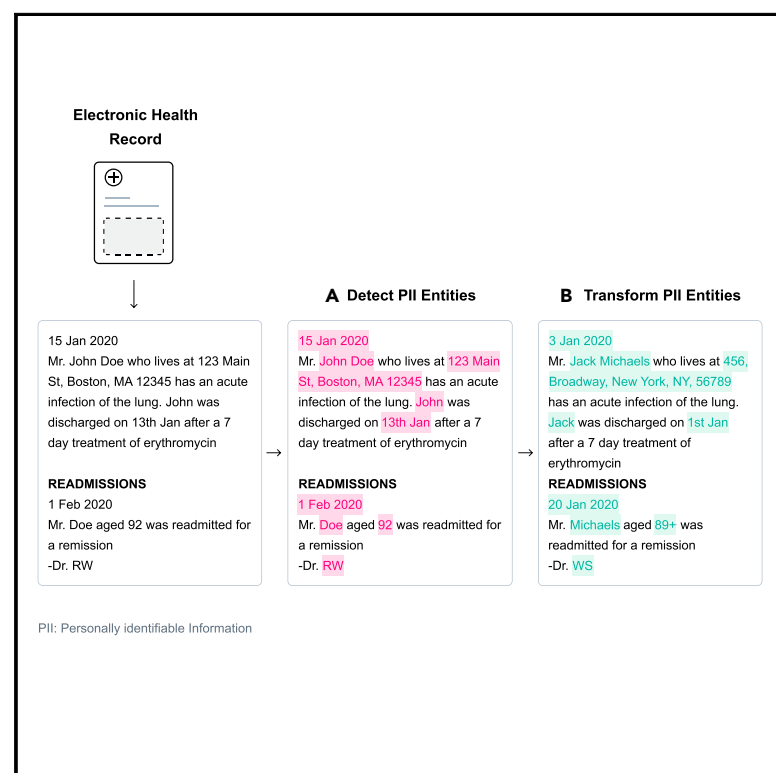


Patterns

Building a best-in-class automated de-identification tool for electronic health records through ensemble learning

Graphical abstract



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In brief

Physician notes contain valuable information about patient health and treatment, but their broad reuse is constrained by the presence of personally identifiable information to protect patient confidentiality. Our approach automatically detects and hides these identifiers in plain sight by replacing them with suitable surrogates. The de-identification system presented outperforms other existing tools, allowing for the generation of de-identified patient data at the scale necessary to help accelerate medical discovery.

Highlights

- An ensemble approach to automated de-identification of unstructured clinical text
- Our approach leverages advances in deep learning along with heuristics
- Detected personally identifiable information is replaced with suitable surrogates
- Patient data are de-identified at scale to accelerate medical discovery

Article

Building a best-in-class automated de-identification tool for electronic health records through ensemble learning

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THE BIGGER PICTURE Clinical notes in electronic health records convey rich historical information regarding disease and treatment progression. However, this unstructured text often contains personally identifiable information such as names, phone numbers, or residential addresses of patients, thereby limiting its dissemination for research purposes. The removal of patient identifiers, through the process of de-identification, enables sharing of clinical data while preserving patient privacy. Here, we present a best-in-class approach to de-identification, which automatically detects identifiers and substitutes them with fabricated ones. Our approach enables de-identification of patient data at the scale required to harness the unstructured, context-rich information in electronic health records to aid in medical research and advancement.



Production: Data science output is validated, understood, and regularly used for multiple domains/platforms

SUMMARY

The presence of personally identifiable information (PII) in natural language portions of electronic health records (EHRs) constrains their broad reuse. Despite continuous improvements in automated detection of PII, residual identifiers require manual validation and correction. Here, we describe an automated de-identification system that employs an ensemble architecture, incorporating attention-based deep-learning models and rule-based methods, supported by heuristics for detecting PII in EHR data. Detected identifiers are then transformed into plausible, though fictional, surrogates to further obfuscate any leaked identifier. Our approach outperforms existing tools, with a recall of 0.992 and precision of 0.979 on the i2b2 2014 dataset and a recall of 0.994 and precision of 0.967 on a dataset of 10,000 notes from the Mayo Clinic. The de-identification system presented here enables the generation of de-identified patient data at the scale required for modern machine-learning applications to help accelerate medical discoveries.

INTRODUCTION

The widespread adoption of electronic health records (EHRs) by health care systems has enabled digitization of patient health journeys. While the structured elements of EHRs (e.g., health insurance billing codes) have been relied upon to support the busi-

ness of health care and front office applications for decades, the unstructured text (e.g., history and physical notes and pathology reports) contains far richer and nuanced information about patient care, supporting novel research.^{1–5} However, this text often contains personally identifiable information (PII) as defined in the Health Insurance Portability and Accountability Act of 1996

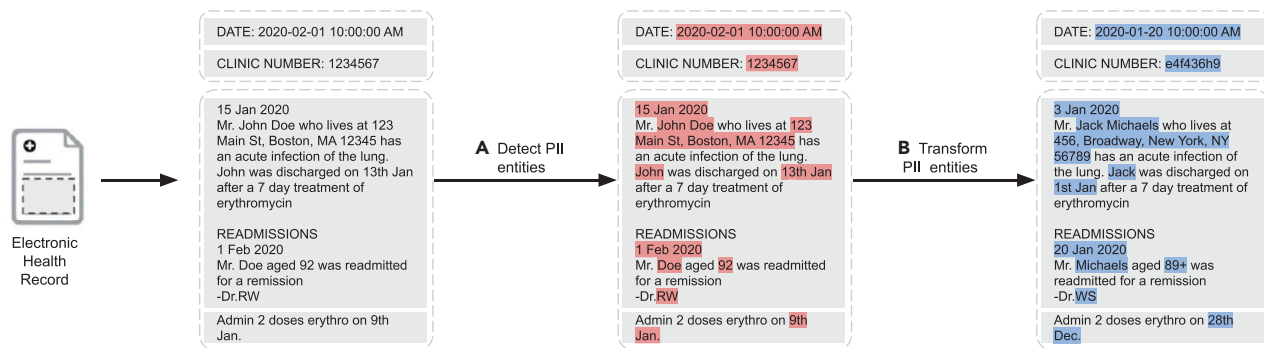


Figure 1. Automated de-identification of electronic health records

Two steps in automated de-identification of EHRs: (A) detecting PII entities and (B) transforming them by replacement with suitable surrogates. PII, personally identifiable information.

(HIPAA), such as personal name, phone number, or residential address.⁶ As a consequence, such data have limited reuse for secondary purposes.⁷ HIPAA permits data derived from EHRs to be widely shared and used when it is de-identified. Under the HIPAA Privacy Rule, de-identification can be accomplished in several ways. The most straightforward is the Safe Harbor implementation, which necessitates removal of an enumerated list of 18 categories of direct (e.g., Social Security number) and quasi-identifiers (e.g., date of service).

Implementing a scalable method for de-identification has several competing requirements. First, from a regulatory perspective, it must achieve extremely high recall, in that it needs to detect nearly all instances of PII. Second, from a clinical utility perspective, it must achieve extremely high precision, so that we maximize the correctness of the biomedical research performed. And, third, the approach needs to be cost effective, so that millions of records can be de-identified in a reasonable amount of time. The traditional approach of manual detection of PII is expensive, time consuming, and prone to human error,^{8,9} which makes automated de-identification a more promising alternative.^{10,11}

Several recent advances in natural language processing (NLP) have created an opportunity to build accurate and scalable automated de-identification systems. First, transfer learning of autoregressive and autoencoder models¹² for a supervised task such as named entity recognition (NER) requires very few labeled data, reducing human effort and error. Second, attention-based deep-learning models, such as transformers,¹³ allow for the non-sequential processing of text and enable the generation of rich contextualized word representations. Third, semantic segmentation algorithms generate a subword-based vocabulary,^{14,15} which can capture out-of-vocabulary words. Finally, the traditional transformer architecture has been improved upon through bidirectional encoder representations from transformers (BERT)¹⁶ and similar technologies that jointly train a *masked language model* (MLM) pre-training objective and a *next sentence prediction* task. BERT has set the stage for learning context-independent representations of terms in text and training context-sensitive models that transform those representations into context-aware representations based on the occurrence of a term in a sentence. We leverage these advances to support de-identification, which we formulate as an NER problem.

In this paper, we integrate a collection of approaches, blending the beneficial aspects of modern deep learning with rules and heuristics, to create a best-in-class approach to automated de-identification. The system transforms each detected PII instance into a suitable surrogate to mitigate the risk that any residual PII can be used to re-identify patients (Figure 1). The inference de-identification tool can be accessed at <https://academia.nferx.com/deid/>.

RESULTS

We first compare the performance of the inference de-identification system with other methods on the i2b2 2014 dataset.¹⁷ The resulting models are evaluated using precision, recall, and F1 scores (formulation provided in the [supplemental methods](#)) for NER on several groups of PII as defined in Table 1. We then compare the performance of these models on a substantially larger and diverse dataset from the Mayo Clinic and perform a deeper dive into the types of errors, distribution of errors per physician note, and distribution of errors per note type. It should be noted that this analysis focuses solely on the performance of detecting PII instances and does not address the risk of re-identification based on the semantics of any instances that the system fails to detect, an issue that is beyond the scope of this study.

Performance on the 2014 i2b2 de-identification dataset

The i2b2 2014 De-identification and Heart Disease Risk Factors challenge¹⁷ is a publicly available dataset of clinical documents with annotated PII elements. This dataset consists of a training set of 792 clinical notes and a test set of 515 clinical notes.

We compared the performance of our approach on the 2014 i2b2 test set with six other established de-identification tools: the method proposed by Dernoncourt et al. that blends conditional random fields (CRFs) and artificial neural networks (ANNs),¹⁸ Scrubber,¹⁹ Physionet,⁸ Philter,²⁰ MIST,²¹ and NeuroNER.²²

The results are provided in Table 2. First, we cite the CRF and ANN approach (CRF + ANN)¹⁸ scores against the group A entities (HIPAA only) as reported in their paper. We also directly report the results for Scrubber, Physionet, and Philter from prior publications²⁰ without performing an empirical analysis because the dataset (2014 i2b2) and the set of PII entities are the same as

Table 1. The list of entities covered by each group of direct and quasi-identifiers

Group name	Included entities
A (entities to be detected for a HIPAA Safe Harbor implementation)	(1) age over 89, (2) phone/fax numbers, (3) email addresses, (4) websites and URLs, (5) IP addresses, (6) dates, (7) Social Security number, (8) medical record numbers, (9) vehicle/device numbers, (10) account/certificate/license numbers, (11) health plan number, (12) street address, (13) city, (14) ZIP code, (15) employer name, and (16) personal names of patients and family members
B	Group A and (17) provider (doctor/nurse) names, (18) user IDs (of care providers)
C	Group B and (19) health care organization/facility names, (20) country, (21) state

It should be noted that groups B and C encompass entities beyond HIPAA Safe Harbor.

that used in our investigation. We trained MIST using sentences from the i2b2 training corpus (see [supplemental methods](#) and [Table S3](#)). We downloaded and used a pre-trained model for NeuroNER (see [supplemental methods](#)). We present the performance of these methods on group B (see [Table 1](#)) entities, which we use as the basis of our comparison.

We present two versions of the nference system. The first version was fine-tuned only on Mayo data and did not utilize any characteristics of the i2b2 training data. When evaluated with group B, this model achieved a precision, recall, and F1 score of 0.961, 0.988, and 0.974, respectively. The second version of our system involved fine-tuning our model with sentences from the i2b2 training set. We could not incorporate inclusion lists and sentence templates associated with the i2b2 data since the dataset is small (see [“methods”](#) for details). The precision, recall, and F1 score increased to 0.979, 0.992, and 0.985, respectively. Precision and recall per identifier type are provided in [Table S4](#).

Performance on the Mayo test dataset

The Mayo Clinic dataset consisted of 10,000 randomly sampled notes from a corpus of 104 million notes corresponding to 477,000 patients' EHR records.

The evaluation performed on the Mayo test dataset was based on identifiers defined by group C since this group best represented the distribution of PII in the dataset. The performance of the de-identification methods (in terms of precision, recall, and F1) are presented in [Table 3](#). The nference method performed best, with precision, recall, and F1 scores of 0.967, 0.994, and 0.979, respectively. Compared with the performance on the i2b2 dataset, we see improved recall (increase of 0.01) and a reduced precision value (decrease of 0.021). NeuroNER achieves precision, recall, and F1 scores of 0.928, 0.933, and 0.931, respectively. The F1 scores of Scrubber, Physionet, and Philter were lower than those achieved on the i2b2 dataset. Among these three methods, Philter demonstrates a relatively

Table 2. Performance of de-identification methods on the 2014 i2b2 test corpus

Method name	Group	Precision	Recall	F1	Basis of results
CRF + ANN (Dernoncourt et al.)	A	0.979	0.978	0.978	Dernoncourt et al. ¹⁸
Physionet	B	0.894	0.698	0.784	Norgeot et al. ²⁰
Scrubber	B	0.762	0.878	0.815	Norgeot et al. ²⁰
Philter	B	0.785	0.999*	0.879	Norgeot et al. ²⁰
MIST (trained on i2b2)	B	0.907	0.879	0.893	N/A
NeuroNER	B	0.979	0.950	0.964	N/A
nference (fine-tuned on Mayo)	B	0.961	0.988	0.974	N/A
nference (fine-tuned on Mayo + i2b2)	B	0.979*	0.992	0.985*	N/A

The results for Scrubber, Physionet, Philter, and the CRF + ANN method are based on previous publications. The MIST method required training and, thus, was trained on the 2014 i2b2 training dataset. We used a pre-trained model for NeuroNER. The two versions of the nference approach were fine-tuned on (1) only the Mayo dataset and (2) both the Mayo and the i2b2 datasets.

*Best performance for the metric.

high recall of 0.918. Closely following Philter, the MIST model achieves a recall of 0.889 with overall performance similar to that on the i2b2 dataset.

Error analysis on the Mayo dataset

We further investigated cases in the Mayo dataset where the nference de-identification model failed to successfully detect the PII element completely (i.e., false negatives). This occurred at a rate of 0.6% (see [Table 4](#)). Across the 10,000 notes considered in the test set, there were 848 error instances that contained these false negative errors. Accounting for duplicate occurrences of the same sentence, there were 797 unique error instances. We grouped these instances based on the type of identifier. The prevalence of the error category is shown in the second column, while the third column in the table represents the contribution of each category to the error in recall (sums to 0.6%).

The most prevalent error was in the recognition of entities pertaining to clinic locations (208 of 797). Many of these were due to partially identified phrases (e.g., “Room 7A” was missed in “out of Southwest Building Room 7A”). The second most prevalent error type was in dates, with 183 false negatives. The third most prevalent error category was in doctor/nurse names and initials, with 169 false negatives. Abbreviations and shorthand used by providers (typically while signing off on a clinical note) contributed to the errors in this category.

Ambiguous instances of PII also resulted in false negatives. These were cases that a human reader would have difficulty/uncertainty in deeming as PII. An example of this is the word “tp” in the phrase “Comment: 03-12-2005 08:04:12—verified tp.” We found that 26% of errors were those in which the nurse abstractors themselves did not agree on the characterization

Table 3. Performance of de-identification methods on the Mayo test dataset

Method	Precision	Recall	F1
Scrubber	0.756	0.677	0.715
Philter	0.709	0.918	0.800
Physionet	0.837	0.772	0.803
MIST (trained on Mayo)	0.818	0.889	0.852
NeuroNER (trained on Mayo)	0.928	0.933	0.931
nference (fine-tuned on Mayo)	0.967*	0.994*	0.979*

These methods were evaluated against group C entities.

*Best performance for the metric.

of PII (Cohen's κ for errors was lower than for non-errors, at 0.7453), pointing to the inherent ambiguity.

Distribution of errors per note

We further investigated the rate at which errors in detecting PII (false negatives) occurred on a per-note level. As shown in [Table 5](#), the error instances were distributed across 637 notes. Furthermore, we see that a majority of false negatives are spread evenly across the notes (525 of 637 notes, or 82.4%, contain a single error). For each subsequent error rate, we computed the coverage of PII entities. Here, coverage represents the fraction of PII present in the subset of notes up to the corresponding error rate.

Even for notes with a large number of errors (more than 6), the number of distinct error types is between 2 and 3. This illustrates that most of the errors are of the same type and an artifact of repetition of text within a note. For example, in the note with 10 errors, eight of the instances were related to location, while the remaining two were related to date. Examples of the errors pertaining to location here are “Location of INR sample: Other:

Smallville Other: Smallville Other: Smallville”, “Recommend Recheck: Other: 04/01/2017 Smallville Other: 04/01/2017 Smallville”, “Recommend Recheck: Other: 04/01/2017 Smallville Other: 04/01/2017 Smallville Other: 04/01/2017 Other: 04/01/2017 Smallville”. Here, the location errors all pertain to the same location “Smallville,” which illustrates how the effective amount of identifiable content is substantially smaller than suggested by the raw count. The date presented (“04/01/2017”) was successfully detected. Both the date and the location have been replaced with synthetic values for the purpose of this example.

Distribution of note types

In the Mayo test set, a physician note is associated with a note type (e.g., progress note, emergency visit, telephone encounter). Given that the structure and semantics of these note types vary greatly from one another, we analyze the enrichment of errors across them. From the 637 notes with errors, we found 134 distinct note types with at least 1 error. The top 14 note types with highest error content are listed in [Table 6](#). Notes of the type “Anti-coag service visit summary” contain the highest rate of errors (22 of 26 sampled notes), followed by “Electrocardiogram” (19 of 30 sampled notes).

Methods

Usage of Mayo clinic dataset

The Mayo EHR dataset is based on data from 477,000 patients that originated from multiple EHR data systems (including Epic and Cerner) spanning over 20 years. The dataset includes 104 million physician notes that capture the health care journey of patients in addition to structured tables containing lab test measurements, diagnosis information, orders, and medicine administration records. This research was conducted with approval from the Mayo Clinic Institutional Review Board.

Table 4. Prevalence and examples of types of false negatives encountered by the nference de-identification system when applied on the Mayo test set

Category	Number of error instances (n = 797)	Contribution to recall error (E = 0.6%)	Example (the PII presented in these examples is fictitious)
Clinic location	208	0.1461%	He had a DWI in January and was required to do treatment through <i>Samson rehab</i> in St. Louis, Missouri
Dates	183	0.1285%	CPL dated 4/27/04
Doctor/nurse name/initial	169	0.1187%	Sent: 2020-10-20 10:00 a.m.. Subject: RE: Consumer/Pat
Pharmacy name	54	0.0379%	S: fax received from <i>Trioki Rx</i> with request for new RX for Viread (tenofovir)
Phone number	50	0.0351%	Phone number patient/caller is calling from or the number of the provider: 724.161.1754
Organization/company	35	0.0246%	Last we talked about her involvement in a group called <i>GO GIRLS!</i>
Health care organization	22	0.0154%	Jane is brought in by a <i>Minerva</i> female attendant and said Jane has been like this for “weeks and weeks.”
Numeric identifier	9	0.0063%	Manufactured by Merck lot number 78-32-DK, expiration date 2020/10/20
Location (address or partial address)	8	0.0056%	500 <i>State Highway 72</i>
Patient name	4	0.0028%	PLOF: X was independent with self cares living

The entities highlighted in italics indicate the word or phrase that the system failed to detect.

Table 5. Distribution of number of errors per note

Errors per note	Number of notes	Total errors	Cumulative errors	PII coverage	Average number of error types
0	9,363	0	0	0.9940	0
1	525	525	525	0.9978	1.00
2	80	160	685	0.9989	1.56
3	10	30	715	0.9991	1.75
4	6	24	739	0.9992	2.30
5	6	30	769	0.9994	2.75
6	2	12	781	0.9995	2.5
7	2	14	795	0.9996	2.25
8	2	16	811	0.9997	2.25
9	3	27	838	0.9999	2.33
10	1	10	848	1.0000	2.00

PII coverage represents the fraction of PII present in the subset of notes up to the corresponding error rate. Average number of error types denotes the number of distinct error types (such as date errors or name errors) per note.

We randomly sampled 10,000 notes, which were reduced to a set of unique sentences. This yielded a test set of 172,102 sentences. These were subsequently annotated by six Mayo Clinic nurse abstractors to create a ground truth label for every word and/or phrase. Each sentence was annotated by at least two different nurse abstractors. The interannotator agreement on labeling a token as PII had a Cohen's κ of 0.9694 (see [supplemental methods](#) and [Table S5](#) for details).

An additional set of 10,000 notes was selected to fine-tune the models. We manually annotated 61,800 unique sentences from these notes to create a tagged fine-tuning set. See [supplemental methods](#) for more details.

Detection of PII entities

The ensemble architecture described in this section leverages state-of-the-art attention-based deep-learning models in

conjunction with rules harvested from the data (each of which is described below) to handle semi-structured text ([Figure 2](#)). There are several salient features of this approach that are worth noting. **Hybrid deep-learning models.** The newer breed of attention-based deep-learning models, in conjunction with transfer learning, allow for faster tuning of these models with significantly smaller sets of labeled data for detecting PII identifiers. We use pre-trained language models based on the BERT¹⁶ architecture that are then fine-tuned for detecting (1) personal names, (2) organizations, (3) locations, and (4) ages. We employed the BERT-base-cased model (<https://huggingface.co/bert-base-cased>) through the HuggingFace/Transformers (<https://github.com/huggingface/transformers>) library. This is a case-sensitive English language pre-trained model based off of the BERT architecture trained using an MLM objective. The fine-tuning process involves training the pre-trained language model on an NER task using a training set of annotated sentences. We used a total of 61,800 tagged example sentences to fine-tune the models. We fine-tuned each transformer model with a maximum sequence length of 256 (after tokenization) over four epochs. We used a training batch size of 32 and a learning rate of 5×10^{-5} with a warm-up proportion of 0.4. We then evaluated the model on a validation dataset and computed the accuracy. We performed the fine-tuning and model validation processes in an iterative manner (see [supplemental methods](#), [Figure S1](#), and [Table S1](#) for complete implementation details). Identifiers such as names, locations, organizations, and ages are well suited to a statistical entity recognition method because they can use the context of the surrounding text to disambiguate the entity type of a word. By contrast, pattern-matching rules are significantly hampered in this respect. It would be hard, for instance, to detect “Glasgow” as a medical term in “He had no helmet and his Glasgow score was 6” and as a location in “Mr. Smith had visited his family in Glasgow” using look-up dictionaries.

However, we use patterns to deterministically tag reasonably well-defined PII identifiers, which are almost entirely context

Table 6. Distribution of number of errors per note type

Note type	No. of error instances	No. of PII instances	No. of notes with at least one error	Total No. of notes	Fraction of notes with at least one error
Phone message/call	60	7,466	54	605	0.09
Ambulatory patient summary	59	14,502	49	334	0.15
Physician office/clinic message	42	8,352	36	661	0.05
Report	50	3,173	36	131	0.27
Medication renewal/refill	36	4,626	31	358	0.09
Progress note, family practice	27	4,975	24	237	0.10
Ambulatory discharge medication list	27	8,109	23	226	0.10
Anti-coag service visit summary	24	1,189	22	26	0.85
Electrocardiogram	19	411	19	30	0.63
Anticoagulation patient intake—text	49	5,777	18	50	0.36
Letter	15	3,519	14	157	0.09
Ambulatory depart summary	12	3,938	12	163	0.07
Progress notes	14	3,943	11	199	0.06
Telephone encounter	12	2,034	11	273	0.04

The proportion of sampled notes for a given type that contain at least one error is presented in the last column. This indicates in which note type an error is more likely to occur.

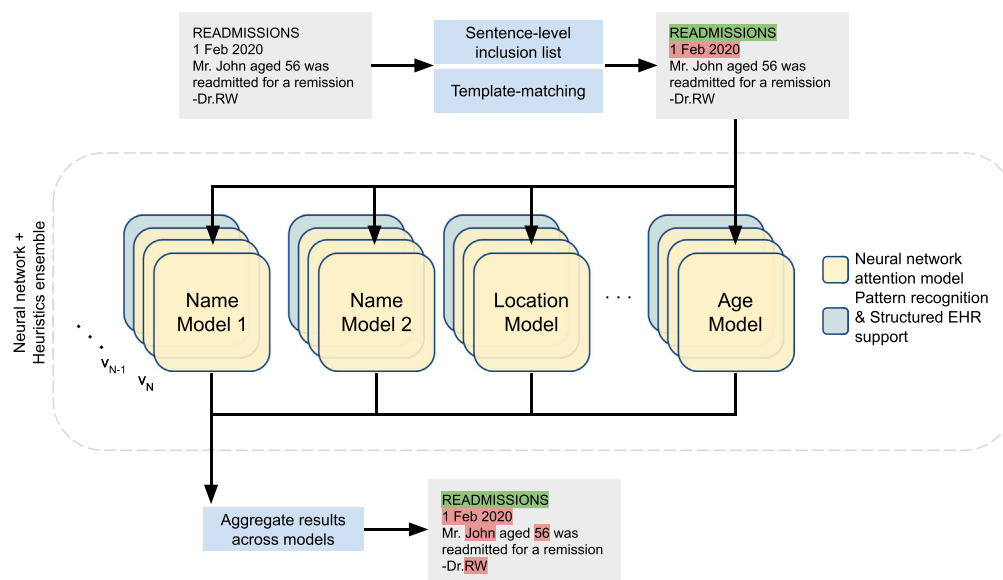


Figure 2. Ensemble architecture for de-identification of unstructured text

Sentence-based inclusion lists and template matching prune out sentences that either (1) lack PII or (2) contain PII in specific well-defined patterns. An ensemble of attention-based neural networks identifies complementary features across different PII types. For each entity type, multiple model versions (v_1, v_2, \dots, v_N) are used in tandem. In addition, pattern recognition modules and structured EHR content from matched patients support the anonymization process. The results from each component of the ensemble are aggregated to yield the original note labeled with PII tags.

independent and unambiguous. This category includes dates and times, phone and pager numbers, clinical IDs and numeric identifiers, email, URLs, IP addresses, and vehicle numbers. In addition, harvested sentence templates (described further below) are relied upon to deterministically tag PII instances matched by the template patterns. Our methods apply to content in both structured (e.g., lab comments) and free-form text (e.g., progress notes).

In addition, it should be noted that we designed our method to detect and transform information about those who provide care, such as physicians, nurses, and pharmacies. Although this is not required by HIPAA Safe Harbor, it allows health care organizations to protect the identities of their employees as well.

Ensemble of models framework and iterative fine-tuning. Given the regulatory necessity of extremely high recall for de-identification, we aggregate the results of multiple models trained for the same PII type. Our ensemble involved employing at least one individual model for names, organizations, locations, and ages (see Table S2). An additional text-normalized model was also trained and utilized for names. In this respect, if a term is detected as PII in any of the models for that type, then it is tagged. A divide-and-conquer approach has been implemented that harnesses the power of multiple models to identify PII or extract meaningful entities (Figure 2). In contrast to a “one size fits all” model, this approach enables each individual model to be fine-tuned to learn different (and complementary) features of the unstructured EHR data, as has been shown to be used in prior de-identification systems²³. For instance, one model focuses on identifying peoples’ names, while another is geared toward addresses and locations.

Furthermore, there are additional models corresponding to cased and uncased variants of the raw data (referred to as

“Name Model 1” and “Name Model 2” in Figure 2). Each model here corresponds to an attention-based deep neural network. One advantage of carving out the entity space to be handled individually by separate models is that each model needs to learn only the distribution of entities of a specific type as opposed to all entities. However, this introduces a challenge in resolving terms in a sentence that have conflicting and/or ambiguous entity types. These conflicts are resolved in the aggregation phase of our ensemble, where a simple voting threshold of one claim is employed (i.e., an entity is considered PII even if one model in the system tags it as such). Since the majority of the components in the ensemble are designed to detect complementary features, we are able to improve recall without much loss of precision.

Integrating databases as part of the core model. We use publicly available databases of names, locations, and addresses to supplement the model fine-tuning process. First names with supporting gender information were downloaded from the US Census database. Cities across the United States as well as lists of hospitals were obtained from Wikipedia. These public databases were used to augment training of our models. In addition, patient-specific information from structured EHRs, including patient names and residential addresses, are used to augment the model training and match against PII in the text.

Sentence-based inclusion list. Clinical note corpora contain a large number of repeated sentences. These stem from various processes, including automated reminders (e.g., “Please let your doctor know if you have problems taking your medications”), repeated phrases in the writing style of physicians (e.g., “Rubella: yes”; “Pain symptoms: no”), or shared elements in the clinical notes, such as section headers (e.g., “History of Present Illness”). From the corpus of physician notes from the Mayo Clinic, a set of 1,600 sentences, which did not contain PII, was incorporated into

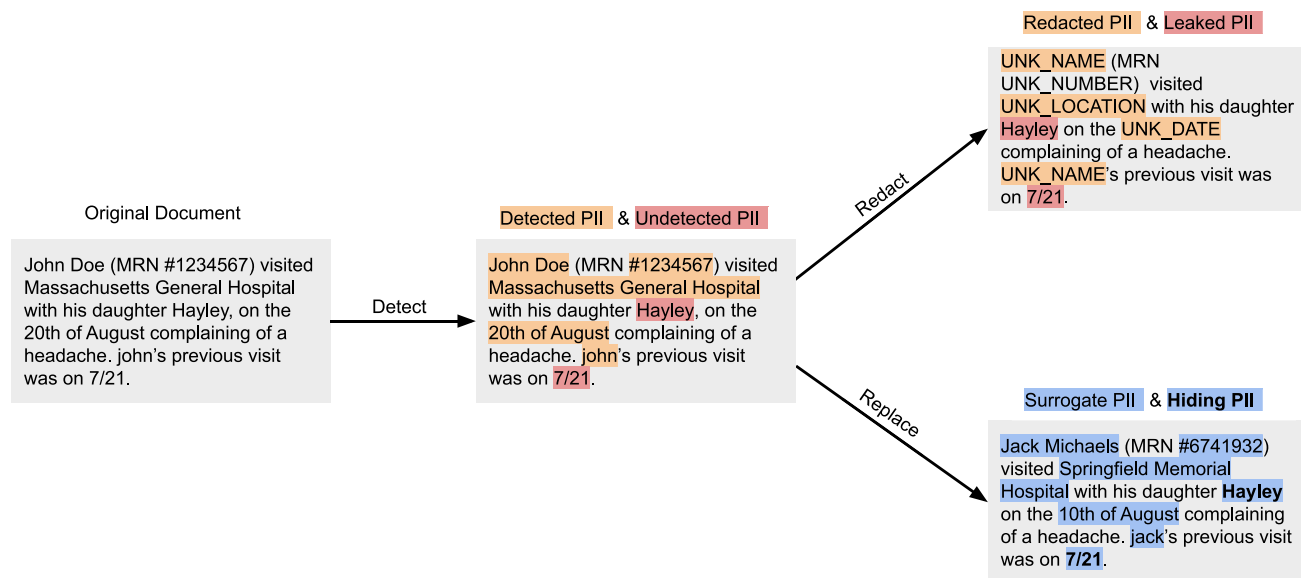


Figure 3. Obfuscation of tagged PII entities

An illustration of the hiding in plain sight mechanism to highlight the utility of the detect → replace strategy. After obfuscation, distinguishing real PII from surrogates is no better than what one would expect by random chance. PII, personally identifiable information.

an “inclusion list.” This inclusion list was further expanded with a set of 25,000 sentences containing medically relevant entities, such as disease or drug names (see [supplemental methods](#) for details on how the inclusion list was constructed). This has the added benefit of improving the precision of the de-identification system because it reduces the risk of misclassifying these important entities as PII by the neural network models. In addition, sentences marked as being devoid of PII during the validation phase in the iterative fine-tuning process were also added to the inclusion list (see [supplemental methods](#)).

Autogenerating templates using statistical NER models. In addition to exact sentences with high prevalence there are also a large number of PII-containing sentences that can be mapped to a template (e.g., “Electronically signed by: SMITH, JOHN C on 01/02/1980 at 12:12 PM CST”) maps to a template of the form “Electronically signed by: <LAST NAME>, <FIRST NAME> <INITIAL> on <DATE> at <TIME>”). While machine-learning NER models can be trained and/or fine-tuned to learn these patterns, there are instances where entity recognition fails. So, although a name of the form “SMITH, JOHN C” might be detected, “DEWEY” in “DEWEY, JONES K” may not be detected. By contrast, regular expression rules faithfully match every PII for these cases.

The problem, however, is that the process of identifying such templates and generating the corresponding regular expressions is an arduous task because it involves manual inspection of a sufficiently large sample of sentences in the corpus. Here, we use the NER ensemble models designed for the detection of PII to aid in the harvesting of these pattern templates. Sentences from our fine-tuning set of 10,000 notes are passed through the ensemble, and detected PII is transformed to its corresponding IOB2 (inside-outside-beginning) mask (e.g., “Electronically signed by: B-PER I-PER I-PER on B-DATE at B-TIME PM

CST”), generating a potential NER template. In addition, a “syntax template” for these sentences is also generated, such that any term that was detected as an entity is mapped to its syntactic representation—one of “W” for alphabets only, “N” for numbers only, and “A” for alphanumeric (e.g., “Electronically signed by: W, W W on N/N/N at N:N PM CST”). Finally, for each unique syntax template, if there exists only one NER template among all instances of the syntax template, a regular expression rule is generated (e.g., “Electronically signed by: [A-Za-z]+, [A-Za-z]+ [A-Za-z]+ on \d+/\d+/\d+ at \d+:\d+ PM CST”) by mapping each syntax token to its corresponding regular expression pattern: “W” to “[A-Za-z]+”, “N” to “\d+”, and “A” to “[w]”.

Transformation of tagged PII entities

The de-identification process is designed to recognize words and phrases that represent PII and other sensitive elements with high recall. However, if the input text is transformed to the de-identified version by *redacting* detected PII, undetected PII (e.g., “Hayley” and the date “7/21” in [Figure 3](#)) is obviously leaked to any person who reads the document. As such, the obfuscation process aims to conceal these residual PII by *replacing* detected PII with suitable surrogates so it is difficult to distinguish between the residual PII and the surrogates.^{21,24,25} This method has been implemented in several de-identification approaches.^{26,27} As highlighted in [Figure 3](#), it is difficult for a human to determine which of “Jack Michaels” or “Hayley” is a leaked instance of PII in the output of the replacement strategy using this mechanism of hiding in plain sight (HIPS).²⁸ Evidence with human readers has shown that when the recall of a NLP tool is high (i.e., when most real identifiers are detected), the rate of distinguishing real from filler identifiers is no better than what one would encounter by random chance. It has further been shown, however, that under highly controlled conditions, it is

possible for a machine-learning system to replicate the behavior of the natural language de-identification tool to remove fillers and leave real identifiers in place.^{28,29}

In addition to employing the HIPS method, we apply entity-specific rules and heuristics to improve the fidelity of the surrogate. We further improve interpretability of the output by ensuring that every unique PII token in all EHR records for a patient has the same transformation.

Consider the input text “John Smith, a pleasant 67 year old presented with his son Jack. John complains of breathing difficulty,” which was transformed to “Jane Kate, a pleasant 67 year old presented with his son Matt. Ryan complains of breathing difficulty.” In this example, “Jane Kate” as a surrogate is an obvious giveaway that it is a fake name and therefore lends itself to be distinguished from any true PII that may have leaked. Furthermore, it appears that a third completely different person is complaining of breathing difficulty. So an ideal transformation would have maintained the format of first name followed by last name, and the gender for “John Smith” and every instance of “John” or “Smith” in the input would be transformed to the same output; something like “Jacob Hamilton, a pleasant 67 year old presented with his son David. Jacob complains of breathing difficulty.”

As discussed, we manage the replacement of surrogates per entity type (see Table S2). Names are transformed in a manner that is consistent with format, gender, and ethnicity of the original (i.e., “Ms. Lopez visited New York General Hospital for her routine checkup” becomes “Ms. Hernandez visited Mass General Hospital for her routine checkup”). Dates are handled in a way to preserve their formatting (i.e., “March 5th, 2014” becomes “February 27th, 2014” and “03-05-2014” becomes “02-27-2014”). The shift in the date is a patient-specific random number. This ensures that dates are shifted consistently for a given patient. Locations and organizations are replaced with suitable surrogates chosen from a predefined dictionary. PII entities that contain numeric digits (such as phone number or patient ID) involve replacing these numbers randomly while maintaining overall length and format.

While the transformation output of an input token is the same for all instances of its occurrence for a given patient, it would be different across patients. That is, while all instances of “John” in one patient might be transformed to “Jacob,” for another patient it could be “Aaron.”

DISCUSSION

Numerous approaches to de-identification have been developed. Automated de-identification systems can broadly be segmented into four categories: (1) rule-based systems, (2) traditional machine-learning systems, (3) deep-learning systems, and (4) hybrid and ensemble systems.

Rule-based systems^{8,19,20,30,31} use pattern matching rules, regular expressions, and dictionary and public database look-ups to identify PII elements. These are simple to implement and usually deterministic; however, these systems have several drawbacks. First, pattern-matching rules for identifiers are typically not robust for handling variance in input due to typographical errors (spelling, punctuation, casing, etc.). A rule that matches “Dr. John” may not be able to match “Dr john.” Second, creating template patterns to match sentence fragments like “Provider Name: Dr. John” that

tag any term after “Provider Name: Dr.” as a name, for example, requires manual effort to understand the data to create these templates. Doing this for large datasets with notes for millions of patients is time consuming and intractable. Third, dictionary-based systems may not be complete, resulting in increased false negatives (i.e., true PII that is not detected). Fourth, blindly using dictionary/database look-ups induces false positives because they tag phrases that are not identifiers in the context in which they are used that need to be disambiguated.³² For example, in “The doctor determined his Braden score as normal”, the term “Braden” might be flagged as PII, when it is only a clinical term.

Traditional machine-learning systems^{21,33–35} use traditional machine-learning algorithms, such as support vector machines and CRFs, to perform NER classification as PII for each word in a sentence. The classification task involves creating labeled data and defining features based on properties like part-of-speech tags, typography (e.g., capitalization, casing, spacing, font weights, or font types), punctuation, and frequency of words and/or their neighbors. These methods, in addition to requiring significant effort in encoding the feature vectors, may not generalize across datasets.

Deep-learning systems¹⁸ have become the state of the art for a wide variety of application domains, including vision (e.g., image classification) and speech (e.g., voice recognition and generation). In language-related tasks (e.g., machine translation), these approaches have surpassed human-level performance.³⁶ Deep learning has proven beneficial in numerous NLP tasks, including predicting the next word (language modeling), tagging tasks such as part of speech tags, entities in a sentence (entity recognition), and dependency parsing. This has enabled applications that traditionally required custom rules and hand-crafted features to be solved without any feature engineering. Modern deep-learning approaches for de-identification have been shown to outperform their predecessors,¹⁸ but they require very large quantities of domain-specific labeled training data to perform well. Specifically, the challenges include, but are not limited to, the presence of long and highly descriptive sentences, usage of clinical shorthand (that varies across physicians and medical specialties), and a variety of semi-structured machine-generated content. Moreover, publicly available datasets for de-identification (including the popular i2b2 2014 dataset)¹⁷ lack diversity, often focusing on only a few types of notes or areas of disease. Training and benchmarking with such datasets are likely to bias the resulting models and fail to capture the nuanced and complex nature of physician notes. Recently, attention-based neural network (transformer) models have also been implemented for de-identification but have shown limited generalizability in the absence of support from encoded rules.³⁷

Hybrid³⁸ and ensemble systems^{39,40} use combinations of rule-based and machine-learning-based components in tandem to improve PII detection efficacy. With these approaches, the choice of components, finding the right split of tasks between them, and the optimal strategy for combining results from them become crucial. Some approaches⁴¹ invoke engineering post-processing layers that fix the errors that are introduced by other (earlier) components. In cases where there is, by design, overlap in the type of PII being predicted (e.g., multiple components detecting people’s names), considerable effort is spent measuring and choosing a method, like a stacked meta classifier or voting

scheme, to pick a winning component.³⁹ The inference de-identification system presented here addresses the limitations of prior methods¹¹ and achieves high levels of recall and precision.

There are several opportunities to further improve the performance of de-identification systems. First, existing knowledge graphs and language models trained on biomedical corpora can be leveraged. For example, if a patient's note contains the sentences "Patient diagnosed with lung cancer" and "ECOG performance status was determined to be 2," ECOG would not be treated as PII, since it has a strong biological association with lung cancer based on the knowledge graph. In the de-identification process, this could be used to recover biological terms incorrectly tagged as PII (false positives). Second, the quality of sentences that are provided to the model can be improved. Unstructured clinical text does not always contain well-formatted text, commonly due to missing punctuation and incorrect casing. A case-sensitive pre-trained model along with an MLM objective can be used to train a system capable of correctly introducing punctuation in the right location. Another challenge with the quality of clinical documents is the prevalence of short fragments and bullet points, giving rise to sentences with poor context. The context of a single sentence can be expanded using preceding and succeeding sentences or employing document-level transformer models such as Transformer-XL.⁴² Third, unsupervised methods can be incorporated to accelerate the annotation process of the NER task. Grouping the word representations generated by a transformer model yields informative clusters (e.g., a cluster of names) that can be annotated according to the nature of words present in the cluster. The NER task can then be formulated as a masked language task, where the overlap of the list of potential candidates for a missing word with the clusters can inform the entity type of the missing word.

Concluding remarks

Overall, this work implemented an ensemble approach to de-identification of unstructured EHR data incorporating transformer models supported by heuristics for automatically identifying PII across diverse clinical note types. Upon detection, suitable surrogates replaced PII in the processed text, thereby concealing residual identifiers (HIPS). The system demonstrates high precision and recall on both publicly available datasets and a large and diverse dataset from the Mayo Clinic.

EXPERIMENTAL PROCEDURES

Resource availability

Lead contact

Sankar Aradhanari is the lead contact for this study and can be reached at sankar@nference.net.

Materials availability

There are no physical materials associated with this study.

Data and code availability

The 2014 i2b2 data are publicly available subject to signed safe usage and for research only. The Mayo EHR clinical notes are not publicly available at this time. The source code is not currently available, but the inference de-identification tool can be accessed at <https://academia.nferx.com/deid/>.

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.patter.2021.100255>.

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AUTHOR CONTRIBUTIONS

Conceptualization, K.M., A.R., and S.A.; methodology, K.M., A.R., V.A., and S.A.; validation, K.M., B.M., V.A., S.B., J.A., J.R., and S.A.; formal analysis, K.M., B.M., and S.A.; data curation, K.M., V.A., and S.B.; writing – original draft, K.M. and A.R.; writing – review & editing, B.M., J.A., J.R., W.F., J.H., V.S., and S.A.; supervision, V.S. and S.A.; resources, V.S. and S.A.

DECLARATION OF INTERESTS

J.A., J.H., and W.F. do not have any competing interests in this project. B.M. is a contracted consultant of the Mayo Clinic. The authors on this article from inference have equity in inference and have a financial interest in inference. A patent application has been submitted by K.M., A.R., and S.A. Mayo Clinic and inference may stand to gain financially from the successful outcome of the research.

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