected through trial and error based on the criterion that translated terms should not sound anomalous or unnatural in Japanese. Translations were prepared for the 10,668 HPO terms as of October 2017.

As the HPO describes the phenotypes of human beings and the MP describes those of mammals, they have many concepts in common. The equivalence of their concepts has already been explored by Mungall [13]. At BLAH6, the relationship between the Japanese translation of the HPO and that of the MP was examined with the goal of harmonizing them so that researchers could easily search for biological resources, using the same expression for the same phenotype. In this comparison, the Japanese translations made by Kenji Naritomi were adopted as the counterparts of the MP Japanese terms.

**Methods**

First, a rule-based method was used to choose the most appropriate translated terms in the following order.

1. If two or more translated terms were the same among the four translations, they were chosen as a unique Japanese term. If there were two sets of words, such that two of the four translations were the same, and the other two were the same, they were labeled as "two appropriate candidates determined by a majority." The rest of these cases were labeled as "a unique Japanese term" and "determined by a majority."

2. For the rest of the HPO terms, a morphological analysis was conducted using Mecab [14], with the MANBYO dictionary [15] as a user dictionary, for all Japanese translation candidates except those based on the LSD. Then, candidates for preferable labels were automatically chosen based on whether the morphological analysis indicated that the terms included anomalous features, defined as below. The MANBYO dictionary contains a large number of medical terms in Japanese. As some of the terms derived from the LSD are combinations of translated words, they were excluded from this analysis.

As no consensus necessarily exists regarding the precise defini-
tion of “anomalous” features, the terms were separately labeled with each feature to make it possible to change the criteria used to identify anomalous terms. The features of anomalous terms were as follows:

1. Terms including verbs or ending with a non-noun word (e.g., 出生時にみられ時間とともに真っすぐなる大腿骨湾曲). These features seem anomalous because HPO terms are supposed to be nouns, and it is preferable for combinations to only involve nouns.

2. Terms including particles or adjectival verbs (e.g., 尺骨の有力な茎状突起), for the same reason as (1).

3. Terms including adjectives (e.g., 幅広い長管骨), also for the same reason as (1).

4. Terms including Japanese commas, which appear much more unusual than English commas when they are used in terms (e.g., 異所性心臓、心臓転位).

5. Terms including untranslated English words (e.g., 角膜stromal浮腫).

6. Cases where English terms were not translated at all for unknown reasons, and the translated terms were blank.

In this analysis, all the anomalous features were adopted. Candidate terms were ranked in the following order:

1. If a translated term included some strange features, it was excluded from consideration.

2. If only one term was left after the exclusion of anomalous terms, it was chosen as the most appropriate one. Such terms were labeled as “a unique Japanese term” and “determined by an exclusion process.”

3. If more than two terms were left, it was difficult to choose which was better, and such cases were labeled as “multiple appropriate candidates determined by an exclusion process.”

4. If all of the terms were excluded, the item was labeled as “no appropriate candidates determined by an exclusion process.” If all the translated terms were initially blank, they were labeled as “BLANK.”

Second, an attempt was made to find out how equivalent concepts between HPO and MP are described in English and Japanese to promote the consistency of translations between these resources. As the equivalence data only contain the IDs of concepts, the English and Japanese terms were collected using the Japanese translation of the HPO [10], with HPO data as of August 2015 and July 2016, and the relationship between HPO and MP was assessed based on the MP data as of October 2012.

Results

The results of labeling all the HPO terms are shown below (Table 1).

In the second phase, the relationships between HPO and MP concepts in Japanese and English were explored, and ways to harmonize the translations were examined. A flow chart is shown below (Fig. 1).

All the HPO and MP terms referring to the same concepts were divided into the four categories described in the flow chart. As the equivalence data were created after the first translation attempt in 2015, some terms had no Japanese translation candidates. The results of a character-string comparison between them are as follows (Table 2).

Discussion

In this trial, about half of the HPO terms were found to have a unique Japanese translation. However, there are three points to consider regarding these labels.

First, those labeled as “determined by a majority” sometimes included anomalous Japanese expressions, as terms were not excluded based on anomalous features if they were identical in a majority of sources (50 percent and more). Therefore, the order of assigning labels should perhaps be reconsidered.

Table 1. Summary table of the labels assigned to all the HPO terms

<table>
<thead>
<tr>
<th>Label</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>All the HPO terms</td>
<td>10,668</td>
</tr>
<tr>
<td>A unique Japanese term</td>
<td>5,678</td>
</tr>
<tr>
<td>Determined by a majority</td>
<td>3,096</td>
</tr>
<tr>
<td>Determined by an exclusion process</td>
<td>2,687</td>
</tr>
<tr>
<td>Two appropriate candidates determined by a majority</td>
<td>105</td>
</tr>
<tr>
<td>Multiple appropriate candidates determined by an exclusion process</td>
<td>2,165</td>
</tr>
<tr>
<td>No appropriate candidates determined by an exclusion process</td>
<td>2,720</td>
</tr>
<tr>
<td>BLANK</td>
<td>5</td>
</tr>
</tbody>
</table>

HPO, Human Phenotype Ontology.

Table 2. Summary table of the categories assigned to HPO/MP terms with the same concepts

<table>
<thead>
<tr>
<th>Category</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>All pairs of HPO/MP terms with the same concepts</td>
<td>1,442</td>
</tr>
<tr>
<td>Both words are the same in both languages</td>
<td>219</td>
</tr>
<tr>
<td>Only the English words are the same</td>
<td>420</td>
</tr>
<tr>
<td>Only the Japanese words are the same</td>
<td>115</td>
</tr>
<tr>
<td>Both words are different in both languages</td>
<td>688</td>
</tr>
<tr>
<td>Japanese translation candidates do not exist yet</td>
<td>128</td>
</tr>
</tbody>
</table>

HPO, Human Phenotype Ontology; MP, Mammalian Phenotype.
Second, HPO terms that had two appropriate candidates determined by a majority were divided into three groups, although they had the same problem as those labeled as “determined by a majority.” The first group included terms with only slight differences, such as whether or not they included “症,” which means “syndrome” in Japanese (e.g., 不眠症|不眠). Therefore, such typical and almost meaningless characters or words should be omitted as stop words in the next matching trial. The second group contained translations that had entirely different meanings (e.g., 硬化症|第1中足骨硬化症). In this case, one of the options must be a mis-translation. A possible reason for this is that some words in the terms were ignored in translation because the translation systems did not contain them in their dictionaries and could not recognize them properly. The last group required manual curation because the order and the selection of translated words were different (e.g., 髄様甲状腺癌|甲状腺髄様癌).

Finally, problems in Japanese translation labeling related to the exclusion process are mainly caused by the definition of anomalous features and the accuracy of the morphological analysis. Therefore, the definitions need to be made more sophisticated in future trials by adding or removing exclusion criteria. It is also important to choose an appropriate morphological analyzer for dealing with medical expressions, such as Juman++ [16] or Sudachi [17].

The relationship between the HPO and MP translations was classified into four categories according to character-string comparisons. First, if the English and Japanese terms are both the same, there is nothing to change. Second, if only the English terms are the same, the HPO translations take precedence over the MP translations, and the latter is unified to follow the former, as the former already seems to be used for more diverse purposes and to be more widespread. There are two reasons for inconsistencies in Japanese translations. One is the same as encountered for Japanese localized terms assigned the label “two appropriate candidates determined by a majority.” The other is that the same terms, especially those that refer to morphological abnormalities of external body parts (instead of abnormal internal situations), are sometimes translated differently depending on the species. For example, the words “male” and “female” are “男性” and “女性” for human beings, respectively. However, for non-human mammals such as mice and rats, these terms are written as “オス” and “メス”, respectively. Therefore, the principle of assigning precedence to the HPO translations is acceptable only in a general sense. Third, if only the Japanese terms are the same, there is no need to change the translation as long as the concepts are similar between the HPO and MP terms. Finally, if both the English and Japanese terms are different, there is no option other than manual curation. Since applying these principles led to the finding that roughly half of the terms need manual curation to be harmonized, another way needs to be found to decrease the necessity for manual curation in further research.

As the HPO includes technical terms, orthodox translations that are generally accepted among health professionals should be adopted. An excellent approach would seem to be to map these terms to other dictionaries for translation and to adapt their translations if doing so is permissible because other dictionaries are thought to be edited according to the same policy. This approach
seems to contribute to external consistency among dictionaries and to reinforce the stability of orthodox translations. Nonetheless, the MP translations can be candidates for replacing the HPO translations, as they sometimes contain better expressions, and a comparison between them enables harmonization and cross-species matching or searching. If translations of the terms cannot be found in other resources, or there are several translation candidates, experts need to translate them manually. Although this task requires extensive work and costs, it is ultimately unavoidable.

To deal with the increase of the number of HPO terms and the excessive dependence on manual curation—despite its inevitability in principle—it may be a good idea to develop a dictionary that contains word-by-word correspondences based on the temporarily completed translations of the HPO and MP. Such a dictionary would enable the generation of translation candidates for new terms consistent with the fixed HPO and MP translations created previously. As some word orders are common in English terms, it is possible to establish fixed Japanese phrases for each of these frequent word orders. Therefore, dictionaries and lists of fixed phrases can reduce the task of manual curation by changing it from translation of terms from scratch to only selection of the most appropriate candidates. These approaches seem to be applicable to HPO localization into other languages.

Conclusion

In this study, an attempt was made to determine a single unique translation for each term in the HPO in a rule-based way. For about half of the terms, only one appropriate Japanese word was identified, and for the rest, manual curation was needed. However, as this approach yielded insufficient accuracy, further consideration is necessary and will be given in venues such as another future hackathon.

The relationship between the HPO and MP was also investigated to evaluate the task of establishing consistency between them. Based on the analysis, the translations of both ontologies should be harmonized to improve their usability for annotating phenotypes of humans and non-human mammals.

It is possible that the number of HPO terms will continue to increase and that there will be more need for manual curation. An effective approach would seem to be to create a dictionary that contains word-by-word correspondences based on the temporary translations and fixed translation phrases for English terms in word orders that frequently appear. These approaches are most likely applicable to HPO localization into other languages.

ORCID

Kota Ninomiya: https://orcid.org/0000-0002-7381-1643
Terue Takatsuki: https://orcid.org/0000-0003-0011-764X
Tatsuya Kushida: https://orcid.org/0000-0002-0784-4113
Yasunori Yamamoto: https://orcid.org/0000-0002-6943-6887
Soichi Ogishima: https://orcid.org/0000-0001-8613-2562

Authors’ Contribution

Conceptualization: KN, TT, TK, YY, SO. Data curation: KN. Formal analysis: KN. Funding acquisition: YY, SO. Methodology: KN, TT, TK. Writing – original draft: KN, TT, TK. Writing – review & editing: KN, TT, TK.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

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References


Introduction

Twitter is one of the leading social media platforms with more than 126 million daily users [1]. Twitter is now regarded by the natural language processing (NLP) community as a valuable source of information and has been the focus of a significant amount of research this last decade. An increasing number of shared-tasks have been organized utilizing data from this platform. Amongst the shared tasks for Twitter data, named entity recognition is well-represented, including the Named Entity Recognition and Linking Challenge series [2] which ran from 2013 to 2016, or the Workshop on Noisy User-generated Text series [3] which organized shared tasks from 2015 to 2017. Aside from named entity recognition, the community has extended its use of Twitter to broader tasks, such as the SemEval tracks on sentiment, opinion and abusive language classification starting in 2013 [4], or for health research with the Social Media Mining for Health (#SMM4H) running since 2016 [5]. Since more than half of tweets are not written in English, shared tasks are also utilizing corpora in various languages: the conference sur l’ Apprentissage Automatique in 2017 in French [6], the Forum for Information Retrieval in 2016 in Indian [7], the Named Entity rEcognition and Linking in 2016 in Italian, a track in Arabic during SemEval 2017 and #SMM4H’20 with a task in French and Russian.

As the foundation for most shared tasks in NLP, and more generally most studies in NLP, the importance of the corpus cannot be overstated. A standardized corpus is essential for the evaluation of the competing systems. The correctness and consistency of the annotations are vital to ensure accurate results and predictions on how the systems will perform on unseen data. Moreover, with the generalization of statistical methods in NLP, annotations are also important for training the systems. Only well-defined, high-quality annotations can ensure that a machine learning-based system will be able to model dis-
Prior to the publication of an extensive review of 78 annotation tools by Neve and Seva [12], we started a review of annotation tools for Twitter data. The inclusion criteria for our review were the availability and the ease of installation of the tools, or otherwise, a demonstration of the tool online. A tool was not easily installed when dependencies were missing, errors occurred, or external software, such as databases, needed. Among the 19 annotation tools we tested, few met the requirements we needed to perform timeline annotations. We had used the brat annotation tool for a previous project involving the annotation of PubMed Central articles; however, we found several problems with it when trying to use it for timeline annotations. Mainly, brat’s user interface was not adapted to annotate adjacent tweets. We reviewed a commercial application, LightTag [13], and though it provided a clean interface and supported many of our requirements, it crashed excessively during use. It also did not allow for subcategories of entity tags and, the tool not being open source, prevented us from modifying it to fit our needs. Other tools tested did not allow for the subcategorization of entity tags, including WebAnno [14], Yedda [15], and Slate [16]. These tools also did not provide support for the normalization of entities extracted. Supplementary Table 1 summarizes our review of the 19 tools. Our review found three possible annotation tools for our project eHost [17], GATE and TextAE, as they met most of our requirements. We chose the GATE and TextAE annotation tools for the hackathon because they were actively supported and updated regularly.

Annotating Twitter Data

When registering for a Twitter account, a user is invited to fill a short description and choose other users to follow. The new user is assigned a unique user ID and each tweet posted by the user is identified by a unique tweet ID. In addition, each tweet is described by metadata such as the posting time or the predicted language of the tweet. The collection of all tweets posted by a given user is called the home timeline.

The four researchers participating in our project during the hackathon were provided with 25 timelines of women that had publicly announced their pregnancy on Twitter. These timelines correspond to a total of 74,016 tweets in English, with an average of 3,000 tweets per timeline. We defined 31 annotation types relevant to these pregnancies and manually pre-annotated the 25 timelines for the event.

With no annotation tool designed for Twitter timelines, we had to adapt an existing tool for this type of data. Before the hackathon, we listed a set of requirements a tool should fulfill to be usable with Twitter timelines and we asked our four participants to evaluate two annotation tools according to those specifications. The specifications are detailed in (Table 1).

Adapting Existing Annotation Tools for Twitter

Prior to the publication of an extensive review of 78 annotation tools by Neve and Seva [12], we started a review of annotation tools for Twitter data. The inclusion criteria for our review were the availability and the ease of installation of the tools, or otherwise, a demonstration of the tool online. A tool was not easily installed when dependencies were missing, errors occurred, or external software, such as databases, needed. Among the 19 annotation tools we tested, few met the requirements we needed to perform timeline annotations. We had used the brat annotation tool for a previous project involving the annotation of PubMed Central articles; however, we found several problems with it when trying to use it for timeline annotations. Mainly, brat’s user interface was not adapted to annotate adjacent tweets. We reviewed a commercial application, LightTag [13], and though it provided a clean interface and supported many of our requirements, it crashed excessively during use. It also did not allow for subcategories of entity tags and, the tool not being open source, prevented us from modifying it to fit our needs. Other tools tested did not allow for the subcategorization of entity tags, including WebAnno [14], Yedda [15], and Slate [16]. These tools also did not provide support for the normalization of entities extracted. Supplementary Table 1 and 2 summarizes our review of the 19 tools. Our review found three possible annotation tools for our project eHost [17], GATE and TextAE, as they met most of our requirements. We chose the GATE and TextAE annotation tools for the hackathon because they were actively supported and updated regularly.

Tuning Gate for Twitter Data Annotation

GATE is an open-source toolkit developed for text annotation and automatic text processing. We used the stand-alone version of GATE to annotate Twitter timelines for prior projects [18]. Although a web-based version of GATE is available, GATE teamware [19], we compared TextAE with the stand-alone version of GATE, as we were already familiar with the tool and it was easier to install during the hackathon than the web-based version.

During the hackathon, we imported our 25 timelines and reviewed the tools with respect to our requirements. We imported a timeline as a unique document in GATE, one tweet per line. We inserted the tweet IDs and the posting dates before the text of the tweets to facilitate the annotation process, all items were separated by tabulations. Tweet IDs and dates were pre-annotated with their tags in the document. We named the file with the user ID. We could have added annotations at the timeline level (metadata), such as the gender or the place of residence of the user, by importing them as pre-annotation and inserting them at the beginning of the document in an empty span.

GATE fulfilled many of our specifications. GATE is actively
The annotation tool should be web-based to support multiple annotators and to enable inter-annotator agreement calculation and disagreement resolution. Web-based tools, such as GATE teamware or brat, make it easier to manage a team of annotators and compute the inter-annotator agreement.

It should be easy to install, to set up the tags and the annotation schema as well as allowing changes to the schema. Twitter data are used for various research projects, each project mining for different types of information requiring their own annotation schemas (e.g., normalizing adverse drug reaction (ADR), extracting reasons of drug non-persistence, etc.)

It should load the tweets composing a timeline in less than 2 seconds and load an external dictionary for normalizing an annotation in less than 3 seconds. A dictionary may be opened several times per tweet to normalize annotations, such as ADRs. A reading time longer than 3 seconds may significantly slow down the annotation of large corpora.

It should not present recurrent bugs preventing or modifying the annotation process. The tool should be actively supported. Active support would ensure the correction of such bugs.

It should periodically save the annotated document and save automatically upon closing the document or, in the absence of automatic saving, warn the annotators to save before closing. When annotating long documents such as timelines, annotators are likely to close a document without saving, losing their annotations.

It should allow the upload of pre-annotated labels and metadata [e.g., tweet IDs or date of post]. The import formats should be standard like XML or JSON. Non-standard formats, such as the XML format used in GATE, required developing conversion scripts to process new corpora.

It should store the annotations in a separate file, leaving the original document intact. Stand-off annotations are preferred because corpora may be used for different projects (e.g., timelines collected to study adverse pregnancy outcomes reused to study topics discussed during pregnancy).

It should allow for nested and crossing annotations. Two annotations are nested if the span of one annotation is included in the span of the second annotation; they cross if they share a common span of text.

It should allow for annotating various levels of a timeline, the timeline itself, and the network of a Twitter user. These levels are annotating spans of a tweet (e.g., the name of a drug), the tweet itself (e.g., the sentiment of the tweet), continuous set of tweets, i.e., an annotation spanning over multiple and adjacent tweets (e.g., all tweets posted by a user in May 2016).

The interface should present a timeline to the annotator in a way that all annotations are easily distinguishable from each other and from the span annotated. Annotations should appear above the span annotated. The metadata should be included in the annotation file but not visible in the timeline during annotation. Most research projects involve annotating multiple types of annotations, e.g., annotating a drug name and annotating if the drug was taken. Annotations are likely to overlap, cluttering up the document without a well-designed user interface.

It should support for defined entity tags to have assignable subcategories. For example, annotating alcohol intake, subcategories could be: intake, possible_intake, no_intake.

It should support the inclusion of a dictionary or ontology for normalizing the annotated entities to standardized terms. For example, normalizing the annotated span ‘sleepy’ by linking it to the MedDRA preferred term ‘Somnolence’.

It should provide a default API to plug in an external classifier implementing an active learning algorithm to assist the annotation process. The classifier could, for example, pre-annotate the sentiments of tweets. Using active learning, it can ask an annotator to correct the labels it assigned with less certainty and retrain its model after the labels are corrected. After some iterations, the classifier should annotate most of the tweets with the correct sentiments, saving manual annotation time compared to manually annotating all tweets [20].

It should calculate the inter-annotator agreement and provide an interface to help adjudication. It should allow annotators to edit these annotations. Due to the time constraint, we did not evaluate the diff tool plugin [21] to compute the inter-annotator agreement in GATE. The format of the output also made it difficult to manually perform these two tasks.

Supported, with its most recent release occurring on January 17, 2020. Written in Java and well documented, it is easy to setup. Pre-annotations and metadata can be imported provided that they are formatted in an XML file following a format specific to GATE. This XML format has been designed to support both nested and crossing annotations. GATE also supports subcategory annotations. Despite the large number of tweets in a timeline, GATE loads a timeline and its annotations in less than a second. It clearly marks completed annotations in the interface and offers the possibility to hide annotations when appropriate. GATE implements interfaces for active learning, but we did not use the service during the hackathon due to lack of time.

GATE appeared to be a valuable tool for annotating timelines but several drawbacks discourage us from using the tool for long-term projects. There are some issues with the stability of the stand-alone version as GATE would crash occasionally. GATE also allows closing a file without saving the annotations and without warning the annotators. The internal XML format, specific to GATE, was difficult to work with and required the development of scripts to convert pre-annotated timelines in order to import them in GATE as well as to export the timeline annotated for further use in external applications. Whereas annotations at the tweet level were well supported, annotating timelines was only possible as pre-annotation and the built-in GATE User Interface would not allow annotators to edit these annotations. Due to the time constraint, we did not evaluate the diff tool plugin [21] to compute the inter-annotator agreement in GATE. The format of the output also made it difficult to manually perform these two tasks.
Tuning PubAnnotation/TextAE for Twitter Data Annotation

TextAE is a web-based interface designed for corpus annotation. The interface is integrated with PubAnnotation [22], a public repository for literature annotation. For the hackathon, we chose the public version of PubAnnotation to create a private project, eliminating the need of a local installation and enabling the storage of our data in the cloud. We imported 5 timelines, representing the timelines in the same way as we did in GATE, one timeline per document, one tweet per paragraph, the document named with the user ID, and tweet IDs/posting dates inserted before the texts of the tweets.

The current versions of PubAnnotation/TextAE do not meet some of our requirements and would be too limited for our usage. However, the tools are still under development and, with the improvements scheduled, they could become standard for annotating Twitter data. An annotation project can be set up in PubAnnotation for multiple annotators as a collection, with a project created for each annotator. Annotation tags are created using a JSON configuration file. The TextAE interface allowed each tweet to be loaded as a paragraph. The annotation interface was not intuitive for all users. However, with documentation online, most annotators were readily able to access text and begin annotating within an hour. Some choices in the ergonomic design of the annotation interface were not optimal for our task and added time to the annotation process. The interface displays the annotated texts with the labels appearing on top of the text. The tool supports nested annotations. Although it is possible to add multiple annotations on the same span of text, this functionality was unstable in the version evaluated. Annotating the span in the wrong order resulted in the loss of the top-level annotation. The tool does not support crossing annotations since it uses HTML to display annotations and HTML does not allow crossing tags. TextAE could annotate a continuous set of tweets but this will require minor changes in the tool. TextAE does not currently support timeline annotations, but plans were made during the hackathon to extend the interface to add and edit this level of annotations. TextAE, combined with PubDictionaries, allows subcategories and normalization of annotations to standardized terms. Despite the size of our timelines (3,828 tweets, 418 KB, on average) and the dictionary used for testing the normalization (two million entries, 132 MB), both tools reacted within the time constraints imposed by our requirements. TextAE also provides an interface for the comparison of documents annotated by multiple annotators for disagreement resolution. TextAE was stable when annotating our timelines and, although the annotations must be manually saved, there is a warning presented to the annotator before closing. Annotations are saved in a separate file that can be exported as JSON or TSV files. Given the time limit of the hackathon, we did not test the import functionality in JSON format. An active learning API for the tool is in development and was not ready during the event.

Conclusion

The need for annotation tools dedicated to Social Media data, such as Twitter, is becoming more apparent as the interest of the NLP community is growing for this data. Since, to the best of our knowledge, there is no annotation tool dedicated to Twitter available, we evaluated during the 6th edition of the Biomedical Linked Annotation Hackathon two generic annotation tools using 25 Twitter timelines as a way to test their functionalities. After defining a catalog of requirements for an annotation tool dedicated to Twitter, we reviewed 19 tools and selected GATE and TextAE/PubAnnotation for our evaluation. Our results show that, whereas neither of them allows the annotation of all information characterizing Twitter timelines, each may be adapted for this purpose, if annotators are willing to compromise on some functionalities.

ORCID

Davy Weissenbacher: https://orcid.org/0000-0001-8331-3675
Karen O’Connor: https://orcid.org/0000-0001-7709-3813
Aiko T. Hiraki: https://orcid.org/0000-0002-7866-286X
Jin-Dong Kim: https://orcid.org/0000-0002-8877-3248
Graciela Gonzalez-Hernandez: https://orcid.org/0000-0002-6416-9556

Authors’ Contribution

Conceptualization: DW, KO, JDK, GGH. Data curation: KO, ATH, DW. Formal analysis: KO, ATH, DW. Funding acquisition: GGH, JDK. Methodology: DW, KO, JDK, GGH. Writing – original draft: DW, KO. Writing – review & editing: DW, KO, GGH, JDK.

Conflicts of Interest

No potential conflict of interest relevant to this article was reported.

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**Supplementary Materials**

Supplementary data can be found with this article online at http://www.genominfo.org/.

**References**


Supplementary Table 1. Requirement review of annotation tools for Twitter

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<tr>
<th>Annotation Tool</th>
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NR = Not Reviewed

We were unable to run 3 of the 19 tools due to installation errors or dependency issues, Argo, Callisto, and Knowtator. Three others, Pubtator, BioQRator, and ezTag, required a specific input file format, such as BioC, and therefore were not suited to annotate tweets. In Supplementary Table 1, we summarize our review of the features of the remaining 13 annotation tools for the most important requirements in our catalog. Note, as we were just reviewing the features of the tools, we did not complete full installations for the tools with external dependencies such as server and/or database installations to run. For those tools, we examined the online demonstrations if available during our assessment.

*Our review was done based on the software demonstration.
*No demonstration available, our review was done based on the documentation.
Supplementary Table 2. Annotation tools reviewed

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- Conceptualization: AB
- Data curation: EFG
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- Funding acquisition: CD
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This journal follows the data sharing policy described in “Data Sharing Statements for Clinical Trials: A Requirement of the International Committee of Medical Journal Editors” (https://doi.org/10.3346/jkms.2017.32.7.1051). As of July 1, 2018, manuscripts submitted to ICMJE journals that report the results of clinical trials must contain a data sharing statement as described below. Clinical trials that begin enrolling participants on or after January 1, 2019 must include a data sharing plan in the trial’s registration. The ICMJE’s policy regarding trial registration is explained at www.icmje.org/recommendations/browse/publishingand-editorial-issues/clinical-trial-registration.html. If the data sharing plan changes after registration, this should be reflected in the statement submitted and published with the manuscript and updated in the registry record. Data sharing statements must indicate the following: whether individual deidentified participant data (including data dictionaries) will be shared; what data in particular will be shared; whether additional, related documents will be available (e.g., study protocol, statistical analysis plan, etc.); and when the data will become available and for how long; by what access criteria data will be shared (including with whom, for what types of analyses, and by what mechanism). Illustrative examples of data sharing statements that would meet these requirements are in Table 1.

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<th>Example 1</th>
<th>Example 2</th>
<th>Example 3</th>
<th>Example 4</th>
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<tr>
<td>Will individual participant data be available (including data dictionaries)?</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>What data in particular will be shared?</td>
<td>All of the individual participant data collected during the trial, after deidentification.</td>
<td>Individual participant data that underlie the results reported in this article, after deidentification (text, tables, figures, and appendices).</td>
<td>Individual participant data that underlie the results reported in this article, after deidentification (text, tables, figures, and appendices).</td>
<td>Not available</td>
</tr>
<tr>
<td>What other documents will be available?</td>
<td>Study protocol, statistical analysis plan, informed consent form, clinical study report, analytic code</td>
<td>Study protocol, statistical analysis plan, analytic code</td>
<td>Study protocol</td>
<td>Not available</td>
</tr>
<tr>
<td>When will data be available (start and end dates)?</td>
<td>Immediately following publication. No end date.</td>
<td>Beginning 3 months and ending 5 years following article publication.</td>
<td>Beginning 9 months and ending 36 months following article publication.</td>
<td>Not applicable</td>
</tr>
<tr>
<td>With whom?</td>
<td>Anyone who wishes to access the data.</td>
<td>Researchers who provide a methodologically sound proposal.</td>
<td>Investigators whose proposed use of the data has been approved by an independent review committee (“learned intermediary”) identified for this purpose.</td>
<td>Not applicable</td>
</tr>
<tr>
<td>For what types of analyses?</td>
<td>Any purpose</td>
<td>To achieve aims in the approved proposal.</td>
<td>For individual participant data meta-analysis.</td>
<td>Not applicable</td>
</tr>
<tr>
<td>By what mechanism will data be made available?</td>
<td>Data are available indefinitely at (link to be included).</td>
<td>Proposals should be directed to xxx@yyy. To gain access, data requestors will need to sign a data access agreement.</td>
<td>Proposals may be submitted up to 36 months following article publication. After 36 months, the data will be available in our University’s data warehouse but without investigator support other than deposited metadata. Information regarding submitting proposals and accessing data may be found at (link to be provided).</td>
<td>Not applicable</td>
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<td>Data are available for 5 years at a third-party website (link to be included).</td>
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ICMJE, International Committee of Medical Journal Editors.  
These examples are meant to illustrate a range of, but not all, data sharing options.

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Taesung Park  
Editor in Chief  
Genomics & Informatics  
Korea Genome Organization (KOGO)
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