

Medical Information Extraction in the Age of Deep Learning

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Summary

Objectives: We survey recent developments in medical Information Extraction (IE) as reported in the literature from the past three years. Our focus is on the fundamental methodological paradigm shift from standard Machine Learning (ML) techniques to Deep Neural Networks (DNNs). We describe applications of this new paradigm concentrating on two basic IE tasks, named entity recognition and relation extraction, for two selected semantic classes—diseases and drugs (or medications)—and relations between them.

Methods: For the time period from 2017 to early 2020, we searched for relevant publications from three major scientific communities: medicine and medical informatics, natural language processing, as well as neural networks and artificial intelligence.

Results: In the past decade, the field of Natural Language Processing (NLP) has undergone a profound methodological shift from symbolic to distributed representations based on the paradigm of Deep Learning (DL). Meanwhile, this trend is, although with some delay, also reflected in the medical NLP community. In the reporting period, overwhelming experimental evidence has been gathered, as illustrated in this survey for medical IE, that DL-based approaches outperform non-DL ones by often large margins. Still, small-sized and access-limited corpora create intrinsic problems for data-greedy DL as do special linguistic phenomena of medical sublanguages that have to be overcome by adaptive learning strategies.

Conclusions: The paradigm shift from (feature-engineered) ML to DNNs changes the fundamental methodological rules of the game for medical NLP. This change is by no means restricted to medical IE but should also deeply influence other areas of medical informatics, either NLP- or non-NLP-based.

Keywords

Neural networks, deep learning, natural language processing, information extraction, named entity recognition, relation extraction

Yearb Med Inform 2020;208-20

<http://dx.doi.org/10.1055/s-0040-1702001>

1 Introduction

The past decade has seen a truly revolutionary paradigm shift for Natural Language Processing (NLP) as a result of which Deep Learning (DL) (for a technical introduction, cf. [1]; for comprehensive surveys, cf. [2] and [3]) became the dominating mind-set of researchers and developers in this field (for surveys, cf. [4, 5]). Yet, DL is by no means a new computational paradigm. Rather it can be seen as the most recent offspring of neural computation in the evolution of computer science (cf. the historical background provided by Schmidhuber [6]). But unlike in previous attempts, it now turns out to be extremely robust and effective for adequately dealing with the contents of unstructured visual [7], audio/speech [8], and textual data [9].

The success of Deep Neural Networks (DNNs) has many roots. Perhaps the most important methodological reason is that, with DNNs, manual feature selection or (semi-)automated *feature engineering* is abandoned. This time-consuming tuning step was at the same time mandatory and highly influential on the performance of earlier generations of ML systems in NLP based on Markov Models (MMs), Conditional Random Fields (CRFs), Support Vector Machines (SVMs), etc. In a DL system, however, the relevant features (and their relative contribution to a classification decision) are automatically computed as a result of thousands of iterative training cycles.

The ultimate reason for the success behind DNNs is a pragmatic criterion though: *system performance*. Compared with results in biomedical Information

Extraction (IE), obtained in previous years with standard ML methods, DL approaches changed profoundly the rules of the game. In a landslide manner, for the same task and domain, performance figures jumped up to levels unprecedented so far and DL systems consistently outperformed by large margins non-DL state-of-the-art (SOTA) systems for different tasks. Section 3 provides ample evidence for this claim and features the new SOTA results with a deeper look at IE, a major application class of medical NLP (for alternative surveys, cf. [10–12]).

Despite specialized hardware at disposal now, training DNNs still requires tremendous computational resources and processing time. Luckily, for general NLP, huge collections of language models (so-called *embeddings*) have already been trained on huge corpora (comprised of hundreds of millions of Web-scraped documents, including newspaper and Wikipedia articles) so that these pre-compiled model resources can be readily reused when dealing with *general-purpose* language. But medical (and biological) language mirrors *special-purpose* language characteristics and comprises a large variety of sublanguages of its own. This becomes obvious in Section 3 where we deal with scholarly scientific writing (with documents typically taken from PubMed). Here, differences to general language are mostly due to the use of a highly specialized technical vocabulary (covered by numerous terminologies, such as MeSH, SNOMED-CT, or ICD). Even more challenging are clinical notes and reports (with documents often taken from the MIMIC¹ (Medical Information Mart

¹ <https://mimic.physionet.org/>

for Intensive Care) clinical database) which typically exhibit syntactically ill-formed, telegraphic language with lots of acronyms and abbreviations as an additional layer of complexity (cf. the seminal descriptive work distinguishing both these sublanguage types by Friedman et al. [13]). Newman-Griffis and Fosler-Lussier [14] investigated different sublanguage patterns for the many varieties of clinical reports (pathology reports, discharge summaries, nurse and Intensive Care Unit notes, etc.), while Nunez and Carenini [15] discussed the portability of embeddings across various fields of medicine reflecting characteristic sublanguage use patterns. These constraints have motivated the medical NLP community to adapt embeddings originally trained on general language to the medical language. Table 1 lists those medically informed embeddings, many of which are the basis for the IE applications discussed in Section 3.

Our survey emphasizes the fundamental methodological paradigm shift of current NLP research from symbolic to distributed representations as the basis of DL. It thus complements earlier contributions to the International Medical Informatics Association (IMIA) Yearbook of Medical Informatics which focused exclusively on the role of social media documents [23], had a balanced view on the relevance of both Electronic Health Records (EHRs) and social media posts [24], or dealt with the importance of shared tasks for the progress in medical NLP [25]. The last

two Yearbook surveys of the NLP section most closely related to medical IE were published in 2015 [26] and 2008 [27]. The survey by Velupillai et al. [28] dealt with opportunities and challenges of medical NLP for health outcomes research, with particular emphasis on evaluation criteria and protocols.

We also refer readers to alternative surveys of DL as applied to medical and clinical tasks. Wu et al. [29] reviewed literature for works using DL for a broader view of clinical NLP, whereas Xiao et al. [30] and Shickel et al. [31] performed systematic reviews on the applications of DL to several kinds of EHR data, not only text. Miotto et al. [32] and Esteva et al. [33] further extended that perspective to include clinical imaging and genomic data beyond the scope of classical EHRs. From an even broader perspective of the huge amounts of biomedical data, Ching et al. [34] examined various applications of DL to a variety of biomedical problems—patient classification, fundamental biological processes, and treatment of patients—and discussed the unique challenges that biomedical data pose for DL methods. In the same vein, Rajkomar et al. [35] used the entire EHR, including clinical free-text notes, for clinical predictive modeling based on DL (targeted, e.g., at the prediction of in-hospital mortality or patient’s final discharge diagnoses). They also demonstrated that DL methods outperformed traditional statistical prediction models.

2 Design and Goals of this Survey

In this survey, we concentrated on publications within the time window from 2017 to early 2020 and screened the contributions from three major scientific communities involved in medical IE:

- Medicine and medical informatics are covered by PubMed;
- Natural language processing is covered by the ACL Anthology, the digital library of the Association for Computational Linguistics;
- Neural networks are covered by the major conference series of the neural network community (Neural Information Processing Systems (NIPS/NeurIPS)) whereas the artificial intelligence community gets in via the Association for the Advancement of Artificial Intelligence (AAAI) Digital Library which keeps the records from the AAAI and IJCAI conferences.

We also included health-related publications from the digital libraries of the Association for Computing Machinery (ACM) and the Institute of Electrical and Electronics Engineers (IEEE). When necessary, we also referred to e-preprint archives such as arXiv.org, since they have become a new, increasingly important distribution channel for the most recent research results in computer science (yet, in that state typically without peer review) and thus foreshadow future directions of research.

We searched these literature repositories with a free-text query that can be approximated as follows: (information extraction OR text mining OR named entity recognition OR relation extraction) AND (deep learning OR neural network) AND (medic* OR clinic* OR health)

For this setting, we found approximately 1,000 unique publications, screened them for relevance, and, finally, included roughly 100 into this survey.

Table 1 An Overview of Common Embeddings—Biomedical Language Models

Model	Base model	Dimensions	Source corpus	Size (in words)
bio.nlpplab.org [16]	Word2vec	200	PubMed + PMC	5.5B
BioWordVec [17]	fastText + node2vec	200	PubMed + MeSH	3.7B
BioSentVec [18]	fastText	200	PubMed + MIMIC-III	4.4B + 539M
Flair('pubmed-X')	Flair	200	PubMed	(5% of PubMed in 2015)
SciBERT [19]	BERT _{BASE}	768	Semantic Scholar	3.1B
BioBERT [20]	BERT _{BASE}	768	PubMed + PMC	4.5B + 13.5B
ClinicalBERT [21]	BioBERT	768	MIMIC-III	539M
BlueBERT [22]	BERT _{BASE} /BERT _{LARGE}	768/1024	PubMed + MIMIC-III	>4B + >500M

3 Deep Neural Networks for Medical Information Extraction

In this section, we introduce applications of DNNs to medical NLP for two different tasks, Named Entity Recognition (NER) and Relation Extraction (REX). The focus of our discussion lies on studies dealing with English as reference language since the vast majority of benchmark and reference data sets are in English². After a brief description of each task, we summarize the current SOTA in tables which generalize often subtle distinctions in experimental design and workflows. Our main goal is to show the diversity of major benchmark datasets, DL approaches, and embeddings being used. For these tables, we extracted all symbolic (e.g., corpus or DL approach) and numerical information (e.g., about annotation metadata, performance scores) directly from the cited papers.

The assessment of different systems for the same task is centered around their performance on gold data in evaluation experiments. We refrain from highlighting minor differences in the reported scores because of different datasets being used for evaluation, changing volumes of metadata, and sometimes even the genres they contain. Hence, from a strict methodological perspective, the reported results have to be interpreted with utmost caution for two main reasons [37]. First, the choice of pre-processing steps, such as tokenization, inclusion/exclusion of punctuation marks, stop word removal, morphological normalization/lemmatization/stemming, n-gram variability, entity blinding strategies, and, second, the calibration of training methods (split bias, pooling techniques, hyperparameter selection (dropout rate, window size, etc.)) have a strong impact on the way a chosen embedding type and DL model finally performs, even within the same experimental setting. However, the data we report give valuable comparative information of the SOTA, though with fuzzy edges. This situa-

² Wu et al. [29, Table 3(b)] found that 71% of the corpora they screened were English, 20% Chinese, 2% Spanish, Japanese or Finnish and all other languages ranked below 1%. For a survey on medical NLP dealing explicitly with languages other than English, see [36].

tion might be remedied by a recently proposed common evaluation framework for biomedical NLP, the BLUE (Biomedical Language Understanding Evaluation) benchmark³ [22], which consists of five different biomedical NLP tasks (including NER and REX) with ten corpora (including BC5CDR, DDI, and i2b2 that also occur in the tables below), or the one proposed by Chauhan et al. [37]⁴ enabling a more lucid comparison of various training methodologies, pre-processing, modeling techniques, and evaluation metrics.

For the tables provided in the next subsections, we used the F_1 score as the main ordering criterion for the cited studies (from highest to lowest)⁵. We usually had to select among a large variety of experimental conditions (with different scores). The final choices we made were led by the criterion to favor comparability among all studies. This means that higher (and lower) outcomes may have been reported in the cited studies for varying experimental conditions. Still, the top-ranked system(s) in each of the following tables defines the current SOTA for a particular application.

3.1 Named Entity Recognition

The task of Named Entity Recognition (NER) is to identify crucial medical named entities (i.e., spans of concrete mentions of semantic types such as diseases or drugs and their attributes) in running text. For a recent survey of DL-based approaches and architectures underlying NER as a generic NLP application, see [38].

3.1.1 Diseases

A primary target of NER in the medical field is the automatic identification of diseases in scientific articles and clinical reports. For instance, textual occurrences of disease mentions (e.g., “*Diabetes II*” or “*cerebral inflammation*”) are mapped to a common

³ https://github.com/ncbi-nlp/BLUE_Benchmark

⁴ <https://github.com/geetickachauhan/relation-extraction>

⁵ We disregard here the common distinction between strict and partial matching; numbers given in the tables typically reflect the strongest condition, i.e., strict (complete) match between system prediction and gold standard data.

semantic type, *Disease*⁶. The crucial role of recognizing diseases in medical discourse is also emphasized by a number of surveys dealing with the recognition of special diseases. For instance, Sheikhalishahi et al. [40] discussed NLP methods targeted at chronic diseases and found that shallow ML and rule-based approaches (as opposed to more sophisticated DL-based ones) prevail. Koleck et al. [41] summarized the use of NLP to analyze symptom information documented in EHR free-text narratives as an indication of diseases and similar to the previous survey found little coverage of DL methods in this application area as well. Savova et al. [42] reviewed the current state of clinical NLP with respect to oncology and cancer phenotyping from EHR. Datta et al. [43] focused on an even more specialized use case—the lexical representation required for the extraction of cancer information from EHR notes in a frame-semantic format.

The research summarized in Table 2 is strictly focused on *Disease* recognition and, for reasons of comparability, based on the use of shared data sets and metadata (gold annotations). Two benchmarks are prominently featured, BC5CDR [44] and NCBI [45]⁷. BC5CDR is a corpus made of 1,500 PubMed articles, with 4,409 annotated chemicals, 5,818 diseases, and 3,116 chemical-disease interactions, created for the *BioCreative V Chemical and Disease Mention Recognition Task* [44]. As an alternative, the NCBI Disease Dataset [45] consists of a collection of 793 PubMed abstracts annotated with 6,892 disease mentions which are mapped to 790 unique disease concepts (thus, this corpus can also be used for grounding experiments).

⁶ Even more ambitious is the task of linking (or grounding) textual mentions and semantic types to unique identifiers of a given terminology or ontology (such as SNOMED-CT, ICD, or the Human Disease Ontology, <https://www.ebi.ac.uk/ols/ontologies/doid>), an issue we will not elaborate on in this survey, cf. e.g., [39].

⁷ Concrete numbers in the column “Number of Mentions,” indicating the number of named entity mentions (possibly split into training, development, and test set, if provided), may slightly differ for the same corpus because of data cleansing (e.g., removal of duplicates), different pre-processing (e.g., tokenization), and other version issues.

Table 2 Medical Named Entity Recognition: Diseases. Benchmark Datasets from BC5CDR [44] and NCBI [45].

Citations	Corpus	# of Mentions train (+ dev) / test	DL Approach	Embeddings (source)	F1-score %
Lee et al. [20]	NCBI	6,881	bidirectional Transformer	BioBERT v1.1 [20]: self-trained BERT on PubMed	89.7
Sachan et al. [47]	BC5CDR	5,818 (all diseases only)	(char) CNN + (word) BiLSTM-CRF	pre-trained embeddings + self-trained on PubMed	89.3
Wang et al. [46]	BC5CDR	12,852 (all mention types)	BiLSTM-CRF	pre-trained: [16]	88.8
Xu et al. [49]	NCBI	6,892	Att-BiLSTM-CRF + self-generated disease dictionary	word embeddings [50] self-trained on PubMed and PMC	88.6
Hong and Lee [51]	NCBI	6,881	BiLSTM + CRF: Unary & Pairwise Network transition model	pre-trained word embeddings (PubMed): [16, 52]	88.6
Beltagy et al. [19]	NCBI	—	bidirectional Transformer	SciBERT [19]: self-trained BERT on biomedical full texts from Semantic Scholar	88.6
Xu et al. [49]	BC5CDR	12,850 (all mention types)	Att-BiLSTM-CRF + self-generated disease dictionary	word embeddings [50] self-trained on PubMed and PMC	88.3
Zhao et al. [39]	BC5CDR	12,852 (all mention types)	CNN + BiLSTM(-CRF)	pre-trained: [16, 53–55]	87.6
Zhao et al. [39]	NCBI	6,881	CNN + BiLSTM(-CRF)	pre-trained: [16, 53–55]	87.4
Sachan et al. [47]	NCBI	6,892	(char) CNN + (word) BiLSTM-CRF	pre-trained embeddings + self-trained on PubMed	87.3
Lee et al. [20]	BC5CDR	12,694 (all mention types)	bidirectional Transformer	BioBERT V1.1 [20]: self-trained BERT on PubMed	87.2
Hong and Lee [51]	BC5CDR	12,852 (all mention types)	BiLSTM + CRF: Unary & Pairwise Network transition model	pre-trained word embeddings (PubMed): [16, 52]	87.2
Lou et al. [48]	BC5CDR	12,864 (all mention types)	non-DL: transition model	n/a	86.2
Wang et al. [46]	NCBI	6,881	BiLSTM-CRF	pre-trained: [16]	86.1
Lou et al. [48]	NCBI	6,892	non-DL: transition model	n/a	82.1

The current top performance for *Disease* recognition comes close to 90% F_1 ⁸. Lee et al. [20] use a Transformer model with in-domain training (BioBERT), but also (attention-based) BiLSTMs perform strongly in the range of 88–89% F_1 score. For the choice of embeddings being used, self-trained ones might be a better choice than pre-trained ones, *e.g.*, those provided by bio.nlplab.org [16]. The incorporation of (large) dictionaries does not provide a

competitive advantage in the experiments reported here. Though multi-task learning and transfer learning seem reasonable choices ([39, 46] and [47], respectively) to combat the sparsity of datasets, they generally do not boost systems to the top ranks.

Interesting though are differences for the *same* approach on *different* evaluation data sets. For the second-best system by Sachan et al. [47], F_1 scores differ for BC5CDR and NCBI by 2.0 (for the third-best [46] by 2.7) percentage points, whereas for the best non-DL approach by Lou et al. [48], this difference amounts to remarkable 4.1 percentage points. This hints at a strong dependence of the results of the same system set-up on

the specific corpus these results have been worked out and, thus, limits generalizability. On the other hand, corpora obviously cannot be blamed for intrinsic analytical hardness since cross-rankings occurs: the system by Lee et al. [20] gets the over-all highest F_1 score for NCBI but underperforms for BC5CDR, whereas for the tagger used by Sachan et al. [47] the ranking is reversed—their system performs better on BC5CDR than on NCBI (differences are in the range of 2 percentage points). The most stable system in this respect is the one by Zhao et al. [39]. Finally, the distance between the best- and second-best-performing DL systems ([20] and [47], respectively) and their best non-DL

⁸ Beltagy et al. [19] report an F_1 score of 90% on the BC5CDR corpus, but it remains unclear whether this result refers to the type *Disease*, *Drug*, or both of them.

Table 3 Medical Named Entity Recognition: Drugs. Benchmark Datasets: n2c2 [56]; i2b2 2009 [57]; MADE 1.0 [59]; DDI [60].

Citations	NE type	Corpus	# of Mentions train (+ dev) / test	DL Approach	Embeddings (source)	F1-score %
Wei et al. [62]	Drug	n2c2	16,225 / 10,575	BiLSTM-CRF	Word2vec embeddings self-trained on MIMIC-III [58]	95.6
Glagic et al. [63]	Drug	i2b2 2009	—	RNN	Word2vec embeddings self-trained on i2b2 2009 [57]	94.6
Zeng et al. [64]	Drug	DDI	11,260 / 3,689	BiLSTM-CRF	word & character embeddings pre-trained on Wikipedia	92.0
Unanue et al. [65]	Drug	DDI: DrugBank	9,715 / 180	BiLSTM-CRF	GloVe [53] pre-trained on CommonCrawl + GloVe self-trained on MIMIC-III [58]	91.8
Li et al. [66]	Drug	MADE 1.0	—	BiLSTM-CRF	pre-trained [16]	91.3
Wunnava et al. [67]	Drug	MADE 1.0	17,008 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	90.4
Dandala et al. [70]	Drug	MADE 1.0	13,507 / 2,395	BiLSTM-CRF	pretrained character & word embeddings	90.0
Tao et al. [71]	Drug	i2b2 2009	7,988 / 8,440	SVM	(as feature for SVM: GloVe [53] embeddings self-trained on MIMIC-III [58])	88.9
Chapman et al. [72]	Drug	MADE 1.0	13,508 / 2,395	CRF	n/a	88.6
Unanue et al. [65]	Drug	DDI: MEDLINE	1,574 / 171	BiLSTM-CRF	GloVe [53] pre-trained on Common Crawl + GloVe self-trained on MIMIC-III [58]	75.6

counterpart [48] amounts to 7.6 percentage points (for NCBI) and 3.1 percentage points (for BC5CDR), respectively.

3.1.2 Medication

The second major medical named entity type we here discuss is related to medication information. NER complexity is increased for this task since it is split into several subtasks, including the recognition of drug names (*Drug*), frequency (*Dr-Freq*) and (manner or) route of drug administration (*Dr-Route*), dosage (*Dr-Dose*), duration of administration (*Dr-Dur*), and adverse drug events (*Dr-ADE*). These subtypes are highly relevant in the context of medication information and are backed up by an international standard, the HL7 Fast Healthcare Interoperability Resources (FHIR)⁹. Tables 3 and 4 provide an overview of the SOTA on this topic.

⁹ See, e.g., the HL7 FHIR Medication Statement at <https://www.hl7.org/fhir/medicationstatement.html#MedicationStatement>

For medication information, four gold standards had a great impact on the field in the past years. The most recent one came out of the 2018 n2c2 Shared Task on Adverse Drug Events and Medication Extraction in Electronic Health Records [56], a successor of the 2009 i2b2 Medication Challenge [57], now with a focus on Adverse Drug Events (ADEs). It includes 505 discharge summaries (303 in the training set and 202 in the test set), which originate from the MIMIC-III clinical care database [58]. The corpus contains nine types of clinical concepts (including drug name), eight attributes (reason, ADE, frequency, strength, duration, route, form, and dosage – from which we chose five for comparison), and 83,869 concept annotations. Relations between drugs and the eight attributes were also annotated and summed up to 59,810 relation annotations (see Section 3.2.1). The third corpus, MADE 1.0 [59], formed the basis for the 2018 Challenge for Extracting Medication, Indication, and Adverse Drug Events (ADEs) from Electronic Health Record (EHR)

Notes and consists of 1,092 de-identified EHR notes from 21 cancer patients. Each note was annotated with medication information (drug name, dosage, route, frequency, duration), ADEs, indication (symptom as reason for drug administration), other signs and symptoms, severity (of disease/symptom), and relations among those entities, resulting in 79,000 mention annotations. Finally, the DDI corpus [60], originally developed for the Drug-Drug Interaction (DDI) Extraction 2013 Challenge [61], is composed of 792 texts selected from the (semi-structured) DrugBank database¹⁰ and other 233 (unstructured) MEDLINE abstracts, summing up 1,025 documents. This fine-grained corpus has been annotated with a total of 18,502 pharmacological substances and 5,028 drug-drug interactions¹¹. Hence,

¹⁰ <https://www.drugbank.ca/>

¹¹ The DDI corpus is actively maintained and enhanced leading to a large number of versions. Hence, comparisons based on DDI have to be carried out very carefully.

Table 4 Medical Named Entity Recognition: Medication Attributes. Benchmark Datasets: n2c2 [56]; i2b2 2009 [57]; MADE 1.0 [59]; DDI [60].

Citations	NE type	Corpus	# of Mentions train (+ dev) / test	DL Approach	Embeddings (source)	F1-score %
Wei et al. [62]	Dr-Freq	n2c2	6,281 / 4,012	BiLSTM-CRF	Word2vec embeddings self-trained on MIMIC-III [58]	97.5
Tao et al. [71]	Dr-Freq	i2b2	3,881 / 3,925	CRF	(as feature for CRF: GloVe [53] embeddings self-trained on MIMIC-III [58])	92.4
Gligic et al. [63]	Dr-Freq	i2b2 2009	—	RNN	Word2vec embeddings self-trained on i2b2 2009 [57]	90.9
Li et al. [66]	Dr-Freq	MADE 1.0	—	BiLSTM-CRF	pre-trained [16]	86.3
Dandala et al. [70]	Dr-Freq	MADE 1.0	4,147 / 659	BiLSTM-CRF	pretrained character & word embeddings	86.3
Chapman et al. [72]	Dr-Freq	MADE 1.0	4,147 / 659	CRF	n/a	85.8
Wunnava et al. [67]	Dr-Freq	MADE 1.0	5,050 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	84.3
Gligic et al. [63]	Dr-Route	i2b2 2009	—	RNN	Word2vec embeddings self-trained on i2b2 2009 [57]	96.9
Wei et al. [62]	Dr-Route	n2c2	5,476 / 3,513	BiLSTM-CRF	Word2vec embeddings self-trained on MIMIC-III [58]	95.6
Tao et al. [71]	Dr-Route	i2b2	3,052 / 3,299	SVM	(as feature for SVM: GloVe [53] embeddings self-trained on MIMIC-III [58])	94.1
Wunnava et al. [67]	Dr-Route	MADE 1.0	2,862 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	92.4
Chapman et al. [72]	Dr-Route	MADE 1.0	2,278 / 389	CRF	n/a	92.1
Li et al. [66]	Dr-Route	MADE 1.0	—	BiLSTM-CRF	pre-trained [16]	91.9
Dandala et al. [70]	Dr-Route	MADE 1.0	2,278 / 389	BiLSTM-CRF	pretrained character & word embeddings	91.7
Wei et al. [62]	Dr-Dose	n2c2	4,221 / 2,681	BiLSTM-CRF	Word2vec embeddings self-trained on MIMIC-III [58]	94.8
Gligic et al. [63]	Dr-Dose	i2b2 2009	—	RNN	Word2vec embeddings self-trained on i2b2 2009 [57]	93.0
Tao et al. [71]	Dr-Dose	i2b2 2009	4,132 / 4,371	CRF	(as feature for CRF: GloVe [53] embeddings self-trained on MIMIC-III [58])	91.5
Wunnava et al. [67]	Dr-Dose	MADE 1.0	5,978 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	88.0
Li et al. [66]	Dr-Dose	MADE 1.0	—	BiLSTM-CRF	pre-trained [16]	88.0
Chapman et al. [72]	Dr-Dose	MADE 1.0	4,893 / 801	CRF	n/a	87.5
Dandala et al. [70]	Dr-Dose	MADE 1.0	4,893 / 801	BiLSTM-CRF	pretrained character & word embeddings	87.1
Wei et al. [62]	Dr-Dur	n2c2	592 / 378	BiLSTM-CRF	Word2vec embeddings self-trained on MIMIC-III [58]	86.2
Li et al. [66]	Dr-Dur	MADE 1.0	—	BiLSTM-CRF	pre-trained [16]	77.6
Wunnava et al. [67]	Dr-Dur	MADE 1.0	926 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	76.7
Dandala et al. [70]	Dr-Dur	MADE 1.0	765 / 133	BiLSTM-CRF	pretrained character & word embeddings	75.0
Chapman et al. [72]	Dr-Dur	MADE 1.0	765 / 133	CRF	n/a	72.2
Gligic et al. [63]	Dr-Dur	i2b2 2009	—	RNN	Word2vec embeddings self-trained on i2b2 2009 [57]	63.0
Tao et al. [71]	Dr-Dur	i2b2	508 / 499	CRF	(as feature for CRF: GloVe [53] embeddings self-trained on MIMIC-III [58])	61.8
Wunnava et al. [67]	Dr-ADE	MADE 1.0	1,807 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	63.5
Yang et al. [73]	Dr-ADE	MADE 1.0	1,807 /	BiLSTM-CRF	pre-trained on Wikipedia, EHR notes, and PubMed [68, 69]	60.5
Dandala et al. [70]	Dr-ADE	MADE 1.0	1,509 / 431	BiLSTM-CRF	pretrained character & word embeddings	57.7
Li et al. [66]	Dr-ADE	MADE 1.0	—	BiLSTM-CRF	pre-trained [16]	55.4
Wei et al. [62]	Dr-ADE	n2c2	959 / 625	BiLSTM-CRF	Word2vec embeddings self-trained on MIMIC-III [58]	53.0
Chapman et al. [72]	Dr-ADE	MADE 1.0	1509 / 431	CRF	n/a	51.1

the medication NER task not only comes with a higher entity type complexity but also with text genres different from the disease recognition task—while the former puts emphasis on clinical reports, the latter focuses on scholarly writing.

Except for route and ADE, all top scores for NER were achieved on the n2c2 corpus. For drug names, the current SOTA exceeds 95% F_1 score established by Wei et al. [62]. As to the subtypes, their system also compares favorably to alternative architectures by a large F_1 margin ranging from 8.6 percentage points (for duration) down to 1.0 (for drug name). For route, the distance to the best system is marginal (around 1 percentage point)¹², whereas for ADE it is huge (more than 10 percentage points, a strong outlier). Overall, frequency, route, and dosage recognition reach outstanding F_1 scores in the range of 95 up to 97%, while for duration information top F_1 scores drop remarkably by at least 10 to 20 percentage points. Still, the recognition of ADEs seems to be the hardest task, with the best system by Wunna et al. [67] peaking at around 64% F_1 on MADE 1.0 data (here the top performing system by Wei et al. [62] plummets down to 53% F_1). Interestingly, ADEs are verbally the least constrained type of natural language utterance compared with all the other entity types considered here.

In terms of DL methodology, BiLSTM-CRFs are the dominating approach. Yet, the type of embeddings used by different DL systems varies a lot ranging from pre-trained Word2vec embeddings and those self-trained on MIMIC-III (for the top performers) to GloVe embeddings pre-trained on CommonCrawl, Wikipedia, EHR notes, and PubMed. There seems to be no generalizable winner for either choice of embeddings given the current state of evaluations, but self-training on medical raw data, such as MIMIC-III, challenge data sets, or, more advisable, using the now available BioSentVec [18] and BlueBERT [22] embeddings pre-trained on MIMIC-III, might be advantageous.

¹² Interestingly, the transfer learning approach advocated by Gligic et al. [63] performs well for some medication NER tasks, but fails to deliver competitive results for the medication relation task (cf. Table 5).

Studies in which the same system configuration was tested on different corpora are still lacking so that corpus effects are unknown (unlike for diseases; see Table 2). Yet, there is one interesting though not so surprising observation: Unanue et al. [65] explored the two slices of the DDI corpus, with a span of F_1 scores of more than 16 percentage points. This obviously witnesses the influence of *a priori* (lack of) structure—DrugBank data is considerably more structured than MEDLINE free texts and, thus, the former gets much higher scores than the latter.

Comparing DL approaches vs. non-DL ones (a CRF architecture) on the same corpus (MADE 1.0), we found that for the core entity type (Drug), the recognition performance differs by almost 3 percentage points, for frequency, route and dose marginally by less than 1, yet for duration and ADE it amounts to roughly 5 and 12 percentage points, respectively—consistently in favor of Deep Neural Networks (DNNs).

3.2 Relation Extraction

Once named entities have been identified, a follow-up question emerges: does some sort of semantic relation hold among these entities? We surveyed this Relation Extraction (REX) task with reference to results that have been achieved for information related to medication attributes and drug-drug interaction.

3.2.1 Medication-Attribute Relations

In Section 3.1.2, we already dealt with single named entity types typically associated with medication information, namely drug names and administration frequency, duration, dosage, route, and ADE, yet in isolated form only. In this subsection, we are concerned with making the close associative ties between *Drugs* and typical conceptual attributes, such as *Frequency*, *Duration*, *Dosage*, *Route*, *ADE*, and *Reason* (for prescription), explicit. Hence, the recognition of the respective named entity types (*Drugs*, *Dr-Freq*, *Dr-Dur*, *Dr-Dose*, *Dr-Route*, *Dr-ADE*, and *Dr-Reason*) turns out to be a good starting point for solving this REX task. Not

surprisingly, the benchmarks for this task are a subset of the ones in Tables 3 and 4 depicting the results for medication-related NER. Table 5 provides an overview of the experimental results for finding medication-attribute relations in medical, in effect, clinical, documents.

The overall results from medication-focused NER are mostly confirmed for the REX task. The n2c2 corpus is the reference dataset for top performance. The group who achieved top F_1 scores for the medication NER problem also performed best for the medication-attribute REX task [62], with extraordinary figures for *Frequency*, *Route*, and *Dosage* relations (in the upper 98% F_1 range), a superior one for the *Duration* relation (93% F_1), and good ones on the (hard to deal with) *Adverse* and *Reason* relations (85% F_1). Still, the distances to the second-best system for the same corpus (n2c2) are not so pronounced in most cases, ranging by 1 percentage point (for *Frequency*, *Route*, *Dosage*, and *Duration*), yet increased up to 3 (for *Adverse*) and 7 (for *Reason*) percentage points.

For the MADE 1.0 corpus, a similar picture emerges. From a lower offset (typically around 3 F_1 percentage points compared with n2c2), differences between the best and second-best systems were on the order of (negligible) 1 percentage point for *Frequency*, *Route*, and *Dosage*, yet increased by roughly 3, 5, and 7 percentage points for *Reason*, *Duration*, and *Adverse events*, respectively. Yet, in 4 out of 6 cases (*Frequency*, *Dosage*, *Duration*, and *Adverse events*) non-DL systems (CRFs, SVMs) outperformed their DL counterparts with small margins (in the range of (again, negligible) 1 percentage point) for *Frequency* and *Dosage*, yet with higher ones for *Duration* and *Adverse events* (5 and 7 percentage points, respectively). In cases where the DL approach ranked higher than a non-DL one, differences ranged between 1 and 3 percentage points (for *Route* and *Reason*, respectively). Thus, the MADE 1.0 corpus constitutes a benchmark where well-engineered standard ML classifiers can still play a competitive role. However, we did not find this pattern of partial supremacy of non-DL approaches for the n2c2 benchmark.

The top performers for the medication attribute REX task [62] employed a joint learning approach based on CNN-RNN

Table 5 Medical Relation Extraction: Medication-Attribute Relations (including ADEs). Benchmark Datasets: n2c2 [56]; MADE 1.0 [59].

Citations	Relation type	NE type 1	NE type 2	Corpus	# of Mentions train (+ dev) / test	DL Approach	Embeddings (source)	F1-score %
Wei et al. [62]	Frequency	Drug	Dr-Freq	n2c2	(6,310 / 4,034)	CNN-RNN	Word2vec embeddings self-trained on MIMIC-III [58]	98.7
Christopoulou et al. [74]	Frequency	Drug	Dr-Freq	n2c2	4,986 (+ 1,312) /	Att-BiLSTM-CRF + Transformer Network	word and relative position embeddings	97.6
Chapman et al. [72]	Frequency	Drug	Dr-Freq	MADE 1.0	4,419 / 730	non-DL: CRF	n/a	94.7
Dandala et al. [70]	Frequency	Drug	Dr-Freq	MADE 1.0	4,419 / 730	Att-BiLSTM	pretrained character & word embeddings	93.7
Wei et al. [62]	Route	Drug	Dr-Route	n2c2	(5,538 / 3,546)	CNN-RNN	Word2vec embeddings self-trained on MIMIC-III [58]	98.6
Christopoulou et al. [74]	Route	Drug	Dr-Route	n2c2	4,327 (+ 1,208) /	Att-BiLSTM-CRF + Transformer Network	word and relative position embeddings	97.8
Dandala et al. [70]	Route	Drug	Dr-Route	MADE 1.0	2,551 / 455	Att-BiLSTM	pretrained character & word embeddings	95.3
Chapman et al. [72]	Route	Drug	Dr-Route	MADE 1.0	2,551 / 455	non-DL: CRF	n/a	94.1
Wei et al. [62]	Dosage	Drug	Dr-Dose	n2c2	(4,225 / 2,695)	CNN-RNN	Word2vec embeddings self-trained on MIMIC-III [58]	98.6
Christopoulou et al. [74]	Dosage	Drug	Dr-Dose	n2c2	3,299 (+ 921) /	Att-BiLSTM-CRF + Transformer Network	word and relative position embeddings	98.0
Chapman et al. [72]	Dosage	Drug	Dr-Dose	MADE 1.0	5,177 / 866	non-DL: CRF	n/a	96.0
Dandala et al. [70]	Dosage	Drug	Dr-Dose	MADE 1.0	5,177 / 866	Att-BiLSTM	pretrained character & word embeddings	95.2
Wei et al. [62]	Duration	Drug	Dr-Dur	n2c2	(643 / 426)	CNN-RNN	Word2vec embeddings self-trained on MIMIC-III [58]	92.9
Chapman et al. [72]	Duration	Drug	Dr-Dur	MADE 1.0	906 / 147	non-DL: CRF	n/a	92.4
Christopoulou et al. [74]	Duration	Drug	Dr-Dur	n2c2	518 (+ 124) /	Att-BiLSTM-CRF + Transformer Network	word and relative position embeddings	91.7
Dandala et al. [70]	Duration	Drug	Dr-Dur	MADE 1.0	906 / 147	Att-BiLSTM	pretrained character & word embeddings	87.8
Wei et al. [62]	Adverse	Drug	Dr-ADE	n2c2	(1,107 / 733)	CNN-RNN	Word2vec embeddings self-trained on MIMIC-III [58]	85.0
Christopoulou et al. [74]	Adverse	Drug	Dr-ADE	n2c2	891 (+ 216) /	Att-BiLSTM-CRF + Transformer Network	word and relative position embeddings	78.3
Chapman et al. [72]	Adverse	Drug	Dr-ADE	MADE 1.0	2,082 / 530	non-DL: CRF	n/a	73.1
Dandala et al. [70]	Adverse	Drug	Dr-ADE	MADE 1.0	2,082 / 530	Att-BiLSTM	pretrained character & word embeddings	66.0
Wei et al. [62]	Reason	Drug	Dr-Reason	n2c2	(5,169 / 3,410)	CNN-RNN	Word2vec embeddings self-trained on MIMIC-III [58]	84.9
Christopoulou et al. [74]	Reason	Drug	Dr-Reason	n2c2	4,069 (+ 1,090) /	Att-BiLSTM-CRF + Transformer Network	word and relative position embeddings	82.2
Dandala et al. [70]	Reason	Drug	Dr-Reason	MADE 1.0	4,554 / 876	Att-BiLSTM	pretrained character & word embeddings	80.9
Yang et al. [73]	Reason	Drug	Dr-Reason	MADE 1.0	4,530 / 871	non-DL: SVM	n/a	77.9

(thus diverging from the most successful architectures for medication NER; see Tables 3 and 4) and rule-based post-processing that outperformed a simple CNN-RNN. Summarizing, the CNN-RNN approach seems more favorable than an (attention-based) BiLSTM, with preferences for self-trained in-domain embeddings.

3.2.2 Drug-Drug Interaction

The second type of medication-focused relation we consider here are drug-drug interactions as featured in the DDI challenge (for surveys on the impact of DL on recent research on drug-drug interactions, cf. [82, 83], for a survey on drug-drug interaction combining progress in data and text mining from EHRs, scientific papers, and clinical reports but lacking in-depth coverage of DL methods, cf. [84], for the NLP-focused recognition of ADEs also lacking awareness of DL contributions to this topic, cf. [85]). Four main types of relations between drugs are considered: pharmacokinetic *Mechanism*, drug *Effect*, recommendation or *Advice* regarding a drug interaction, and *Interaction* between drugs without providing any additional information. Overall, the DDI corpus on which these evaluations were run is divided into 730 documents taken from DrugBank and 175 abstracts from MEDLINE and contains 4,999 relation annotations (4,020 train, 979 test).

Recognition rates for these relations (cf. Table 6) are considerably lower than for the medication-related attributes when linked to drugs (cf. Table 5). The best systems peak at 85% F_1 score for *Advice* (a distance of more than 13 percentage points to the top recognition results for medication-attributes), they slip to 78%¹³ and 77% for *Mechanism* and *Effect*, respectively, and plummet to 59% for *Interaction*¹⁴. Differences between the first

and second-ranked systems are typically small, yet become larger on subsequent ranks (roughly between 3 to 4 percentage points relative to the top-ranked system). As with medication attributes, drug-drug interactions can also be recognized in a competitive way by CNN-RNN architectures, but attention-based LSTMs perform also considerably well. Again, self-trained embeddings using in-domain corpora seem to be advantageous for this relation class. Reflecting the drop in performance, one may conclude that drug-drug interactions constitute a markedly harder task than the conceptually much closer medication-attribute relations.

Finally, Table 6 most drastically supports our claim that DL approaches outperform non-DL ones. The difference between both approaches amounts to 5 percentage points for *Mechanism*, 7 for *Effect* and *Interaction*, and 8 for *Advice*.

4 Conclusions

We have presented various forms of empirical evidence that (with one exception only) Deep Learning-based neural networks outperform non-DL, feature engineered, approaches for several information extraction tasks. However, despite their success, Deep Neural Networks and their embedding models have their shortcomings as well.

One of the most problematic issues is their dependence on huge amounts of training data: SOTA embedding models are currently trained on hundreds of billions of tokens [89]. This magnitude of data volume is out of reach for any training effort in the medical/clinical domain [90]. Also, embeddings are very vulnerable to malicious attacks or adversarial examples—small changes at the input level may result in severe misclassification [5]. Another well-known problem relates to the instability of

word embeddings. Word embeddings depend on their random initialization and the processing order of the underlying examples and therefore they do not necessarily converge on exactly the same embeddings even after several thousands of training iterations [91, 92]. Finally, although DL is celebrated for not requiring manual feature engineering, the effects of proper hyperparameter tuning on DNNs [93] remain an issue for DL [94]. Apart from these intrinsic problems, Kalyan and Sangeetha [95] and Khattak et al. [96] refer to extrinsic drawbacks of neural networks, such as opaque encodings (resulting in lacking interpretability) or limited transferability of large models (hindering knowledge distillation for smaller models).

Still, the sparsity of corpora and special linguistic phenomena of the medical (clinical) sublanguage(s) create intrinsic problems for data-greedy DL approaches that have to be overcome by special learning strategies for neural systems, such as transfer learning or domain adaptation. Research on adapting general language models to medical language constraints is just in its beginning. Yet, there is no simple solution to this problem. Wang et al. [97] evaluated Word2vec embeddings trained on private clinical notes, PMC, Wikipedia, and the Google News corpus both qualitatively and quantitatively and showed that the ones trained on Electronic Health Record data performed better on most of the tested scenarios. However, they also found that word embeddings trained on biomedical domain corpora do not necessarily have better performance than those trained on general domain corpora for any downstream biomedical NLP task (other experimental evidence of the effects of in- and out-of-domain corpora and further parameters, such as corpus size, on word embedding performance is reported by Lai et al., [98]).

While this survey focused on the application domain of medical IE to demonstrate the outstanding role of DL for medical Natural Language Processing, one might be tempted to generalize this trend to other applications as well. There is, indeed, plenty of evidence in the literature that other application fields, such as question answering (and the closely related area of machine reading), summarization, machine translation, and speech pro-

¹³ Xu et al. [86] even reach slightly more than 79% F_1 score for *Mechanism* (using UMLS-based concept embeddings with a Bi-LSTM approach), but substantially fall below the results for the other three relation types in comparison with all the systems mentioned in Table 6.

¹⁴ Dewi et al. [87] and Sun et al. [88] report on 86.3% and 84.5% F_1 scores, respectively, for the overall relation classification task both using a multi-layered CNN archi-

ture, yet unfortunately fail to provide details on each of the four single relations under scrutiny here. Both results exceed the overall result of the best-performing system depicted in Table 6 [76] (77.3%) by a large margin of 9 and 7 percentage points, respectively.

Table 6 Medical Relation Extraction: Drug-Drug Interaction. Benchmark Dataset: DDI [60].

Citation	Relation type	NE type 1	NE type 2	Corpus	# of Mentions train (+dev) / test	DL Approach	Embeddings (source)	F1-score %
Sun et al. [75]	Mechanism	Drug	Drug	DDI	1,319 / 299	Recurrent hybrid CNN with focal loss function	pre-trained on PubMed & Wikipedia: [16] + semantic & position embeddings self-trained on DDI	78.3
Zheng et al. [76]	Mechanism	Drug	Drug	DDI	1,319 / 302	Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed + DDI corpus	77.5
Wang et al. [77]	Mechanism	Drug	Drug	DDI	1,319 / 302	BiLSTM	word embeddings [78] self-trained on PubMed + dependency layers	75.4
Lim et al. [79]	Mechanism	Drug	Drug	DDI	1,260 / 301	binary tree-LSTM	pre-trained on PubMed & Wikipedia: [16]	75.1
Zhang et al. [80]	Mechanism	Drug	Drug	DDI	1,319 / 302	hierarchical Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed	74.0
Raihani and Laachfoubi [81]	Mechanism	Drug	Drug	DDI	1,319 / 302	non-DL: SVM	n/a	73.5
Zheng et al. [76]	Effect	Drug	Drug	DDI	1,687 / 360	Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed + DDI corpus	76.6
Sun et al. [75]	Effect	Drug	Drug	DDI	1,669 / 360	Recurrent hybrid CNN with focal loss function	pre-trained on PubMed & Wikipedia: [16] + semantic & position embeddings self-trained on DDI	73.5
Lim et al. [79]	Effect	Drug	Drug	DDI	1,592 / 357	binary tree-LSTM	pre-trained on PubMed & Wikipedia: [16]	72.9
Zhang et al. [80]	Effect	Drug	Drug	DDI	1,687 / 360	hierarchical Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed	71.8
Wang et al. [77]	Effect	Drug	Drug	DDI	1,687 / 360	BiLSTM	word embeddings [78] self-trained on PubMed + dependency layers	69.5
Raihani and Laachfoubi [81]	Effect	Drug	Drug	DDI	1,687 / 360	non-DL: SVM	n/a	69.5
Zheng et al. [76]	Advice	Drug	Drug	DDI	826 / 221	Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed + DDI corpus	85.1
Lim et al. [79]	Advice	Drug	Drug	DDI	814 / 221	binary tree-LSTM	pre-trained on PubMed & Wikipedia: [16]	82.7
Wang et al. [77]	Advice	Drug	Drug	DDI	826 / 221	BiLSTM	word embeddings [78] self-trained on PubMed + dependency layers	80.9
Sun et al. [75]	Advice	Drug	Drug	DDI	822 / 221	Recurrent hybrid CNN with focal loss function	pre-trained on PubMed & Wikipedia: [16] + semantic & position embeddings self-trained on DDI	80.5
Zhang et al. [80]	Advice	Drug	Drug	DDI	826 / 221	hierarchical Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed	80.3
Raihani and Laachfoubi [81]	Advice	Drug	Drug	DDI	826 / 221	non-DL: SVM	n/a	77.4

Table 6 Medical Relation Extraction: Drug-Drug Interaction. Benchmark Dataset: DDI [60]. (continued)

Citation	Relation type	NE type 1	NE type 2	Corpus	# of Mentions train (+ dev) / test	DL Approach	Embeddings (source)	F1-score %
Sun et al. [75]	Interaction	Drug	Drug	DDI	188 / 96	Recurrent hybrid CNN with focal loss function	pre-trained on PubMed & Wikipedia: [16] + semantic & position embeddings self-trained on DDI	58.9
Zheng et al. [76]	Interaction	Drug	Drug	DDI	188 / 96	Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed + DDI corpus	57.7
Zhang et al. [80]	Interaction	Drug	Drug	DDI	188 / 96	hierarchical Att-BiLSTM	word embeddings self-trained on Drug subset of PubMed	54.3
Raihani and Laachfoubi [81]	Interaction	Drug	Drug	DDI	188 / 96	non-DL: SVM	n/a	52.3
Wang et al. [77]	Interaction	Drug	Drug	DDI	188 / 96	BiLSTM	word embeddings [78] self-trained on PubMed + dependency layers	51.0
Lim et al. [79]	Interaction	Drug	Drug	DDI	188 / 92	binary tree-LSTM	pre-trained on PubMed & Wikipedia: [16]	43.9

cessing, reveal the same pattern. However, for text categorization (in the sense of mapping free text to some pre-defined medical category system, such as ICD, SNOMED, or MeSH) this preference is less obvious, since traditional Machine Learning or rule-based models still play an important role here and, more often than for the IE application scenario, show competitive performance against DL approaches. Whether this exception will persist or will be swept away by future research remains an open issue.

Acknowledgments

The first author was partially funded by the German Bundesministerium für Bildung und Forschung (BMBF) within the SMITH project under grant no. 01ZZ1803G. We thank all six reviewers for their insightful and helpful comments.

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