Introduction (TBD)

The International Workshop on Health Text Mining and Information Analysis (LOUHI) provides an interdisciplinary forum for researchers interested in automated processing of health documents. Health documents encompass electronic health records, clinical guidelines, spontaneous reports for pharmacovigilance, biomedical literature, health forums/blogs or any other type of health-related documents. The LOUHI workshop series fosters interactions between the Computational Linguistics, Medical Informatics and Artificial Intelligence communities. The eight previous editions of the workshop were co-located with SMBM 2008 in Turku, Finland, with NAACL 2010 in Los Angeles, California, with Artificial Intelligence in Medicine (AIME 2011) in Bled, Slovenia, during NICTA Techfest 2013 in Sydney, Australia, co-located with EACL 2014 in Gothenburg, Sweden, with EMNLP 2015 in Lisbon, Portugal, with EMNLP 2016 in Austin, Texas; in 2017 was held in Sydney, Australia; and in 2018 was co-located with EMNLP 2018 in Brussels, Belgium. This year the workshop is co-located with EMNLP 2019 in Hong Kong.

The aim of the LOUHI 2019 workshop is to bring together research work on topics related to health documents, particularly emphasizing multidisciplinary aspects of health documentation and the interplay between nursing and medical sciences, information systems, computational linguistics and computer science. The topics include, but are not limited to, the following Natural Language Processing techniques and related areas:

- Techniques supporting information extraction, e.g. named entity recognition, negation and uncertainty detection
- Classification and text mining applications (e.g. diagnostic classifications such as ICD-10 and nursing intensity scores) and problems (e.g. handling of unbalanced data sets)
- Text representation, including dealing with data sparsity and dimensionality issues
- Domain adaptation, e.g. adaptation of standard NLP tools (incl. tokenizers, PoS-taggers, etc) to the medical domain
- Information fusion, i.e. integrating data from various sources, e.g. structured and narrative documentation
- Unsupervised methods, including distributional semantics
- Evaluation, gold/reference standard construction and annotation
- Syntactic, semantic and pragmatic analysis of health documents
- Anonymization/de-identification of health records and ethics
- Supporting the development of medical terminologies and ontologies
- Individualization of content, consumer health vocabularies, summarization and simplification of text
- NLP for supporting documentation and decision making practices
- Predictive modeling of adverse events, e.g. adverse drug events and hospital acquired infections
- Terminology and information model standards (SNOMED CT, FHIR) for health text mining
- Bridging gaps between formal ontology and biomedical NLP
The call for papers encouraged authors to submit papers describing substantial and completed work but also focus on a contribution, a negative result, a software package or work in progress. We also encouraged to report work on low-resourced languages, addressing the challenges of data sparsity and language characteristic diversity.

This year we received a high number of submissions (50), therefore the selection process was very competitive. Due to time and space limitations, we could only choose a small number of the submitted papers to appear in the program.

Each submission went through a double-blind review process which involved three program committee members. Based on comments and rankings supplied by the reviewers, we accepted 23 papers. Although the selection was entirely based on the scores provided by the reviewers, we regretfully had to set a relatively high threshold for acceptance. The overall acceptance rate is 46%. After the decision about acceptance, 2 papers were withdrawn by the authors. During the workshop, 11 papers will be presented orally, and 10 papers will be presented as posters.

Finally, we would like to thank the members of the program committee for providing balanced reviews in a very short period of time, and the authors for their submissions and the quality of their work.
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Table of Contents

Cross-document coreference: An approach to capturing coreference without context
   Kristin Wright-Bettner, Martha Palmer, Guergana Savova, Piet de Groen and Timothy Miller . . . 1

Comparing the Intrinsic Performance of Clinical Concept Embeddings by Their Field of Medicine
   John-Jose Nunez and Giuseppe Carenini ................................................................. 11

On the Effectiveness of the Pooling Methods for Biomedical Relation Extraction with Deep Learning
   Tuan Ngo Nguyen, Franck Dernoncourt and Thien Huu Nguyen .................................. 18

Syntax-aware Multi-task Graph Convolutional Networks for Biomedical Relation Extraction
   Diya Li and Heng Ji .................................................................................................. 28

BioReddit: Word Embeddings for User-Generated Biomedical NLP
   Marco Basaldella and Nigel Collier ........................................................................... 34

Leveraging Hierarchical Category Knowledge for Data-Imbalanced Multi-Label Diagnostic Text Understanding
   Shang-Chi Tsai, Ting-Yun Chang and Yun-Nung Chen ........................................... 39

Experiments with ad hoc ambiguous abbreviation expansion
   Agnieszka Mykowiecka and Malgorzata Marciniak ................................................... 44

Multi-Task, Multi-Channel, Multi-Input Learning for Mental Illness Detection using Social Media Text
   Prasadith Kirinde Gamaarachchige and Diana Inkpen ............................................. 54

Extracting relevant information from physician-patient dialogues for automated clinical note taking
   Serena Jeblee, Faiza Khan Khattak, Noah Crampton, Muhammad Mamdani and Frank Rudzicz 65

Biomedical Relation Classification by single and multiple source domain adaptation
   Sinchani Chakraborty, Sudeshna Sarkar, Pawan Goyal and Mahanandeeshwar Gattu ........... 75

Assessing the Efficacy of Clinical Sentiment Analysis and Topic Extraction in Psychiatric Readmission Risk Prediction
   Elena Alvarez-Mellado, Eben Holderness, Nicholas Miller, Fyonn Dhang, Philip Cawkwell, Kirsten Bolton, James Pustejovsky and Mei-Hua Hall ...................................................... 81

What does the language of foods say about us?
   Hoang Van, Ahmad Musa, Hang Chen, Stephen Kobourov and Mihai Surdeanu ............... 87

Dreaddit: A Reddit Dataset for Stress Analysis in Social Media
   Elsbeth Turcan and Kathy McKeown ........................................................................ 97

Towards Understanding of Medical Randomized Controlled Trials by Conclusion Generation
   Alexander Te-Wei Shieh, Yung-Sung Chuang, Shang-Yu Su and Yun-Nung Chen ............ 108

Building a De-identification System for Real Swedish Clinical Text Using Pseudonymised Clinical Text
   Hanna Berg, Taridzo Chomutare and Hercules Dalianis .......................................... 118

Automatic rubric-based content grading for clinical notes
   Wen-wai Yim, Ashley Mills, Harold Chun, Teresa Hashiguchi, Justin Yew and Bryan Lu ..... 126

Dilated LSTM with attention for Classification of Suicide Notes
   Annika M Schoene, George Lacey, Alexander P Turner and Nina Dethlefs .................... 136
Writing habits and telltale neighbors: analyzing clinical concept usage patterns with sublanguage embeddings
Denis Newman-Griffis and Eric Fosler-Lussier ................................................................. 146

Recognizing UMLS Semantic Types with Deep Learning
Isar Nejadgholi, Kathleen C. Fraser, Berry De Bruijn, Muqun Li, Astha LaPlante and Khaldoun Zine El Abidine ................................................................. 157

Ontological attention ensembles for capturing semantic concepts in ICD code prediction from clinical text
Matus Falis, Maciej Pajak, Aneta Lisowska, Patrick Schrempf, Lucas Deckers, Shadia Mikhael, Sotirios Tsaftaris and Alison O’Neil ................................................................. 168

Neural Token Representations and Negation and Speculation Scope Detection in Biomedical and General Domain Text
Elena Sergeeva, Henghui Zhu, Amir Tahmasebi and Peter Szolovits ........................................ 178
Conference Program

November 3, 2019

9:00–10:30 Session 1

9:00  Introduction

9:05  Cross-document coreference: An approach to capturing coreference without context
Kristin Wright-Bettner, Martha Palmer, Guergana Savova, Piet de Groen and Timothy Miller

09:30  Poster booster

09:45  Poster session

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Elena Alvarez-Mellado, Eben Holderness, Nicholas Miller, Fyonn Dhang, Philip Cawkwell, Kirsten Bolton, James Pustejovsky and Mei-Hua Hall

10:30–11:00 Break

11:00–12:30 Session 2

11:00 What does the language of foods say about us?
Hoang Yan, Ahmad Musa, Hang Chen, Stephen Kobourov and Mihai Surdeanu

11:25 Dreaddit: A Reddit Dataset for Stress Analysis in Social Media
Elsbeth Turcan and Kathy McKeown

11:50 Towards Understanding of Medical Randomized Controlled Trials by Conclusion Generation
Alexander Te-Wei Shieh, Yung-Sung Chuang, Shang-Yu Su and Yun-Nung Chen

12:15 Building a De-identification System for Real Swedish Clinical Text Using Pseudonymised Clinical Text
Hanna Berg, Taridzo Chomutare and Hercules Dalianis
November 3, 2019 (continued)

12:40–14:00 Lunch

14:00–15:30 Session 3

14:00  Invited Talk
       TBA

14:40  Automatic rubric-based content grading for clinical notes
       Wen-wai Yim, Ashley Mills, Harold Chun, Teresa Hashiguchi, Justin Yew and
       Bryan Lu

15:05  Dilated LSTM with attention for Classification of Suicide Notes
       Annika M Schoene, George Lacey, Alexander P Turner and Nina Dethlefs

15:30–16:00 Break

16:00–17:40 Session 4

16:00  Writing habits and telltale neighbors: analyzing clinical concept usage patterns
       with sublanguage embeddings
       Denis Newman-Griffis and Eric Fosler-Lussier

16:25  Recognizing UMLS Semantic Types with Deep Learning
       Isar Nejadgholi, Kathleen C. Fraser, Berry De Bruijn, Muqun Li, Astha LaPlante
       and Khaldoun Zine El Abidine

16:50  Ontological attention ensembles for capturing semantic concepts in ICD code pre-
       diction from clinical text
       Matus Falis, Maciej Pajak, Aneta Lisowska, Patrick Schrempf, Lucas Deckers, Sha-
       dia Mikhael, Sotirios Tsafarlis and Alison O’Neil

17:15  Neural Token Representations and Negation and Speculation Scope Detection in
       Biomedical and General Domain Text
       Elena Sergeeva, Henghui Zhu, Amir Tahmasebi and Peter Szolovits
Cross-document coreference: An approach to capturing coreference without context

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Abstract

In this paper, we discuss a cross-document coreference annotation schema that we developed to further automatic extraction of timelines in the clinical domain. Lexical senses and coreference choices are determined largely by context, but cross-document work requires reasoning across contexts that are not necessarily coherent. We found that an annotation approach that relies less on context-guided annotator intuitions and more on schematic rules was most effective in creating meaningful and consistent cross-document relations.

1 Introduction

The ability to learn cross-document coreference and temporal relationships in clinical text is crucial for the automatic extraction of comprehensive patient timelines of events (Raghavan et al., 2014). To that end, we present a gold corpus of 198 clinical-narrative document sets, where each set consists of three notes for a given patient (594 individual notes total). Each file is annotated with intra-document temporal, coreference, and bridging relations (SET-SUBSET, WHOLE-PART, CONTAINS-SUBEVENT), and each set is annotated with cross-document coreference and bridging relations.

The goal of the current project was to leverage the inherited, intra-document annotations from two prior projects (discussed in Section 2) to capture longer, more developed timelines of patient information. We did this by creating human-annotated cross-document coreference and bridging links and then using inference to combine this information with the knowledge already gained from the intra-document temporal and coreference/bridging links.

In this paper, we discuss the impacts of cross-document-specific phenomena on human annotation and machine learning, most notably the effect of disjunct narratives on cross-document coreference judgments. Cohesive discourse is a crucial linguistic tool for determining coreference, yet the cross-document relations annotation task fundamentally takes place across discontinuous narratives. We found an approach that is governed more by annotation rules than annotator intuition to be most effective, producing an inter-annotator agreement score of 93.77% for identical relations. While an approach that moves away from linguistically-intuitive judgments may seem surprising at first, it is in fact quite fitting for a task that is inherently void of the discourse-level linguistic cues that humans employ to make those intuitive associations.

We also discuss other cross-document phenomena, inter-annotator agreement, and, briefly, areas for future work. Related work is discussed throughout.

2 The THYME colon cancer corpus

This annotation effort merged and expanded on document-level annotations created by two prior projects – a temporal relations project¹ (Styler et al., 2014), and a coreference and bridging relations project.² These two projects will be referred to as THYME 1 (Temporal History of Your Medical Events) and Clinical Coreference.

---

The corpus consists of de-identified physicians’ notes on colon cancer patients. The examples used throughout this paper are artificially created; however, we have done our best to replicate the relevant linguistic contexts. Each set of three notes consists of a clinical report, a pathology report, and a second clinical report, in that chronological order and spanning a period of weeks or months. Capturing such temporally-extensive information gives us the ability to track the status of the disease over time and responses (or not) to treatment.

The THYME colon cancer corpus now includes: a) intra-document gold annotations for all markables (events, entities, and temporal expressions) and several types of temporal, coreference, and bridging relations; and b) cross-document gold annotations for four coreference and bridging relation types, which represent a subset of the intra-document types and are described in Table 1.

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<table>
<thead>
<tr>
<th>Relation Type</th>
<th>Description</th>
<th>Link</th>
</tr>
</thead>
<tbody>
<tr>
<td>IDENTICAL (IDENT)</td>
<td>M1 refers to the same event/entity as M2</td>
<td>[M1] IDENT [M2]</td>
</tr>
<tr>
<td>SET-SUBSET (S-SS)</td>
<td>M2 refers to one or more members of a larger group, represented by M1.</td>
<td>[M1] SET - [M2] SUBSET</td>
</tr>
<tr>
<td>CONTAINS-SUBEVENT (CON-SUB)</td>
<td>M1 temporally contains M2, and M2 is inherently part of the structure of M1.</td>
<td>[M1] CON-SUB [M2]</td>
</tr>
<tr>
<td>WHOLE-PART (W-P)</td>
<td>M2 is compositionally part of a larger entity, represented by M1.</td>
<td>[M1] WHOLE - [M2] PART</td>
</tr>
</tbody>
</table>

Table 1: Gold-annotated cross-document relation types in the THYME colon cancer corpus. M1 refers to Markable 1 and M2 refers to Markable 2. Markables include events, entities, or temporal expressions. W-P was used only for entities; CON-SUB only for events. All four relation types are coreference or bridging links rather than temporal links, except for CON-SUB, which conveys both temporal and structural information and is represented as a temporal link (TLINK) in our annotation tool. This TLINK type is discussed in Section 3.1.

Many prior studies have noted the intractability of creating cross-document gold annotations on large corpora (Day et al., 2000, for example). Each cross-document effort has therefore restricted the scope of their annotations in some way (e.g., Song et al., 2018; Cybulskia and Vossen, 2014) and/or developed machine-produced annotations for cross-document relations, rather than human-produced (Raghavan et al., 2014; Dutta and Weikum, 2015; Baron and Freedman, 2008; Go and Allen 2004; etc.). We likewise restricted our approach by limiting the cross-document relations to the groups of three files which represent each patient, and by limiting the number of annotated relation types. However, the THYME corpus is the largest dataset of gold-annotated clinical narratives to-date that we are aware of, in terms of types of markables and relations annotated.

We are indebted to the contributions of the projects that preceded us. Much of the technical and conceptual groundwork had already been laid for our task. In particular, the notion of narrative containers (Styler et al., 2014; Pustejovsky and Stubbs, 2011) informed our addition of the CONTAINS-SUBEVENT temporal link and our cross-document annotation process.

However, we found that the segregation of tasks during the creation of the single-file gold annotations caused a variety of technical and conceptual conflicts once their outputs were merged. Furthermore, aspects of the temporal task suffered from focusing only on local-context relations; a global grasp of the text, which coreference annotation facilitates, reveals...
temporally significant information that may be otherwise missed or misinterpreted.

Early experiments showed that conflicts in the merged annotations rendered meaningful cross-document annotation untenable. To reconcile these conflicts, we therefore introduced an intra-document corrections-style manual annotation pass prior to cross-document double-annotation and adjudication.

3 Cross-document annotation: Process, assumptions, phenomena

It has been well-attested that determining cross-document relations poses a unique set of challenges for both systems and annotators. Song et al. (2018) discuss the cognitive strain on annotators, and others have observed the decrease in linguistic cues that occurs cross-document (Raghavan et al., 2014; Hong et al., 2016). In this paper, we are most interested in the latter, particularly the impacts of cross-document mode on identical relations.

Many coreference annotation guidelines, including ours, use a straightforward definition of coreference, which may be summarized as two different mentions in a text having the same real-or hypothetical-world referent (e.g., Cybulska and Vossen, 2014; Richer Event Description Annotation Guidelines, 2016 5; Cohen et al., 2017). This definition leads to a binary approach to identical judgments – two mentions either refer to the same thing or they do not. Annotators are forced to make a polar choice about representations of meaning, when those representations in fact exist on a spectrum. This is not a new discovery: “There are cases where variant readings of a single lexical form would seem to be more appropriately visualized as points on a continuum – a single fabric of meaning with no clear boundaries” (Cruse, 1986). However, the natural language processing community is still learning how to deal with this.

Others have identified the problems that this oversimplified definition creates for annotation: “Degrees of referentiality as well as relations that do not fall neatly into either coreference or non-coreference—or that accept both interpretations—are a major reason for the lack of inter-coder agreement in coreference annotation” (Recasens, 2010). Hovy et al. (2013) also recognized the need for a more nuanced approach and introduced membership and subevent relations as a result.

Furthermore, both Recasens and Hovy discuss the role that pragmatics plays in determining coreference:

- “Two mentions fully corefer if their activity/event/state DE [discourse element] is identical in all respects, as far as one can tell from their occurrence in the text” (Hovy et al., 2013, emphasis added).
- “We redefine coreference as a scalar relation between two (or more) linguistic expressions that refer to discourse entities considered to be at the same granularity level relevant to the linguistic and pragmatic context” (Recasens et al., 2011, emphasis added).

Context, therefore, contributes in a crucial way to determining sense for a given lexical unit – and therefore also to determining coreference relations for that unit. We agree with Cruse, Recasens, and Hovy and observe the unique challenge this poses for cross-document annotation, since distinct narratives do not share a coherent discourse context. Recasens et al. (2011) propose that categorization and meaning are constructed in a temporary, active process; in cross-document work, we are attempting to create meaningful relations between temporally disconnected discourses. Put differently, the coherence of context is decreased while the number of contexts for lexical senses is increased.

While not surprising, this phenomenon does have interesting consequences for annotation. In fact, by the definitions given above, “doing” coreference between disjunct linguistic and pragmatic contexts could be viewed, on some level, as impossible.

But not all hope is lost. Particularly for our corpus, texts are very closely related and it is possible to create meaningful relations. However, the phenomenon just described requires an approach to cross-document coreference annotation that is unique from within-document. We dealt with this primarily by adding a subevent relation that was governed more by annotation rules and less by annotators’ intuitions. We present the reasons for and outcome of this approach in the next section, followed by discussion of other cross-document phenomena and our technical cross-document linking process.

---

5 https://github.com/timjogorman/RicherEventDescription/blob/master/guidelines.md
3.1 An approach to coreference across disjunct contexts

Consider the following single-file example:

(1) October 15th, 2015 – Dr. Wu performed resection of the primary tumor. Ms. Smith’s recovery from surgery has been without complication.

The choice here about whether to link resection and surgery as coreferential is likely to produce a disagreement. Annotator A may decide they are IDENTICAL (IDENT) since they clearly refer on some level to the same cancer treatment procedure; a significant semantic relationship would be lost if we did not link them. Annotator B, however, may decide resection refers only to the literal act of removing the tumor, while surgery points to the entire procedure. Essentially, the annotators disagree about whether the two terms are “close enough” on the meaning spectrum to warrant an IDENT link. More precisely, the disagreement stems from different interpretations of semantic granularity – Annotator A’s identity “lens” is more coarse-grained, while Annotator B’s is more fine-grained.

Consider a second example:

(2) PLAN: Resection of primary tumor and gallbladder removal. Patient is scheduled for surgery on October 15th, 2015.

Here, the finer-grained approach to Resection and surgery is supported – required, in fact – by the context. No coreference relationship is possible since the surgery clearly consists of two subprocedures, the tumor resection and the gallbladder removal.

Now consider the two examples together, where (2) is from the chronologically earlier note and (1) is from the later note in a single set:

- No coreference link
- surgery CONTAINS Resection
- surgery CONTAINS removal

Note B: October 15th, 2015 – Dr. Wu performed resection of the primary tumor. Ms. Smith’s recovery from surgery has been without complication.
- resection IDENTICAL surgery

The IDENT link shown for Note B represents the original gold annotation in our data, i.e., the more coarse-grained approach to identity described above. This is arguably the better perspective here, based on Recasens’ definition of coreference above (“discourse entities considered to be at the same granularity level relevant to the linguistic and pragmatic context”); Note B’s narrative is quite broad-brushed and supports what Hovy terms a “wide reading” of resection (Hovy et al., 2013). Pragmatically, resection and surgery are the same in Note B; pragmatically, they are not the same in Note A.

The predicament for cross-document linking is obvious. If we link resectionA to resectionB, this entails that resectionA is IDENT to surgeryB; if we then link surgeryA to surgeryB, this now entails that the procedure temporally contains itself. If we leave resectionA unlinked to resectionB to avoid this conflict, problematically, we miss the relation between identical strings that refer to the same event (Resection of primary tumor IDENT resection of primary tumor), not to mention that leaving these unlinked would be extremely counterintuitive for annotators.

This type of situation is common in cross-document work. Since identity judgments are based on granularity levels that are in turn determined by the pragmatics of the narrative, and since the pragmatic contexts of two or more disjunct narratives are not necessarily coherent, cross-document mode frequently forces annotators to choose between: (a) not linking two mentions that are obviously and significantly semantically related, or (b) linking these mentions and thereby forcing logically-conflicting information as in (3), which in turn renders the existing temporal links much less meaningful.

To account for this variation in context-determined granularity, we introduced the CONTAINS-SUBEVENT (CON-SUB) link, which says that EVENT B is both temporally contained by EVENT A and it composes part of EVENT A’s essential structure (modeled after the subevent relation in O’Gorman et al., 2016). We added this new relation intra-document in the corrections pass, as well as in the later cross-document pass. For examples like (3), this meant the Note B IDENT relation was re-interpreted as a subevent relation: surgery CON-SUB resection. This allowed us to preserve the close semantic connection between the two EVENTS in both narratives, while avoiding the logical conflicts that would have rendered our output much less meaningful and informative. We can also assume the inter-annotator agreement...
achieved (discussed in section 4) is much higher than it would have been had we left annotators in the predicament shown in (3).

The consistency noted above was achieved by an approach that relied less on discourse cues and more on general semantic distinctions. Instead of allowing annotators to intuitively judge between wide and narrow readings (borrowing Hovy’s terms again) of lexical items based on the context, we required IDENT and CON-SUB relations to be based more on the dictionary definitions of the terms. This is because we could not predict the granularity distinctions that cross-document information would expose, as shown in (3). For example, annotators were required to differentiate between “general” surgery terms (e.g., surgery, procedure, operation, etc.) and “specific” surgery terms (colectomy, resection, excision, etc.), such that the general term nearly always contained the specific term as a subevent. This compensated for the majority of granularity distinctions that cross-document information would expose, as shown in (3). For example, annotators were required to differentiate between “general” surgery terms (e.g., surgery, procedure, operation, etc.) and “specific” surgery terms (colectomy, resection, excision, etc.), such that the general term nearly always contained the specific term as a subevent. This compensated for the majority of granularity distinctions that cross-document information would expose, as shown in (3).

Song et al. (2018) took an opposite approach to cross-document coreference linking through their use of event hoppers, which permit “coreference of two events that are intuitively the same although certain features may differ” (emphasis added). We found this approach did not suit our needs since the ultimate goal was to capture a coherent timeline of clinical events, and intuitive coreference linking produced temporal conflicts, as shown above.

While coreference linking is not possible on the cross-document level in the same nuanced and intuitive way that it is within-document, there is still a great deal of important information we can capture. The texts in our corpus are topical very similar and there are typically a lot of corroborating details, such as dates and locations (again, these have been de-identified, but in a consistent fashion). Additionally, the clinically-delineated sections and the note types and structure provide clues about how to interpret the events; for example, due to the date and descriptive details, we can know which procedure in the clinical note the pathology note refers to, even if the overall procedure is not explicitly mentioned in the pathology note.

Time constraints prevented us from adding CON-SUB for all event types. We annotated it for four event categories, chosen based on clinical significance and demonstrated need due to cross-document conflicts like the one in (3): (a) patient treatment events, including surgical procedures and chemotherapy/radiation treatments; (b) cancer events (cancer, adenocarcinoma, tumor, etc.); (c) medications; and (d) chronic disease events.

Due to other conflicts arising from the disconnected contexts that the subevent relation was not able to reconcile, we permitted the cross-document adjudicators (but not annotators) to make within-document annotation changes when absolutely necessary.

In summary, we found that it is possible to capture meaningful cross-document coreference relations, but the approach must differ from intra-document annotation because pragmatically-directed within-document intuitions may conflict in unpredictable ways on the cross-document level.

3.2 Other cross-document phenomena

We have discussed in depth the way identical judgments are affected by disjunct contexts. We discuss two more cross-document phenomena here: (1) the use of inference in linking stative events; and (2) how cross-document work exposes typos and misinformation. Notably, this could be leveraged to identify mistakes in the text, which may contribute to current efforts to reduce medical errors in patient treatment.

Inference and stative events

Cross-document coreference is typically easier for punctual events (such as tests and procedures) and harder for durative events that can change in value over time (for example, a mass that is initially benign but becomes malignant). As with many other cross-document challenges, this issue is also present within-document, but is exacerbated in the cross-document setting because context is reduced. Consider the following example:

(4) Note A (March 24 2012 SECTIONTIME):

**Pulse Rate:** Regular

Note B (March 26 2012 SECTIONTIME):

**Heart:** Regular rate

Here we have two clinically-relevant states associated with two different times: the regularity of the patient’s heart rate on March 24, 2012, and the regularity of the patient’s heart rate on March 26, 2012. The question for a cross-doc annotator is whether these two EVENTs are IDENTICAL.

For the current example, it is likely the regular condition has continued, but the fact is we do not
know, especially since the patient may have a medical condition that causes sporadic irregularity. Furthermore, we might be initially inclined to infer sameness due to the close temporal proximity of the two measurements (two days apart), but that thought trajectory quickly leads to problems: When are two continuous events not temporally near enough to infer sameness? A week? A month? How do we decide?

Song et al. (2018) discuss a similar example across four notes, in which they corefer the first three events because they occur in “about the same time period and same place” (occurring over the timespan of a month), but they do not corefer the fourth event “as it happened at a different time” (about four months after the most recent other mention). However, it is not clear how they determined that a month is a reasonably close enough timespan to infer sameness, while four months is not.

Our approach, therefore, was that when condition or attributive EVENTS – events that vary in value – are measured or identified at two different times, they should not be linked, unless there is explicit linguistic evidence (e.g., use of the present perfect tense) they are the same event. Essentially, we decided that temporal proximity alone was not enough to infer an identical relation for two condition/value events.

Of course, inference is a source of inter-annotator disagreement for other cross-document choices as well. A comprehensive analysis is outside the scope of this paper, but the topic is discussed further in the following point.

**How cross-document annotation exposes mistakes in the data**

We discuss this in detail not only because it has implications for discovering misinformation in the text, but also because it demonstrates two more significant challenges to cross-document clinical annotation: the heavy cognitive burden on annotators, and the need for clinical knowledge. Consider the following example:

(5) **Note A** (DOCTIME: August 21, 2012):

We have ordered a *CT* abdomen and pelvis to rule out liver metastases prior to surgery. Mr. Olson will also need an EKG and bloodwork. *Testing was negative.*

- *CT* assigned DocTimeRel of *AFTER*, i.e., it occurs after DOCTIME (Aug 21, 2012).

**Note B** (DOCTIME: September 30, 2012):

*CT* abdomen and pelvis was compared to the prior *study* of August 20, 2012, Mr. Olson had low-anterior resection.

- August 20, 2012 CONTAINS study

CT$_A$ and *study$_B$* are in fact IDENTICAL. Combined with the temporal information noted above, this entails that the same event both occurs after Aug. 21, 2012, and is temporally contained by Aug. 20, 2012 – a logical impossibility.

We know they are the same event based primarily on real-world knowledge of the standard order of medical procedures, as follows: It is clear in Note B that there are two different CT scans. The question facing a cross-document annotator is which one, if either, is IDENT to CT$_A$? We know explicitly from the text that CT$_A$ occurred prior to the patient’s surgery. CT$_B$ occurred after the patient’s surgery, since, however cryptically, it references observation of the surgery (“Mr. Olson had low-anterior resection”). Therefore, CT$_A$ and CT$_B$ are not referring to the same scan.

Now the question is whether *study$_B$* is IDENT to CT$_A$. The initial evidence is to the contrary – *study$_B$* is explicitly said to occur on Aug. 20, while CT is inferable after (or later in the day on) Aug. 21. However: (a) it is unusual to have two CT scans back-to-back, without further discussion; (b) an Aug. 20 CT is not discussed in the Aug. 21 note; (c) in Note A, immediately after noting that several tests have been ordered, the text says, “Testing was negative.” Based on the verb tenses in the paragraph, the assumption would likely be that *Testing* here refers to other tests, not the ones just ordered. However, the flow of discourse suggests otherwise, along with the fact that no other prior testing is referred to in the same section. With the additional information we have from Note B, a more reasonable interpretation presents itself: *study$_B$* is IDENT to CT$_A$, and Aug. 20 is the correct date of the scan. The note was likely originally written on Aug. 20, prior to the scan that was done later that day, and was later updated with the test results but without any indication of the update being written at a later time. This analysis was confirmed by review of all notes by our medical expert consultant.

There are several noteworthy observations about this: First, there is quite a bit of oncological knowledge required to notice the conflict above. Furthermore, the non-standard syntax in Note B would make it easy for an annotator to miss the fact that CT$_B$ is after the resection.
Second, even armed with the necessary clinical knowledge, there is still a fair amount of inference involved in making the above choice. However, note that all of the annotation options here, including the option to not link at all, require a lot of inference (as is the nature of many cross-document analyses). There are different types of inference based on different kinds of information. While we decided that temporal closeness is not enough by itself to infer a relation for condition/value events, we decided here that medical knowledge of standard processes is enough to infer a relation.

Third, assuming the above observations were made, an impossible annotation choice presents itself: Do we make the coreference link even though it forces a temporal conflict, or do we keep the timeline clean and lose the coreference relation? We decided on the former, and kept track of the noted temporal conflicts in order to inform systems training.

Finally, note the time, attention, and careful thought process required for determining this single cross-document link. While certainly not all decisions are this demanding, the amount of time necessary to produce high-caliber annotations should be apparent. It took highly-experienced annotators about 1.5 hours on average to complete one document set, or an estimated 891 hours total for two annotators and one adjudicator to produce 198 gold sets with a total of 10,560 cross-document links. This does not include time spent on initial annotation experiments, process and guidelines development, annotator training, and post-processing steps.

### 3.3 Cross-document annotation process

To manage the potentially vast number of cross-document links, we established a set of assumptions about inferable relations that guided the following process and are further discussed in Table 2 (note: “structural links” refers to links that have a hierarchical rather than identical relationship: CON-SUB, S-SS, W-P):

- (a) Link topmost mention to topmost mention. We assume the other relations can be inferred from within-document chains.
- (b) If there is a within-document structural link between two markables, do not create that same link cross-document for the same two events/entities. Put differently, create cross-document structural links only when both components of the relation do not have a cross-document IDENT link. Again, we assume that other relations can be inferred.
- (c) Always create IDENT links whenever appropriate.

### 4 Inter-annotator agreement

We scored inter-annotator agreement (IAA) only for annotation categories that were new to the current project, i.e., intra-document CON-SUB links and all cross-document links. Furthermore, we only scored annotator-annotator agreement (not annotator-gold), since adjudicators were permitted to change single-file annotations while annotators were not. The total number of gold markables and relations are shown in Table 3; IAA results are shown in Table 4 and are averaged over all the documents (both tables shown on following page).

The IDENT score is much higher than the structural linking scores because the structural links were only created in cases where neither component of the link had a cross-document IDENT relation (see Section 3.3). These relations were therefore brand-new and had to be identified...
without the benefit of a single coherent discourse, as discussed in depth above. On the other hand, annotators were able to draw on the information conveyed in intra-document relations when determining cross-document IDENT relations.

The WHOLE-PART (W-P) IAA score is zero because there were very few cross-document W-P relations in the corpus, under our guidelines. W-P is used only for entities, and we did not do W-P cross-document linking for anatomical entities (due to the massive amount of mentions, the spider-webbed relations, and the number of vague terms – tissue portions, etc. – we only created IDENT anatomy relations at the cross-document level). Therefore, the only cross-narrative W-P relations were between organizations/departments and members of those entities, which were only rarely knowable from the text.

The CONTAINS-SUBEVENT (CON-SUB) agreement score is likely higher than the SET-SUBSET (S-SS) score because we applied it to four specific event categories (see Section 3.1) that consist of oft-repeated terms. S-SS, on the other hand, had no such constraints, making this relation much more challenging to identify over the scope of three often-lengthy documents. Furthermore, while some set-member relations are obvious, others are not. For example:

(6) **Note A:** Pt denies alcohol or tobacco use.

**Note B:** He denies drinking.

- use\_NEG S-SS drinking\_NEG

<table>
<thead>
<tr>
<th>Markables (594 documents)</th>
<th>143,147 total</th>
<th>Relations, within-doc and cross-doc (594 documents)</th>
<th>70,572 total</th>
<th>Cross-doc relations (198 documents)</th>
<th>10,762 total</th>
</tr>
</thead>
<tbody>
<tr>
<td>TIMEX3s</td>
<td>7,796</td>
<td>Temporal links</td>
<td>35,428 total</td>
<td>IDENTICAL</td>
<td>9,102</td>
</tr>
<tr>
<td>Entities*</td>
<td>47,355</td>
<td>CONTAINS</td>
<td>14,037</td>
<td>SET-SUBSET</td>
<td>405</td>
</tr>
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<td>EVENTSs</td>
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<td>CON-SUB**</td>
<td>4,718</td>
<td>WHOLE-PART</td>
<td>405</td>
</tr>
<tr>
<td>SECTIONTIME</td>
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<td>BEFORE</td>
<td>4,217</td>
<td>CON-SUB</td>
<td>1,242</td>
</tr>
<tr>
<td>DOCTIME</td>
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<td>OVERLAP</td>
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<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>BEGINS-ON</td>
<td>1,200</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>ENDS-ON</td>
<td>557</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>NOTED-ON</td>
<td>5,608</td>
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<tr>
<td></td>
<td></td>
<td>Aspectual links</td>
<td>873 total</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>IDENTICAL</td>
<td>23,827</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>SET-SUBSET</td>
<td>5,907</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>WHOLE-PART</td>
<td>3,885</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>CON-SUB**</td>
<td>4,718</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Entities are referred to as MARKABLEs in our guidelines, due to the naming practice of the prior Clinical Coreference project.

Table 3: Total gold markables and relations for the THYME colon cancer corpus.

<table>
<thead>
<tr>
<th>Intra-document IAA</th>
<th>Cross-document IAA</th>
</tr>
</thead>
<tbody>
<tr>
<td>CON-SUB: 34.14%</td>
<td>IDENTICAL: 93.77%</td>
</tr>
<tr>
<td></td>
<td>CON-SUB: 36.43%</td>
</tr>
<tr>
<td></td>
<td>SET-SUBSET: 6.88%</td>
</tr>
<tr>
<td></td>
<td>WHOLE-PART: 0.00%</td>
</tr>
</tbody>
</table>

Table 4: Intra-document and cross-document inter-annotator agreement scores in terms of percentage agreement.

One of our annotators identified the S-SS link shown, while the other did not. In the future, more examples and/or constraints of fringe S-SS relations in the annotation guidelines could be developed to improve S-SS agreement.

5 Conclusion

As demonstrated, developing an extensive timeline of patient events that occur over multiple weeks and months is an extremely complicated process. Understanding the breadth of complexity and the heavy demands on annotators is necessary for projecting annotation budgets and timelines, and for understanding the nature and quality of the resulting data for predicting machine learning performance. Two of the most pressing areas for future research include: (a) further development and testing of our approach to cross-document linking presented in section 3.1; and (b) development of a comprehensive methodology for incorporating medical expertise, as alluded to in section 3.2 (building on but extending beyond the light-annotation tasks methodology proposed
by Stubbs, 2013). It is critical that wherever possible the annotation process is based on clear rules rather than annotator intuition as the former lends itself to automation whereas the latter at best results in a non-scalable solution with a narrow field of implementation. Developing these rules requires medical domain expertise.

Our results for cross-document coreference annotation leave ample room for improvement. Yet we believe that the approaches discussed here will serve as another significant step in the development of automatic extraction of event timelines in medical data.

Acknowledgments

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References


Comparing the Intrinsic Performance of Clinical Concept Embeddings by Their Field of Medicine

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Abstract
Pre-trained word embeddings are becoming increasingly popular for natural language-processing tasks. This includes medical applications, where embeddings are trained for clinical concepts using specific medical data. Recent work continues to improve on these embeddings. However, no one has yet sought to determine whether these embeddings work as well for one field of medicine as they do in others. In this work, we use intrinsic methods to evaluate embeddings from the various fields of medicine as defined by their ICD-9 systems. We find significant differences between fields, and motivate future work to investigate whether extrinsic tasks will follow a similar pattern.

1 Introduction
The application of natural language processing (NLP) and machine learning to medicine presents an exciting opportunity for tasks requiring prediction and classification. Examples so far include predicting the risk of suicide or accidental death after a patient is discharged from general hospitals (McCoy et al., 2016) or classifying which patients have peripheral vascular disease (Afzal et al., 2017). A common resource across NLP for such tasks is to use high-dimensional vector word representations. These word embedding include the popular \textit{word2vec} system (Mikolov et al., 2013) which was initially trained on general English text, using a skip-gram model on a Google News corpus.

Due to considerable differences between the language of medical text and general English writing, prior work has trained medical embeddings using specific medical sources. Generally, these approaches have trained embeddings to represent medical concepts according to their ‘clinical unique identifiers’ (CUIs) in the Unified Library Management System (ULMS) (Bodenreider, 2004). Words in a text can then be mapped to these CUIs (Yu and Cai, 2013). Various sources have been used, such as medical journal articles, clinical patient records, and insurance claims (De Vine et al., 2014), (Minarro-Giménez et al., 2014), (Choi et al., 2016).

Prior authors have sought to improve the quality of these embeddings, such as using different training techniques or more training data (Beam et al., 2018). In order to judge the quality of these embeddings, they have primarily used evaluation methods quantifying intrinsic qualities, such as their ability to predict drug-disease relations noted in the National Drug File - Reference Terminology (NDF-RT) ontology (Minarro-Giménez et al., 2014), or whether similar types of clinical concepts had cosine similar vectors (Choi et al., 2016).

To date these embeddings have been both trained and evaluated on general medical data. That is, no fields of medicine were specified or excluded; data could be from an obstetrician delivering a baby, a cardiologist placing a stent, or a dermatologist suggesting acne treatment. It is unclear how well such embeddings perform for a specific field of medicine. For example, we can consider psychiatry, the field of medicine concerned with mental illnesses such as depression or schizophrenia. Prior work has shown that psychiatric symptoms are often described in a long, varied, and subjective manner (Forbush et al., 2013) which may present a particular challenge for training these embeddings and NLP tasks generally.

As these pre-trained embeddings may increasingly be used for down-stream NLP tasks in spe-
cific fields of medicine, we seek to determine whether embeddings from one field perform relatively well or poorly relative to others. Specifically, we aim to follow prior work using intrinsic evaluation methods, comparing the geometric properties of embedding vectors against others given known relationships. This will offer a foundation for future work that may compare the performance on extrinsic NLP tasks in different medical fields. Finding relative differences may support that certain medical fields would benefit from embeddings trained on data specific to their field, or using domain adaptation techniques as sometimes used in the past (Yu et al., 2017).

2 Methods

2.1 Sets of Embeddings

We sought to compare a variety of clinical concept embeddings trained on medical data. Table 1 contains details of the sets compared in this project, all of which are based on word2vec. We obtained DeVine200 (De Vine et al., 2014), ChoiClaims300, and ChoiClinical300 (Choi et al., 2016) all from the latter’s Github. We downloaded BeamCui2Vec500 (Beam et al., 2018) from this site. Unfortunately, we were unable to obtain other sets of embeddings mentioned in the literature (Minarro-Giménez et al., 2014), (Zhang et al., 2018) (Xiang et al., 2019).

2.2 Determining a Field of Medicine’s Clinical Concepts

A clinical concept’s corresponding field of medicine is not necessarily obvious. In order to have an objective and unambiguous classification, we utilized the ninth revision of the International Statistical Classification of Diseases and Related Health Problems (ICD-9) (Slee, 1978). This is a widely used system of classifying medical diseases and disorders, dividing them into seventeen chapters representing medical systems/categories such as mental disorders, or disease of the respiratory system. While the 10th version is available, we chose this version based on prior work using it, and the pending release of the 11th version. We will use these ICD9 systems to define the different medical fields.

We determined a CUI’s field of medicine according to a CUI-to-ICD9 dictionary available from the UMLS (Bodenreider, 2004). We consider pharmacological substance related to a field of medicine system if it treats or prevents a disease with an ICD9 code within a particular ICD9 system. We determined this by using the NDF-RT dictionary, which maps CUIs of substances to the CUIs of conditions they treat or prevent, and then convert these CUIs to the ICD9 systems as before. As such, A CUI representing a drug may have multiple ICD9 systems and therefore medical fields.

2.3 Evaluation Methods

We sought to compare multiple methods for evaluating the quality of a medical field’s embeddings based on prior work. We were unable to use Yu et al’s (2017) method, based on comparing the correlation of vector cosine similarity against human judgements from the UNMSRS-Similarity dataset (Pakhomov, 2018) due to there being too few examples across many medical fields. The code for all implemented methods will be publicly available upon publication of this work from the first author’s GitHub.

Medical Relatedness Measure (MRM) This method from Choi et al (2016) is based on quantifying whether concepts with known relations are neighbours of each other. They use known relationships between drugs and the diseases they treat or prevent, and also the relations between diseases that are grouped together in the Clinical Classifications Software (CCS) hierarchical groupings, a classification from the Agency for Healthcare Research and Quality (Cli). The scoring utilizes Discounted Cumulative Gain, which attributes a diminishing score the further away a known relationship is found if within k neighbours.

In our implementation, we calculate the Medical Relatedness Measure (MRM) based on the ‘coarse’ groupings from the CCS hierarchies. Scores are calculated for CUIs that represent diseases with a known ICD9 code. The mean MRM is then calculated for all CUIs within a given ICD9 system. The implementation was adapted from Python 2.7 code available from the original author’s Github. We calculate MRM as:

$$\text{MRM}(V, F, k) = \frac{1}{|V(F)|} \sum_{v \in V(F)} \frac{1}{|V(v)|} \sum_{i=1}^{k} \frac{1}{\log(2^{r+1})}$$

Where $V$ are medical conditions, $F$ a field of medicine, $V(F)$ the medical conditions within an ICD-9 system/field of medicine, $G$ the CCS group that medical condition $v \in V(F)$ is part of, and $V(G)$ the subset of medical conditions found in
### Table 1: Characteristics of the embeddings compared, including the name referred, the embedding dimensions, the number of embeddings in the dataset, and the type of data used to train them.

<table>
<thead>
<tr>
<th>Name</th>
<th>Dimension</th>
<th>Number</th>
<th>Number of Training Data</th>
<th>Type of Training Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>DeVine200</td>
<td>200</td>
<td>52,102</td>
<td>17k + 348k clinical narratives</td>
<td>journal abstracts</td>
</tr>
<tr>
<td>ChoiClaims300</td>
<td>300</td>
<td>14,852</td>
<td>4m health insurance claims</td>
<td></td>
</tr>
<tr>
<td>ChoiClinical300</td>
<td>300</td>
<td>22,705</td>
<td>20m clinical narratives</td>
<td></td>
</tr>
<tr>
<td>BeamCui2Vec500</td>
<td>500</td>
<td>109,053</td>
<td>60m + 20m + 1.7m health insurance claims clinical narratives full journal texts</td>
<td></td>
</tr>
</tbody>
</table>

Table 2: Illustrative example showing how System Vector Accuracy (SysVec) would be calculated for the medical field “Mental Disorders” if it contained only six drugs. Predicted medical field is the medical field/ICD9 system vector closest to the drug, or \( n \) closest fields if a drug treats conditions in \( n \) multiple fields. System vectors are the normalized mean vector of that system’s medical conditions.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Actual Medical Field</th>
<th>Predicted Medical Field</th>
<th>Correct?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluoxetine</td>
<td>Mental Disorders</td>
<td>Mental Disorders</td>
<td>Yes</td>
</tr>
<tr>
<td>Sertraline</td>
<td>Mental Disorders</td>
<td>Neoplasms</td>
<td>No</td>
</tr>
<tr>
<td>Risperidone</td>
<td>Mental Disorders</td>
<td>Mental Disorders</td>
<td>Yes</td>
</tr>
<tr>
<td>Olanzapine</td>
<td>Mental Disorders</td>
<td>Mental Disorders</td>
<td>Yes</td>
</tr>
<tr>
<td>Valproic Acid</td>
<td>Diseases of the Nervous System</td>
<td>Mental Disorders</td>
<td>Congenital Abnormalities</td>
</tr>
<tr>
<td>Lamotrigine</td>
<td>Diseases of the Nervous System</td>
<td>Diseases of the Skin</td>
<td>No</td>
</tr>
</tbody>
</table>

**Mental Disorders SysVec Score: 4/6 = 0.67**

Medical Conceptual Similarity Measure (MCSM) The other method used by Choi et al’s work evaluates whether embeddings known to be of a particular set are clustered together. They use conceptual sets from the UMLS such as ‘pharmacologic substance’ or ‘disease or syndrome’. Discounted Cumulative Gain is again used, based on whether a CUI has other CUIs of its set within \( k \) neighbours.

We reimplement this method, but instead of using the UMLS conceptual sets, we create sets from the ICD9 systems, again giving a score to neighbours that are diseases or drugs from the same field of medicine/ICD9 system. Again, this was adapted from code from Choi et al’s Github. The Medical Conceptual Similarity Measure (MCSM) can be represented as:

\[
\text{MCSM}(V, F, k) = \frac{1}{|V(F)|} \sum_{v \in V(F)} \sum_{i=1}^{k} \frac{1_{F(v(i))}}{\log_2(i+1)}
\]

Similar to MRM, \( F \) is a medical field/ICD9 system, \( V(F) \) the medical conditions within a system, and \( 1_{F} \) 0 or 1 depending on whether neighbour \( v(i) \) is also in this medical field.

For illustration, consider an example calculating the MCSM for the medical field/system (\( F \)) “Infectious and Parasitic Diseases”. This involves calculating the score for the medical condition (\( v \)) primary tuberculosis infection. If rifampin, an an-
tibiotic, was found to be nearby, it would con-
tribute to the MCSM, as it treats conditions in “In-
fectious and Parasitic Diseases” and so would be
classified as being part of this system. On the other
hand, if the respiratory illness asthma was one of
the k nearest neighbours, it would add nothing to
the MCSM score, as it is a disease in a different
system, “Diseases of the Respiratory System”.

**Significance against Bootstrap Distribution**

**Bootstrap** Beam et al (2018) also evaluate
how well known relationships between concepts
are represented by embedding vector similarity.
For a given known relation, they generate a boot-
strap distribution by randomly calculating cosine
similarities between embedding vectors of the
same class (e.g. a random drug and disease when
evaluating drug-disease relations). For a given
known relation, they consider that the embeddings
produced an accurate prediction if their cosine
similarity is within the top 5%, the equivalent of
p < 0.05 for a one-sided t-test.

Our implementation considers the may-treat or
may-prevent known relationships from the NDF-
RT dataset. We calculate the percentage of known
relations for drug-disease pair within each medici-
field. Beam et al have not yet made their code
publicly available, so we reimplemented this tech-
inique in Python.

**System Vector Accuracy (SysVec)** We imple-
ment a new, simple method to evaluate a medici-
field’s embeddings. A representative vector is
calculated for each medical field/ICD9 system
by taking the mean of the normalized embedding
vectors of a field’s diseases. We then consider all
of the drugs known to treat or prevent a disease
of a given medical field. A field’s **System Vec-
tor Accuracy** is then the percentage of these
drugs whose most similar (by cosine similarity) system
vector is this field’s. A higher score indicates bet-
ter performance. We implemented this method in
Python.

For example, a system vector for “Mental Dis-
orders” would be calculated from the embeddings
during diseases such as schizophrenia and major de-
pressive disorder. “Mental Disorders” **System
Vector Accuracy** is the percentage of its medi-
cations (e.g. fluoxetine, risperidone, paroxetine)
whose embedding vectors are more similar to the
“Mental Disorders” system vector than all others.
Fluoxetine is an anti-depressant medication solely
used to treat “Mental Disorders”, so we would ex-
pect its vector to be more similar to this system
vector than, say, the system vector representing
“Diseases of the Skin and Subcutaneous Tissue”.

Some drugs treat or prevent diseases in n mul-
iple medical field. For a field, such a drug is clas-
sified as being accurately predicted if the field’s
system vector is amongst the n most similar sys-
tem vectors. For instance, valproic acid is an anti-
convulsant used to treat both mental disorders and
those of the nervous system. “Mental Disorders”’
**System Vector Accuracy** would take into account
whether its system vector was one of the n=2 most
similar system vectors. For further illustration, Ta-
ble 2 shows an example SysVec calculation.

### 2.4 Comparing Scores

**Comparing Sets of Embeddings** We calculated
the mean scores for an embedding set, only includ-
ing embeddings with corresponding ICD9 values
and present in all of the compared sets. For the
MCSM and MRM scores, we conducted two-way
paired t-tests between the scores from each em-
bedding set, adjusted with the Bonferroni correc-
tion. For the binary Bootstrap and SysVec scores,
we judged statistical significance by calculating
z-scores and their corresponding Bonferroni cor-
corrected p-values.

A negative control set of embeddings was con-
structed by taking the embeddings from Beam et
al (2018) and randomly arranging which clinical
concepts an embedding corresponds to.

**Comparing Fields of Medicine** As the embed-
dings from Beam et al (2018) are most recent,
trained on the most data, and have significantly
higher scores than the other embeddings com-
pared, we used these embeddings to compare
scores from the different fields of medicine. This
set also contained the most embeddings, allowing
more embeddings from each field to be compared.

We sought to determine whether a field of
medicine’s embeddings were significantly worse
or better than the average. As such, for each field
of medicine we calculated the mean score from
each evaluation method. We then used statisti-
cal tests to compare a field’s scores from a given
evaluation method with the same scores from all
other fields. For MCSM and MRM scores we used
two-tailed t-tests, and for Bootstrap and SysVec,
z-scores, all corrected with the Bonferroni correc-
tion.
To aggregate a medical field’s results, we calculated a ‘Net Significance’ metric by taking how many of the four method’s scores were significantly above the mean, minus how many were significantly below. We found this more interpretable than other methods such as aggregating normalized scores.

3 Results

3.1 Differences Between Sets of Embeddings
Comparing the sets of embeddings (Table 3) shows consistent differences. BeamCui2Vec500’s scores are the highest across all methods, and this difference is very significant, with p-value $\ll 10^{-5}$ after Bonferroni correction. The ChoiClaims300 embeddings seem next best, and the remaining sets still have much higher scores than those of the negative control.

3.2 Differences Between Medical Systems
Differences are also observed between embeddings from the various fields of medicine as represented by the ICD-9 systems (Table 4). For instance, embeddings related to the medical field Mental Disorders have scores significantly above the mean score across all systems for two evaluation methods, while those of the field Symptoms, Signs, and Ill-defined Conditions are significantly below for three. Due to a smaller number of documented drug-disease relationships across two medical fields, scores were not calculated with those methods using these relationships.

4 Discussion and Future Direction

To our knowledge, this is the first investigation into whether clinical concept embeddings from a given field of medicine perform relatively well or poorly compared to others. We conducted this investigation comparing available sets of such embeddings, using a variety of previously described intrinsic evaluation methods in addition to a new one. Given that one set of embeddings performed better than others, we used this set to compare the different fields of medicine, and found significant results between various fields.

The superior performance of one set of embeddings - those from Beam et al (2018) - are consistent with the depth and breadth of data used to train these embeddings. Training used three different types of data, including that from health insurance claims, clinical narratives, and full texts from medical journals. The size of the dataset was also much larger than that of the others. Our work validates their findings that their embeddings offer the best performance. However, it would be interesting to also consider the recent clinical concept embeddings developed by (Xiang et al., 2019). They use a similar amount of data (50 million) as Beam et al, using a large dataset from electronic health records, and apply a novel method to incorporate time-sensitive information. At the time of submission, we were unable to obtain their embeddings, and so leave this comparison to future work.

Examining the differences between fields of medicine, we note that the poor performance of embeddings from the system “Symptoms, Signs, and Ill-defined Conditions” may support validity of the results. This collection of miscellaneous medical conditions would not be expected to have the intrinsic vector similarity and cohesion evaluated by our evaluation methods.

Further work may explore why the other systems have varied performance. We wonder if the observed results correlate with possible distinction of the various medical fields. For example, one of the best performing systems was “Neoplasms”. The conditions in this field are often unambiguous - a cancer like non-small cell lung cancer has little other meaning - and the drugs used for these diseases tend to be similarly specific. On the other hand, poorly performing systems such as “Diseases of the Skin and Subcutaneous Tissue” and ”Diseases of the Musculoskeletal Systems and Connective Tissue” often utilize immunosuppressant medications that are used across many fields of medicine. Future work could investigate this conjecture by comparing scores when restricting what clinical concepts are compared, such as only common or distinct medications.

This work evaluated embeddings using intrinsic measures of embedding quality. This presents some advantages, but also the most obvious limitation and direction for future work. These intrinsic methods allowed a consistent evaluation to be carried out between medical fields, and allowed a wide variety of embedding sets to be compared. The methods all evaluate qualities that well-trained embeddings should have, though still represent artificial use-cases. Evaluating these embeddings on extrinsic, down-stream tasks may provide more practically relevant comparisons. However, these tasks will need to be comparable...
<table>
<thead>
<tr>
<th>Embedding Set</th>
<th>MRM</th>
<th>MCSM</th>
<th>Bootstrap</th>
<th>SysVec</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative Control</td>
<td>0.02</td>
<td>1.24</td>
<td>0.05</td>
<td>0.35</td>
</tr>
<tr>
<td>DeVine200</td>
<td>0.24</td>
<td>5.14</td>
<td>0.27</td>
<td>0.79</td>
</tr>
<tr>
<td>ChoiClaims300</td>
<td>0.43</td>
<td>5.34</td>
<td>0.42</td>
<td>0.80</td>
</tr>
<tr>
<td>ChoiClinical300</td>
<td>0.33</td>
<td>4.49</td>
<td>0.42</td>
<td>0.74</td>
</tr>
<tr>
<td>BeamCui2Vec500</td>
<td>0.52</td>
<td>6.39</td>
<td>0.67</td>
<td>0.90</td>
</tr>
</tbody>
</table>

Table 3: Mean scores for embedding sets for each evaluation method. See Methods section for abbreviations.

<table>
<thead>
<tr>
<th>ICD-9 Systen</th>
<th>MRM</th>
<th>MCSM</th>
<th>Bootstrap</th>
<th>SysVec</th>
<th>Net Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>All Systems (Negative Control)</td>
<td>0.05</td>
<td>1.08</td>
<td>0.04</td>
<td>0.25</td>
<td>-</td>
</tr>
<tr>
<td>All Systems</td>
<td>0.55</td>
<td>8.07</td>
<td>0.89</td>
<td>0.63</td>
<td>-</td>
</tr>
<tr>
<td>Infectious and Parasitic Diseases</td>
<td>0.45</td>
<td>7.72</td>
<td>0.93</td>
<td>0.92</td>
<td>0</td>
</tr>
<tr>
<td>Neoplasms</td>
<td>0.62</td>
<td>9.69</td>
<td>0.94</td>
<td>0.55</td>
<td>+2</td>
</tr>
<tr>
<td>Endocrine, Nutritional and Metabolic Diseases, and Immunity Disorders</td>
<td>0.44</td>
<td>5.64</td>
<td>0.89</td>
<td>0.53</td>
<td>-2</td>
</tr>
<tr>
<td>Diseases of the Blood and Blood-forming Organs</td>
<td>0.31</td>
<td>4.36</td>
<td>0.82</td>
<td>0.79</td>
<td>-2</td>
</tr>
<tr>
<td>Mental Disorders</td>
<td>0.53</td>
<td>9.34</td>
<td>0.96</td>
<td>0.83</td>
<td>+2</td>
</tr>
<tr>
<td>Diseases of the Nervous System and Sense Organs</td>
<td>0.76</td>
<td>8.44</td>
<td>0.87</td>
<td>0.33</td>
<td>+1</td>
</tr>
<tr>
<td>Diseases of the Circulatory System</td>
<td>0.59</td>
<td>8.12</td>
<td>0.96</td>
<td>0.72</td>
<td>+2</td>
</tr>
<tr>
<td>Diseases of the Respiratory System</td>
<td>0.36</td>
<td>5.85</td>
<td>0.94</td>
<td>0.82</td>
<td>+1</td>
</tr>
<tr>
<td>Diseases of the Digestive System</td>
<td>0.61</td>
<td>7.93</td>
<td>0.77</td>
<td>0.62</td>
<td>0</td>
</tr>
<tr>
<td>Diseases of the Genitourinary System</td>
<td>0.61</td>
<td>6.82</td>
<td>0.86</td>
<td>0.58</td>
<td>0</td>
</tr>
<tr>
<td>Complications of Pregnancy, Childbirth, and the Puerperium</td>
<td>0.51</td>
<td>10.27</td>
<td>-</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>Diseases of the Skin and Subcutaneous Tissue</td>
<td>0.37</td>
<td>5.1</td>
<td>0.81</td>
<td>0.58</td>
<td>-2</td>
</tr>
<tr>
<td>Diseases of the Musculoskeletal System and Connective Tissue</td>
<td>0.47</td>
<td>8.22</td>
<td>0.88</td>
<td>0.29</td>
<td>-2</td>
</tr>
<tr>
<td>Congenital Anomalies</td>
<td>0.5</td>
<td>6.24</td>
<td>0.73</td>
<td>0.73</td>
<td>-1</td>
</tr>
<tr>
<td>Certain Conditions Originating in the Perinatal Period</td>
<td>0.48</td>
<td>9.84</td>
<td>-</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>Symptoms, Signs, and Ill-defined Conditions</td>
<td>0.26</td>
<td>2.68</td>
<td>0.77</td>
<td>0.56</td>
<td>-3</td>
</tr>
<tr>
<td>Injury and Poisoning</td>
<td>0.59</td>
<td>9.09</td>
<td>0.75</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 4: Comparison of mean scores using different evaluation methods for the fields of medicine as represented by their ICD-9 system. The row All Systems shows the mean score for each method across embeddings from all systems. A bold score indicates that a system’s score was significantly above the All Systems score, while an italic score indicates it was below. Significance is judged by having a p-value <0.05 after Bonferroni correction. Net Significance is the number of these significant differences above the All Systems score minus the number below. A system’s score is not calculated if there are fewer than ten examples for a method. See Methods section for evaluation method abbreviations. All scores in this table are calculated using the embeddings from Beam et al.

and available for multiple medical fields. For instance, the recent work by Xiang et al (2019) compared embeddings trained by different methodologies on a task predicting the onset of heart failure (Rasmy et al., 2018). This would be an appropriate task to judge embeddings from “Diseases of the Circulatory System”; others would be needed for other systems. We also plan to investigate the validity of these intrinsic evaluation methods by comparing them to extrinsic results.
Another future direction could be to investigate what could be done to improve performance in the fields with lower scores. For instance, Zhang et al (2018) used domain adaptation techniques for psychiatric embeddings, and this could also be carried out for those systems we identified as doing poorly. Alternatively, one could train embeddings solely on data from one field of medicine and investigate how this affects performance.

References

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José Antonio Minarro-Giménez, Oscar Marín-Alonso, and Matthias Samwald. 2014. Exploring the application of deep learning techniques on medical text corpora. Studies In Health Technology And Informatics, 205:584–588.


On the Effectiveness of the Pooling Methods for Biomedical Relation Extraction with Deep Learning

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Abstract

Deep learning models have achieved state-of-the-art performances on many relation extraction datasets. A common element in these deep learning models involves the pooling mechanisms where a sequence of hidden vectors is aggregated to generate a single representation vector, serving as the features to perform prediction for RE. Unfortunately, the models in the literature tend to employ different strategies to perform pooling for RE, leading to the challenge to determine the best pooling mechanism for this problem, especially in the biomedical domain. In order to answer this question, in this work, we conduct a comprehensive study to evaluate the effectiveness of different pooling mechanisms for the deep learning models in biomedical RE. The experimental results suggest that dependency-based pooling is the best pooling strategy for RE in the biomedical domain, yielding the state-of-the-art performance on two benchmark datasets for this problem.

1 Introduction

In order to analyze the entities in text, it is crucial to understand how the entities are related to each other in the documents. In the literature, this problem is formalized as relation extraction (RE), an important task in information extraction. RE aims to identify the semantic relationships between two entity mentions within the same sentences in text. Due to its important applications on many areas of natural language processing (e.g., question answering, knowledge base construction), RE has been actively studied in the last decade, featuring a variety of feature-based or kernel-based models for this problem (Zelenko et al., 2002; Zhou et al., 2005; Bunescu and Mooney, 2005; Sun et al., 2011; Chan and Roth, 2010; Nguyen et al., 2009). Recently, the introduction of deep learning has produced a new generation of models for RE with the state-of-the-art performance on many different benchmark datasets (Zeng et al., 2014; dos Santos et al., 2015; Xu et al., 2015; Liu et al., 2015; Zhou et al., 2016; Wang et al., 2016; Zhang et al., 2017, 2018b). The advantage of deep learning over the previous approaches for RE is the ability to automatically learn effective features for the sentences from data via various network architectures. The same trend has also been observed for RE in the biomedical domain where deep learning is gaining more and more attention from the research community (Mehryary et al., 2016; Björne and Salakoski, 2018; Nguyen and Verspoor, 2018; Verga et al., 2018).

The typical deep learning models for RE have involved Convolutional Neural Networks (CNN) (Zeng et al., 2014; Nguyen and Grishman, 2015b; Zeng et al., 2015; Lin et al., 2016; Zeng et al., 2017), Recurrent Neural Networks (RNN), (Miwa and Bansal, 2016; Zhang et al., 2017), Transformer (self-attention) networks (Verga et al., 2018), and Graph Convolutional Neural Networks (GCNN) (Zhang et al., 2018b). There are two major common components in such deep learning models for RE, i.e., the representation component and the pooling component. First, in the representation component, some deep learning architectures are employed to compute a sequence of vectors to represent an input sentence for RE for which each vector tends to capture the specific context information for a word in that sentence. Such word-specific representation sequence is then fed into the second pooling component (e.g., max pooling) that aggregates the representation vectors to obtain an overall vector to represent the whole input sentence for the classification problem in RE.

While there have been many work in the literature to compare different deep learning architectures for the representation component, the pos-
sible methods for the pooling component of the deep learning models have not been systematically benchmarked for RE in general and for the biomedical domain in particular. Specifically, the prior work on relation extraction with deep learning has only assumed one form of pooling in the model without considering the possible alternatives for this component. In this work, we argue that the pooling mechanisms also have significant impact on the performance of the deep learning models for RE and it is important to understand how well different pooling methods perform in this case. Consequently, in this work, we conduct a comprehensive investigation on the effectiveness of different max pooling methods for the deep learning models of RE, focusing on the biomedical domain as the case study. Our goal is to determine the best pooling methods for the deep learning models in biomedical RE. We also want to emphasize the experiments where the pooling methods are compared in a compatible manner with the same representation components and resources for the biomedical RE models in this work. Such compatible comparison is unfortunately very rare in the current literature about deep learning for RE as new models are being intensively proposed, employing a diversity of options and resources (i.e., pre-trained word embeddings, optimizers, etc.). Therefore, this is actually the first work to compare different pooling methods for deep relation extraction on the same setting.

In the experiments, we find that syntactic information (i.e., dependency parsing) can be exploited to provide the best pooling strategies for biomedical RE. In fact, our experiments also suggest that it is more beneficial to apply the syntactic information in the pooling component of the deep learning models for biomedical RE than that in the representation component. This is different from most of the prior work on relation extraction that has only employed the syntactic information in the representation component of the deep learning models (Xu et al., 2016; Miwa and Bansal, 2016). Based on the syntax-based pooling mechanism, we achieve the state-of-the-art performance on two benchmark datasets for biomedical RE.

2 Model

Relation Extraction can be seen as a multi-class classification problem that takes a sentence and two entity mentions of interest in that sentence as the input. The goal is to predict the semantic relation between these two entity mentions according to some predefined set of relations. Formally, let \( W = [w_1, w_2, \ldots, w_n] \) be the input sentence where \( n \) is the number of tokens and \( w_i \) is the \( i \)-th word/token in \( W \). As entity mentions can span multiple consecutive words/tokens, let \([s_1, e_1]\) be the span of the first entity mention \( M_1 \) where \( s_1 \) and \( e_1 \) are the indexes for the first and last token of \( M_1 \) respectively. Similarly, we define \([s_2, e_2]\) as the span for the second entity mention \( M_2 \). For convenience, we assume that the entity mentions are not nested, i.e., \( 1 \leq s_1 \leq e_1 < s_2 \leq e_2 \leq n \).

2.1 Input Vector Representation

In order to encode the positions and the entity types of the two entity mentions in the input sentence, following (Zhang et al., 2018b), we first replace the tokens in the entity mentions \( M_1 \) and \( M_2 \) with the special tokens of format \( \text{M}_1\text{-Type}_1 \) and \( \text{M}_2\text{-Type}_2 \) respectively (\( \text{Type}_1 \) and \( \text{Type}_2 \) represent the entity types of \( M_1 \) and \( M_2 \) respectively). The purpose of this replacement is to help the models to abstract from the specific tokens/words of the entity mentions and only focus on their positions and entity types, the two most important pieces of information of the entity mentions for RE.

Given the enriched input sentence, the first step in the deep learning models for RE is to convert each word in the input sentence into a vector to facilitate the real-valued computation of the models. In this work, the vector \( v_i \) for \( w_i \) is obtained by concatenating the following two vectors:

1. The word embeddings of \( w_i \): The embeddings for the special tokens are initialized randomly while the embeddings for the other words are retrieved from the pre-trained word embedding table provided by the Word2Vec toolkit with 300 dimensions (Mikolov et al., 2013).

2. The embeddings for the part-of-speech (POS) tag of \( w_i \) in \( W \): We assign a POS tag for each word in the input sentence using the Stanford CoreNLP toolkit. The embedding for each POS tag is also randomly initialized in this case.

Note that both the word embeddings and the POS embeddings are updated during the training time of the models in this work. The word-to-vector conversion transforms the input sentence \( W = [w_1, w_2, \ldots, w_n] \) into a sequence of vectors \( V = [v_1, v_2, \ldots, v_n] \) (respectively) that would be used as the input for all the deep learning mod-
els considered in this work to ensure a compatible comparison. As mentioned in the introduction, the deep learning models for RE involves two major components, i.e., the representation component and the pooling component. We describe the options for such components in the following sections.

### 2.2 The Representation Component for RE

Given the input sequence of vectors $V = [v_1, v_2, \ldots, v_n]$, the next step in the deep learning models for RE is to transform this vector sequence into a more abstract vector sequence $A = [a_1, a_2, \ldots, a_n]$ so $a_i$ would capture the underlying representation for the context information specific to the $i$-th word in the sentence. In this work, we examine the following typical architectures to obtain such an abstract sequence $A$ for $V$:

1. **CNN** (Zeng et al., 2014; Nguyen and Grishman, 2015b; dos Santos et al., 2015): CNN is one of the early deep learning models for RE. It involves an 1D convolution layer over the input vector sequence $V$ with multiple window sizes for the filters. CNN produces a sequence of vectors in which each vector contains some $n$-grams specific to a word in the sentence. This sequence of vectors is used as $A$ for our purpose.

2. **BiLSTM** (Nguyen and Grishman, 2015a): In **BiLSTM**, two Long-short Term Memory Networks (LSTM) are run over the input vector sequence $V$ in the forward and backward direction. The hidden vectors generated at the position $i$ by the two networks are then concatenated to constitute the abstract vector $a_i$ for this position. Due to the recurrent nature, $a_i$ involves the context information over the whole input sentence $W$ although a greater focus is put on the context of the current word.

3. **BiLSTM-CNN**: This model resembles the MASS model presented in (Le et al., 2018). It first applies a bidirectional LSTM layer over the input sequence $V$ whose results are further processed by a Convolutional Neural Network (CNN) layer as in CNN. We also use the output of the CNN layer as the abstract vector sequence $A$ for this model.

4. **BiLSTM-GCNN** (Zhang et al., 2018b): Similar to BiLSTM-CNN, BiLSTM-GCNN also first employs a bidirectional LSTM network to abstract the input vector sequence $V$. However, in the second step, different from BiLSTM-CNN, BiLSTM-GCNN introduces a Graph Convolutional Neural Network (GCNN) layer that consumes the LSTM hidden vectors and augments the representation for a word with the representation vectors of the surrounding words in the dependency trees. The output of the GCNN layer is also a sequence of vectors to represent the contexts for the words in the sentence and functions as the abstract sequence $A$ in our case. **BiLSTM-GCNN** (Zhang et al., 2018b) is one of the current state-of-the-art models for RE in the literature.

Note that there are many other variants of such models for RE in the literature (Xu et al., 2016; Zhang et al., 2017; Verga et al., 2018). However, as our goal in this paper is to evaluate different pooling mechanisms for RE, we focus on these standard representation learning methods to avoid the confounding effect of the complicated models, thus better revealing the effectiveness of the pooling methods.

### 2.3 The Pooling Component for RE

The goal of the pooling component is to aggregate the representation vectors in the abstract sequence $A$ to constitute an overall vector $F$ to represent the whole input sentence $W$ and the two entity mentions of interest (i.e., $F = \text{aggregate}(A)$). The overall representation vector should be able to capture the most important features induced in $A$. The typical method to achieve such aggregation in the RE models is to apply the element-wise max-pooling operation over subsets of vectors in $A$ whose results are combined to obtain the overall representation vector. While there are different methods to select the vector subsets for the max-pooling operation, the prior work for RE has only employed one particular selection method in their deep learning models (Nguyen and Grishman, 2015a; Zhang et al., 2018b; Le et al., 2018). This raises the question about the impact of the other subset selection methods for such prior RE models. Can these methods benefit from different pooling mechanisms? What are the best pooling methods for the deep learning models in RE? In order to answer these questions, besides the architectures for the representation component in the previous section, we investigate the following subset selection methods for the pooling component of the RE models in this work:

1. **ENT-ONLY**: In this pooling method, we use the subsets of the vectors corresponding to the words in the two entity mentions of interest in
for the max-pooling operations (i.e., $M_1$ with the words in the range $[s_1, e_1]$ and $M_2$ with the words in the range $[s_2, e_2]$). This is motivated by the utmost importance of the two entity mentions of interest for RE and employed in some prior work (Nguyen and Grishman, 2015a; Zhang et al., 2018b):

\[
F_{M_1} = \text{max-pool} (a_{s_1}, a_{s_1+1}, \ldots, a_{e_1}) \\
F_{M_2} = \text{max-pool} (a_{s_2}, a_{s_2+1}, \ldots, a_{e_2}) \\
F_{\text{ENT-ONLY}} = [F_{M_1}, F_{M_2}]
\]

2. ENT-SENT: Besides the entity mentions, the other context words in the sentence might also involve important information for the relation prediction in RE. For instance, in the sentence “Acetazolamide can elevate cyclosporine levels.”, the context word “elevate” is crucial to determine the semantic relations between the two entity mentions of interest “Acetazolamide” and “cyclosporine”. In order to capture such important contexts for pooling, the typical approach in the prior work for RE is to perform the max-pooling operation over the abstract vectors for every word in the sentence (i.e., the whole set $A$) (Zeng et al., 2014; dos Santos et al., 2015; Le et al., 2018). The rationale is to select the features of the abstract vectors in $A$ with the highest values in each dimension to reveal the most important context for RE. The max-pooled vector over the whole set $A$ is combined with the $F_{\text{ENT-ONLY}}$ vector in this method:

\[
F_{\text{SENT}} = \text{max-pool} (a_1, a_2, \ldots, a_n) \\
F_{\text{ENT-SENT}} = [F_{\text{ENT-ONLY}}, F_{\text{SENT}}]
\]

3. ENT-DYM: Similar to ENT-SENT, this method also seeks the important context information beyond the two entity mentions of interest. However, instead of taking the whole vector sequence $A$ for the pooling, ENT-DYM divides $A$ into three separate vector subsequences based on the start and end indexes of the first and second entity mentions (i.e., $s_1$ and $e_2$) respectively. The max-pooling operation is then applied over these three subsequences and the resulting vectors are combined to form an overall vector (i.e., dynamic pooling) (Zeng et al., 2015):

\[
F_{\text{LEFT}} = \text{max-pool} (a_1, a_2, \ldots, a_{s_1-1}) \\
F_{\text{MIDDLE}} = \text{max-pool} (a_{s_1}, a_{s_1+1}, \ldots, a_{e_2}) \\
F_{\text{RIGHT}} = \text{max-pool} (a_{e_2+1}, a_{e_2+2}, \ldots, a_n) \\
F_{\text{ENT-DYM}} = [F_{\text{LEFT}}, F_{\text{MIDDLE}}, F_{\text{RIGHT}}, F_{\text{ENT-ONLY}}]
\]

4. ENT-DEP0: The previous pooling methods have only relied on the sequential structures of the sentence where the chosen subsets of $A$ for pooling always contain vectors for the consecutive words in the sentence. Unfortunately, such sequential pooling might introduce irrelevant words into the selected subsets of $A$, potentially causing noise in the pooling features and impeding the performance of the RE models. For instance, in the previous sentence example “Acetazolamide can elevate cyclosporine levels.”, the ENT-SENT and ENT-DYM methods would also include the word “levels” in the pooling subsets that is not very important for the relation prediction in this case. Consequently, in ENT-DEP0, we explore the possibility to use the dependency parse tree of the input sentence $W$ to filter out the irrelevant words for the pooling operation. In particular, instead of considering every word in the input sentence, ENT-DEP0 only pools over the abstract vectors in $A$ that correspond to the words along the shortest dependency path (SDP) between the two entity mentions $M_1$ and $M_2$ in the dependency tree for $W$ (called $SDP_0(M_1, M_2)$). Note that the shortest dependency paths have been shown to be able to select the important context words for RE in many previous work (Zhou et al., 2005; Chan and Roth, 2010; Xu et al., 2016). Similar to ENT-SENT and ENT-DYM, we also include $F_{\text{ENT-ONLY}}$ in this method:

\[
F_{\text{DEP0}} = \text{max-pool}_{a_i \in SDP_0(M_1, M_2)} (a_i) \\
F_{\text{ENT-DEP0}} = [F_{\text{DEP0}}, F_{\text{ENT-ONLY}}]
\]

5. ENT-DEP1: This method is similar to ENT-DEP0. However, instead of directly pooling over the words in the shortest dependency path $SDP_0(M_1, M_2)$, ENT-DEP1 extends this path to also include every word that is connected to some word in $SDP_0(M_1, M_2)$ via an edge in the dependency tree for $W$ (i.e., one edge distance from $SDP_0(M_1, M_2)$). We denote this extended word set by $SDP_1(M_1, M_2)$ for which the corresponding abstract vectors in $A$ would be chosen for
the max-pooling operation. The motivation for $SDP_1(M_1, M_2)$ is that the representations of the words close to the shortest dependency path between $M_1$ and $M_2$ might also provide useful information to improve the performance for RE. In our experiments, we find that one edge is the optimal distance to enlarge the shortest dependency paths. Using larger distance for the pooling mechanism would hurt the performance of the deep learning models for RE:

$$F_{DEP1} = \max_{a_i \in SDP_1(M_1, M_2)} (a_i)$$

$$F_{ENT-DEP1} = [F_{DEP1}, F_{ENT-ONLY}]$$

Once the overall representation vector $F$ for the input sentence $W$ and the two entity mentions of interest has been produced, we feed it into a feed-forward neural network with a softmax layer in the end to obtain the probability distribution $P(y|W, M_1, M_2) = \text{feed-forward}(F)$ over the possible relation types for our RE problem. This probability distribution would then be used for both making prediction (i.e., by taking the relation type with the highest probability) and training models (i.e., by optimizing the negative log-likelihood function).

3 Experiments

3.1 Datasets

In order to evaluate the performance of the models in this work, we employ the following biomedical datasets for RE in the experiments:

DDI-2013 (Herrero-Zazo et al., 2013): This dataset contains 730 documents from the Drugbank database, involving about 25,000 examples for the training and test sets (each example consists of a sentence and two entity mentions of interest for classification). There are 4 entity types (i.e., drug, brand, group and brand+n) and 5 relation types (i.e., mechanism, advise, effect, int, and no_relation) in this dataset. The no_relation is to indicate any example that does not belong to any relation types of interest. This dataset is severely imbalanced, containing 85% negative examples in the training dataset. In order to deal with such imbalanced data, we employ weighted sampling that equally distributes the selection probability for the positive and negative examples.

BB3 (Deléger et al., 2016). This dataset contains 95 documents; each of them involves a title and abstract from a document from the PubMed database. There are 800 examples in this dataset divided into two separate sets (i.e., the training set and the validation set). BB3 also include a test set; however, the relation types for the examples in this test set are not provided. In order to obtain the performance of the models on the test set, the performers need to submit their system outputs to an official API that would evaluate the output and return the model performance. We train the models in this work on the training data and employ the official API to obtain their test set performance to be reported in the experiments for this dataset.

Following the prior work on these datasets (Chowdhury and Lavelli, 2013; Lever and Jones, 2016; Zhou et al., 2018; Le et al., 2018), we use the micro-averaged F1 scores as the performance measure in the experiments to ensure a compatible comparison.

3.2 Parameters and Resources

As the DDI-2013 dataset does not involve a development set, we tune the parameters for the models in this work based on the validation data of the BB3 dataset and use the selected parameters for both datasets in the experiments. The best parameters from this tuning process include the learning rate of 0.5 and momentum of 0.8 for the stochastic gradient descent (SGD) optimizer with nesterov’s momentum to optimize the models. In order to regularize the models, we apply dropout between layers with the drop rate for word embeddings set to 0.7 and other drop rates set to 0.5. We also employ the weight dropout DropConnect in (Wan et al., 2013) to regularize the hidden-to-hidden transition matrix within each bidirectional LSTM in the models (Merity et al., 2017). For all the models that involve bidirectional LSTMs (i.e., BiLSTM, BiLSTM-CNN, and BiLSTM-GCNN), two layers of bidirectional LSTMs are utilized with 300 hidden units for each LSTM network. For the models with CNN components (i.e., CNN and BiLSTM-CNN), we use one CNN layer with multiple window sizes of 2, 3, 4, and 5 for the filters (200 filters for each window size). For the BiLSTM-GCN model, two GCNN layers are employed with 300 hidden units in each layer. Finally, for the final feed-forward neural network to compute the probability distribution (i.e., feed-forward), we utilize two hidden layers for which 1000 hidden units are used for the first layer and the number of hidden units for the sec-
ond layer is determined by the number of relation types in the datasets.

3.3 Evaluating the Pooling Methods for RE

This section evaluates the performance of different pooling methods when they are applied to the deep learning models for RE on the two datasets DDI-2013 and BB3. In particular, we integrate each of the pooling methods in Section 2.3 (i.e., ENT-ONLY, ENT-SENT, ENT-DYM, END-DEP0, and END-DEP1) into each of the deep learning models in Section 2.2 (i.e., CNN, BiLSTM, BiLSTM-CNN, and BiLSTM-GCN), resulting in 20 different model combinations to be investigated in this section. For each model combination, we train five versions of the model with different random seeds for parameter initialization over the training datasets. The performance of such versions over the test sets is averaged to serve as the overall model performance on the corresponding dataset. Tables 1 and 2 report the performance of the models on the DDI-2013 dataset and BB3 dataset respectively.

<table>
<thead>
<tr>
<th>Model</th>
<th>P</th>
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<th>F1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CNN</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>52.7</td>
<td>43.1</td>
<td>47.4</td>
</tr>
<tr>
<td>+ ENT-SENT</td>
<td>75.8</td>
<td>60.7</td>
<td>67.3</td>
</tr>
<tr>
<td>+ ENT-DYM</td>
<td>66.5</td>
<td>70.6</td>
<td>68.5</td>
</tr>
<tr>
<td>+ ENT-DEP0</td>
<td>59.8</td>
<td>61.5</td>
<td>60.6</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>67.6</td>
<td>65.1</td>
<td>66.3</td>
</tr>
<tr>
<td><strong>BiLSTM</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>74.0</td>
<td>69.4</td>
<td>71.6</td>
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<tr>
<td>+ ENT-SENT</td>
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<td>73.1</td>
</tr>
<tr>
<td>+ ENT-DYM</td>
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<td>73.4</td>
<td>72.4</td>
</tr>
<tr>
<td>+ ENT-DEP0</td>
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<td>69.4</td>
<td>71.1</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
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<td>76.4</td>
<td>73.9</td>
</tr>
<tr>
<td><strong>BiLSTM-CNN</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>69.6</td>
<td>72.3</td>
<td>70.9</td>
</tr>
<tr>
<td>+ ENT-SENT</td>
<td>69.4</td>
<td>74.9</td>
<td>72.0</td>
</tr>
<tr>
<td>+ ENT-DYM</td>
<td>71.0</td>
<td>69.7</td>
<td>71.8</td>
</tr>
<tr>
<td>+ ENT-DEP0</td>
<td>72.2</td>
<td>69.5</td>
<td>70.8</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>71.0</td>
<td>74.3</td>
<td>72.6</td>
</tr>
<tr>
<td><strong>BiLSTM-GCN</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
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<td>71.4</td>
<td>70.4</td>
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<td>71.1</td>
<td>70.6</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>72.7</td>
<td>72.9</td>
<td>72.8</td>
</tr>
</tbody>
</table>

Table 1: Results on DDI 2013

<table>
<thead>
<tr>
<th>Model</th>
<th>P</th>
<th>R</th>
<th>F1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CNN</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>54.2</td>
<td>65.7</td>
<td>59.1</td>
</tr>
<tr>
<td>+ ENT-SENT</td>
<td>55.0</td>
<td>62.5</td>
<td>59.1</td>
</tr>
<tr>
<td>+ ENT-DYM</td>
<td>54.6</td>
<td>53.3</td>
<td>53.5</td>
</tr>
<tr>
<td>+ ENT-DEP0</td>
<td>55.9</td>
<td>65.8</td>
<td>60.6</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>55.7</td>
<td>67.7</td>
<td>61.1</td>
</tr>
<tr>
<td><strong>BiLSTM</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>58.9</td>
<td>59.6</td>
<td>59.2</td>
</tr>
<tr>
<td>+ ENT-SENT</td>
<td>60.7</td>
<td>59.2</td>
<td>59.9</td>
</tr>
<tr>
<td>+ ENT-DYM</td>
<td>50.2</td>
<td>66.0</td>
<td>56.9</td>
</tr>
<tr>
<td>+ ENT-DEP0</td>
<td>51.6</td>
<td>78.0</td>
<td>61.9</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>54.7</td>
<td>72.6</td>
<td>62.4</td>
</tr>
<tr>
<td><strong>BiLSTM-CNN</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>56.4</td>
<td>66.2</td>
<td>60.8</td>
</tr>
<tr>
<td>+ ENT-SENT</td>
<td>53.6</td>
<td>69.2</td>
<td>60.5</td>
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<tr>
<td>+ ENT-DYM</td>
<td>47.1</td>
<td>78.0</td>
<td>58.7</td>
</tr>
<tr>
<td>+ ENT-DEP0</td>
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<td>71.4</td>
<td><strong>62.5</strong></td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>54.1</td>
<td>74.7</td>
<td>62.4</td>
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<tr>
<td><strong>BiLSTM-GCN</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>+ ENT-ONLY</td>
<td>62.7</td>
<td>56.1</td>
<td>58.9</td>
</tr>
<tr>
<td>+ ENT-SENT</td>
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<td>58.7</td>
<td>58.5</td>
</tr>
<tr>
<td>+ ENT-DYM</td>
<td>56.8</td>
<td>58.4</td>
<td>56.6</td>
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<tr>
<td>+ ENT-DEP0</td>
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<td>67.4</td>
<td>60.8</td>
</tr>
<tr>
<td>+ ENT-DEP1</td>
<td>54.4</td>
<td>71.1</td>
<td>61.5</td>
</tr>
</tbody>
</table>

Table 2: Results on BioNLP BB3

From the tables, we have the following observations about the effectiveness of the pooling methods for RE with deep learning:

1. Comparing ENT-SENT, ENT-DYM and ENT-ONLY, we see that the pooling methods over the whole sentence (i.e., ENT-SENT and ENT-DYM) are significantly better than ENT-ONLY that only focuses on the two entity mentions of interest in the DDI-2013 dataset. This is true across different deep learning models in this work. However, this comparison is reversed for the BB3 dataset where ENT-ONLY is in general better or comparable to ENT-SENT and ENT-DYM over different deep learning models. We attribute such phenomena to the fact that the BB3 dataset often contains many entity mentions and relations within a single sentence (i.e., overlapping contexts) while the sentences in DDI-2013 tend to involve only a single relation with few entity mentions. This makes ENT-SENT and ENT-DYM ineffective for BB3 as the pooling mechanisms over the whole sentence are likely to involve the contexts for the other entity mentions and relations in the sentences, causing the low quality of the resulting representations and the confusion of the model for the relation prediction. This problem is less severe in DDI-2013 as the context of the whole sentence (with a single relation) is more aligned with the important context for the relation prediction. We call the many entity mentions and relations in a single sentence of BB3 as the multiple relation effect for convenient discussion in this paper.

2. Comparing ENT-SENT and ENT-DYM, their performance are comparable in DDI-2013 (except for CNN where ENT-DYM is better); however, in the BB3 dataset, ENT-SENT significantly outperforms ENT-DYM over all the models. This suggests the amplification of the multiple relation ef-
fect in BB3 due to $\textit{ENT-DYM}$ where the separation of the sentence context for pooling encourages the emergence of context information for multiple relations in the final representation vector and increases the confusion of the models.

3. Comparing the syntax-based pooling methods and the non-syntax pooling methods, the pooling based on dependency paths (i.e., $\textit{ENT-DEP0}$) is worse than the non-syntax pooling methods (i.e., $\textit{ENT-SENT}$ and $\textit{ENT-DYM}$) and perform comparably with $\textit{ENT-ONLY}$ in the DDI-2013 dataset over all the models (except for the CNN model where $\textit{ENT-ONLY}$ is much worse). These evidences suggest that the dependency paths themselves are not able to capture effective contexts for the pooling operation beyond the entity mentions for biomedical RE in DDI-2013. However, when we switch to the BB3 dataset, it turns out that $\textit{ENT-DEP0}$ is significantly better than all the non-syntax pooling methods (i.e., $\textit{ENT-ONLY}$, $\textit{ENT-SENT}$ and $\textit{ENT-DYM}$) for all the comparing models. This can be explained by the multiple relation effect in BB3 for which the dependency paths help to identify the most related context words for the two given entity mentions and filter out the confusing context words for the other relations in the sentences. The models would thus become less confused with different contexts for multiple relations as those in $\textit{ENT-SENT}$ and $\textit{ENT-DYM}$ for better performance in this case.

4. Finally, among all the pooling methods, we find that $\textit{ENT-DEP1}$ significantly outperforms the other pooling methods across different models and datasets (except the CNN model on DDI-2013 and BiLSTM on BB3). In particular, the performance improvement is substantial over the non-syntax pooling methods in BB3 where $\textit{ENT-DEP1}$ is up to 2% better than $\textit{ENT-SENT}$, $\textit{ENT-DYM}$ and $\textit{ENT-ONLY}$ on the absolute F1 scores. This helps to demonstrate the benefits of $\textit{ENT-DEP1}$ for biomedical RE to both recognize the important context words for pooling in DDI-2013 and reduce the confusion effect of the multiple relations in single sentences for the models in BB3.

3.4 Comparing the Deep Learning Models for RE

Regarding the comparison among different deep learning models, the major observations from Tables 1 and 2 include:

1. The performance of CNN is in general worse than the other models with the bidirectional LSTM components (i.e., BiLSTM, BiLSTM-CNN and BiLSTM-GCNN) over different pooling methods and datasets. This illustrates the importance of bidirectional LSTMs to capture the effective feature representations for biomedical RE.

2. Comparing BiLSTM and BiLSTM-CNN, we find that BiLSTM is better in DDI-2013 while BiLSTM-CNN achieves better performance in BB3 (over different pooling methods). In other words, the CNN layer is only helpful for the BiLSTM model in the BB3 dataset. This can also be attributed to the multiple relation effect in BB3 where the CNN layer helps to further abstract the representations from BiLSTM to better reveal the underlying structures in such confusing and complicated contexts in the sentences of BB3 for RE.

3. Graph convolutions over the dependency trees are not effective for biomedical RE as incorporating it into the BiLSTM model hurts the performance significantly. In particular, BiLSTM-GCNN is significantly worse than BiLSTM no matter which pooling methods are applied and which datasets are used for evaluation.

4. Interestingly, comparing the BiLSTM model with the $\textit{ENT-DEP1}$ pooling method (i.e., BiLSTM + $\textit{ENT-DEP1}$) and the BiLSTM-GCN model with the non-syntax pooling methods (i.e., $\textit{ENT-ONLY}$, $\textit{ENT-SENT}$ and $\textit{ENT-DYM}$), we see that BiLSTM + $\textit{ENT-DEP1}$ is significantly better with large performance gaps over both datasets DDI-2013 and BB3. For example, BiLSTM + $\textit{ENT-DEP1}$ is 1.9% better than BiLSTM-GCNN + $\textit{ENT-SENT}$ in the DDI-2013 dataset and 3.5% better than BiLSTM-GCNN + $\textit{ENT-ONLY}$ in BB3 with respect to the absolute F1 scores. In fact, BiLSTM + $\textit{ENT-DEP1}$ also achieves the best performance among the compared models in this section for both datasets. The major difference between BiLSTM + $\textit{ENT-DEP1}$ and BiLSTM-GCN with the non-syntax pooling methods lies at the specific component of the models where the syntactic information (i.e., the dependency trees) is applied. In BiLSTM-GCN with the non-syntax pooling methods, the syntactic information is employed in the representation learning component while in BiLSTM + $\textit{ENT-DEP}$, the application of the syntactic information is postponed all the way to the pooling component. Our experiments thus demonstrate that it is more effective to utilize the syntactic information in the pooling component.
than in the representation learning component of the deep learning models for biomedical RE. This is an interesting and unique observation given that the prior work for RE has only focused on using the syntactic information in the representation component and never explicitly investigated the effectiveness of the syntactic information for the pooling component of the deep learning models.

3.5 Comparing to the State-of-the-art Models

In order to further demonstrate the advantage of the syntactic information for the pooling component for biomedical RE, this section compares BiLSTM + ENT-DEP1 (i.e., the best model with the ENT-DEP1 pooling in this work) with the best reported models on the two datasets DDI-2013 and BB3. For a fair comparison between models, we select the previous single (non-ensemble) models for the comparison in this section. Tables 3 and 4 presents the model performance.

<table>
<thead>
<tr>
<th>Models</th>
<th>P</th>
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<th>F1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raihani and Laachfouti, 2017</td>
<td>73.6</td>
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<td>71.8</td>
</tr>
<tr>
<td>Zhang et al., 2018a</td>
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<td>72.9</td>
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<tr>
<td>Zhou et al., 2018</td>
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<td>73.0</td>
</tr>
<tr>
<td>Björne and Salakoski, 2018</td>
<td>75.3</td>
<td>66.3</td>
<td>70.5</td>
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<td><strong>BiLSTM + ENT-DEP1</strong></td>
<td><strong>71.6</strong></td>
<td><strong>76.4</strong></td>
<td><strong>73.9</strong></td>
</tr>
</tbody>
</table>

Table 3: Comparison with the state-of-the-art systems on the DDI-2013 test set

<table>
<thead>
<tr>
<th>Models</th>
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</tr>
</thead>
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</tr>
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<td>Mehriry et al., 2016</td>
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<td>52.1</td>
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<td>Li et al., 2016</td>
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<td>Le et al., 2018</td>
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<td><strong>BiLSTM + ENT-DEP1</strong></td>
<td><strong>54.7</strong></td>
<td><strong>72.6</strong></td>
<td><strong>62.4</strong></td>
</tr>
</tbody>
</table>

Table 4: Comparison with the state-of-the-art systems on the BB3 test set

The most important observation from the tables is that the BiLSTM model, once combined with the ENT-DEP1 pooling method, significantly outperforms the previous models on DDI-2013 and BB3, establishing new state-of-the-art performance for these datasets. In particular, in the DDI-2013 dataset, BiLSTM + ENT-DEP1 is 0.9% better than the current state-of-the-art model in (Zhou et al., 2018) while the performance improvement over the best reported model for BB3 in (Li et al., 2016) is 5.3% (over the absolute F1 scores). Such substantial improvement clearly demonstrates the advantages of the syntactic information and its delayed application in the pooling component of the deep learning models for biomedical RE.

4 Related Work

Traditional work on RE has mostly used feature engineering with syntactical information for statistical or kernel based classifiers (Zelenko et al., 2002; Zhou et al., 2005; Bunescu and Mooney, 2005; Sun et al., 2011; Chan and Roth, 2010). Recently, deep learning has been shown to advance many benchmark datasets for this RE problem due to its representation learning capacity. The typical architectures for such deep learning models involve CNN, LSTM, the attention mechanism and their variants (Zeng et al., 2014; dos Santos et al., 2015; Zhou et al., 2016; Nguyen and Grishman, 2015a; Miwa and Bansal, 2016; Zhang et al., 2017, 2018b). Deep learning has also been applied to biomedical RE in the last couple of years and started to demonstrate much potentials for this area (Mehryary et al., 2016; Björne and Salakoski, 2018; Nguyen and Verspoor, 2018; Verga et al., 2018).

Pooling is a common and crucial component in most of the deep learning models for RE. (Nguyen and Grishman, 2015b; dos Santos et al., 2015) apply the pooling operation over the whole sentence for RE while Zeng et al. (2015) proposes the dynamic pooling mechanism in the CNN models. However, none of these prior work systematically examines different pooling mechanisms for deep learning in RE as we do in this work.

5 Conclusion

We conduct a comprehensive study on the effectiveness of different pooling mechanisms for the deep learning models in biomedical relation extraction. Our experiments suggest that the pooling mechanisms have a significant impact on the performance of the deep learning models and a careful evaluation should be done to decide the appropriate pooling mechanism for the biomedical RE problem. From the experiments, we also find that syntactic information (i.e., dependency parsing) provides the best pooling methods for the models and biomedical RE datasets we investigate in this work (i.e., ENT-DEP1). We achieve the state-of-the-art performance for biomedical RE over the two datasets DDI-2013 and BB3 with such syntax-based pooling methods.
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Ang Sun, Ralph Grishman, and Satoshi Sekine. 2011. Semi-supervised relation extraction with large-scale word clustering. In ACL.


26


Syntax-aware Multi-task Graph Convolutional Networks for Biomedical Relation Extraction

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Abstract

In this paper we tackle two unique challenges in biomedical relation extraction. The first challenge is that the contextual information between two entity mentions often involves sophisticated syntactic structures. We propose a novel graph convolutional networks model that incorporates dependency parsing and contextualized embedding to effectively capture comprehensive contextual information. The second challenge is that most of the benchmark data sets for this task are quite imbalanced because more than 80% mention pairs are negative instances (i.e., no relations). We propose a multi-task learning framework to jointly model relation identification and classification tasks to propagate supervision signals from each other and apply a focal loss to focus training on ambiguous mention pairs. By applying these two strategies, experiments show that our model achieves state-of-the-art F-score on the 2013 drug-drug interaction extraction task.

1 Introduction

Recently relation extraction in biomedical literature has attracted increasing interests from medical language processing research community as an important stage for downstream tasks such as question answering (Hristovski et al., 2015) and decision making (Agosti et al., 2019). Biomedical relation extraction aims to identify and classify relations between two entity mentions into pre-defined types based on contexts. In this paper we aim to extract drug-drug interactions (DDIs), which occur when taking two or more drugs within a certain period of time that alters the way one or more drugs act in human body and may result in unexpected side effects (Figure 1). Extracting DDI provides important clues for research in drug safety and human health care.

Dependency parses are widely used in relation extraction task due to the advantage of shortening the distance of words which are syntactically related. As shown in Figure 1, the partial dependency path \{iron ← cobalt ← interactions\} reveals that these two drugs are interactive, and the path \{interactions → absorption → retention\} further indicates the mechanism relation between these two mentions. Therefore capturing the syntactic information involving the word interaction on the dependency path \{iron ← cobalt ← interactions → absorption → retention\} can effectively help on the classification of the relation between these two mentions \{cobalt, iron\}. In order to capture indicative information from wide contexts, we adopt the graph convolutional networks (GCN) (Kipf and Welling, 2016; Marcheggiani and Titov, 2017) to obtain the syntactic information by encoding the dependency structure over the input sentence with graph convolution operations. To compensate the loss of local context information in GCN, we incorporate the contextualized word representation pre-trained by the BERT model (Devlin et al., 2019) in large-scale biomedical corpora containing over 200K abstracts from PubMed and over 270K full texts from PMC (Lee et al., 2019).

Moreover, we notice that data imbalance is another major challenge in biomedical text as the distribution of relations among biomedical mentions

Figure 1: Example of drug-drug interaction on dependency tree.
are usually very sparse. Over 80% candidate mention pairs have no relation in DDI 2013 (Herrero-Zazo et al., 2013) training set. To tackle this problem, we propose a binary relation identification task as an auxiliary task to facilitate the main multi-classification task. For instance, the detection of drug interaction on dependency path \{iron \leftarrow cobalt \leftarrow interactions \rightarrow absorption \rightarrow retention\} will assist the prediction of the relation type mechanism by using the signals from binary classification as an inductive bias to avoid misclassifying it as no relation. We also exploit the focal loss (Lin et al., 2017) to potentially help the multi-class relation classification task by forcing the loss implicitly focus on ambiguous examples.

To recap, our contributions are twofold: First, we adopt the syntax-aware graph convolutional networks incorporating contextualized representation. Second, we further design an auxiliary task to solve the data imbalance problem, which achieves the state-of-the-art micro F-score on the DDIExtraction 2013 shared task.

2 Methods

![Figure 2: Framework of syntax-aware multi-task graph convolutional networks.](image)

2.1 Contextual and Syntax-aware GCN

As a variant of the convolutional neural networks (LeCun et al., 1998), the graph convolutional networks (Kipf and Welling, 2016) is designed for graph data and it has been proven effective in modeling text data via syntactic dependency graphs (Marcheggiani and Titov, 2017).

We encode the tokens in a biomedical sentence as \( x = \{x_1, \ldots, x_n\} \), where \( x_i \) is a vector which concatenates the representation of the token \( i \) and the position embeddings corresponding to the relative positions from candidate mention pairs. We feed the token vectors into a \( L \)-layer GCN to obtain the hidden representations of each token which are directly influenced by its neighbors no more than \( L \) edges apart in the dependency tree. We apply the Stanford dependency parser (Chen and Manning, 2014) to generate the dependency structure:

\[
    h_i^{(l)} = \sigma\left(\sum_{j=1}^{n} \tilde{A}_{ij} W_j^{(l)} h_j^{l-1} / d_i + b_j^{(l)}\right)
\]

where \( \tilde{A} = A + I \) with \( A \) is the adjacent matrix of tokens in dependency tree, \( I \) is the identity matrix. \( W_j^{(l)} \) is a linear transformation, \( b_j^{(l)} \) is a bias term, and \( \sigma \) is a nonlinear function. Following Zhang et al. (2018), \( d_i \) is the degree of the token \( i \) in dependency tree with an additional self-loop.

We notice that some token representations are more informative by gathering information from syntactically related neighbors through GCN. For example, the representation of the token interactions from a 2-layer GCN operating on its two edges apart neighbors provides inductive information for predicting a mechanism relation. Thus, we adopt attentive pooling (Zhou et al., 2016) to achieve the optimal pooling:

\[
    \alpha = \text{softmax}(w^T \tanh(h))
\]

\[
    h_{\text{attentive}} = h_{\alpha}^T
\]

where \( w \) is a trained parameter to assign weights based on the importance of each token representation.

We obtain the final representation by concatenating the sentence from attentive pooling and the mention representations from max pooling. We finally obtain the prediction of relation type by feeding the final representations into a fully connected neural network followed by a softmax operation.

Graph neural networks (Zhou et al., 2018b) can learn effective representations but suffer from the loss of local context information. We believe the local context information is also crucial for biomedical relation extraction. For example, in the following sentence “The response to [Factrel]drug may be blunted by [phenothiazines]drug and [dopamine antagonists]drug”, it’s intuitive to tell Factrel and phenothiazines are interactive while phenothiazines and dopamine antagonists
have no interaction according to the sentence order. However, GCNs treat the three drugs as interacting with each other as they are close in dependency structure with no order information.

BERT (Devlin et al., 2019) is a recently proposed model based on a multi-layer bidirectional Transformer (Vaswani et al., 2017). Using pre-trained BERT has been proven effective to create contextualized word embeddings for various NLP tasks (Han et al., 2019; Wang et al., 2019). The BioBERT (Lee et al., 2019) is a biomedical language representation model pre-trained on large-scale biomedical corpora. The output of each encoder layer of the input token can be used as a feature representation of that token. As shown in Figure 2, we encode the input tokens as contextualized embeddings by leveraging the last hidden layer of the corresponding token in BioBERT.

As the BERT model uses WordPiece (Wu et al., 2016) to decompose infrequent words into frequent subwords for unsupervised tokenization of the input token, if the token has multiple BERT subword units, we use the first one. After getting the contextualized embedding of each token, we feed them into the GCN layer to make our model context-aware.

2.2 Auxiliary Task Learning with Focal Loss

In the DDIExtraction 2013 task, all possible interactions between drugs within one sentence are annotated, which means a single sentence with multiple drug mentions will lead to separate instances of candidate mention pairs (Herrero-Zazo et al., 2013). There are 21,012 mention pairs generated from 3,790 sentences in training set and over 80% have no relations. This data imbalance problem due to sparse relation distribution is a main reason for low recall in DDI task (Zhou et al., 2018a; Sun et al., 2019).

Here we address this relation type imbalance problem by adding an auxiliary task on top of the syntax-aware GCN model. To conduct the auxiliary task learning, we add a separate binary classifier for relation identification as shown in Figure 2. All classifiers share the same GCN representation and contextualized embeddings, and thus they can potentially help each other by propagating their supervision signals.

Additionally, instead of setting the objective function as the negative log-likelihood loss, here we optimize the parameters in training by minimizing a focal loss (Lin et al., 2017) which focuses on hard relation types. For instance, the int relation indicates drug interaction without providing any extra information (e.g., Some [anticonvulsants]drug may interact with [Mephenytoin]drug).

This relation type only accounts for 0.82% in training set and is often misclassified into other relation types. We denote \( t_i \) and \( p_i \) as the ground truth and the conditional probability value of the type \( i \) in relation types \( C \), the focal loss can be defined as:

\[
\mathcal{L} = - \sum_{i}^{C} (\alpha(1 - p_i)^\gamma t_i \log(p_i)) + \lambda||\theta||^2
\]

where \( \alpha \) is a weighting factor to balance the importance of samples from various types, \( \gamma \) is the focusing parameter to reduce the influence of well-classified samples in the loss. \( \lambda \) is the \( L2 \) regularization parameter and \( \theta \) is the parameter set.

The auxiliary task along with the focal loss enhances our model’s ability to handle imbalance data by leveraging the inductive signal from the easier identification task and meanwhile down-weighting the influence of easy classified instances thus directing the model to focus on difficult relation types.

3 Experiments

3.1 Datasets and Task Settings

<table>
<thead>
<tr>
<th>System</th>
<th>Prec</th>
<th>Rec</th>
<th>F1</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNN (Liu et al., 2016)</td>
<td>75.70</td>
<td>64.66</td>
<td>69.75</td>
</tr>
<tr>
<td>Multi Channel CNN (Quan et al., 2016)</td>
<td>75.99</td>
<td>65.25</td>
<td>70.21</td>
</tr>
<tr>
<td>GRU (Yi et al., 2017)</td>
<td>73.67</td>
<td>70.79</td>
<td>72.20</td>
</tr>
<tr>
<td>AB-LSTM (Sahu and Anand, 2018)</td>
<td>74.47</td>
<td>64.96</td>
<td>69.39</td>
</tr>
<tr>
<td>CNN-GCNN (Asada et al., 2018)</td>
<td>73.31</td>
<td>71.81</td>
<td>72.55</td>
</tr>
<tr>
<td>Position-aware LSTM (Zhou et al., 2018a)</td>
<td>75.80</td>
<td>70.38</td>
<td>72.99</td>
</tr>
<tr>
<td>RHCNN (Sun et al., 2019)</td>
<td>77.30</td>
<td>73.75</td>
<td>75.48</td>
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<td>LSTM baseline</td>
<td>69.34</td>
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<td>65.88</td>
</tr>
<tr>
<td>GCN baseline</td>
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<td>67.14</td>
<td>69.47</td>
</tr>
<tr>
<td>–without attentive pooling</td>
<td>77.12</td>
<td>75.03</td>
<td>76.06</td>
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<tr>
<td>–without BioBERT</td>
<td>76.51</td>
<td>73.56</td>
<td>75.01</td>
</tr>
<tr>
<td>–without multi-task learning</td>
<td>76.01</td>
<td>71.92</td>
<td>73.91</td>
</tr>
<tr>
<td>Our Model</td>
<td>77.62</td>
<td>75.69</td>
<td>76.64</td>
</tr>
</tbody>
</table>

Table 1: Precision (Prec), recall (Rec) and micro F-score (F1) results on DDI 2013 corpus.

We evaluate our model on the DDIExtraction 2013 relation dataset (Herrero-Zazo et al., 2013). The corpus is annotated with drug mentions and their four types of interactions: Mechanism (pharmacokinetic mechanism of a DDI), Effect (effect
of a DDI), Advice (a recommendation or advice regarding a DDI) and Int (a DDI simply occurs without extra information). We randomly choose 10% from the training dataset as the development set. Following previous work (Liu et al., 2016; Quan et al., 2016; Zhou et al., 2018a; Sun et al., 2019), we use a negative instance filtering strategy to filter out some negative drug pairs based on manually-formulated rules. Instances containing drug pair referring to the same thing and drug pair appearing in the same coordinate structure with more than two drugs (e.g., drug1, drug2, and drug3) will be filtered. Entity mentions are masked with DRUG for better generalization and avoiding overfitting.

We train the model with GCN hidden state size of 200, the SGD optimizer with a learning rate of 0.001, a batch size of 30, and 50 epochs. Dropout is applied with a rate of 0.5 for regularization. The contextual embedding size from BioBERT is 768. The focusing parameter $\gamma$ is set as 1. All hyper-parameters are tuned on the development set.

3.2 Results and Analysis

The experiment results are reported from a 2-layer GCN which achieves the best performance and shown in Table 1. Our model significantly outperforms all previous methods at the significance level of 0.05. To analyze the contributions and effects of the various components in our model, we also perform ablation tests. The ablated GCN model outperforms the LSTM baseline by 3.6% F1 score, which demonstrates the effectiveness of GCN on modeling mention relations through dependency structure. The utilization of contextualized embedding from BioBERT which encodes the contextual information involving sequence order and word disambiguation implicitly helps the model to learn contextual relation patterns, therefore the performance is further improved. We obtain a significant F-score improvement (2.7%) by applying multi-task learning. As over 80% mention pairs are negative samples, the multi-task learning effectively solves the problem by jointly modeling relation identification and classification tasks and applying focal loss to focusing on ambiguous mention pairs, and thus we also gain 3.8% absolute score on recall. Specifically, the F1 score of Int type is increased from 54.38% to 59.79%.

For the remaining errors, we notice that our model often fails to predict relations when the sentence are parsed poorly due to the complex content which suggests us to seek for more powerful parser tools. Besides, we also observe some errors occurring in extremely short sentences. For example, in the following sentence “[Calcium]supplements/[Antacids]”, our model cannot capture informative representations as the mentions are masked with DRUG and the sentence is too concise to offer indicative evidence.

4 Related Work

Traditional feature/kernel-based models for biomedical relation extraction rely on engineered features which suffer from low portability and generalizability (Kim et al., 2015; Zheng et al., 2016; Raihani and Laachfoubi, 2017). To tackle this problem, recent studies apply Convolutional Neural Networks (CNNs) and Recurrent Neural Networks (RNNs) to automatically learn feature representations with input words encoded as pre-trained word embeddings (Zhao et al., 2016; Liu et al., 2016; Quan et al., 2016; Zhang et al., 2017; Zhou et al., 2018a; Sun et al., 2019). Learning representations of graphs are widely studied and several graph neural networks have been applied in the biomedical domain. Lim et al. (2018) proposed recursive neural network based model with a subtree containment feature. Asada et al. (2018) encoded drug pairs with CNNs and used external knowledge base to encode their molecular pairs with two graph neural networks. Here we directly apply syntax-aware GCNs on biomedical text to extract drug-drug interaction.

5 Conclusions and Future Work

We propose a syntax-aware multi-task learning model for biomedical relation extraction. Our model can effectively extract the drug-drug interactions by capturing the syntactic information through graph convolution operations and modeling context information via contextualized embeddings. An auxiliary task with focal loss is designed to mitigate the data imbalance by leveraging the inductive signal from binary classification and increasing the influence of decisive relation types. In the future, we plan to explore more informative parsers like the abstract meaning representation parser to create graph structure and consider leveraging external knowledge to further enhance the extraction quality.
Acknowledgments

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References


Abstract

Word embeddings, in their different shapes and evolutions, have changed the natural language processing research landscape in the last years. The biomedical text processing field is no stranger to this revolution; however, researchers in the field largely trained their embeddings on scientific documents, even when working on user-generated data. In this paper we show how training embeddings from a corpus collected from user-generated text from medical forums heavily influences the performance on downstream tasks, outperforming embeddings trained both on general purpose data or on scientific papers when applied to user-generated content.

1 Introduction

In the Natural Language Processing community, user-generated content, i.e. data from social media, user forums, review websites, and so on, has been the subject of many studies in the past years; the same holds for the biomedical domain, where there has been a great effort on the applications of NLP techniques for biomedical scientific publications, patient records, and so on. However, the intersection of the two fields is still in its infancy, even when dealing with relatively basic NLP tasks. For instance, in the field of user-generated biomedical natural language processing (hence UG-BioNLP), to the best of our knowledge there are no publicly available corpora for Named Entity Recognition (NER) akin in size and purpose e.g. to the CoNLL 2003 dataset. (Tjong Kim Sang and De Meulder, 2003), making it hard to compare systems effectively. Moreover, while there have been experiments on training word embeddings with biomedical data, we are not aware of any publicly available word embeddings trained on UG-BioNLP data.

For this reason, we decided to investigate the impact of using purpose-trained word embeddings in the Bio-UG field. In order to train such embeddings, we collected a dataset from Reddit, scraping posts from medical-themed subreddits, both on general health topics such as ‘r/AskDocs’, or on disease-specific subreddits, such as ‘r/cancer’, ‘r/asthma’, and so on. We then trained word embeddings on this corpus using different off-the-shelf techniques. Then, to evaluate the embeddings, we collected a second dataset of 4800 threads from the health forum HealthUnlocked, which was annotated for the NER task. Then, we analyzed the performance of the embeddings on the tasks of NER and of adverse effect mention detection. For NER, we used Conditional Random Fields as a baseline. We compared them against Bidirectional LSTM-CRFs (Lample et al., 2016), on which we analyzed the impact of using our custom-trained word embeddings against embeddings trained on general purpose data and scientific biomedical publications when evaluating on our purpose-built HealthUnlocked dataset and on the PsyTAR and CADEC corpora. Finally, we evaluated the performance of a simple architecture for adverse reaction mention detection on the PsyTAR corpus. We conclude the paper explaining our intentions for future research, in other to obtain other results that confirm the preliminary findings we present in this work.

2 Related Work

The benefit of using in-domain embeddings for the biomedical domain has already been proven effective. For example, (Pakhomov et al., 2016) and (Wang et al., 2018) found that using clinical notes or biomedical articles for training word embeddings has generally a positive impact on down-
stream NLP tasks. (Nikfarjam et al., 2015) trained embeddings on user-generated medical content and used them successfully on the pharmacovigilance task; however, they trained the embeddings on an adverse reaction mining corpus, hence making them too task-specific to be considered useful on generic UG-BioNLP tasks.

3 Datasets

3.1 BioReddit

To train our embeddings on user-generated biomedical text, we choose to scrape data from the discussion website Reddit. The website is organized by forums, called subreddits, where the discussion is restricted to a topic, e.g. general news, computer science, and so on. There is a great number of health-themed subreddits, where users from all around the world discuss their health problems or ask for medical advice, which is ideal for training our embeddings.

We also evaluated the micro-blogging platform Twitter as a possible source for the embeddings, but we quickly discarded it due to its unstructured nature. On Twitter, in fact, information is not pre-aggregated by subject, and one has to search for the required posts by searching for keyword or hashtag. This, along with the restrictive limits imposed by Twitter APIs, makes it hard to find relevant content, so we decided to continue with Reddit instead.

We designed a scraping script that downloaded discussions from 68 health themed subreddits. We selected subreddits where users

- could ask for advice, e.g. /r/AskDocs, /r/DiagnoseMe, r/AskaPharmacist,
- discuss a specific illness, e.g. /r/cancer, r/migraine, r/insomnia,
- can discuss on any health-related topic, e.g. /r/health, r/HealthIT, r/HealthInsurance.

We collected all the posts from these subreddits from the beginning of 2015 to the end of 2018. After that, we cleaned the corpus for bot-generated content, e.g. bots automatically suggesting to seek professional medical advice. We obtained a corpus with 300 million tokens and a vocabulary size of 780,000 words. While the number of tokens is considerably lower than the size of other word embedding training datasets, which could be two orders of magnitude bigger, the vocabulary is

3.2 HealthUnlocked

In order to evaluate our embeddings, as a first step, we decided to focus on the Named Entity Recognition task. We obtained 4800 forum threads from HealthUnlocked\(^1\), a British social network for health where users can discuss their health with people with similar conditions and obtain advice from professionals.

We annotated the dataset by marking the entities belonging to seven categories, namely: Phenotype, Disease, Anatomy, Molecule, Gene, Device, and Procedure. We describe in detail the categories in Table 1.

Since the dataset is collected from patients’ discussions, the language used is far from technical. For example\(^2\),
- an user describes paresthesia of arm as “a tickling sensation in my arms”;
- another patient, to describe her swollen abdomen, writes that she “looked six months pregnant”;
- another user writes that “her mood is low”, to explain her depression.

All these phrases, while expressed in layman’s language, describe very specific symptoms. For this reason, we developed a set of annotation guidelines where the annotators were asked to mark any possible mention of an entity belonging to the seven categories above, even if not expressed with technical language. After running a pilot annotation task on a small set of discussions, we fine tuned the annotation guidelines, and we asked PhD-qualified biomedical experts to annotate 4800 threads from the forums. After the annotation, the files were shuffled and split in train, test, and development set, obtaining 8750, 2526, and 1250 sentences respectively. The number of annotations per category and per set is described in Table 1.

3.3 PsyTAR

The PsyTAR dataset “contains patients expression of effectiveness and adverse drug events as-

\(^1\)https://healthunlocked.com/
\(^2\)Please note that we use feminine pronouns to preserve the privacy of the patients.
associated with psychiatric medications.” (Zolnoori et al., 2019). The dataset contains 6000 sentences annotated for mentions and spans (i.e. NER) of Adverse Drug Reactions, Withdrawal Symptoms, Drug Effectiveness, Drug Ineffectiveness, Sign/Symptoms/Illness, and Drug Indications. Each entity is grounded against the Unified Medical Language System (UMLS) and SNOMED Clinical Terms. The source of the corpus is the drug review website Ask a Patient. The language used is very simple, without the use of specialist terms, and with no guarantee of grammatical/spelling correctness.

### 3.4 CADEC

The CADEC corpus (Karimi et al., 2015) is a corpus of consumer reviews for pharmacovigilance. It is sourced from Ask a Patient too and it is annotated for mentions of concepts such as drugs, adverse reactions, symptoms and diseases, which are linked against SNOMED and MedDRA.

### 4 Experiments

#### 4.1 Embeddings

Using the dataset described in Section 3.1, we trained three word embedding models, namely GloVe (Pennington et al., 2014), ELMo (Peters et al., 2018), and Flair (Akbik et al., 2018). We choose these models due to their popularity, performance, and relative low resource requirements. In particular, GloVe requires just hours to be trained on a CPU, while ELMo and Flair obtained state-of-the-art results in the NER task at the time of their publication, and both models can be trained in relatively short time (~1 week) using 1 or 2 GPUs. As general purpose and PubMed embeddings, we use the ones provided or recommended by the respective architecture authors; unfortunately, we are not aware of any GloVe PubMed pre-trainer embeddings available in the public domain. Using our BioReddit dataset, we trained all the embeddings with their default parameters, as described in their respective papers.

#### 4.2 Named Entity Recognition

In order to evaluate our embeddings we use Conditional Random Fields and as a baseline, and then we evaluate our embeddings using a Bidirectional LSTM-CRF sequence tagging neural network (Lample et al., 2016). We refer the reader to the original paper for an explanation on how this architecture works, as the details are outside to the scope of the present paper.

We present our results in Table 2. As expected, all the neural architectures largely improve the results obtained by the CRF and, in line with the literature, Flair performs slightly better than ELMo, which in turn performs better than GloVe. Using our purpose-built embeddings, called BioReddit in the Table, we always obtain an improvement with respect to using embeddings trained on general-purpose data (Default in Table) or on PubMed, barring the smallest GloVe vectors.

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>P</th>
<th>R</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRF</td>
<td>69.7</td>
<td>60.1</td>
<td>64.5</td>
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<tr>
<td>GloVe-Default</td>
<td>69.6</td>
<td>68.3</td>
<td>68.9</td>
</tr>
<tr>
<td>GloVe-BioReddit-50</td>
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<td>67.2</td>
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<td>GloVe-BioReddit-100</td>
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<td>70.9</td>
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<tr>
<td>GloVe-BioReddit-200</td>
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<td>71.2</td>
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<td>ELMo-Default</td>
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<td>ELMo-PubMed</td>
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<td>75.8</td>
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<td>Flair-PubMed</td>
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<td>75.1</td>
<td>75.4</td>
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<td>Flair-BioReddit</td>
<td>76.5</td>
<td>76.2</td>
<td>76.4</td>
</tr>
</tbody>
</table>

Table 2: Performance of different embeddings technique on NER, when trained and evaluated on the dataset described in Section 3.2.
Table 3: Performance on the NER task of the Flair-BioReddit on the HealthUnlocked dataset on the seven categories defined in Section 3.2.

<table>
<thead>
<tr>
<th>Category</th>
<th>P</th>
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</thead>
<tbody>
<tr>
<td>Anatomy</td>
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<td>74.3</td>
</tr>
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<td>Device</td>
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<td>88.6</td>
<td>88.5</td>
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<td>Phenotype</td>
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<td>66.9</td>
<td>68.6</td>
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<tr>
<td>Procedure</td>
<td>76.6</td>
<td>80.2</td>
<td>78.4</td>
</tr>
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</table>

Table 4: Performance of the Flair embeddings on the NER and Adverse Reaction Mention Detection on the PsyTAR and CADEC corpora.

<table>
<thead>
<tr>
<th>Corpus</th>
<th>Embedding</th>
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<th>F</th>
</tr>
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<tr>
<td>Default</td>
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<td>62.4</td>
<td></td>
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<td>PsyTAR NER</td>
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<tr>
<td>BioReddit</td>
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<td>63.7</td>
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<tr>
<td>PsyTAR ADR</td>
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<td>69.2</td>
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</tr>
<tr>
<td>PubMed</td>
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<td>72.6</td>
<td>75.0</td>
<td></td>
</tr>
<tr>
<td>BioReddit</td>
<td>79.5</td>
<td>73.7</td>
<td>76.5</td>
<td></td>
</tr>
<tr>
<td>Default</td>
<td>77.1</td>
<td>76.0</td>
<td>76.5</td>
<td></td>
</tr>
<tr>
<td>CADEC NER</td>
<td>77.2</td>
<td>76.1</td>
<td>76.7</td>
<td></td>
</tr>
<tr>
<td>Bioreddit</td>
<td>78.6</td>
<td>77.4</td>
<td>78.0</td>
<td></td>
</tr>
</tbody>
</table>

4.3 Adverse Reaction Mention Detection

The task of Adverse Reaction Mention Detection (hence ADR) consists in detecting whether in a sentence a user mentions that he is experiencing/experienced an adverse reaction to a drug. For this task, we designed a simple neural architecture, where a bidirectional GRU (Cho et al., 2014) reads a sentence, and a softmax layer on its top performs the binary classification task of detecting whether the input sentence contains an ADR or not. When evaluating on the PsyTAR corpus we again obtain the best performance when using our BioReddit embeddings, followed by the PubMed trained ones and the default ones.

5 Conclusions

In this paper we showed how training ad-hoc embeddings for the task of user-generated biomedical text processing improves the results in the tasks of named entity recognition and adverse reaction mention detection. While preliminary, our results show a strong indication that embeddings trained on biomedical scientific literature only are not guaranteed to be effective when used on user-generated data, since people use “layman terms” which are seldom, if ever, used in scientific literature. As future work, we acknowledge the need to better investigate the results we present here. A good starting point would be to analyze other embedding techniques, in order to investigate if the performance improvement is due to embedding techniques themselves or to the datasets used. Moreover, we need to analyze the performance of our BioReddit embeddings on non-user generated content, as e.g. scientific abstracts, in order to investigate whether they are able to perform effectively on this domain too. Finally, we think that a manual investigation of the results of the downstream tasks is important, to investigate e.g. if the improvement in the ADR task is due to the embeddings helping to classify sentences with more colloquial language. Unfortunately, due to licensing and privacy issues, we are not allowed to release the HealthUnlocked corpus. However, we make available our BioReddit embeddings trained on GloVe, ELMo and Flair at [https://github.com/basaldella/bioreddit](https://github.com/basaldella/bioreddit). For the sake of reproducibility, we also make available our PsyTAR preprocessed splits online at [https://github.com/basaldella/psytarpreprocessor](https://github.com/basaldella/psytarpreprocessor).
Acknowledgements

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References


Leveraging Hierarchical Category Knowledge for Data-Imbalanced Multi-Label Diagnostic Text Understanding

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Abstract
Clinical notes are essential medical documents to record each patient’s symptoms. Each record is typically annotated with medical diagnostic codes, which means diagnosis and treatment. This paper focuses on predicting diagnostic codes given the descriptive present illness in electronic health records by leveraging domain knowledge. We investigate various losses in a convolutional model to utilize hierarchical category knowledge of diagnostic codes in order to allow the model to share semantics across different labels under the same category. The proposed model not only considers the external domain knowledge but also addresses the issue about data imbalance. The MIMIC3 benchmark experiments show that the proposed methods can effectively utilize category knowledge and provide informative cues to improve the performance in terms of the top-ranked diagnostic codes which is better than the prior state-of-the-art. The investigation and discussion express the potential of integrating the domain knowledge in the current machine learning based models and guiding future research directions.

1 Introduction
Electronic health records (EHR) usually contain clinical notes, which are free-form text generated by clinicians during patient encounters, and a set of metadata diagnosis codes from the International Classification of Diseases (ICD), which represent the diagnoses and procedures in a standard way. ICD codes have a variety of usage, ranging from billing to predictive modeling of the patient state (Choi et al., 2016). Automatic diagnosis prediction has been studied since 1998 (de Lima et al., 1998). Mullenbach et al. (2018) pointed out the main challenges of this task: 1) the large label space, with over 15,000 codes in the ICD-9 taxonomy, and over 140,000 codes in the newer ICD-10 taxonomies (Organization et al., 2007), and 2) noisy text, including irrelevant information, misspellings and non-standard abbreviations, and a large medical vocabulary. Several recent work attempted at solving this task by neural models (Shi et al., 2017; Mullenbach et al., 2018).

However, most prior work considered the output labels independently, so that the codes with few samples are difficult to learn (Shi et al., 2017). Therefore, Mullenbach et al. (2018) proposed an attentional model to effectively utilize the textural forms of codes to facilitate learning. In addition to textual definitions of codes, the category domain knowledge may provide additional cues to allow the codes under same category to share parameters, so the codes with few samples can benefit from it. To effectively utilize the category knowledge from the ICD codes, this paper proposes several refined category losses and incorporate them into convolutional models and then evaluate the performance on both MIMIC-3 (Johnson et al., 2016) and our internal dataset. The experiments on MIMIC shows that the proposed knowledge integration model significantly improves the previous methods and achieves the state-of-the-art performance, and the improvement can also be observed in our internal dataset. The idea is similar to the prior work (Singh et al., 2018), which considered the keyword hierarchy for information extraction from medical documents, but our work focuses on leveraging domain knowledge for clinical code prediction. Our contributions are three-fold:

- This paper first leverages external domain knowledge for diagnostic text understanding.
- The paper investigates multiple ways for incorporating the domain knowledge in an end-to-end manner.
- The proposed mechanisms improve all prior
models and achieves the state-of-the-art performance on the benchmark MIMIC dataset.

2 Methodologies

Given each clinical record in EHR, the goal is to predict the corresponding diagnostic codes with the external hierarchical category information. This task is framed as a multi-label classification problem. The proposed mechanism is built on the top of various convolutional models to further combine with the category knowledge. Below we introduce the previously proposed convolutional models which are used for latter comparison in the experiment and detail the mechanism that leverages hierarchical knowledge.

2.1 Convolutional Models

There are various models for sequence-level classification, and this paper focuses on two types of convolutional models for investigation. The models are described as follows. Note that the proposed mechanism is flexible for diverse models.

TextCNN Let $x_i \in \mathbb{R}^k$ be the $k$-dimensional word embedding corresponding to the $i$-th word in the document, represented by the matrix $X = [x_1, x_2, ..., x_N]$, where $N$ is the length of the document. TextCNN (Kim, 2014) applies both convolution and max-pooling operations in one dimension along the document length. For instance, a feature $c_i$ is generated from a window of words $x_i, x_{i+1}, ..., x_{i+h}$, where $h$ is the kernel size of the filters. The pooling operation is then applied over $c = [c_1, c_2, ..., c_{n-h+1}]$ to pick the maximum value $\hat{c} = \max(c)$ as the feature corresponding to this filter. We implement the model with kernel size $= 3, 4, 5$, considering different window sizes of words.

Convolutional Attention Model (CAML) Because the number of samples of each code is highly unbalanced, it is difficult to train each label with very few samples. To resolve this issue, the CAML model utilizes the descriptive definition of diagnosis codes, which additionally applies a per-label attention mechanism, where the additional benefit is that it selects the $n$-grams from the text that are most relevant to each predicted label (Mullenbach et al., 2018).

2.2 Knowledge Integration Mechanism

Considering the hierarchical property of ICD codes, we assume that using the higher level labels could learn more general concepts and thus improve the performance. For instance, the definitions of ICD-9 codes 301.2 and 307.1 are “Schizoid personality disorder” and “Anorexia nervosa” respectively. If we only use the labels given by the dataset, they are seen as two independent labels; however, in the ICD structure, both 301.2 and 307.1 belong to the same high-level category “mental disorders”. The external knowledge shows that category knowledge provides additional cues to know code relatedness. Therefore, we propose four types of mechanisms that incorporate hierarchy category knowledge to improve the ICD prediction below.

Cluster Penalty Motivated by Nie et al. (2018), we compute two constraints to share the parameters of the ICD codes under the same categories. The between-cluster constraint, $\Omega_{\text{between}}$, indicates the total distance of parameters between
mean of all ICD codes and the mean of each category.

\[ \Omega_{between} = \sum_{k=1}^{K} \| \bar{\theta}_k - \bar{\theta} \|^2, \]  

(1)

where \( \bar{\theta} \) is the mean vectors of all ICD codes, \( \bar{\theta}_k \) is the mean vector of the \( k \)-th category. The within-cluster constraint, \( \Omega_{within} \), is the distance of parameters between the mean of each category and its low-level codes.

\[ \Omega_{within} = \sum_{k=1}^{K} \sum_{i \in J(k)} \| \theta_i - \bar{\theta}_k \|^2, \]  

(2)

where \( J(k) \) is a set of labels that belong to the \( k \)-th category. \( \Omega_{between} \) and \( \Omega_{within} \) are formulated as additional losses to enable the model to share parameters across codes with the same categories.

**Multi-Task Learning** Considering that the high-level category can be treated as another task, we apply a multi-task learning approach to leverage the external knowledge. This model focuses on predicting the low-level codes, \( y_{low} \), as well as its high-level category, \( y_{high} \), individually illustrated in Figure 1.

\[ y_{high} = W_{high} \cdot h + b_{high} \]  

(3)

where \( W_{high} \in \mathbb{R}^{N_{high} \times d} \), \( N_{high} \) means the number of high-level categories, and \( d \) is the dimension of hidden vectors derived from CNN.

**Hierarchical Learning** We build a dictionary for mapping our low-level labels to the corresponding high-level categories illustrated in Figure 1. To estimate the weights for high-level categories, \( y_{high} \), two mechanisms are proposed:

- Average meta-label: The probability of the \( k \)-th high-level category can be approximated by the averaged weights for low-level codes that belong to the \( k \)-th category.

\[ y_{high} = \frac{1}{K} \sum_{k} y_{low}^k \]  

(4)

- At-least-one meta-label: Motivated by Nie et al. (2018), meta labels are created by examining whether any disease label for the \( k \)-th category has been marked as tagged, where the high-level probability is derived from the low-level probability of disease labels.

\[ y_{high} = 1 - \prod_{k} (1 - y_{low}^k) \]  

(5)

### 2.3 Training

The knowledge integration mechanisms are built on top of the multi-label convolutional models, which treat each ICD label as a binary classification. The predicted values for high-level categories come from the proposed mechanisms. Considering that learning low-level labels directly is difficult due to the highly imbalanced label distribution, we add a loss term indicating the high-level category in order to learn the general concepts in addition to the low-level labels, and train the model in an end-to-end fashion. Note that the high-level loss is set as \( loss_{high} = \Omega_{between} + \Omega_{within} \) for cluster penalty and the binary log loss for other methods.

\[ loss = loss_{low} + \lambda \cdot loss_{high}, \]  

(6)

where \( \lambda \) is the parameter to control the influence of the knowledge category and we choose \( \lambda = 0.1 \).

### 3 Experiments

In order to measure the effectiveness of the proposed methods, the following experiments are conducted.

#### 3.1 Setup

We evaluate our model on two datasets, one is the benchmark MIMIC-3 data and another is the dataset collected by National Taiwan University Hospital (NTUH). MIMIC-3 (Johnson et al., 2016) is a benchmark dataset, where the text and structured records from a hospital ICU. We use the same setting as the prior work (Mullenbach et al., 2018), where 47,724 discharge summaries is for training, with 1,632 summaries and 3,372 summaries for validation and testing, respectively. We also obtain a subdataset from original MIMIC3-Full, called MIMIC3-50, which has the top 50 high frequency labels. NTUH dataset is collected

### Table 1: Dataset comparison and statistics. From the full set of the internal data (1495 labels) to 200, only 6.0% of data points are discarded.

<table>
<thead>
<tr>
<th># training documents</th>
<th>MIMIC-3</th>
<th>Internal</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Full</td>
<td>50</td>
</tr>
<tr>
<td>mean length of texts</td>
<td>47,424</td>
<td>8,067</td>
</tr>
<tr>
<td>vocabulary size</td>
<td>51,917</td>
<td>51,917</td>
</tr>
<tr>
<td>OOV rate</td>
<td>0.137</td>
<td>0.137</td>
</tr>
<tr>
<td># labels</td>
<td>8,922</td>
<td>50</td>
</tr>
<tr>
<td>mean number of labels</td>
<td>15.9</td>
<td>5.7</td>
</tr>
</tbody>
</table>
## Table 2: The results on MIMIC-3 data (%). † indicates the improvement over the baseline.

<table>
<thead>
<tr>
<th>MIMIC-3-50</th>
<th>P@1</th>
<th>P@3</th>
<th>P@5</th>
<th>MAP</th>
<th>Macro-F</th>
<th>Micro-F</th>
<th>Macro-AUC</th>
<th>Micro-AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNN (Shi et al., 2017)</td>
<td>82.8</td>
<td>71.2</td>
<td>61.4</td>
<td>57.9</td>
<td>63.0</td>
<td>82.2</td>
<td>91.2</td>
<td></td>
</tr>
<tr>
<td>+ Cluster Penalty</td>
<td>83.5</td>
<td>71.9</td>
<td>62.4</td>
<td>58.3</td>
<td>63.7</td>
<td>88.5</td>
<td>91.3</td>
<td></td>
</tr>
<tr>
<td>+ Multi-Task</td>
<td>83.5</td>
<td>71.3</td>
<td>61.9</td>
<td>57.6</td>
<td>62.8</td>
<td>88.1</td>
<td>91.1</td>
<td></td>
</tr>
<tr>
<td>+ Hierarchical avg</td>
<td>84.5</td>
<td>72.1</td>
<td>62.4</td>
<td>58.6</td>
<td>64.3</td>
<td>88.9</td>
<td>91.4</td>
<td></td>
</tr>
<tr>
<td>at-least-one</td>
<td>83.4</td>
<td>72.1</td>
<td>62.4</td>
<td>58.5</td>
<td>63.8</td>
<td>88.4</td>
<td>91.3</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MIMIC3-Full</th>
<th>P@1</th>
<th>P@3</th>
<th>P@5</th>
<th>MAP</th>
<th>Macro-F</th>
<th>Micro-F</th>
<th>Macro-AUC</th>
<th>Micro-AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNN (Shi et al., 2017)</td>
<td>80.5</td>
<td>73.6</td>
<td>59.6</td>
<td>45.4</td>
<td>3.8</td>
<td>42.9</td>
<td>81.8</td>
<td>97.1</td>
</tr>
<tr>
<td>+ Cluster Penalty</td>
<td>80.9</td>
<td>74.0</td>
<td>59.5</td>
<td>45.2</td>
<td>3.3</td>
<td>40.5</td>
<td>82.1</td>
<td>97.0</td>
</tr>
<tr>
<td>+ Multi-Task</td>
<td>82.8</td>
<td>75.8</td>
<td>61.5</td>
<td>46.6</td>
<td>3.6</td>
<td>43.9</td>
<td>83.3</td>
<td>97.3</td>
</tr>
<tr>
<td>+ Hierarchical avg</td>
<td>79.0</td>
<td>73.1</td>
<td>59.2</td>
<td>45.2</td>
<td>4.3</td>
<td>42.7</td>
<td>83.0</td>
<td>97.1</td>
</tr>
<tr>
<td>at-least-one</td>
<td>82.1</td>
<td>74.3</td>
<td>59.7</td>
<td>44.9</td>
<td>2.6</td>
<td>42.0</td>
<td>80.3</td>
<td>96.7</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CAML (Mullenbach et al., 2018)</th>
<th>P@1</th>
<th>P@3</th>
<th>P@5</th>
<th>MAP</th>
<th>Macro-F</th>
<th>Micro-F</th>
<th>Macro-AUC</th>
<th>Micro-AUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>+ Cluster Penalty</td>
<td>88.4</td>
<td>82.4</td>
<td>68.8</td>
<td>54.0</td>
<td>5.4</td>
<td>51.2</td>
<td>87.5</td>
<td>98.3</td>
</tr>
<tr>
<td>+ Multi-Task</td>
<td>89.7</td>
<td>83.4</td>
<td>69.7</td>
<td>54.8</td>
<td>6.9</td>
<td>52.3</td>
<td>88.8</td>
<td>98.5</td>
</tr>
<tr>
<td>+ Hierarchical avg</td>
<td>89.6</td>
<td>83.5</td>
<td>70.9</td>
<td>56.1</td>
<td>8.2</td>
<td>53.9</td>
<td>89.5</td>
<td>98.6</td>
</tr>
<tr>
<td>at-least-one</td>
<td>89.4</td>
<td>83.3</td>
<td>69.5</td>
<td>54.8</td>
<td>6.2</td>
<td>51.7</td>
<td>88.3</td>
<td>98.4</td>
</tr>
</tbody>
</table>

### Table 3: The results on NTUH data.

<table>
<thead>
<tr>
<th>Data-200</th>
<th>Macro-F1</th>
<th>Micro-F1</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNN</td>
<td>7.6</td>
<td>39.8</td>
</tr>
<tr>
<td>+ Multi-Task</td>
<td>11.7†</td>
<td>41.6†</td>
</tr>
<tr>
<td>+ Hierarchical (avg)</td>
<td>9.3†</td>
<td>44.1†</td>
</tr>
<tr>
<td>CAML</td>
<td>6.2</td>
<td>42.6</td>
</tr>
<tr>
<td>+ Multi-Task</td>
<td>14.5†</td>
<td>44.7†</td>
</tr>
<tr>
<td>+ Hierarchical (avg)</td>
<td>18.4†</td>
<td>45.7†</td>
</tr>
</tbody>
</table>

## 3.2 Results

The baseline and the results of adding the proposed mechanisms are shown in Table 2. For MIMIC3-50, all proposed mechanisms achieve the improvement for almost all metrics, and the best one is from the hierarchical learning with average meta-label. The consistent improvement indicates that category knowledge provides informative cues for sharing parameters across low-level codes under the same categories. For MIMIC3-Full, our proposed mechanisms still outperform the baseline CNN model, and the best performance comes from the one with multi-task learning. The reason may be that multi-task learning has more flexible constraints compared with hierarchical learning, and it is more suitable for this more challenging scenario due to data imbalance. In addition, the proposed knowledge integration mechanisms using multi-task learning or hierarchical learning with average meta-label are able to improve the prior state-of-the-art model, CAML (Mullenbach et al., 2018), demonstrating the superior capability and the importance of domain knowledge.

To further investigate the model effectiveness, we perform the experiments on the NTUH dataset in Table 3. Due to shorter clinical notes and higher OOV rate, this dataset is more challenging and the results are lower than the ones in MIMIC-3. Nevertheless, the proposed methods still improve the performance by integrating category knowledge using multi-task learning or hierarchical learning with average meta-label. In sum, our proposed category knowledge integration mechanisms are...
capable of improving the text understanding performance by combining the domain knowledge with neural models and achieve the state-of-the-art results.

3.3 Qualitative Analysis
From our prediction results, we find that our proposed mechanisms tend to predict more labels than the baseline models for both CNN and CAML. Specifically, our methods can assist models to consider more categories from shared information in the hierarchy. The additional codes often contain the right answers and sometimes are in the correct categories but not exactly matched. Moreover, our mechanisms have the capability of correcting the wrong codes to the correct ones which are under the same category. The appendix provides some examples for reference.

4 Conclusion
This paper proposes multiple mechanisms using the refined losses to leverage hierarchical category knowledge and share semantics of the labels under the same category, so the model can better understand the clinical texts even if the training samples are limited. The experiments demonstrate the effectiveness of the proposed knowledge integration mechanisms given the achieved state-of-the-art performance and show the great generalization capability for multiple datasets. In the future, we plan to analyze the performance of each label, investigating which label can benefit more from the proposed approaches.

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References


Experiments with ad hoc ambiguous abbreviation expansion

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Abstract

The paper addresses experiments to expand ad hoc ambiguous abbreviations in medical notes on the basis of morphologically annotated texts, without using additional domain resources. We work on Polish data but the described approaches can be used for other languages too. We test two methods to select candidates for word abbreviation expansions. The first one automatically selects all words in text which might be an expansion of an abbreviation according to the language rules. The second method uses clustering of abbreviation occurrences to select representative elements which are manually annotated to determine lists of potential expansions. We then train a classifier to assign expansions to abbreviations based on three training sets: automatically obtained, consisting of manual annotation, and concatenation of the two previous ones. The results obtained for the manually annotated training data significantly outperform automatically obtained training data. Adding the automatically obtained training data to the manually annotated data improves the results, in particular for less frequent abbreviations. In this context the proposed a priori data driven selection of possible extensions turned out to be crucial.

1 Introduction

Saving time and effort is a crucial reason for using abbreviations and acronyms in all types of texts. In informal texts like e-mails, communicator messages, and notes, it is very common to create ad hoc abbreviations, which are easy for the author (in the case of personal notes) or a reader to interpret in the context of a topic discussed by a group of people.

The time/effort saving principle is also valid for medical notes prepared by physicians during patient visits, hospital examinations, and for nursing notes. They are often written in a hurry, but have to be understandable for other people involved in the treatment of a patient. They cannot be completely hermetic, but usually, they are difficult to interpret both for patients and nonspecialists (Mowery et al., 2016b). Ad hoc abbreviations are also difficult for automatic data processing systems to handle. But proper understanding and normalization of all types of abbreviations is indispensable for the correct operation of information extraction systems (Pradhan et al., 2014), data classification (Névéol et al., 2016; Mowery et al., 2016a), question answering (Kakadiaris et al., 2018), and many other applications.

The interpretation of an abbreviation consists of two aspects: its recognition and expansion. The recognition of well established abbreviations and acronyms is usually done with the help of dictionaries. For the English medical domain, several dictionaries such as the resources of the U.S. National Library of Medicine are available. Ad hoc abbreviations are not present in dictionaries and they are mainly recognized as unknown words. Sometimes, they are ambiguous with correct full word forms listed in general language dictionaries. For example, dept might be an abbreviation of ‘department’ or ‘deputy’, in Polish medical notes temp ‘rate’ is an abbreviation of temperatura ‘temperature’. In informal texts, the period after abbreviations, required after some of them in Polish, English and many other languages, is often omitted, which makes the distinction between word and abbreviation more difficult. Ad hoc abbreviations are also ambiguous with standard language abbreviations, and their interpretation is different from those used in standard language: literature, papers or everyday use. For example, in many languages (e.g. English, German, Polish) the abbreviation op means the opus number in musical composition. In Polish medical notes, it can be opa-
Our study focuses on expanding word abbreviations in medical notes in Polish. We are not systematically considering phrase abbreviations, usually called acronyms, as selecting candidates for their expansions requires different methods. In the paper we test two approaches for selecting candidates for word abbreviation expansions which are used for training a classifier to assign the appropriate expansion to an abbreviation, see (Pakhomov, 2002). In the first method, we check a hypothesis that full forms of ad hoc abbreviations are represented in texts of the same domain and type. So, for each candidate for abbreviation, we select all words which might be expansions for the abbreviation according to the language rules. We test if the occurrences of potential expansions in text can be sufficient for training the classifier. This method provides us with many suggestions which might never be used. To limit this number, we modified the method by selecting words with distributed representation close to the representation of the abbreviation. In the second method, we select candidates for abbreviation expansions based on annotation of selected elements of abbreviation occurrences clusters. Clustering is done by the Chinese whispers algorithm (Biemann, 2006) according to their contexts. For each cluster, we expand a manually randomly selected 2 to 6 elements of each cluster. This procedure gives us a short list of potential expansions.

2 Related Work

The problem of abbreviation recognition and expansion, has so far been addressed mainly for English data, e.g. (Nadeau and Turney, 2005) and (Moon et al., 2012) where supervised machine learning algorithms are used, and (Du et al., 2019) who describes a complex system for English data that recognizes many types of abbreviations. But, there are papers describing the problem for other languages too, e.g. Swedish (Kvist and Velupillai, 2014) – SCAN system based on lexicon, and German, e.g. (Kreuzthaler et al., 2016) where abbreviation identification linked to the disambiguation of period characters in clinical narratives is addressed.

Methods of dealing with acronyms are described among others in (Schwartz and Hearst, 2003) where the authors look for acronym definitions in data and identify them as a text in parentheses adjacent to the acronym/abbreviation; (Tengstrand et al., 2014) where experiments for Swedish are described; and (Spasic, 2018) where terminology extraction methods are applied.

Experiments in which similar to our data driven approach is tested, are described in (Oleynik et al., 2017). They used a method of abbreviation expansion based on N-grams and achieved an F1 score of 0.91, evaluated on a test set of 200 text excerpts.

3 Data

Medical reports which we used to carry out the experiments are an anonymized sample of data collected by the company providing electronic health record services. The research is a part of a project, the purpose of which is, among other things, automatic preparation of statistics concerning data on diseases, symptoms and treatments. The statistics should be based on information extracted automatically from descriptions of patients visits. The data was collected in many clinics and concerned visits to doctors of various specialties. Identification information was removed from texts, and only descriptions of interrogation, examination and recommendations were processed.

As Polish is an inflectional language we preprocessed text to obtain base word forms and POS tags. Medical reports usually contain a limited dictionary but a lot of words are not present in general dictionaries, thus specialized medical taggers would be the most appropriate for performing this task. However, manually annotated data to train the medical tagger is not available for Polish, thus we had to processed texts with the general purpose morphological tagger. In this work we used Concraft2 – the new version of the Concraft (Waszczuk, 2012) tagger which cooperates with the general purpose morphological analyzer Morfeusz2 (Woliński, 2014) and also performs tokenization and sentence identification. Additionally, we ensured that line breaks were treated
as sentence delimiters, as often a dot was not used at the end of a line, while the topic was changed. The quality of automatic tagging of medical texts in Polish is not high, see (Marciniak and Mykowiecka, 2011). Medical notes contain a large number of spelling errors, there are many acronyms/abbreviations and specialized words not present in the general purpose morphological analyzer. Thus, we performed our experiments using both exact word forms and lemmas.

The entire dataset consists of about 10 million tokens and 15,000 different word forms. This number is larger than for English data of the same size as Polish is an inflectional language. It means that one word can have several forms, e.g.: kropla ‘drop’ is represented in our data as: krople, kropli, kroplach, kroplami. As we differentiate between capital and small letters we additionally have the following forms: Krople, KROPLE. The latter decision resulted from a desire to preserve information about acronyms for future work. For example, the form PSA is rather the acronym of an examination while the form psa might be interpreted as a ‘dog’ in the genitive (in medical texts it mainly occurs in the phrase sierść psa ‘dog fur’ in the context of allergens).

Around 7% of tokens are recognized as unknown words and this group of tokens consists of about 91,000 different elements: abbreviations; acronyms; proper names such as medications and illnesses containing proper names (e.g. Hashimoto’s thyroiditis); and typos that occur in large numbers in medical notes. Some abbreviations are represented in Morfeusz2 but often their meaning is not appropriate for medical texts, e.g.: a string ‘por’ is recognized as an abbreviation of ‘lieutenant’ or ‘compare’ while in medical data it is ‘clinic’.

Tokens which are not recognized by dictionaries, are natural candidates for being abbreviations. In many papers addressing the problem of abbreviation recognition, the authors limit themselves to considering such tokens, see (Park and Byrd, 2001), (Kreuzthaler et al., 2016). In our approach, when selecting potential abbreviations, we took into account all forms out of the dictionary, and short words (up to 5 letters) which were in the dictionary. As we wanted to use contexts in our experiment, we decided to consider forms which occurred in the data more than 15 times. This limited the list of unknown tokens to 2808 and the list of word forms considered as potential abbreviations to 3152.

The data set was divided into ten parts, one was left for evaluation purposes and the remaining 9 were used as a training set and a source of information on the number and types of abbreviations used.

The test set consists of about 996,000 tokens and thousands of abbreviation occurrences. To make manual checking of the results feasible we decided to perform our experiment on a small subset of 15 abbreviations. This short list consists of abbreviations which seem to be ambiguous (a few likely interpretations) and are rather common – 3069 occurrences in test data which means 0.3% of tokens. Their proper recognition is, therefore, important for correct text interpretation. All occurrences of these 15 abbreviations in the test set were manually expanded by a person with experience in medical data processing. All difficult cases were consulted with a specialist. A fragment of the list together with exemplary variations is given in Table 1.

### 4 Language Models

On the basis of the entire data set, we trained four word2vec (Mikolov et al., 2013) versions of language models (the choice of specialized data seems to be straightforward, but was also supported by (Charbonnier and Wartena, 2018)). One pair of models was trained on the original (tokenized) texts – inflected forms of words. The second pair of models was trained on the lemmatized text (in Polish, nouns, verbs and adjectives have many different inflectional variants). In both pairs we calculated vectors of length 100; one model was trained on all tokens which occurred at least 5 times and the second one was trained on text in which all numbers were replaced by one symbol. In the final experiments form based models turned out to be the most efficient.

### 5 Baseline

We solve the problem of abbreviation expansion as the task of word sense disambiguation where a classifier is trained on all expansions represented in the data. As it is difficult to compare our approach to other work as the assumptions of the tasks related to abbreviation expansion were dif-

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1Marciniak and Mykowiecka (2011) reported that around 6% of tokens in hospital records are acronyms and abbreviations.
Table 1: Four from the list of 15 abbreviations with variations, the number of all different longer words found in the training data.

<table>
<thead>
<tr>
<th>Abr.</th>
<th>Variations</th>
<th>All</th>
<th>Possible Meanings in Test Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>fiz</td>
<td>FIZ, fiz, Fiz</td>
<td>15</td>
<td>fizjologiczny, fizycznie, fizyczny, fizykalnie, fizykoterapia Physiological, Physically, Physical, Physical Therapy</td>
</tr>
<tr>
<td>cz</td>
<td>CZ, Cz</td>
<td>44</td>
<td>czerwien, czołowy, czynnik, czynność, częstość, część head, head(adj), mainly, main, deep</td>
</tr>
<tr>
<td>gl</td>
<td>gl</td>
<td>22</td>
<td>głowa, głęboki, głęboko, głowy, głowni</td>
</tr>
<tr>
<td>op</td>
<td>OP Op</td>
<td>30</td>
<td>opak, opakowanie, opatrunek, opera, operacja, operacyjnie, operacyjny, oko prawy, opis, opuszek, opór operated, care, description, pad, resistance</td>
</tr>
</tbody>
</table>

Table 2: Number of occurrences in train and test data. The three potential extensions lists for simulated training sets: AL – all words being potential expansions, SL – all the possible words in our distributional model whose similarity to a particular abbreviation was higher than 0.2 for a language model created on forms, CL – annotations of randomly selected cluster elements.

<table>
<thead>
<tr>
<th>abbr.</th>
<th>simulated train data</th>
<th>annotated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>AL</td>
<td>SL</td>
</tr>
<tr>
<td>cz</td>
<td>51022</td>
<td>46172</td>
</tr>
<tr>
<td>fiz</td>
<td>10769</td>
<td>10684</td>
</tr>
<tr>
<td>gl</td>
<td>15591</td>
<td>14460</td>
</tr>
<tr>
<td>kr</td>
<td>37381</td>
<td>24349</td>
</tr>
<tr>
<td>mies</td>
<td>9021</td>
<td>8949</td>
</tr>
<tr>
<td>op</td>
<td>24677</td>
<td>21673</td>
</tr>
<tr>
<td>poj</td>
<td>4386</td>
<td>4035</td>
</tr>
<tr>
<td>pow</td>
<td>22517</td>
<td>5037</td>
</tr>
<tr>
<td>pr</td>
<td>88312</td>
<td>20386</td>
</tr>
<tr>
<td>rodz</td>
<td>6459</td>
<td>6459</td>
</tr>
<tr>
<td>sr</td>
<td>3894</td>
<td>2922</td>
</tr>
<tr>
<td>wz</td>
<td>9942</td>
<td>6914</td>
</tr>
<tr>
<td>zab</td>
<td>8670</td>
<td>8085</td>
</tr>
<tr>
<td>zaw</td>
<td>3826</td>
<td>1296</td>
</tr>
<tr>
<td>zal</td>
<td>1657</td>
<td>1544</td>
</tr>
<tr>
<td>total</td>
<td>298149</td>
<td>182965</td>
</tr>
</tbody>
</table>

6 Methods for Determining Expansions

When we look for potential expansions of a selected token without any additional resources, we have to consider two cases. The first, is that we should leave the token unchanged as it could be a correct word or acronym. We do not address this problem. The second, is that we should select all words from the data that can be abbreviated to the considered token according to language rules. So, the list of potential expansions consists of all forms from the data which met the conditions of being an abbreviation in Polish. We analyse cases in which a token x might be an abbreviation of a word y if:

- the beginning of y is equal to x;
- the POS tag does not indicate an abbreviation or an unknown word (to avoid using incorrectly written words as potential extensions);
- the abbreviation does not cross Polish two-letter compounds (‘rz’, ‘sz’, ‘cz’, ‘ch’).

The first potential extensions list (AL) contains all words meeting the above conditions. It consists of 1345 elements. The AL list contains forms which are never shortened. Their usage should thus be different from that of the abbreviation itself. To eliminate such unlikely expansions and to limit the number of potential labels, we selected
Table 3: Test set abbreviation expansions in numbers

<table>
<thead>
<tr>
<th>abbr.</th>
<th>test anot.</th>
<th>expansions</th>
</tr>
</thead>
<tbody>
<tr>
<td>cz</td>
<td>137</td>
<td>czerwień(1), czynnik(3), czynność(1), częstość(14), częstotliwość(102) redness, factor, activity, frequency, part</td>
</tr>
<tr>
<td>fiz</td>
<td>59</td>
<td>fizjologiczny(2), fizycznie(1), fizyczny(fizyczność(5), fizykalnie(45), fizykalny(6) physical, physically, physical, physically, physical</td>
</tr>
<tr>
<td>gł</td>
<td>48</td>
<td>gladki(1), głowa(16), głowów(4), główie(17), głowy(10) smooth, head, head, mainly, main</td>
</tr>
<tr>
<td>kr</td>
<td>224</td>
<td>krawędź(1), kreatynina(2), kropla(68), krople(4), kręgosłup(149) edge, creatinine, drop, drops, spine</td>
</tr>
<tr>
<td>mies</td>
<td>206</td>
<td>miesiąc(187), miesiączka(18), miesięczny(1) month, menstruation, monthly</td>
</tr>
<tr>
<td>op</td>
<td>1785</td>
<td>oko prawe(349), ostatni poród(1), opakowanie(1384), opatrunek(6), operacja(22), operacyjnie(3), operacyjny(10), operować(1), opieka(7), opuszek(2) right eye, last delivery, package, dressing, surgery, surgically, surgical, operate, care, fingertip</td>
</tr>
<tr>
<td>poj</td>
<td>147</td>
<td>pojawić(2), pojedynczy(127), pojemnik(18) appear, single, container</td>
</tr>
<tr>
<td>pow</td>
<td>65</td>
<td>powierzchnia(17), powiększony(6), powiększyć(8), powlecąć(9), powyżej(24), powód(1) surface, enlarged, enlarge, coated, above, reason</td>
</tr>
<tr>
<td>pr</td>
<td>100</td>
<td>Pogotowie Ratunkowe(5), public relations(3), prawa ręka(1), PR(1), per rectum(25), prawidłowo(17), prawidłowy(34), prawy(20), preparat(2), prostata(4), przewód(1), przychodnia(1), próba(6) Emergency Service, public relations, right hand, PR(in ECG), per rectum, properly, normal, right, preparation, prostate, tract, clinic, test</td>
</tr>
<tr>
<td>rodz</td>
<td>52</td>
<td>rodzzeństwo(8), rodzice(3), rodzina(4), rodzinnie(1), rodzinnie(20), rodzinny(16) sibling, parents, family, family, family, family</td>
</tr>
<tr>
<td>wz</td>
<td>31</td>
<td>wzorzec(9), wzrost(4) speculum, high</td>
</tr>
<tr>
<td>śr</td>
<td>65</td>
<td>średni(3), średnia(47), średnio(10), średni(1), układ(3), średkowy(1) medium, diameter, medium, Wednesday, middle, middle</td>
</tr>
<tr>
<td>zab</td>
<td>90</td>
<td>zabieg(14), zaburzenie(76) surgery, disorder</td>
</tr>
<tr>
<td>zaw</td>
<td>29</td>
<td>zawiesina(23), zawód(6) suspension, profession</td>
</tr>
<tr>
<td>zal</td>
<td>31</td>
<td>załamek(10), założyć(5), załączyć(16) crinkle, put on, attach</td>
</tr>
</tbody>
</table>

Table 3: Test set abbreviation expansions in numbers

from all the possible word forms in our distributional model, those whose similarity to a particular abbreviation was higher than 0.2 for a language model created on forms, see Section (4). These candidates form the second expansion list of 259 elements (SL). The numbers of occurrences of all expansions of these three lists in the training data are given in Table 2.

6.2 Clustering and Manual Annotation

To check whether abbreviation usages form any differentiable clusters, we identified all their occurrences in the training data. For each such occurrence, we determined the context vector, which was equal to the average of vectors of surrounding tokens. In the experiment, we set the context as three tokens before and after each abbreviation. Then, we clustered occurrences of the abbreviation via the Chinese whispers algorithm (Biemann, 2006) which does not impose defining a priori a number of clusters. As we aimed to select examples of various interpretations of the same abbreviation and various usage of the same interpretation of the abbreviation, we established quite a high level of similarity between nodes in the initial graph. The similarity was counted as the cosine between vectors and we set it experimentally to 0.7 (it had no theoretical justification). Increasing the parameter of similarity we obtain more clusters and they represent higher granularity of abbreviation contexts.

For each cluster, we randomly selected from 2 to 6 elements (depending on the cluster size) and manually annotated them and the representative elements of the cluster pointed out by the algorithm, with proper expansions in the data. 85 elements used in this manual annotation constitutes the third list (CL) used in our experiments. In Table 2 the number of annotated examples in both train and test data are give. In test data a very high variance of abbreviations occurrences caused mainly by the big number of clusters obtained for the most frequent abbreviation (op) can be seen (from 29 to 1785).
6.3 Training Data

The core training set (SIM) is constructed via simulation by shortening word forms beginning with any of the abbreviation from the appropriate list processed in the exact experiment: AL, SL, and CL. The longest list AL contains 1345 potential expansions, SL limits the number of potential expansions to 259 elements while the manually created list CL has 85 elements. However, the SIM set may be biased as some of the words from these lists might be never shortened. What is more, in some typical places in which the chosen abbreviations occur, the full form may never or almost never be used. To check the real value of such simulated training set and, to test if a much smaller training set could be sufficient for this task, we also prepared manually annotated training set. It was built from all the manually annotated examples (a procedure described in 6.2). In Table 2, there is a comparison of the numbers of considered abbreviations in simulated (depending on the chosen expansion list) and manually annotated training and test data. As it turned out that nearly every abbreviation can also be an acronym, and one of them oko prawy – op occurs many times in our annotated data, to make comparisons more complete we also prepared a version of our training data (SIM-ac) in which two consecutive words recognized during manual annotation as a possible full form of an acronym, are abbreviated to the sequence of their first letters.

7 Neural Net Architecture

In the experiment, we used bidirectional LSTM nets as being most frequently judged as good for sequence processing. We formulated our task as a prediction task in which we predict a word on the basis of its context (and, optionally, on the basis of a representation of the abbreviation used instead of it). As clinical notes are short, concise and frequently change subject, we assign a label (which is a full word form) to a word on the basis of its left and right contexts of 3 or 5 words.

Input to a net consists of a subset of the following data (names given after features descriptions are used in Table 4 headings):

- word vectors form the models trained on the entire dataset,
- POS tags encoded as one-hot vector (pos) (31 most frequent categories),
- vector representing an abbreviation itself (padded with zeros if needed),
- all possible longer versions of the particular abbreviations coded in a vector representing all possible forms of all the abbreviations taken into account. This vector was added as the additional information which was combined with the LSTM output layer (only one output value is considered),

Two net architectures tested are given in Figure 1. On the left side, a basic BiLSTM net with input consisting of seven word representation is shown (the central word is an abbreviation – being actually in the data or inserted in place of a full word form in the simulation variant). Representation consisted either of word embeddings only, or of embeddings concatenated with the POS one-hot encodings. We also tested variants in which only context words were used. The architecture given on the right takes only context words as input. The additional input vector represents all valid extensions of a given abbreviation (cd). In both cases, the last layer is a standard classification dense layer with a sigmoid activation function. Its size depends on the length of a particular extensions list. The implementation is done in Keras with Tensorflow backend. The Adam optimizer was used and the other settings are standard values used in Keras implementation.

8 Results

The net architecture for further experiments was chosen on the basis of the 10-fold cross validation results for one configuration, see Table 4.

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2The picture was obtained using Netron https://lutzroeder.github.io/netron/
The number of epochs was established on the basis of validation on 1/10th of the training data while learning on the remaining training set to 2 for models which use big simulated data, and 5 for models which use only small annotated data. The batch size was equal to 32 and 1 respectively. Apart from the first model which does not take into account either the abbreviation representation (c) or the list of possible expansions (cd), other results do not vary much. We decided to choose the second best variant with the list of possible expansions added as an additional layer but with the three not five word context. As our annotated data is not big, we preferred a comparable model with fewer parameters.

The chosen set of features (second column in Table 4) was used for building models for all abbreviation lists and four variants of training sets (only annotated data – ANOT, only simulated – SIM (only word abbreviations) and SIM-ac (word abbreviations and acronyms), and the sum of both.

The results on cross validation (Table 5) are the best for simulated data, probably because of their size and repetitiveness. Only for the longest expansion list the results are the best on the smallest annotated set. The small number of examples could have just been memorized more easily. With this one exception it is also generally the rule that the bigger the training set the better, although adding annotated examples lowered the results slightly for SL list.

The results for the test set (Table 6) are better than those obtained for the cross validation on the training set in many cases. This is probably due to the small size of the test data set and many occurrences of the easy to resolve cases, for example the frequent occurrences of the ‘op’ abbreviation, which was correctly identified as ‘opakowanie’ ‘package’. However, models trained on the simulated set alone, performed significantly worse in terms of the recall (precision only deteriorated a little).

Using simulated data for training models has one important advantage – it saves time and effort. But there are also some disadvantages which have to be carefully analyzed. A few examples of miss-interpretation of ‘pr’ strings are given in Table 7. In our particular task, the possible problems can have different sources. First, some abbreviations are never (or almost never) expanded within the corpus. These are for example very common acronyms (like OP – ‘right eye’) which are rarely written in the full form, or an abbreviation meta which is never used in its full form metastaza ‘metastasis’ in our corpus. We did not fully address the problem in this work and phrase extensions which were recognized during manual annotation were added manually to the expansion lists. The second problem is that some words are never abbreviated, but we automatically added them to our expansion lists making the problem harder to solve. However, the good results obtained for the AL list show that this situation was not very confusing for our models (which have access to annotated data). The third problem is the fact that the contexts in which the abbreviated form are used may differ from the contexts where the full form occurs. If in some contexts, only abbreviations are used and the full form never occurs, it is not possible to learn this pattern. For example, when prescribing the number of medicine packages, doctors always use op instead of opakowanie ‘package’, e.g. Lantus (1 op. 30%). Our experiment confirmed that this is really the case. Results obtained by the models trained on simulated data only, although having very good cross validation results, have much worse recall on test data than models trained on annotated examples. However, adding annotated data to the simulated train set improved the results. For all but the AL list, the results obtained on the entire data even outperformed those obtained for the annotated data.

9 Conclusions

In the paper, we wanted to test if simulated abbreviations can be used to expand ambiguous ad hoc abbreviations in medical notes. Although simulation of the training data is a very useful practice, as manual data annotation is an expensive and time consuming process, our work shows that the obtained results are not always satisfactory. The F1 measure we obtained is below the artificially established baseline (the F1 measure equal to 0.64). Moreover, the experiments show that annotation of a small number of thoroughly selected examples of abbreviation occurrences gives satisfactory results for the task with the F1 measure equal to 0.92. It significantly outperforms the artificial baseline – the most common expansion e.g. the standard baseline for the word sense disambiguation task. However, the best results are obtained when the simulated data are combined with man-
Table 4: Results for 10-fold cross validation for different bidirectional LSTM settings for one training set (a subset of randomly selected cluster elements) and a chosen extension list. In all but the sixth case, word embeddings based on word forms were used. Additional information used in the models: pos – part of speech, c – vector representing an abbreviation itself, cd – vector coding possible extensions of the particular abbreviations (architecture from Figure 1b).

<table>
<thead>
<tr>
<th>Weighted Precision</th>
<th>Recall</th>
<th>F1</th>
<th>Weighted Precision</th>
<th>Recall</th>
<th>F1</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.562</td>
<td>0.690</td>
<td>0.662</td>
<td>0.691</td>
<td>0.689</td>
<td>0.677</td>
</tr>
<tr>
<td>0.652</td>
<td>0.770</td>
<td>0.757</td>
<td>0.764</td>
<td>0.764</td>
<td>0.764</td>
</tr>
<tr>
<td>0.597</td>
<td>0.721</td>
<td>0.702</td>
<td>0.719</td>
<td>0.717</td>
<td>0.711</td>
</tr>
<tr>
<td>0.367</td>
<td>0.469</td>
<td>0.458</td>
<td>0.471</td>
<td>0.472</td>
<td>0.475</td>
</tr>
<tr>
<td>0.391</td>
<td>0.502</td>
<td>0.493</td>
<td>0.493</td>
<td>0.505</td>
<td>0.492</td>
</tr>
<tr>
<td>0.368</td>
<td>0.476</td>
<td>0.465</td>
<td>0.472</td>
<td>0.478</td>
<td>0.473</td>
</tr>
</tbody>
</table>

Table 5: Results for 10-fold cross validation of the selected net architecture for all extension lists and training set variants (notation explained in the text). The three potential extensions lists for simulated training sets: AL – all words being potential expansions, SL – all the possible words in our distributional model whose similarity to a particular abbreviation was higher than 0.2 for a language model created on forms, CL – annotations of randomly selected cluster elements. The best results for each expansion list are shown in bold.

<table>
<thead>
<tr>
<th>Trainset</th>
<th>List</th>
<th>AL</th>
<th>SL</th>
<th>CL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>P</td>
<td>R</td>
<td>F1</td>
<td>P</td>
</tr>
<tr>
<td>ANOT-rd</td>
<td>0.874</td>
<td>0.885</td>
<td>0.875</td>
<td>0.809</td>
</tr>
<tr>
<td>ANOT</td>
<td>0.854</td>
<td>0.869</td>
<td>0.853</td>
<td>0.855</td>
</tr>
<tr>
<td>SIM</td>
<td>0.864</td>
<td>0.872</td>
<td>0.866</td>
<td>0.896</td>
</tr>
<tr>
<td>ANOT+SIM</td>
<td>0.864</td>
<td>0.872</td>
<td>0.866</td>
<td>0.893</td>
</tr>
</tbody>
</table>

Table 6: Results for the test set of the models trained on different datasets for all extension lists (notation explained in the text). The best results for each expansion list are shown in bold. The artificial baseline results, when we consider only those expansions which really occurred in the data and the most frequent expansion is taken as a solution, are (weighted) $P=0.568$, $R=0.742$, $F1=0.64$. Most of our results are well above this baseline and only models trained on simulated data gave lower results on two expansions lists.

<table>
<thead>
<tr>
<th>Model trained on</th>
<th>List</th>
<th>AL</th>
<th>SL</th>
<th>CL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>P</td>
<td>R</td>
<td>F1</td>
<td>P</td>
</tr>
<tr>
<td>ANOT</td>
<td>0.914</td>
<td>0.926</td>
<td>0.917</td>
<td>0.891</td>
</tr>
<tr>
<td>SIM</td>
<td>0.800</td>
<td>0.482</td>
<td>0.556</td>
<td>0.770</td>
</tr>
<tr>
<td>SIM-ac</td>
<td>0.907</td>
<td>0.386</td>
<td>0.441</td>
<td>0.888</td>
</tr>
<tr>
<td>ANOT+SIM</td>
<td>0.947</td>
<td>0.749</td>
<td>0.824</td>
<td>0.911</td>
</tr>
<tr>
<td>ANOT+SIM-ac</td>
<td>0.947</td>
<td>0.715</td>
<td>0.798</td>
<td>0.915</td>
</tr>
</tbody>
</table>

As the results obtained for the SL expansion list (a list of all words from the data whose dis-
tributional similarity was higher than 0.2) and the ANOT+SIM training data are very good, it would be interesting to test how important the selection of annotated examples is and to test how many manually annotated data is necessary for obtaining satisfactory results. In the future work we want to test our method on a large set of abbreviations and include strings which are ambiguous between words and abbreviations.

Acknowledgments

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References


Multi-Task, Multi-Channel, Multi-Input Learning for Mental Illness Detection using Social Media Text

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Abstract

We investigate the impact of using emotional patterns identified by the clinical practitioners and computational linguists to enhance the prediction capabilities of a mental illness detection (in our case depression and post-traumatic stress disorder) model built using a deep neural network architecture. Over the years, deep learning methods have been successfully used in natural language processing tasks, including a few in the domain of mental illness and suicide ideation detection. We illustrate the effectiveness of using multi-task learning with a multi-channel convolutional neural network as the shared representation and use additional inputs identified by researchers as indicative in detecting mental disorders to enhance the model predictability. Given the limited amount of unstructured data available for training, we managed to obtain a task-specific AUC higher than 0.90. In comparison to methods such as multi-class classification, we identified multi-task learning with multi-channel convolutional neural network and multiple-inputs to be effective in detecting mental disorders.

1 Introduction

Social media platforms have revolutionized the way people interact as a society and have become an integral part of everyday life where many people have started sharing their day to day activities on these platforms. Such real-time data portraying one’s daily life could reveal invaluable insights into one’s cognition, emotion, and behavioral aspects. With its rapid growth among different demographics and being a source enriched with valuable information, social media can be a significant contributor to the process of mental disorder and suicide ideation detection.

In the domain of mental illness detection and especially when using social media text, lack of sufficiently-large annotated data and the inability to extract explanation on the derived outcome have restricted researchers to use traditional machine learning algorithms other than state-of-the-art methods such as deep neural networks. The proposed research explores the feasibility of applying the state-of-the-art processes in combination with features identified using manual feature engineering methods to enhance the prediction accuracy while maintaining low false negative and false positive rates. Also, this research looks into detecting multiple mental disorders at the same time by sharing lower level features among the different tasks. Intuitively, learning multiple mental disorders using a single neural network architecture in comparison to using a single network to identify only one mental illness is a logical approach considering the psychological and linguistic characteristics shared among the individuals susceptible to being diagnosed with different mental disorders.

Mental illness detection in social media using Natural Language Processing (NLP) methods is considered as a difficult task due to the complex nature of mental disorders. According to the American Psychiatric Association (2013), a mental disorder is a “syndrome characterized by a clinically significant disturbance in an individual’s cognition, emotion regulation, or behavior that reflects a dysfunction in the psychological, biological, or developmental processes underlying mental functioning”. Mental disorders have become and continue to be a global health problem. More than 300 million people from varied demographics suffer from depression (World Health Organization, 2018a), and have broader implications where 23% deaths in the world were caused due to mental and substance use disorders (World Health Organization, 2014). In Canada, one in every five Canadians has experienced some form of mental ill-
The adverse impact of mental disorders is prominent when looking into the number of Canadians who have committed suicide, where 90% of them were suffering from some form of a mental disorder (Mental Health Commission of Canada, 2016).

When considering the current treatment procedures for mental illnesses, the first step is to screen the individual for symptoms using questionnaires. With such an approach, the interviewee could be more vulnerable to memory bias and also could adapt to the guidelines prescribed by the assessor. Using such screening procedures might not expose the actual mental state of an individual, and hence, the prescribed treatments could be inadequate. Also, due to various socio-economic aspects, people with mental disorders have not been able to receive adequate treatments. The lack of sufficient treatments can be seen in countries with all types of income levels. For example, between 76% to 85% of the people from countries in low to middle-class income do not receive sufficient treatments for their mental illnesses, while 35% to 50% of the people from high-income countries do not receive adequate treatments (World Health Organization, 2018b). In addition to insufficient treatments, social stigma and discrimination have prohibited people from getting proper treatment and social support (World Health Organization, 2018b). Due to the rapid growth in social media users within different demographics (Statista, 2017), and the abundance of information that can be extracted about the users could bring invaluable insights that can be used to detect signs of mental illnesses and suicide ideation that can be challenging to obtain using structured questionnaires. Taking the factors mentioned above into consideration, we have proposed a solution that incorporates certain profound features manually engineered by researchers into a multi-task learning architecture, to enhance the model predictability to distinguish neurotypical users from users susceptible to mental disorders. We hope that our research will encourage other researchers to investigate further the possibilities of incorporating verified manually engineered features into architectures similar to the proposed one, to enhance the prediction accuracies in identifying users susceptible to mental disorders.

Key Contributions

- We demonstrate the applicability of a convolutional neural network with a multi-channel architecture on different classification tasks using unstructured and limited social media data.
- We illustrate the use of multi-task learning to predict users susceptible to depression and PTSD (Post Traumatic Stress Disorder).
- We built an emotion classifier to identify the emotion category (i.e., sad, anger, fear, joy) associated with the tweets posted by the users and used those categories as multiple inputs within the deep neural network architecture. We also explore the impact of using metadata (age and gender) as multiple inputs to enhance the model predictability.

2 Ethical Considerations

It is of greater importance to follow strict guidelines on ethical research conduct when the research data resembles vulnerable users who could be compromised. The researchers working with data that could be used to single out individuals must take adequate precautions to avoid further psychological distress. During our research, we have given thorough considerations to these ethical facets and have adopted strict guidelines to ensure the anonymity and privacy of the data. Similar to the guidelines proposed by Benton et al. (2017a), we have exercised strict hardware and software security measures. Our research does not involve any intervention and has focused mainly on the applicability of machine learning models in determining users susceptible to mental disorders.

3 Related work

As social media has become an integral part of ones’ day-to-day-life, it will be insightful to identify to what extent an individual has disclosed her/his personal information and whether accurate and sufficient information is being published to determine whether or not a person has a mental disorder. Considering the Twitter platform, rather than just sharing depressed feelings, users...
are more likely to self-disclose to the extent where they reveal detailed information about their treatment history (Park et al., 2013). The same level of self-disclosure can be identified in the Reddit forums (Balani and De Choudhury, 2015) and specifically by users with anonymous accounts (Pavalanathan and De Choudhury, 2015). Also, it was identified that personality traits and meta-features such as age and gender could have a positive impact on the model performances when detecting users susceptible to PTSD and depression (Preot et al., 2015). Similarly, we have also identified that age and gender as multiple inputs have positively impacted model predictability when used with multi-task learning.

Text extracted from social media platforms such as Twitter, Facebook, Reddit, and other similar forums has been successfully used in various natural language processing (NLP) tasks to identify users with different mental disorders and suicide ideation. Social media text was used to classify users with insomnia and distress (Jamison-Powell et al., 2012; Lehrman et al., 2012), postpartum depression (De Choudhury et al., 2013a,b, 2014), depression (Resnik et al., 2015a, 2013, 2015b; Schwartz et al., 2014; Tsugawa et al., 2015), Post-Traumatic Stress Disorder (Coppersmith et al., 2014a,b), schizophrenia (Loveys et al., 2017) and many other mental illnesses such as Attention Deficit Hyperactivity Disorder (ADHD), Generalized Anxiety Disorder, Bipolar Disorder, Eating Disorders and obsessive-compulsive disorder (OCD) (Coppersmith et al., 2015a).

With the advancements in neural network-based algorithms, more research has been conducted successfully in detecting mental disorders, despite the limited amount of data. Kshirsagar et al. (2017) have used recurrent neural networks with attention to detect social media posts resembling crisis. Husseini Orabi et al. (2018) demonstrated that using convolution neural network-based architectures produces better results compared to recurrent neural network-based architectures when detecting users susceptible to depression. Even though our experiments are to categorize users into three classes (i.e., control, depression, PTSD), the proposed multi-channel architecture have produced comparable results to the ones presented by Husseini Orabi et al. (2018) using binary classification to distinguish users susceptible to depression.

4 Proposed solution

The proposed solution consists of two key components. The first identifies the type of emotion expressed by each user using the model trained on the WASSA 2017 shared task dataset. The identified emotion categories are used as multiple inputs within the multi-task learning environment. The second component is the model that predicts users susceptible to PTSD or depression. When structuring the two neural network models (i.e., for emotion classification and mental illness detection), a common base architecture is used. The base architecture is a multi-channel Convolutional Neural Network (CNN) with three different kernel sizes (i.e., 1, 2, and 3). Through experiments, we identified that the multi-channel CNN architecture manages to produce better validation accuracies compared to the accuracies produced by Recurrent Neural Network (RNN) based models, which are commonly used with sequence data.

4.1 Data

Emotion Classification We use the data from the 8th Workshop on Computational Approaches to Subjectivity, Sentiment & Social Media Analysis (WASSA-2017). The data was used in the shared task to identify emotion intensity (Mohammad and Bravo-Marquez, 2017). The tweets in the dataset were assigned with the labels: anger, fear, joy and sadness, and their associated intensities. Table 1 presents the detailed statistics of the dataset.

<table>
<thead>
<tr>
<th>Emotion</th>
<th>Train</th>
<th>Test</th>
<th>Dev</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anger</td>
<td>857</td>
<td>760</td>
<td>84</td>
<td>1701</td>
</tr>
<tr>
<td>Fear</td>
<td>1147</td>
<td>995</td>
<td>110</td>
<td>2252</td>
</tr>
<tr>
<td>Joy</td>
<td>823</td>
<td>714</td>
<td>79</td>
<td>1616</td>
</tr>
<tr>
<td>Sadness</td>
<td>786</td>
<td>673</td>
<td>74</td>
<td>1533</td>
</tr>
<tr>
<td>Total</td>
<td>3613</td>
<td>3142</td>
<td>347</td>
<td>7102</td>
</tr>
</tbody>
</table>

Table 1: The number of tweets under each emotion category

The dataset contains 194 tweets that belong to multiple emotion categories. For example, the tweet: “I feel like I am drowning. #depression #anxiety #failure #worthless” is associated with the labels ‘fear’ and ‘sadness’. When training the model, we created a single training dataset by combining both the train and test data and tested the trained model on the development dataset. The main reason for using such an approach is to im-
prove the neural network model training by providing as much data as possible so that model overfitting will be reduced while increasing the model generalization. During our training, we did not take into consideration the emotional intensity and expect to use it in our future research as an additional input.

**Mental Illness Detection** To detect whether a user is a neurotypical user or if the user is susceptible to having either PTSD or depression, we used the dataset from the Computational Linguistics and Clinical Psychology (CLPsych) 2015 shared task (Coppersmith et al., 2015b). Table 2 presents the detailed statistics of this dataset.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>PTSD</th>
<th>Depressed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of users</td>
<td>572</td>
<td>246</td>
<td>327</td>
</tr>
<tr>
<td>Average age</td>
<td>24.4</td>
<td>27.9</td>
<td>21.7</td>
</tr>
<tr>
<td>Gender (female)</td>
<td>74%</td>
<td>67%</td>
<td>80%</td>
</tr>
<tr>
<td>distribution per class</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2: CLPsych 2015 shared task dataset statistics

**Preprocessing:** All the URLs, @mentions, hashtags, RTweets, emoticons, emojis, and numbers were removed. We removed a selected set of stopwords but kept first, second, and third person pronouns. It was discovered that users susceptible to mental disorders such as depression have frequently used the first-person singular pronouns compared to neurotypical users (Pennebaker et al., 2007). Also, the punctuation marks except for a selected few were removed. The full stop, comma, exclamation point, and the question mark were kept while removing all the other punctuation marks. The NLTK (Natural Language Toolkit) tweet tokenizer was used to tokenize the tweets. We selected 200,000 unique tokens to build the vocabulary, rather than choosing all the unique words, which could lead to sparse word vectors with high dimensionality.

**Vocabulary Generation:** To obtain an enriched dictionary containing the most relevant terms, we introduced a novel approach instead of the traditional approach used in many deep learning APIs (e.g., Keras deep learning high-level API\(^1\)). Our approach takes into account the top ‘K’ terms based on their term frequency and inverse document frequency (TF-IDF) scores. To build the dictionary, first, we calculated the TF-IDF values under each user (i.e., by considering all the train/validation tweets of a single user as one single document). Then we took the maximum score out from all the assigned TF-IDF scores given the word. The reason for taking the maximum is to extract the words identified as closely related to a given user. Based on the computed TF-IDF scores, we picked the top ‘K’ words (K=200,000) to construct the vocabulary. A dictionary created using the above approach allows the model to capture the underlying relationships between the critical words. In comparison to the word frequency-based approach, using the vocabulary based on the TF-IDF scores has produced relatively better results for the recorded matrices (refer Table 4). When analyzing the model’s prediction accuracy and loss (i.e., on training and validation data) over five-fold cross-validation, we identified that the model trained using the TF-IDF based vocabulary has been more stable with less randomness compared to the model trained using the vocabulary based on word frequencies.

When choosing the maximum sequence length for the input data, it is essential to capture as much information as possible from each user, especially given consideration to the research domain of mental illness detection. Since we have concatenated all the individual tweets belonging to one user as a single string, a high variance in the sequence length was identified among users. On average, a single user has used around 15,800 tokens, where the maximum number of tokens used by an individual user is nearly 64,800. Rather than experimenting with different sequence lengths, we selected the maximum length for the sequence by adding three standard deviations to the average sequence length covering 99% of users with a sequence length of 46,200 tokens. The shorter sequences were padded with zeros (to the end of the sequence), and the longer sequences were truncated (from the end of the sequence).

**Model Architecture** The selected model architecture consists of three main components: multi-task learning, CNN with multi-channel, and multi-inputs. Multi-task learning is known to be successful when the data is noisy and limited so that when trying to learn one task, it could gain additional information from the other tasks to identify the most relevant features. Learning a shared representation so that individual tasks can benefit

\(^1\)https://keras.io/preprocessing/text/
from one another (Caruana, 1997) can be considered as one of the most appropriate architectures when trying to detect multiple mental illnesses. Benton et al. (2017b) demonstrated the successful use of multi-task learning to recognize mental illnesses and suicide ideation. Different from their approach, we add multiple features discovered by researchers in the fields of computational linguistics and psychology, to enhance the model performances. We consider that it is important to identify the impact of manually engineered features on the model’s predictability. We also recognized that using a CNN multi-channel architecture is best suited for tasks dealing with limited unstructured data compared to RNN architectures or multilayer perceptrons (MLP).

We used a multi-channel model as the base model in both emotion classification (i.e., to detect anger, sadness, joy, fear) and mental illness detection (i.e., to detect PTSD and depression). The multi-channel model uses three versions of a standard CNN architecture with different kernel sizes. We identified that using different kernel sizes (different n-grams sizes) with Global Maximum Pooling produces better results compared to a standard CNN architecture. The optimal validation accuracies for both emotion and mental illness detection models were derived using three channels with kernel sizes 1, 2, and 3. Increasing the kernel sizes or the number of channels reduced the validation accuracies.

For the emotion classification task, the CNN in each channel was tested with 64 filters, same padding and a stride of 1 (distance between successive sliding windows). We used Rectified Linear Unit (ReLU) as the activation function. To normalize the data and to reduce the impact of model overfitting, we used the batch normalization layer and used a dropout (Srivastava et al., 2014) as the regularization technique with a probability of 0.2. As the final layer in each channel, we used global maximum pooling to reduce the number of parameters needed to learn so that it could further reduce the impact of model overfitting. The outputs from each global maximum pooling layers (from each channel) were concatenated and fed into a fully connected layer with four hidden units that use sigmoid activation to generate the output. All the inputs were sent through trainable embedding layers (randomly initialized) with a dimension of 300 for the emotion classification task and 100 for the mental illness detection task.

Throughout our research, we did not emphasize much on word embeddings as our primary objective was to identify the impact of merging features derived using deep learning methods with few of the notable features that were identified over the years by researchers on detecting mental illnesses. Even though our best results were obtained by instantiating the embedding layer weights with random numbers (refer Table 4), we conducted several preliminary experiments using word embeddings trained on the fastText (Joulin et al., 2017) algorithm. We decided to use fastText because given the unstructured nature of the twitter messages we could obtain a more meaningful representation by expressing a word as a vector constructed out of the sum of several vectors for different parts of the word (Bojanowski et al., 2017). One of the reasons for the low measurements could be due to the reason that we used fewer data to train our embeddings.

**Mental illness detection** When building the multi-task learning model to detect mental illnesses, the base model architecture (i.e., for the shared representation) has used a structure similar to the one used in emotion classification. The fundamental changes to the base model include: using 256 filters instead of 64 and using L1 kernel regularization in each convolution layer. We used the trained model on the emotion data to predict the emotion category of the individual Twitter messages in the CLPISych 2015 dataset. We grouped the predicted probabilities for each user under the different emotion categories by calculating the standard deviation. We used the predicted probabilities for each emotion category as multiple inputs when detecting neurotypical and depressed users, while age and gender were used as inputs when predicting users with PTSD. Before concatenating the multiple-inputs with the output from the multi-channel architecture, the multiple-inputs were transformed using a fully connected layer with 128 hidden units and ReLU activation. The output from the shared layers and the transformed multiple inputs were merged before being used as the input to the fully connected layer with three individual hidden units and sigmoid activation. Before applying multiple-inputs to the neural network architecture, all the relevant inputs were normalized using a minimum, maximum scaler initialized within the range 0 and 1. The neu-
ral network architecture used for the multi-task, multi-channel, multi-input model for mental illness detection is shown in Figure 1.

Model Training When training both the emotion and the mental illness detection models, we minimized the validation loss to learn the optimal neural network parameters. To train both the models, we used minibatch gradient descent with smaller batch sizes. Using a smaller batch size is known to stabilize model training while increasing the model generalizability. In many cases, the optimal results were obtained by using batch sizes smaller or equal to 32 (Masters and Luschi, 2018). In our experiments, we used batch sizes 32 and 8 respectively for training the emotion and mental illness detection models. Both models were trained for 15 epochs and used early stopping when the validation loss has stopped improving. The Adam optimizer (Kingma and Ba, 2014) was used when training both the models with the default learning rate of 0.001.

5 Results

5.1 Emotion classification

The emotion detection task was implemented as a multi-class, multi-label classification because the same Twitter message can belong to multiple classes. Since we would like to have independent probability values for each class rather than a probability distribution over the four classes, we used binary cross-entropy as the loss function. Having independent probability values is better-oriented towards the identification of independent emotion categories. Table 3 reports the emotion classification results obtained using multi-channel CNN (MCCNN), and in addition, results from several other experiments: CNN with max-pooling (CNNMax) and bidirectional Long Short Term Memory module (biLSTM) were reported for comparison.

<table>
<thead>
<tr>
<th></th>
<th>Acc (%)</th>
<th>F1 (%)</th>
<th>P (%)</th>
<th>R (%)</th>
<th>P rank (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCCNN</td>
<td>88.88</td>
<td>77.41</td>
<td>79.68</td>
<td>75.67</td>
<td>84.85</td>
</tr>
<tr>
<td>CNNMax</td>
<td>85.82</td>
<td>68.95</td>
<td>76.97</td>
<td>62.70</td>
<td>81.39</td>
</tr>
<tr>
<td>biLSTM</td>
<td>85.07</td>
<td>68.66</td>
<td>75.97</td>
<td>63.49</td>
<td>80.97</td>
</tr>
</tbody>
</table>

Table 3: Emotion multi-class, multi-label classification results

In Table 3, the recorded accuracy is based on the Keras API\(^2\) accuracy calculation on the multi-class, multi-label models where it takes into account the individual label predictions rather than a complete match (i.e., if there is more than one label per instance). The F1-score, precision, and recall measures are calculated based on the ‘macro’ averaging on the exact match and hence the low percentages. We have also reported the label ranking average precision score, which averages over the individual ground truth label per instance. This metric is mainly used in tasks that involve multi-label ranking. From the reported results, it can be seen that the multi-channel CNN model has given the best results compared to the standard CNN model and the RNN based model. Based on the outcome, we have used the above trained multi-channel CNN model to make predictions on the CLPsych 2015 individual tweets.

5.2 Mental illness detection

For detecting mental illnesses, we used binary cross-entropy loss as the loss function and sigmoid activation as the final layer activation. The data was sampled using Stratified Shuffle Split to maintain class distribution between 80%/20% of training and validation data. Our models were evaluated only on the validation data because the CLPSych 2015 shared task test data labels were not made available. To ascertain the model reliability, we performed 5-fold cross-validation and discovered that having multi-inputs on a multi-task, multi-channel architecture does increase the model performances. The recorded model accuracies were averaged over five folds with a standard deviation of 0.01.

Table 4 demonstrates the model performances according to different combinations of the multi-task, multi-channel, and multi-input architectures. To demonstrate the effectiveness of the proposed approach, we conducted several experiments using variants of two deep learning architectures; Convolutional Neural Networks, and Recurrent Neural Networks and also to measure a baseline, we used the shallow learning approach; Support Vector Machine. The experiments: MtMcMi (Multi-task Multi-channel, Multi-input), MtMc (Multi-task, Multi-channel), MtMcMiFT (Multi-task, Multi-channel, Multi-input, using FastText word representations), MtMcMiFr (Multi-task, Multi-channel, Multi-input, using word Frequen-
The metrics used for the evaluation are accuracy, precision, recall, F1-score, and Area Under the Receiver Operating Characteristic curve (AUC). The AUC score is used to compare the model performances where the average AUC is calculated with standard deviation. The standard deviation is used as a mechanism to identify variance among model performances, which could bring insight into the reliability of the trained model. For each experiment, we recognized that the standard deviation is approximately around 0.01, which provides an empirical confirmation that the sampling using stratified shuffle splits provides an accurate representation of the complete dataset.

The “MtMcMi” architecture uses features based on emotion as multi-inputs on the control and depressed users while age and gender were used on the users with PTSD. In comparison to “MtMc” which is multi-task, multi-channel without multi-inputs, we could see that using multiple inputs have increased the average AUC score, as well as most of the other evaluation matrices (i.e., precision, recall, and F1-score). Even though the increase could not be considered as significant, the potential room for improvement is high if provided with more accurate emotion prediction and additional profound features identified by researchers. Concerning the emotion detection task, we could have obtained considerable improvements in prediction accuracies if provided with additional data when training the deep learning model.

When analyzing the result for “MtMcMiFr”, which is using the Multi-task, Multi-channel, Multi-input architecture with the vocabulary created using the default word frequencies; we could identify that our proposed approach, which uses the vocabulary constructed using the weighted TF-IDF words has produced comparatively better results. Even though the gained improvements could

![Figure 1: Multi-task, multi-channel, multi-input model for mental illness detection](image)

<table>
<thead>
<tr>
<th>Model</th>
<th>Accuracy (%)</th>
<th>Avg. F1 (%)</th>
<th>Avg. Precision (%)</th>
<th>Avg. Recall (%)</th>
<th>Avg. AUC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MtMcMi</td>
<td>89.08</td>
<td>87.59</td>
<td>91.35</td>
<td>89.07</td>
<td>86.61</td>
</tr>
<tr>
<td>MtMc</td>
<td>88.55</td>
<td>86.89</td>
<td>91.96</td>
<td>88.53</td>
<td>83.06</td>
</tr>
<tr>
<td>MtMcMiFr</td>
<td>85.50</td>
<td>86.28</td>
<td>85.45</td>
<td>85.90</td>
<td>85.91</td>
</tr>
<tr>
<td>McMclass</td>
<td>92.00</td>
<td>75.69</td>
<td>76.73</td>
<td>89.05</td>
<td>78.25</td>
</tr>
<tr>
<td>bilstMMtMi</td>
<td>51.35</td>
<td>71.61</td>
<td>78.60</td>
<td>41.27</td>
<td>41.73</td>
</tr>
<tr>
<td>bilstMt</td>
<td>52.48</td>
<td>72.31</td>
<td>78.60</td>
<td>47.48</td>
<td>46.40</td>
</tr>
<tr>
<td>svmMc class</td>
<td>81.73</td>
<td>52.91</td>
<td>42.85</td>
<td>73.82</td>
<td>57.01</td>
</tr>
</tbody>
</table>

Table 4: Mental illness detection using multi-task, multi-channel, multi-input architecture. The labels ‘C’, ‘D’ and ‘P’ denotes ‘Control’, ‘Depressed’ and ‘PTSD’
not be considered as significant, the proposed method can be regarded as an effective approach when initiating the vocabulary, that provides a balance between the rare and frequently used words.

In the experiment “MtMcMiFT”, where we used the fastText embeddings layer with the number of dimensions equal to 100 as an input to the Multi-channel convolutional neural network, it was observed that the results obtained using the randomly initialized embedding layer are higher than with the fastText pre-trained embeddings. This could be due to embeddings not being trained on a sufficiently large dataset (we trained them only on the CLPSych 2015 dataset). In future work, we will conduct further research to enhance the embedding layer word representation by using state-of-the-art language modeling approaches trained on larger datasets.

The effectiveness of using convolutional neural network models can be identified when evaluating the results obtained using Recurrent Neural Network (RNN) based architectures. The “biLSTM-MmiMi” method uses a bidirectional Long Short Term Memory (“biLSTM”) model in a Multi-task, Multi-input design and comparatively has produced poor results for different combinations of hyperparameters. This could be due to several reasons such as the unstructured nature of the Twitter text as well as the non-existence of long-term dependencies. For example, our best results were obtained when using the kernel sizes (i.e., the number of consecutive tokens) one, two, and three and ones the kernel sizes are increased the overall model predictability decreases. When using a “biLSTM” model as the shared layer in multi-task learning without multiple inputs (“biLSTM-Mmi”), the results are somewhat better compared to when using multi-inputs.

To demonstrate the effectiveness of using multi-task learning to detect multiple mental disorders, we compared the proposed approach with multiclass classification to distinguish neurotypical users from users susceptible to having either PTSD or depression. First, we used a multi-channel convolutional neural network to predict the three classes (i.e., control, depress, and PTSD). In comparison to our proposed approach, we can identify that multiclass classification using CNN have produced slightly better results on two occasions, which is for average accuracy and recall under the control class. Through further analysis, we see that average precision, F1-score, and AUC scores are higher for all three classes when using the proposed approach. Overall the multiclass classification task has produced low scores (especially for precision, F1-score, and AUC) when detecting users susceptible to depression and PTSD while the proposed approach has contributed significantly better results. The better results could be due to the reason that depression is commonly identified among individuals with PTSD, and the shared layer has managed to learn such common characteristics while the task-specific layers have learned the individual features unique to each disorder.

As a baseline, we used the linear SVM classifier with TF-IDF features (200,000 features) in a multiclass classification task (i.e., svmMclass). When sampling the data, five splits of 80% training and 20% testing were created using the Stratified Shuffle Split method to maintain class distribution. We also computed a majority class baseline, which classifies everything in the largest class (the control class in the training data). It achieved an accuracy of 50.21% on the test data. Overall, we can see that using limited unstructured data with an architecture based on CNN have produced better results compared to the solution based on RNN. Notably, the multi-task, multi-channel architecture with multiple-inputs has provided the best results and confirms that using multiple-inputs has a positive influence on the overall model performances. Also, the appropriateness of using multi-task learning instead of multiclass classification to detect multiple mental disorders is highlighted. Similar to the fact that certain mental disorders share specific common symptoms (American Psychiatric Association, 2013), multi-task learning has managed to learn such characteristics through a shared representation followed with task-specific layers to identify the unique attributes to differentiate multiple mental disorders.

6 Comparison to related work

Even though our work could not be directly compared with (Benton et al., 2017b), we can identify that our model has produced competitive results, especially when comparing the AUC score for detecting users with PTSD and depression. Our best model has scored an AUC score >0.90 in identifying all three individual classes (control, depression, PTSD) in comparison to an AUC
score <0.80 for detecting PTSD and depression and an AUC score >0.90 for detecting the neurotypical users (Benton et al., 2017b). In the CLPSych 2015 shared task (Coppersmith et al., 2015b), Resnik et al. (2015a) have reported AUC scores of 0.86 (depression vs. control), 0.84 (depression vs. PTSD) and 0.89 (PTSD vs. control) and similarly Preatiuc-Pietro et al. (2015) have reported an average AUC score around 0.86 in differentiating neurotypical users from users susceptible to PTSD and depression. Even though we have produced better results using the validation dataset, we could not directly compare our results with the shared task participants as they have evaluated their models against the test dataset which was not made available to us. In our future work, we will conduct experiments using public forum post data extracted from platforms such as Reddit\(^3\). In comparison, the proposed approach can be tested with adequate adjustments to detect multiple mental disorders such as depression, anxiety, PTSD, and six others using the dataset introduced by Cohan et al. (2018). The authors have achieved an F1-score of 27.83% for multi-class classification and 53.56% and 57.60% respectively, when detecting depression and PTSD as binary classification tasks.

7 Conclusion

In this paper, we investigated the impact of merging features derived using deep neural network architectures with profound manually engineered features identified by researchers over the years using shallow learning to detect mental disorders using social media text. In particular, we have identified that by using a multi-channel convolutional neural network as a shared layer in a multi-task learning architecture with multiple-inputs (e.g., different emotion categories, age, and gender) have produced comparatively competitive results in detecting multiple mental disorders (in our case depression and PTSD). For future work, we will continue our research on suicide risk detection, and the temporal impact different mental disorders have on suicide ideation.

References


\(^3\)https://www.reddit.com/


Saif M. Mohammad and Felipe Bravo-Marquez. 2017. WASSA-2017 Shared Task on Emotion Intensity. In 8th Workshop on Computational Approaches to Subjectivity, Sentiment and Social Media Analysis.


Abstract

We present a system for automatically extracting pertinent medical information from dialogues between clinicians and patients. The system parses each dialogue and extracts entities such as medications and symptoms, using context to predict which entities are relevant. We also classify the primary diagnosis for each conversation. In addition, we extract topic information and identify relevant utterances. This serves as a baseline for a system that extracts information from dialogues and automatically generates a patient note, which can be reviewed and edited by the clinician.

1 Introduction

In recent years, electronic medical record (EMR) data have become central to clinical care. However, entering data into EMRs is currently slow and error-prone, and clinicians can spend up to 50% of their time on data entry (Sinsky et al., 2016). In addition, this results in inconsistent and widely variable clinical documentation, which present challenges to machine learning models.

Most existing work on information extraction from clinical conversations does not differentiate between entities that are relevant to the patient (such as experienced symptoms and current medications), and entities that are not relevant (such as medications that the patient says were taken by someone else).

In this work, we extract clinically relevant information from the transcript of a conversation between a physician and patient (and sometimes a caregiver), and use that information to automatically generate a clinical note, which can then be edited by the physician. This automated note-taking will save clinicians valuable time and allow them to focus on interacting with their patients rather than the EMR interface. We focus on linguistic context and time information to determine which parts of the conversation are medically relevant, in order to increase the accuracy of the generated patient note. In addition, the automatically generated notes can provide cleaner and more consistent data for downstream machine learning applications, such as automated coding and clinical decision support.

Figure 1 shows a synthetic example of the kind of medical conversation where context and time information are important.

![Figure 1: Synthetic conversation example.](image)

2 Related Work

Previous studies have shown that current EMR data are difficult to use in automated systems because of variable data quality (Weiskopf and Weng, 2013; Thiru et al., 2003; Roth et al., 2003). Weiskopf and Weng (2013) showed that EMR data is frequently incomplete, and is often not evaluated for quality. In addition, the variance in documentation style, abbreviations, acronyms, etc. make it difficult for algorithms to interpret the text.

Some recent work on machine learning methods for EMR data includes predicting mortality
and discharge diagnoses (Rajkomar et al., 2018), predicting unplanned hospital readmissions for 5k
patients by encoding EMR data with a convolutional neural network (Nguyen et al., 2018), and
predicting diagnosis codes along with text explanations (Mullenbach et al., 2018).

Although there is some existing work on generating text from structured data (Dou et al., 2018;
Lebret et al., 2016), very little work has been done in the clinical domain. Liu (2018) generated pa-
tient note templates with a language model, which was able to approximate the organization of the
note, but no new information from the patient encounter was used.

Du et al. (2019) introduced a system for extracting symptoms and their status (experienced or not)
from clinical conversations using a multi-task learning model trained on 3,000 annotated conver-
sations. However, their model was trained on a limited set of 186 symptoms and did not address
other medically relevant entities.

A latent Dirichlet allocation (LDA) model (Blei et al., 2003) is a topic modeling technique and has
been applied to clinical text to extract underlying useful information. For example, Bhattacharya
et al. (2017) applied LDA on structured EMR data such as age, gender, and lab results, showing that
the relevance of topics obtained for each medical diagnosis aligns with the co-occurring conditions.
Chan et al. (2013) applied topic modeling on EMR data including clinical notes and provided an em-
pirical analysis of data for correlating disease topics with genetic mutations.

3 Dataset

<table>
<thead>
<tr>
<th>Primary diagnosis</th>
<th>Dyads</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADHD</td>
<td>100</td>
</tr>
<tr>
<td>Depression</td>
<td>100</td>
</tr>
<tr>
<td>COPD</td>
<td>101</td>
</tr>
<tr>
<td>Influenza</td>
<td>100</td>
</tr>
<tr>
<td>Osteoporosis</td>
<td>87</td>
</tr>
<tr>
<td>Type II diabetes</td>
<td>86</td>
</tr>
<tr>
<td>Other</td>
<td>226</td>
</tr>
</tbody>
</table>

Table 1: Data distribution (ADHD: Attention Deficit Hyperactivity Disorder; COPD: Chronic Obstructive Pulmonary Disorder)

For training and testing our models, we use a dataset of 800 patient-clinician dialogues (dyads)
purchased from Verilogue Inc.1, which includes demographic information about the patient as well
as the primary diagnosis. The data consist of audio files and human-generated transcripts with speaker
labels. Table 1 shows the distribution of diagnoses in the dataset.

Since these data are proprietary, we also use a few transcripts of staged clinical interviews from
YouTube as examples. 2

4 Annotation

First, the conversation transcripts are automatically annotated for time phrases using Heidel-
Time, a freely available rule-based time phrase tagger (Strötegen and Gertz, 2010), as well as a lim-
ited set of common medical terms.

Two physicians then conduct manual annotation by correcting the automatic annotations and
making any necessary additions, using a custom-developed annotation interface. The following
types of entities are annotated: anatomical locations, diagnoses, symptoms, medications, reasons
for visit, referrals, investigations/therapies, and time phrases. A total of 476 conversations are an-
notated by a unique physician, and inter-annotator agreement is calculated using DKPro Statistics3
on 30 conversations which were annotated by both physicians. The agreement across all entity types
is 0.53 Krippendorffs alpha (Krippendorff, 2004) and 0.80 F1 (partial match).

We developed a custom annotation interface for labeling entities and their attributes in the tran-
scripts, shown in Figure 3. The software includes the ability to add new annotation types and at-
tributes, edit and delete previous annotations, and view the entire conversation for context.

1http://www.verilogue.com
2YouTube videos of simulated patient encounters were sourced by searching for the following terms: “medical
history”, “patient interview”, and “clinical assessment”. Our clinician team member watched potential videos in the search
list and selected only the ones that met the following criteria: 1) clinician asking a patient questions in simulated clinical
scenarios; 2) a subjective perception of adequate fidelity to real clinical encounters. The audio for these dialogues were
transcribed by a professional transcriptionist. Examples used in this paper:
1: https://www.youtube.com/watch?v=O2qYU8n4VsA, 2: https://www.youtube.com/watch?v=CUSxC-XHT2A, 3: https://www.youtube.com/watch?v=s_jIcAk1xeA
3https://dkpro.github.io/dkpro-statistics
5 Methods and experiments

The automated pipeline currently includes the following components: preprocessing, utterance type classification (questions, answers, statements, etc.), entity extraction (medications, symptoms, diagnoses, etc.), attribute classification (modality and pertinence), primary diagnosis classification, SOAP classification, and note generation. In this section we discuss each component in detail, including methods and results. See Figure 4 for a diagram of the system components.

5.1 Preprocessing and data splitting

Before passing the data to our models, the text of the transcripts is lowercased, and punctuation is separated from words using WordPunctTokenizer from NLTK (Steven Bird and Loper, 2009). For the utterance type and attribute classification tasks, each word in an utterance is represented as a word embedding. In this work, we use publicly available ELMo embeddings (Peters et al., 2018) trained on PubMed abstracts, as well as word2vec embeddings trained on PubMed.

Of the 476 annotated conversations, we randomly select 50 to use as a test set for entity extraction and attribute classification.

5.2 Utterance type classification

In order to understand the conversational context, it may be useful to know whether an utterance is a question or answer. To this end, we classify each utterance as one of the following types: question, statement, positive answer, negative answer, backchannel (such as ‘uh-huh’ or ‘yeah’) or excluded (incomplete or vague utterance).

The utterance type classification model is a two-layer bidirectional gated recurrent unit (GRU) neural network (Cho et al., 2014), implemented in PyTorch, with the architecture shown in Figure 5. We augment the training data with two external, publicly available datasets: the Switchboard corpus (Calhoun et al., 2010), and the AMI corpus⁵. We map the utterance labels from the AMI and Switchboard corpora to our set of labels, and add these data to our training set.

We evaluate the utterance type classifier on a set of 20 conversations, annotated independently by 2 annotators with inter-annotator agreement of 0.77 (Cohen’s kappa).

Table 2 shows the classification results by utterance type. As the most frequent type, statements are the easiest for the model to identify. The low performance of infrequent classes indicates that we could potentially improve performance by using an oversampling or regularization method.

5.3 Entity extraction

5.3.1 Time phrase extraction

In order to determine clinical relevance, it is important to know the time and duration of events in the patient history. We use HeidelTime to identify

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⁴http://evexdb.org/pmresources/vec-space-models/

⁵http://groups.inf.ed.ac.uk/ami/corpus/
time phrases in the transcripts, including times, dates, durations, frequencies, and quantities.

5.3.2 Clinical entity extraction

In addition to time phrases, we identify the following clinical concept types: anatomical locations, signs and symptoms, diagnoses, medications, referrals, investigations and therapies, and reasons for visit. To identify these entities, we search the transcript text for entities from a variety of medical lexicons, including the BioPortal Symptom lexicon ⁶, SNOMED-CT ⁷, the Consumer Health Vocabulary (CHV) ⁸, and RxNorm (a database of

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⁶https://bioportal.bioontology.org/ontologies
⁷http://www.snomed.org/
Table 2: Utterance type classification results

<table>
<thead>
<tr>
<th>Type</th>
<th>Instances</th>
<th>P</th>
<th>R</th>
<th>F_1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Question</td>
<td>539</td>
<td>0.72</td>
<td>0.49</td>
<td>0.59</td>
</tr>
<tr>
<td>Statement</td>
<td>2,347</td>
<td>0.82</td>
<td>0.83</td>
<td>0.82</td>
</tr>
<tr>
<td>AnswerPositive</td>
<td>195</td>
<td>0.36</td>
<td>0.41</td>
<td>0.38</td>
</tr>
<tr>
<td>AnswerNegative</td>
<td>82</td>
<td>0.74</td>
<td>0.34</td>
<td>0.47</td>
</tr>
<tr>
<td>Backchannel</td>
<td>494</td>
<td>0.56</td>
<td>0.76</td>
<td>0.64</td>
</tr>
<tr>
<td>Excluded</td>
<td>131</td>
<td>0.20</td>
<td>0.16</td>
<td>0.18</td>
</tr>
<tr>
<td>Average</td>
<td>3,788</td>
<td>0.72</td>
<td>0.72</td>
<td>0.71</td>
</tr>
</tbody>
</table>

Table 3: Entity extraction results

<table>
<thead>
<tr>
<th>Type</th>
<th>Instances</th>
<th>P</th>
<th>R</th>
<th>F_1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anatomical locations</td>
<td>328</td>
<td>0.79</td>
<td>0.45</td>
<td>0.57</td>
</tr>
<tr>
<td>Diagnosis</td>
<td>346</td>
<td>0.88</td>
<td>0.62</td>
<td>0.72</td>
</tr>
<tr>
<td>Investigation or therapy</td>
<td>239</td>
<td>0.42</td>
<td>0.24</td>
<td>0.31</td>
</tr>
<tr>
<td>Medication</td>
<td>579</td>
<td>0.55</td>
<td>0.79</td>
<td>0.65</td>
</tr>
<tr>
<td>Referral</td>
<td>61</td>
<td>0.11</td>
<td>0.11</td>
<td>0.11</td>
</tr>
<tr>
<td>Sign/symptom</td>
<td>650</td>
<td>0.82</td>
<td>0.38</td>
<td>0.52</td>
</tr>
<tr>
<td>Time expression</td>
<td>1286</td>
<td>0.98</td>
<td>0.64</td>
<td>0.77</td>
</tr>
<tr>
<td>Average</td>
<td>3,489</td>
<td>0.80</td>
<td>0.56</td>
<td>0.64</td>
</tr>
</tbody>
</table>

Table 4: Modality classification results

<table>
<thead>
<tr>
<th>Type</th>
<th>Instances</th>
<th>P</th>
<th>R</th>
<th>F_1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Actual</td>
<td>504</td>
<td>0.87</td>
<td>0.80</td>
<td>0.83</td>
</tr>
<tr>
<td>Negative</td>
<td>144</td>
<td>0.63</td>
<td>0.64</td>
<td>0.64</td>
</tr>
<tr>
<td>Possible</td>
<td>5</td>
<td>0.09</td>
<td>0.40</td>
<td>0.14</td>
</tr>
<tr>
<td>None</td>
<td>91</td>
<td>0.59</td>
<td>0.71</td>
<td>0.65</td>
</tr>
<tr>
<td>Average</td>
<td>744</td>
<td>0.78</td>
<td>0.76</td>
<td>0.77</td>
</tr>
</tbody>
</table>

Table 5: Pertinence classification results

<table>
<thead>
<tr>
<th>Type</th>
<th>Instances</th>
<th>P</th>
<th>R</th>
<th>F_1</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADHD</td>
<td>126</td>
<td>0.54</td>
<td>0.41</td>
<td>0.28</td>
</tr>
<tr>
<td>COPD</td>
<td>22</td>
<td>0.20</td>
<td>0.45</td>
<td>0.28</td>
</tr>
<tr>
<td>Depression</td>
<td>32</td>
<td>0.27</td>
<td>0.81</td>
<td>0.41</td>
</tr>
<tr>
<td>Influenza</td>
<td>246</td>
<td>0.72</td>
<td>0.83</td>
<td>0.77</td>
</tr>
<tr>
<td>Other</td>
<td>312</td>
<td>0.79</td>
<td>0.51</td>
<td>0.62</td>
</tr>
<tr>
<td>None</td>
<td>6</td>
<td>0.32</td>
<td>1.00</td>
<td>0.48</td>
</tr>
<tr>
<td>Average</td>
<td>744</td>
<td>0.68</td>
<td>0.61</td>
<td>0.62</td>
</tr>
</tbody>
</table>

5.3.3 Attribute classification

After extracting relevant entities, we classify them according to several attributes, including modality (i.e., whether the events were actually experienced or not) and pertinence (i.e., to which disease the entities are relevant, if any). For example, a patient might mention a medication that they have not actually taken, so we would not want to record that medication as part of the patient’s history. In these cases, the context of the conversation, as well as time information, is crucial to recording the patient’s information accurately.

The attribute classifier is a support vector machine (SVM) trained with stochastic gradient descent using scikit-learn (Pedregosa et al., 2011). Each entity is represented as the average word embedding, concatenated with the word embeddings for the previous and next 5 words. We also include the speaker code of the utterance in which the entity appears. We train the model on 252 conversations and test on 50 for which we have human-assigned modality and pertinence labels.

We classify entities into the following modality categories: actual, negative, possible, or none. Table 4 shows the results of modality classification on the test set of 50 conversations. Since the majority of entities have a modality of ‘actual’, the model performs the best on this class. Entities are also classified as pertinent to one of the disease categories, or none. Table 5 shows the results of pertinence classification. Again we see that the classifier performs the best on the classes with more examples.

Modality classification performs fairly well with a context window of 5, likely because the relevant information can be found nearby in the text. However, pertinence classification is not as accu-
rate, perhaps because it requires more global information about what conditions the patient has. In some cases, pertinence may be purely determined by a clinician's medical knowledge, not the information present in the text.

In the future we hope to have more annotated data on which to train, which should improve the overall performance, especially for the smaller classes.

5.4 Clinical note generation

In the note generation phase, we convert the structured data from the previous steps (i.e., entities and their attributes) into a free text clinical note that resembles what a physician would have written. This involves organizing the entities according to a structured note organization and, finally, generating the text of the note.

5.4.1 SOAP entity classification

After extracting clinical entities, we classify them according to the traditional four sections of a clinical note: subjective (S), objective (O), assessment (A), plan (P) (Bickley and Szilagyi, 2013). We also add a ‘none’ category, which means that the given entity should not be included in the note.

The SOAP classifier is a neural network trained on each word of the entity, the previous and next five words, the speaker code of the corresponding utterance, and the type of entity. The text and context are represented as word embeddings using the PubMed word2vec model. Since the note generation requires special annotations, we currently only have 58 conversations for training, and 20 for test.

Table 6 shows the results of SOAP classification. The model is the most accurate at determining which information to exclude from the note.

<table>
<thead>
<tr>
<th>Type</th>
<th>Instances</th>
<th>P</th>
<th>R</th>
<th>F₁</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td>299</td>
<td>0.52</td>
<td>0.56</td>
<td>0.54</td>
</tr>
<tr>
<td>O</td>
<td>51</td>
<td>0.44</td>
<td>0.43</td>
<td>0.44</td>
</tr>
<tr>
<td>A</td>
<td>55</td>
<td>0.35</td>
<td>0.16</td>
<td>0.22</td>
</tr>
<tr>
<td>P</td>
<td>66</td>
<td>0.22</td>
<td>0.15</td>
<td>0.18</td>
</tr>
<tr>
<td>None</td>
<td>708</td>
<td>0.69</td>
<td>0.72</td>
<td>0.70</td>
</tr>
<tr>
<td>Average</td>
<td>1189</td>
<td>0.59</td>
<td>0.61</td>
<td>0.60</td>
</tr>
</tbody>
</table>

Table 6: SOAP classification results

5.4.2 SOAP note generation

Our current note generation step organizes the entities into the SOAP sections, and lists them along with their attributes. Actually generating full sentences that more closely resemble a physician-generated note is the next step for our future work.

5.5 Primary diagnosis classification

We classify the primary diagnosis for each conversation. The purpose of this task is to automatically identify the main diagnosis for billing codes. We train and test the models on a 5-fold cross validation of the 800 dyads. We apply 𝑡𝑓-𝑖𝑑𝑓 on the cleaned text of each dyad and then use logistic regression, SVMs with various parameter settings, and random forest models for classification. The 𝐹₁ score is calculated based on the human-assigned labels available in the transcription.

The primary diagnosis classifier performs reasonably well even without labeled entity features. The results for influenza achieve almost 0.90 𝐹₁ score, while the results for COPD and depression obtain an 𝐹₁ score of approximately 0.70. By inspecting the conversations, we find that visits with a primary diagnosis of depression mostly consist of general discussions related to daily routine, family life, and mood changes, which often result in misclassification probably because no medical terms are mentioned. By contrast, in patient visits where the primary diagnosis is influenza, the discussion is more focused on the disease.

The top words used by the classifier were H1N1, ache, temperature, sore, sick, symptom, swine, body, and strep, which possibly makes it easier to classify. On the other hand, COPD is misclassified mostly as the category ‘other’, which includes
diseases such as asthma, CHF (Congestive heart failure), hypercholesterolemia, atopic dermatitis, HIV/AIDS, prenatal visit, hypercholesterolemia. That is, the COPD dyads may be misclassified because of the presence of other respiratory diseases in the ‘other’ category. We plan to extend the diagnosis classifier to multi-label classification.

5.6 Topic modeling

Topic modeling is an unsupervised machine learning technique used to form \( k \) topics (i.e., clusters of words) occurring together, where \( k \) is usually chosen empirically. We perform topic modeling with LDA using the open-source gensim package (Řehůřek and Sojka, 2010) with varying numbers of topics \( k = (5, 10, 12, 15, 20, 25, 30, \text{ and } 40) \).

Due to their colloquial nature, patient-clinician conversations contain many informal words and non-medical terms. We remove common words, including stop words from NLTK (Steven Bird and Loper, 2009), backchannel words (like ‘uh-huh’), and words with frequencies above 0.05% of the total number words in all the documents (to reduce the influence of more generic words).

The topic modeling results are shown in Table 8; we choose \( k=12 \) topics because they provided the best topic distribution and coherence score. The words in each topic are reported in decreasing order of importance.

A manual analysis shows that topic 0 captures words related to ADHD and depression, while topic 1 is related to asthma and flu, and topic 3 is related to women’s health, and so on. This qualitative evaluation of topics shows that topic modeling can be helpful in extracting important information and identifying the dominant topic of a conversation. In our future work, we also plan to do a quantitative evaluation of topic modeling results using state-of-the-art methods such as the methodology proposed by Wallach et al. (2009).

We see the potential use of topic modeling to keep track of the focus of each visit, the distribution of word usage, categorization, and to group patients together using similarity measures. We also use it for relevant text extraction in the next section.

5.7 Relevant utterance extraction

Identifying the key parts of the doctor-patient conversation can be helpful in finding the relevant information. In the previous section, we observe that topic modeling can be helpful in identifying the underlying topics of the dyads. We also use topic modeling to extract the utterances relevant to the primary disease diagnosis.

We apply the following steps adapted from a publicly available text summarization method:\footnote{https://github.com/g-deoliveira/TextSummarization}

1. Fit the LDA model to all dyads.
2. Pass the dyads for each class to the LDA model to determine the class-wise topic distribution.
3. Select the dominant topics for each class using the topic weight matrix.
4. For each dyad within this subset:
We conduct experiments on all 800 dyads and the 11 dyads from YouTube. Topic modeling is performed exactly as described in the previous section, with 12 topics. The results are shown in Table 6 and 7. The three dyads shown are from open-source YouTube data focusing on (a) cough and rib pain, (b) women’s health and contraception, and (c) anxiety, respectively.

The results indicate a reasonable quality of relevant text extraction despite the limited amount of data. We can see that many of the utterances discussing the presenting problem are extracted. Since we do not have labels for the true relevance of the sentences to the disease, we are unable to provide any quantitative metrics, which is the subject of future work.

6 Conclusion & future work

The cumulative output of these models constitutes the initial automated system. Although for these experiments we used manual transcriptions, in practice the input would be from automatic speech recognition (ASR). Future research will include using ASR to record transcripts in real time, as well as expanding the types of entities we extract, identifying quantity, quality, and severity.

Diagnosis classification currently handles 6 classes only, and does not account for conditions other than the primary diagnosis that may be discussed in the conversation. We will also expand diagnosis classification to handle more classes, and to predict multiple diagnoses.
We have presented a system for extracting clinically relevant entities from physician-patient dialogues using linguistic context. The results show that clinical note-taking can be at least partially automated, saving clinicians valuable time. This system can result in a streamlined data entry process and a cleaner EMR note that can be used for analytics and automated decision making.

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Biomedical Relation Classification by Single and Multiple source Domain Adaptation

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Abstract

Relation classification is crucial for inferring semantic relatedness between entities in a piece of text. These systems can be trained given labelled data. However, relation classification is very domain-specific and it takes a lot of effort to label data for a new domain. In this paper, we explore domain adaptation techniques for this task. While past works have focused on single source domain adaptation for bio-medical relation classification, we classify relations in an unlabeled target domain by transferring useful knowledge from one or more related source domains. Our experiments with the model have shown to improve state-of-the-art F1 score on 3 benchmark biomedical corpora for single domain and on 2 out of 3 for multi-domain scenarios. When used with contextualized embeddings, there is further boost in performance outperforming neural-network based domain adaptation baselines for both the cases.

1 Introduction

In the biomedical domain, a relation can exist between various entity types like protein-protein, drug-drug, chemical-protein etc. Detecting relationships is a fundamental sub-task for automatic Information Extraction to overcome efforts of manual inspection, especially for growing biomedical articles. However, existing supervised systems are highly data-driven. This poses a challenge since manual labelling is a costly and time-consuming process and there is a dearth of labelled data in the biomedical domain covering all tasks and for new datasets. A system trained on a specific dataset1 may perform poorly on another, for the same task (Mou et al., 2016), due to dataset variance which can arise owing to sample selection bias (Rios et al., 2018).

Domain Adaptation aims at adapting a model trained on a source domain to another target domain that may differ in their underlying data distributions. Past work on domain adaptation for bio-medical relation classification has focused on single-source adaptation (Rios et al., 2018). However, multiple sources from related domains can prove to be beneficial for classification in a low-resource scenario.

In this paper, we perform domain adaptation for biomedical binary relation classification at the sentence-level. For single-source single target (SSST) we transfer between different datasets of protein-protein interaction, along with drug-drug interaction. We also explore multi-source single target (MSST) adaptation to incorporate more richness in the knowledge transferred by using additional smaller corpora for protein-protein relation and multiple labels for chemical-protein relation respectively. Given an unlabeled target domain, we transfer common useful features from related labelled source domains using adversarial training (Goodfellow et al., 2014). It helps to overcome the sampling bias and learn common indistinguishable features, promoting generalization, using min-max optimization. We adopt the Multinomial Adversarial Network integrated with the Shared-Private model (Chen and Cardie, 2018) which was originally proposed for the task of Multi-Domain Text Classification. It can handle multiple source domains at a time which is in contrast to traditional binomial adversarial networks. The Shared-Private model (Bousmalis et al., 2016) consists of a split representation where the private space learns specific features related to a particular domain while a shared space learns features common to all the domains. Such representation promotes non-contamination of the two spaces preserving their uniqueness. The contributions of our approach are as follows:
1) We show that using a shared-private model along with adversarial training improves SSST adaptation compared to neural network baselines. When multiple source corpora from similar domains are used it leads to further performance enhancement. Moreover, use of contextualized sentential embeddings leads to better performance than existing baseline methods for both MSST and SSST.

2) We explore the generalizability of our framework using two prominent neural architectures: CNN (Nguyen and Grishman, 2015) and Bi-LSTM (Kavuluru et al., 2017), where we find the former to be more robust across our experiments.

2 Methodology

For every labeled sources and a single unlabeled target we have set of NER tagged sentences, each of which is represented as: 
\[ X = \{ e_1, e_2, w_1, \ldots w_n \} \]
where \( e_1 \) and \( e_2 \) are two tagged entities and \( w_j \) is the \( j^{th} \) word in the sentence. A labelled source instance is accompanied by the relation label (True or False). In this section we discuss the input representation followed by model description.

2.1 Input Representation

We form word and position embeddings for every word in an NER tagged sentence. We use the PubMed-and-PMC-w2v\(^2\) to generate word embeddings. The size being \(|V| \cdot d_w\), where \( d_w \) is the word embedding dimension which is 200 and \(|V|\) is the vocabulary size. The position embedding vector for the \( j^{th} \) word in a sentence relative to two tagged entities \( e_1 \) and \( e_2 \) is represented as a tuple: 
\[ (p_{e1(j)}, p_{e2(j)}) \]
where, \( p_{e1(j)} \) and \( p_{e2(j)} \) ∈ \( \mathbb{R}^e \).

2.2 Model

Fig 1 shows the adaptation of MAN framework whose various components are discussed below.

**Shared & Domain feature extractor** (\( F_s, F_{di} \))

The input representation is fed to both \( F_{di} \) and \( F_s \) for labeled source domains whereas for unlabeled target instances it is fed only to \( F_s \). For SSST the model is trained on a single labeled source domain and tested on a unlabeled target domain. For MSST we do not combine the sources as a single corpus since that leads to a number of false negatives. We make two different assumptions to consider multiple sources: 1) Following Nguyen et al., (2014) we consider multiple labels from single corpus as different sources, 2) We use additional smaller corpora from a similar domain as multi-source. The **Shared Feature space** (\( F_s \)) learns domain agnostic representations and **Private Feature space** (\( F_{di} \)) learns domain specific features for every \( i^{th} \) labeled domain.

We apply two different architectures for both \( F_s \) and \( F_{di} \) to analyze the changes in performance of the approach for the task : Convolutional neural network (Nguyen and Grishman, 2015) (MAN CNN) and Bi-LSTM (Kavuluru et al., 2017) (MAN Bi-LSTM). We have performed detailed experiments on each of these in Section 5.

**Domain discriminator**\(^3\), \( D \) is a fully-connected layer with softmax that predicts multiple domain probabilities using Multinomial Adversarial Network. The output from \( F_s \) is fed to \( D \) which is adversarially trained separately from the entire network using L2 loss described as follows:

\[ L_D(\hat{d}, d) = \sum_{i=1}^{N} (\hat{d}_i - 1 \{d = i\})^2 \]

where, \( d \) is the index assigned for a domain and \( \hat{d} \) is the prediction. It is generalized as \( \sum_{i=1}^{N} \hat{d}_i = 1 \) and \( \forall i \neq 0, \hat{d}_i \geq 0 \). \( F_s \) tries to fool \( D \) so that it can not correctly guess the domain from where a sample instance is coming from. Thus \( F_s \) learns indistinguishable features in the process.

**Relation Classifier** \( C \) is a fully-connected layer with a softmax, used to predict the class probabilities. We use Bio-BERT (Lee et al., 2019) embeddings for every sentence as features (Geeticka Chauhan, 2019) BERT[CLS] that have shown to improve performance in many downstream tasks. This is concatenated with the fixed size sentence representation from \( F_s \) and \( F_{di} \), together

![Figure 1: MAN for Domain Adaptation of Binary Relation Classification. The figure shows the training flow given a sentence from a labeled source domain. \( D \) is trained separately than rest of the network.](http://evexdb.org/pmresources/vec-space-models/)

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\(^2\)http://evexdb.org/pmresources/vec-space-models/

\(^3\)Forward and backward passes of \( F_s \), \( F_{di} \), and \( C \) are followed by Domain Discriminator, \( D \).
Datasets Entity Pair # of Sent # of Positive # of Negative

AiMed 1995 1000 4834
BioInfer 1100 2534 7132
LLL PPI 77 164 166
HPRD50 145 163 270
IEPA 486 355 482

Table 1: Protein Protein Interaction Dataset statistics.

Datasets Entity Pair # of Train # of Valid # of Test

DDI Drug-Drug 27779 - 5713
CPR: 3 Chem-Prot 768 550 665
CPR: 4 2254 1094 1661
CPR: 5 173 116 195
CPR: 6 235 199 293
CPR: 9 727 457 644

Table 2: Drug-Drug Interaction and Chemical-Protein Dataset statistics.

they serve as input to C. For unlabeled target, during test no domain specific features are generated from $F_d_i$ and that part is set to zero vector. For binary classification we adopt Negative Log Likelihood Loss for C described below:

$$L_c(\hat{y}, y) = -\log P(\hat{y} = y)$$

where, $y$ is the true relation label and $\hat{y}$ is the softmax label. The objective of $F_{di}$ is same as that of C and it relies only on labeled data. On the other hand the objective of the Shared Feature Extractor $F_s$ is represented as follows:

$$Loss \ of \ F_s = Classifier \ loss + \lambda \ Domain \ loss$$

It consists of two loss components: improve performance of C and enhance learning of invariant features across all domains. A hyper parameter $\lambda$ is used to balance both of them.

3 Datasets

The dataset statistics is summarized in Table 1 and Table 2. A 10-fold cross validation was performed for the Protein-Protein Interaction dataset. For given set of entities E in a sentence, it is split into $\binom{E}{2}$ instances. All positive instances of datasets with more than two relation types are merged and assigned True labels while negative instances are assigned False labels. Unlabeled data is formed by removing labels from development and test datasets.

4 Experiments

Pre-processing: We anonymize the named entities in the sentence by replacing them with predefined tags like @PROT1$, @DRUG$ (Bhasuran and Natarajan, 2018).

4.1 Single source single target (SSST)

A thorough experiment is conducted using all possible combinations of the three benchmark data-sets AiMed (Bunescu et al., 2005), BioInfer (Pyysalo et al., 2006), DDI (Herrero-Zazo et al., 2013) whose results are discussed in Table 3

4.2 Multi-source single target (MSST)

The experiments with two different assumptions to consider multiple sources are as follows:

Multiple smaller corpora from similar domain: For Protein Protein Interaction there are three smaller standard corpora in literature, namely, LLL (Nedellec, 2005), IEPA (Ding et al., 2001), HPRD50 (Fundel et al., 2007). All three were considered as additional sources to transfer knowledge. AiMed (AM) and BioInfer (BI) were alternately selected as the unlabeled target in 2 different experiments while the remaining 4 denoted as 4P are considered as source corpus.

Multiple labels from single corpora: For ChemProt corpora we consider various labels as different sources following Nguyen et al., (2014) The five positive labels of ChemProt are: CPR: 3, CPR: 4, CPR: 5, CPR: 6, CPR: 9 which stand for upregulator, downregulator, agonist, antagonist and substrate, respectively. We predict the classification performance for unlabeled targets CPR:6 and CPR:9 taking multi-source labeled input denoted as 3C from three sources- CPR: 3, CPR: 4, CPR: 5 as positive instances and remaining as negative.

4.3 Baselines

We compare our approach with different baselines which are mentioned as follows:

- BioBERT (Rios et al., 2018): For SSST we train it on one dataset and test on another. For MSST we combine the multiple sources as a single source and test on labeled target.

- CNN+DANN (Lisheng Fu, 2017): A variant of adversarial training which is gradient reversal (RevGrad) is used with CNN (Nguyen and Grishman, 2015).
### Results and Discussions

In Table 3 we see that BioInfer generalizes well to AiMed and DDI corpora using vanilla LSTM or CNN architecture. However, with MAN and contextual embeddings, we do not see prominent gains as much as the other datasets. This can be due to the class imbalance in data (positive to negative instance ratio 1:5.9) (Hsu et al., 2015; Rios et al., 2018). For AiMed and BioInfer, we find that the knowledge transfer among themselves gives the best performance thus strengthening the fact that datasets from the same domain can contribute to performance enhancement justifying the performance gains in MSST experiments. Our model outperforms other baselines just with the use of adversarial training which might be attributed to joint learning better representation from shared and private feature extractors. The use of contextual BERT[CLS] tokens leads to increase in performance scores since they encode important relations between words in a sentence (Vig, 2019; Hewitt and Manning, 2019).

In Table 4, BioBERT is seen to perform well for ChemProt. We hypothesize that this may be due to the same underlying dataset being used during train and test. Though we use different labels as multi-source, that may not contribute to generating enough variance in sources since they were from the same dataset. For AiMed and BioInfer, however, three different smaller corpora were used, where the proposed method outperforms BioBERT. When compared across all the six SSST experiments, the Bi-LSTM based model lacks in performance may be due to absence of any attention mechanism which would have helped in selecting more relevant context (Chen and Cardie, 2018). We observe that adversarial training along with contextualized BERT sentence embeddings leads to performance gains across all datasets.
6 Conclusions
Our proposed model significantly outperformed the existing neural network based domain adaptation baselines for SSST. Among the two MSST experiments, we showed that the system gains when multiple source corpora are used. We also experimented with two architectures out of which CNN is seen to perform marginally better compared to Bi-LSTM. Our analysis on Section 5 further explains the effect of sources, adversarial training and use of contextualized BERT sentential embeddings.

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References


Assessing the Efficacy of Clinical Sentiment Analysis and Topic Extraction in Psychiatric Readmission Risk Prediction

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Abstract

Predicting which patients are more likely to be readmitted to a hospital within 30 days after discharge is a valuable piece of information in clinical decision-making. Building a successful readmission risk classifier based on the content of Electronic Health Records (EHRs) has proved, however, to be a challenging task. Previously explored features include mainly structured information, such as sociodemographic data, comorbidity codes and physiological variables. In this paper we assess incorporating additional clinically interpretable NLP-based features such as topic extraction and clinical sentiment analysis to predict early readmission risk in psychiatry patients.

1 Introduction and Related Work

Psychotic disorders affect approximately 2.5-4% of the population (Periä et al., 2007) (Bogren et al., 2009). They are one of the leading causes of disability worldwide (Vos et al., 2015) and are a frequent cause of inpatient readmission after discharge (Wiersma et al., 1998). Readmissions are disruptive for patients and families, and are a key driver of rising healthcare costs (Mangalore and Knapp, 2007) (Wu et al., 2005). Assessing readmission risk is therefore critically needed, as it can help inform the selection of treatment interventions and implement preventive measures.

Predicting hospital readmission risk is, however, a complex endeavour across all medical fields. Prior work in readmission risk prediction has used structured data (such as medical comorbidity, prior hospitalizations, sociodemographic factors, functional status, physiological variables, etc) extracted from patients’ charts (Kansagara et al., 2011). NLP-based prediction models that extract unstructured data from EHR have also been developed with some success in other medical fields (Murff et al., 2011). In Psychiatry, due to the unique characteristics of medical record content (highly varied and context-sensitive vocabulary, abundance of multiword expressions, etc), NLP-based approaches have seldom been applied (Vigod et al., 2015; Tulloch et al., 2016; Greenwald et al., 2017) and strategies to study readmission risk factors primarily rely on clinical observation and manual review (Olfson et al., 1999) (Lorine et al., 2015), which is effort-intensive, and does not scale well.

In this paper we aim to assess the suitability of using NLP-based features like clinical sentiment analysis and topic extraction to predict 30-day readmission risk in psychiatry patients. We begin by describing the EHR corpus that was created using in-house data to train and evaluate our models. We then present the NLP pipeline for feature extraction that was used to parse the EHRs in our corpus. Finally, we compare the performances of our model when using only structured clinical variables and when incorporating features derived from free-text narratives.

2 Data

The corpus consists of a collection of 2,346 clinical notes (admission notes, progress notes, and discharge summaries), which amounts to 2,372,323 tokens in total (an average of 1,011 tokens per note). All the notes were written in English and extracted from the EHRs of 183 psychosis patients from McLean Psychiatric Hospital in Belmont, MA, all of whom had in their history at least one instance of 30-day readmission.

The age of the patients ranged from 20 to 67 (mean = 26.65, standard deviation = 8.73). 51% of the patients were male. The number of admis-
sions per patient ranged from 2 to 21 (mean = 4, standard deviation = 2.85). Each admission contained on average 4.25 notes and 4,298 tokens. In total, the corpus contains 552 admissions, and 280 of those (50%) resulted in early readmissions.

3 Feature Extraction

The readmission risk prediction task was performed at the admission level. An admission consists of a collection of all the clinical notes for a given patient written by medical personnel between inpatient admission and discharge. Every admission was labeled as either ‘readmitted’ (i.e. the patient was readmitted within the next 30 days of discharge) or ‘not readmitted’. Therefore, the classification task consists of creating a single feature representation of all the clinical notes belonging to one admission, plus the past medical history and demographic information of the patient, and establishing whether that admission will be followed by a 30-day readmission or not.

45 clinically interpretable features per admission were extracted as inputs to the readmission risk classifier. These features can be grouped into three categories (See Table 1 for complete list of features):

- Sociodemographics: gender, age, marital status, etc.

- Past medical history: number of previous admissions, history of suicidality, average length of stay (up until that admission), etc.

- Information from the current admission: length of stay (LOS), suicidal risk, number and length of notes, time of discharge, evaluation scores, etc.

The Current Admission feature group has the most number of features, with 29 features included in this group alone. These features can be further stratified into two groups: ‘structured’ clinical features and ‘unstructured’ clinical features.

3.1 Structured Features

Structure features are features that were identified on the EHR using regular expression matching and include rating scores that have been reported in the psychiatric literature as correlated with increased readmission risk, such as Global Assessment of Functioning, Insight and Compliance:

Global Assessment of Functioning (GAF): The psychosocial functioning of the patient ranging from 100 (extremely high functioning) to 1 (severely impaired) (AAS, 2011).

Insight: The degree to which the patient recognizes and accepts his/her illness (either Good, Fair or Poor).

Compliance: The ability of the patient to comply with medication and to follow medical advice (either Yes, Partial, or None).

These features are widely-used in clinical practice and evaluate the general state and prognosis of...
the patient during the patient’s evaluation.

3.2 Unstructured Features

Unstructured features aim to capture the state of the patient in relation to seven risk factor domains (Appearance, Thought Process, Thought Content, Interpersonal, Substance Use, Occupation, and Mood) from the free-text narratives on the EHR. These seven domains have been identified as associated with readmission risk in prior work (Holderness et al., 2018).

These unstructured features include: 1) the relative number of sentences in the admission notes that involve each risk factor domain (out of total number of sentences within the admission) and 2) clinical sentiment scores for each of these risk factor domains, i.e. sentiment scores that evaluate the patients psychosocial functioning level (positive, negative, or neutral) with respect to each of these risk factor domain.

These sentiment scores were automatically obtained through the topic extraction and sentiment analysis pipeline introduced in our prior work (Holderness et al., 2019) and pretrained on in-house psychiatric EHR text. In our paper we also showed that this automatic pipeline achieves reasonably strong F-scores, with an overall performance of 0.828 F1 for the topic extraction component and 0.5 F1 on the clinical sentiment component.

The clinical sentiment scores are computed for every note in the admission. Figure 1 details the data analysis pipeline that is employed for the feature extraction.

First, a multilayer perceptron (MLP) classifier is trained on EHR sentences (8,000,000 sentences consisting of 340,000,000 tokens) that are extracted from the Research Patient Data Registry (RPDR), a centralized regional data repository of clinical data from all institutions in the Partners HealthCare network. These sentences are automatically identified and labeled for their respective risk factor domain(s) by using a lexicon of clinician identified domain-related keywords and multiword expressions, and thus require no manual annotation. The sentences are vectorized using the Universal Sentence Encoder (USE), a transformer attention network pretrained on a large volume of general-domain web data and optimized for greater-than-word length sequences.

The outputs of each clinical sentiment model are then averaged across notes to create a single value for each risk factor domain that corresponds to the patient’s level of functioning on a -1 to 1 scale (see Figure 2).
4 Experiments and Results

We tested six different classification models: Stochastic Gradient Descent, Logistic Regression, C-Support Vector, Decision Tree, Random Forest, and MLP. All of them were implemented and fine-tuned using the scikit-learn machine learning toolkit (Pedregosa et al., 2011). Because an accurate readmission risk prediction model is designed to be used to inform treatment decisions, it is important in adopting a model architecture that is clinically interpretable and allows for an analysis of the specific contribution of each feature in the input. As such, we include a Random Forest classifier, which we also found to have the best performance out of the six models.

To systematically evaluate the importance of the clinical sentiment values extracted from the free text in EHRs, we first build a baseline model using the structured features, which are similar to prior studies on readmission risk prediction (Kansagara et al., 2011). We then compare two models incorporating the unstructured features. In the "Baseline+Domain Sentences" model, we consider whether adding the counts of sentences per EHR that involve each of the seven risk factor domains as identified by our topic extraction model improved the model performance. In the "Baseline+Clinical Sentiment" model, we evaluate whether adding clinical sentiment scores for each risk factor domain improved the model performance. We also experimented with combining both sets of features and found no additional improvement.

Each model configuration was trained and evaluated 100 times and the features with the highest importance for each iteration were recorded. To further fine-tune our models, we also perform three-fold cross-validated recursive feature elimination 30 times on each of the three configurations and report the performances of the models with the best performing feature sets. These can be found in Table 2.

Our baseline results show that the model trained using only the structured features produce equivalent performances as reported by prior models for readmission risk prediction across all healthcare fields (Artetxe et al., 2018). The two models that were trained using unstructured features produced better results and both outperform the baseline results. The "Baseline+Clinical Sentiment" model produced the best results, resulting in an F1 of 0.72, an improvement of 14.3% over the baseline.

In order to establish what features were not relevant in the classification task, we performed recursive feature elimination. We identified 13 feature values as being not predictive of readmission (they were eliminated from at least two of the three feature sets without producing a drop in performance) including: all values for marital status (Single, Married, Other, and Unknown), missing values for GAF at admission, GAF score difference between admission & discharge, GAF at discharge, Veteran status, Race, and Insight & Mode of Past Insight values reflecting a clinically positive change (Good and Improving). Poor Insight values, however, are predictive of readmission.

5 Conclusions

We have introduced and assessed the efficacy of adding NLP-based features like topic extraction and clinical sentiment features to traditional structured-feature based classification models for early readmission prediction in psychiatry patients. The approach we have introduced is a hybrid machine learning approach that combines deep learning techniques with linear methods to ensure clinical interpretability of the prediction model.

Results show not only that both the number of sentences per risk domain and the clinical sentiment analysis scores outperform the structured-feature baseline and contribute significantly to better classification results, but also that the clinical sentiment features produce the highest results in all evaluation metrics (F1 = 0.72).

These results suggest that clinical sentiment features for each of seven risk domains extracted from free-text narratives further enhance early readmission prediction. In addition, combining state-of-art MLP methods has a potential utility in generating clinical meaningful features that can be be used in downstream linear models with interpretable and transparent results. In future work, we intend to increase the size of the EHR corpus, increase the demographic spread of patients, and extract new features based on clinical expertise to
increase our model performances. Additionally, we intend to continue our clinical sentiment annotation project from (Holderness et al., 2019) to increase the accuracy of that portion of our NLP pipeline.

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References


What does the language of foods say about us?

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Abstract

In this work we investigate the signal contained in the language of food on social media. We experiment with a dataset of 24 million food-related tweets, and make several observations. First, the language of food has predictive power. We are able to predict if states in the United States (US) are above the median rates for type 2 diabetes mellitus (T2DM), income, poverty, and education – outperforming previous work by 4–18%. Second, we investigate the effect of socioeconomic factors (income, poverty, and education) on predicting state-level T2DM rates. Socioeconomic factors do improve T2DM prediction, with the greatest improvement coming from poverty information (6%), but, importantly, the language of food adds distinct information that is not captured by socioeconomics. Third, we analyze how the language of food has changed over a five-year period (2013 – 2017), which is indicative of the shift in eating habits in the US during that period. We find several food trends, and that the language of food is used differently by different groups such as different genders. Last, we provide an online visualization tool for real-time queries and semantic analysis.

1 Introduction

With an average of 6,000 new tweets posted every second, Twitter\(^1\) has become a digital footprint of everyday life for a representative sample of the United States (US) population (Mislove et al., 2011). Previously, Fried et al. (2014) demonstrated that the language of food on Twitter can be used to predict health risks, political orientation, and geographic location. Here, we use predictive models to extend this analysis – exploring the ways in which the language of food can shed insight on health and the changing trends in both food culture and language use in different communities over time. We apply this methodology to the particular use case of predicting communities which are risk for type 2 diabetes mellitus (T2DM), a serious medical condition which affects over 30 million Americans and whose diagnosis alone costs $327 billion each year\(^2\). We refer to T2DM as diabetes in the rest of the paper. We show that by combining knowledge from tweets with other social characteristics (e.g., average income, level of education) we can better predict risk of T2DM. The contributions of this work are four-fold:

1. We use the same methods proposed by Fried et al. (2014) with a much larger (7 times) tweet corpus gathered from 2013 – 2017 to predict the risk of T2DM. We collected over 24 million tweets with meal-related hashtags (e.g., \#breakfast, \#lunch) and localized 5 million of them to states within the US. We show that more data helps, and that by training on this larger dataset the state-level T2DM risk prediction accuracy is improved by 4–18%, compared to the results in Fried et al. (2014). We also apply the same models to predict additional state-level indicators: income, poverty, and education levels in order to further investigate the predictive power of the language of food. On these prediction tasks, our model outperforms the majority baseline by 12–34%. We believe that this work may drive immediate policy decisions for the communities deemed at risk without awaiting for similar results from major health organizations, which take months or years to be generated and disseminated.\(^3\) Equally as important, we believe that this state-level T2DM risk prediction task may improve predicting risks

\(^1\)https://twitter.com/
\(^3\)https://www.cdc.gov/nchs/nhis/about_nhis.htm
for *individuals* from their social media activity, a task which often suffers from sparsity (Bell et al., 2018).

2. Unlike (Fried et al., 2014), we also investigate the effect of socioeconomic factors on the diabetes prediction task itself. We observe that aggregated US social demographic information from average income\(^4\), poverty\(^5\), and education\(^6\) is complementary to the information gained from tweet language used for predicting diabetes risk. We add the correlation between each of these socioeconomic factors and the diabetes\(^7\) rate in US states as additional features in the models in (1). We demonstrate that the T2DM prediction model strongly benefits from the additional information, as prediction accuracy further increases by 2–6%. However, importantly, the model that relies solely on these indicators performs considerably worse than the model that includes features from the language of food, which demonstrates that the language of food provides distinct signal from these indicators.

3. Furthermore, with a dataset that spans nearly five years, we also analyze language trends over time. Specifically, using pointwise mutual information (PMI) and a custom-built collection of healthy/unhealthy food words, we investigate the strength of healthy/unhealthy food references on Twitter, and observe a downward trend for healthy food references and an upward trend for unhealthy food words in the US.

4. Lastly, we provide a visualization tool to help understand and visualize semantic relations between words and categories such as how different genders refer to vegetarian vs. low-carb diets.\(^8\) Our tool is based on semantic axes plots (Heimerl and Gleicher, 2018).

2 Related Work

Many previous efforts have shown that social media can serve as a source of data to detect possible health risks. For example, Akbari et al. (2016) proposed a supervised learning approach that automatically extracts public wellness events from microblogs. The proposed method addresses several problems associated with social media such as insufficient data, nosiness and variance, and inter-relations among social events. A second contribution of Akbari et al. (2016) is an automatically-constructed large-scale diabetes dataset that is extended with manually handcrafted ground-truth labels (positive, negative) for wellness events such as diet, exercise and health.

Bell et al. (2016) proposed a strategy that uses a game-like quiz with data and questions acquired semi-automatically from Twitter to acquire relevant training data necessary to detect individual T2DM risk. In following work, Bell et al. (2018) predicted individual T2DM risk using a neural approach, which incorporates tweet texts with gender information and information about the recency of posts.

Sadeque et al. (2018) discussed several approaches for predicting depression status from a user’s social media posts. They proposed a new latency-based F1 metric to measure the quality and speed of the model. Further, they re-implemented some of the common approaches for this task, and analyzed their results using their proposed metric. Lastly, they introduced a window-based technique that trades off between latency and precision in predicting depression status.

Our work is closest to (Fried et al., 2014). Similar to us, Fried et al. (2014) predicted latent population characteristics from Twitter data such as overweight rate or T2DM risk in US states. Our work extends (Fried et al., 2014) in several ways. First, in addition of tweets, we incorporate state-level indicators such as poverty, education, and income in our risk classifier, and demonstrate that language provides distinct signal from these indicators. Second, we use the much larger tweet dataset to infer language-of-food trends over a five-year span. Third, we provide a visualization tool to explore food trends over time, as well as semantic relations between words and categories in this context.

3 Data

We collected tweets along with their meta data with Twitter’s public streaming API\(^9\). Tweets have been filtered by a set of seven hashtags to make the dataset more relevant to food (see distribution in Table 1). We stored the tweets and their metadata...
### Table 1: Seven meal-related hashtags and their corresponding number of tweets filtered from Twitter. The right-most column indicates the number of tweets we could localize to a US state or Washington D.C.

<table>
<thead>
<tr>
<th>Term</th>
<th># of tweets</th>
<th># of tweets localized in US</th>
</tr>
</thead>
<tbody>
<tr>
<td>#dinner</td>
<td>5,455,890</td>
<td>1,367,745</td>
</tr>
<tr>
<td>#breakfast</td>
<td>5,125,014</td>
<td>1,183,462</td>
</tr>
<tr>
<td>#lunch</td>
<td>4,969,679</td>
<td>1,094,681</td>
</tr>
<tr>
<td>#brunch</td>
<td>1,910,950</td>
<td>681,978</td>
</tr>
<tr>
<td>#snack</td>
<td>797,676</td>
<td>220,697</td>
</tr>
<tr>
<td>#meal</td>
<td>495,073</td>
<td>101,976</td>
</tr>
<tr>
<td>#supper</td>
<td>124,979</td>
<td>22,154</td>
</tr>
<tr>
<td>Total</td>
<td>24,493,223</td>
<td>4,362,940</td>
</tr>
</tbody>
</table>

4. State-level prediction tasks

We investigate the predictive power of the language of food through four distinct prediction tasks: T2DM rate, income, poverty, and education level. We use the tweets from the above dataset as the only input for our prediction models.

#### T2DM rate prediction

We use the diabetes rate from the Kaiser Commission on Medicaid and Uninsured (KCMU)'s analysis of the Center for Disease Control’s Behavioral Risk Factor Surveillance System (BRFSS) 2017 Survey (its most recent year). The state-level diabetes rate is defined as the percentage of adults in each state who have been told by a doctor that they have diabetes. The median diabetes rate for the US is 10.8%. For each state, we convert the diabetes rate into a binary variable with a value of 1 if the state diabetes rate is greater than or equal to the national median rate, and a value of 0 if it is below. For example, the state with highest diabetes rate, West Virginia (15.2%), is assigned a binary variable of 1 (high T2DM rates). On the other hand, states with below-national-median rate, like Arizona (10.4%), are assigned a label of 0 (low T2DM rates).

#### Income rate prediction

We collect income data from the United States Census Bureau (USCB)'s analysis of the American Community Survey (ACS)'s Income and Poverty in the United States: 2017. The data shows that national median household income is $60,336. Similarly to above, we convert the household median income of the state into a binary variable with a value of 0 (low income) if its median household income is lower than national median, and a value of 1 (high income) if its median household income is equal or greater. For example, Alabama ($48,193) is labeled as low-income and Alaska ($74,058) is labeled as high-income.

#### Poverty rate prediction

To predict poverty rates, we also collect poverty data from the USCB’s analysis of the ACS’s Income and Poverty in the United States: 2017, which shows that national median poverty rate is 13.4%. Again, we assign each state a binary variable indicating whether its rate is above or below this national median.

#### Education rate prediction

For predicting education rate, we use the higher education attain-
ment rate (HEAR) data from the Center of American Progress (CAP)\(^6\). The data shows that national median HEAR is 43.2%. Once again, the state-level HEAR is converted to a binary variable in the same manner as above.

Because each of these binary variables is at the state level, we group the tweets by state before feature extraction. We use leave-one-out cross-validation (LOOCV) as proposed by Fried et al. (2014). This approach is necessary because even though we have a large tweet corpus, we only have 51 aggregate data points (one for each state plus Washington, D.C.). For classification, we use Support Vector Machines (SVM) (Vapnik, 2013) for feature-based classification. To avoid overfitting, we tuned the classifier’s hyper-parameters during training using the tweets from 2013 to 2016. We tested the tuned prediction models for each task using solely tweets from 2017.

We use two sets of features: lexical (words from tweets) and topical (sets of words appearing in similar contexts). For lexical features, we compare open (all unique tweet words or hashtags) and closed (800 food words) vocabularies, using the token counts as the tweet features. These experiments help us to determine the predictive power of the specific language of food versus the broader context in the full tweets (or socially compact hashtag). For topic model features, we use Latent Dirichlet Allocation (LDA) (Blei et al., 2003), to learn a set of topics from food tweets. Because tweets are very short in nature (up to 140 characters), this approach allows us to analyze correlations that could go beyond individual words. We chose 200 as the number of topics for LDA to learn. After LDA is trained using MALLET\(^13\), we use it to create the set of topics for each tweet, and the topic with highest probability is then assigned to each tweet as an additional feature. Topics are counted across all tweets in a state in the same manner as the lexical features.

We also experimented with Deep Averaging Network (DAN) (Iyyer et al., 2015), a simple but robust bag-of-words model based on averaging word embeddings that has been shown to perform well in sentiment analysis and factoid question answering with little training data. In our case, we implemented DAN with embeddings generated using Word2Vec (Mikolov et al., 2013) trained over all 24 million tweets (including the ones that were not localized). We compute the embedding for each token in our dataset, and pass them to the network; see Figure 1. Again using LOOCV, in each pass we leave out one state, train the network on tweets from the 50 other states and predict the T2DM rate for the left out state.

4.2 Impact of socioeconomic factors

Previous work has shown that the T2DM rate can be predicted by socioeconomic factors such as poverty (Chih-Cheng et al., 2012), income (Yelenas et al., 2015), and education (Ayyagari et al., 2011). Therefore, we incorporate these factors into our prediction models (Section 4.1) to assess their contribution. We represent each socioeconomic factor and its correlation with the T2DM rate in the corresponding state as a feature, and include these new features alongside the lexical and topic-based ones. Even though in general, the correlations are relatively low (see Table 2), we will show that the model strongly benefits from the additional information leading to accuracy increases of 2–6% (see Section 5). This indicates that the language of food captures different signal and reflects distinct information from these indicators. However, because these indicators are represented as single features, as opposed to the other features (e.g., there are tens of thousands of food word features, each of which is represented as an integer count), they tended to be ignored by the classifier. To account for this, we empirically explored a series of multipliers to increase the weights of the values of these indicator features\(^14\). For this task, we use the same SVM classifier from Section 4.1, as well as a Random Forest (RF) classifier (Breiman, 2001)\(^15\).

\(^{13}\)http://mallet.cs.umass.edu/

\(^{14}\)For these correlation multipliers, we experimented with powers of 10, from \(10^1\) to \(10^6\).

\(^{15}\)To avoid overfitting, we do not fine-tune the RF classifier’s hyperparameters.
Table 2: Correlation between socioeconomic factors (education, income, poverty) and type 2 diabetes mellitus (T2DM) in 2017. Each correlation is calculated from the binary data described in Section 4.1. For the correlation values we used Pearson correlation (Boslaugh, 2012).

<table>
<thead>
<tr>
<th>Socioeconomic factor</th>
<th>Correlation with T2DM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Education</td>
<td>-0.37</td>
</tr>
<tr>
<td>Income</td>
<td>-0.14</td>
</tr>
<tr>
<td>Poverty</td>
<td>0.18</td>
</tr>
</tbody>
</table>

4.3 Exploring food trends

We use pointwise mutual information (PMI) between food words/hashtags and years to analyze food trends over time. We divide our corpus of tweets into four parts, each containing a complete year’s set of tweets (from 2014 to 2017) and then calculate PMI for pairs (food term $t$, year $y$) using the formula:

$$PMI(t, y) = \frac{C(t, y)}{C(t) \ast C(y)}, \quad (1)$$

where, $C(t, y)$ is the number of occurrences of term $t$ in year $y$, $C(t)$ is the total number of occurrences of the term, and $C(y)$ is the number of tweets in year $y$. Intuitively, the higher the PMI value of a term in a given year, $PMI(t, y)$, the more that term is associated with tweets from that year in particular.

4.4 Semantic axes analysis

Word vector embeddings are a standard tool used to analyze text, as they capture similarity relationships between the different words. However, interpreting such embeddings and understanding the encoded grammatical and semantic relations between words can be difficult due to the high dimensionality of the embedding space (typically 50-300 dimensions).

Semantic axes visualizations allow us to view specific low dimensional projections where the new dimensions can be used to explore different semantic concepts (Heimerl and Gleicher, 2018). For our task, we generate several word embeddings from our dataset using the CBOW Word2Vec model of (Mikolov et al., 2013). Different than other visualization tools (e.g., t-SNE, PCA), when using semantic axes we need to define two semantic axes by two opposite concepts (e.g., man vs. woman and breakfast vs. dinner) and project a collection of vectors (words in embedding) based on the specific 2D space. The result is a 2D scatter plot with respect to two different concepts.

We first create a word embedding for all the tweets in our dataset. This allows us to explore the correlations between different concepts.

We further augment the semantic axes tool\(^{16}\) provided by Heimerl and Gleicher (2018), to allow a concept axis to be defined by two sets of words (rather than exactly two words). For example, instead of having one axis defined by the pair (vegetables, meat) we can now use two sets of words (vegetables, fruit, vegetarian, vegan, etc., and meat, fish, chicken, beef, etc.). This allows us to capture more complex concepts such as “meat-eaters” that are not captured by individual words.

5 Results

We present the results for all prediction tasks of state level characteristics, as well as the evaluation of the contribution of socioeconomic factors alongside food language in predicting T2DM rate. We also investigate the shifts in eating habits over time (i.e., food trends), as well as the trends in different groups through our semantic axes experiments.

5.1 State-level characteristics prediction

In Table 3, we show the results for predicting state-level socioeconomic characteristics using various sets of features. We compare the results from our dataset with the results of Fried et al. (2014) for predicting T2DM rates. However, since Fried et al. (2014) do not experiment with predicting poverty, income, and education level, for these we compare against a majority baseline. As there are 51 states (including Washington D.C.), and each binary socioeconomic factor is based on the national median, this means that for each factor there will be 26 states either above or below (resulting in a majority baseline of 50.98%).

Comparing the effects of each type of lexical features and their combination with LDA topic features on these prediction tasks, we make several observations.

**Performance comparison by feature set:** First and foremost, the results demonstrate that the language of food can be used to predict health and social characteristics such as diabetes risk, income, poverty, and education level. The highest overall performance is achieved by using all tweet words (both with and without LDA). This suggests that

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\(^{16}\) http://embvis.flovis.net/
Table 3: Results from using various feature sets to predict state-level characteristics: whether a given state is above or below the national median for diabetes, poverty, income, and education. We also show the average performance across all characteristics. We compare against Fried et al. (2014) as well as the majority baseline. Note that Fried et al. do not predict poverty, income, or education level. The low number of data points (51 states) is responsible for the same accuracy value in multiple experiments.

we can capture significant predictive signal from tweets when capturing food words in context.

The highest prediction performance is seen when predicting the state-level income rate, demonstrating a high correlation between food-related words and income. When predicting state-level diabetes rate, we also see strong predictive power from the language of food – all models perform above 70%, up to 78.43%. This confirms our hypothesis that there is a strong correlation between food-related words (and presumably food behaviors) and diabetes rate, one indicator of public health.

Amount and recency of data: For diabetes prediction, with our larger dataset, we improve upon the results of Fried et al. (2014) (ranging from 4 to 18%). In particular, when we use the food-word features combined with LDA topics, we increase prediction accuracy by almost 18%. These results suggest that more data matters in this type of analysis, as evidenced by the learning curves shown in Figure 2, where we compare performance against amount of training data (by year).

We also created learning curves for prediction of T2DM, but from the opposite direction, i.e., starting from tweets from 2017 only, and then adding tweets from earlier years one year at a time. We observe that the more recent the data, the more useful it is for prediction. We hypothesize that in terms of the utility of increased data, the performance of food-word features is improved only as the amount of relevant data increases. For the first part of the curve (only from 17, combined 17–16, combined 17–15), the classifier’s performance is improved with additional tweets. However, after this peak, additional older tweets decrease performance, suggesting that people change their eating behavior over a period spanning multiple years.

The importance of recency of tweet data is also discussed in (Bell et al., 2016).

Comparison to previous work: The best performing model of Fried et al. (2014) relies on hash-tags (see Table 3, lines 5 and 7) and the worst performing model use food words (lines 9 and 11). However, with more data we find that we get the best performance with food words (line 12). We hypothesize that with smaller data, the concise semantics of hashtags are more informative, but with more data the model is able to learn the relative semantics of the food words themselves. Further, while LDA topics do not benefit any model of Fried et al. in terms of predicting diabetes, here we find that with additional data, LDA topics benefit the food words model (compare lines 10 and 12), perhaps because additional data leads to more representative LDA topics.
The Deep Averaging Network from Figure 1 consistently underperformed the results reported in Table 3. This approach obtained an accuracy of 60.78% for the T2DM rate prediction task, considerably lower than the 78% obtained by the best SVM configuration in Table 3. We hypothesize that the reason behind this low performance is the small number of data points (51), which is insufficient to train a neural network.

5.2 Impact of socioeconomic factors

In Table 5, we show the SVM results for predicting T2DM rate from extending the feature matrix from 5.1 with one additional feature based on the correlation between each socioeconomic factor (education, income, and poverty) and T2DM. For each factor, we compare several multipliers (see Section 4.2) to amplify the impact of the socioeconomic correlations. Consistently, we find that models with more features benefit from larger multipliers. For example, the extended food word features that have several hundred features perform best with a multiplier of $10^2$, while the other extended models, which all have tens of thousands of features, perform best with a multiplier of $10^3$.

The best multiplier for each model, according to SVM performance, is used in our Random Forest models (Table 4).

From these extended models we see that using poverty information as an additional feature improves our SVM performance by a range of 2–8% and our RF performance by up to 6%. The other socioeconomic factors, i.e., income and education, do not help when using an SVM classifier (Table 5), but when using a RF classifier we see up to 6% improvement (Table 4). Overall, our highest T2DM prediction performance is obtained with SVM using Food + LDA + poverty. This performance surpasses 80% accuracy and is the highest value reported for this task. Further, to the best of our knowledge, the effect of using poverty information to improve T2DM rate prediction is novel and suggests a potential avenue for improving classifiers with socioeconomic correlation information.

Importantly, predicting the T2DM below/above median labels from the poverty indicator alone has an accuracy of 58.82%. This value is considerably lower than that of the classifier that uses poverty coupled with the extended word features from tweets, which obtained 80% accuracy. This demonstrates that the language of food provides signal that is distinct from this indicator, which suggests that there is value in social media mining for the monitoring of health risks.

5.3 Food trends

Given our dataset that spans nearly five years, we are also able to investigate whether changes in food habits over time can be detected in social media language. To this end, we explored a list of 800 food words and their change in PMI values in the different years. To understand which food words...

Table 4: Results for predicting T2DM rate using a random forest classifier with our additional socioeconomic correlation features. For each feature set, we use the best performing multiplier, as determined in the previous experiment that used a SVM classifier (Table 5). That is, the best performing multiplier for food word features is $10^2$, while other features’ multipliers are $10^3$. # Features Results from best performing multiplier

| All Words+LDA with RF   | 62.75 | Fried et al. (2014) 64.71 |
| Food+LDA with RF       | 70.59 | Fried et al. (2014) 60.78 |
| Food+Hashtags+LDA with RF | 66.67 | Fried et al. (2014) 62.75 |
| Hashtags+LDA with RF   | 68.63 | Fried et al. (2014) 64.71 |
| Food+Hashtags+LDA with RF | 66.67 | Fried et al. (2014) 62.75 |
| Food+Hashtags+LDA with RF | 66.67 | Fried et al. (2014) 62.75 |
| Food+Hashtags+LDA with RF | 66.67 | Fried et al. (2014) 62.75 |
| Food+Hashtags+LDA with RF | 66.67 | Fried et al. (2014) 62.75 |

Figure 3: The learning curve using food word features, on the diabetes rate prediction task for 2017. In this figure, the data portion used for each point is in reverse order compared to Figure 2, that is, starting from most recent tweets and going back in time. The horizontal axis is labeled based on the year(s) from which the tweets used for prediction were used.
indicate healthy vs. unhealthy diets, we manually classified the 800 food words into three categories – healthy, unhealthy and neutral – using reliable online resources\(^{17}\). The annotations were independently performed by three annotators. The inter-annotator Kappa agreement scores\(^{18}\) shown in Table 6 indicate fair to good agreement between the three annotators.

We computed PMI values for each of these 800 words and each year in our US dataset. We also computed the PMI values for the three categories and each year (here all words from each category are treated as one). The category trends in our US dataset indicate a slight increase of mentions of unhealthy food words and a slight decrease in mentions of healthy food words in US tweets; see Figure 4. These results suggest a continued decline in dietary patterns in the US, despite seemingly increased interest in health benefits from food\(^{19}\).

### 5.4 Semantic axes visualization

As discussed in Section 4.4, visualizations can help discover correlations between different concepts, as well as look at trends over time. In Figure 5, we consider the two axes defined by man vs. woman, and breakfast vs. dinner. Exploring the four corners we can identify particular types of foods representative of those coordinates. In the top-left we see words associated with women and breakfast (yogurt, cupcake, pastry), whereas in the bottom-left we see words associated with men and breakfast (sausage, bacon, ham). Similarly, in the top-right we see words associated with women and dinner (mussels, halibut, eggplant) whereas the bottom right we see words associated with men and dinner (lasagna, lamb, teriyaki). This data confirms common stereotypes, e.g., (1) men tend to eat more meat, whereas women often prefer fish, and (2) women are more health-conscious compared to men.

We also consider topics (defined by a collection of words) as axes, as illustrated in Table 7. The two axes now are man vs. woman, and vegetarian vs. low-carb diets. To represent the vegetarian topic we use the words vegan, vegetarian, tofu, and to represent the low-carb topic we use keto, paleo, and atkins. We then average the word embedding vectors for all words in the topic to create the 2D projection.

We list the 4 corners in the projection as 4 rows in Table 7, where the left column corresponds to the concepts and the right column contains the words. Several patterns emerge: vegetarian words associated with women tend to be soups, salads,
and gourmet types of foods (saffron, fennel). In contrast, the vegetarian words associated with men tend to be vegetables (spinach, kale, carrot). In the low-carb and women corner we find breakfast words and deserts (cupcake, pastry, caramel, wheat) whereas in the low-carb and men corner we see more hearty foods (spaghetti, hamburgers, buns).

<table>
<thead>
<tr>
<th>Concepts</th>
<th>Foods</th>
</tr>
</thead>
<tbody>
<tr>
<td>woman, vegetarian diet</td>
<td>mint, saffron, fennel, squash, soup, tomato, eggplant</td>
</tr>
<tr>
<td>man, vegetarian diet</td>
<td>beet, onion, coconut, spinach, kale, carrot</td>
</tr>
<tr>
<td>woman, low-carb diet</td>
<td>hazelnut, nut, cupcake, pastry, grain, caramel, wheat</td>
</tr>
<tr>
<td>man, low-carb diet</td>
<td>cereal, spaghetti, buns, hamburger, pepperoni, crunch</td>
</tr>
</tbody>
</table>

Table 7: The four corners in the man vs. woman and vegetarian words vs. low-carb diets plot. Each row represents one corner. The left column contains the pair of concepts; the right column contains the foods associated with those concepts.

### 6 Conclusion

We showed that the language of food has predictive power for non-trivial state-level health tasks such as predicting if a state has higher/lower diabetes risk than the median. When augmented with socio-economic data such as poverty indicators, performance improves further, but we demonstrate that the language of food captures different signal and reflect distinct information from these socio-economic data. We also provide visualization tools to analyze the underlying data and visualize patterns and trends. This work may have immediate use in public health, e.g., by driving rapid policy decisions for the communities deemed at health risk. Further, we hope that this work complements predicting health risk for individuals, a task that is plagued by sparsity, and which could potentially benefit from additional community-level information.

### Acknowledgments

Mihai Surdeanu declares a financial interest in lum.ai. This interest has been properly disclosed to the University of Arizona Institutional Review Committee and is managed in accordance with its conflict of interest policies.

### References


Dreaddit: A Reddit Dataset for Stress Analysis in Social Media

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Abstract

Stress is a nigh-universal human experience, particularly in the online world. While stress can be a motivator, too much stress is associated with many negative health outcomes, making its identification useful across a range of domains. However, existing computational research typically only studies stress in domains such as speech, or in short genres such as Twitter. We present Dreaddit, a new text corpus of lengthy multi-domain social media data for the identification of stress. Our dataset consists of 190K posts from five different categories of Reddit communities; we additionally label 3.5K total segments taken from 3K posts using Amazon Mechanical Turk. We present preliminary supervised learning methods for identifying stress, both neural and traditional, and analyze the complexity and diversity of the data and characteristics of each category.

1 Introduction

In our online world, social media users tweet, post, and message an incredible number of times each day, and the interconnected, information-heavy nature of our lives makes stress more prominent and easily observable than ever before. With many platforms such as Twitter, Reddit, and Facebook, the scientific community has access to a massive amount of data to study the daily worries and stresses of people across the world.1

Stress is a nearly universal phenomenon, and we have some evidence of its prevalence and recent increase. For example, the American Psychological Association (APA) has performed annual studies assessing stress in the United States since 20072 which demonstrate widespread experiences of chronic stress. Stress is a subjective experience whose effects and even definition can vary from person to person; as a baseline, the APA defines stress as a reaction to extant and future demands and pressures,3 which can be positive in moderation. Health and psychology researchers have extensively studied the connection between too much stress and physical and mental health (Lupien et al., 2009; Calcia et al., 2016).

In this work, we present a corpus of social media text for detecting the presence of stress. We hope this corpus will facilitate the development of models for this problem, which has diverse applications in areas such as diagnosing physical and mental illness, gauging public mood and worries in politics and economics, and tracking the effects of disasters. Our contributions are as follows:

• Dreaddit, a dataset of lengthy social media posts in five categories, each including stressful and non-stressful text and different ways of expressing stress, with a subset of the data annotated by human annotators;4
• Supervised models, both discrete and neural, for predicting stress, providing benchmarks to stimulate further work in the area; and
• Analysis of the content of our dataset and the performance of our models, which provides insight into the problem of stress detection.

In the remainder of this paper, we will review relevant work, describe our dataset and its annotation, provide some analysis of the data and stress detection problem, present and discuss results of some supervised models on our dataset, and finally conclude with our summary and future work.

2 Related Work

Because of the subjective nature of stress, relevant research tends to focus on physical sig-

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3https://www.apa.org/helpcenter/stress-kinds
4Our dataset will be made available at http://www.cs.columbia.edu/~eturcan/data/dreaddit.zip.
nals, such as cortisol levels in saliva (Allen et al., 2014), electroencephalogram (EEG) readings (Al-Shargie et al., 2016), or speech data (Zuo et al., 2012). This work captures important aspects of the human reaction to stress, but has the disadvantage that hardware or physical presence is required. However, because of the aforementioned proliferation of stress on social media, we believe that stress can be observed and studied purely from text.

Other threads of research have also made this observation and generally use microblog data (e.g., Twitter). The most similar work to ours includes Winata et al. (2018), who use Long Short-Term Memory Networks (LSTMs) to detect stress in speech and Twitter data; Guntuku et al. (2018), who examine the Facebook and Twitter posts of users who score highly on a diagnostic stress questionnaire; and Lin et al. (2017), who detect stress on microblogging websites using a Convolutional Neural Network (CNN) and factor graph model with a suite of discrete features. Our work is unique in that it uses data from Reddit, which is both typically longer and not typically as conducive to distant labeling as microblogs (which are labeled in the above work with hashtags or pattern matching, such as “I feel stressed”). The length of our posts will ultimately enable research into the causes of stress and will allow us to identify more implicit indicators. We also limit ourselves to text data and metadata (e.g., posting time, number of replies), whereas Winata et al. (2018) also train on speech data and Lin et al. (2017) include information from photos, neither of which is always available. Finally, we label individual parts of longer posts for acute stress using human annotators, while Guntuku et al. (2018) label users themselves for chronic stress with the users’ voluntary answers to a psychological questionnaire.

Researchers have used Reddit data to examine a variety of mental health conditions such as depression (Choudhury et al., 2013) and other clinical diagnoses such as general anxiety (Cohan et al., 2018), but to our knowledge, our corpus is the first to focus on stress as a general experience, not only a clinical concept.

3 Dataset
3.1 Reddit Data
Reddit is a social media website where users post in topic-specific communities called subreddits, and other users comment and vote on these posts. The lengthy nature of these posts makes Reddit an ideal source of data for studying the nuances of phenomena like stress. To collect expressions of stress, we select categories of subreddits where members are likely to discuss stressful topics:

- **Interpersonal conflict:** abuse and social domains. Posters in the abuse subreddits are largely survivors of an abusive relationship or situation sharing stories and support, while posters in the social subreddit post about any difficulty in a relationship (often but not exclusively romantic) and seek advice for how to handle the situation.

- **Mental illness:** anxiety and Post-Traumatic Stress Disorder (PTSD) domains. Posters in these subreddits seek advice about coping with mental illness and its symptoms, share support and successes, seek diagnoses, and so on.

- **Financial need:** financial domain. Posters in the financial subreddits generally seek financial or material help from other posters.

We include ten subreddits in the five domains of abuse, social, anxiety, PTSD, and financial, as detailed in Table 1, and our analysis focuses on the domain level. Using the PRAW API,\(^5\) we scrape all available posts on these subreddits between January 1, 2017 and November 19, 2018; in total, 187,444 posts. As we will describe in subsection 3.2, we assign binary stress labels to 3,553 segments of these posts to form a supervised and semi-supervised training set. An example segment is shown in Figure 1. Highlighted phrases are in-

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\(^5\)https://github.com/praw-dev/praw

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I have this feeling of dread about school right before I go to bed and I wake up with an upset stomach which lasts all day and makes me feel like I’ll throw up. This causes me to lose appetite and not wanting to drink water for fear of throwing up. I’m not sure where else to go with this, but I need help. If any of you have this, can you tell me how you deal with it? I’m tired of having this every day and feeling like I’ll throw up.

Figure 1: An example of stress being expressed in social media from our dataset, from a post in r/anxiety (reproduced exactly as found). Some possible expressions of stress are highlighted.
Table 1: Data Statistics. We include ten total subreddits from five domains in our dataset. Because some subreddits are more or less popular, the amount of data in each domain varies. We endeavor to label a comparable amount of data from each domain for training and testing.

<table>
<thead>
<tr>
<th>Domain</th>
<th>Subreddit Name</th>
<th>Total Posts</th>
<th>Avg Tokens/Post</th>
<th>Labeled Segments</th>
</tr>
</thead>
<tbody>
<tr>
<td>abuse</td>
<td>r/domesticviolence</td>
<td>1,529</td>
<td>365</td>
<td>388</td>
</tr>
<tr>
<td></td>
<td>r/survivorsofabuse</td>
<td>1,372</td>
<td>444</td>
<td>315</td>
</tr>
<tr>
<td></td>
<td><strong>Total</strong></td>
<td><strong>2,901</strong></td>
<td><strong>402</strong></td>
<td><strong>703</strong></td>
</tr>
<tr>
<td>anxiety</td>
<td>r/anxiety</td>
<td>58,130</td>
<td>193</td>
<td>650</td>
</tr>
<tr>
<td></td>
<td>r/stress</td>
<td>1,078</td>
<td>107</td>
<td>78</td>
</tr>
<tr>
<td></td>
<td><strong>Total</strong></td>
<td><strong>59,208</strong></td>
<td><strong>191</strong></td>
<td><strong>728</strong></td>
</tr>
<tr>
<td>financial</td>
<td>r/almosthomeless</td>
<td>547</td>
<td>261</td>
<td>99</td>
</tr>
<tr>
<td></td>
<td>r/assistance</td>
<td>9,243</td>
<td>209</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>r/food_pantry</td>
<td>343</td>
<td>187</td>
<td>43</td>
</tr>
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<td></td>
<td>r/homeless</td>
<td>2,384</td>
<td>143</td>
<td>220</td>
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<tr>
<td></td>
<td><strong>Total</strong></td>
<td><strong>12,517</strong></td>
<td><strong>198</strong></td>
<td><strong>717</strong></td>
</tr>
<tr>
<td>PTSD</td>
<td>r/ptsd</td>
<td>4,910</td>
<td>265</td>
<td>711</td>
</tr>
<tr>
<td>social</td>
<td>r/relationships</td>
<td>107,908</td>
<td>578</td>
<td>694</td>
</tr>
<tr>
<td></td>
<td><strong>Total</strong></td>
<td><strong>187,444</strong></td>
<td><strong>420</strong></td>
<td><strong>3,553</strong></td>
</tr>
</tbody>
</table>

dicators that the writer is stressed: the writer mentions common physical symptoms (nausea), explicitly names fear and dread, and uses language indicating helplessness and help-seeking behavior.

The average length of a post in our dataset is 420 tokens, much longer than most microblog data (e.g., Twitter’s character limit as of this writing is 280 characters). While we label segments that are about 100 tokens long, we still have much additional data from the author on which to draw. We feel this is important because, while our goal in this paper is to predict stress, having longer posts will ultimately allow more detailed study of the causes and effects of stress.

In Table 2, we provide examples of labeled segments from the various domains in our dataset. The samples are fairly typical; the dataset contains mostly first-person narrative accounts of personal experiences and requests for assistance or advice. Our data displays a range of topics, language, and agreement levels among annotators, and we provide only a few examples. Lengthier examples are available in the appendix.

### 3.2 Data Annotation

We annotate a subset of the data using Amazon Mechanical Turk in order to begin exploring the characteristics of stress. We partition the posts into contiguous five-sentence chunks for labeling; we wish to annotate segments of the posts because we are ultimately interested in what parts of the post depict stress, but we find through manual inspection that some amount of context is important. Our posts, however, are quite long, and it would be difficult for annotators to read and annotate entire posts. This type of data will allow us in the future not only to classify the presence of stress, but also to locate its expressions in the text, even if they are diffused throughout the post.

We set up an annotation task in which English-speaking Mechanical Turk Workers are asked to label five randomly selected text segments (of five sentences each) after taking a qualification test; Workers are allowed to select “Stress”, “Not Stress”, or “Can’t Tell” for each segment. In our instructions, we define stress as follows: “The Oxford English Dictionary defines stress as ‘a state of mental or emotional strain or tension resulting from adverse or demanding circumstances’. This means that stress results from someone being uncertain that they can handle some threatening situation. We are interested in cases where that someone also feels negatively about it (sometimes we can find an event stressful, but also find it exciting and positive, like a first date or an interview).”.

We specifically ask Workers to decide whether the author is expressing both stress and a negative attitude about it, not whether the situation itself seems stressful. Our full instructions are available in the appendix.

We submit 4,000 segments, sampled equally from each domain and uniformly within domains,
I only get it when I have a flashback or strong reaction to a trigger. I notice it sticks around even when I feel emotionally calm and can stick around for a long time after the trigger, like days or weeks. Its a new symptom I think. Also been having lots of nightmares again recently. Not sure what to do as Im not currently in therapy, but I am waiting to be seen at a mental health clinic.

Regardless, that didn't last long, maybe half a year. I released that apartment, and most of my belongings (I kept a few boxes of my things from the military, personal effects, but little else). Looking back, there were some signs of emotional manipulation here, but it was subtle... and you know how it is, love is blind. We got engaged. It was quite the affair.

Our dog Jett has been diagnosed with diabetes and is now in the hospital to stabilize his blood sugar. Luckily, he seems to be doing well and he will be home with us soon. Unfortunately, his bill is large enough that we just can’t cover it on our own (especially with our poor financial situation). We’re being evicted from our home soon and trying to find a place with this bill is just too much for us by ourselves. To help us pay the bill we’ve set up a GoFundMe.
in each domain (“coverage” of the domain).

Results of the analysis are highlighted in Table 3. We first note that variety of expression depends on domain and topic; for example, the variety in the expression of negative emotions is particularly low in the financial domain (with 1.54% of words being negative emotion (“negemo”) words and only 31% of “negemo” words used). We also see clear topic shifts among domains: the interpersonal domains contain roughly 1.5 times as many social words, proportionally, as the others; and domains are stratified by their coverage of the anxiety word list (with the most in the mental illness domains and the least in the financial domain).

We also examine the overall lexical diversity of each domain by calculating Yule’s I measure (Yule, 1944). Figure 2 shows the lexical diversity of our data, both for all words in the vocabulary and for only words in LIWC’s “negemo” word list. Yule’s I measure reflects the repetitiveness of the data (as opposed to the broader coverage measured by our LIWC analysis). We notice exceptionally low lexical diversity for the mental illness domains, which we believe is due to the structured, clinical language surrounding mental illnesses. For example, posters in these domains discuss topics such as symptoms, medical care, and diagnoses (Figure 1, Table 2). When we restrict our analysis to negative emotion words, this pattern persists only for anxiety; the PTSD domain has comparatively little lexical variety, but what it does have contributes to its variety of expression for negative emotions.

By label. We perform similar analyses on data labeled stressful or non-stressful by a majority of annotators. We confirm some common results in the mental health literature, including that stressful data uses more first-person pronouns (perhaps reflecting increased self-focus) and that non-stressful data uses more social words (perhaps reflecting a better social support network).
Table 4: LIWC Analysis by Label. Results from our analysis using LIWC word lists, with the same definitions as in Table 3. First-person pronouns (“1st-Person”) use the LIWC “I” word list.

<table>
<thead>
<tr>
<th>Measure</th>
<th>Stress</th>
<th>Non-Stress</th>
</tr>
</thead>
<tbody>
<tr>
<td>% Conjunctions</td>
<td>0.88%</td>
<td>0.74%</td>
</tr>
<tr>
<td>Tokens/Segment</td>
<td>100.80</td>
<td>93.39</td>
</tr>
<tr>
<td>Clauses/Sentence</td>
<td>4.86</td>
<td>4.33</td>
</tr>
<tr>
<td>F-K Grade</td>
<td>5.31</td>
<td>5.60</td>
</tr>
<tr>
<td>ARI</td>
<td>4.39</td>
<td>5.01</td>
</tr>
</tbody>
</table>

Table 5: Complexity by Label. Measures of syntactic complexity for stressful and non-stressful data.

Figure 3: Lexical Diversity by Agreement. Yule’s I measure (on the y-axis) is plotted against domain size (on the x-axis) for each level of annotator agreement. Perfect means all annotators agreed; High, 4/5 or more; Medium, 3/5 or more; and Low, everything else.

Additionally, we calculate measures of syntactic complexity, including the percentage of words that are conjunctions, average number of tokens per labeled segment, average number of clauses per sentence, Flesch-Kincaid Grade Level (Kincaid et al., 1975), and Automated Readability Index (Senter and Smith, 1967). These scores are comparable for all splits of our data; however, as shown in Table 5, we do see non-significant but persistent differences between stressful and non-stressful data, with stressful data being generally longer and more complex but also rated simpler by readability indices. These findings are intriguing and can be explored in future work.

By agreement. Finally, we examine the differences among annotator agreement levels. We find an inverse relationship between the lexical variety and the proportion of annotators who agree, as shown in Figure 3. While the amount of data and lexical variety seem to be related, Yule’s I measure controls for length, so we believe that this trend reflects a difference in the type of data that encourages high or low agreement.

5 Methods

In order to train supervised models, we group the labeled segments by post and randomly select 10% of the posts (≈10% of the labeled segments) to form a test set. This ensures that while there is a reasonable distribution of labels and domains in the train and test set, the two do not explicitly share any of the same content. This results in a total of 2,838 train data points (51.6% labeled stressful) and 715 test data points (52.4% labeled stressful). Because our data is relatively small, we train our traditional supervised models with 10-fold cross-validation; for our neural models, we break off a further random 10% of the training data for validation and average the predictions of 10 randomly-initialized trained models.

In addition to the words of the posts (both as bag-of-n-grams and distributed word embeddings), we include features in three categories:

Lexical features. Average, maximum, and minimum scores for pleasantness, activation, and imagery from the Dictionary of Affect in Language (DAL) (Whissel, 2009); the full suite of 93 LIWC features; and sentiment calculated using the Pattern sentiment library (Smedt and Daelemans, 2012).

Syntactic features. Part-of-speech unigrams and bigrams, the Flesch-Kincaid Grade Level, and the Automated Readability Index.

Social media features. The UTC timestamp of the post; the ratio of upvotes to downvotes on the post, where an upvote roughly corresponds to a reaction of “like” and a downvote to “dislike” (upvote ratio); the net score of the post (karma) (calculated by Reddit, $n_{\text{upvotes}} - n_{\text{downvotes}}$); and

https://www.reddit.com/wiki/faq
the total number of comments in the entire thread under the post.

5.1 Supervised Models

We first experiment with a suite of non-neural models, including Support Vector Machines (SVMs), logistic regression, Naïve Bayes, Perceptron, and decision trees. We tune the parameters for these models using grid search and 10-fold cross-validation, and obtain results for different combinations of input and features.

For input representation, we experiment with bag-of-n-grams (for \( n \in \{1..3\} \)), Google News pre-trained Word2Vec embeddings (300-dimensional) (Mikolov et al., 2013), Word2Vec embeddings trained on our large unlabeled corpus (300-dimensional, to match), and BERT embeddings trained on our unlabeled corpus (768-dimensional, the top-level [CLS] embedding) (Devlin et al., 2019). We experiment with subsets of the above features, including separating the features by category (lexical, syntactic, social) and by magnitude of the Pearson correlation coefficient (\( r \)) with the training labels. Finally, we stratify the training data by annotator agreement, including separate experiments on only data for which all annotators agreed, data for which at least 4/5 annotators agreed, and so on.

We finally experiment with neural models, although our dataset is relatively small. We train both a two-layer bidirectional Gated Recurrent Neural Network (GRNN) (Cho et al., 2014) and Convolutional Neural Network (CNN) (as designed in Kim (2014)) with parallel filters of size 2 and 3, as these have been shown to be effective in the literature on emotion detection in text (e.g., Xu et al., 2018; Abdul-Mageed and Ungar (2017)). Because neural models require large amounts of data, we do not cull the data by annotator agreement for these experiments and use all the labeled data we have. We experiment with training embeddings with random initialization as well as initializing with our domain-specific Word2Vec embeddings, and we also concatenate the best feature set from our non-neural experiments onto the representations after the recurrent and convolutional/pooling layers respectively.

Finally, we apply BERT directly to our task, fine-tuning the pretrained BERT-base\(^8\) on our classification task for three epochs (as performed in Devlin et al. (2019) when applying BERT to any task). Our parameter settings for our various models are available in the appendix.

6 Results and Discussion

We present our results in Table 6. Our best model is a logistic regression classifier with Word2Vec embeddings trained on our unlabeled corpus, high-correlation features (\( \geq 0.4 \) absolute Pearson’s \( r \)), and high-agreement data (at least 4/5 annotators agreed); this model achieves an F-score of 79.8 on our test set, a significant improvement over the majority baseline, the n-gram baseline, and the pre-trained embedding model, (all by the approximate randomization test, \( p < 0.01 \)). The high-correlation features used by this model are LIWC’s clout, tone, and “I” pronoun features, and we investigate the use of these features in the other model types. Particularly, we apply different architectures (GRNN and CNN) and different input representations (pretrained Word2Vec, domain-specific BERT).

We find that our logistic regression classifier described above achieves comparable performance to BERT-base (approximate randomization test, \( p > 0.5 \)) with the added benefits of increased interpretability and less intensive training. Additionally, domain-specific word embeddings trained on our unlabeled corpus (Word2Vec, BERT) significantly outperform n-grams or pretrained embeddings, as expected, signaling the importance of domain knowledge in this problem.

We note that our basic deep learning models do not perform as well as our traditional supervised models or BERT, although they consistently, significantly outperform the majority baseline. We believe this is due to a serious lack of data; our labeled dataset is orders of magnitude smaller than neural models typically require to perform well. We expect that neural models can make good use of our large unlabeled dataset, which we plan to explore in future work. We believe that the superior performance of the pretrained BERT-base model (which uses no additional features) on our dataset supports this hypothesis as well.

In Table 7, we examine the impact of different feature sets and levels of annotator agreement on our logistic regressor with domain-specific Word2Vec embeddings and find consistent patterns supporting this model. First, we
<table>
<thead>
<tr>
<th>Model</th>
<th>P</th>
<th>R</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td>Majority baseline</td>
<td>0.5161</td>
<td>1.0000</td>
<td>0.6808</td>
</tr>
<tr>
<td>CNN + features*</td>
<td>0.6023</td>
<td>0.8455</td>
<td>0.7035</td>
</tr>
<tr>
<td>CNN*</td>
<td>0.5840</td>
<td>0.9322</td>
<td>0.7182</td>
</tr>
<tr>
<td>GRNN w/ attention + features*</td>
<td>0.6792</td>
<td>0.7859</td>
<td>0.7286</td>
</tr>
<tr>
<td>GRNN w/ attention*</td>
<td>0.7020</td>
<td>0.7724</td>
<td>0.7355</td>
</tr>
<tr>
<td>n-gram baseline*</td>
<td>0.7249</td>
<td>0.7642</td>
<td>0.7441</td>
</tr>
<tr>
<td>n-grams + features*</td>
<td>0.7474</td>
<td>0.7940</td>
<td>0.7700</td>
</tr>
<tr>
<td>LogReg w/ pretrained Word2Vec + features</td>
<td>0.7346</td>
<td>0.8103</td>
<td>0.7706</td>
</tr>
<tr>
<td>LogReg w/ fine-tuned BERT LM + features*</td>
<td>0.7704</td>
<td>0.8184</td>
<td>0.7937</td>
</tr>
<tr>
<td>LogReg w/ domain Word2Vec + features*</td>
<td>0.7433</td>
<td>0.8320</td>
<td>0.7980</td>
</tr>
<tr>
<td>BERT-base*</td>
<td>0.7518</td>
<td>0.8699</td>
<td>0.8065</td>
</tr>
</tbody>
</table>

Table 6: Supervised Results. Precision (P), recall (R), and F1-score (F) for our supervised models. Our best model achieves 79.80 F1-score on our test set, comparable to the state-of-the-art pretrained BERT-base model. In this table, “features” always refers to our best-performing feature set (≥ 0.4 absolute Pearson’s r). Models marked with a * show a significant improvement over the majority baseline (approximate randomization test, $p < 0.01$).

<table>
<thead>
<tr>
<th>Agreement Threshold for Data</th>
<th>Any Majority</th>
<th>60% (3/5)</th>
<th>80% (4/5)</th>
<th>100% (5/5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>75.40</td>
<td>76.31</td>
<td>78.48</td>
<td>77.69</td>
</tr>
<tr>
<td>All</td>
<td>76.90</td>
<td>77.12</td>
<td>77.10</td>
<td>78.28</td>
</tr>
<tr>
<td>LIWC</td>
<td>77.91</td>
<td>78.91</td>
<td>78.16</td>
<td>77.66</td>
</tr>
<tr>
<td>DAL</td>
<td>75.58</td>
<td>77.06</td>
<td>78.05</td>
<td>77.06</td>
</tr>
<tr>
<td>Lexical</td>
<td>76.42</td>
<td>77.92</td>
<td>77.54</td>
<td>77.88</td>
</tr>
<tr>
<td>Syntactic</td>
<td>74.63</td>
<td>75.49</td>
<td>76.66</td>
<td>76.19</td>
</tr>
<tr>
<td>Social</td>
<td>76.67</td>
<td>76.45</td>
<td>78.38</td>
<td>78.06</td>
</tr>
<tr>
<td>$</td>
<td>r</td>
<td>\geq 0.4$</td>
<td>77.44</td>
<td>78.76</td>
</tr>
<tr>
<td>$</td>
<td>r</td>
<td>\geq 0.3$</td>
<td>77.01</td>
<td>78.28</td>
</tr>
<tr>
<td>$</td>
<td>r</td>
<td>\geq 0.2$</td>
<td>77.53</td>
<td>78.61</td>
</tr>
<tr>
<td>$</td>
<td>r</td>
<td>\geq 0.1$</td>
<td>76.61</td>
<td>77.07</td>
</tr>
</tbody>
</table>

Table 7: Feature Sets and Data Sets. The results of our best classifier trained on different subsets of features and data. Features are grouped by type and by magnitude of their Pearson correlation with the train labels (no features had an absolute correlation greater than 0.5); data is separated by the proportion of annotators who agreed. Our best score (corresponding to our best non-neural model) is shown in bold.

We see a tradeoff between data size and data quality, where lower-agreement data (which can be seen as lower-quality) results in worse performance, but the larger 80% agreement data consistently outperforms the smaller perfect agreement data. Additionally, LIWC features consistently perform well while syntactic features consistently do not, and we see a trend towards the quality of features over their quantity; those with the highest Pearson correlation with the train set (which all happen to be LIWC features) outperform sets with lower correlations, which in turn outperform the set of all features. This suggests that stress detection is a highly lexical problem, and in particular, resources developed with psychological applications in mind, like LIWC, are very helpful.

Finally, we perform an error analysis of the two best-performing models. Although the dataset is nearly balanced, both BERT-base and our best logistic regression model greatly overclassify stress, as shown in Table 8, and they broadly overlap but do differ in their predictions (disagreeing with one another on approximately 100 instances).

We note that the examples misclassified by both models are often, though not always, ones with low annotator agreement (with the average percent agreement for misclassified examples being 0.55 for BERT and 0.61 for logistic regression). Both models seem to have trouble with less explicit expressions of stress, framing negative ex-
Hello everyone. A very close friend of mine was in an accident a few years ago and deals with PTSD. He has horrific nightmares that wake him up and keep him in a state of fright. We live in separate provinces, so when he does have his dreams it is difficult to comfort him. Each time he calls, and I struggle with what to say on the phone.

I asked the other day if they’ve set a date. He laughed in my face and said ‘no’ as if it were the most ridiculous thing he’s ever heard. He comes home late, and showers immediately. Then, he showers every morning before he leaves. He doesn’t talk to my mum and I, at all, and he’s cagey and secretive about everything, to the point of hostility towards my sister.

If he’s the textbook abuser, she is the textbook victim. She keeps giving him chances and accepting his apologies and living in this cycle of abuse. She thinks she’s the one doing something wrong. I keep telling her that the only thing she is doing wrong is staying with this guy and thinking he will change. I tell her she does not deserve this treatment.

We include some examples of these errors in Table 9, and further illustrative examples are available in the appendix.

7 Conclusion and Future Work

In this paper, we present a new dataset, Dreaddit, for stress classification in social media, and find the current baseline at 80% F-score on the binary stress classification problem. We believe this dataset has the potential to spur development of sophisticated, interpretable models of psychological stress. Analysis of our data and our models shows that stress detection is a highly lexical problem benefitting from domain knowledge, but we note there is still room for improvement, especially in incorporating the framing and intentions of the writer. We intend for our future work to use this dataset to contextualize stress and offer explanations using the content features of the text. Additional interesting problems applicable to this dataset include the development of effective distant labeling schemes, which is a significant first step to developing a quantitative model of stress.

Acknowledgements

We would like to thank Fei-Tzin Lee, Christopher Hidey, Diana Abagyan, and our anonymous reviewers for their insightful comments during the writing of this paper. This research was funded in part by a Presidential Fellowship from the Fu Foundation School of Engineering and Applied Science at Columbia University.

Table 8: Confusion Matrices. Confusion matrices of our best models and the gold labels. 0 represents data labeled not stressed while 1 represents data labeled stressed.

<table>
<thead>
<tr>
<th>Text</th>
<th>Gold Label</th>
<th>Agreement</th>
<th>Subreddit Name</th>
<th>Models Failed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hello everyone. A very close friend of mine was in an accident a few years ago and deals with PTSD. He has horrific nightmares that wake him up and keep him in a state of fright. We live in separate provinces, so when he does have his dreams it is difficult to comfort him. Each time he calls, and I struggle with what to say on the phone.</td>
<td>Not Stress</td>
<td>60%</td>
<td>ptsd</td>
<td>Both</td>
</tr>
<tr>
<td>I asked the other day if they’ve set a date. He laughed in my face and said ‘no’ as if it were the most ridiculous thing he’s ever heard. He comes home late, and showers immediately. Then, he showers every morning before he leaves. He doesn’t talk to my mum and I, at all, and he’s cagey and secretive about everything, to the point of hostility towards my sister.</td>
<td>Stress</td>
<td>60%</td>
<td>domesticviolence</td>
<td>BERT</td>
</tr>
<tr>
<td>If he’s the textbook abuser, she is the textbook victim. She keeps giving him chances and accepting his apologies and living in this cycle of abuse. She thinks she’s the one doing something wrong. I keep telling her that the only thing she is doing wrong is staying with this guy and thinking he will change. I tell her she does not deserve this treatment.</td>
<td>Not Stress</td>
<td>100%</td>
<td>domesticviolence</td>
<td>LogReg</td>
</tr>
</tbody>
</table>

Table 9: Error Analysis Examples. Examples of test samples our models failed to classify correctly. “BERT” refers to the state-of-the-art BERT-base model, while “LogReg” is our best logistic regressor described in section 6.
References


George Udny Yule. 1944. The statistical study of literary vocabulary. Cambridge Univ. Pr.

Towards Understanding of Medical Randomized Controlled Trials by Conclusion Generation

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Abstract
Randomized controlled trials (RCTs) represent the paramount evidence of clinical medicine. Using machines to interpret the massive amount of RCTs has the potential of aiding clinical decision-making. We propose a RCT conclusion generation task from the PubMed 200k RCT sentence classification dataset to examine the effectiveness of sequence-to-sequence models on understanding RCTs. We first build a pointer-generator baseline model for conclusion generation. Then we fine-tune the state-of-the-art GPT-2 language model, which is pre-trained with general domain data, for this new medical domain task. Both automatic and human evaluation show that our GPT-2 fine-tuned models achieve improved quality and correctness in the generated conclusions compared to the baseline pointer-generator model. Further inspection points out the limitations of this current approach and future directions to explore.

1 Introduction
Randomized controlled trials (RCTs) are the most rigorous method to assess the effectiveness of treatments, such as surgical procedures and drugs, in clinical medicine (Sibbald and Roland, 1998). A typical RCT often constitutes of two randomized groups of patients receiving either the “intervention” (new treatment) or “control” (conventional treatment). Then, a statistical analysis is done after the experiments to determine whether the intervention has a significant effect (i.e. actually making patients better or worse). The results from various RCTs contribute to the medical decisions made by physicians every day. However, analyzing these large amounts of data could be overwhelming for clinicians (Davidoff and Miglus, 2011). With the help of machine readers, we can alleviate the burden for providing correct information that contributes to critical clinical decisions.

In this work, we aim to evaluate the capabilities of deep learning models on understanding RCTs by generating the conclusions of RCT abstracts. We achieve this by transforming the PubMed 200k RCT abstract sentence classification dataset (Dernoncourt and Lee, 2017) into a RCT conclusion generation task. Generating a correct and coherent conclusion requires the model to 1) identify the objectives of the trial, 2) understand the result and 3) generate succinct yet comprehensible texts. Therefore, this task can be a preliminary goal toward a more thorough understanding of clinical medicine literature.

To tackle this task, we first build a pointer-generator model (See et al., 2017) as the baseline. This model is widely used in abstractive summarization, which is similar to our conclusion generation task. We then leverage the high quality text generation capability of the Open AI GPT-2 (Radford et al., 2019) language model by fine-tuning the general domain GPT-2 model into a medical domain conclusion generator.

Because the correctness of RCT understanding is essential for supporting clinical decisions and neural summarization models could inaccurately present facts from the source document, we incorporate human evaluation on the correctness and quality of the generated in addition to standard ROUGE score (Lin, 2004) for automated summarization scoring. Evaluation results show the fine-tuned GPT-2 models score higher for both correctness and quality. However, there is still quite a large room for improvement both on the diversity and accuracy of the generated conclusions, providing a guidance for future research directions.

*These authors contribute this paper equally.
*The code is available at: https://github.com/MiuLab/RCT-Gen
2 Related Work

The paper focuses on generating RCT conclusions, which is related to natural language generation. We describe the related work below and emphasize the difference between the prior work and our work. In our proposed method, we exploit the state-of-the-art language model representations for understanding the complex medical literature, and related work is then briefly described below.

2.1 Medical Natural Language Generation

Several medical domain natural language generation tasks have been studied using machine learning models, including generating radiology reports from images (Jing et al., 2018; Vaswani et al., 2017) and summarizing clinical reports (Zhang et al., 2018; Pivovarov and Elhadad, 2015) or research literature (Cohan et al., 2018). Recently, Gulden et al. (2019) studied extractive summarization on RCT descriptions.

Abstractive summarization, in which the model directly generates summaries from the source document, is closely related to our conclusion generation task. Most neural approaches for abstractive summarization were based on sequence-to-sequence recurrent neural networks (RNNs) with attention mechanisms (Devlin et al., 2019). The pointer-generator network (See et al., 2017) combined a copy mechanism that directly copies words from the source document and a coverage mechanism to avoid repetition caused by the RNN-based decoder, achieving good performance by handling unseen information. Devlin et al. (2019) further combined intra-encoder and intra-decoder attention with policy learning by using ROUGE-L score as the reward and improved the performance in terms of the evaluation metric. Hsu et al. (2018) combined an extractive model that provided attention on the sentence level and the pointer-generator architecture, and Cohan et al. (2018) also worked on abstractive summarization of long documents, including medical papers from the PubMed database, based on the pointer-generator network.

However, our goal to generate conclusions is different from abstractive summarization in that summarization is to shorten the source document while preserving most of the important information, whereas our conclusion generation model gives one or two sentences describing the main outcome of the given trial. Given the superior performance of pointer-generation networks from the above related summarization work, this paper uses the pointer-generation model as baseline and focuses on RCT conclusion generation instead of abstractive summarization.

2.2 Contextualized Representations

Recent advances of contextualized representation models, such as ELMo (Peters et al., 2018), Open AI GPT (Radford et al., 2018) and BERT (Devlin et al., 2019) achieved remarkable results across different natural language understanding tasks, such as question answering, entailment classification and named entity recognition. At the core of these models was language modeling, with either forward prediction used in GPT, bidirectional prediction used in ELMo, or masked prediction used by BERT. Variants of BERT also improved the performance of bio-medical natural language understanding tasks (Xu et al., 2019; Pugaliya et al., 2019). Peng et al. (2019) further proposed a new benchmark to evaluate the performance of contextualized models in the bio-medical domain.

Particularly, the Open AI GPT-2 model (Radford et al., 2019) has demonstrated rudimentary zero-shot summarization capabilities with only language modeling training. Its forward prediction architecture made it suitable for autoregressive generation in a sequence-to-sequence task. Most benchmarks on contextualized representation were based on sequence classification tasks such as natural language inference and multiple choice question answering (Wang et al., 2018; Peng et al., 2019). Our work, on the other hand, focuses on exploring GPT-2’s capability of generating goal-directed sentences in the medical domain. Note that to our knowledge, this paper is the first attempt that investigates GPT-2 towards the medical document understanding and interpretation.

3 Task Formulation

The PubMed 200k RCT dataset was originally constructed for sequential short text classification, with each sentence labeled as “background”, “objective”, “methods”, “results” and “conclusions”. We concatenated the “background”, “objective” and “results” sections of each RCT paper abstract as the model input and the goal of the model is to generate the “conclusions”. Table 1 illustrates
Varenicline is believed to work, in part, by reducing craving responses to smoking cues and by reducing general levels of craving; however, these hypotheses have never been evaluated with craving assessed in the natural environments of treatment-seeking smokers.

**Objectives:** Ecological momentary assessment procedures were used to assess the impact of varenicline on cue-specific and general craving in treatment-seeking smokers prior to quitting.

**Results:** During all phases, smoking cues elicited greater craving than neutral cues; the magnitude of this effect declined after the first week. General craving declined across each phase of the study. Relative to the placebo condition, varenicline was associated with a greater decline in general craving over the drug manipulation phase. Varenicline did not significantly attenuate cue-specific craving during any phase of the study.

Target (True Negative):
Smoking cues delivered in the natural environment elicited strong craving responses in treatment-seeking smokers, but cue-specific craving was not affected by varenicline administered prior to the quit attempt. These findings suggest that the clinical efficacy of varenicline is not mediated by changes in cue-specific craving during the pre-quit period of treatment-seeking smokers.

Table 1: An example of the GPT-2 model generating a false negative conclusion (Varenicline did reduce general craving), while the GPT-2 model generated a better true negative one. The “(BACKGROUND)”, “(OBJECTIVE)” and “(RESULTS)” tags denote the sentence classifications according to the original PubMed RCT dataset and are not included in the actual input of our conclusion generation task.

### 4 Models

Language model pre-training has achieved a great success among language understanding tasks with different model architectures. Because training language models requires a large amount of text data, and it is relatively difficult to acquire a lot of RCT documents, this work focuses on first pre-training language models with the transformer architecture (Vaswani et al., 2017) and then adapts the model to support the medical domain by fine-tuning. The language model pre-training from general texts is described below.

#### 4.1 Transformer Encoder in GPT-2

We first introduce the transformer encoder (Vaswani et al., 2017) used as the backbone of the GPT-2 model. The transformer encoder is a stack of $N$ transformer encoder blocks, where the $l$-th block takes a sequence of hidden representations $X^l = \{X^l_1, \cdots, X^l_n\}$ as the input and outputs an encoded sequence $X^{l+1} = \{X^{l+1}_1, \cdots, X^{l+1}_n\}$.
A transformer encoder block consists of a multi-head self-attention layer and a position-wise fully connected feed-forward layer. A residual connection (He et al., 2016) is employed around each of the two layers followed by layer normalization (Ba et al., 2016). In GPT-2, however, the layer normalization step is moved to the front of the multi-head self-attention layers and the feed-forward layers. An illustration of a GPT-2 transformer encoder block is presented in Figure 1. Each component is briefly described as follows.

**Byte-Pair Encoding** GPT-2 uses a special byte pair encoding (BPE) for input and output representations. It can cover essentially all Unicode strings, which is useful in processing the medical texts due to the significant out-of-vocabulary problems such as distinct nomenclature and jargon. This special BPE prevents merging characters from different categories and preserves word-level segmentation properties with a space exception.

**Positional Encoding** Because the transformer model relies on a self-attention mechanism with no recurrence, the model is unaware of the sequential order of inputs. To provide the model with positional information, positional encodings are applied to the input token embeddings

\[ X^l_i = \text{embed}_{\text{token}}[w_i] + \text{embed}_{\text{pos}}[i], \]

where \( w_i \) denotes the \( i \)-th input token, \( \text{embed}_{\text{token}} \) and \( \text{embed}_{\text{pos}} \) denote a learned token embedding matrix and a learned positional embedding matrix respectively.

**Multi-Head Self-attention** An attention function can be described as mapping a query to an output with a set of key-value pairs. The output is a weighted sum of values. We denote queries, keys and values as \( Q, K \) and \( V \), respectively. Following the original implementation (Vaswani et al., 2017), a scaled dot-product attention is employed as the attention function. Hence, the output can be calculated as

\[ \text{Attention}(Q, K, V) = \text{softmax}\left( \frac{QK^T}{\sqrt{d_k}} \right)V, \]

where \( d_k \) denotes the dimension of key vectors.

The idea of multi-head attention is to compute multiple independent attention heads in parallel, and then concatenate the results and project again.

The multi-head self-attention in the \( l \)-th block can be calculated as

\[ \text{MultiHead}(X^l) = \text{Concat}(\text{head}_1, \cdots, \text{head}_h)W^O, \]

\[ \text{head}_i = \text{Attention}(X^lW^Q_i, X^lW^K_i, X^lW^V_i), \]

where \( X^l \) denotes the input sequence of the \( l \)-th block, \( h \) denotes the number of heads, \( W^Q_i, W^K_i, W^V_i \) and \( W^O \) are parameter matrices.

**Position-Wise Feed-Forward Layer** The second sublayer in a block is a position-wise feed-forward layer, which is applied to each position separately and independently. The output of this layer can be calculated as

\[ \text{FFN}(x) = \max(0, x \cdot W_1 + b_1)W_2 + b_2, \]

where \( W_1 \) and \( W_2 \) are parameter matrices, \( b_1 \) and \( b_2 \) are parameter biases.

**Residual Connection and Layer Normalization** As shown in Figure 1, layer normalization is first applied on the input to the multi-head attention and feed-forward sublayers. The residual connection is then added around the two sublayers. The output of the \( l \)-th block can be calculated as

\[ H^l = \text{MultiHead}(\text{LayerNorm}(X^l)) + X^l, \]

\[ X^{l+1} = \text{FFN}(\text{LayerNorm}(H^l)) + H^l. \]

### 4.2 GPT-2 Pre-Training

The generative pre-training (GPT) via a language model objective is shown to be effective for learning representations that capture syntactic and semantic information without supervision (Peters et al., 2018; Radford et al., 2018; Devlin et al., 2019). The GPT model proposed by Radford et al. (2018) employs the transformer encoder with 12 encoder blocks. It is pre-trained on a large generic corpus that covers a wide range of topics. The training objective is to minimize the negative log-likelihood:

\[ \mathcal{L} = \sum_{t=1}^{T} - \log P(w_t | w_{<t}, \theta), \]

where \( w_t \) denotes the \( t \)-th word in the sentence, \( w_{<t} \) denotes all words prior to \( w_t \), and \( \theta \) are parameters of the transformer model.

To avoid seeing the future contexts, a masked self-attention is applied to the encoding process.
In the masked self-attention, the attention function is modified into

$$\text{Attention}(Q, K, V) = \text{softmax}(\frac{QK^T}{\sqrt{d_k}} + M)V,$$

where $M$ is a matrix representing masks. $M_{ij} = -\infty$ indicates that the $j$-th token has no contribution to the output of the $i$-th token, so it is essentially “masked out” when encoding the $i$-th token. Therefore, by setting $M_{ij} = -\infty$ for all $j > i$, we can calculate all outputs simultaneously without looking at future contexts. It was pre-trained on the WebText dataset consisting of 40 GB high-quality text crawled from internet sources. We use the small version (12 layers and 117 M parameters) of the released GPT-2 models.

4.3 GPT-2 Fine-Tuning

After the model is pre-trained with a language model objective, it can be fine-tuned on downstream tasks with supervised data. In our task, we adapt the GPT-2 to the target domain by fine-tuning using RCT data. Figure 2 illustrates the learning procedure. By fine-tuning on the target data, the GPT-2 model may have the potential of understanding and generating medical texts.

In the fine-tuning stage, we modify the attention masking of the GPT-2 model so that source byte pairs are fully aware of the entire context of the source sentence, while the target byte pairs are aware of the entire source sentence plus the generated byte pairs that precede itself. That is, for context token pairs $(c_i, c_j) \in c_1, \cdots, c_m$, we set $M_{ij} = -\infty$ for all $j > i$, while for context and source token pairs $(c_i, s_j)$, where $c_i \in c_1, \cdots, c_m$ and $s_j \in s_1, \cdots, s_n$, we set $M_{ij} = 0$. For all source token pairs $(s_i, s_j) \in s_1, \cdots, s_n$, we also set $M_{ij} = 0$. This setting is illustrated in Figure 3.

5 Experiments

Here we describe experimental details of the baseline pointer-generator model and the GPT-2 fine-tuned models.

5.1 Experimental Setup

The baseline model is a pointer-generator network (See et al., 2017) with both copy and coverage mechanisms, and is trained with a coverage loss. We adopt the implementation of Zhang et al. (2018). The vocabulary size is about 50,000, with uncased word embeddings pretrained from the PubMed RCT 200k training set and the abstracts from the PubMed dataset of long documents (Cohan et al., 2018). We concatenate $n \in \{0, 1, 3, 5\}$ hint words following the source sentences, where the hint words are first $n$ words of the target conclusion. Our pointer-generator model uses beam search with beam size 5 to decode the final output conclusion.

In our GPT-2 models, we conduct conclusion generation using $n \in \{0, 1, 3, 5\}$ hint words. For $n = 0$, we append “In conclusion , ” to the in-
### 5.2 Automatic Evaluation

Table 2 shows the best validation ROUGE scores of baselines and our models. Note that the hint words are not considered in score calculation and the output of all models are lower-cased. The GPT-2 fine-tuned model significantly outperforms the pointer-generator (PGNet) baseline on all ROUGE scores, where the best performing model is GPT-2 with hint word $n = 1$, demonstrating the effectiveness of generating good conclusions in our model. However, more hint words do not bring additional gain in ROUGE scores, probably because more constraints hinder the GPT-2 model to explore potentially good conclusions. Moreover, the ablation result shows the significant drop in all ROUGE scores, indicating the importance of including the “background” and “objective” sections in the input for better content understanding.

### 5.3 Human Evaluation

We recruited 10 medical students with prior training in bio-statistics and epidemiology to annotate and rate the generated conclusions. Our questionnaire contains two types of questions: the annotation question and the rating question.

- **Annotation**: A annotation question contains a source document and four conclusions, namely the target conclusion written by human, the GPT-2 $n = 0$, the GPT-2 $n = 1$ and the PGNet generated conclusions. The raters are asked to classify each generated conclusions as either true positive, true negative, false positive, false negative or not applicable. We define true / false as whether the generated conclusion corresponds to the given document, and positive / negative as whether the intervention studied has a statistically significant effect, regardless of the effect being favourable or detrimental to the patients. This is to explicitly examine whether the generated conclusion is precise in terms of RCT content understanding.

- **Rating**: Rating questions use the same set-up except the question is a 5 point Likert scale for correctness, quality and overall impression. Each rater is given 5 annotation questions and 5 rating questions, with each source document randomly chosen from the validation set. This is to judge the generated conclusions both regarding to and regardless of the source document.
To mitigate bias, we do not inform which conclusion was generated or written by human, and the conclusions are lower-cased and randomly ordered in each question for fair comparison.

Table 3 presents the results from the annotation question, where the number of true positive and true negative generations from the GPT-2 fine-tuned models increase when compared to the PGNet baseline. It is clear that the proposed GPT-2 achieves better performance in terms of accuracy (the ratio of true samples). We also include the performance of human-written conclusions in the last row, which serves as the upper bound of this task. However, there is still a gap between human-written conclusions and the generated ones.

In the rating questions depicted in Table 4, the human written conclusions obtain a score nearly 4 out of 5 on all three dimensions. The GPT-2 models have comparable scores in overall impression, both scoring around 3.5 out of 5. The most significant improvement of the GPT-2 generated conclusions is the text quality, with the correctness improvement to a lesser extent. The correctness of GPT-2 $n=1$ is slightly better than that of GPT-2 $n=0$ in the annotation question, yet in the rating question, GPT-2 $n=0$ has a higher averaged score. In sum, the human evaluation results demonstrate that our models significantly outperform the baseline pointer generator and tell that the proposed RCT conclusion generation task is not the same as typical summarization task. However, errors occur when the study design becomes more complicated or the outcomes are complex. Therefore, future work should aim at enhancing the language understanding capabilities of generation models. Methods such as pre-training the GPT-2 models with medical domain literature or using external background knowledge might fill the missing gap in the correctness performance. This is very crucial regarding to our RCT understanding task and other tasks that require precise and reliable language generation.

Here we select 3 examples to better illustrate our evaluation methods and the discussed limitations of the current models. The example in Table 5 show two successful generations from the GPT-2 models. Table 6 shows a false positive example by the GPT-2 $n=1$ model. On the other hand, a false negative example generated by the GPT-2 $n=0$ can be seen in Table 7. The generated conclusions in Table 7 is also much shorter than the target conclusion written by human. Other factors that could cause this issue may include that the human authors mention information not included in the preceding source document, additional comments on the results and background knowledge and they paraphrase the same concept in different ways.

Given the above results, this paper opens a new research direction by formulating the RCT conclusion generation task and investigates the potential of language generation models towards better understanding of medical documents.

<table>
<thead>
<tr>
<th>System</th>
<th>Correctness</th>
<th>Quality</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>PGnet $n=1$</td>
<td>3.02</td>
<td>2.86</td>
<td>2.86</td>
</tr>
<tr>
<td>GPT-2 $n=0$</td>
<td>3.42</td>
<td>3.66</td>
<td>3.52</td>
</tr>
<tr>
<td>GPT-2 $n=1$</td>
<td>3.30</td>
<td>3.94</td>
<td>3.50</td>
</tr>
<tr>
<td>Target</td>
<td>3.92</td>
<td>4.08</td>
<td>3.98</td>
</tr>
</tbody>
</table>

Table 4: Human evaluation results for generation quality on the rating questions.

The generated conclusions in Table 7 is also much shorter than the targets. The average length generated by GPT-2 $n=0$ and GPT-2 $n=1$ are 19.4 and 21.0, while that of human written conclusions is 41.4. This could be caused by the limitation of greedy decoding, but the examples generated by PGNet, which applies beam search, only gives an average length of 22.6. This suggests investigation of additional measures to enrich and lengthen the generated conclusions in future work.

Another important issue is the correctness of the generation model. The GPT-2 models are able to identify simple patterns and generate conclusions with the correct relationship. However, errors occur when the study design becomes more complicated or the outcomes are complex. Therefore, future work should aim at enhancing the language understanding capabilities of generation models. Methods such as pre-training the GPT-2 models with medical domain literature or using external background knowledge might fill the missing gap in the correctness performance. This is very crucial regarding to our RCT understanding task and other tasks that require precise and reliable language generation.

Given the above results, this paper opens a new research direction by formulating the RCT conclusion generation task and investigates the potential of language generation models towards better understanding of medical documents.
Proton pump inhibitor (PPI) therapy is considered as the first choice for treatment of non-erosive reflux disease (NERD). However, NERD is less sensitive to PPIs than erosive gastroesophageal reflux disease (GERD) and the differences between PPIs and H2 receptor antagonists are less evident in NERD than in erosive GERD. Since gastric acid secretions are lower in the Japanese population than in Western populations, we aimed to investigate whether PPI therapy is really necessary for NERD patients in Japan. Both roxatidine and omeprazole significantly improved the heartburn score at 4 and 8 weeks. The clinical response rates did not differ between roxatidine and omeprazole. Both roxatidine and omeprazole significantly relieved not only reflux but also abdominal pain and indigestion. The degrees of improvement did not differ between the two groups.

Table 5: An example of GPT-2 models generating true positive conclusions.

Table 6: An example of GPT-2 \( n = 0 \) model generating a true negative conclusion, while the GPT-2 \( n = 1 \) model generated a false positive one.

Table 7: An example of GPT-2 \( n = 0 \) model generating a false negative conclusion, while the GPT-2 \( n = 1 \) model generated a true positive one.

### 7 Conclusion and Future Work

This work introduces the RCT paper conclusion generation task as a stepping stone to the automatic understanding of clinical research literature. Our results show the general domain pre-trained GPT-2 language model can be fine-tuned to generate medical domain conclusions. The evaluation results show improvements regarding to both quality and correctness in conclusions generated by the
fine-tuned GPT-2 model compared to the pointer-generator summarization model. Further study is needed to enhance the generation quality by reducing repetition errors and increasing the generation length, and to improve the correctness through better language understanding for practical clinical scenarios.

Beyond generating conclusions for RCT papers, generative language models in the medical domain with improved correctness and quality can open up new opportunities to tasks that require profound domain knowledge. For example, automatic generation of systemic review and meta-analysis articles.

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References


Abstract
This article presents experiments with pseudonymised Swedish clinical text used as training data to de-identify real clinical text with the future aim to transfer non-sensitive training data to other hospitals. Conditional Random Fields (CFR) and Long Short-Term Memory (LSTM) machine learning algorithms were used to train de-identification models. The two models were trained on pseudonymised data and evaluated on real data. For benchmarking, models were also trained on real data, and evaluated on real data as well as trained on pseudonymised data and evaluated on pseudonymised data. CRF showed better performance for some PHI information like Date Part, First Name and Last Name; consistent with some reports in the literature. In contrast, poor performances on Location and Health Care Unit information were noted, partially due to the constrained vocabulary in the pseudonymised training data. It is concluded that it is possible to train transferable models based on pseudonymised Swedish clinical data, but even small narrative and distributional variation could negatively impact performance.

1 Introduction
Electronic health records (EHR) are produced in a steady stream, with the potential of advancing future medical care. Research on EHR data holds the potential to improve our understanding of patient care, care processes, and disease characteristics and progression. However, much of the data is sensitive, containing Protected Health Information (PHI) such as personal names, addresses, phone numbers, that can identify particular individuals and thus cannot be available to the public for general scientific inquiry. Although good progress has been made in the general sub-field of de-identifying clinical text, the problem is still not fully resolved (Meystre et al., 2010; Yogarajan et al., 2018).

This study examines the use of pseudonymised health records as training data for de-identification tasks. Several ethical and scientific issues arise regarding the balance between maintaining patient confidentiality and the need for wider application of trained models. How will a de-identification system be constructed and used in a cross hospital setting without risking the privacy of patients? Is it possible to obscuring the training data by pseudonymising it and then use it for the training of a machine learning system?

De-identification and pseudonymisation are two related concepts. In this paper de-identification is used as a more general term to describe the process of finding personal health information to be able to conceal identifying information. A pseudonymised text is a text where the personal health information has been identified either manually or automatically and then replaced with realistic surrogates.

The research question in this study is whether it is possible to use de-identified and pseudonymised clinical text in Swedish as training data for de-identifying real clinical text, and hence make it possible to transfer the system cross hospital. We highlight whether learning from the exist-
ing, non-sensitive, pseudonymised Swedish clinical text can be useful in a new and different context; considering the normal variations in the distribution and nature of PHI information, and potential effects of scrubbing (Berman, 2003), that is, removing and modifying PHIs that was carried out to patient records during the de-identification process.

2 Previous research

The identification of PHI is a type of named entity recognition task where sensitive named entities specifically are identified. The first study with CRF-based de-identification for Swedish was on the gold standard Stockholm EPR PHI Corpus. The distribution of PHIs is shown in Table 1. In this instance, manual annotation with expert consensus was used to create the gold standard (Dalianis and Velupillai, 2010).

De-identification tasks based on the CRF machine learning algorithm has been carried out on this data set previously with precision scores ranging between 85% and 95%, recalls ranging between 71% and 87% and F1-scores between 0.76 and 0.91 (Dalianis and Velupillai, 2010; Berg and Dalianis, 2019).

One approach previously used for concealing the training set’s sensitive data was carried out by (Dalianis and Boström, 2012), using the Stockholm EPR PHI Corpus. In the study, the textual part of the data were used to create 14 different features and part of speech tags. The textual part was then removed, and only the features and part of speech tags were used for training a Random Forest model. Fairly high precision of 89.1 % was obtained, but with a recall of 54.3 % and F1-score of 64.8.

In contrast to using only the sensitive EHR data for training, McMurry et al. (2013) integrated both publicly available scientific, medical publications and private sensitive clinical notes to develop a de-identification system. While considering the term frequencies and part of speech tags between the two data sources, they used both rule lists and decision trees for their system. This was an interesting approach since it raised the prospect of using non-sensitive data in building useful de-identification models. However, it is not clear whether medical journals have significant advantages over any other public text, like news corpora, for detecting PHI. A study similar to McMurry et al. (2013), by Berg and Dalianis (2019), showed few benefits of combining non-medical public text and sensitive clinical notes to build a de-identification system for medical records.

More recently, deep learning approaches using recurrent neural networks seem to yield significant improvements over traditional rules-based methods or statistical machine learning (Dernoncourt et al., 2017). Still, recent studies indicate that combining several approaches will yield the best results. For instance, the best system in a recent de-identification shared task was a combination of bidirectional LSTM, CRF and a rule-based sub-system (Liu et al., 2017).

Significant domain variation, such as a different language, is an important factor that was not considered in the discussed shared task. Domain differences were cited as the reason for poor performance on psychiatric notes de-identification (Stubbs et al., 2017), compared with the previous de-identification task on general clinical narratives (Stubbs et al., 2015).

Within the same language and similar clinical settings, the change of domain is likely not substantial. While in future research it may be worth considering domain adaption techniques to work towards a system meant to be used between hospitals, they were not considered in this study, beyond the use of non-sensitive dictionaries for names and location.

3 Data and methods

In this study, machine learning approaches are used since the best de-identification systems appear to be machine learning-based (Kushida et al., 2012). While rule-based methods such as using dictionaries and pattern-matching were previously more prevalent than machine learning methods for solving text-based de-identification problems (Meystre et al., 2010), today it is more typical to have both approaches used, since rule-based methods still yield better results for some PHI information (Neamatullah et al., 2008b). Dictionaries and patterns were therefore used as features within one of the models.

3.1 Data

Two different data sets for de-identification were used: Stockholm EPR PHI Psuedo Corpus (Pseudo) as well as the Stockholm EPR PHI Cor-
The Stockholm EPR PHI Pseudo Corpus was produced from the Stockholm EPR PHI Corpus by automatically pseudonymising all PHIs. This process is described by Dalianis (2019). The Stockholm EPR PHI Corpus is described by Dalianis and Velupillai (2010). An example is shown in Figure 1 from the Stockholm EPR PHI Pseudo Corpus.

The Stockholm EPR PHI Corpus and the Stockholm EPR PHI Pseudo Corpus are both parts of Swedish Health Record Research Bank (HEALTH BANK). HEALTH BANK encompasses structured and unstructured data from 512 clinical units from Karolinska University Hospital collected from 2006 to 2014 (Dalianis et al., 2015).

The number of entities and types of entities in both the Stockholm EPR PHI Pseudo Corpus and the Stockholm EPR PHI Corpus is shown in Table 2. From Table 2, it can be observed that the distribution of PHI instances between the two data sets is somewhat similar, but there is a significant difference when it comes to unique instances between the two data sets. In total, the Real data set contains proportionally more unique instances than the Pseudo data set. The entities in the Real data set also tend to have more tokens.

### Table 2: The distribution of PHI instances between the Stockholm EPR PHI Corpus (Pseudo) and the Stockholm EPR PHI Pseudo Corpus (Real)

<table>
<thead>
<tr>
<th>PHI classes</th>
<th>Pseudo Unique</th>
<th>Real Unique</th>
</tr>
</thead>
<tbody>
<tr>
<td>First Name</td>
<td>885</td>
<td>938</td>
</tr>
<tr>
<td>Last Name</td>
<td>911</td>
<td>957</td>
</tr>
<tr>
<td>Age</td>
<td>51</td>
<td>64</td>
</tr>
<tr>
<td>Phone Number</td>
<td>310</td>
<td>327</td>
</tr>
<tr>
<td>Location</td>
<td>159</td>
<td>229</td>
</tr>
<tr>
<td>Full Date</td>
<td>726</td>
<td>770</td>
</tr>
<tr>
<td>Date Part</td>
<td>1897</td>
<td>2079</td>
</tr>
<tr>
<td>Health Care Unit</td>
<td>1278</td>
<td>2277</td>
</tr>
<tr>
<td>Total PHI instances</td>
<td>6217</td>
<td>7647</td>
</tr>
</tbody>
</table>

The two models were evaluated on both the real data set that is annotated for PHI, but not pseudonymised, 'Pseudo-Real', as well as on the pseudonymised data set, 'Pseudo-Pseudo'. For additional comparison basis models trained on the real data set were evaluated on test sets from the same data set, 'Real-Real'.

### 3.2.1 CRF

In this study, the CRF algorithm implemented in CRFSuite (Okazaki, 2007) is used with the sklearn-crfsuite wrapper and the LSTM architecture described by Lample et al. (2016), based on an open-source implementation with Tensorflow is used.

The linear-chain Conditional Random Fields model, implemented with sklearn-CRFSuite4, was used.
Discharge letter **Huddinge hospital**
Resp. specialist/chief physician **Caroline Berg**
Journal author **Marianne Lindgren**
Discharge Date 20120325
Care episode 20120311-20120318
Main diagnosis acc to ICD-10 D073
Medical history 52-year-old woman, familiar at the clinic. Goes to **Karin Lundgren** and to the pain clinic.

**Figure 1:** Example of a pseudonymised record. The original Swedish pseudonymised record is to the right and the translated version is to the left. The underlined words are the surrogates, where real data has been replaced with pseudonyms.

uses lexical, orthographic, syntactic and dictionary features. The CRF is based on trial-and-error experiments with feature sets described by Berg and Dalianis (2019), and uses the same features except for section features.

### 3.2.2 LSTM

The long short-term memory (LSTM) needs word embeddings as features for the training. Word2vec\(^5\) was used to produce word embeddings using shallow neural networks, based on two corpora: a clinical corpus and medical journals. For the training using real clinical data, word embeddings were produced using a clinical corpus of 200 million tokens that produced 300,824 vectors with a dimension of 300.

For the training with pseudo clinical data, word embeddings were produced using Läkartidningen corpus (The Swedish scientific medical journals from 1996 to 2005) containing 21 million tokens that produced 118,662 vectors with a dimension of 300. The reason for using Läkartidningen is that the corpus does not contain sensitive data and hence is also more easily usable for transferable cross hospital training.

### 4 Results

The results of the experimental work are summarised in Figure 2. As can be observed in the figure, the CRF algorithm seems to generally outperform the LSTM algorithm on all metrics; precision, recall and F1 measure.

This result is not consistent with repeated reports in the literature, where deep learning approaches such as LSTM have been shown to out-perform most other methods, including CRF. Since deep learning approaches normally require very large amounts of data, one explanation for this result could be that the word embeddings used in this study did not contain sufficient context variations required for more robust performance or an insufficient training set of annotated data.

The ability to identify date part and age entities are similar when training on pseudonymised data and real data for the CRF. In contrast, Location, Health Care Unit and Full Date were negatively affected when using pseudonymised training data regardless of using a CRF or LSTM model.

### 4.1 CRF - Results

Experimental results of the CRF algorithm are shown in Table 3. Not presented in the table is the combination of training on real data and evaluation of pseudo data (Real-Pseudo), but the results of this combination gave a precision of 86.37 and recall of 77.80% and an F1-score of 81.86.

### 4.2 LSTM - Results

The experimental results of the LSTM algorithm are shown in Table 4 and again, not presented in the table is the combination of training on real data and evaluation of pseudo data (Real-Pseudo). The result of this combination is a precision of 65.83% and recall of 74.79% and F1-score of 70.03.

### 5 Analysis

The training set used in this study has a substantially constrained vocabulary compared to the evaluation set, which may partially explain the overall performance achieved when evaluating on real data (Pseudo-Real). The pseudo (training)
version of the data has less PHI tokens and the entities are more often single tokens.

The Full Date structure yyyyddmm - yyyyddmm is commonly occurring in the pseudo data, and the dash between the dates, ",", is often incorrectly identified. For example, using the CRF algorithm on real-data training and pseudo-data testing (Real-Pseudo), of the 159 instances not identified as full dates tokens, sixty contain ",". The pseudo data uses the structure yyyyddmm while the real data uses yyddmm, which leads to errors. For these kinds of errors on standard data formats such as dates, it is easy to see how rule-based approaches using regular expressions could significantly improve the overall performance of the system.

The weakest performance area was for location information. There is a large variety of locations in the pseudo-data. These are also fairly specific and unlikely to occur in the real data, for example, locations with very few inhabitants. These uncommon rural places have names similar to residential homes (äldreboenden). There are multiple instances of the suffix 'gården' (yard) in the location pseudo-PHI, whereas, in the real data, the same suffix is common for care units.

In the pseudo-data, the care units are more general than in the real data, often too general to be annotated in the real data set. Infirmaries are fairly common in the real data but non-existent in the pseudo data. This lack of variation in the pseudo is partially responsible for the drop in performance.

There are at least two ways to think about mitigating this poor performance. First, location and care unit could be combined as one entity type since they are conceptually very similar, and sometimes have interchangeable entity names. Secondly, using more detailed municipality street and location mapping databases as dictionaries could be considered.

6 Discussion

There is one similar study to ours but for English by Yeniterzi et al. (2010), where the authors train their de-identification system with all combinations of pseudonymised textual data (or what they call resynthesized records) and real data and their results are in line with ours. However, there are some studies on cross-domain adaptation. In cross-domain adaption there is, however,
In this study, in addition to narrative differences between the training data and the target evaluation data, the number of care units and locations involved, as well as personal names, are widely varied. With large amounts of out of vocabulary variation, training on limited data will likely yield poor results. In practice, these data types exist in other non-sensitive sources such as city and rural location and street mapping data.

Except for location and care unit, evaluation on pseudo-data (Pseudo-Pseudo) produced better outcomes compared to performance on real-data (Pseudo-Real), which can be expected. What was a bit unexpected was the lower performance of the LSTM algorithm. The algorithm’s results would potentially have been improved by larger vector data or more labelled data (Dernoncourt et al., 2017). While clinical notes have unique linguistic structures and grammatical peculiarities, non-clinical data sources could still provide important contextual information for constructing a useful vector space. Additional sources using non-sensitive data, such as public corpora in the general domain, hold a potential to improve performance on the de-identification task, therefore this line of inquiry will be followed up on in future work. In the same vein, factoring in part of speech tags from other sources of clinical data could be useful in this case. For instance, there are de-identification databases of clinical text, such as MIMIC (Neamatullah et al., 2008a; Goldberger et al., 2000), which could be used as additional information for training purposes, and using only the part of speech tags reduces security risks (Boström and Dalianis, 2012).

Current results are calculated as exact matches, and the partial match is not factored in, which may affect the result. As mentioned in the analysis the CRF algorithm rarely classifies the '-' in between dates as a part of the dates, and these are therefore not counted as matches despite the most identifying parts of the entity being identified.

To improve the general performance, a combination of both the LSTM and CRF algorithms could be performed instead of testing them independently. Combining high-performance algorithms and the use of ensemble methods seem to produce the best results as reported in the literature (Dernoncourt et al., 2017; Liu et al., 2017), and these techniques will be investigated in future work on the data sets.

Figure 2: Line graphs visualising the results of both CRF and LSTM, and the outcomes of the evaluations. The x-axis have the PHI entities; Age, Date Part (Dp), First Name (Fn), Full Date (Fd), Health Care Unit (Hcu), Last Name (Ln), Location (L) and the average result (Avg).

a substantial domain change between the training and testing data, unlike in this study. Martinez et al. (2014) used models trained in one hospital on pathology reports in another hospital. Their system only required minor feature normalisation, and the reported results were comparable across the hospitals. Although this demonstrates feasibility, it is important to note that the pathology reports were from the same medical sub-speciality with only some narrative differences.
7 Conclusions and future directions

The results of this study suggest that although it is possible to train models on pseudonymised data for use in different contexts, there is severe deterioration in performance for some PHI information. Even small narrative and distributional variation could negatively impact performance. Transferring a system from one set of clinical text to a different set could result in the performance of the system deteriorating; in this study the Pseudo-Real case. This problem, what we call the cross pseudo-real text adaptation problem, is an issue that could happen due to the pseudonymisation/de-identification processes on the training data due to the narrative and distributional variation as well as other differences in the nature of the PHI between the training data and the target.

In the future, we will try to improve the pseudonymisation module described in Dalianis (2019) to produce a larger variation in the vocabulary as the lack of variation may affect the current result negatively. We will also apply the learned models to other Nordic languages such as Norwegian clinical text and use the system as a pre-annotation system to assist the manual annotators in their work to create a Norwegian gold standard.

Acknowledgments

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References


Automatic rubric-based content grading for clinical notes

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Abstract
Clinical notes provide important documentation critical to medical care, as well as billing and legal needs. Too little information degrades quality of care; too much information impedes care. Training for clinical note documentation is highly variable, depending on institutions and programs. In this work, we introduce the problem of automatic evaluation of note creation through rubric-based content grading, which has the potential for accelerating and regularizing clinical note documentation training. To this end, we describe our corpus creation methods as well as provide simple feature-based and neural network baseline systems. We further provide tagset and scaling experiments to inform readers of plausible expected performances. Our baselines show promising results with content point accuracy and kappa values at 0.86 and 0.71 on the test set.

1 Introduction
Clinical notes, essential aspects of clinical care, document the principal findings of the visit, hospital stay or treatment episode, including complaints, symptoms, relevant medical history, tests performed, assessments and plans. During an encounter or soon after, notes are created based on subjective history, objective observations, as well as clinician assessment and care plans. Although this is a regular aspect of clinical care in all institutions, there is a large variability in the details taken. Clinical documentation training is often informal and institution-dependent, as systematic training of clinical documentation can be time-consuming and expensive. Training involves continued monitoring of note quality and completeness.

In this work, we present the problem of clinical note grading and provide several baseline systems for automatic content grading of a clinical note given a predefined rubric. To solve the problem, we built a simple feature-based system and a simple BERT-based system. We additionally provide training size experiments and tagset experiments for our baseline systems, as well as experiments with the relevance of using Unified Medical Language System (UMLS) and similarity features.

2 Background
Clinical notes serve as critical documentation for several purposes: medical, billing, and legal requirements. At the same time, a clinical note is written, updated, and consumed for the purpose of communication between clinicians for clinical care over a period of time. Too much irrelevant information can become an impediment to clinical care. Therefore, the identification of a base level of information is required to assess a note.

At our institution, clinical note documentation (or medical scribe) trainees are assessed via quizzes and exams in which trainees produce clinical notes based on mock patients visits (written to mimic outpatient encounters). Clinical note responses are graded against a grading rubric by training specialists, who have medical scribing experience as well as specialized training in determining note quality. The goal is to train scribes to produce a clinical note based on listening to the patient-clinician conversation during a clinical visit. Thus, the scribe expected to actively producing content, not just merely transcribe dictations from a clinician.

The purpose of a note rubric is to encapsulate the base requirements of a clinical note. Rubrics contain 40-60 rubrics items which reflect the information that needs to be captured during a medical encounter. A rubric item is typically written as a phrase and includes medically relevant attributes. For example, a rubric item discussing a
symptom will typically require information about duration and severity. A rubric item discussing medication will often include dosage information. Each rubric is associated with a section of the note where it needs to be placed. For training purposes, standard note sections include: History of Present Illness (HPI), a detailed subjective record of the patient’s current complaints; Review of Systems (ROS), the patient’s subjective complaints grouped by organ system; Physical Exam (PE), the clinician’s objective findings grouped by organ system; and Assessment and Plan (AP), the clinician’s diagnosis and the next steps in treatment for each diagnosis. Figure 1 gives an example of a clinical note with these sections.

If the note contains text that satisfies a rubric item, then a content point for that rubric item is awarded. If the note contains an incorrect statement (e.g., the wrong medication dosage), then that rubric point is not awarded, regardless of a correct statement appearing elsewhere. If the note lacks the inclusion of a rubric point, then that rubric point is not awarded. At most, one content point can be awarded per rubric item. Examples of several rubric items with corresponding portions of a clinical note are shown in below.

**Rubric item examples**

- frequent_bm_3-4.times_per_day (HPI section), documents relevant symptom history – "The patient complains of frequent bowel movements, 3-4 times daily."
- pe_skin_intact_no_clubbing_cyanosis (PE section), documents physical exam performed during visit – "Skin: Skin intact. No clubbing or cyanosis. Warm and dry."
- plan_advise_brat_diet (AP section), documents that the provider recommended the BRAT diet to the patient – "Recommended that the patient follow the BRAT diet for the next few days."

### 3 Related Work

Most community efforts in automatic grading have been in the context of automatic essay grading (AEG) and automatic short answer grading (ASAG), both of which harbor significant differences than our rubric-based content grading task.

AEG involves rating the quality of an essay in terms of coherence, diction, and grammar variations. Typically, an essay is given a score, e.g., from 1-6. Applications include essay grading for standardized tests such as for the GMAT (Graduate Management Admission Test) or TOEFL (Test of English as Foreign Language) (Valenti et al., 2003). Key architects for these systems are often commercial organizations. Examples of commercial computer-assisted scoring (CAS) systems include E-rater and Intelligent Essay Assessor. Systems such as E-rater use a variety of linguistic features, including grammar, diction, and as well as including discourse level features (Attali and Burstein, 2006; Zesch et al., 2015). In another approach, the Intelligent Essay Assessor uses latent semantic analysis to abstract text to lower-rank dimension-cutting representations of documents. Scores are assigned based on similarity of new text to be graded to a corpus of previously graded text (Foltz et al., 1999). The release of the Kaggle dataset has made this type of data more available to the public (kaggle.com/c/asap aes). A key difference of AEG task from our grading task is that our efforts focus on specific content item grading and feedback, over a single holistic document level rating.

In ASAG, free text answers to a prompt are graded categorically or numerically. It is very closely related to paraphrasing, semantic similarity, and textual entailment. System reporting for this task has often been on isolated datasets with a wide range of topics and setups. Often, these systems require extensive feature engineering (Burrows et al., 2015). One example system is C-rater, produced by ETS, which grades based on the presence or absence of required content (Lea-
Each required piece of content, similar to our rubric, in the text is marked as absent, present, negated, with a default of not_scored. Text goes through several processing steps, including spelling correction, concept recognition, pronoun resolution, and parsing. These features are then sent through a maximum entropy model for classification. Semantic similarity approaches apply a mixture of deep processing features, e.g. shortest path between concepts or concept similarities (Mohler and Mihalcea, 2009). In the SemEval 2013 Task 7 Challenge, the task involved classification of student answers to questions, given a reference answer, and student answers. Student answers are judged to be correct, partially_correct_incomplete, contradictory, irrelevant, or non_domain (Dzikovska et al., 2013). Although it has much in common to our rubric-based content grading setup, short answer grading has less document level issues to contend with. Moreover, specifically for our case, document-level scoring has some non-linearity with the individual classification of sub-document level text, e.g. finding one contradictory piece of evidence negates a finding of a positive piece of evidence.

The work of (Nehm et al., 2012), which attempts to award content points for specific items for college biology evolution essays, most closely resembles our task. In the task, students are awarded points based on whether they articulate key natural selection concepts, e.g. familiar plant/trait gain (mutation causing snail to produce poisonous toxin would increase fitness). The authors experimented with configuring two text analytic platforms for this task: SPSS Text Analysis 3.0 (SPSSTA) and Summarization Integrated Development Environment (SIDE). SPSSTA requires building hand-crafted vocabulary and attached rules. SIDE uses a bag-of-words representation run through a support vector machine algorithm for text classification. Key differences from our task are that rubric items are more numerous and specific; furthermore, our medium is a clinical note, which has documented linguistic and document style differences than normal essays; finally, our goal is not only to grade but to give automated in-text feedback.

In this work, we present our system for grading a clinical note given a grading rubric, which also gives subdocument level feedback. To our knowledge, there has been no previous attempt at clinical note automatic content grading.

4 Corpus Creation

We created a corpus by grading training notes by multiple different trainees quizzed on the same mock patient visit quiz, which included 40 rubric items. Annotation was carried out using using the ehost annotation tool (South et al., 2012). Annotators were asked to highlight sentences for if they were relevant to a rubric item. Furthermore, they were to mark whether a highlight had one of four attributes: correct, incorrect, contrary, incorrect_missing_item, and incorrect_section.

frequent_bm_3-4_times_per_day attribute examples

- correct – "The patient reports having 3-4 bowel movements a day." (Located in the HPI)
- incorrect contrary – "The patient has been having bowel movements every 30 minutes." (Located in the HPI) Explanation: The frequency is much higher than what would be expected for 3-4 times per day. Thus the information content is considered to be inaccurate or contrary to what is given by the rubric.
- incorrect_missing_item – "The patient reports having many bowel movements each day." (Located in the HPI) Explanation: This statement does not give any inaccurate information but is missing a crucial element that is required to earn this rubric point, which is the frequency value is 3-4 times per day.
- incorrect_section – "The patient reports having 3-4 bowel movements a day." (Located in the AP) Explanation: This statement is correct, but is located in the wrong section of the note.

Parts of the note were marked for exclusion, such as short hand text (excluded because they are just notes taken by the scribe in training of the conversation) and the review of systems (ROS) part of the note (this was excluded because grading of that section was not enforced at its time of creation). Entire notes were marked for exclusion in cases where the note was intended for a different exam or in cases when the note contained all short hand and templated sections (e.g. only notes and section headers such as “Chief Complaint”). Discontinuous rubric item note contexts were linked together with a relation. The final corpus after included 338
Table 1: Content-point inter-annotator agreement (percent identity/kappa).

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Table 2: Offset-level inter-annotator agreement (label/label-attribute).

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Inter-annotator agreement was measured among 5 annotators for 9 files. We measured agreement at several levels. At a note level, we measured pairwise percent identity and Cohen’s kappa (McHugh, 2012) by content points. At the text offset level, we measured precision, recall, and f1 (Hripcsak and Rothschild, 2005) for inexact overlap of highlights both at the label level (e.g. cc_frequent_urination) and at a label attribute level (e.g. cc_frequent_urination:correct). Since incorrect_section is not counted in content-based grading, for both inter-annotator agreement and for subsequent analysis, incorrect_section highlights were counted as correct unless overlapping with a incorrect_missingitem highlight, which would make it count as incorrect_missingitem. The agreement scores are shown in Table 1 and 2. Fleiss kappa (Fleiss, 1971) at the content point level was at 0.714. The rest of the corpus was divided among 5 annotators.

5 Methods

We performed classification for two types of systems: a feature-based and a simple BERT based neural network system for text classification. Since discontinuous text highlights accounted for less than 10% of items, we choose not to model this nuance. Both systems used the same pre-processing pipeline configurations shown in Figure 2.

5.1 Pipeline configurations

We split 338 files into 270 training, 68 test set. Tuning was performed on the training set in 5-fold cross validation.

We split 338 files into 270 training, 68 test set. Tuning was performed on the training set in 5-fold cross validation.

5.2 Feature based system

The feature based system includes an n-gram feature extraction, which then passed through a chi-squared statistical feature selection, before using a support vector machine implemented by scikit learn svc.

If the umls configuration is turned on, then Unified Medical Language System (UMLS) concept with its negation value, extracted using MetaMap, concept-grams are also added (Aronson and Lang, 2010). If turned on, similarity features for n-grams

1Includes highlights for excluding the note, as well as excluding short hand text and ROS sections
Whether or not to turn off the feature extraction, instances were transformed into BERT word vector representations. We used the output for CLS position of the embeddings to represent the whole sequence similar to that of the original paper (Devlin et al., 2018).

To mimic the feature-based system’s case of simfeats_on, we include a switch to an architecture that also feeds in the CLS position output from a paired BERT classification setup. When simfeats_on is turned off, the architecture becomes that of a simple BERT sentence classification (bert). When text_off is turned on, then the architecture becomes that of a simple BERT sentence pair classification (bertpair). When simfeats_on is turned on and text_off is turned off, we have a system with both types of representations (bert+bertpair). A figure of the BERT classifier setup is show in Figure 3.

Because certain medical vocabulary may not be available with the general English trained cor-

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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plan_bp_goal_13080</td>
<td>153 123</td>
<td>11</td>
<td>18 3</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>plan_may_notice_log_swelling_notif_if_unbearable</td>
<td>216 93</td>
<td>114</td>
<td>5 4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plan_prescribed_amiodipine_5mg</td>
<td>268 205</td>
<td>40</td>
<td>18 5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plan_recommend_30_mins_physical_activity_3-5_times_per_week</td>
<td>296 128</td>
<td>131</td>
<td>34 3</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plan_reduce_stress_levels</td>
<td>119 100</td>
<td>13</td>
<td>1 5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plan_rnr_1_month_with_continuous_bp_log</td>
<td>280 172</td>
<td>85</td>
<td>16 7</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>plan_trial_today_to_infiltrate_out_info</td>
<td>302 202</td>
<td>91</td>
<td>3 6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>side_effects_of_difficulty_breathing_with_metoprolol</td>
<td>164 104</td>
<td>38</td>
<td>21 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>stress_work_related</td>
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<td>7</td>
<td>1 0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>takes_blood_pressure_every_morning</td>
<td>222 145</td>
<td>66</td>
<td>11 0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>tried_yogurt_and_jersey_poor_improvement</td>
<td>176 66</td>
<td>106</td>
<td>4 0</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>was_seen_by_pl_reynoldsesterday</td>
<td>154 44</td>
<td>96</td>
<td>13 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>weight_normal</td>
<td>61 52</td>
<td>5</td>
<td>3 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3: Label frequencies for full corpus

and umls concept grams (if the umls flag is on) are also added. We used jaccard similarity between 1-, 2-, and 3- grams. The full configurations include the following:

- **top_n**: top number of significant features to keep according to chi-squared statistic feature selection (integer)
- **sec_feat**: setting to determine how section information is encoded. If “embed” is set, then each feature with be concatenated with its section, e.g. “sect[hp]=patient”. If “sep” is turned on, “sect=hp” is added as a feature.
- **umls**: whether or not to use umls features (binary)
- **simfeat_on**: whether or not to turn on similarity features (binary)
- **text_off**: whether or not to turn off the features not related to similarity
- **umls_text_off**: whether or not to turn off the umls features not related to similarity
- **sent_win**: window for which surrounding sentence features should be added, e.g. sent[-1]=the would be added as a feature from previous sentence unigram feature “the” if sent_win=1. (integer)
Figure 3: BERT with additional sentence pair classifier

Poror, we used pre-trained BERT embedding and vocabulary from bio-bert, which is fine-tuned on pubmed data (Lee et al., 2019).

5.4 Simple baseline

For a further comparison, we have included a feature-based document-based baseline. This baseline largely follows the previously mentioned feature-based baseline though performed at the document level. Because it is performed at a document level, some attribution to a sub-document level unit becomes necessary. (Recall, we wish to be able to identify which part of the document is relevant to a rubric item as well as if we believe it is correct or otherwise.) To identify a corresponding offset labeling for this setting, we attributed a document level classification to a sentence which contained the maximum amount of important features. We defined features to be important according to their learned feature weight magnitude crossing a configurable threshold value. Thus, for a document classification, based on this logic we are able to assign a sentence related to that classification, for which we can use all our previously mentioned metrics for evaluation. We found a threshold of 10 to work well in our experiments.

5.5 Evaluation

Similar to inter-annotator agreement, we measured performance using several metrics. At the note level, we measured distance to target full document score by mean absolute error (MAE). We also measured content point accuracy and kappa to get a sense of the performance in point assignment.

At the offset level, we measured precision, recall, and f1 for rubric item label-attribute value. For minichunk classification, offsets were translated according to the start and end of the minichunk in the document.

6 Results

Evaluations in cross-validation are reported as the average of all 5 folds. Consistent with this, graphical error bars in the learning curve figures represent the standard deviation across folds.

6.1 Experimental configurations

Feature-based parameter tuning. We started with tag gran at 4lab, simfeats on true, and top n set at 1000, sent_win=0, and neg samp=2. We then greedily searched for an optimal solution varying one parameter at a time to optimize precision, recall and f1 measure. Our final configurations were set to neg samp=10, sect feat set to “sep”, top n=4000, sent_win=0. We kept the same configurations for the other feature-based systems.

Neural network hyperparameter tuning. For the neural network models, we mainly experimented with negative sampling size, dropout, and context length. We found a neg samp=100, dropout=0.5, epochs=2 and context len=300 to work well.

6.2 Cross-validation and test results

Table 4 shows performances for the feature-based system using different tagsets in cross-validation. Unsurprisingly the most detailed label (4lab) at the more granular level (minichunk) showed the best performance. Tables 4 and 5 shows the comparison of different system configurations in evaluated in cross validation. Table 6 shows results for the test set for different systems. In general, the feature-based system with the full feature-set out performed for the cross-validation and test experiments. Among the BERT systems, the simple BERT system did better than the other two configurations.

6.3 Scaling and tagset experiments

Figure 4 shows the effect of increasing training size for several metrics, under several select systems. Interestingly, 2lab and 3lab settings show similar behaviors across different metrics. For the document level baseline, tagset does not make any large difference across all three metrics. Different from other systems, 2lab and 3lab setting,
Figure 4: Learning curve experiments. 2lab, 3lab, 4lab demarks the tagset configuration. doc-* identifies the simple document classification baseline. The left column shows the performance of different n-gram configurations for 3 performance metrics. The right column shows BERT system performances for the 3 metrics along with two feature-based systems for comparison.
when trained to maximize both precision and recall equally for label attributes, MAE will rise instead of lower such as for the other system setups. This makes sense, as both tagset settings miss crucial examples that exhibit confusing features. For example, on the 2lab setting, only positive examples are shown not those that have incorrect information or those that have missing information (i.e. partially correct information). Likewise, the 3lab setting does not have evidence for partially correct items. We found experimentally, when tuning for higher precision and lower recall, that MAE also tends to lower– suggesting that these two settings can be better maximized by tuning for MAE instead.

Like the document-based baseline, the BERT systems’ performances showed that tagset did not make as big of a difference across all three metrics at different training size levels. This is possibly because there were not enough examples to even properly fine-tune for these two systems which require more training data. At higher levels of training sizes, tagsets may again come into effect. Though the BERT systems at lower training sizes start at lower performances, it quickly catches up to the document classification baseline for the MAE and f1 metrics, though never gets close to the 4lab feature-based baseline.

### Table 4: Feature-based system results for 5-fold cross-validation, varying tagsets.

<table>
<thead>
<tr>
<th>tagset</th>
<th>mae</th>
<th>acc</th>
<th>kappa</th>
<th>p</th>
<th>r</th>
<th>f</th>
</tr>
</thead>
<tbody>
<tr>
<td>4lab</td>
<td>2.0</td>
<td>0.85</td>
<td>0.70</td>
<td>0.71</td>
<td>0.75</td>
<td>0.73</td>
</tr>
<tr>
<td>3lab</td>
<td>4.5</td>
<td>0.82</td>
<td>0.63</td>
<td>0.69</td>
<td>0.64</td>
<td>0.66</td>
</tr>
<tr>
<td>2lab</td>
<td>5.4</td>
<td>0.80</td>
<td>0.60</td>
<td>0.69</td>
<td>0.63</td>
<td>0.66</td>
</tr>
<tr>
<td>doc-4lab</td>
<td>3.1</td>
<td>0.80</td>
<td>0.59</td>
<td>0.58</td>
<td>0.53</td>
<td>0.55</td>
</tr>
<tr>
<td>doc-3lab</td>
<td>2.5</td>
<td>0.80</td>
<td>0.58</td>
<td>0.65</td>
<td>0.43</td>
<td>0.52</td>
</tr>
<tr>
<td>doc-2lab</td>
<td>2.9</td>
<td>0.78</td>
<td>0.55</td>
<td>0.67</td>
<td>0.42</td>
<td>0.52</td>
</tr>
</tbody>
</table>

### Table 5: Results for 5-fold cross-validation, tag_gran=4lab.

<table>
<thead>
<tr>
<th>system</th>
<th>mae</th>
<th>acc</th>
<th>kappa</th>
<th>p</th>
<th>r</th>
<th>f</th>
</tr>
</thead>
<tbody>
<tr>
<td>ngram+simfeats+umls</td>
<td>2.0</td>
<td>0.85</td>
<td>0.70</td>
<td>0.71</td>
<td>0.75</td>
<td>0.73</td>
</tr>
<tr>
<td>ngram+umls</td>
<td>2.2</td>
<td>0.85</td>
<td>0.70</td>
<td>0.71</td>
<td>0.75</td>
<td>0.73</td>
</tr>
<tr>
<td>ngram+simfeats</td>
<td>2.1</td>
<td>0.84</td>
<td>0.69</td>
<td>0.70</td>
<td>0.74</td>
<td>0.72</td>
</tr>
<tr>
<td>simfeats+umls</td>
<td>3.7</td>
<td>0.75</td>
<td>0.50</td>
<td>0.63</td>
<td>0.53</td>
<td>0.57</td>
</tr>
<tr>
<td>bert+bert_pair</td>
<td>2.6</td>
<td>0.76</td>
<td>0.51</td>
<td>0.59</td>
<td>0.62</td>
<td>0.60</td>
</tr>
<tr>
<td>bert</td>
<td>2.8</td>
<td>0.76</td>
<td>0.51</td>
<td>0.60</td>
<td>0.61</td>
<td>0.60</td>
</tr>
<tr>
<td>bert_pair</td>
<td>2.9</td>
<td>0.74</td>
<td>0.48</td>
<td>0.57</td>
<td>0.56</td>
<td>0.56</td>
</tr>
</tbody>
</table>

### Table 6: Detailed results for the test set, tag_gran=4lab.

<table>
<thead>
<tr>
<th>system</th>
<th>mae</th>
<th>acc</th>
<th>kappa</th>
<th>p</th>
<th>r</th>
<th>f</th>
</tr>
</thead>
<tbody>
<tr>
<td>ngram+simfeats+umls</td>
<td>2.5</td>
<td>0.85</td>
<td>0.70</td>
<td>0.68</td>
<td>0.78</td>
<td>0.72</td>
</tr>
<tr>
<td>ngram+umls</td>
<td>2.5</td>
<td>0.85</td>
<td>0.70</td>
<td>0.68</td>
<td>0.78</td>
<td>0.72</td>
</tr>
<tr>
<td>ngram+simfeats</td>
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<td>0.86</td>
<td>0.71</td>
<td>0.68</td>
<td>0.79</td>
<td>0.73</td>
</tr>
<tr>
<td>simfeats+umls</td>
<td>3.8</td>
<td>0.77</td>
<td>0.54</td>
<td>0.63</td>
<td>0.55</td>
<td>0.59</td>
</tr>
<tr>
<td>bert+bert_pair</td>
<td>3.2</td>
<td>0.77</td>
<td>0.53</td>
<td>0.57</td>
<td>0.60</td>
<td>0.58</td>
</tr>
<tr>
<td>bert</td>
<td>2.8</td>
<td>0.79</td>
<td>0.57</td>
<td>0.59</td>
<td>0.63</td>
<td>0.61</td>
</tr>
<tr>
<td>bert_pair</td>
<td>3.1</td>
<td>0.76</td>
<td>0.51</td>
<td>0.58</td>
<td>0.60</td>
<td>0.59</td>
</tr>
</tbody>
</table>

### 6.4 UMLS and similarity features

The addition of similarity features did not provide a significant boost for the ngram feature-based system. Similarity features alone for the feature-based system underperformed at all training size levels compared to the ngram models. On the other hand, the addition of UMLS features increased performance across three metrics for all training size levels.

The BERT based system using only the simple BERT representation (without paired features), outperformed the other two settings across the three metrics at most training size levels in cross-validation. However, near higher levels of training data, BERT with BERT pair becomes comparable. The BERT pair system underperforms across all three metrics and at all training sizes.

### 6.5 Error Analysis

One challenging aspect of the classification task was the imbalanced categories across notes for different rubric items. Some labels were inherently less frequent, e.g. weight_normal had a total of 61 compared to cc_frequent_urination with 579 highlights. Indeed the performance amongst all rubric item scores was highly variable, with 13% f1 standard variation for the label-attribute measure. Moreover, the distribution of classes per label was also highly variable, as shown in Figure 3. For example, when no_recent_antibiotics or stress_work_related appears, in labeled data they are often correct. As a consequence, accurately predicting less populated classes becomes more difficult. For the best performing system, for example, there were instances where “Patients weight is not normal” was considered correct despite the rubric specifying the opposite. Similarly, “Patient denies feeling loopy” would be marked correct when the rubric says otherwise. When measuring at the label level for highlights instead, the performance on the test was higher by more than 10% f1, as shown in Table 7. This
<table>
<thead>
<tr>
<th>eval</th>
<th>tp</th>
<th>fp</th>
<th>fn</th>
<th>p</th>
<th>r</th>
<th>f</th>
</tr>
</thead>
<tbody>
<tr>
<td>label-att</td>
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<td>700</td>
<td>424</td>
<td>0.68</td>
<td>0.78</td>
<td>0.72</td>
</tr>
<tr>
<td>label</td>
<td>1708</td>
<td>413</td>
<td>186</td>
<td>0.81</td>
<td>0.90</td>
<td>0.85</td>
</tr>
</tbody>
</table>

Table 7: Results for the test set, ngram+simfeats+umls
tag_gran=4lab.

indicates that many errors are due to confusion between classification categories. Contradictions, labeled incorrect, contrary, for this reason was a large problem.

Manually studying errors in the test set for the best performing system, we found that rubric items frequently identified in the training, were broadly correctly classified. However, there were some rubric items that had more inconsistencies in how they were being tagged or graded. Some errors were partly due to human grading error. For example, checks_blood_sugar_regularly_110s_before_meals was a rubric item that scribes frequently missed when creating notes. Due to this, some sentences with just “checks” were sometimes labeled for checks_blood_sugar_regularly_110s_before_meals despite the fact that the sentences were about checking blood pressure. This leads to cases where synonymous phrases to “blood sugar”, “blood glucose”, did not get labeled as instances with “blood glucose” by the human graders.

7 Conclusions

In this paper we present the problem of clinical note grading and provide several baseline system evaluations at different levels of training data. We show experimentally that the choice of labeling has large effects upon the system performance. Furthermore, though neural network systems may relieve a lot of feature-engineering, this may not be plausible for smaller corpora.

Further improvements can be made by rubric-item specific pipeline specialization, as well as further augmentation of specific feature extraction modules, e.g. better negation handling. Deeper processing methods and features, including use of lemma-ization and dependency structures, would make features more generalizable. On the other hand, to maximize performance, feature-extraction can also be made more rubric-item specific, for example by hand-crafting features. For this work, we used a linear support vector machine for our classifier, but further experimentation with different classifiers for each rubric item would lead to higher performances.

The BERT systems can be improved by increasing training size and adding more feed-forward layers.

Our proposed system can be used to expedite and formalize clinical note creation in a training setting. For example, instead of having a human grader view all training notes, a simple pass of the automated grading system can eliminate those that will fail with some confidence. For others, a human grader can correct the output of the system, which would speed the grading process then if the grader had to mark highlights alone. In this work, we focus on cases for which we have many examples of clinical notes generated for the same encounter with a fixed rubric. Future work will investigate grading for arbitrary notes and rubrics.

Acknowledgments

We would like to thank the Augmedix training specialist team as well as all those involved with creating the source dataset for this work.

Very special thanks to Kevin Holub and Sonny Siddhu for their efforts in initiating Augmedix’s efforts in AI-assisted in-text grading for which is the motivation behind this project.

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Dilated LSTM with attention for Classification of Suicide Notes

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Abstract

In this paper we present a dilated LSTM with attention mechanism for document-level classification of suicide notes, last statements and depressed notes. We achieve an accuracy of 87.34% compared to competitive baselines of 80.35% (Logistic Model Tree) and 82.27% (Bi-directional LSTM with Attention). Furthermore, we provide an analysis of both the grammatical and thematic content of suicide notes, last statements and depressed notes. We find that the use of personal pronouns, cognitive processes and references to loved ones are most important. Finally, we show through visualisations of attention weights that the Dilated LSTM with attention is able to identify the same distinguishing features across documents as the linguistic analysis.

1 Introduction

Over recent years the use of social media platforms, such as blogging websites has become part of everyday life and there is increasing evidence emerging that social media can influence both suicide-related behaviour (Luxton et al., 2012) and other mental health conditions (Lin et al., 2016). Whilst there are efforts to tackle suicide and other mental health conditions online by social media platforms such as Facebook (Facebook, 2019), there are still concerns that there is not enough support and protection, especially for younger users (BBC, 2019). This has led to a notable increase in research of suicidal and depressed language usage (Coppersmith et al., 2015; Pestian et al., 2012) and subsequently triggered the development of new healthcare applications and methodologies that aid detection of concerning posts on social media platforms (Calvo et al., 2017). More recently, there has also been an increased use of deep learning techniques for such tasks (Schoene and Dethlefs, 2018), however there is little evidence which features are most relevant for the accurate classification. Therefore we firstly analyse the most important linguistic features in suicide notes, depressed notes and last statements. Last Statements have been of interest to researchers in both the legal and mental health community, because an inmates last statement is written, similarly to a suicide note, closely before their death (Texas Department of Criminal Justices, 2019). However, the main difference remains that unlike in cases of suicide, inmates on death row have no choice left in regards to when, how and where they will die. Furthermore there has been extensive analysis conducted on the mental health of death row inmates where depression was one of the most common mental illnesses. Work in suicide note identification has also compared the different states of mind of depressed and suicidal people, because depression is often related to suicide (Mind, 2013). Secondly, we introduce a recurrent neural network architecture that enables us to (1) model long sequences at document level and (2) visualise the most important words to accurate classification. Finally, we evaluate the results of the linguistic analysis against the results of the neural network visualisations and demonstrate how these features align. We believe that by exploring and comparing suicide notes with last statements and depressed notes, both qualitatively and quantitatively it could help us to find further differentiating factors and aid in identifying suicidal ideation.

2 Related Work

The analysis and classification of suicide notes, depression notes and last statements has traditionally been conducted separately. Work on suicide notes has often focused on identifying suicidal ideation online (O’dea et al., 2017) or distinguish-
ing genuine from forged suicide notes (Coulthard et al., 2016), whilst the main purpose of analysing last statements has been to identify psychological factors or key themes (Schuck and Ward, 2008).

Suicide Notes Recent years have seen an increase in the analysis of suicidal ideation on social media platforms, such as Twitter. Shahreem et al. (2018) searched the Twitter API for specific keywords and analysed the data using both traditional machine learning techniques as well as neural networks, achieving an accuracy of 97.6% using neural networks. Research conducted by Burnap et al. (2017) have developed a classifier to distinguish suicide-related themes such as the reports of suicides and casual references to suicide. Work by Just et al. (2017) used a dataset annotated for suicide risks by experts and a linguistic analysis tool (LIWC) to determine linguistic profiles of suicide-related twitter posts. Other work by Pestian et al. (2010) has looked into the analysis and automatic classification of sentiment in notes, where traditional machine learning algorithms were used. Another important area of suicide note research is the identification of forged suicide notes from genuine ones. Jones and Bennell (2007) have used supervised classification model and a set of linguistic features to distinguish genuine from forged suicide notes, achieving an accuracy of 82%.

Depression notes Work on identifying depression and other mental health conditions has become more prevalent over recent years, where a shared task was dedicated to distinguishing depression and PTSD (Post Traumatic Stress Disorder) on Twitter using machine learning (Coppersmith et al., 2015). Morales et al. (2017) have argued that changes in cognition of people with depression can lead to different language usage, which manifests itself in the use of specific linguistic features. Research conducted by Resnik et al. (2015) also used linguistic signals to detect depression with different topic modelling techniques. Work by Rude et al. (2004) used LIWC to analyse written documents by students who have experienced depression, currently depressed students as well as student who never have experienced depression, where it was found that individuals who have experienced depression used more first-person singular pronouns and negative emotion words. Nguyen et al. (2014) used LIWC to detect differences in language in online depression communities, where it was found that negative emotion words are good predictors of depressed text compared to control groups using a Lasso Model (Tibshirani, 1996). Research conducted by Morales and Levitan (2016) showed that using LIWC to identify sadness and fatigue helped to accurately classify depression.

Last statements Most work in the analysis of last statements of death row inmates has been conducted using data from The Texas Department of Criminal Justice, made available on their website (Texas Department of Criminal Justices, 2019). Recent work conducted by Foley and Kelly (2018) has primarily focused on the analysis of psychological factors, where it was found that specifically themes of ‘love’ and ‘spirituality’ were constant whilst requests for forgiveness declined over time. Kelly and Foley (2017) have identified that mental health conditions occur often in death row inmates with one of the most common conditions being depression. Research conducted by Heflick (2005) studied Texas last statements using qualitative methods and have found that often afterlife belief and claims on innocence are common themes in these notes. Eaton and Theuer (2009) studied qualitatively the level of apology and remorse in last statements, whilst also using logistic regression to predict the presence of apologies achieving an accuracy of 92.7%. Lester and Gunn III (2013) used the LIWC program to analyse last statements, where they have found nine main themes, including the affective and emotional processes. Also, Foley and Kelly (2018) found in a qualitative analysis that the most common themes in last statements were love (78%), spirituality (58%), regret (35%) and apology (35%).

3 Data

For our analysis and experiments we use three different datasets, which have been collected from different sources. For the experiments we use standard data preprocessing techniques and remove all identifying personal information.¹

Last Statements Death Row This dataset has been made available by the Texas Department of Criminal Justices (2019) and contains 545 records of prisoners who have received the death penalty between 1982 and 2017 in Texas, U.S.A. A total of 431 prisoners wrote notes prior to their death.

¹The authors are happy to share the datasets upon request
Due to the information available on this data we have done a basic analysis on the data available, hereafter referred to as *LS*.

**Suicide Note** The data for this corpus has mainly been taken from Schoene and Dethlefs (2016), but has been further extended by using notes introduced by The Kernel (2013) and Tumbler (2013). There are total of 161 suicide notes in this corpus, hereafter referred to as *GSN*.

**Depression Notes** We used the data collected by Schoene and Dethlefs (2016) of 142 notes written by people identifying themselves as depressed and lonely, hereafter referred to as *DL*.

### 4 Linguistic Analysis

To gain more insight into the content of the datasets, we performed a linguistic analysis to show differences in structure and contents of notes. For the purpose of this study we used the Linguistic Inquiry and Word Count software (LIWC) (Tausczik and Pennebaker, 2010), which has been developed to analyse textual data for psychological meaning in words. We report the average of all results across each dataset.

**Dimension Analysis** Firstly, we looked at the word count and different dimensions of each dataset (see Table 1). It has previously been argued by Tausczik and Pennebaker (2010) that the words people use can give insight into the emotions, thoughts and motivations of a person, where LIWC dimensions correlate with emotions as well as social relationships. The number of *words per sentences* are highest in DL writers and lowest in last statement writers. Research by Osgood and Walker (1959) has suggested that people in stressful situations break their communication down into shorter units. This may indicate alleviated stress levels in individuals writing notes prior to receiving the death sentence. *Clout* stands for the social status or confidence expressed in a person’s use of language (Pennebaker et al., 2014). This dimension is highest for people writing their last statements, whereas depressed people rank lowest on this. Cohan et al. (2018) have noted that this might be due to the fact that depressed individuals often have a lower socio-economic status. The *Tone* of a note refers to the emotional tone, including both positive and negative emotions, where numbers below 50 indicate a more negative emotional tone (Cohn et al., 2004). The tone for LS is highest overall and the lowest in DL, indicating a more overall negative tone in DL and positive tone in LS.

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tokens per note</td>
<td>110.65</td>
<td>109.72</td>
<td>98.58</td>
</tr>
<tr>
<td>Word per Sent</td>
<td>14.87</td>
<td>11.42</td>
<td>16.88</td>
</tr>
<tr>
<td>Clout</td>
<td>47.73</td>
<td>67.68</td>
<td>19.94</td>
</tr>
<tr>
<td>Tone</td>
<td>54.83</td>
<td>75.43</td>
<td>25.51</td>
</tr>
</tbody>
</table>

Table 1: LIWC Dimension Analysis

**Function Words and Content Words** Next, we looked at selected function words and grammatical differences, which can be split into two categories called *Function Words* (see Table 2), reflecting how humans communicate and *Content words* (see Table 2), demonstrating what humans say (Tausczik and Pennebaker, 2010). Previous studies have found that whilst there is an overall lower amount of function words in a person’s vocabulary, a person uses them more than 50% when communicating. Furthermore it was found that there is a difference in how human brains process function and content words (Miller, 1991). Previous research has found that function words have been connected with indicators of people’s social and psychological worlds (Tausczik and Pennebaker, 2010), where it has been argued that the use of function words require basic skills. The highest amount of function words were used in DL notes, whilst both GSN and LS have a similar amount of function words. Rude et al. (2004) has found that high usage, specifically of first-person singular pronouns (“I”) could indicate higher emotional and/or physical pain as the focus of their attention is towards themselves. Overall Just et al. (2017) has also identified a larger amount of personal pronouns in suicide-related social media content. Previous work by Hancock et al. (2007) has found that people use a higher amount of negations when also expressing negative emotions and used fewer words overall, compared to more positive emotions. This seem to be also true for the number of negations used in this case where amount of *Negations* were also highest in the DL corpus and lowest in the LS corpus, whilst the overall words count was lowest for DL and negative emotions highest. Furthermore, it was found that *Verbs, Adverb and Adjectives* are often used to communicate content, however previous studies have found (Jones and Bennell, 2007; Gregory,
that individuals that commit suicide are under a higher drive and therefore would reference a higher amount of objects (through nouns) rather than using descriptive language such as adjectives and adverbs.

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Function</td>
<td>56.35</td>
<td>56.33</td>
<td>60.20</td>
</tr>
<tr>
<td>Personal pronouns</td>
<td>16.23</td>
<td>20.44</td>
<td>15.19</td>
</tr>
<tr>
<td>I</td>
<td>11.04</td>
<td>12.65</td>
<td>12.8</td>
</tr>
<tr>
<td>Negations</td>
<td>2.71</td>
<td>1.71</td>
<td>4.06</td>
</tr>
<tr>
<td>Verb</td>
<td>19.29</td>
<td>19.58</td>
<td>21.65</td>
</tr>
<tr>
<td>Adjective</td>
<td>4.45</td>
<td>2.58</td>
<td>4.98</td>
</tr>
<tr>
<td>Adverb</td>
<td>4.43</td>
<td>3.14</td>
<td>7.69</td>
</tr>
</tbody>
</table>

Table 2: LIWC Function and Content Words

**Affect Analysis** The analysis of emotions in suicide notes and last statements has often been addressed in research (Schoene and Dethlefs, 2018; Lester and Gunn III, 2013). The number of Affect words is highest in LS notes, whilst they are lowest in DL notes, this could be related to the emotional Tone of a note (see Table 1). This also applies to the amount of Negative emotions as they are highest in DL notes and Positive emotions as these are highest in LS notes. Previous research has analysed the amount of Anger and Sadness in GSN and DL notes and has shown that it is more prevalent in DL note writers as these are typical feelings expressed when people suffer from depression (Schoene and Dethlefs, 2016).

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Affect</td>
<td>9.1</td>
<td>11.58</td>
<td>8.44</td>
</tr>
<tr>
<td>Positive emotion</td>
<td>5.86</td>
<td>8.99</td>
<td>3.15</td>
</tr>
<tr>
<td>Negative emotion</td>
<td>3.15</td>
<td>2.58</td>
<td>5.21</td>
</tr>
<tr>
<td>Anger</td>
<td>0.61</td>
<td>0.65</td>
<td>1.03</td>
</tr>
<tr>
<td>Sadness</td>
<td>1.09</td>
<td>1.08</td>
<td>2.53</td>
</tr>
</tbody>
</table>

Table 3: LIWC Affect Analysis

**Social and Psychological Processes** Social Processes highlights the social relationships of note writers, where it can be seen in Table 4 that the highest amount of social processes can be found in LS and the lowest in DL. Furthermore LS notes tend to speak most about family relations and least about friends, this was also found by Kelly and Foley (2017) who found a low frequency in interpersonal relationships.

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Social processes</td>
<td>12.21</td>
<td>18.19</td>
<td>8.33</td>
</tr>
<tr>
<td>Family</td>
<td>1.17</td>
<td>2.17</td>
<td>0.47</td>
</tr>
<tr>
<td>Friends</td>
<td>0.77</td>
<td>0.38</td>
<td>0.73</td>
</tr>
</tbody>
</table>

Table 4: LIWC Social Processes

The term **Cognitive processes** encompasses a number of different aspects, where we have found that the highest amount of cognitive processes was in DL notes and the lowest in LS notes. Boals and Klein (2005) have found that people who use more cognitive mechanisms to cope with traumatic events such as break ups by using more causal words to organise and explain events and thoughts for themselves. Arguably this explains why there is a lower amount in LS notes as LS writers often have a long time to organise their thoughts, events and feelings whilst waiting for their sentence (Death Penalty Information Centre, 2019). **Insight** encompasses words such as *think* or *consider*, whilst *Cause* encompasses words that express reasoning or causation of events, e.g.: *because* or *hence*. These terms have previously been coined as **cognitive process words** by (Gregory, 1999), who argued that these words are less used in GSN notes as the writer has already finished the decision making process whilst other types of discourse would still try to justify and reason over events and choices. This can also be found in the analysis of our own data, where both GSN and LS notes show similar, but lower frequency of terms in those to categories compared to DL writers. **Tentativeness** refers to the language use that indicates a person is uncertain about a topic and uses a number of filler words. A person who use more tentative words, may have not expressed an event to another person and therefore has not processed an event yet and it has not been formed into a story (Tausczik and Pennebaker, 2010). The amount of tentative words used in DL notes is highest, whilst it is lowest in LS words. This might be due to the fact that LS writers already had to reiterate over certain events multiple times as they go through the process of prosecution.

**Personal Concerns** Personal Concerns refers to the topics most commonly brought up in the different notes, where we note that both Money and Work are most often referred to in GSN notes and lowest in LS notes. This might be due to the the fact that (Mind, 2013) lists these two topics as
some of the most common reasons for a person to commit suicide. Religion is most commonly referenced in LS notes, which confirms previous analysis of such notes (Foley and Kelly, 2018; Kelly and Foley, 2017) and lowest in DL notes. (Just et al., 2017) has found that the topic of Death is commonly referenced in suicide-related communication on Twitter. This was also found in this dataset, where GSN notes most commonly referenced death, whilst DL notes were least likely to reference this topic.

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cognitive Processes</td>
<td>12.19</td>
<td>10.85</td>
<td>16.77</td>
</tr>
<tr>
<td>Insight</td>
<td>2.37</td>
<td>2.3</td>
<td>4.07</td>
</tr>
<tr>
<td>Cause</td>
<td>0.95</td>
<td>0.8</td>
<td>1.94</td>
</tr>
<tr>
<td>Tentativeness</td>
<td>2.57</td>
<td>1.5</td>
<td>3.23</td>
</tr>
</tbody>
</table>

Table 5: LIWC Psychological Processes

Time Orientation and Relativity

Looking at the Time Orientation of a note can give interesting insight into the temporal focus of attention and differences in verb tenses can show psychological distance or to which extend disclosed events have been processed (Tausczik and Pennebaker, 2010). Table 7 shows that the focus of LS letters is primarily in the past whilst GSN and DL letters focus on the present. The high focus on the past in DL notes as well as GSN notes could be, because these notes might draw on their past experiences to express the issues of their current situation or problems. The most frequent use of future tense is in LS letters which could be due to a LS notes writers common focus on afterlife (Heflick, 2005).

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Work</td>
<td>1.24</td>
<td>0.41</td>
<td>0.99</td>
</tr>
<tr>
<td>Money</td>
<td>0.68</td>
<td>0.18</td>
<td>0.31</td>
</tr>
<tr>
<td>Religion</td>
<td>0.82</td>
<td>2.7</td>
<td>0.09</td>
</tr>
<tr>
<td>Death</td>
<td>0.76</td>
<td>0.68</td>
<td>0.64</td>
</tr>
</tbody>
</table>

Table 6: LIWC Personal Concerns

<table>
<thead>
<tr>
<th>Type</th>
<th>GSN</th>
<th>LS</th>
<th>DL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Focus past</td>
<td>3.24</td>
<td>2.86</td>
<td>3.32</td>
</tr>
<tr>
<td>Focus present</td>
<td>14.39</td>
<td>1.43</td>
<td>16.11</td>
</tr>
<tr>
<td>Focus future</td>
<td>2.1</td>
<td>2.27</td>
<td>1.51</td>
</tr>
</tbody>
</table>

Table 7: LIWC Time orientation

Overall it was noted that for most analysis GSN falls between the two extremes of LS and DL.

5 Learning Model

The primary model is the Long-short-term memory (LSTM) given its suitability for language and time-series data (Hochreiter and Schmidhuber, 1997). We feed into the LSTM an input sequence \( x = (x_1, \ldots, x_N) \) of words in a document alongside a label \( y \in \mathbb{Y} \) denoting the class from any of the three datasets. The LSTM learns to map inputs \( x \) to outputs \( y \) via a hidden representation \( h_t \) which can be found recursively from an activation function.

\[
    f(h_{t-1}, x_t),
\]

where \( t \) denotes a time-step. During training, we minimise a loss function, in our case categorical cross-entropy as:

\[
    L(x, y) = -\frac{1}{N} \sum_{n \in N} x_n \log y_n.
\]

LSTMs manage their weight updates through a number of gates that determine the amount of information that should be retained and forgotten at each time step. In particular, we distinguish an ‘input gate’ \( i \) that decides how much new information to add at each time-step, a ‘forget gate’ \( f \) that decides what information not to retain and an ‘output gate’ \( o \) determining the output. More formally, and following the definition by Graves (2013), this leads us to update our hidden state \( h \) as follows (where \( \sigma \) refers to the logistic sigmoid function and \( c \) is the ‘cell state’):

\[
    i_t = \sigma(W_{xi}x_t + W_{hi}h_{t-1} + W_{ci}c_{t-1} + b_i) \quad (3)
\]

\[
    f_t = \sigma(W_{xf}x_t + W_{hf}h_{t-1} + W_{cf}c_{t-1} + b_f) \quad (4)
\]

\[
    c_t = f_t c_{t-1} + i_t \tanh(W_{xc}x_t + W_{hc}h_{t-1} + b_c) \quad (5)
\]

\[
    o_t = \sigma(W_{xo}x_t + W_{ho}h_{t-1} + W_{co}c_t + b_o) \quad (6)
\]

\[
    h_t = o_t \tanh(c_t) \quad (7)
\]

A standard LSTM definition solves some of the problems of vanilla RNNs have (Hochreiter and
Schmidhuber, 1997), but it still has some shortcomings when learning long dependencies. One of them is due to the cell state of an LSTM; the cell state is changed by adding some function of the inputs. When we backpropagate and take the derivative of $c_t$ with respect to $c_t - 1$, the added term would disappear and less information would travel through the layers of a learning model.

For our implementation of a Dilated LSTM, we follow the implementation of recurrent skip connections with exponentially increasing dilations in a multi-layered learning model by Chang et al. (2017). This allows LSTMs to better learn input sequences and their dependencies and therefore temporal and complex data dependencies are learned on different layers. Whilst dilated LSTM alleviates the problem of learning long sequences, it does not contribute to identifying words in a sequence that are more important than others. Therefore we extend this network by (1) an embedding layer and (2) an attention mechanism to further improve the network’s ability. A graph illustration of our learning model can be seen in Figure 2.

**Dilated LSTM with Attention** Each document $D$ contains $i$ sentences $S_i$, where $w_i$ represents the words in each sentence. Firstly, we embed the words to vectors through an embedding matrix $W_e$, which is then used as input into the dilated LSTM.

The most important part of the dilated LSTM is the dilated recurrent skip connection, where $c_t^{(l)}$ is the cell in layer $l$ at time $t$:

$$c_t^{(l)} = f(x_t^{(l)}, c_{t-1}^{(l)}).$$  

(8)

$s^{(l)}$ is the skip length; or dilation of layer $l$; $x_t^{(l)}$ as the input to layer $l$ at time $t$; and $f(\cdot)$ denotes a LSTM cell; $M$ and $L$ denote dilations at different layers:

$$s^{(l)} = M^{(l-1)}, l = 1, \ldots, L.$$  

(9)

The dilated LSTM alleviates the problem of learning long sequences, however not every word in a sequence has the same meaning or importance.

**Attention layer** The attention mechanism was first introduced by Bahdanau et al. (2015), but has since been used in a number of different tasks including machine translation (Luong et al., 2015), sentence pairs detection (Yin et al., 2016), neural image captioning (Xu et al., 2016) and action recognition (Sharma et al., 2015). Our implementation of the attention mechanism is inspired by Yang et al. (2016), using attention to find words that are most important to the meaning of a sentence at document level. We use the output of the dilated LSTM as direct input into the attention layer, where $O$ denotes the output of final layer $L$ of the Dilated LSTM at time $t+1$.

The attention for each word $w$ in a sentence $s$ is computed as follows, where $u_{it}$ is the hidden representation of the dilated LSTM output, $\alpha_{it}$ represents normalised alpha weights measuring the importance of each word and $S_i$ is the sentence vector:

$$u_{it} = \tanh(O + b_w)$$  

(10)

$$\alpha_{it} = \frac{\exp(u_{it}^T u_w)}{\sum_t \exp(u_{it}^T u_w)}$$  

(11)

$$s_i = \sum_t \alpha_{it} O_t.$$  

(12)

Figure 1: A 2-layer dilated LSTM with Attention.

6 **Experiments and Results**

For our experiments we use all three datasets. Table 8 shows the results for the experiments series. We establish three performance baselines on the datasets by using three different algorithms previously used on similar datasets. Firstly, we use the ZeroR and LMT (Logistic Model Tree) previously used by (Schoene and Dethlefs, 2016). Additionally we chose to benchmark our algorithm also against the originally proposed Bidirectional LSTM with attention proposed by Yang et al. (2016), which was also used on similar existing datasets before (Schoene and Dethlefs, 2018).
Furthermore we benchmark the Dilated Attention LSTM against two other types of recurrent neural networks. We use 200-dimensional word embeddings as input into each network and all neural networks share the same hyper-parameters, where learning rate = 0.001, batch size = 128, dropout = 0.5, hidden size = 150 units and the Adam optimizer is used. For our proposed model - the Dilated LSTM with Attention - we establish the number of dilations empirically. There are 2 dilated layers with exponentially increasing dilations starting at 1. Due to the size of the dataset we have split the data into 70% training, 15% validation and 15% test data. We report results based on the test accuracy of the prediction results. It can be seen in Table 8 that the dilated LSTM with an attention layer outperforms the BiLSTM with Attention by 5.07%. Furthermore it was found that both the LMT and a vanilla bi-directional LSTM outperform a standard LSTM on this task. Previous results on similar tasks have yielded an accuracy of 69.41% using BiLSTM with Attention (Schoene and Dethlefs, 2018) and 86% using a LMT (Schoene and Dethlefs, 2016).

7 Evaluation

In order to evaluate the DLSTM with attention we look in more detail at the predicted labels and visualise examples of each note to show which features are assigned the highest attention weights.

7.1 Label Evaluation

In Figure 2 we show the confusion matrix over the DLSTM with attention. It can be seen that LS notes are most often correctly predicted and DL notes are least likely to be accurately predicted.

The same applies to results of the main competing model (Bi-directional LSTM with Attention), Figure 3 shows that this model still misclassifies LS notes with DL notes.

7.2 Visualisation of attention weights

In order to see which features are most important to accurate classification we visualise examples from the test set of each dataset, where Figures 4, 5 and 6 show the visualisation of attention weights in the GSN, LS and DL datasets respectively. Furthermore, we also show three examples of the test data with typical errors the learning model makes in Figures 7, 8 and 9. Words highlighted in darker shades have a higher attention weight.

The most important words highlighted in a last statement note (see Figure 4) are personal pronouns as well as an apology and expression of love towards friends and family members. This corresponds with the higher amount of personal pronouns, positive emotions and references to Family in LS notes compared to GSN and DL notes. Furthermore it can be seen that there is a low amount of cognitive process words and more action verbs such as killing or hurt, which could confirm that inmates have had more time to process events and thoughts and don’t need cognitive words as a coping mechanism anymore (Boals and Klein, 2005).

Figure 5 shows a GSN note, where the most important words are also pronouns, references to family, requests for forgiveness and endearments. Previous research has shown that forgiveness is an important feature as well as the giving instructions such as help or phrases like do not follow are key
Table 8: Test accuracy and F1-score of different learning models in %

<table>
<thead>
<tr>
<th>Model</th>
<th>Test Accuracy</th>
<th>Aver. Precision</th>
<th>Aver. Recall</th>
<th>Aver. F1-score</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZeroR</td>
<td>42.85</td>
<td>0.43</td>
<td>0.41</td>
<td>0.42</td>
</tr>
<tr>
<td>LMT</td>
<td>80.35</td>
<td>0.81</td>
<td>0.79</td>
<td>0.80</td>
</tr>
<tr>
<td>LSTM</td>
<td>62.16</td>
<td>0.63</td>
<td>0.61</td>
<td>0.62</td>
</tr>
<tr>
<td>BiLSTM</td>
<td>65.82</td>
<td>0.66</td>
<td>0.64</td>
<td>0.65</td>
</tr>
<tr>
<td>BiLSTM with Attention</td>
<td>82.27</td>
<td>0.85</td>
<td>0.83</td>
<td>0.84</td>
</tr>
<tr>
<td>DLSTMAttention</td>
<td>87.34</td>
<td><strong>0.88</strong></td>
<td><strong>0.87</strong></td>
<td><strong>0.87</strong></td>
</tr>
</tbody>
</table>

8 Conclusion

In this paper we have presented a new learning model for classifying long sequences. We have shown that the model outperforms the baseline by 6.99 % and by 5.07 % a competitor model. Furthermore we have provided an analysis of the linguistic features on three datasets, which we have later compared in a qualitative evaluation by visualising the attention weights on examples of each dataset. We have shown that the neural network pays attention to similar linguistic features as provided by LIWC and found in human evaluated related research.
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Writing habits and telltale neighbors: analyzing clinical concept usage patterns with sublanguage embeddings

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Abstract

Natural language processing techniques are being applied to increasingly diverse types of electronic health records, and can benefit from in-depth understanding of the distinguishing characteristics of medical document types. We present a method for characterizing the usage patterns of clinical concepts among different document types, in order to capture semantic differences beyond the lexical level. By training concept embeddings on clinical documents of different types and measuring the differences in their nearest neighborhood structures, we are able to measure divergences in concept usage while correcting for noise in embedding learning. Experiments on the MIMIC-III corpus demonstrate that our approach captures clinically-relevant differences in concept usage and provides an intuitive way to explore semantic characteristics of clinical document collections.

1 Introduction

Sublanguage analysis has played a pivotal role in natural language processing of health data, from highlighting the clear linguistic differences between biomedical literature and clinical text (Friedman et al., 2002) to supporting adaptation to multiple languages (Laippala et al., 2009). Recent studies of clinical sublanguage have extended sublanguage study to the document type level, in order to improve our understanding of the syntactic and lexical differences between highly distinct document types used in modern EHR systems (Feldman et al., 2016; Grön et al., 2019).

However, one key axis of sublanguage characterization that has not yet been explored is how domain-specific clinical concepts differ in their usage patterns among different document types. Established biomedical concepts may have multiple, often non-compositional surface forms (e.g., “ALS” and “Lou Gehrig’s disease”), making them difficult to analyze using lexical occurrence alone. Understanding how these concepts differ between document types can not only augment recent methods for sublanguage-based text categorization (Feldman et al., 2016), but also inform the perennial challenge of medical concept normalization (Luo et al., 2019): “depression” is much easier to disambiguate if its occurrence is known to be in a social work note or an abdominal exam.

Inspired by recent technological advances in modeling diachronic language change (Hamilton et al., 2016; Vashisth et al., 2019), we characterize concept usage differences within clinical sublanguages using nearest neighborhood structures of clinical concept embeddings. We show that overlap in nearest neighborhoods can reliably distinguish between document types while controlling for noise in the embedding process. Qualitative analysis of these nearest neighborhoods demonstrates that these distinctions are semantically relevant, highlighting sublanguage-sensitive relationships between specific concepts and between concepts and related surface forms. Our findings suggest that the structure of concept embedding spaces not only captures domain-specific semantic relationships, but can also identify a “fingerprint” of concept usage patterns within a clinical document type to inform language understanding.

2 Related Work

Sublanguage analysis historically focused on describing the characteristic grammatical structures of a particular domain (Friedman, 1986; Grishman, 2001; Friedman et al., 2002). As methods for automated analysis of large-scale data sets have improved, more studies have investigated lexical and semantic characteristics, such as usage patterns of different verbs and semantic categories.
Table 1: Document type subcorpora in MIMIC-III. Tokenization was performed with SpaCy; Matches and Concepts refer to number of terminology string match instances and number of unique concepts embedded, respectively, using SNOMED-CT and LOINC vocabularies from UMLS 2017AB release. The number of high-confidence concepts identified for each document type is given with their mean consistency.

(Denecke, 2014), as well as more structural information such as document section patterns and syntactic features (Zeng et al., 2011; Temnikova et al., 2014). The use of terminologies to assess conceptual features of a sublanguage corpus was proposed by Walker and Amsler (1986), and Drouin (2004); Grön et al. (2019) used sublanguage features to expand existing terminologies, but large-scale characterization of concept usage in sublanguage has remained a challenging question.

Word embedding techniques have been utilized to describe diachronic language change in a number of recent studies, from evaluating broad changes over decades (Hamilton et al., 2016; Vashisth et al., 2019) to detecting fine-grained shifts in conceptualizations of psychological concepts (Vylomova et al., 2019). Embedding techniques have also been used as a mirror to analyze social biases in language data (Garg et al., 2018). Similar to our work, Ye and Fabbri (2018) investigate document type-specific embeddings from clinical data as a tool for medical language analysis. However, our approach has two significant differences: Ye and Fabbri (2018) used word embeddings only, while we utilize concept embeddings to capture concepts across multiple surface forms; more importantly, their work investigated multiple document types as a way to control for specific usage patterns within sublanguages in order to capture more general term similarity patterns, while our study aims to capture these sublanguage-specific usage patterns in order to analyze the representative differences in language use between different expert communities.

3 Data and preprocessing

We use free text notes from the MIMIC-III critical care database (Johnson et al., 2016) for our analysis. This includes approximately 2 million text records from hospital admissions of almost 50 thousand patients to the critical care units of Beth Israel Deaconess Medical Center over a 12-year period. Each document belongs to one of 15 document types, listed in Table 1.

As sentence segmentation of clinical text is often optimized for specific document types (Griffis et al., 2016), we segmented our documents at linebreaks and tokenized using SpaCy (version 2.1.6; Honnibal and Montani 2017). All tokens were lowercased, but punctuation and deidentifier strings were retained, and no stopwords were removed.

4 Experiments

Methods for learning clinical concept representations have proliferated in recent years (Choi et al., 2016; Mencia et al., 2016; Phan et al., 2019), but often require annotations in forms such as billing codes or disambiguated concept mentions. These annotations may be supplied by human experts such as medical coders, or by adapting medical NLP tools such as MetaMap (Aronson and Lang, 2010) or cTAKES (Savova et al., 2010) to perform concept recognition (De Vine et al., 2014).

For investigating potentially divergent usage patterns of clinical concepts, these strategies face
serious limitations: the full diversity of MIMIC data has not been annotated for concept identifiers, and the statistical biases of trained NLP tools may suppress underlying differences in automatically-recognized concepts. We therefore take a distant supervision approach, using JET (Newman-Griffis et al., 2018). JET uses a sliding context window to jointly train embedding models for words, surface forms, and concepts, using a log-bilinear objective with negative sampling and shared embeddings for context words. It leverages known surface forms from a terminology as a source of distant supervision: each occurrence of any string in the terminology is treated as a weighted training instance for each of the concepts that string can represent. As terminologies are typically many-to-many maps between surface forms and concepts, this generally leads to a unique set of contexts being used to train the embedding of each concept, though any individual context window may be used as a sample for training multiple concepts. We constrain the scope of our analysis to only concepts and strings from SNOMED-CT and LOINC,1 two popular high-coverage clinical vocabularies.

4.1 Identifying concepts for comparison

For each document type, we concatenate all of its documents (maintaining linebreaks), identify all occurrences of SNOMED-CT and LOINC strings in each line, and use these occurrences to train word, term, and concept embeddings with JET. Due to the size of our subcorpora, we used a window size of 5, minimum frequency of 5, embedding dimensionality of 100, initial learning rate of 0.05, and 10 iterations over each corpus.

Prior research has noted instability of nearest neighborhoods in multiple embedding methods (Wendlandt et al., 2018). We therefore train 10 sets of embeddings from each of our subcorpora, each using the same hyperparameter settings but a different random seed. We then use all 10 replicates from each subcorpus in our analyses, in order to control for variation in nearest neighborhoods introduced by random initialization and negative sampling. To evaluate the baseline reliability of concept embedding neighborhoods from each subcorpus, we calculated per-concept consistency by measuring, over all pairs of embedding sets within the 10 replicates, the average set membership overlap between the top 5 nearest neighbors by cosine similarity for each concept embedding.

As shown in Figure 1a, these consistency scores vary widely both within and between document types, with some document types producing no concept embeddings with consistency over 40%. Interestingly, as illustrated in Figure 1b, there is no linear relationship between log corpus size and mean concept consistency ($R^2 \approx 0.011$), suggesting that low consistency is not solely due to limited training data.

To mitigate concerns about the reliability of embeddings for comparison, a set of high-confidence concepts is identified for each document type by...
4.2 Cross-corpus analysis

Our key question is what concept embeddings reveal about clinical concept usage between document types. To maintain a sufficient sample size, we restrict our comparison to the 7 document types with at least 50 high confidence concepts: Case Management, Discharge Summary, Echo, Nursing/Other, Nutrition, Physician, and Radiology. Physician, ECG, and Nursing were also used by Feldman et al. (2016) for their lexicosyn-

3 We found in our analysis that most concept consistency numbers clustered roughly bimodally, between 0-30% or 60-90%; this is reflected at a coarse level in the overall distributions in Figure 1a. Varying the threshold outside of these ranges did not have a significant impact on the number of concepts retained; the 50% threshold was chosen for simplicity. With larger corpora, yielding higher concept coverage, a higher threshold could be chosen for a stricter analysis.

tactic analysis, although they combined Nursing documents (longer narratives) and Nursing/Other (which tend to be much shorter) into a single set, while we retain the distinction. Interestingly, the fourth type they analyzed, ECG, produced only 14 high-confidence concepts in our analysis, suggesting high semantic variability despite the large number of documents.

As learned concept sets differ between document types, the first step for comparing a document type pair is to identify the set of concepts embedded for both. For reference type A and comparison type B, we identify high-confidence concepts from A that are also present in B, and calculate four distributions using this shared set:

Reference consistency: self-consistency across each of the shared concepts, using only other shared concepts to identify nearest neighborhoods in embeddings for the reference set.

Comparison consistency: self-consistency of each shared concept in embeddings for the comparison document type, again using only shared concepts for neighbors. As the shared set is based on high-confidence concepts from the reference
set, this is not symmetric with reference consistency (as the high-confidence sets may differ).

**Cross-type consistency**: average consistency for each shared concept calculated over all pairs of replicates (i.e., comparing the nearest neighbors of all 10 reference embedding sets to the nearest neighbors in all 10 comparison embedding sets).

**Consistency deltas**: the difference, for each shared concept, between its reference self-consistency and its cross-type consistency. This provides a direct evaluation of how distinct the concept usage is between two document types, where a high delta indicates highly distinct usage.

Mean values for these distributions are provided for each pair of our 7 document types in Figure 2. Comparing Figures 2b and 2c, it is clear that high-confidence concepts for one document type are typically not high-confidence for another. Most document type pairs show fairly strong divergence, with deltas ranging from 30-60%. **Physician** notes have comparatively high cross-set consistency of around 20% for their high-confidence concepts, likely reflecting the all-purpose nature of these documents, which include patient history, medications, vitals, and detailed examination notes. Interestingly, **Case Management** and **Nutrition** are starkly divergent from other document types, with near-zero cross-set consistency and comparatively high self-consistency of over 70% in the compared concept sets, despite a relatively high overlap between their high-confidence sets and concepts learned for other document types.

In order to control for the low overlap between high-confidence sets in different document types, we also re-run our consistency analysis restricted to only concepts that are high-confidence in both the reference and comparison sets. As shown in Figure 3, this yields considerably smaller concept

---

**Figure 3**: Concept neighborhood consistency statistics, restricted to concepts that are high-confidence in both reference and comparison sets. In this case, reference self-consistency and target self-consistency are symmetric, so only reference self-consistency is presented in Figure 3b.

![Figure 3](image-url)
<table>
<thead>
<tr>
<th>Query</th>
<th>Discharge Summary</th>
<th>Nursing/Other</th>
<th>Radiology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diabetes Mellitus (C0011849)</td>
<td>Gestational Diabetes (C0085207)</td>
<td>Poorly controlled (C3853134)</td>
<td></td>
</tr>
<tr>
<td>Type 2 (C0441730)</td>
<td>A2 immunologic symbol (C1443036)</td>
<td>Insulin (C0021641)</td>
<td>Diabetes Mellitus, Insulin-Dependent (C0011854)</td>
</tr>
<tr>
<td>Type 1 (C0441729)</td>
<td>Diabetes Mellitus, Insulin-Dependent (C0011854)</td>
<td>Diabetes Mellitus, Non-Insulin-Dependent (C0011860)</td>
<td></td>
</tr>
<tr>
<td>Gestational Diabetes</td>
<td>Factor V (C0015498)</td>
<td>Stage level 5 (C0441777)</td>
<td></td>
</tr>
<tr>
<td>(C0085207)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diabetes Mellitus, Insulin-</td>
<td>A1 immunologic symbol (C1443035)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dependent (C0011854)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2: 5 nearest neighbor concepts to Diabetes Mellitus and Mental state from 3 high-confidence document types, averaging cosine similarities across all replicate embedding sets within each document type. †The two nearest neighbors to Mental state for all three document types were two LOINC codes using the same “mental status” string; they are omitted here for brevity.

4.3 Qualitative neighborhood analysis

Analysis of neighborhood consistency enables measuring divergence in the contextual usage patterns of clinical concepts; however, this divergence could be due to spurious or semantically uninformative correlations instead of clinically-relevant distinctions in concept similarities. To confirm that our methodology captures informative distinctions in concept usage, we qualitatively review example neighborhoods. To mitigate variability of nearest neighborhoods in embedding spaces, we identify a concept’s qualitative nearest neighbors for a given document type by calculating its pairwise cosine distance vectors for all 10 replicates in that document type and taking the \( k \) neighbors with lowest average distance.

As with our consistency analyses, we focus on the neighborhoods of high-confidence concepts, although we do not filter the neighborhoods themselves. Of all high-confidence concepts identified in our embeddings, only two were high-confidence in 5 different document types, and these were highly generic concepts: Interventional procedure (C0184661) and a corresponding LOINC code (C0945766). Seven concepts were high-confidence for 4 document types; of these, two were generic procedure concepts, two were concepts for the broad gastrointestinal category, and three were versions of body weight. For a diversity of concepts, we therefore turned to the 75 concepts that were high-confidence within 3 document types. We reviewed each of these concepts, and describe our findings for three of the most broadly clinically-relevant below.

**Diabetes Mellitus (C0011849)**  
*Diabetes Mellitus* (search strings: “diabetes mellitus” and “diabetes mellitus dm”) was high-confidence in Discharge Summary, Nursing/Other, and Radiology document types; Table 2 gives the top 5 neighbors from each type. These neighbors are semantically consistent across document types: more specific diabetes-related concepts, related biological fac-
tors; continuing down the nearest neighbors list yields related symptoms and comorbidities such as Irritable Bowel Syndrome (C0022104) and Gastroesophageal reflux disease (C0017168).

**Memory loss (C0751295)** Memory loss (search string: “memory loss”) was also high-confidence in Discharge Summary, Nursing/Other, and Radiology documents. For brevity, its nearest neighbors are omitted from Table 2, as there is little variation among the top 5. However, the next neighbors (at only slightly greater cosine distance) vary considerably across document types, while remaining highly consistent within each individual type. In Discharge Summary, more high-level concepts related to overall function emerge, such as Functional status (C0598463), Relationships (C0439849), and Rambling (C4068735). Radiology yields more symptomatically-related neighbors: Aphagia (C0221470) is present in both, but Radiology includes Disorientation (C0233407), Delusions (C0011253), and Gait, Unsteady (C0231686). Finally, Nursing/Other finds concepts more related to daily life, such as Cigars (C0678446) and Multifocals (C3843228), though at a greater cosine distance than the other document types (Figure 4).

**Mental state (C0278060)** Mental state (search strings: “mental status”, “mental state”) was high-confidence in Discharge Summary, Echo, and Radiology, and highlighted an unexpected consequence of relying on the Distributional Hypothesis (Harris, 1954) for semantic characterization in sublanguage-specific corpora. The top 5 nearest neighbors (excluding two trivial LOINC codes for the same concept, also using the “mental status” search string) are given in Table 2. In Discharge Summary documents, “mental status” is typically referred to in detailed patient narratives, medication lists, and the like, and this yields semantically-reasonable nearest neighbors such as Confusion (C0009676) and Coherent (C4068804).

In Echo documents, however, “mental status” occurs most frequently within an “Indication” field of the “PATIENT/TEST INFORMATION” section. Two common patterns emerge in “Indication” texts: references to altered or reduced mental status, or patients who are vegetative and being evaluated for organ donor eligibility. Though “mental status” and “organ donor” do not co-occur, their consistent occurrence in the same contextual structures leads to extremely similar embeddings (see Figure 4). A similar issue occurs in Radiology notes, where the “MEDICAL CONDITION” section includes several instances of elderly patients presenting with either hypothermia or altered mental status; as a result, two hypothermia concepts (C1963170 and C0020672) are in the 10 nearest neighbors to Mental state.

Results from Radiology also highlight one limitation of distant supervision for learning concept embeddings: as the word “state” is polysemous, including a geopolitical entity, geographical concepts such as Mississippi (C0026221) end up with similar embeddings to Mental state. A similar issue occurs in the neighbors for Memory loss; due to string polysemy, the concept CIGAR string - sequence alignment (C4255278) ends up with a similar embedding to Cigars (C0678446).

### 4.4 Nearest surface form embeddings

As JET learns embeddings of concepts and their surface forms jointly in a single vector space, we also analyzed the surface forms embeddings nearest to different concepts. This enabled us both to evaluate the semantic congruence of surface form and concept embeddings, and to further delve into corpus-specific contextual patterns that emerge in the vector space. As with our concept neighborhood analysis, for each of our 10 replicate embeddings in each document type, we calculated the cosine distance vector from each high-confidence concept to all of the term embeddings in the same replicate, and then averaged these distance vectors to identify neighbors robust to embedding noise.

Table 3 presents surface form neighbors identified...
for three high-confidence clinical concepts chosen for clinical relevance and wide usage; these concepts are discussed in the following paragraphs.

**Blood pressure (C0005823)** Blood pressure is high-confidence in *Discharge Summary, Echo,* and *Radiology* documents. It is a key concept that is measured frequently in various settings; intuitively, it is a sufficiently core concept that it should exhibit little variance. Its neighbor surface forms indeed indicate fairly consistent use across the three document types, referencing both related measurements (“heart rate”) and related concepts (“exercise” and “stress”).

**Echocardiogram (C0013798)** Echocardiogram is high-confidence in *Discharge Summary, Echo,* and *ECG* documents. It is a frequent concept used in clinical settings, and its neighbor surface forms reflect its usage. In *Discharge Summary* embeddings, the neighbors are more general surface forms evaluating the findings (“fair”) and relevant history/symptoms that led to the ECG (“exercise”, “stress”). *ECG* embeddings reflect their more technical nature, with surface forms such as “no change” and “abnormal” yielding high similarity.

**Community (C0009462)** Community is a very broad concept and a common word, and is discussed primarily in documents concerned with whole-person health; it is high confidence in *Discharge Summary, Nutrition,* and *Case Management* documents. Each of these document types reflects different usage patterns. The nearest surface forms in *Discharge Summary* embeddings reflect a focus on living conditions, referring to “health center”, “residence”, and “nursing facility”. In *Nutrition* documents, *Community* is discussed primarily in terms of “community-acquired pneumonia”, likely leading to more treatment-oriented neighbor surface forms. Finally, in *Case Management* embeddings, nearby surface forms reflect discussion of specific risk factors or resources (“substance”, “monitoring”) to consider in maintaining the patient’s health and responding to their specific needs (e.g., “hearing”, “speech”).

### 5 Discussion

We have shown that concept embeddings learned from different clinical document type corpora reveal characteristics of how clinical concepts are used in different settings. This suggests that sublanguage-specific embeddings can help profile distinctive usage patterns for text categorization, offering greater specificity than latent topic distributions while not relying on potentially brittle lexical features. In addition, such profiles could also assist with concept normalization by providing more-informed prior probability distributions for medical vocabulary senses that are conditioned on the document or section type that they occur in.

A few limitations of our study are important to note. The embedding method we chose offers flexibility to work with arbitrary corpora and vocabularies, but its use of distant supervision...
introduces some undesirable noise. The example given in Section 4.3 of the similar embeddings learned for the concept cigars and the concept of the CIGAR string in genomic sequence editing illustrates the downside of not leveraging disambiguation techniques to filter out noisy matches. On the other hand, our restriction to strings from SNOMED-CT and LOINC provided a high-quality set of strings intended for clinical use, but also removed many potentially helpful strings from consideration. For example, the UMLS also includes the non-SNOMED/LOINC strings “diabetes” and “diabete mellitus” [sic] for Diabetes Mellitus (C0011849), both of which occur frequently in MIMIC data. Misspellings are also common in clinical data; leveraging well-developed technologies for clinical spelling correction would likely increase the coverage and confidence of sublanguage concept embeddings.

At the same time, the low volume of data analyzed in many document types introduces its own challenges for the learning process. First, though JET can in principle learn embeddings for every concept in a given terminology, this is predicated on the relevant surface forms appearing with sufficient frequency. For a small document sample, many such surface forms that would otherwise be present in a larger sample will either be missing entirely or insufficiently frequent, leading to effectively “missed” concepts. While we are not aware of another concept embedding method compatible with arbitrary unannotated corpora that could help avoid these issues, some strategies could be used to reduce the potential impact of both training noise and low sample sizes. One approach, which might also help improve concept consistency in the document types that yielded few or no high-confidence concepts, would be pretraining a shared base embedding on a large corpus such as PubMed abstracts, which could then be tuned on each document type-specific subcorpus. While this could introduce its own noise in terms of the differences between biomedical literature language and clinical language (Friedman et al., 2002), it could help control for some degree of sampling error and provide a linguistically-motivated initialization for the concept embedding models.

Finally, as we observed with Mental state (C0278060), relying on similarity in contextual patterns can lead to capturing more corpus-specific features with embeddings, as opposed to (sub)language-specific features, as target corpora become smaller and more homogeneous. If a particular concept or set of concepts are always used within the same section of a document, or in the same set phrasing, the “similarity” captured by organization of an embedding space will be more informed by this writing habit endemic to the specific corpus than by clinically-informed semantic patterns that can generalize to other corpora.

6 Conclusion

Analyzing nearest neighborhoods in embedding spaces has become a powerful tool in studying diachronic language change. We have described how the same principles can be applied to sublanguage analysis, and demonstrated that the structure of concept embedding spaces captures distinctive and relevant semantic characteristics of different clinical document types. This offers a valuable tool for sublanguage characterization, and a promising avenue for developing document type “fingerprints” for text categorization and knowledge-based concept normalization.

Acknowledgments

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Extracting UMLS Concepts from Medical Text Using General and Domain-Specific Deep Learning Models

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Abstract

Entity recognition is a critical first step to a number of clinical NLP applications, such as entity linking and relation extraction. We present the first attempt to apply state-of-the-art entity recognition approaches on a newly released dataset, MedMentions. This dataset contains over 4000 biomedical abstracts, annotated for UMLS semantic types. In comparison to existing datasets, MedMentions contains a far greater number of entity types, and thus represents a more challenging but realistic scenario in a real-world setting. We explore a number of relevant dimensions, including the use of contextual versus non-contextual word embeddings, general versus domain-specific unsupervised pre-training, and different deep learning architectures. We contrast our results against the well-known i2b2 2010 entity recognition dataset, and propose a new method to combine general and domain-specific information. While producing a state-of-the-art result for the i2b2 2010 task (F1 = 0.90), our results on MedMentions are significantly lower (F1 = 0.63), suggesting there is still plenty of opportunity for improvement on this new data.

1 Introduction

Entity recognition is a widely-studied task in clinical NLP, and has been the focus of a number of shared tasks, including the i2b2 2010 Shared Task (Uzuner et al., 2011), SemEval 2014 Task 7 (Pradhan et al., 2014), and SemEval 2015 Task 14 (Elhadad et al., 2015). Most previous work has focused on identifying only a few broad types of entities, such as ‘problems’, ‘tests’, and ‘treatments’ in the i2b2 task, and ‘diseases’ in the SemEval tasks. Even when corpora have been annotated for more entity types, as in the GENIA corpus of biological annotations (Ohta et al., 2002), entity recognition tasks typically focus on only a small subset of those (Kim et al., 2004).

However, in some downstream applications it would be useful to identify all terms in a document which exist as concepts in the Unified Medical Language System (UMLS) Metathesaurus (Bodenreider, 2004). This resource comprises a much wider range of biomedical entity types than has previously been considered in clinical entity recognition. Additionally, the UMLS Metathesaurus defines important relationships between entity types (and the lower-level concepts associated with them) via its Semantic Network. Therefore, extracting and labelling entities with respect to their UMLS semantic type, rather than more generic types such as ‘problem’ or ‘test’, can be an important first step in many practical clinical NLP applications.

In this work, we present the first attempt to apply existing clinical entity recognition approaches to a new dataset called MedMentions, which is annotated for all UMLS semantic types (Mohan and Li, 2019). We compare the effectiveness of these approaches with reference to a well-known baseline dataset (i2b2 2010) and analyze the errors that occur when applying such techniques to new problems. On the basis of this error analysis, we propose an adaptation to the BERT architecture to better combine the general and clinical knowledge learned in the pre-training phase, and show that this improves over the more basic approaches.

2 Background

Early successes in clinical/biomedical entity extraction employed approaches such as conditional random fields (Jonnalagadda et al., 2012; Fu and Ananiadou, 2014; Boag et al., 2015) and semi-Markov models (De Bruijn et al., 2011), requiring numerous engineered features. In recent years,
such approaches have been surpassed in performance by deep learning models (Habibi et al., 2017). However, there is a wide range of variation possible within this set of techniques. We briefly discuss some of the parameters of interest in the following sections.

2.1 General vs. Domain-Specific Word Embeddings

Since words may have one dominant meaning in common use, and a different meaning in the medical domain, some work has explored whether word embeddings trained on medical text (e.g. clinical notes, medical journal articles) are more effective in medical entity recognition than those trained on general text sources (e.g. news, Wikipedia).

Roberts (2016) examined the effect of training word embeddings on different corpora for the task of entity extraction on the i2b2 2010 dataset. He compared six corpora: the i2b2 dataset itself, the clinical notes available in the MIMIC database (Johnson et al., 2016), MEDLINE article abstracts, WebMD forum posts, and generic text corpora from Wikipedia and Gigaword. It was found that the best F1 score was obtained by training on the MIMIC corpus, and that combining corpora also led to strong results. Si et al. (2019) also compared training embeddings on MIMIC data versus general domain data, and similarly found that pre-training on the MIMIC data led to better performance on both the i2b2 2010 and SemEval tasks. Alsentzer et al. (2019) trained embeddings only on the discharge summaries from MIMIC, and reported a marginal improvement on the i2b2 2010 task over using the entire MIMIC corpus. Peng et al. (2019) found that pre-training a BERT model on PubMed abstracts led to better performance for biomedical entity extraction, while pre-training on a combination of PubMed abstracts and MIMIC notes led to better performance when extracting entities from patient records.

2.2 Contextual vs. Non-Contextual Word Embeddings

For many years, word embeddings were non-contextual; that is, a word would have the same embedding regardless of the context in which it occurred. Popular word embeddings of this type include GloVe (Pennington et al., 2014), word2vec (Mikolov et al., 2013), and FastText (Bojanowski et al., 2017). Peters et al. (2018) popularized the idea of contextualized word embeddings, which allowed the same word to have a different representation, depending on the context. The character-based ELMo word embeddings introduced by Peters et al. (2018) can be used just as the non-contextual word embeddings were. Sheikhshabbafghi et al. (2018) trained ELMo word embeddings on a dataset of biomedical papers and achieved a new state of the art in gene mention detection on the BioCreative II gene mention shared task. This work showed that domain-specific contextual embeddings improve various types of biomedical named entities recognition. Later in 2018, BERT embeddings were also introduced (Devlin et al., 2019). The BERT architecture improved over ELMo by using a different training objective to better take into account both left and right contexts of a word, and made it possible to make use of the entire pre-trained network in the downstream task, rather than simply extracting the embedding vectors.

Si et al. (2019) compared word2vec, GloVe, FastText, ELMo, and BERT embeddings on the i2b2 2010 dataset. When using the pre-trained vectors (trained on general-domain corpora), BERT-large performed the best and word2vec performed the worst, but there was no clear advantage to the contextualized embeddings (e.g. GloVe performed better than ELMo). When the embeddings were pre-trained on MIMIC data, the contextualized embeddings did perform appreciably better than the non-contextualized embeddings.

2.3 Classifier Architecture

Much of the recent work on medical entity extraction has made use of the Long Short-Term Memory (LSTM) architecture (Hochreiter and Schmidhuber, 1997), with some variations and modifications: (1) most work uses a bi-directional LSTM (bi-LSTM), so the prediction for any word in the sequence can take into account information from both the left and right contexts, (2) some work additionally feeds the output of the bi-LSTM layer into a CRF classifier (Huang et al., 2015a; Chalapathy et al., 2016a; Lample et al., 2016; Habibi et al., 2017; Tourille et al., 2018), to predict the most likely sequence of labels, rather than just the most likely label for each word independently, and (3) some models incorporate additional information (e.g. character embeddings, or traditionally engineered features) at various points in the model (Unanue et al., 2017).
In contrast, the BERT model makes use of the Transformer architecture, an attention-based method for sequence-to-sequence modelling (Vaswani et al., 2017). Once the model has been pre-trained, in the entity extraction stage it is only necessary to add a simple classification layer on the output. However, others have also experimented with feeding the output of the BERT model to a bi-LSTM (Si et al., 2019).

3 Methods

3.1 Data

We consider three datasets in this study: the i2b2 2010 dataset, the ‘full’ MedMentions dataset, and the ‘st21pv’ MedMentions dataset. Details are shown in Table 1.

The i2b2 2010 corpus\(^1\) consists of de-identified clinical notes (discharge summaries), annotated for three entity types: problems, tests, and treatments (Uzuner et al., 2011). The original shared task also included subtasks on assertion classification and relation extraction, but we focus here on entity extraction. In the original data release for the shared task, the training set contained 394 documents; however, the current release of the dataset contains only 170 training documents. Therefore, it is unfortunately not possible to directly compare results across the two versions of the dataset. However, a majority of the recent works are using the current release of the dataset (Zhu et al., 2018; Bhatia et al., 2019; Chalapathy et al., 2016b).

The MedMentions corpus was released earlier this year\(^2\), and contains 4,392 abstracts from PubMed, annotated for concepts and semantic types from UMLS (2017AA release). UMLS concepts are fine-grained biomedical terms, with approximately 3.2 million unique concepts contained in the metathesaurus (Mohan and Li, 2019). Each concept is linked to a higher-level semantic type, such as ‘Disease or syndrome’, ‘Cell component’, or ‘Clinical attribute’. In this work we focus on identifying the semantic type for each extracted text span, leaving the concept linking/normalization for future work. The creators of the dataset have defined an official 60%-20%-20% partitioning of the corpus into training, development, and test sets.

There are 127 semantic types in UMLS. Of these, there is only one (‘Carbohydrate sequence’) which never appears in the full MedMentions dataset. Approximately 8% of the concepts in UMLS can be linked to more than one semantic type (Mohan and Li, 2019); in such cases the dataset contains a comma-separated list of all these type IDs corresponding to alphabetical order of semantic types. Where a text span has been labelled with more than one label, we select only the first one. As a result of this, there is one other type (‘Enzyme’) which appears in MedMentions, but only doubly-labelled with ‘Amino acid, peptide, or protein’, and thus does not occur in our singly-labelled training or test data. Finally, there is an extra class (‘UnknownType’), for a total of 126 semantic types or classes in the ‘full’ training data. Of these, there are three (‘Amphibian’, ‘Drug delivery device’, and ‘Vitamin’) which never occur in the test data.

The full MedMentions dataset suffers from high class imbalance (e.g. there are 31,485 mentions for the semantic type ‘Qualitative concept’ and only two mentions for ‘Fully formed anatomical structure’). Furthermore, many of the semantic types are not particularly useful in downstream clinical NLP tasks, either due to being too broad or too specialized. As a result, the creators of the MedMentions dataset also released an alternate version called ‘st21pv’, which stands for ‘21 semantic types from preferred vocabularies’. The details of how this subset was constructed are given by Mohan and Li (2019), but essentially it contains only 21 semantic types, from specific vocabularies most relevant to biomedical researchers. The raw abstracts, and partitions into training, development, and test sets are the same as in the full dataset – only the set of annotations differs.

The i2b2 and MedMentions datasets differ across a number of important dimensions: the discharge summaries in the i2b2 dataset tend to be hastily written or dictated, with short, incomplete sentences and numerous acronyms and abbreviations, compared to the academic writing style of the MedMentions abstracts. The discharge summaries also tend to be longer, averaging approximately 980 tokens per document, compared to 267 tokens per document in MedMentions. The semantic content of the documents is also different, with the discharge summaries focused exclusively on a single patient and their history, disease progression, treatment, and outcomes, while the MedMentions abstracts typically summarize

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\(^1\)www.i2b2.org/NLP/DataSets

\(^2\)github.com/chanzuckerberg/MedMentions
the results of a scientific study, covering a wide range of biomedical topics. Finally, there are clearly far more entity types in MedMentions than in i2b2, and greater imbalance between the different classes. Therefore, there is no guarantee that methods which perform well on the i2b2 data will also be effective on the MedMentions dataset.

3.2 Entity Recognition

Based on our review of the literature (Section 2) we experimented with two basic architectures: bi-LSTM+CRF (with pre-trained contextual and non-contextual word embeddings as input), and BERT (with a simple linear classification layer and a bi-LSTM classification layer). The details of these classifiers and their training are described below.

3.2.1 Text pre-processing

For the bi-LSTM+CRF models, input text retained casing information, but all numerical tokens were normalized to a single NUM token.

BERT uses WordPiece tokenization (Wu et al., 2016), which breaks longer words into frequently occurring sub-word units to improve handling of rare words and morphological variation. This requires additional pre- and post-processing for the entity recognition task, since the data is labelled at the word level. Following the recommendation of Devlin et al. (2019), we first re-tokenize the text using the WordPiece tokenizer, assign the given label to the first piece of each word, and assign any subsequent pieces a padding label.

In all cases, we convert the text and labels to CoNLL IOB format for input to the classifiers.

3.2.2 bi-LSTM+CRF

We use a standard bi-LSTM+CRF architecture (e.g., see (Huang et al., 2015b)), implemented in PyTorch. The bi-LSTM component has 2 bi-directional layers with hidden size of 1536 nodes. The 100-dimensional character embeddings are learned through the training process and concatenated with pre-trained GloVe embeddings (Pennington et al., 2014) as proposed by Chalapathy et al. (2016a). We compare the performance of general GloVe embeddings, trained on Wikipedia and Gigaword, and clinical GloVe embeddings, trained on the MIMIC-III corpus (Johnson et al., 2016). In both cases the GloVe embeddings have 300 dimensions. For pre-training on MIMIC, we used a minimum frequency cut-off of 5, and a window size of 15.

We also experimented with contextual ELMo embeddings (Peters et al., 2018) and the bi-LSTM+CRF architecture, comparing general ELMo embeddings with clinical ELMo embeddings. The clinical ELMo embeddings were released by Zhu et al. (2018) and trained on Wikipedia pages whose titles are medical concepts in the SNOMED-CT vocabulary, as well as MIMIC-III.

The bi-LSTM+CRF models were trained using the Adam optimizer with a learning rate of 0.001 and a batch size of 32 for 10 epochs.

3.2.3 BERT

The BERT (Bidirectional Encoder Representations from Transformers) model is described by Devlin et al. (2019) and proposes to address some of the limitations observed in LSTM models. In our experiments, we use the BERT-base architecture, which has 12-layers, hidden size 768, and 12 self-attention heads. To perform the entity recognition, we added a linear layer and a softmax layer on top of the last BERT layer to determine the most probable label for each token. While this is the approach taken by Alsentzer et al. (2019), others suggest using a more complex classification model in conjunction with BERT (Si et al., 2019), and so we also experiment with a bi-LSTM layer with input and output size of $4 \times 768$ on top of the concatenation of the last four layers of BERT.

We consider four pre-trained BERT models:

- **BERT-base**

Table 1: Properties of the datasets. For MedMentions, we combine the training and validation sets into ’Train’.

<table>
<thead>
<tr>
<th></th>
<th>i2b2 2010</th>
<th>MedMentions (full)</th>
<th>MedMentions (st21pv)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Train</td>
<td>Test</td>
<td>Train</td>
</tr>
<tr>
<td># entity types</td>
<td>3</td>
<td>3</td>
<td>126</td>
</tr>
<tr>
<td># documents</td>
<td>170</td>
<td>256</td>
<td>3513</td>
</tr>
<tr>
<td># tokens</td>
<td>149,743</td>
<td>267,837</td>
<td>936,247</td>
</tr>
<tr>
<td># entities</td>
<td>16,520</td>
<td>31,161</td>
<td>281,719</td>
</tr>
</tbody>
</table>

5 github.com/google-research/bert
released by Google, pre-trained on Wikipedia and BookCorpus (Devlin et al., 2019).

- **bioBERT (v1.1)** The bioBERT model is initialized with BERT-base, and then further pre-trained on biomedical abstracts from PubMed (Lee et al., 2019).
- **clinicalBERT** The clinicalBERT model is initialized with bioBERT, and then further pre-trained on clinical notes from the MIMIC corpus (Alsentzer et al., 2019).
- **NCBI BERT** The NCBI BERT model is initialized with BERT-base, and then further pre-trained on PubMed abstracts and MIMIC notes (Peng et al., 2019).

In the fine-tuning stage, we generally follow the recommendations in Devlin et al. (2019), and use an Adam optimizer with $\beta_1 = 0.9$, $\beta_2 = 0.999$, L2 weight decay of 0.01, and a dropout probability of 0.1 on all layers. We use a learning rate warmup over the first 10% of steps, and linear decay of the learning rate thereafter. Before training the final models, we conducted a series of hyper-parameter optimization experiments using 10-fold cross-validation on the training set. In this optimization stage we considered combinations of batch sizes in \{16, 32\}, learning rates in \{0.00002, 0.00003, 0.00005, 0.0001\}, and number of training epochs in \{1...10\}. We also determined that the uncased BERT-base model led to marginally better results, and so we use that in our final evaluation (bioBERT and clinicalBERT are cased, while NCBI BERT is uncased). For the BERT-bi-LSTM model, we also experimented with training only the bi-LSTM component and not fine-tuning the pre-trained layers, but found that fine-tuning led to better results in development.

3.3 Evaluation

To evaluate the systems, we use micro-averaged strict precision, recall, and F-score. This means that for any given recognized entity, it is only counted as a true positive if both the span and the label match exactly with the gold standard annotation. Note also that these metrics are computed on the entity-level, not the token level. For example, given the following gold and predicted label sequences:

GOLD: O O B-prob I-prob I-prob
PRED: O O B-prob I-prob I-prob
A token-level evaluation would identify two true positives, but a strict entity-level evaluation identifies zero true positives.

4 Results

Table 2 shows the results of the entity recognition experiments for each model and dataset.

4.1 Effect of Contextual vs. Non-Contextual Word Embeddings

If we first consider the bi-LSTM+CRF results for the i2b2 dataset, we observe that the contextual ELMo embeddings lead to better results than the non-contextual GloVe embeddings, and in both cases, better results are obtained by pre-training the embeddings on domain-specific text. For MedMentions, however, for both versions of the dataset we observe that the general-domain GloVe embeddings outperform the clinical GloVe embeddings, but the clinical ELMo embeddings outperform the general ELMo embeddings. Si et al. (2019) also observed a greater benefit to using contextual embeddings when pre-training on domain-specific corpora. Here, this may be due in part to differences between the training corpora; for example, clinical GloVe was trained only on MIMIC notes, while clinical ELMo was trained on a combination of MIMIC notes and Wikipedia articles about medical concepts, which may be more similar to the biomedical abstracts contained in MedMentions.

The BERT models offer a substantial improvement in F1 over the models based on Glove or ELMo embeddings for each of the three datasets. For the i2b2 dataset, the best results are obtained using clinicalBERT (F1 = 0.88) and NCBI BERT (F1 = 0.89), each of which involved pre-training on clinical notes from MIMIC. This demonstrates the importance of pre-training on documents which are similar in nature to those seen in the labelled dataset. Consistent with this, the bioBERT model (pre-trained on biomedical abstracts) leads to the best result on both MedMentions datasets.

4.2 Effect of Classifier Structure

Finally, comparing the effectiveness of a simple linear model versus a bi-LSTM model as the top BERT layer, we observe that this change makes
## Table 2: Results of entity recognition for each dataset and model.

<table>
<thead>
<tr>
<th>Model</th>
<th>Domain</th>
<th>i2b2 2010</th>
<th>MedMentions (full)</th>
<th>MedMentions (st21pv)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>P</td>
<td>R</td>
<td>F1</td>
</tr>
<tr>
<td>Glove + bi-LSTM+CRF</td>
<td>general</td>
<td>0.81</td>
<td>0.76</td>
<td>0.79</td>
</tr>
<tr>
<td></td>
<td>clinical</td>
<td>0.83</td>
<td>0.77</td>
<td>0.80</td>
</tr>
<tr>
<td>ELMo + bi-LSTM+CRF</td>
<td>general</td>
<td>0.80</td>
<td>0.80</td>
<td>0.80</td>
</tr>
<tr>
<td></td>
<td>clinical</td>
<td>0.86</td>
<td>0.86</td>
<td>0.86</td>
</tr>
<tr>
<td>BERT-base + linear</td>
<td>general</td>
<td>0.85</td>
<td>0.87</td>
<td>0.86</td>
</tr>
<tr>
<td></td>
<td>biomed</td>
<td>0.86</td>
<td>0.88</td>
<td>0.87</td>
</tr>
<tr>
<td>clinicalBERT + linear</td>
<td>biomed + clinical</td>
<td>0.87</td>
<td>0.88</td>
<td>0.88</td>
</tr>
<tr>
<td>NCBI BERT + linear</td>
<td>biomed + clinical</td>
<td>0.88</td>
<td>0.90</td>
<td>0.89</td>
</tr>
<tr>
<td>bioBERT + bi-LSTM</td>
<td>biomed</td>
<td>0.86</td>
<td>0.88</td>
<td>0.87</td>
</tr>
<tr>
<td>NCBI BERT + bi-LSTM</td>
<td>biomed + clinical</td>
<td>0.88</td>
<td>0.90</td>
<td>0.89</td>
</tr>
</tbody>
</table>

### 4.3 MedMentions vs i2b2 dataset

Comparing across datasets, it is clear that performance is worse on the MedMentions data than the i2b2 data. One obvious reason for this is that the MedMentions datasets have many more entity types, or classes, than the i2b2 dataset. This means that the number of examples per class seen in the training data is much lower (in some cases, only a handful), and also that the classes tend to be more easily confused. The ambiguity between classes arises at the level of the training data where, depending on context, the same text span will be associated with different labels (e.g. *neurocognitive function* is sometimes labelled as a ‘biologic function’ and sometimes as a ‘mental process’; *PSA levels* is labelled as both a ‘laboratory procedure’ and a ‘laboratory or test result’; and an even more highly-ambiguous term such as *probe* is variously labelled as a ‘medical device’, ‘indicator reagent or diagnostic aid’, ‘nucleic acid nucleoside or nucleotide’, ‘functional concept’, ‘diagnostic procedure’, ‘research activity’, and ‘chemical viewed functionally’). Thus in MedMentions, the context becomes extremely important, and fine-grained distinctions between entity types must be learned.

### 5 Error Analysis

In the following section, we examine the errors made by the entity recognition systems from two different perspectives: first, we compare generally the types of errors made on the two datasets; then, we consider the role of general versus domain-specific pre-training and examine some of the specific errors that occur in each case.

#### 5.1 Types of Errors by Dataset

Depending on the downstream application of entity recognition, different types of errors may be associated with different costs. For example, if a company is using this model in practice, the cost associated with having human annotators adjust label boundaries may be different from the cost associated with having them search for entities which have been missed altogether. Our evaluation metrics, however, do not reflect the differences among various types of errors. To further investigate the nature of the errors being made by the system, we investigated three specific types of ‘partial errors’. These cases are counted as *false* in calculating the evaluation metrics, but the model actually gets at least part of the information correct:

- **Right span, wrong label:** the text span associated with an entity is correctly identified, but assigned the wrong label.
- **Right label, overlapping span:** the entity is correctly labelled, but the text span associated with the entity is not exactly the same as that indicated in the gold transcripts.
- **Wrong label, overlapping span:** the entity overlaps with one of the gold entities, but is assigned the wrong label.

In addition to these categories, errors can be complete false positives (model extracts an entity which does not overlap at all with any gold entities), or complete false negatives (model completely misses an entity from the gold transcripts).
In the i2b2 dataset, over half of the errors belong to one of the ‘partial’ error types (Figure 1a), with the biggest error category overall being right label, overlapping span. In many cases, the model identifies the main noun phrase associated with the entity, but not adjectives or prepositional phrases that further describe the entity: e.g., *patient feeling weaker than usual*, where the model labels *weaker* as the problem but the gold entity is *weaker than usual*, or *her glucose remained somewhat low*, where the model labels *low* as the problem, but the gold entity is *somewhat low*. One of the reasons that might lead to this kind of error is the inconsistency that exists in the human annotations. The task of identifying the span of entities can be very subjective and there is always some level of disagreement among human annotators. Also, sometimes annotation guidelines are interpreted differently by various annotators. As a result of this inconsistency, in the training stage, the model sees examples that the spans of the same entities has been labeled differently. This type of error may occur more frequently in the i2b2 dataset, due to the difficulty in annotating hurriedly written notes compared to academically written abstracts.

For MedMentions-st21pv, we again observe that roughly half the errors are ‘partial’ errors, but with a sizable increase in errors that involve the wrong label, with either a correct or overlapping span (Figure 1b). As discussed previously, the increase in right span, wrong label is likely due to the higher ambiguity between entity types in this dataset. In many cases, errors of the type wrong label, overlapping span appear to be due to the inability of the annotation scheme to handle overlapping entities. For example, in *co-expression analysis*, where the model labels *co-expression* as a ‘biologic function’, rather than extracting the entire phrase as a ‘research activity’, or in *this region of the brain*, where the model simply labels *brain* as an ‘anatomical structure’ rather than *region of the brain* as a ‘spatial concept’.

### 5.2 General vs Domain Knowledge

We then performed an exploratory error analysis to identify and compare the type of errors made by models using general and domain-specific embeddings. For this analysis, we considered BERT-base and NCBI BERT. BERT-base is pre-trained on Wikipedia and the Google Books database, so it transfers the general knowledge of language to the model. NCBI BERT starts its training from BERT-base, and then continues training on PubMed (biomedical abstracts) and MIMIC (clinical notes). Generally, embeddings like NCBI BERT are assumed to be more effective since they transfer both general and domain-specific knowledge. We analyze the errors of the two models to test this assumption. For simplicity, we concentrate our analysis on the i2b2 dataset, although we observe similar patterns in MedMentions.

NCBI BERT results in a higher overall F1-score than BERT-base on the i2b2 dataset, and there are 2027 entities that are correctly recognized by NCBI BERT and incorrectly recognized by BERT-
base. However, there are also 1209 entities that are correctly recognized by BERT-base but incorrectly recognized by NCBI BERT. Therefore, it is not the case that NCBI BERT encodes the same knowledge as BERT-base, plus more that it has learned from PubMed and MIMIC; rather, the two systems have different strengths and weaknesses.

Qualitatively, we observed that some entities correctly recognized by NCBI BERT and missed by BERT-base involve common words that have a specialized meaning in medicine, for example in the sentence: *Suck, root, gag, grasp, and morrow were normal*. The BERT-base model does not extract any entities, while the NCBI BERT model recognizes *suck, root, gag, and grasp* as standard tests of infant reflexes. NCBI BERT also appears to be better at recognizing specialized acronyms and abbreviations, particularly when there is very little context, as in *Brother died 64 / MI*, where only NCBI BERT recognizes *MI* as a problem (myocardial infarction) in this brief family history, or *No CPR / No defib*, where NCBI BERT correctly labels *CPR and defib* as treatments, while BERT-base mis-labels them as problems.

In cases where BERT-base does better than NCBI BERT, it may be partially due to a better knowledge of well-formed text. We observed several examples where BERT-base appeared to be better at identifying the appropriate qualifiers to attach to a given entity: e.g., in *no interval development of effusion*, BERT-base correctly extracts the entire phrase *internal development of effusion*, while NCBI BERT only extracts *effusion*. Similarly, in *Right ventricular chamber size and free wall motion are normal*, BERT-base extracts *Right ventricular chamber size* as a single entity of the type ‘test’, while NCBI BERT splits it into *Right ventricular* and *size*.

Of course, these observations are purely anecdotal at this point, and will require future work and annotation to fully quantify the nature of the differences between the models. However, given the fact that the two models make different errors, it is at least reasonable to assume that predictions from the two models can be combined in a complementary fashion to improve the overall performance. We explore one possible architecture for doing so in the following section.

### 6 Concatenated Model

As a result of our error analysis, we propose a concatenated BERT model, to better combine the general knowledge from BERT-base and the clinical knowledge from the more specialized BERT models. To build such a model we concatenate the last encoding layer of a domain-specific BERT model with the last encoding layer of the general BERT model and feed this concatenation to a linear or bi-LSTM classification layer. During training we jointly fine-tune both BERT models and the classification layer. We implemented this model with both NCBI BERT and bioBERT models, since they previously led to the optimal results for i2b2 and MedMentions, respectively. NCBI BERT is concatenated with the uncased BERT-base model and bioBERT is concatenated with the cased BERT-base model.

Results for the concatenated models are given in Table 3. For all three datasets, we observe a small improvement over the best performing models in Table 2. The best result for i2b2 is achieved by concatenating the NCBI BERT and BERT-base models, with either a linear or bi-LSTM classifier on top. The resulting F1 score of 0.90 beats the previously reported state-of-the-art of 0.89 on the current release of the dataset with 170 training documents (Zhu et al., 2018). For MedMentions, concatenating bioBERT and BERT-base leads to the best results, with MedMentions-full attaining the best result using a linear classifier and MedMentions-st21pv attaining the best result with the bi-LSTM. To our knowledge, there are no prior results reported on entity (i.e. semantic type) extraction on this dataset.

Regarding the classifier layer, we observe that replacing a linear classifier layer with a bi-LSTM does not improve the results on i2b2 dataset. This is consistent with the results shown in Table 2 and indicates that a simple linear classifier is *enough* to learn the entity recognition task for i2b2 dataset. In the case of the MedMentions dataset, a bi-LSTM classifier improves the F1 score on MedMentions-st21pv but worsens it on MedMentions-full. These results show that there is room for more rigorous investigation about the classifier layer for extracting entities in MedMentions dataset. More complex neural structures with optimized hyperparameters may be needed to improve these results.

Although the improvements that we see by con-
catering the models are relatively small, they are consistent across the three datasets. This suggests that explicitly inputting both general and domain-specific information to the entity recognizer, rather than sequentially pre-training on different domains and hoping that the model ‘remembers’ information from each domain, can be a promising direction for future research.

### 7 Conclusion

We have presented the results of a set of medical entity recognition experiments on a new dataset, MedMentions. We contrasted these results with those obtained on the well-studied i2b2 2010 dataset. We explored a number of relevant dimensions, including the use of various embedding models (contextual versus non-contextual, general versus domain-specific, and LSTM versus attention-based) as well as linear versus bi-LSTM classifier layers. We also proposed a new modification to the previous BERT-based named entity recognition architectures, which allows the classifier to incorporate information from both general and domain-specific BERT embeddings. Our results on i2b2 are state-of-the-art, and our results on MedMentions set a benchmark for future work on this new dataset.

As popular public datasets become more and more studied over time, there is a chance that even if individual researchers follow good train-validate-test protocols, we eventually overfit to the datasets as a community, since there is so much published information available about what works well to improve performance on the test set. One goal of this work was to explore the gap in performance between a well-known clinical entity recognition dataset and a new, unstudied dataset. The same models and training procedures lead to significantly lower performance on the MedMentions dataset, for a variety of reasons: greater number of entity types, more class ambiguity, higher class imbalance, etc. Ultimately, we find that the model which performs best on i2b2 2010 is not the model that performs best on MedMentions, and that results on MedMentions can be improved by pre-training on more similar documents (biomedical abstracts), and by using more complex models (BERT + bi-LSTM rather than BERT + linear). We hope that other researchers will continue to advance the state-of-the-art on this new dataset.

### Acknowledgement

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### References


Ontological attention ensembles for capturing semantic concepts in ICD code prediction from clinical text

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Abstract

We present a semantically interpretable system for automated ICD coding of clinical text documents. Our contribution is an ontological attention mechanism which matches the structure of the ICD ontology, in which shared attention vectors are learned at each level of the hierarchy, and combined into label-dependent ensembles. Analysis of the attention heads shows that shared concepts are learned by the lowest common denominator node. This allows child nodes to focus on the differentiating concepts, leading to efficient learning and memory usage. Visualisation of the multi-level attention on the original text allows explanation of the code predictions according to the semantics of the ICD ontology. On the MIMIC-III dataset we achieve a 2.7% absolute (11% relative) improvement from 0.218 to 0.245 macro-F1 score compared to the previous state of the art across 3,912 codes. Finally, we analyse the labelling inconsistencies arising from different coding practices which limit performance on this task.

1 Introduction

Classification of clinical free-text documents poses some difficult technical challenges. One task of active research is the assignment of diagnostic and procedural International Classification of Diseases (ICD) codes. These codes are assigned retrospectively to hospital admissions based on the medical record, for population disease statistics and for reimbursements for hospitals in countries such as the United States. As manual coding is both time-consuming and error-prone, automation of the coding process is desirable. Coding errors may result in unpaid claims and loss of revenue (Adams et al., 2002).

Automated matching of unstructured text to medical codes is difficult because of the large number of possible codes, the high class imbalance in the data, and the ambiguous language and frequent lack of exposition in clinical text. However, the release of large datasets such as MIMIC-III (Johnson et al., 2016) has paved the way for progress, enabling rule-based systems (Farkas and Szarvas, 2008) and classical machine learning methods such as support vector machines (Suominen et al., 2008), to be superseded by neural network-based approaches (Baumel et al., 2017; Karimi et al., 2017; Shi et al., 2018; Duarte et al., 2018; Rios and Kavuluru, 2018). The most successful reported model on the ICD coding task is a shallow convolutional neural network (CNN) model with label-dependent attention introduced by Mullenbach et al. (2018) and extended by Sadoughi et al. (2018) with multi-view convolution and a modified label regularisation module.

One of the common features of the aforementioned neural network models is the use of attention mechanisms (Vaswani et al., 2017). This mirrors advances in general representation learning. In the text domain, use of multi-headed attention has been core to the development of Transformer-based language models (Devlin et al., 2018; Radford et al., 2019). In the imaging domain, authors have had success with combining attention vectors learned at the global and local levels with Double Attention networks (Chen et al., 2018). In the domain of structured (coded) medical data, Choi et al. (2017) leveraged the ontological structure of the ICD and SNOMED CT coding systems in their Gram model, to combine the attention vectors of a code and its ancestors in order to predict the codes for the next patient visit based on the codes assigned in the previous visit.

Our contributions are:

1. A structured ontological attention ensemble mechanism which provides improved accuracy, efficiency, and interpretability.
2. An analysis of the multi-level attention weights with respect to the text input, which allows us to interpret the code predictions according to the semantics of the ICD ontology.

3. An analysis of the limitations of the MIMIC-III dataset, in particular the labelling inconsistencies arising from variable coding practices between coders and between timepoints.

2 Dataset

We used the MIMIC-III dataset (Johnson et al., 2016) (“Medical Information Mart for Intensive Care”) which comes from the intensive care unit of the Beth Israel Deaconess Medical Center in Boston. We concatenated the hospital discharge summaries associated with each admission to form a single document and combined the corresponding ICD-9 codes. The data was split into training, development, and test patient sets according to the split of Mullenbach et al. (2018) (see Table 1).

<table>
<thead>
<tr>
<th>Dataset</th>
<th># Documents</th>
<th># Unique patients</th>
<th># ICD-9 Codes</th>
<th># Unique ICD-9 codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Training</td>
<td>47,719</td>
<td>36,997</td>
<td>758,212</td>
<td>8,692</td>
</tr>
<tr>
<td>Development</td>
<td>1,631</td>
<td>1,374</td>
<td>28,896</td>
<td>3,012</td>
</tr>
<tr>
<td>Test</td>
<td>3,372</td>
<td>2,755</td>
<td>61,578</td>
<td>4,085</td>
</tr>
<tr>
<td>Total</td>
<td>52,722</td>
<td>41,126</td>
<td>848,686</td>
<td>8,929</td>
</tr>
</tbody>
</table>

Table 1: Distribution of documents and codes in the MIMIC-III dataset.

3 Methods

We formulate the problem as a multi-label binary classification task, for which each hospital discharge summary is labelled with the presence or absence of the complete set of ICD-9 codes for the associated admission. Our model is a CNN similar to those of (Mullenbach et al., 2018; Sadoughi et al., 2018). Inspired by the graph-based attention model of (Choi et al., 2017), we propose a hierarchical attention mechanism (mirroring the ICD ontology) which yields a multi-level, label-dependent ensemble of attention vectors for predicting each code. Our architecture is shown in Figure 1 and described below.

3.1 Embedding

Documents were pre-processed by lower-casing the text and removing punctuation, followed by tokenisation during which purely numeric tokens were discarded. We used a maximum input length of 4500 tokens and truncated any documents longer than this (260 training, 16 development, and 22 test). Tokens were then embedded with a 100-dimensional word2vec model. For each document, token embeddings were concatenated to give a $100 \times N$ document embedding matrix $D$, where $N$ is the document length.

We pre-trained the word2vec model on the training set using continuous bag-of-words (CBOW) (Mikolov et al., 2013). The vocabulary comprises tokens which occur in at least 3 documents (51,847 tokens). The embedding model was fine-tuned (not frozen) during subsequent supervised training of the complete model.

3.2 Convolutional module

The first part of the network proper consists of a multi-view convolutional module, as introduced by Sadoughi et al. (2018). Multiple one-dimensional convolutional kernels of varying size with stride $= 1$ and weights $W$ are applied in parallel to the document embedding matrix $D$ along the $N$ dimension. The outputs of these kernels are padded at each end to match the input length $N$. This yields outputs of size $C \times M \times N$ where $C$ is the number of kernel sizes (“views”), $M$ is the number of filter maps per view, and $N$ is the length of the document. The outputs are max-pooled in the $C$ dimension i.e., across each set of views, to yield a matrix $E$ of dimensions $M \times N$:

$$E = \text{tanh} ( \max_{C=0,3} W_C * D)$$

(1)

Optimal values were $C = 4$ filters of lengths $\{6, 8, 10, 12\}$ with $M = 256$ filter maps each.

3.3 Prediction via label-dependent attention

Label-specific attention vectors are employed to collapse the variable-length $E$ document representations down to fixed-length representations. For each label $l$, given the matrix $E$ as input, a token-wise linear layer $u_l$ is trained to generate a vector of length $N$. This is normalised with a softmax operation, resulting in an attention vector $a_l$:

$$a_l = \text{softmax}(E^T u_l)$$

(2)
The attention vector is then multiplied with the matrix $E$ which yields a vector $v_l$ of length $M$, a document representation specific to a label:

$$v_l = a_l E \quad (3)$$

If multiple linear layers $u_{l,0}, u_{l,1}, \ldots$ are trained for each label at this stage, multiple attention vectors (or “heads”) will be generated. Thus, multiple document representations $v_l$ could be made available, each of length $M$, and concatenated together to form a longer label-specific representation for the document. We experimented with multiple attention vectors and found two vectors per label to be optimal. To make a prediction of the probability of each label, $P(l)$, there is a final dense binary classification layer with sigmoid activation. This is shown for two attention vectors:

$$P(l) = \sigma(W_l[v_{l,0}; v_{l,1}] + \beta_l) \quad (4)$$

### 3.4 Prediction via label-dependent ontological attention ensembles

The ICD-9 codes are defined as an ontology, from more general categories down to more specific descriptions of diagnosis and procedure. Rather than simply training two attention heads per code as shown in Section 3.3, we propose to exploit the ontological structure to train shared attention heads between codes on the same branch of the tree, thus pooling information across labels which share ancestry. In this work, we use two levels of ancestry, where the first level corresponds to the pre-floating-point portion of the code. For instance, for the code 425.11 Hypertrophic obstructive cardiomyopathy, the first-degree ancestor is 425 Cardiomyopathy and the second-degree ancestor is 420-429 Other forms of heart disease (the chapter in which the parent occurs). This is illustrated in Figure 2.

![Figure 2: Illustration of inheritance of the linear layers $u_l$. This yields label-specific ontological attention ensembles of the attention heads $a_l$ and subsequently the document representations $v_l$.](image)

For the entire set of 8929 labels, we identi-
fied 1167 first-degree ancestors and 179 second-degree ancestors. Compared to two attention vectors per code, this reduces the parameter space and memory requirements from 17,858 attention heads (8929 x 2) to 10,275 attention heads (8929 + 1167 + 179) as well as increasing the number of training samples for each attention head.

The label prediction for each code is now derived from the concatenated child \((c)\), parent \((p)\) and grandparent \((gp)\) document representations:

\[
P(l_{\text{child}}) = \sigma(W_l[c; v_l;p; v_{l, gp}] + \beta_l) \tag{5}\]

In order to facilitate learning of multiple attention heads, we employ deep supervision using the ancestral labels, adding auxiliary outputs for predicting the parent and grandparent nodes:

\[
P(l_{\text{parent}}) = \sigma(W_l[p; v_{l, gp}] + \beta_p) \tag{6}\]

\[
P(l_{\text{grandparent}}) = \sigma(W_l[gp; v_{l, gp}] + \beta_{gp}) \tag{7}\]

### 3.5 Training process

We trained our model with weighted binary cross entropy loss using the Adam optimiser (Kingma and Ba, 2014) with learning rate 0.0005.

**Stratified shuffling:** The network accepts input of any length but all instances within a single batch need to be padded to the same length. To minimise the amount of padding, we used length-stratified shuffling between epochs. For this, documents were grouped by length and shuffled only within these groups; groups were themselves then shuffled before batch selection started.

**Dampened class weighting:** We employed the standard practice of loss weighting to prevent the imbalanced dataset from affecting performance on rare classes. We used a softer alternative to empirical class re-weighting, by taking the inverse frequencies of positive (label=1) and negative (label=0) examples for each code \(c\), and adding a damping factor \(\alpha\). In the equations below, \(n_{\text{label}_c=1}\) stands for the number of positive examples for the ICD code \(c\), and \(n\) stands for the total number of documents in the dataset.

\[
\omega_{(c,1)} = \left( \frac{n}{n_{\text{label}_c=1}} \right)^\alpha, \quad \text{for } n_{\text{label}_c=1} > 5 \tag{9}
\]

\[
\omega_{(c,0)} = \left( \frac{n}{n_{\text{label}_c=0}} \right)^\alpha, \quad \text{otherwise}
\]

**Deep supervision:** The loss function was weighted in favour of child codes, with progressively less weight given to the codes at higher levels in the ICD ontology. A weighting of 1 was used for the child code loss, a weighting \(w_h\) for the parent code auxiliary loss, and \(w_{gp}^2\) for the grandparent code auxiliary loss, i.e.,

\[
\text{Loss} = L_c + w_h L_p + w_{gp}^2 L_{gp} \tag{10}
\]

Optimal values were \(\alpha = 0.25\) and \(w_h = 0.1\).

### 3.6 Implementation and hyperparameters

The word2vec embedding was implemented with Gensim (Rehůrek and Sojka, 2010) and the ICD coding model was implemented with PyTorch (Paszke et al., 2017). Experiments were run on Nvidia V100 16GB GPUs. Hyperparameter values were selected by maximising the development set macro-F1 score for codes with more than 5 training examples.

### 4 Experiments

#### 4.1 Results

In our evaluation, we focus on performance across all codes and hence we prioritise macro-averaged metrics, in particular macro-averaged precision, recall, and F1 score. Micro-averaged F1 score and Precision at \(k\) \((P@K)\) are also reported in order to directly benchmark performance against previously reported metrics. All reported numbers are the average of 5 runs, starting from different random network initialisations.

We compare our model to two previous state-of-the-art models: Mullenbach et al. (2018), and Sadoughi et al. (2018) (published only on arXiv). We trained these models with the hyperparameter values quoted in the respective publications, and used the same early stopping criteria as for our model. Both Mullenbach et al. and Sadoughi et al. use label regularisation modules, at the output and at the attention layer respectively. In line with their published results, we found that only the method of Sadoughi et al. gave an improvement and thus it
Table 2: Benchmark results for the models trained with \( F_1^{\text{macro}} \) stopping criterion.

<table>
<thead>
<tr>
<th>Method</th>
<th>( R_{\text{macro}} )</th>
<th>( P_{\text{macro}} )</th>
<th>( F_1^{\text{macro}} )</th>
<th>( F_1^{\text{micro}} )</th>
<th>( P_{@8} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mullenbach et al. (2018)</td>
<td>0.218</td>
<td>0.195</td>
<td>0.206</td>
<td>0.499</td>
<td>0.651</td>
</tr>
<tr>
<td>Sadoughi et al. (2018)</td>
<td>0.261</td>
<td>0.186</td>
<td>0.218</td>
<td>0.498</td>
<td>0.662</td>
</tr>
<tr>
<td>Ontological Attention</td>
<td>0.341</td>
<td>0.192</td>
<td>0.245</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3: Benchmark results for the models trained with \( F_1^{\text{micro}} \) stopping criterion.

<table>
<thead>
<tr>
<th>Method</th>
<th>( R_{\text{micro}} )</th>
<th>( P_{\text{micro}} )</th>
<th>( F_1^{\text{macro}} )</th>
<th>( F_1^{\text{micro}} )</th>
<th>( P_{@8} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mullenbach et al. (2018)</td>
<td>0.469</td>
<td>0.593</td>
<td>0.172</td>
<td>0.523</td>
<td>0.685</td>
</tr>
<tr>
<td>Sadoughi et al. (2018)</td>
<td>0.516</td>
<td>0.560</td>
<td>0.173</td>
<td>0.537</td>
<td>0.695</td>
</tr>
<tr>
<td>Ontological Attention</td>
<td>0.514</td>
<td>0.617</td>
<td>0.206</td>
<td>0.560</td>
<td>0.727</td>
</tr>
</tbody>
</table>

Table 4: Ablation study of individual components of the final method. All models are trained with the \( F_1^{\text{macro}} \) stopping criterion. Experiments 2 and 3 do not use the ontological attention mechanism, and instead have one or two attention heads respectively per code-level label. For experiment 4, child-parent and parent-grandparent connections were randomised, removing shared semantics between codes across the full 3 levels.

<table>
<thead>
<tr>
<th>Method</th>
<th>( F_1^{\text{macro}} )</th>
<th>Relative ( F_1^{\text{macro}} ) change (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ontological Attention</td>
<td>0.245</td>
<td>0.0</td>
</tr>
<tr>
<td><strong>Efficacy of ontological attention ensemble</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. No deep supervision</td>
<td>0.243</td>
<td>-0.82</td>
</tr>
<tr>
<td>2. No ontology: One attention head for each label</td>
<td>0.234</td>
<td>-4.5</td>
</tr>
<tr>
<td>3. No ontology: Two attention heads for each label</td>
<td>0.242</td>
<td>-1.2</td>
</tr>
<tr>
<td>4. Partial ontology: Randomised ontological connections</td>
<td>0.231</td>
<td>-5.7</td>
</tr>
<tr>
<td><strong>Efficacy of additional modifications</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5. No class weighting</td>
<td>0.232</td>
<td>-5.3</td>
</tr>
<tr>
<td>6. Reduced convolutional filters (70, as in Sadoughi et al. (2018))</td>
<td>0.236</td>
<td>-3.7</td>
</tr>
</tbody>
</table>

is included in the model reported here. However, this regularisation is not used in our own model where we observed no benefit.

Overall results are shown in Table 2. Our method significantly outperforms the benchmarks on macro-F1 and \( P_{@8} \).

Previous models have optimised for \( F_1^{\text{micro}} \) average. Different target metrics require different design choices: after removal of the class weighting in the loss function and when using \( F_1^{\text{micro}} \) as our stopping criterion, we are able to surpass previous state-of-the-art results on macro-F1. The results are presented in Table 3; our method achieves the highest \( F_1^{\text{micro}} \) score, as well as the highest \( P_{@8} \) score. We note that \( P_{@8} \) score is consistently higher for models stopped using the \( F_1^{\text{micro}} \) criterion.

In Table 4 we present an ablation study. It can be seen that the improvement in performance of the ontological attention model is not simply due to increased capacity of the network, since even with 73% greater capacity (17,858 compared to 10,275 attention vectors), the two-vector multi-headed model has a 1.2% drop in performance. Experiments with deep supervision and randomisation of the ontology graph connections show the benefit of each component of the ontological architecture. We also measure the effect of additional changes made during optimisation of the architecture and training.

**Levels of the ontology:** Three levels of the ontology (including the code itself) were found to be optimal for the Ontological Attention model (see Figure 3). Adding parent and grandparent levels provide incremental gains in accuracy. Adding a level beyond the grandparent node (i.e., the great-grandparent level) does not provide further improvement. Since we identified only 22 ancestral nodes at the level directly above the grandparent, we hypothesise that the grouping becomes too coarse to be beneficial. In fact, all procedure codes share the same ancestor at this level; the remaining
21 nodes are split between diagnostic codes.

Figure 3: $F_{1\text{macro}}$ for models using attention ensembles across different levels of the ontological tree. Error bars represent the standard deviation across 5 different random weight initialisations. The model with 1 level has only the code-level attention head, the model with 2 levels also includes the shared parent attention heads; the model with 3 levels adds the shared grandparent attention heads (this is our reported Ontological Attention model), and finally, the model with 4 levels adds shared great-grandparent attention heads.

4.2 Analysis of the attention weights

In Figure 4 we show how the weights of code-level $u_l$ vectors (which give rise to the attention heads) change when the ontological attention ensemble mechanism is introduced. As expected, we observe that in the case of a single attention head, the weights for different codes largely cluster together based on their position in the ontology graph. Once the parent and grandparent attention heads are trained, the ontological similarity structure on the code level mostly disappears. This suggests that the common features of all codes within a parent group are already extracted by the parent attention. Thus, the capacity of the code-level attention is spent on the representation of the differences between the descendants of a single parent.

4.3 Interpretability of the attention heads

In Section 4.2, we showed the links between the ontology and the attention heads within the space of the $u_l$ vector weights. We can widen this analysis to links between the predictions and the input, by examining which words in the input documents are attended by the three levels of attention heads for a given label. A qualitative visual example is shown in Figure 5. We performed quantitative frequency analysis of high-attention terms (keywords) in the training set. A term was considered a keyword if its attention weight in a document surpassed the threshold $t_{kw}$:

$$t_{kw}(N, \gamma_{kw}) = \gamma_{kw} \frac{1}{N},$$

where $N$ is the length of a document and $\gamma_{kw}$ is a scalar parameter controlling the strictness of the threshold. With $\gamma_{kw} = 1$, a term is considered a keyword if its attention weight surpasses the uniformly distributed attention. In our analysis we chose $\gamma_{kw} = 17$ for all documents.

We aggregated these keywords across all predicted labels in the training set, counting how many times a term is considered a keyword for a label. The results of this analysis are in line with our qualitative analysis of attention maps. The most frequent keywords for the labels presented in the example in Figure 5 include “cancer”, “ca”, “tumor”, at the grandparent level (focusing on the concept of cancer); “metastatic”, “metastases” and “metastasis” at the parent level (focusing on the concept of metastasis); and “brain”, “craniotomy”, “frontal” at the code-level (focusing on terms relating to specific anatomy). A sibling code (198.5 Secondary malignant neoplasm of bone and bone marrow) displays similar behaviour in focusing on anatomy, with “bone”, “spine”, and “back” being among the most frequent keywords.

Not all codes display such structured behaviour. For instance, the grandparent 401-405 Hypertensive disease attended to the term “hypertension” most frequently. The parent code 401 Essential hypertension, does not attend to “hypertension”, but neither does it attend to any useful keywords — this may be due to the code being simple compared to its sibling codes, which are more specific (e.g., 402 Hypertensive heart disease). Interestingly, the children of 401 Essential hypertension attend to the word “hypertension” again, while also focusing on terms that set them apart from each other — e.g., 401.0 Malignant essential hypertension focuses on terms implying malignancy, such as “urgency”, “emergency”, and “hemorrhage”.

5 Limitations due to labelling variability

Since performance on this task appears to be much lower than might be acceptable for real-world use, we investigated further. Figure 6 shows the per-label F1 scores; it can be seen that there is high
Figure 4: Two-dimensional t-SNE (Maaten and Hinton, 2008) representation of the $u_l$ vectors (which give rise to the attention heads) for a subset of 182 codes with at least 100 occurrences each, in the data belonging to 16 different parent nodes. (a) Legend for the annotation of data points according to their parent node. (b) $u_l$ vectors from the model with only a single attention head for each code (i.e., no ontology). It can be seen that codes naturally cluster by their parent node. Selected higher-level alignments are indicated by additional contours — for grandparent nodes (3 nodes) and for diagnoses/procedure alignment (in the case of cardiovascular disease). (c) $u_l$ vectors in the ontological attention ensemble model for the same set of codes (and the same t-SNE hyperparameters). In most cases the clustering disappears, indicating that the attention weights for the ancestral codes have extracted the similarities from descendants’ clusters.

<table>
<thead>
<tr>
<th>Method</th>
<th>$R_{macro}$</th>
<th>$P_{macro}$</th>
<th>$F1_{macro}$</th>
<th>$F1_{micro}$</th>
<th>$P@8$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mullenbach et al. (2018)</td>
<td>0.226</td>
<td>0.200</td>
<td>0.212</td>
<td>0.500</td>
<td>0.651</td>
</tr>
<tr>
<td>Sadoughi et al. (2018)</td>
<td>0.272</td>
<td>0.187</td>
<td>0.222</td>
<td>0.497</td>
<td>0.662</td>
</tr>
<tr>
<td>Ontological Attention</td>
<td>0.347</td>
<td>0.199</td>
<td><strong>0.252</strong></td>
<td>0.507</td>
<td>0.686</td>
</tr>
</tbody>
</table>

Table 5: Benchmark results for the models trained with $F1_{macro}$ stopping criterion.

variability in accuracy, that is only partially correlated with the number of training examples.

Inspection of examples for some of the poorly performing codes revealed some variability in coding policy, described further below.

### 5.1 Misreporting of codes

The phenomenon of human coding errors is reported in the literature; for instance, Kokotailo
Figure 5: Discharge summary snippet with highlights generated from attention heads for (a) the grandparent code (190-199 Malignant neoplasm of other and unspecified sites), (b) the parent code (198 Secondary malignant neoplasm of other specified sites), and (c) the specific code (198.3 Secondary malignant neoplasm of brain and spinal cord). Different words and phrases are attended at each level.

Figure 6: Per-code frequency of training examples v.s. $F_1^{macro}$ score from the ontological attention model and Hill estimated sensitivity and specificity to be 80% and 100% respectively for ICD codes relating to stroke and its risk factors (Kokotailo and Hill, 2005). In the MIMIC-III dataset, we inspected the assignment of smoking codes (current smoker 305.1, past smoker V15.82, or never smoked i.e., no code at all), using regular expression matching to identify examples of possible miscoding, followed by manual inspection of 60 examples (10 relating to each possible miscoding category) to verify our estimates. We estimated that 10% of patients had been wrongly assigned codes, and 30% of patients who had a mention of smoking in their record had not been coded at all. We also observed that often the “correct” code is not clear-cut. For instance, many patients had smoked in the distant past or only smoke occasionally, or had only recently quit; in these cases, where the narrator reliability may be questionable, the decision of how to code is a matter of subjective clinical judgement.

5.2 Revisions to the coding standards

Another limitation of working with the MIMIC-III dataset is that during the deidentification process, information about absolute dates was discarded. This is problematic when we consider that the MIMIC-III dataset contains data that was collected between 2001 and 2012, and the ICD-9 coding standard was reviewed and updated annually between 2006 and 2013 (Centers for Medicare & Medicaid Services) i.e., each year some codes were added, removed or updated in their meaning.

To investigate this issue, we took the 2008 standard and mapped codes created post-2008 back to this year. In total, we identified 380 codes that are present in the dataset but were not defined in the 2008 standard. An example can be seen in Figure 7. We report our results on the 2008 codeset in Table 5. It can be seen that there is an improvement to the metrics on this dataset, which we expect would increase further if all codes were mapped back to the earliest date of 2001. Without time data, it is an unfair task to predict codes which are fundamentally time-dependent. This is an interesting example of conflicting interests between (de)identifiability and task authenticity.

During real-world deployment, codes should be assigned according to current standards. In order to use older data, codes should be mapped forwards rather than backwards. The backwards operation was possible by automated re-mapping of the codes, however the forwards operation is more arduous. Newly introduced codes may require annotation of fresh labels or one-to-many conversion — both operations requiring manual inspection of the original text. A pragmatic approach would be to mask out codes for older documents where they cannot be automatically assigned.

6 Conclusions

We have presented a neural architecture for automated clinical coding which is driven by the ontological graph of relationships between codes. This model establishes a new state-of-the-art result for the task of automated clinical coding with MIMIC-III dataset. Compared to simply doubling the number of attention heads, our ontological attention ensemble mechanism provides improve-
ments in accuracy, in memory efficiency, and in interpretability. Our method is not specific to an ontology, and in fact could be used for a graph of any formation. If we were to exploit further connections within the ICD ontology e.g., between related diagnoses and procedures, and between child codes which share modifier digits, we would expect to obtain a further performance boost.

We have illustrated that labels may not be reliably present or correct. Thus, even where plenty of training examples are available, the performance may (appear to) be low. In practice, the most successful approach may be to leverage a combination of automated techniques and manual input. An active learning setup would facilitate adoption of new codes by the model as well as allowing endorsement of suggested codes which might otherwise have been missed by manual assignment, and we propose this route for future research.

References


Adam Paszke, Sam Gross, Soumith Chintala, Gregory Chanan, Edward Yang, Zachary DeVito, Zeming Lin, Alban Desmaison, Luca Antiga, and Adam


Neural Token Representations and Negation and Speculation Scope Detection in Biomedical and General Domain Text

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Abstract
Since the introduction of context-aware token representation techniques such as Embeddings from Language Models (ELMo) and Bidirectional Encoder Representations from Transformers (BERT), there have been numerous reports on improved performance on a variety of natural language tasks. Nevertheless, the degree to which the resulting context-aware representations can encode information about morpho-syntactic properties of the tokens in a sentence remains unclear.

In this paper, we investigate the application and impact of state-of-the-art neural token representations for automatic cue-conditional speculation and negation scope detection coupled with the independently computed morpho-syntactic information. Through this work, we establish a new state-of-the-art for the BioScope and NegPar corpora.

Furthermore, we provide a thorough analysis of neural representations and additional features interactions, cue-representation for conditioning, discussing model behavior on different datasets and, finally, address the annotation-induced biases in the learned representations.

1 Introduction
In 2018, a set of new state-of-the-art results were established for a variety of Natural Language Processing tasks, the majority of which can be attributed to the introduction of context aware token representations, learned from large amounts of data with Language-modeling like tasks as a training goal (Devlin et al., 2018; Peters et al., 2018).

It is, however, unclear to what degree the computed representations capture and encode high-level morphological/syntactic knowledge about the usage of a given token in a sentence. One way of exploring the potential of the learned representation would be through investigating the performance on a task that would require the representation to acquire some notion of syntactic units such as phrases and clauses, as well as the relationship between the syntactic units and other tokens in the model. An example of such a task is Speculation or Negation Scope Detection.

The main contributions of this work can be summarized as follows:

- We achieve and report a new state-of-the-art for the negation and speculation scope detection on several biomedical and general domain datasets, which were created using different definitions of what constitutes a scope of a given negation/speculation.¹

- We investigate different ways of incorporating additional automatically-generated syntactic features into the model and explore the potential improvements resulting from the addition of such features.

- Following Fancellu et al. (2017), we provide a thorough comparison of our proposed model with other state-of-the-art models and analyze their behaviour in the absence of potential “linear clues”, the presence of which might result in highly accurate predictions even for syntax-unaware token representations.

2 The Task
In general, speculation or negation scope detection can be constructed as the following conditional token classification task: given a negation or speculation cue (i.e., a word or phrase that expresses negation or speculation such as ‘No’ and ‘May’). ¹

¹ An implementation of our model together with the pre-trained models for scope detection will be available later.
identify which tokens are affected by the negation or represent an event that is speculative in nature (referred to as the scope of the negation or speculation). Consider the following example:

(1) These findings that *(may be from an acute pneumonia)* include minimal bronchiectasis as well.

In this case, the speculation cue is “*may*” and the string of tokens that contains the speculative information is “*may be from an acute pneumonia*”.

Each data point, as such, is a string of tokens paired with the corresponding negation or speculation cue. Note that nested negations in the same sentence would be distinguished only by the associated cue.

From the syntactic structure point of view, it is clear that in most cases, the boundaries of a given scope strongly correlate with the clausal structure of the sentence *(Morante and Sporleder, 2012)*. There is also a strong connection between the fine-grained part-of-speech (POS) of the cue and the scope boundaries.

Consider the following examples where the type of possible adjectives (either attributive or predicative) results in different scope boundaries (scope highlighted as italic):

(2) This is a patient who had *possible pyelonephritis* with elevated fever.

(3) *Atelectasis in the right mid zone is, however, possible.*

Such a property of the task requires a well-performing model to be able to determine cue-types and the corresponding syntactic scope structure from a learned representation of cue-sentence pairs. As such, it can be used as an (albeit imperfect) proxy for assessing the knowledge about the structure of the syntax that a sentence aware token representation potentially learns during training.

### 2.1 Datasets

There are no universal guidelines on what constitutes a scope of a given negation or speculation; different definitions might affect a given model’s performance. To take this ambiguity into account, we report our results on two different datasets: BioScope *(Vincze et al., 2008)* and NegPar *(Liu et al., 2018)*.

- The BioScope corpus *(Vincze et al., 2008)* consists of three different types of text: Biological publication abstracts from Genia Corpus (1,273 abstracts), Radiology reports from Cincinnati Children’s Hospital Medical Center (1,954 reports), and full scientific articles in the bioinformatics domain (nine articles in total). In this work, we focus on two of the sub-corpora: Abstracts and Clinical reports. One should note that BioScope corpus does not allow discontinuous scopes.

- NegPar *(Liu et al., 2018)* is a corpus of Conan Doyle stories annotated with negation cues and the corresponding scopes. The corpus is available both in English and Chinese. In this work, we only use the English part of the corpus. Unlike BioScope, NegPar provides a canonical split as training (981 negation instances), development (174 instances) and test sets (263 negation instances). NegPar annotation guidelines allows for discontinuous scopes.

### 3 Previous Work

Negation scope detection algorithms can be classified into two categories: (1) rule-based approaches that rely on pre-defined rules and grammar; and (2) statistical machine learning approaches that utilize surface level features of the input strings to detect the scope of the negation.

**Rule-based approaches** Due to the somewhat restricted nature of clinical texts syntax, a pre-defined rule-based key-word triggered negation scope detection system achieves competitive performance on a variety of clinical-notes derived data-sets *(Chapman et al., 2001; Harkema et al., 2009; Elkin et al., 2005)*.

**Machine learning approaches** While rule-based approaches might achieve high performance on medical institution specific datasets, they do not generalize well for other dataset types and they may require customization of the rules to adapt to the new corpus and/or domain. By contrast, machine learning-based systems do not require active human expert participation to adapt to a new dataset/domain. Earlier works utilizing the statistical approaches for negation scope detection include Support Vector Machines (SVM), Conditional Random Fields based models (CRF) *(Agarwal and Yu, 2010; Councill et al., 2010)* as well
as hybrid CRF-SVM ensemble models (Zhu et al., 2010) (Morante and Daelemans, 2009)

Recently, Neural Network-based approaches have been proposed for such tasks, including Convolutional Neural Network (CNN)-based (Qian et al., 2016) and Long Short Term Memory (LSTM)-based (Fancellu et al., 2017; Sergeeva et al., 2019) models.

The work on specifically speculation scope detection is less varied and mainly confined to CONLL-2010 Shared-Task2 submissions (Farkas et al., 2010). It is, however, important to note that due to the similarity in the formulation of the task, the majority of the negation-specific machine learning approaches can be directly applied to the speculation scope detection problem provided the speculation annotated data is available for training.

We also draw inspiration from a large body of work (Linzen et al., 2016; Gulordava et al., 2018; Marvin and Linzen, 2018) examining the nature of modern context aware representations from a linguistic perspective.

4 Model Training and Evaluation

4.1 Neural Token Representation

The use of pre-trained continuous word representations has been ubiquitous in modern statistical natural language processing. The importance of an appropriate word-level representation is especially noticeable in per-token prediction tasks: in such a set-up the model goal is to fine-tune or modify the existing input token representation in such a way that it contains the necessary information to make a correct classification decision at prediction time.

In this work, we consider the following approaches for generating the input token representation:

• Global Vectors (GloVe) (Pennington et al., 2014): A pre-trained token representation that relies on the direct matching of tokens and the corresponding ratios of token co-occurrences with their neighbours. Note that the definition of the neighbour in this setup is static (that is, the ultimate representation would incorporate an averaged notion of context) and relies on the bag-of-words representation of the context.

• Embeddings from Language Models (ELMo) (Peters et al., 2018): A bi-directional LSTM model-based token representation, pre-trained on the language modeling task. Instead of modeling the bag-of-words neighborhood co-occurrence probabilities directly, this model approximates the conditional probability of a token given the ordered linear context of the token usage.

• Bidirectional Encoder Representations from Transformers (BERT) (Devlin et al., 2018): A transformer-based token representation trained on the modified language modeling task together with a broader context next sentence prediction task. In this model, the context of a token is continuously incorporated into the representation of the token itself as a weighted sum of the neighboring token representations through the use of the multi-head-attention mechanism. The linear order of the token information is provided at input time as an additional positional embedding, since the unmodified transformer architecture does not encode any notion of the linear order.

Despite the performance gains achieved by the widespread use of contextual word embeddings like ELMo and BERT, the questions about the nature of the learned representation remain unanswered. Both ELMo and BERT were introduced to incorporate the wider structure of the given input into individual token representation at the time of training; however, both models only have access to the linear order of the context.

The question then arises: To what degree does the word embedding trained on a language modeling like task and computed using the whole linear context of a sentence encode the broader syntax-related characteristics of a token used within a context?

In order to gain insight into the nature of the learned representations and their potential use for negation and speculation scope detection, we introduce the following syntax-informed features to be used together with the token embedding:

POS : Part-Of-Speech of a given token as defined by the Penn Treebank tagging scheme (Marcus et al., 1993).

DEP : Type of dependency between the token and its parent, representing limited dependency tree information of a given token.
**PATH**: A string of Google Universal POS tags (Petrov et al., 2012) of the three direct ancestors of the token in the dependency tree; this feature captures local constituent-like information of a given token.

**LPATH**: Depth of the token in the syntactic dependency tree.

**CP**: The distance between a given token and the negation cue in the constituency parse tree generated using (Kitaev and Klein, 2018). If a negation cue has multiple tokens, the minimum of the distances is used.

Note that all features were automatically generated, and as a result, represent a “noisy” source of information about the syntactic characteristic of a token. If adding syntactic features as additional inputs would not affect or would significantly degrade the model’s performance, it is reasonable to assume that the information represented by such features is already present in the token representation in some way.

### 4.2 Modes of Evaluation

To provide a fair comparison of different types of embeddings, we introduce two different modes of evaluation. The first mode (referred to as Feature-based embeddings later in the paper) is designed to test the embeddings in the same setup as previously used to get the state-of-the-art performance on the dataset. The second mode (referred to as BERT fine-tuning later in the paper) is designed to test BERT embeddings in their native direct fine-tuning setting.

**Feature-based Embeddings using Bi-directional LSTM**: Figure 1 demonstrates the proposed framework for the desired task. One should note that the factor that differentiates the two experiments from one another is the embeddings. The task specific layers (two-layer Bi-directional LSTM) remains the same across all experiments. To properly condition each scope on a given cue, we concatenate a specific cue embedding to the input embedding, before computing the final representation for each token. Additional syntactic information is also provided by concatenating the input embedding with all of the syntactic feature embeddings.

**BERT Fine-tuning**: The original setup for the use of BERT embedding does not require an elaborate task-specific layer; the task specific model is a copy of the original transformer-based BERT architecture with the corresponding pre-trained model-parameters, and the top prediction layer swapped for a new task specific layer that predicts the probability of a given label for a token representation. Crucially, the token representation is allowed to change during the fine-tuning. For this particular setup, it is unclear how to account for the conditional nature of the scope prediction task. In other words, a sentence can potentially contain more than one negation/piece of speculative information.

We consider two different testing scenarios to evaluate the different ways of providing the cue information to the model:

1. Providing the embedded cue at the top layer of the model by concatenating it to the learned token embedding.
2. Providing the embedded cue at the bottom as a part of the input to the transformer layer before the fine-tuning by adding the cue embeddings (initialized randomly at the fine-tuning stage) to the initial token representation.

To test if the additional syntactic information provides any additional benefit to our framework, we also add the mean of all of the syntactic feature embeddings to the initial pre-transformer representation of the input.

### 4.3 Hyperparameter Settings

**Feature-based Embeddings** For the aforementioned set of experiments, the following architecture parameters have been considered:
Table 1: Performance of the negation scope detection task on BioScope and NegPar corpora using different approaches. Results are reported as the percentage of number of predicted scopes that exactly match the golden scope (PCS).

<table>
<thead>
<tr>
<th>Model</th>
<th>BioS_Abstracts</th>
<th>BioS_Clinical</th>
<th>NegPar</th>
<th>NegPar(CV)</th>
</tr>
</thead>
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<tr>
<td>Fancellu et al. (2017)</td>
<td>81.38%</td>
<td>94.21%</td>
<td>68.93%</td>
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<tr>
<td>Fancellu et al. (2018)</td>
<td>N/A</td>
<td>N/A</td>
<td>61.98%</td>
<td>N/A</td>
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<tr>
<td>Bi-LSTM_{GloVe}</td>
<td>63.24%(1.80%)</td>
<td>90.46%(3.64%)</td>
<td>51.48%(4.45%)</td>
<td>49.18%(4.97%)</td>
</tr>
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<td>Bi-LSTM_{ELMo}</td>
<td>81.62%(1.87%)</td>
<td>93.10%(2.18%)</td>
<td>71.52%(1.98%)</td>
<td>75.29%(3.35%)</td>
</tr>
<tr>
<td>Bi-LSTM_{BERT}</td>
<td>79.29%(3.06%)</td>
<td>91.26%(2.82%)</td>
<td>66.78%(3.50%)</td>
<td>69.45%(3.55%)</td>
</tr>
<tr>
<td>Bi-LSTM_{GloVe} + AF</td>
<td>79.00%(2.07%)</td>
<td>94.02%(1.98%)</td>
<td>69.70%(2.81%)</td>
<td>73.11%(3.19%)</td>
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<tr>
<td>Bi-LSTM_{ELMo} + AF</td>
<td>83.30%(3.16%)</td>
<td>94.25%(2.86%)</td>
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</tr>
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<td>Bi-LSTM_{BERT} + AF</td>
<td>80.68%(3.23%)</td>
<td>93.10%(2.77%)</td>
<td>67.42%(2.12%)</td>
<td>73.39%(4.12%)</td>
</tr>
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<td>BERT (c-top)</td>
<td>74.63%(3.23%)</td>
<td>92.87%(2.04%)</td>
<td>63.14%(2.08%)</td>
<td>—</td>
</tr>
<tr>
<td>BERT (c-bottom)</td>
<td>86.97%(2.24%)</td>
<td>93.68%(2.37%)</td>
<td>76.78%(2.04%)</td>
<td>81.91%(3.04%)</td>
</tr>
<tr>
<td>BERT (c-bottom) + AF</td>
<td><strong>87.03%(2.38%)</strong></td>
<td><strong>94.25%(2.86%)</strong></td>
<td><strong>79.00%(1.37%)</strong></td>
<td><strong>80.64%(2.57%)</strong></td>
</tr>
</tbody>
</table>

The number in the parenthesis indicates the standard deviation of the score.

- These results are generated using an older version of the corpus annotation.
- Since this work is aimed at cross-lingual negation detection, the reported results are based on using cross-language word embeddings, which are likely to degrade a single-language model performance.

Figure 2: A diagram of the proposed BERT-based architecture for negation and speculation scope detection with inclusion of additional features.

- Word embedding dimension: GloVe: 300; ELMo, BERT: 1024
- Syntactic feature embedding dimension: 10 per feature
- Task-specific LSTM embedding dimension: 400

During training, a dropout rate of 0.5 (Gal and Ghahramani, 2016) was used to prevent overfitting. The Adam optimizer (Kingma and Ba, 2014) was used with step size of $10^{-3}$ and batch size of 32 for 50 epochs for BioScope and 200 epochs for NegPar. The reason why we use different epochs is that there are fewer training examples for NegPar than BioScope. Therefore, it takes more epochs for the NegPar models to converge.

**BERT Fine-tuning** The BERT models have the following architecture parameters:

- Word embedding dimension: 1024
- BERT_{LARGE} layer transformer configuration (Devlin et al., 2018)
- Syntactic features embedding dimension: 1024 for each feature
- Cue embedding dimensions: 1024

We perform fine tuning on the negation/speculation task for 20 epochs. The Adam optimizer was used with learning rate of $10^{-5}$ and batch size of 2 for 10 epochs for the BioScope corpus and 50 epochs for the NegPar corpus.

### 4.4 Evaluation Procedure

We report our results in terms of the percentage of number of predicted scopes that exactly match the golden scopes (PCS). Since pre-trained BERT models use their own tokenization algorithm, it results in inconsistent final number of tokens in the dataset across evaluation modes. As a result, other traditional evaluation metrics such as precision, recall and F1 are inappropriate to be used in this study as they depend on the number of tokens.
Since the BioScope dataset does not have a canonical training/development/test set split, we report 10-fold cross-validation results together with the standard deviation of the resulting scores.

For the NegPar dataset, we report the result on the test set as well as 10-fold cross-validation results. To overcome the possible random initialization influences on the results, we report the average score for 10 random seeds on the test set together with the associated standard deviation.

5 Results

The performance of different approaches on BioScope and NegPar corpora for the negation scope detection and the speculating scope detection are shown in Table 1 and Table 2, respectively. BiLSTM-marked entries of the table correspond to Feature-based and BERT-marked entries correspond to BERT fine-tuning approaches.

5.1 Feature-based Approach

Effect of embedding on performance: Except for the negation scope detection task on BioScope clinical notes, ELMo embeddings significantly outperformed GloVe embeddings as well as the feature-based use of BERT embeddings, but not the fine-tuned version of BERT. While the former is expected, the latter is noteworthy: for NER task (Devlin et al., 2018), for example, the difference in performance between the fine-tuning and feature-based approach results is 1.5% of the F1 score. For negation scope detection the difference is a striking 7.68% on BioScope-abstracts and 10% on a test set of the NegPar dataset. For speculation scope detection the difference remains as large (7.93%). We theorize that this differences comes from the different syntactic nature of the target strings of tokens: NER systems are concerned with finding named entities in text, where the majority of the named entities are represented by relatively short (token-wise) noun phrases, negation/speculation scope detection requires recognition of a much more diverse set of syntactic phenomena. This suggests an important difference between the featurized and fine-tuned approaches for highly syntax-dependent token classification tasks.

Syntactic features induced gains: In general, we observe consistent small gains in performance for all types of embedding on BioScope (both speculation and negation detection modalities) but not on the NegPar dataset. The only exception to this pattern is in non-context aware GloVe embeddings. Adding syntactic features embeddings has inconsistent effects on standard deviations over modalities and datasets.

5.2 BERT fine-tuning approach

Cue-conditioning influence on the results: The way to condition a given instance on a particular cue greatly influences the model performance: providing cue information at the top layer of the model results in poor performance of the model for all datasets and both negation and speculation modalities.

Syntactic features induced gains and the importance of Cross Validation evaluation: Adding features to the best performing BERT fine-tuned models does not result in any significant differences on the BioScope dataset. We observe a significant gain in performance on NegPar: note that in this case the gain is purely train/test set split induced and disappears entirely in a cross-validation mode of evaluation.

Artificial noise and the model performance: Even though the experimental results suggest no to minimal contribution of the additional features to the best model performance, natural questions to ask are: “Does the feature enriched model rely on the provided features during the prediction phase?” and “Do the final learned representations differs significantly for feature-enriched and featureless inputs?” We introduce noise into the trained model inputs to check if artificial noise undermines its performance. In particular, we consider the model BERT(cue-bottom) + AF, as it provides the best performance out of all feature-enriched models.

With a given probability, which we call the noise level, we replace a given feature value with a random value: for categorical features (POS, DEP, PATH), we replace it with a random category, and for numerical features (LPATH,CP), we replace it with a random integer drawn from a uniform distribution bounded by the feature’s possible minimum and maximum values. We observe a consistent and significant decrease in performance as the probability of seeing the incorrect features increases (see Figure 3). This suggests that the additional features introduced in this paper play an important role in decision making. This is supported by the fact that the performance on clini-
natural reports negation detection remains almost un-
affected by the change, since the majority of the
negation scopes in this dataset can be captured by
structure-independent heuristics.

![Figure 3: Performance of the BERT + AF models with respect to the noise level, averaged for 10 fold CV](image)

**5.3 Linear punctuation cues and model performance**

Even though the scope boundaries correlate to the
syntactic structures of the sentence, a good per-
formance on a given dataset does not necessarily
prove the model acquired any kind of a structural
knowledge: as was noted in Fancellu et al. (2017),
the majority of scopes in the BioScope corpus
consist of cases where the punctuation boundaries
match the scope boundaries directly. For those
cases, the model does not have to learn any kinds
of underlying syntactic phenomena: learning a
simple heuristic to mark everything between a cue
and the next punctuation mark as a scope would
produce an illusion of a more complex syntax-
informe performance.

To see if our model’s performance is signifi-
cantly affected by the punctuation clues, we re-
move all the punctuation from the training cor-
pus, re-train all the models on the modified data-
sset and evaluate the learned models on the test set.
We also report the performance on “hard” (non-
punctuation bound) instances of scopes separately.

As can be seen in Table 3, removing punctua-
tion affects all models’ behaviour similarly: model
performance degrades by losing 2-3 percent of
PCS on average. Interestingly, the performance on the
non-punctuation boundaries scopes declines similarly, which suggest that punctuation plays an
important role in computing a given token repre-
sentation, and not only as a direct linear cue that
signifies the scope’s start and end.

**5.4 Error overlap**

Given the difference in the model architectures, a
natural question to ask is: “Is the best perform-
ing model strictly better than the others, or do
they make different types of errors?” We compute
the error overlap between BERT and ELMo on
the negation detection task as shown in Figure 4.
About half of ELMo and slightly more than a quar-
ter of BERT errors appear to be model specific,
suggesting the potential for ensemble-induced
improvements.

Table 2: Performance of speculation scope detection task on BioScope corpus using different approaches. Results
are reported as the percentage of number of the predicted scopes that exactly match the golden scope (PCS).

<table>
<thead>
<tr>
<th>Model</th>
<th>BioScope_Abstracts</th>
<th>BioScope_Clinical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Qian et al. (2016)</td>
<td>85.75%</td>
<td>73.92%</td>
</tr>
<tr>
<td>Bi-LSTM_{GloVe}</td>
<td>47.99% (4.07%)</td>
<td>46.90% (2.87%)</td>
</tr>
<tr>
<td>Bi-LSTM_{ELMo}</td>
<td>84.62% (2.33%)</td>
<td>81.82% (2.74%)</td>
</tr>
<tr>
<td>Bi-LSTM_{BERT}</td>
<td>81.35% (1.95%)</td>
<td>78.75% (3.24%)</td>
</tr>
<tr>
<td>Bi-LSTM_{GloVe} + AF</td>
<td>85.07% (2.66%)</td>
<td>80.73% (3.01%)</td>
</tr>
<tr>
<td>Bi-LSTM_{ELMo} + AF</td>
<td>86.57% (2.65%)</td>
<td>81.55% (2.74%)</td>
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<tr>
<td>Bi-LSTM_{BERT} + AF</td>
<td>84.43% (1.08%)</td>
<td>81.37% (4.32%)</td>
</tr>
<tr>
<td>BERT (cue-top)</td>
<td>57.32% (2.14%)</td>
<td>60.49% (4.77%)</td>
</tr>
<tr>
<td>BERT (cue-bottom)</td>
<td><strong>89.28% (1.65%)</strong></td>
<td><strong>83.71% (2.77%)</strong></td>
</tr>
<tr>
<td>BERT (cue-bottom) + AF</td>
<td>88.91% (1.65%)</td>
<td>82.36% (4.27%)</td>
</tr>
</tbody>
</table>

The number within the parentheses indicates the standard deviation of the score.
Table 3: Performance on percentage of correct span on BioScope Abstracts sub-corpus trained under different schemes.

<table>
<thead>
<tr>
<th></th>
<th>Trained w/ punctuation</th>
<th>Trained w/o punctuation</th>
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<td></td>
<td>all hard cases</td>
<td>all hard cases</td>
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<tr>
<td>Bi-LSTM GloVe</td>
<td>63.24%(1.80%)</td>
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<td>Bi-LSTM ELMo</td>
<td>81.62%(1.87%)</td>
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<td>Bi-LSTM BERT</td>
<td>79.29%(3.06%)</td>
<td>77.54%(3.18%)</td>
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<td>Bi-LSTM GloVe + AF</td>
<td>79.00%(2.07%)</td>
<td>76.21%(1.76%)</td>
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<td>Bi-LSTM ELMo + AF</td>
<td>83.30%(3.16%)</td>
<td>82.31%(2.95%)</td>
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<tr>
<td>Bi-LSTM BERT + AF</td>
<td>80.68%(3.23%)</td>
<td>80.45%(3.24%)</td>
</tr>
<tr>
<td>BERT (c-bottom)</td>
<td>86.97%(2.24%)</td>
<td>83.48%(3.22%)</td>
</tr>
<tr>
<td>BERT (c-bottom + AF)</td>
<td>87.03%(2.38%)</td>
<td>84.58%(3.58%)</td>
</tr>
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6 Conclusions and Future Work

This work presents a comparison among different context-aware neural token representations and the corresponding performance on the negation and speculation scope detection tasks. Furthermore, we introduce a new state-of-the-art BERT-based cue-conditioned feature-enriched framework for negation/speculation scope detection. Based on the empirical results, we are inclined to recommend BERT fine-tuning over using a feature-based approach with BERT for syntax-dependent tasks.

We used two commonly used publicly available datasets, BioScope and NegPar for our evaluation. Despite the observed gains on the test set of the NegPar corpus, the effect of the syntactic features on BERT (fine-tuned) performance remains largely inconclusive.

It is also important to note that the syntactic information we have been trying to incorporate into the model was generated automatically; one of the possible avenues of research would be comparing the possible golden annotation induced gains with the imperfect information gain we observe when incorporating silver syntactic features.

We were unable to find any consistent grammatical explanation for the errors context-aware models result in on the test data; however, this does not conclusively mean that such an explanation does not exist. An appropriate next step would be annotating a smaller set of sentences, grouped by the corresponding syntactic construction and see if a given token representation yields improved performance on such a construction.

Figure 4: Distribution of error overlaps: BERT vs. ELMo on BioScope Abstracts dataset.

Figure 5: Distribution of error overlaps: BERT vs. BERT with features (BERT + AF) vs. ELMo on NegPar test set.
Acknowledgments

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References


<table>
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<td>Schoene, Annika M</td>
<td>136</td>
</tr>
<tr>
<td>Schrempf, Patrick</td>
<td>168</td>
</tr>
<tr>
<td>Sergeeva, Elena</td>
<td>178</td>
</tr>
<tr>
<td>Shieh, Alexander Te-Wei</td>
<td>108</td>
</tr>
<tr>
<td>Su, Shang-Yu</td>
<td>108</td>
</tr>
<tr>
<td>Surdeanu, Mihai</td>
<td>87</td>
</tr>
<tr>
<td>Szolovits, Peter</td>
<td>178</td>
</tr>
<tr>
<td>Tahmasebi, Amir</td>
<td>178</td>
</tr>
<tr>
<td>Tsaftaris, Sotirios</td>
<td>168</td>
</tr>
<tr>
<td>Tsai, Shang-Chi</td>
<td>39</td>
</tr>
<tr>
<td>Turcan, Elsbeth</td>
<td>97</td>
</tr>
<tr>
<td>Turner, Alexander P</td>
<td>136</td>
</tr>
</tbody>
</table>
Van, Hoang, 87
Wright-Bettner, Kristin, 1
Yew, Justin, 126
Yim, Wen-wai, 126
Zhu, Henghui, 178
Zine El Abidine, Khaldoun, 157