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# Explainable query generation for cohort discovery and biomedical reasoning using natural language

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A dissertation  
submitted in partial fulfillment of the  
requirements for the degree of

Doctor of Philosophy

University of Washington

2023

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## Abstract

Explainable query generation for cohort discovery  
and biomedical reasoning using natural language

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Clinical trials serve a critical role in the generation of medical evidence and furthering of biomedical research. In order to identify potential participants, investigators publish eligibility criteria, such as past history of certain conditions, treatments, or laboratory tests. Patients meeting a trial's eligibility criteria are considered potential candidates for recruitment. Recruitment of participants remains, however, a major barrier to successful trial completion, and manual chart review of hundreds or thousands of patients to determine a candidate pool can be prohibitively labor- and time-intensive.

At the same time, the amount and variety of data contained in Electronic Health Records (EHRs) is increasing dramatically, creating both challenges and opportunities for patient recruitment. While more granular and potentially useful data are captured and stored in EHRs now than in the past, the process of accessing and leveraging these data often requires technical expertise and extensive knowledge of biomedical terminologies and data models.

This thesis focuses on the development of an integrated system for identifying patients in clinical databases using a natural language interface. Humans use natural language nearly effortlessly, and thus automated means of leveraging natural language to identify patients in databases hold great potential in time and cost savings. The primary contributions of this work include a novel database schema annotation and mapping method enabling data

model agnostic query generation, a method for generating intermediate logical representations of eligibility criteria, exploration of dynamic reasoning upon non-specific criteria, and development of an integrated graph-based knowledge base of biomedical concepts.

This work also introduces two new annotated corpora, the Leaf Clinical Trials (LCT) corpus and Leaf Logical Forms (LLF) corpus. The LCT corpus is unique in the granularity with which it represents complex eligibility criteria, while the LLF corpus is the most extensive annotated corpus of eligibility criteria logical representations at the time of this writing. Both corpora are valuable contributions to the biomedical informatics and natural language processing communities.

To evaluate the viability of our methods, both our system and a human database programmer generated queries to identify patients eligible for 8 past clinical trials at our institution. We then compared actual participant enrollments to those found eligible. We demonstrate that our system rivals and sometimes surpasses an experienced human programmer in finding eligible patients.

## TABLE OF CONTENTS

	Page
List of Figures . . . . .	iv
List of Tables . . . . .	vii
List of Abbreviations . . . . .	ix
Chapter 1: Introduction . . . . .	1
1.1 Problem . . . . .	1
1.2 Contributions . . . . .	3
1.3 Overview . . . . .	4
Chapter 2: Background . . . . .	6
2.1 The Importance of Clinical Trials . . . . .	6
2.2 Challenges in Recruitment for Clinical Trials . . . . .	7
2.3 The Case for Software in Matching Patients for Clinical Trials . . . . .	7
2.4 Challenges in Electronic Screening and NLP in Clinical Trials . . . . .	9
2.5 NLP Methods for Clinical Trial Recruitment . . . . .	11
2.6 Corpora . . . . .	14
2.7 Clinical Knowledge Bases . . . . .	15
2.8 Summary . . . . .	16
Chapter 3: Leaf Clinical Trials Corpus . . . . .	17
3.1 Overview . . . . .	17
3.2 Motivation . . . . .	17
3.3 Annotation schema . . . . .	18
3.4 Evaluation . . . . .	29
3.5 Limitations . . . . .	34

3.6 Summary . . . . .	34
Chapter 4: Leaf Logical Forms Corpus . . . . .	35
4.1 Overview . . . . .	35
4.2 Motivation . . . . .	35
4.3 Related Work . . . . .	37
4.4 Methods . . . . .	38
4.5 Results . . . . .	41
4.6 Summary . . . . .	41
Chapter 5: Knowledge Base . . . . .	43
5.1 Overview . . . . .	43
5.2 Motivation . . . . .	43
5.3 Methods . . . . .	44
5.4 Limitations . . . . .	49
5.5 Summary . . . . .	50
Chapter 6: Semantic Metadata Mapping . . . . .	51
6.1 Overview . . . . .	51
6.2 Related Work . . . . .	51
6.3 Methods . . . . .	52
6.4 Limitations . . . . .	55
6.5 Summary . . . . .	55
Chapter 7: Query Generation . . . . .	57
7.1 Overview . . . . .	57
7.2 Motivation . . . . .	57
7.3 Related Work . . . . .	58
7.4 Methods . . . . .	59
7.5 Evaluation . . . . .	65
7.6 Results . . . . .	67
7.7 Discussion . . . . .	70
7.8 Conclusions . . . . .	71

Chapter 8: Web Application . . . . .	72
8.1 Overview . . . . .	72
8.2 Related Work . . . . .	72
8.3 Methods . . . . .	73
8.4 Future Work . . . . .	80
8.5 Conclusion . . . . .	81
Chapter 9: Conclusions . . . . .	82
9.1 Contributions . . . . .	82
9.2 Limitations . . . . .	84
9.3 Future Work . . . . .	85
Bibliography . . . . .	86

## LIST OF FIGURES

Figure Number	Page
2.1 An example screenshot of the Leaf user interface. The user has generated a query to find patients who identify as female, have had at least two Hemoglobin A1c laboratory results greater than or equal to 6.5 in the past 6 months, and are 65 years of age or older. . . . .	8
2.2 An example screenshot of the Criteria2Query application. Users type eligibility criteria in free text or enter a clinical trials "NCT" identifier. The application then generates a SQL query and executes it against an OMOP database in order to identify eligible patients. . . . .	13
3.1 Example eligibility criteria annotated used the LCT corpus annotation schema (left) and corresponding example SQL queries (right) using a hypothetical database table and columns. Annotations were done using the Brat annotation tool. The ICD-10 codes shown are examples and not intended to be exhaustive. . . . .	19
3.2 Examples of clinical trials eligibility criteria annotated with Chia and LCT annotation schemas. Each example shows a criterion from a Chia annotation (above) and an LCT annotation of the same text for purposes of comparison (below). . . . .	27
4.1 An example semantic parse using the LLF corpus annotation schema. The hypothetical input sentence is on the left, while the corresponding logical form is shown on the right. Indentations and colors have been added for readability. . . . .	37
4.2 An example LLF corpus annotation. The annotation file is saved in JavaScript (.js) format, which enables syntax highlighting and validation to assist annotators. Whether a given criterion was an inclusion or exclusion criteria is indicated at the top, followed by the original raw text, then augmented text. The final annotated logical forms are shown last. . . . .	40

6.1	The LeafAI query engine’s SQL query generation process using two hypothetical database schema to generate queries for platelet counts (shown in logical form after normalization). This example illustrates the flexibility of LeafAI’s semantic metadata mapping system (represented here in JSON format) in adapting to virtually any data model. On the left, ”Tall Table Structure”, platelet counts must be filtered from within a general purpose ”labs” table. The LeafAI KB recognizes that labs may be stored as LOINC codes, and the corresponding SMM indicates that records in this table can be filtered to LOINC values. On the right, ”Pivoted Table Structure”, platelet counts are stored as a specific column in a ”complete_blood_counts” table, and thus can be directly queried without further filtering. Additional metadata, columns, tables, types and so on needed in SMMs are omitted for brevity. . . . .	56
7.1	LeafAI query architecture. Inter-module communication is performed using the gRPC framework. Individual modules are deployed as Docker containers and communicate solely with the central API, which orchestrates query generation and handles query generation requests. . . . .	60
7.2	LeafAI query generation processes . . . . .	62
7.3	Longitudinal results for four trials. The blue line indicates recall for LeafAI and orange the human programmer. The X axis represents the line number for each eligibility criteria. Dots indicate that a query was executed for a given line. On the right, boxes represent the text of a criteria, with comments below discussing strategies and findings. . . . .	69
8.1	The web application user interface. User-entered criteria are shown in the above right, while system responses are shown in the lower left. User criteria are displayed and executed in order, with each count of patients representing a subset of the preceding count. . . . .	74
8.2	Normalized named entities identified in user input before query execution. On the left, the system correctly identified ”BMI”. On the right, ”Diabetes Mellitus” is misspelled, and the user is notified. . . . .	76
8.3	A time-lapse representation asynchronously reporting of incremental query results, from top to bottom. This is performed using two streaming interfaces, one from the clinical database to the API, and a second from the API to the web application. . . . .	77
8.4	An iterative workflow using an example case for adults with Diabetes Mellitus. Users are able to directly edit results and re-execute their queries while preserving query history, saving user time and preserving previous user actions and findings. . . . .	78

8.5 An example of a user editing concepts discovered using LeafAI's reasoning system. Users are able to enable or disable reasoned concepts. . . . .	79
8.6 An example of a user disabling a named entity from a system response. Named entities can be enabled and disabled by clicking directly on them. Disabled entities are shown with gray colored text and struck through by a line. Upon re-execution, logic for disabled entities is removed from the generated query.	80

## LIST OF TABLES

Table Number	Page
3.1 Examples of representative LCT annotation schema entities. A full listing of all entities can be found in the LCT annotation guidelines at <a href="https://github.com/uw-bionlp/clinical-trials-gov-annotation/wiki">https://github.com/uw-bionlp/clinical-trials-gov-annotation/wiki</a> . . . . .	21
3.2 Examples of representative relations. Direction of arrows indicates role, i.e., subject → target entity. . . . .	23
3.3 Annotation statistics for EliIE, Chia, and LCT corpora. . . . .	29
3.4 Hyperparameters and pre-trained embeddings used for named entity recognition and relation extraction baseline results. For the NER task, the same architecture and hyperparameters were used for both general and fine-grained entity models. For the relation extraction task, the same hyperparameters were used with both the BERT and R-BERT architectures. . . . .	30
3.5 Baseline entity prediction scores (% , Precision / Recall / F <sub>1</sub> ). Corpus-level micro-averaged scores are shown in the bottom row. For brevity a representative sample of entities is shown. Count refers to the total count of unique spans annotated in the entire corpus. Entities included in the total count and scores but omitted for brevity are Acuteness, Allergy, Condition-Type, Code, Coreference, Ethnicity, Eq-Operator, Eq-Unit, Indication, Immunization, Insurance, Life-Stage-And-Gender, Organism, Other, Specimen, Study and Provider. . . . .	31
3.6 Baseline relation prediction scores (% , Precision / Recall / F <sub>1</sub> ). Corpus-level micro-averaged scores are shown in the bottom row. For brevity a representative sample of relations is shown. Count refers to the total count annotated in the entire corpus, including relations not shown. The count total excludes general to fine-grained entity relations, which as overlapping spans are not used for relation prediction. Relations included in the total count and scores but omitted for brevity are Acuteness, Code, Criteria, Except, From, Indication-For, Is-Other, Max-Value, Min-Value, Polarity, Provider, Refers-To, Specimen, Stage, Study-Of and Type. . . . .	32

<b>3.7 Results of NER experiments using the manually annotated and semi-automated portions of the corpus.</b>	The manually annotated portion includes 513 documents while the semi-automatically annotated portion is 493 documents. . . . .	33
<b>4.1</b>	Example inputs and logical form syntax styles with fine-tuning performance results using the T5 <sub>base</sub> model. . . . .	42
<b>6.1</b>	Hypothetical database table for patient diagnoses. Codes from a variety of different vocabularies are shown under the <i>coding-system</i> column. . . . .	54
<b>6.2</b>	Example SMM configuration for the <i>code</i> column. . . . .	54
<b>7.1</b>	Statistics for each clinical trial evaluated by the LeafAI query engine and human programmer. The number of enrolled and matched patients were determined by cross-matching enrollments listed within our EHR. The <i>Time</i> column indicates the number of hours the human programmer spent developing queries for each trial. . . . .	67
<b>7.2</b>	The LeafAI query engine's handling of eligibility criteria for each trial. The column <i>No Patients</i> indicates the count of criteria which would, if executed, cause no patients to be eligible. The column <i>Not Computable</i> indicates the count of criteria which LeafAI could not generate a query for, for various reasons. Both of these types of criteria were ignored by the system. . . . .	68

## **LIST OF ABBREVIATIONS**

AST: Abstract syntax tree

BI-LSTM: Bi-directional long-short term memory

CRF: Conditional random field

EHR: Electronic health record

FHIR: Fast healthcare interoperability resources

ICD: International classification of disease

KB: Knowledge base

LCT: Leaf clinical trials (corpus)

LLF: Leaf logical forms (corpus)

NLP: Natural language processing

OMOP: Observational medical outcomes partnership

SEQ2SEQ: Sequence to Sequence

SNOMED: Systematized nomenclature of medicine

SMM: Semantic metadata mapping

UMLS: Unified medical language system

## ACKNOWLEDGMENTS

I want to begin by thanking my advisor, Meliha Yetisgen. At a time when I knew next-to-nothing about natural language processing, she kindly and patiently met with me and explained the basics and suggested what I should focus on learning. She was among the first to suggest I get a PhD, wrote not one but two recommendations for my graduate school application (first as non-matriculated, then as a formal PhD student), and has provided ongoing constructive feedback, advice, support, and opportunities for much of my career. I think there are few people in life that we can objectively say we have been changed by, but in mine she is certainly one of them. I'm eternally grateful.

I also want to thank the members of my committee, Nina Kim, Trevor Cohen, and Fei Xia. Nina was among the first physicians I interacted with at UW, and her feedback and suggestions over the years have been extremely helpful. Trevor and Fei are both truly excellent teachers and passionate about their work.

Sean Mooney and Peter Tarczy-Hornoch have also been critical to my PhD journey. I'll never forget the day that I showed Peter—who happened to be passing by—a demo of the prototype of Leaf. That Peter was willing to sit and talk with me about how it worked for over an hour meant the world to me (particularly as a former History major with never-ending imposter syndrome). Sean has been unerringly supportive and a wonderful mentor.

Last, I'm incredibly grateful to my wife, Akiko. Through the years of working full-time while pursuing my PhD, she has helped me stay grounded and focused on our family and children, who I am intensely proud of and who will always be my greatest joy and priority. I can only hope that they grow to enjoy the same love of learning and imagining *what could be?* that I think make life interesting and worth living.

## **DEDICATION**

To my children, Elly, Ami, and Sage.

# Chapter 1

## INTRODUCTION

### 1.1 *Problem*

Randomized controlled trials serve a critical role in the generation of medical evidence and furthering of biomedical research. In order to identify patients for clinical trials, investigators publish eligibility criteria, such as past history of certain conditions, treatments, or laboratory tests. These eligibility criteria are typically composed in free-text, consisting of inclusion and exclusion criteria. Patients meeting a trial's eligibility criteria are considered potential candidates for recruitment.

Recruitment of participants remains a major barrier to successful trial completion [1], so generating a large pool of potential candidates is often necessary. Manual chart review of hundreds or thousands of patients to determine a candidate pool, however, can be prohibitively labor- and time-intensive. Cohort discovery tools such as Leaf [2] and i2b2 [3] may be used, providing researchers with a relatively simple drag-and-drop graphical interface in their web browser to create database queries to find potential patients in electronic health records (EHR) [4]. Learning how to use such tools nevertheless presents a challenge, as graphically-represented concepts may not align with researchers' understanding of biomedical phenomena or trial eligibility criteria. In addition, certain complex queries may simply be impossible to execute due to structural limitations on the types of possible queries presented in these tools, such as complex temporal or nested sequences of events [5].

An alternative approach which holds promise is the use of natural language processing (NLP) to automatically analyze eligibility criteria and generate database queries to find patients in EHRs. NLP-based approaches have the advantage of obviating potential learning curves of tools such as Leaf, while leveraging existing eligibility criteria composed in a format

researchers are already familiar with. Previous research in the use of NLP for trial recruitment [6, 7, 8, 9, 10, 11, 12, 13, 14] has shown promise but also limitations, preventing more widespread use for recruitment. For example, eligibility criteria often refer to non-specific criteria [15, 16], such as "conditions affecting respiratory function", which save investigator time by avoiding the need for exhaustive lists of conditions, but also must be reasoned upon to determine. Past research has also largely supported only single, static data models, limiting potential usage to only institutions or researchers with data in particular formats, and preventing future additions or changes to data structure necessitated by changes in clinical workflow, research needs, and so on. In terms of evaluation, past research has also largely evaluated solutions based on metrics such as concept normalization performance and so on, which while important, miss addressing the more important question of how well such tools find patients that an investigator may reasonably expect them to find given certain criteria.

This work explores the development of a user-friendly, explainable, and generalizable application for cohort discovery from natural language by querying virtually any database schema. We demonstrate that this system is capable of state-of-the-art query generation enabling significantly faster discovery of eligible patients for clinical trials and biomedical research in general.

The importance of improving efficiencies in trial recruitment is difficult to overstate. Clinical trials are considered the gold standard of evaluation of new treatments but also extremely costly; for example, the average cost of a Phase 3 trial in the United States ranges from US\$11.5 million to US\$52.9 million [17]. Much of those costs are due to difficulties in finding and keeping patients enrolled. Systems which stand to improve efficiencies in finding patients eligible for trials using a medium researchers already know well and frequently use - natural language - therefore may also contribute to reducing costs and simplifying processes for enrolling patients.

The potential impact of this work extends beyond clinical trials. Biomedical research in general very often relies on finding patients meeting certain criteria, whether for retrospective analysis, sample size estimates, preparation to research, and so on. We hope that the

methods and application we develop for creating a natural language interface to databases may simplify and streamline this common critical step for a variety of research purposes.

## 1.2 Contributions

This work explores the development of a novel database query-generation system to identify patients eligible for clinical trials using a natural language interface. The primary contributions of this work are:

1. *Annotated Corpora*: this work introduces two new annotated clinical corpora: the LCT corpus and LLF corpus. The LCT corpus contains over 1,000 annotated eligibility criteria documents and captures annotated phenomena important for the task of clinical query generation at a highly granular level [18]. These phenomena include clinical events, such as diagnoses, procedures, laboratory tests, vitals, and medications, demographic facets such as sex, ethnicity, and age, as well as abstract, temporal, and qualifying values such as contraindications, indications, temporal sequences, negations, risks, and severity. The LCT corpus is unique in the granularity of its annotation schema and number of annotated documents. The LLF corpus is a subset of 2,000 lines of eligibility criteria from the LCT corpus which includes the original criteria sentence alongside a machine-readable intermediate logical representations. To the best of our knowledge, the LLF corpus is the only large publicly available data set of human-annotated eligibility criteria as logical forms.
2. *Model-agnostic query generation*: We introduce a novel database annotation schema and mapping method to enable query generation on virtually any clinical data model. Our system uses a subset of Unified Medical Language System (UMLS) [19] concepts to flexibly represent database resources using concepts related to metadata, such as patient identifiers, demographics, and vocabularies.
3. *Knowledge Base*: Various tasks related to query generation, such as hierarchy navigation

tion (e.g., finding types of surgeries of cancers), vocabulary and code translation (e.g., conversion to SNOMED, LOINC, or ICD-10 codes), and reasoning (explained next) necessitate the use of a Knowledge Base (KB). We introduce a graph-based KB which incorporates the UMLS, the Disease Ontology [20], Symptom Ontology [21], COVID-19 Ontology [22], Potential Drug-Drug Interactions [23], LOINC2HPO [24], and the Disease-Symptom Knowledge Base [25], as well as various equivalency mappings not present in the UMLS [26, 27, 28, 29].

4. *Reasoning upon non-specific criteria:* Many criteria within eligibility criteria are non-specific, and instead need to be first reasoned upon in order to be computable. For example, a criterion may refer to patients "indicated for bariatric surgery", or "who have a disease affecting respiratory function". Using our logical form representations and KB, we demonstrate a system for dynamically reasoning on non-specific criteria.
5. *Evaluation against a human programmer and actual trial enrollments:* Past systems for automatically identifying patients eligible for clinical trials have largely limited their evaluations to sub-systems such as normalization performance, and thus it is difficult to assess how well such systems actually perform when used with real-world data and trials. In contrast, we evaluate our system using enrollments linked to clinical data for 8 past clinical trials at the University of Washington. We compare the patients found by our generated queries to actual enrolled patients. Finally, we compare our results to those of an human clinical database programmer.

### **1.3 Overview**

*Chapter 2, Background*, provides a general overview of clinical trials, as well as past work in cohort discovery and eligibility prediction in relation to NLP. Literature specific to given tasks is also presented briefly in each chapter where applicable.

*Chapter 3, Leaf Clinical Trials corpus*, introduces the first annotated corpus, including motivation, annotation schema, comparison to other corpora, and baseline named entity

recognition (NER) and relation extraction performance using multiple neural architectures.

*Chapter 4, Leaf Logical Forms corpus*, presents the second annotated corpus and our logical forms annotation schema. The corpus is evaluated using various syntax styles suggested from the literature.

*Chapter 5, Knowledge Base*, explores the motivation and development of the KB. We discuss each source at a high level and SPARQL queries used for reasoning.

*Chapter 6, Semantic Metadata Mapping*, describes the motivation and methods we use for dynamically generating queries for eligibility criteria in a data model-agnostic fashion.

*Chapter 7, Query Generation*, explores the development of our methods for query generation, including models used for NER and logical form transformation, normalization, and reasoning. Our system is evaluated in comparison to that of a human database programmer.

*Chapter 8, Web Application*, examines ongoing and future research related to our user-facing web application for interactive cohort discovery.

*Chapter 9, Conclusions*, summarizes this work's primary contributions and possible directions for future research.

## Chapter 2

# BACKGROUND

This chapter provides an introduction to clinical trials and their importance to biomedical research. Section 2.1 provides background for and discusses the motivation and reasoning in focusing on clinical trials. Section 2.2 examines challenges in trial recruitment. Section 2.3 discusses past work in patient matching for clinical trials using electronic software. Section 2.4 examines challenges in electronic screening for trials using NLP. Section 2.5 discusses research in NLP methods for clinical trial recruitment, while Sections 2.6 and 2.7 examine published corpora related to clinical trials and clinical knowledge bases. Section 2.8 provides a summary of the chapter.

### **2.1    *The Importance of Clinical Trials***

A clinical trial is a prospective study comparing the effects and value of an intervention (typically a medication, biologic, or procedure) against a control group without it [30]. Clinical trials are considered "controlled" when a control group not receiving an interventional treatment is used for comparison, and "randomized" when participants are randomly placed in said treatment or control groups. Randomization is considered ideal for reducing risk of investigator bias and producing study groups closely in proportion to known risk factors. Randomized controlled trials (RCTs) are widely recognized as the best available method to determine if a given intervention is safe and effective [30].

Eligibility for a clinical trial is determined by a trial's *eligibility criteria*, which are free-text descriptions of required conditions, treatments, laboratory tests and so on. Eligibility criteria are composed of *inclusions*, which patients *must meet*, and *exclusions*, which patients *must not meet* in order to be eligible. The extent to which trial results can be assumed to

be generalizable to patients not participating in a trial but with comparable health status is determined by *external validity* [31]. External validity can be influenced by a number of factors, first and foremost the selection of patients potentially eligible for a trial. Finding patients appropriately meeting eligibility criteria in as unbiased way as possible is thus critical to drawing meaningful conclusions from clinical trials, and ultimately scientific progress and improvement of human health.

## **2.2 Challenges in Recruitment for Clinical Trials**

Many clinical trials fail to meet their expected number of enrollments [32, 33, 34, 35, 36, 37]. The reasons for this are varied, including overly restrictive inclusion criteria [33], a lack of awareness on the part of patients, particularly in underserved and historically disadvantaged communities [35], fear or apprehension of medical research due to past abuses [32], and uncertainty of risk on the part of providers leading to withheld offers to participate [37]. In addition, patients who do agree to participate in clinical trials tend to be wealthier, have greater access to healthcare resources, be members of ethnic majorities, and often unrepresentative of overall populations of patients suffering from a given condition [33, 35, 37, 38, 39, 40]. Beyond questions of generalizability, recruitment challenges also cause delays to clinical trials, with an estimated 86% of trials delayed between 1 and 6 months, and some for even longer [41, 42].

These challenges often have severe effects on the outcomes of clinical trials, and to a certain extent new treatments available to patients. These effects can include inadequate statistical analysis of outcomes, cost overruns, extended duration of trials, increased costs of new medications, and treatments that potentially do not exhibit expected beneficial outcomes in understudied populations. [43, 39, 44, 45].

## **2.3 The Case for Software in Matching Patients for Clinical Trials**

While computer software and NLP alone can likely not solve many of these challenges, research suggests that in many cases they can add significant value, time- and cost-savings

toward trial recruitment [39, 42]. For example, Thadani *et al* found electronic screening methods to significantly reduce the burden of manual chart review in one study by approximately 81% [42]. Examining multiple clinical trials, Penberthy *et al* similarly found up to a 20-fold decrease in staff time spent reviewing eligible patient records by using electronic screening software [39]. More recently, Ni *et al* used a combination of NLP techniques and structured data analysis to screen for potential clinical trial candidates and compared the results to a gold standard data set reviewed by medical doctors. The authors found their highest performing methods achieved an approximate 90% workload reduction in chart review and 450% increase in trial screening efficiency [46]. Drag and drop tools web-based tools such as Leaf [2], shown in Figure 2.1, and i2b2 [3] are also capable of determining eligible patients for clinical trials.

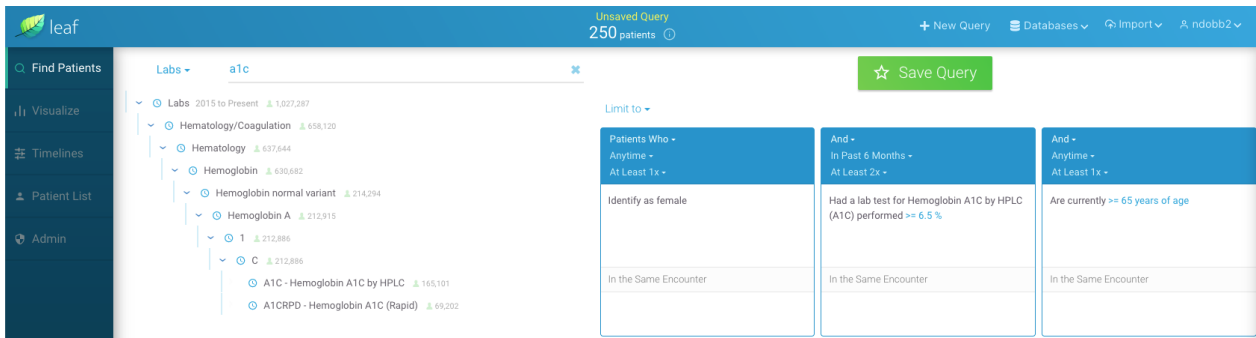


Figure 2.1: An example screenshot of the Leaf user interface. The user has generated a query to find patients who identify as female, have had at least two Hemoglobin A1c laboratory results greater than or equal to 6.5 in the past 6 months, and are 65 years of age or older.

Thus though the aim of this project is to produce an application capable of general purpose cohort discovery - rather than solely for the purposes of clinical trial recruitment - clinical trials are a meaningful and valuable means by which to **gather data, evaluate performance, and measure potential real-world impact** of solutions for cohort discovery. Moreover, screening software and NLP have been demonstrated to dramatically improve trial recruitment efficiency in many scenarios. In terms of data, the web-

site <https://clinicaltrials.gov>, maintained by the United States National Library of Medicine, hosts freely accessible descriptions of hundreds of thousands of clinical trials from around the world. Because clinical trials enrollments are in many cases recorded in EHRs (which also include the same patients' clinical data), they also can serve as a uniquely objective means of measuring the effectiveness of NLP systems in matching actual enrolled participants based on eligibility criteria. Put another way, an NLP-based system for matching patients to real-world eligibility criteria should reasonably be expected to find many or most patients enrolled in a given clinical trial - with the assumption that patients enrolled in those trials correctly met the necessary criteria as determined by study investigators. Thus however imperfect (e.g., a lack of diagnosis data for an existing condition may cause certain patients to be inappropriately deemed ineligible), clinical trials are thus well-suited for evaluation of NLP-based cohort discovery systems and thus a focus of much of this project.

#### **2.4 Challenges in Electronic Screening and NLP in Clinical Trials**

Using NLP to determine patients potentially eligible for a clinical trial has numerous challenges. Consider, for example, a list of eligibility criteria such as<sup>1</sup>:

1. *Newly diagnosed with breast cancer and scheduled for surgery*
2. *18 years or above*
3. *Those who experience high psychological stress will enter the RCT whereas those with low stress will be followed in an observational questionnaire study*
4. *No severe psychiatric disease requiring treatment, e.g., schizophrenia*

While perhaps appearing deceptively simple, this example demonstrates many of the difficulties of this task. In criterion 1, "Newly" in "Newly diagnosed with breast cancer", suggests that diagnoses occurring further in the past (though how far is unclear) should

---

<sup>1</sup>Adapted from <https://clinicaltrials.gov/ct2/show/NCT03254875>

not be included. Meanwhile, "surgery" in "scheduled for surgery" likely refers to surgery in relation to breast cancer, though this is not explicitly stated. In criterion 2, "18 years" likely refers to participants' age, but this too is not explicitly stated. Criterion 3, meanwhile, is a description of processes which will take place during the trial, but is not actually an eligibility criterion (i.e., participants may be eligible whether their actual stress levels are high or low). In criterion 4, "psychiatric disease" is non-specific and may refer to a large number of unstated conditions, aside from schizophrenia which is given as an example.

Appropriately interpreting the semantics and unstated requirements of these criteria are challenging for NLP systems. For example, an NLP system may correctly determine that "breast cancer" refers to a condition and "surgery" refers to a procedure, but may still fail if "Newly" is not determined to refer to breast cancer or to mean something occurring for the first time. In a subsequent step, an NLP system may normalize (i.e., determine a coded representation of a concept, for example a Unified Medical Language System [UMLS] code) "breast cancer" incorrectly as "Malignant Neoplasms" (C0006826) rather than "Malignant neoplasm of breast" (C0006142). In criterion 3, an NLP system may attempt to limit eligibility to patients with high stress levels, despite the criterion not being a formal restriction as such. In criterion 4, an NLP system may fail to reason that other unstated conditions, such as hysteria or hallucinations, should also be excluded.

Beyond challenges in interpreting eligibility criteria, certain criteria may simply be absent or even incorrect in the data source the NLP system generates a query for. For example, Eastern Cooperative Oncology (ECOG) performance status scores [47] are frequently listed in eligibility criteria but often absent in structured clinical databases.

Last, an NLP system capable of identifying patients eligible for clinical trials should also be able to explain *why* patients are eligible and *what it did* to determine eligibility. This step is key to gaining user trust in the system [48, 49], but also challenging as many systems tend to prioritize performance over interpretability.

## 2.5 NLP Methods for Clinical Trial Recruitment

Various methods for matching eligibility criteria to cohorts of patients using NLP have been put forth by the research community [6, 7, 8, 9, 10, 11, 12, 13, 14]. NLP-based cohort discovery methods hold unique potential and appeal, as they are theoretically able to leverage existing eligibility criteria described in natural language, a medium researchers and investigators already use and are comfortable with. Recent methods which utilize NLP in some form can generally be grouped into 5 categories:

1. **Database query generation** to Structured Query Language (SQL) or similar systems using either (a) rules, (b) neural network-based encoder-decoder architectures, or both.
2. **Document ranking and classification** using clinical notes in terms of relevancy vis-à-vis a given eligibility criteria.
3. **Projection into embeddings** of patient medical history and trial eligibility criteria in a shared vector space and matching via similarity measurement or entailment.
4. **Logical representations and reasoning** to represent eligibility criteria and patient records, matching by combinations of Semantic Web technologies, ontologies, Description Logics, and rule-based reasoning.
5. A combination of the above.

**Database query generation** - SQL-based relational databases are widely used both commercially and within academic institutions, and as such SQL is perhaps unsurprisingly often the target language in natural language to database query research [50]. Yuan *et al* developed Criteria2Query (shown in Figure 2.2), a hybrid information extraction (IE) pipeline and application which uses both rules and machine learning to generate database queries on an Observational Medical Outcomes (OMOP) [51] database. This work was expanded by Fang *et al*, who added functionality for iterative query generation via human

correction and adjustment [8]. Although not specific to RCTs, other highly relevant recent work on query generation in the biomedical domain has been done using Encoder-Decoder neural architectures for transforming clinical natural language questions into SQL queries [52, 53, 54, 55, 12]. Park *et al* [53] experimented with transforming medical questions generated in the MIMICSQL data set [56, 54] using both SQL and SPARQL queries with varying database schema representations. Bae *et al* similarly experimented with methods for handling typos, misspellings, and abbreviations in generating SQL queries from natural language questions. Pan *et al* [55] leveraged intermediate abstract syntax tree-based representations and a SQL grammar-based Decoder architecture for dynamic database schema matching.

**Document ranking and classification** - Focusing on clinical notes, Chen *et al* [10] used hybrid rule-based heuristics and sentence pattern-matching to detect criteria structure, as well as a combination of neural network-based bi-directional long short-term and conditional random field (biLSTM+CRF) architecture and knowledge graphs using the UMLS for determining condition, lab, procedure and drug relationships. Soni and Roberts [7] utilized the BERT Transformer architecture [57] and Lucene [58] to summarize, rank and classify clinical notes as relevant to a given eligibility criterion, with the most relevant notes predicted to be eligible.

**Embedding projections** - Dhayne *et al* [12] experimented with treating patient-to-RCT matching as a joint embedding and similarity measurement problem while also incorporating the SNOMED-CT ontology to infer basic "is-a" and "has-type" relations between concepts. Similarly, Zhang *et al* [9] used joint patient and eligibility criteria embeddings for entailment prediction, where predicting that a patient can be inferred from a given eligibility criteria equates to eligibility.

**Logical representations and reasoning** - Patrao *et al* developed Recruit [11], an ontology-driven trial recruitment system which transformed SQL relational data to Resource Description Framework (RDF) graph-based triples. The RDF triples in turn were made query-able by use of an OWL-based reasoning system [59] and normalization techniques to infer cancer staging. Building upon earlier work [60, 61, 62], Baader *et al* [63] explored

**Criteria2Query**

**Support**

**✓ Inclusion Criteria**

Please input criteria line by line.

Evidence of the AD pathological process, by a positive amyloid assessment either on CSF A $\beta$ 1-42 as measured on Elecsys  $\beta$ -Amyloid(1-42) Test System OR amyloid PET scan  
AD dementia of moderate severity, as defined by a screening MMSE score of 16-21 points, inclusive, and a CDR-GS of 1 or 2

**✗ Exclusion Criteria**

Please input criteria line by line.

Pregnant or breastfeeding  
Inability to tolerate MRI procedures or contraindication to MRI  
Contraindication to PET imaging  
Residence in a skilled nursing facility

<input checked="" type="checkbox"/> #	Inclusion Criteria:	Delete All Tags
<input checked="" type="checkbox"/> 1	Evidence of the <b>AD[Adrenal adenoma]</b> <small>CONDITION</small> pathological process , by a positive amyloid assessment either on <b>CSF A<math>\beta</math>1 - 42[Cerebrospinal fluid index]</b> <small>MEASUREMENT</small> as measured on <b>Elecsys <math>\beta</math> - Amyloid ( 1 - 42 ) Test System OR amyloid[Amyloid A level]</b> <small>MEASUREMENT</small> <b>PET scan[Positron emission tomography]</b> <small>PROCEDURE</small>	
<input checked="" type="checkbox"/> 2	<b>AD dementia[Dementia]</b> <small>CONDITION</small> of moderate severity , as defined by a screening <b>MMSE score[Rating of perceived exertion [Score]]</b> <small>MEASUREMENT</small> of <b>16 - 21 points</b> , inclusive <small>VALUE</small> , and a <b>CDR - GS[Clinical dementia rating scale]</b> <small>MEASUREMENT</small> of <b>1 or 2</b> <small>VALUE</small>	

<input checked="" type="checkbox"/> #	Exclusion Criteria:	Delete All Tags
<input checked="" type="checkbox"/> 1	<b>Pregnant[Pregnant]</b> <small>CONDITION</small> or <b>breastfeeding[Breastfeeding painful]</b> <small>CONDITION</small>	
<input checked="" type="checkbox"/> 2	Inability to tolerate <b>MRI[Magnetic resonance imaging]</b> <small>PROCEDURE</small> procedures or <b>contraindication[Medical contraindication]</b> <small>CONDITION</small> to <b>MRI[Magnetic resonance imaging]</b> <small>PROCEDURE</small>	
<input checked="" type="checkbox"/> 3	<b>Contraindication[Medical contraindication]</b> <small>CONDITION</small> to <b>PET imaging[Positron emission tomography (PET) imaging: limited area (eg, chest, head/neck)]</b> <small>PROCEDURE</small>	
<input checked="" type="checkbox"/> 4	Residence in a skilled <b>nursing[Caries of infancy associated with breast feeding]</b> <small>CONDITION</small> facility	

Figure 2.2: An example screenshot of the Criteria2Query application. Users type eligibility criteria in free text or enter a clinical trials "NCT" identifier. The application then generates a SQL query and executes it against an OMOP database in order to identify eligible patients.

the use of Description Logics and ontologies in matching patients in the MIMIC data set to logical representations of eligibility criteria, for example representing "Diabetes mellitus

type 1” as ” $\exists_y.\text{diagnosed\_with}(x, y) \wedge \text{Diabetes\_mellitus\_type\_1}(y)$ ”. Liu *et al* [13] used domain experts to manually translate criteria into a custom syntax parsable by software. For example, the criterion ”Patients more than 18 years old when they received the treatment” would be represented as ”#Inclusion features[’StartDate’]  $i =$  demographics[’BirthDate’] + @YEARS(18)”. Parsed eligibility criteria were then executed on a proprietary database schema to determine eligible patients.

## 2.6 Corpora

A key element for any NLP-based approach in identifying patients is robust corpora which capture eligibility criteria semantics sufficiently for high-accuracy query generation. Such corpora can serve as reliable benchmarks for purposes of comparing NLP methods as well as training data sets. A number of corpora have been published. Weng *et al* created EliXR [64], a pipeline of rule-based methods for syntactic parsing of eligibility criteria using pattern matching. EliXR was validated using a subset of 1,000 randomly selected eligibility criteria annotated by human annotators and compared to syntax trees extracted by their system. This work was expanded by Boland *et al* to support normalization of complex temporal patterns [65]. Neither data set was publicly released. Kang *et al* developed EliIE [66], a hybrid machine-learning and rule-based system for eligibility criteria extraction and normalization. EliIE was evaluated against a human-annotated corpus of eligibility criteria from 230 trials related to Alzheimer’s Disease. EliIE was the first system for eligibility criteria extraction which used machine-learning methods (a Conditional Random Field, or CRF [67]) and the first to use an extraction schema based on the Observational Medical Outcomes Partnership (OMOP) [51] common data model. The EliIE corpus was similarly not publicly released, and also limited in terms of generalizability by focusing on Alzheimer’s Disease alone.

More recently, Kury *et al* released Chia [68], a human-annotated corpus of eligibility criteria from 1,000 Phase IV clinical trials across various disease domains. Like EliIE, Chia used an annotation schema based on the OMOP common data model. Following Chia, Sun *et al* created a corpus of COVID-19-related trials’ eligibility criteria [69]. The authors used

a similar annotation schema to that of Chia, but included separate annotation layers for cohorts, criterion, named entities, and concepts, which they referred to as a hierarchical annotation model.

Yu *et al* [70] released a corpus designed for direct text-to-query generation with semantic parsing, however given the relative simplicity of generated queries to date compared to the complexity of clinical databases, it's not clear this approach is yet viable for real-world clinical trials recruitment.

## 2.7 Clinical Knowledge Bases

A system capable of reasoning on non-specific eligibility criteria necessarily must store some representation of clinical phenomena and their relations. Such information is often described as a knowledge base (KB) [71], or an information store often composed of questions and answers, entities and relations, or factoids. Large-scale KBs, such as DBpedia [72] are represented as graphs of tuples, where a tuple represents two entities and a relation between them (also often called nodes and edges), such as { "asthma", "is-a", "disorder of the respiratory system" }. Chains of tuple-based entities joined by relations can therefore represent fairly complex phenomena in a compositional fashion. In medicine, the most widely used KB is the Unified Medical Language System, or UMLS [19]. The UMLS is a metathesaurus of controlled vocabularies and terminologies such as ICD-9, ICD-10, LOINC, SNOMED, CPT, and others, with mappings between them. While often accessed as a relational database, the UMLS can also be transformed and represented tuple form [73].

Within the clinical domain, KBs (often including the UMLS) have been used for various purposes related to retrieval, reasoning, disambiguation, and question answering. For example, Martinez *et al* and Sankhavara *et al* demonstrated use of the UMLS for query expansion, enabling patient record retrieval via use of synonyms [74, 75]. For reasoning, Shulz and Hahn demonstrated a technique for leveraging the UMLS to reasoning upon anatomical structure [76]. Kazi *et al* demonstrated the use of the UMLS for basic question-answering of medical students [77].

More recent work has explored alternative representations of KBs within neural networks. Lin *et al* explored neural network-based inference techniques for inferring missing relations among entities based on other information present in the UMLS and other KBs [78]. Hao *et al* and Huang *et al* examined approaches for integrating the UMLS into pre-training of BERT [57] embeddings [79, 80]. Petroni *et al* point out that even without fine-tuning for a specific domain or task, pre-training of language models with vast data from the internet exhibit reasonable performance on many tasks which necessitate basic reasoning [81].

## 2.8 Summary

This chapter discussed the importance of clinical trials to biomedical research and human health, as well as challenges in cost and recruitment. We next discussed motivation for and challenges in the use of NLP in patient matching for clinical trials, as well as related research in clinical KBs and published corpora related to clinical trials.

## Chapter 3

# LEAF CLINICAL TRIALS CORPUS

### 3.1 Overview

The NLP tasks involved in transforming eligibility criteria into database queries may include **named entity recognition** (NER) to tag meaningful spans of text as named entities, **relation extraction** to classify relations between named entities, **normalization** to map named entities to common coded representations (e.g., ICD-10), **negation detection** to detect negated statements (e.g., "not hypertensive") and so on. Gold standard corpora quality can thus directly affect performance and the validation of each of these tasks. Such corpora can serve as reliable benchmarks for purposes of comparing NLP methods as well as training data sets.

In this chapter we describe the creation of a gold standard corpus of human-annotated clinical trial eligibility criteria and performance data measured during our evaluation. Predictive models trained on this corpus enable and are used in subsequent aims of this project. Section 3.2 discusses motivations and need for the corpus. Section 3.3 describes the LCT annotation schema. We discuss our evaluation methods in Section 3.4, limitations in Section 3.5, and provide a summary in Section 3.6. This work was published as Dobbins *et al* in 2022 [18].

### 3.2 Motivation

We aimed to develop an expressive, task-oriented annotation schema which could capture a wide range of medical concepts and logical constructs present in eligibility criteria. To accomplish this, we first analyzed previously published corpora [64, 65, 66, 68] and expanded the list of included biomedical phenomena to fully capture the context and logic present in

real clinical trials criteria. As one example, we introduced an entity called *Contraindication* to reflect where use of a given treatment is inadvisable due to possible harm to the patient.

Figure 3.1 illustrates the importance of corpora structure for the task of query generation, using examples of eligibility criteria annotated using our annotation schema and corresponding hypothetical Structured Query Language (SQL) queries. In the first eligibility criterion, "preeclampsia" is explicitly named, and thus can be directly normalized to an ICD-10 or other coded representation. However, eligibility criteria involving non-specific drugs, conditions, procedures, contraindications, and so on are used frequently in clinical trials. In the second criterion in Figure 3.1, "diseases" in "diseases that affect respiratory function" is non-specific, and must be reasoned upon in order to determine appropriate codes, such as asthma, chronic obstructive pulmonary disease (COPD), or emphysema. Programmatically reasoning to generate queries in such cases would be challenging and often impossible if the underlying semantics were not captured appropriately. With this in mind, we developed the Leaf Clinical Trials, or LCT, annotation schema in order to enable reasoning and ease query generation for real-world clinical trials use. As the second example in Figure 3.1 shows, the LCT annotation captures the semantics of complex criteria, with changes to "respiratory function" annotated using a *Stability/change* entity and *Stability* relation, and the cause, "diseases" annotated with a *Caused-By* relation.

### 3.3 Annotation schema

The LCT annotation schema is designed with the following goals and assumptions:

1. The annotation schema should be **practical** and **task-oriented** with a focus on facilitating ease of query generation.
2. A greater number of **more specific, less ambiguous** annotated phenomena should be favored over a smaller number of possibly ambiguous ones.
3. Annotations should be **easily transformable** into composable, interconnected pro-

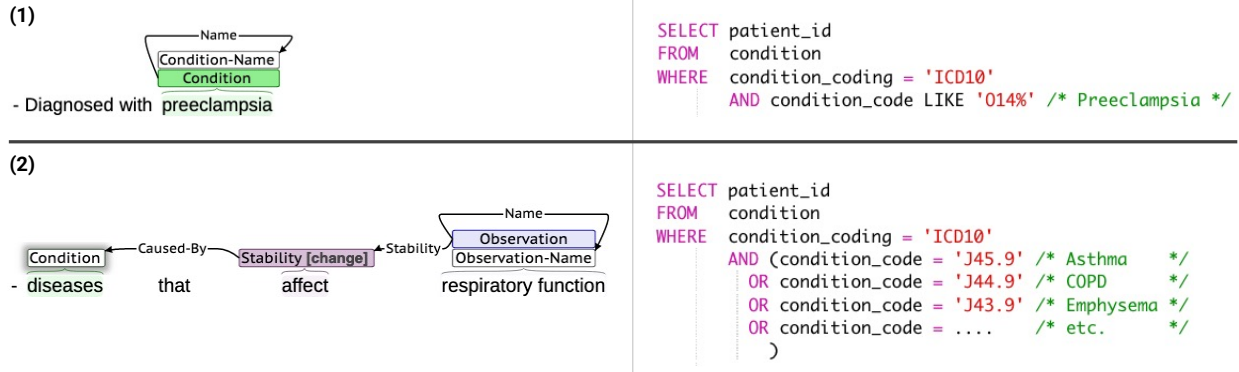


Figure 3.1: Example eligibility criteria annotated used the LCT corpus annotation schema (left) and corresponding example SQL queries (right) using a hypothetical database table and columns. Annotations were done using the Brat annotation tool. The ICD-10 codes shown are examples and not intended to be exhaustive.

grammatic objects, trees, or node-edge graph representations.

4. The annotation schema should **model eligibility criteria intent and semantics** as closely as possible in order to ensure generated queries can do the same.

The LCT annotation schema is composed of **entities** and **relations**. Entities refer to biomedical, demographic, or other named entities relevant to eligibility criteria, and are annotated as a span of one or more tokens. We organized LCT entities into the following categories:

- **Clinical** - *Allergy, Condition, Condition-Type, Code, Contraindication, Drug, Encounter, Indication, Immunization, Observation, Organism, Specimen, Procedure, Provider.*
- **Demographic** - *Age, Birth, Death, Ethnicity, Family-Member, Language, Life-Stage-And-Gender.*
- **Logical** - *Exception, Negation.*

- **Qualifiers** - *Acuteness, Assertion, Modifier, Polarity, Risk, Severity, Stability.*
- **Comparative** - *Criteria-Count, Eq-Comparison* (an abbreviation of "Equality Comparison"), *Eq-Operator, Eq-Temporal-Period, Eq-Temporal-Recency, Eq-Temporal-Unit, Eq-Unit, Eq-Value.*
- **Other** - *Ccoreference, Insurance, Location, Other, Study.*

The LCT corpus also includes 7 *Name* entities: *Allergy-Name, Condition-Name, Drug-Name, Immunization-Name, Observation-Name, Organism-Name* and *Procedure-Name*. *Name* entities serve a special purpose in the LCT corpus, as they indicate that a span of text refers to a *specific* condition, drug, etc., as opposed to *any* condition or drug. *Name* entities overlap with their respective general entities. For example, the span "preeclampsia" refers to a specific condition, and would thus be annotated as both a *Condition* and *Condition-Name*, while the span "diseases" is non-specific and would be annotated as only *Condition*. A full listing of the LCT annotation guidelines can be found at <https://github.com/uw-bionlp/clinical-trials-gov-annotation/wiki>.

We defined a total of 50 entities in the LCT corpus. Examples of selected representative entities are presented in Table 3.1. In our representation, a subset of entities have **values** as well. For example, an *Encounter* may have a value of *emergency, outpatient* or *inpatient*. Values are optional in some entities (such as *Encounters* or *Family-Member*, where they may not always be clear or are intentionally broad) and always present in others. In the example annotations presented below, values are denoted using brackets ("[...]"") following entity labels.

Relations serve as semantically meaningful connections between entities, such as when one entity acts upon, is found by, caused by, or related in some way to another. We categorize relations into the following:

- **Alternatives and Examples** - *Abbrev-Of, Equivalent-To, Example-Of.*

Category	Entity	Values	Example Text
Clinical	Condition	–	Diagnosed with <u>hypertension</u> in past year <i>Condition</i>
	Contraindication	–	any <u>contraindications</u> to vaginal delivery <i>Contraindication</i>
	Drug	–	on <u>beta blockers</u> <i>Drug</i>
	Encounter	emergency, outpatient, inpatient	recently <u>admitted</u> to a hospital <i>Encounter[inpatient]</i>
	Observation	lab, vital, clinical-score, survey, social-habit	Platelet count less than 500 <i>Observation[lab]</i>
Demographic	Procedure	–	Undergoing or scheduled for a <u>colonoscopy</u> <i>Procedure</i>
	Age	–	43 years <u>old</u> <i>Age</i>
Logical	Birth	–	<u>born</u> within the past 6 months <i>Birth</i>
	Family-Member	mother, father, sibling, etc.	history of <u>maternal</u> breast cancer <i>Family-Member[mother]</i>
	Language	–	Speaks <u>English</u> or <u>Spanish</u> <i>Language Language</i>
Negation	–	–	with <u>no</u> systemic disease <i>Negation</i>
Qualifier	Assertion	intention, hypothetical, possible	which <u>may</u> cause conditions <i>Assertion[hypothetical]</i>
	Modifier	–	<u>alcohol</u> or <u>substance</u> abuse <i>Modifier Modifier</i>
	Polarity	low, high, positive, negative	showing <u>elevated</u> serum creatinine <i>Polarity[high]</i>
	Risk	–	at heightened <u>potential</u> for suicide <i>Risk</i>
	Severity	mild, moderate, severe	with <u>serious</u> complications from surgery <i>Severity[severe]</i>
	Stability	stable, change	conditions known to <u>affect</u> mood <i>Stability[change]</i>
Comparative	Criteria-Count	–	at least 3 of the <u>following</u> conditions: <i>Criteria-Count</i>
	Eq-Comparison	–	greater than 50ml <i>Eq-Comparison</i>
	Eq-Temporal-Period	past, present, future	<u>Active</u> illness <i>Eq-Temporal-Period[present]</i>
	Eq-Temporal-Recency	first-time, most-recent	<u>Latest</u> BMI <i>Eq-Temporal-Recency[most-recent]</i>
Other	Location	residence, clinic, hospital, unit, emergency-department	Seen at <u>diabetes care clinic</u> <i>Location[clinic]</i>

Table 3.1: **Examples of representative LCT annotation schema entities.** A full listing of all entities can be found in the LCT annotation guidelines at <https://github.com/uw-bionlp/clinical-trials-gov-annotation/wiki>.

- **Clinical** - *Code, Contraindicates, Indication-For, Name, Provider, Specimen, Stage, Type.*
- **Dependent** - *Caused-By, Found-By, Treatment-For, Using.*
- **Logical** - *And, If-Then, Negates, Or.*
- **Qualifier** - *Acuteness, Asserted, Dose, Modifies, Polarity, Risk-For, Severity, Stability.*
- **Comparative** - *After, Before, Criteria, Duration, During, Max-Value, Min-Value, Minimum-Count, Numeric-Filter, Operator, Per, Temporal-Period, Temporal-Recency, Temporal-Unit, Temporality, Unit, Value.*
- **Other** - *From, Except, Has, Is-Other, Location, Refers-To, Study-Of.*

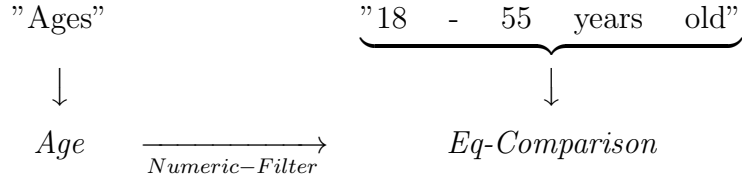
We defined a total of 51 relations in the LCT corpus. Examples of relation criteria are shown in Table 3.2.

In our annotations, some entity spans overlap with other entity spans in order to fully capture complex underlying semantics. Consider for example, the expression "Ages 18-55 years old". While an *Age* entity may be assigned to token "Ages", if an *Eq-Comparison* entity alone were assigned to the span "18-55 years old", the underlying semantics of the tokens "18", "-", "55", and "years" would be lost. In the following examples, we use the term **fine-grained entity** to refer to entities which are sub-spans of other **general entities**. Fine-grained entities are linked to general entities by relations. We use down arrow symbols ( $\downarrow$ ) to denote entity annotation and left and right arrow symbols ( $\leftarrow$  and  $\rightarrow$ ) to denote relations. The (+) symbols denote overlapping entities on the same span.

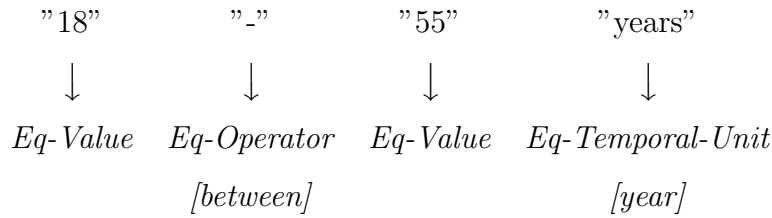
The example expression "Ages 18-55 years old" would be annotated in three layers. In the first layer, the expression is annotated with *Age* and *Eq-Comparison* general entities with a relation between them:

Category	Relation	Example Annotation
Alternatives and Examples	Abbrev-Of	<u>Post Concussion Syndrome</u> $\xleftarrow{\text{Abbrev-Of}}$ ( <u>PCS</u> ) <u>Condition</u>
	Equivalent-To	<u>Thrombocytopenia</u> $\xleftarrow{\text{Equivalent-To}}$ platelets <u>Condition</u> $\xrightarrow{\text{Observation[lab]}}$
	Example-Of	<u>skin condition</u> $\xleftarrow{\text{Example-Of}}$ (e.g. <u>eczema</u> ) <u>Condition</u>
Clinical	Contraindicates	conditions <u>contraindicating</u> $\xrightarrow{\text{Contraindicates}}$ <u>MRI</u> <u>Contraindication</u> $\xrightarrow{\text{Procedure}}$
Dependent	Caused-By	<u>swellings</u> $\xrightarrow{\text{Caused-By}}$ due to <u>trauma</u> <u>Observation</u> $\xrightarrow{\text{Condition}}$
	Found-By	<u>lesion</u> $\xrightarrow{\text{Found-By}}$ seen on standard <u>imaging</u> <u>Observation</u> $\xrightarrow{\text{Procedure}}$
	Treatment-For	<u>coronary bypass surgery</u> $\xrightarrow{\text{Treatment-For}}$ for <u>atherosclerosis</u> <u>Procedure</u> $\xrightarrow{\text{Condition}}$
	Using	<u>total knee arthroplasty</u> $\xrightarrow{\text{Using}}$ with <u>spinal anesthesia</u> <u>Procedure</u> $\xrightarrow{\text{Procedure}}$
	If-Then	BMI $\xrightarrow{\text{Eq-Comparison}}$ greater than 38 <u>women</u> <u>Life-Stage-And-Gender[female]</u> $\xleftarrow{\text{If-Then}}$ for
Qualifier	Risk-For	<u>risk</u> $\xrightarrow{\text{Risk-For}}$ of <u>death</u> <u>Risk</u> $\xrightarrow{\text{Death}}$
	Severity	<u>mild</u> $\xleftarrow{\text{Severity}}$ symptoms <u>Severity[mild]</u> $\xrightarrow{\text{Observation}}$
	Stability	<u>hemodynamically</u> $\xrightarrow{\text{Stability}}$ unstable <u>Observation</u> $\xrightarrow{\text{Stability[change]}}$
Temporal and Comparative	After	<u>infected</u> $\xrightarrow{\text{After}}$ following <u>admission</u> <u>Condition</u> $\xrightarrow{\text{Encounter[inpatient]}}$
	Before	diagnosis of <u>aortic stenosis</u> $\xrightarrow{\text{Before}}$ prior to <u>visit</u> <u>Condition</u> $\xrightarrow{\text{Encounter}}$
	Duration	<u>type 1 diabetes</u> $\xrightarrow{\text{Duration}}$ for <u>at least 1 year</u> <u>Condition</u> $\xrightarrow{\text{Eq-Comparison}}$
	During	<u>mechanically ventilated</u> $\xrightarrow{\text{During}}$ while <u>admitted</u> <u>Procedure</u> $\xrightarrow{\text{Encounter[inpatient]}}$
Other	Numeric-Filter	<u>body weight</u> $\xrightarrow{\text{Numeric-Filter}}$ less than 110 pounds <u>Observation[vital]</u> $\xrightarrow{\text{Eq-Comparison}}$
	Minimum-Count	<u>admitted</u> $\xrightarrow{\text{Minimum-Count}}$ at least twice <u>Encounter[inpatient]</u> $\xrightarrow{\text{Eq-Comparison}}$
	Temporality	<u>seen</u> $\xrightarrow{\text{Temporality}}$ within past 6 months <u>Encounter</u> $\xrightarrow{\text{Eq-Comparison}}$
	Location	<u>admitted</u> $\xrightarrow{\text{Location}}$ to the <u>ICU</u> <u>Encounter[inpatient]</u> $\xrightarrow{\text{Location[unit]}}$

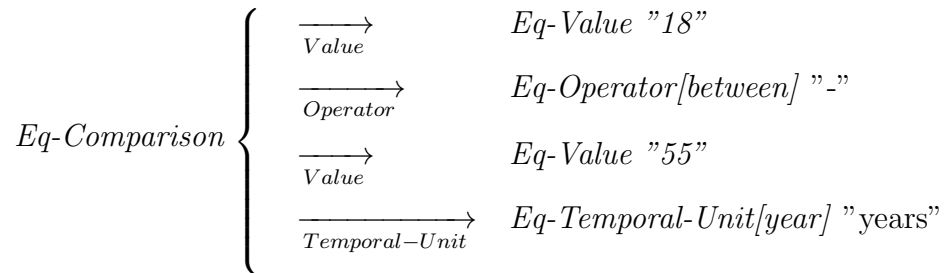
Table 3.2: **Examples of representative relations.** Direction of arrows indicates role, i.e., subject → target entity.



In the second layer, fine-grained entities with respective values are annotated:



In the third layer, relations connecting fine-grained entities to the general *Eq-Comparison* entity are added:



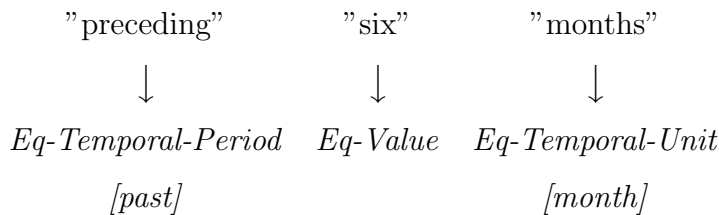
This multilayered annotation strategy allows significant flexibility in capturing entities and relations in a slot-filling fashion, simplifying the task of downstream query generation.

The LCT annotation schema contributes the following novel features: (1) deep granularity in entities and relations, which enables (2) rich semantic representation, closely capturing the intent of complex clinical trial eligibility criteria and facilitating accurate query generation.

### 3.3.1 Deep Entity and Relation Granularity

We assume that more specific annotation labels are generally more straightforward to generate accurate queries with. For example, within the span, "preceding six months", annotating the token "preceding" as *Temporal* (an entity type in Chia) may appear to be adequate, given that an English-speaking human would understand that this refers to the past. Without further information, however, a naïve algorithm would be unable to determine (1) whether such a entity refers to the past, present, or future, (2) that the token "six" refers to a numeric value, and (3) that "months" refers to a unit of temporal measurement. In such cases, most query generation algorithms introduce additional rule-based or syntactic parsing modules, such as SuTime [82] to further normalize the phrase to a value [64, 6]. This ambiguity in label semantics creates unnecessary complexity in downstream systems, requiring that the same text be processed a second time.

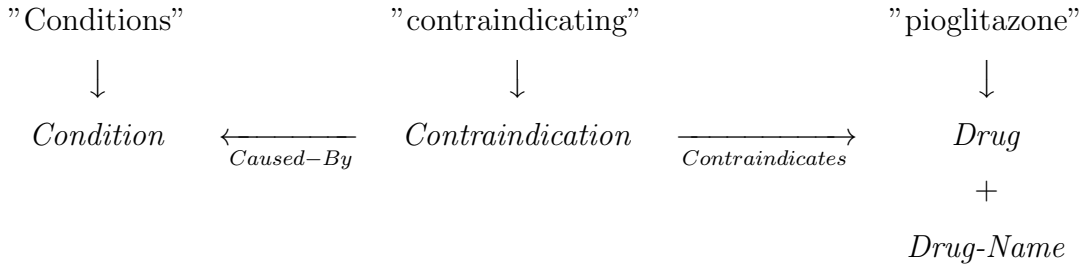
In contrast, we designed the LCT annotation schema to favor discrete, explicit entities and relations where possible, with an aim toward reducing the need for additional normalization steps needed for query generation. In our annotation schema, this example would be annotated with the following fine-grained entities:



As shown in the example, each token is uniquely annotated, with the values *[past]* and *[month]* serving to more clearly disambiguate semantics of the temporal phrase to a normalized temporal value. Moreover, as fine-grained entities are connected by relations to general entities, which can in turn have relations to other general entities, the LCT annotation schema is able to capture eligibility criteria semantics at a deeper level than other corpora.

### 3.3.2 Rich Semantic Representation

Certain eligibility criteria cannot be directly translated into queries, but instead must first be reasoned upon. For example, a query to find patients meeting the criterion of "conditions contraindicating pioglitazone" requires reasoning to first answer the question, *What* conditions contraindicate use of pioglitazone? Such reasoning may be performed by a knowledge base or other methods, but cannot be done unless the contraindicative relation is first detected:



As the span "Conditions" is labeled *Condition* but does not have an overlapping *Condition-Name* entity, it is considered unnamed and thus would need to be reasoned upon to determine. "[P]ioglitazone", on the other hand, includes a *Drug-Name* entity and is thus considered named. The absence of overlapping *Name* entities serves as an indicator to downstream applications that reasoning may be needed to determine relevant conditions or drugs.

### 3.3.3 Comparison to Chia

We designed the LCT annotation schema by building upon the important previous work of EliIE and Chia. Chia builds upon EliIE and is more recent. Figure 3.2 shows comparison examples of annotations of the same eligibility criteria using the two corpora in the Brat annotation tool [83].

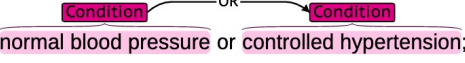
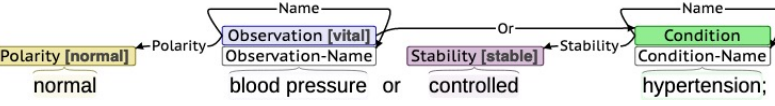
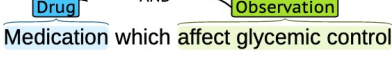
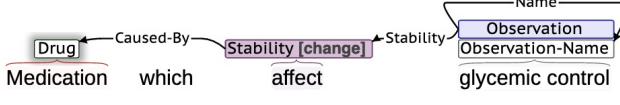
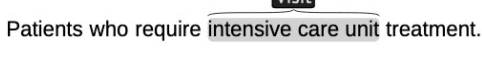
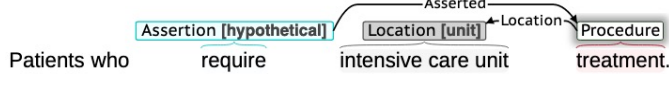
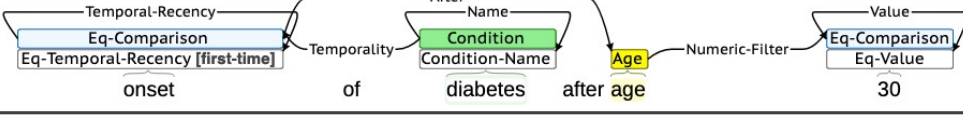
Chia		(1) <a href="https://clinicaltrials.gov/ct2/show/NCT01803828">https://clinicaltrials.gov/ct2/show/NCT01803828</a>
LCT		
Chia		(2) <a href="https://clinicaltrials.gov/ct2/show/NCT02315287">https://clinicaltrials.gov/ct2/show/NCT02315287</a>
LCT		
Chia		(3) <a href="https://clinicaltrials.gov/ct2/show/NCT01743755">https://clinicaltrials.gov/ct2/show/NCT01743755</a>
LCT		
Chia		(4) <a href="https://clinicaltrials.gov/ct2/show/NCT02624908">https://clinicaltrials.gov/ct2/show/NCT02624908</a>
LCT		

Figure 3.2: Examples of clinical trials eligibility criteria annotated with Chia and LCT annotation schemas. Each example shows a criterion from a Chia annotation (above) and an LCT annotation of the same text for purposes of comparison (below).

### 3.3.4 Annotation process

We extracted 1,020 randomly selected clinical trials eligibility descriptions from <https://clinicaltrials.gov> from 2018 to 2021, 20 for training and inter-annotator comparison and 1,000 for post-training annotation.

During annotation, 14 documents were found to be information poor (often with no spans to annotate) and discarded, resulting in 1,006 total annotated eligibility descriptions.

Annotation was performed by two annotators, the first a biomedical informatician and the second a computer scientist. For initial annotation training, 20 documents were distributed to both annotators. Annotation was done in the following steps:

1. Annotation meetings were held bi-weekly for 3 months following initial annotation training in which the annotation guidelines were introduced. Initial meetings focused on discussion of annotation guideline implementation and revision.
2. After annotation guideline revisions and annotation training were completed, eligibility criteria were assigned to each annotator, with each clinical trial eligibility criteria annotated by a single annotator using the BRAT annotation tool [83]. Due to differences in time availability for annotation, roughly 90% (887 documents) of the annotation task was performed by the first annotator, and 99 documents by the second annotator.
3. At the point in which 50% of the corpus was annotated, we trained two neural networks (one for general entities and another for fine-grained entities) using the biLSTM+CRF-based NeuroNER tool [84] on our manually annotated eligibility criteria to predict annotations for the remaining 50%.
4. Manual annotation was completed on the remaining 50% of eligibility descriptions by editing and correcting the predicted entities from NeuroNER in (3).

The resulting corpus included 887 single-annotated and 119 double-annotated total notes. Summary statistics for the corpus are shown in Table 3.3.

### 3.3.5 *Inter-annotator agreement*

Inter-annotator agreement was calculated using  $F_1$  scoring for entities and relations with 20 double-annotated documents. Entity annotations were considered matching only if entity types and token start and end indices matched exactly. Relations annotations were similarly

Measure	EliIE [66]	Chia [68]	<b>LCT Corpus</b>
Disease domain	Alzheimer’s Disease	All	<b>All</b>
No. of Eligibility Descriptions	230	1,000	<b>1,006</b>
No. of Annotations	15,596	68,174	<b>105,816</b>
No. of Entity types	8	15	<b>50</b>
No. of Relation types	3	12	<b>51</b>
Mean Entities per doc.	-	46	<b>105</b>
Mean Relations per doc.	-	19	<b>49</b>

Table 3.3: **Annotation statistics for EliIE, Chia, and LCT corpora.**

considered matching only if relation type and token start and end indices of both the subject and target matched exactly.

Initial inter-annotator agreement using the 20 training documents was 76.1% for entities and 60.3% for relations. Inter-annotator agreement improved slightly to 78.1% (+2%) for entities and 60.9% (+0.6%) for relations in the 99 additional double-annotated documents, indicating reasonably high annotator agreement considering the complexity of the annotation task.

### 3.4 Evaluation

To evaluate baseline predictive performance on the LCT corpus, we first created a randomly assigned 80/20 split of the corpus, with 804 documents used for the training set and 202 for the test set. For entity prediction, we trained NER models using biLSTM+CRF and BERT [57] neural architectures. For BERT-based prediction, we used two pretrained models trained on published medical texts, SciBERT [85] and PubMedBERT [86]. For both biLSTM+CRF and BERT predictions, we trained one model to predict general entities and another for fine-grained entities.

For relation extraction, we evaluated SciBERT for sequence classification as well as a

modified BERT architecture, R-BERT, following methods developed by Wu & He [87], also using the pretrained SciBERT model. Table 3.4 shows hyperparameters used for each task.

Task	Architecture	Hyperparameter / Embeddings	Training Value
Named Entity Recognition	biLSTM+CRF	Character Dimensions	25
		Token Embedding Dimensions	100
		Learning Rate	0.005
		Dropout	0.5
		Pretrained Embeddings	GloVe [88]
Relation Extraction	BERT & R-BERT	Pretrained Model	SciBert
		Learning Rate	0.00003

Table 3.4: Hyperparameters and pre-trained embeddings used for named entity recognition and relation extraction baseline results. For the NER task, the same architecture and hyperparameters were used for both general and fine-grained entity models. For the relation extraction task, the same hyperparameters were used with both the BERT and R-BERT architectures.

We achieved the highest micro-averaged  $F_1$  score of 81.3% on entities using SciBERT and 85.2% on relations using the R-BERT architecture with SciBERT<sup>1</sup>. Results of representative entities and relations are shown in Tables 3.5 and 3.6.

### 3.4.1 Annotation quality evaluation

To determine the quality of single-annotated documents compared to those which were double-annotated, we trained NER models (one for general and another for fine-grained entities, as in earlier experiments) using SciBERT with the 887 single-annotated documents

---

<sup>1</sup>A full listing of baseline prediction results can be found with the annotation guidelines at <https://github.com/uw-bionlp/clinical-trials-gov-annotation/wiki/Named-Entity-Recognition-and-Relation-Extraction-performance>.

Category	Entity	Count	biLSTM+CRF	PubMedBERT	SciBERT
Clinical	Condition	7,087	78.6 / 78.1 / 78.3	76.1 / 79.4 / 77.7	78.4 / 83.3 / 80.8
	Contraindication	142	93.7 / 78.9 / 85.7	77.4 / 80.0 / 78.6	100 / 96.6 / 98.3
	Drug	1,404	76.8 / 81.3 / 79.0	74.1 / 80.9 / 77.4	73.4 / 80.9 / 77.0
	Encounter	302	64.1 / 58.1 / 60.9	51.7 / 61.7 / 56.3	58.3 / 74.4 / 65.4
	Observation	2,558	74.3 / 66.1 / 69.9	67.9 / 73.5 / 70.6	72.1 / 77.6 / 74.7
Demographic	Procedure	3,016	68.4 / 75.5 / 71.9	67.0 / 75.9 / 71.2	71.3 / 79.4 / 75.1
	Age	708	91.3 / 95.4 / 93.3	82.4 / 88.5 / 85.3	99.1 / 98.3 / 98.7
	Birth	27	100 / 80.0 / 88.8	100 / 62.5 / 76.9	100 / 62.5 / 76.9
	Death	35	33.3 / 33.3 / 33.3	0.0 / 0.0 / 0.0	100 / 20.0 / 33.3
	Family-Member	147	40.0 / 19.0 / 25.8	33.3 / 55.5 / 41.6	44.9 / 61.1 / 51.7
Logical	Language	194	92.5 / 96.1 / 94.3	73.8 / 100 / 84.9	96.6 / 93.5 / 95.0
	Negation	952	74.3 / 82.7 / 78.2	60.9 / 73.1 / 66.4	73.5 / 82.9 / 77.9
	Assertion	1,157	66.6 / 62.8 / 64.7	56.1 / 58.9 / 57.5	62.1 / 65.8 / 63.9
	Modifier	3,464	65.0 / 58.3 / 61.5	59.2 / 64.0 / 61.5	58.5 / 65.4 / 61.8
	Polarity	360	82.5 / 88.0 / 85.1	74.6 / 67.4 / 70.8	81.4 / 79.5 / 80.4
Qualifier	Risk	117	93.1 / 96.4 / 94.7	91.3 / 91.3 / 91.3	95.4 / 91.3 / 93.3
	Severity	569	86.8 / 90.8 / 88.7	76.7 / 79.5 / 78.1	86.5 / 94.1 / 90.2
	Stability	397	84.2 / 67.6 / 75.0	79.4 / 75.0 / 77.1	75.3 / 84.7 / 79.7
	Criteria-Count	33	50.0 / 66.6 / 57.1	28.5 / 40.0 / 33.3	12.5 / 20.0 / 15.5
	Eq-Comparison	5,298	83.1 / 83.8 / 83.4	81.4 / 85.0 / 83.2	85.3 / 89.3 / 87.3
Comparative	Eq-Temporal-Period	2,057	88.7 / 89.2 / 88.9	70.0 / 73.9 / 71.9	82.6 / 86.3 / 84.4
	Eq-Temporal-Recency	131	68.7 / 84.6 / 75.8	43.4 / 55.5 / 48.7	50.0 / 66.6 / 57.1
	Eq-Temporal-Unit	1,808	95.1 / 97.6 / 96.4	97.4 / 98.1 / 97.8	98.2 / 99.4 / 98.8
	Eq-Value	3,835	91.8 / 95.3 / 93.5	95.5 / 96.2 / 95.9	96.4 / 97.1 / 96.7
Other	Location	371	68.5 / 58.7 / 63.2	65.4 / 71.6 / 68.3	73.4 / 78.3 / 75.8
-	Total	56,146	80.2 / 79.6 / 79.9	75.3 / 78.7 / 77.0	79.0 / 83.7 / 81.3

Table 3.5: **Baseline entity prediction scores (%)**, Precision / Recall / F<sub>1</sub>). Corpus-level micro-averaged scores are shown in the bottom row. For brevity a representative sample of entities is shown. *Count* refers to the total count of unique spans annotated in the entire corpus. Entities included in the total count and scores but omitted for brevity are *Acuteness*, *Allergy*, *Condition-Type*, *Code*, *Coreference*, *Ethnicity*, *Eq-Operator*, *Eq-Unit*, *Indication*, *Immunization*, *Insurance*, *Life-Stage-And-Gender*, *Organism*, *Other*, *Specimen*, *Study* and *Provider*.

Category	Relation	Count	SciBERT	R-BERT+SciBERT
Alternatives and Examples	Abbrev-Of	462	95.2 / 90.9 / 93.0	92.3 / 93.1 / 94.2
	Equivalent-To	516	61.5 / 69.5 / 65.3	59.6 / 67.3 / 63.2
	Example-Of	1,497	94.8 / 92.9 / 93.8	90.5 / 91.7 / 91.1
Clinical	Contraindicates	153	90.9 / 90.9 / 90.9	90.9 / 90.9 / 90.9
	Caused-By	726	63.0 / 86.4 / 72.9	78.6 / 86.4 / 82.3
	Found-By	293	90.4 / 59.3 / 71.7	79.3 / 71.8 / 75.4
	Treatment-For	457	69.2 / 69.2 / 69.2	61.7 / 74.3 / 67.4
	Using	405	73.8 / 83.7 / 78.4	66.6 / 64.8 / 65.7
Logical	And	821	54.1 / 60.0 / 56.9	53.8 / 53.8 / 53.8
	If-Then	261	57.6 / 65.2 / 61.2	55.5 / 65.2 / 60.0
	Negates	984	74.3 / 91.0 / 81.8	74.5 / 88.7 / 81.0
	Or	4,156	85.1 93.2 89.0	88.4 / 92.2 / 90.2
Qualifier	Asserted	1,184	83.7 / 89.0 / 86.3	85.9 / 89.0 / 87.5
	Modifies	3,400	90.9 / 94.2 / 92.5	92.2 / 95.4 / 93.8
	Risk-For	90	92.3 / 85.7 / 88.8	92.8 / 92.8 / 92.8
	Severity	529	80.2 / 96.6 / 87.6	86.3 / 96.6 / 91.2
	Stability	395	76.0 / 92.6 / 83.5	76.4 / 95.1 / 84.7
Temporal and Comparative	After	166	75.0 / 70.5 / 72.7	72.2 / 76.4 / 74.2
	Before	320	70.2 / 86.6 / 77.6	78.1 / 83.3 / 80.6
	Duration	243	59.3 / 79.1 / 67.8	64.5 / 83.3 / 72.7
	During	350	66.6 / 68.7 / 67.6	63.6 / 65.6 / 64.6
	Numeric-Filter	1,957	84.6 / 93.3 / 88.7	85.7 / 92.3 / 88.8
	Minimum-Count	173	64.2 / 69.2 / 66.7	71.4 / 76.9 / 74.0
Other	Temporality	2,645	80.7 / 90.7 / 85.4	81.8 / 92.2 / 86.7
	Location	207	64.2 / 94.7 / 76.6	69.2 / 94.7 / 80.0
-	Total	24,379	80.2 / 88.2 / 84.0	82.5 / 88.0 / 85.2

Table 3.6: **Baseline relation prediction scores (%, Precision / Recall / F<sub>1</sub>)**. Corpus-level micro-averaged scores are shown in the bottom row. For brevity a representative sample of relations is shown. *Count* refers to the total count annotated in the entire corpus, including relations not shown. The count total excludes general to fine-grained entity relations, which as overlapping spans are not used for relation prediction. Relations included in the total count and scores but omitted for brevity are *Acuteness*, *Code*, *Criteria*, *Except*, *From*, *Indication-For*, *Is-Other*, *Max-Value*, *Min-Value*, *Polarity*, *Provider*, *Refers-To*, *Specimen*, *Stage*, *Study-Of* and *Type*.

and evaluated on the 119 double-annotated documents. The results were a precision of 79.7%, recall of 82.5%, and an  $F_1$  score of 81.4%, which are very close to the highest performance of our randomly split train/test set results shown in Table 3.5. These results indicate relative uniformity and consistency in the corpus across both single- and double-annotated documents.

<b>Training Set</b>	<b>Test Set</b>	<b>Precision</b>	<b>Recall</b>	<b><math>F_1</math></b>
Manual	Semi-automated	75.4	82.1	78.6
Semi-automated	Manual	80.1	79.9	80.0

Table 3.7: **Results of NER experiments using the manually annotated and semi-automated portions of the corpus.** The manually annotated portion includes 513 documents while the semi-automatically annotated portion is 493 documents.

As the latter near-half (493 documents) of the LCT corpus was automatically annotated, then manually corrected, we also evaluated the quality of the manually annotated portion versus the semi-automatically annotated portion to ensure consistency. We first trained NER models with SciBERT using the manually annotated portion and tested on the semi-automated portion, then reversed the experiment and trained on the semi-automated portion and tested on the manually annotated portion. Results are shown in Table 3.7.

Results of the experiments when training on both the manually and semi-automatically annotated halves of the corpus show comparable results, with the greatest difference being in precision, with the manual annotation-trained model performing slightly worse (-4.7%) in prediction versus the semi-automated annotation-trained model. Overall  $F_1$  scores were similar at 78.6% and 80.0%, suggesting reasonable consistency across the corpus.

### **3.5 Limitations**

The LCT corpus is designed as a granular and robust resource of annotated eligibility criteria to enable models for entity and relation prediction as means of query generation. The corpus does have a number of limitations however which should be recognized. First, the corpus is largely singly annotated, with 119 of 1,006 documents (11%) double annotated and reconciled, while double annotation is generally considered to be the gold standard in the NLP research community. However, the reasonably high  $F_1$  score from experiments to evaluate NER when training on the singly annotated portion of the corpus suggests relative consistency of annotation across both single and double annotated documents. Additionally, entities in roughly half of the LCT corpus (493 documents) were automatically predicted, then manually corrected. This can potentially lead to data bias if predicted entities are not thoroughly reviewed and corrected by human annotators. Similar results from our experiments to detect differences in performance by training on the manually annotated portion versus the semi-automatically annotated portion ( $F_1$  scores of 78.6% and 80.0%) suggest this may not be not a significant issue.

### **3.6 Summary**

We created the LCT corpus, a gold standard human-annotated corpus of highly granular clinical trial eligibility criteria annotations. The NER and relation extraction models trained on this corpus achieved high performance and enable our subsequent aims.

## Chapter 4

# LEAF LOGICAL FORMS CORPUS

### **4.1 Overview**

This chapter examines the motivation for and creation of the LLF corpus, which we train a Seq2Seq model on for transforming input free-text eligibility criteria into logical forms used for generating database queries. Section 4.2 discusses complications in query generation and motivation for logical form structures. Section 4.3 examines related work in eligibility criteria logical representation. In Section 4.4 we describe steps in creating the corpus annotations. Section 4.5 shows results of our experiments, and Section 4.6 provides a summary of this chapter.

### **4.2 Motivation**

The LCT corpus, as examined in Chapter 3, is a human-annotated corpus of entities and relations within clinical trial eligibility criteria. Entities connected by relations can be imagined as a graph of nodes (entities) and edges (relations) between them. For example, the criterion, "*Diagnosed with type 2 diabetes mellitus within the past 3 years*", may be represented as a *Condition* node (diabetes mellitus) connecting to a *Equality-Comparison* node (past 3 years) by a *Temporality* edge.

We found SQL query generation using LCT-based graphs to be in many cases error prone and difficult for a number of reasons. First, nested *Or* and *And* relations tend to be problematic when represented as edges within a graph. For example in a criterion such as:

*Either of the following:*

1. *BMI > 35 in past 6 months*

## 2. Previous diagnosis of morbid obesity or obesity with one or more comorbidities

An *Or* relation would exist between "*BMI > 35*" and "*morbid obesity or obesity with one or more comorbidities*", as well as a nested *Or* between "*morbid obesity*" and "*obesity*" as well as a nested *And* between "*obesity*" and "*comorbidities*". In such cases, query generation is challenging as a system must recursively evaluate dependencies between entities to determine the scope of Boolean relations. This problem can be doubly challenging as the complexity of the underlying statement means that predictions of entities and relations by models will also necessarily be imperfect.

An alternative to this approach is to use a so-called intermediate representation (IR), which transforms the original natural language by removing "noise" unnecessary to a given task and which more logically represents underlying semantics, also known as a Semantic Parse [89] (see Herzig *et al* [90] for an examination of IR-based SQL generation approaches). Similar to earlier work using Description Logics, Roberts and Demner-Fushman [91] proposed a representation of questions on EHR databases using a comparatively compact but flexible format using first order logic expressions, for example, representing "Is she wheezing this morning?" as

$$\delta(\lambda x. has\_problem(x, C0043144, status) \wedge time\_within(x, "this morning"))$$

This style of representation is powerfully generalizable, but also difficult to translate directly into SQL statements as multiple predicates (e.g., *has\_problem* and *time\_within*) may correspond to one or many SQL statements, depending on context, complicating direct transformation into queries.

We thus chose a similar intermediate representation (hereafter simply "logical forms") as proposed by Roberts and Demner-Fushman but more closely resembling a nested functional structure in programming languages such as Python or JavaScript and more amenable to SQL generation. A criterion such as "Diabetic women and men over age 65" would be represented by our logical forms as

```

intersect(
    cond("Diabetic"),
    union(female(), male()),
    age().num_filter(eq(op(GT), val("65")))
)

```

An example input criterion and logical form output is shown in Figure 4.1.

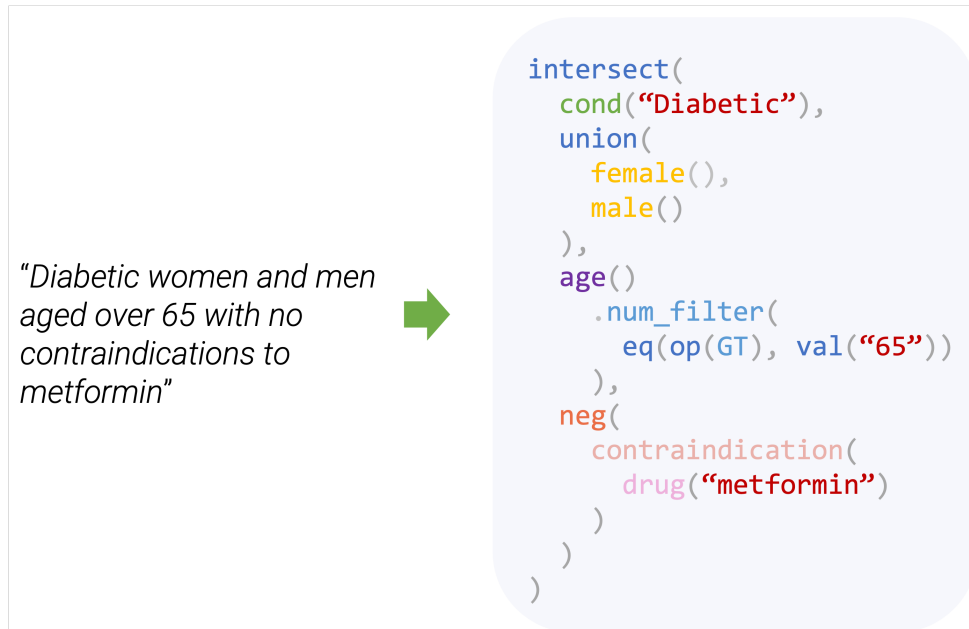


Figure 4.1: An example semantic parse using the LLF corpus annotation schema. The hypothetical input sentence is on the left, while the corresponding logical form is shown on the right. Indentations and colors have been added for readability.

### 4.3 Related Work

A number of logical (sometimes called "formal") representations of eligibility criteria have been explored by researchers. These include ad hoc expression languages such as EON [92], SAGE [93], ERGO [94], and EliXR [64], typically in tree-like dependency structure, Arden

Syntax, a prescribed syntax of operators and operations related to health data [95], logic-based languages such as Protégé [96] and SQL, and others [97, 98, 99]. See Weng *et al* [100] for a comprehensive review. As discussed in the previous section, more recent work by Roberts and Demner-Fushman proposed the use of a First Order Logic-based forms [91] which have been demonstrated to work well in tasks such as question answering [101].

#### 4.4 Methods

We developed annotation guidelines for the LLF corpus using a simplification of entities and relations from the preceding LCT corpus[18]. Generally speaking, LCT *entities* correspond to logical form *functions*, while LCT *relations* correspond to logical form *predicates*. For example, the LCT *Condition* entity has a corresponding *cond()* function, while the *Num-Filter* relation has a corresponding *.num\_filter()* predicate. The LLF annotation guidelines can be found at <https://github.com/ndobb/clinical-trials-seq2seq-annotation/wiki>.

We also hypothesized that the performance of predicting logical forms could likely be improved by replacing "raw" tokens in each eligibility criteria with corresponding logical form names derived from named entities from the LCT corpus. For example, given the eligibility criterion:

"*Diabetics* who *smoke*",

we would replace the named entities for "*Diabetics*" and "*smoke*":

*cond("Diabetics") who obs("smoke")*

using *Condition* and *Observation* annotations in the LCT corpus. We call this substituted text an "augmented" eligibility criteria. The augmented criteria syntax reshapes named entities to more closely resemble expected logical form syntax and allows us to leverage the LCT corpus for logical form transformation.

Creation and annotation of the LLF corpus proceeded in the following steps:

1. We randomly chose 2,000 lines of eligibility criteria from the LCT corpus, limited to only criteria which included at least one named entity and which were not annotated as hypothetical criteria. 30% of the 2,000 lines (600) were randomly chosen among lines with particularly complex entity and relation types, such as *If-Then*, *Before-After*, *Contraindication*, etc.
2. Each annotation file consists of the text "EXC" if exclusion or "INC" if inclusion (line 1), an original "raw" eligibility criteria (line 3), an augmented eligibility criteria (line 5), and an (initially blank) expected logical form equivalent to annotate (line 7). An example annotation is shown in Figure 4.2.
3. 3 informatics graduate students met weekly for 2 months to review annotations. Annotators were initially trained on 20 triple-annotated training annotations.
4. After training, each annotator was assigned a batch of 100 sentences (one per file) and tasked with writing a logical form version of each.
5. After each batch was completed, we executed a quality control script to parse each logical form annotation to ensure consistency. Any syntax errors were reported to and corrected by the annotators.
6. Annotators received additional batches of files to annotate until all 2,000 single-annotated annotations had been completed.

The pair-wise mean inter-annotator agreement by BLEU score was 82.4%. After annotations were completed, we experimented with predicting logical forms by fine-tuning T5 [102] Seq2Seq models. The T5 architecture and pre-trained models are widely used for and achieve at or near state-of-the-art for many machine translation and semantic parsing tasks.

```
'INC'

'– women age 20 – 34 years ;'

'– female() age() eq(val("20"), op(BETWEEN), val("34"), temporal_unit(YEAR)) ;'

intersect(
    female(),
    age()
        .num_filter(
            eq(val("20"), op(BETWEEN), val("34"), temporal_unit(YEAR))
        )
)
```

Figure 4.2: An example LLF corpus annotation. The annotation file is saved in JavaScript (.js) format, which enables syntax highlighting and validation to assist annotators. Whether a given criterion was an inclusion or exclusion criteria is indicated at the top, followed by the original raw text, then augmented text. The final annotated logical forms are shown last.

Following earlier work on task-oriented dialog semantic parsing structures in the domain of digital assistants, we also experimented using various alternative input-output syntax styles from our original logical forms:

1. **Shift-Reduce.** Einolghozatic *et al.* [103] used square brackets instead of parentheses and blank spaces instead of commas. We followed Rongali *et al.*'s suggestion to add a trailing repeat of function names to improve performance.
2. **Pointer.** Rongali *et al.* [104] found that replacing input tokens with " $@ptr_{index}$ ", where  $index$  corresponds to a token's sequential position in the input text improved performance in their semantic parsing task. We modified this approach by omitting the characters "ptr" and using the sequential position of the quoted span as our index rather than individual token positions.

## 4.5 Results

We used a randomly sorted 70/20/10 train/test/validation split of the LFF corpus to fine-tune the pretrained T5<sub>base</sub> model using combinations of these syntax styles. We call our gold standard annotated logical form syntax "Standard" style. Example inputs, outputs, and training results are shown in Table 4.1.

We found that our Standard logical forms achieved the highest performance using both BLEU [105] and ROUGE-L [106] scores, two commonly used metrics in measuring Seq2Seq performance. Replacing raw tokens with function names corresponding to named entities also significantly improved performance (+14.7%, comparing raw text to Standard input styles), demonstrating that leveraging the LCT corpus to generate augmented text achieved relatively high performance (> 93% BLEU score) for this task. As it was the highest-performing syntax style and also the most straightforward to parse, we chose to use the Standard logical form style as our IR for this project.

## 4.6 Summary

We created the LLF corpus, a gold standard human-annotated corpus of 2,000 eligibility criteria and corresponding logical forms. The LLF syntax was developed based on study of past work and entities and relations present in the LCT corpus. Our best performing prediction method using a fine-tuned T5 model achieved > 93% BLEU score.

Syntax Style	Example Input	Example Logical Form	BLEU	ROUGE-L
Raw-text→ Standard	Diabetics who smoke	$\begin{aligned} & \text{intersect}( \\ & \quad \text{cond}("Diabetics"), \\ & \quad \text{obs}("smoke") \\ & ) \end{aligned}$	78.7	79.1
Standard	cond("Diabetics") obs("smoke")	$\begin{aligned} & \text{intersect}( \\ & \quad \text{cond}("Diabetics"), \\ & \quad \text{obs}("smoke") \\ & ) \end{aligned}$	<b>93.5</b>	<b>92.3</b>
Standard+ Pointer	cond(@1) who obs(@2)	$\begin{aligned} & \text{intersect}( \\ & \quad \text{cond}(@1), \\ & \quad \text{obs}(@2) \\ & ) \end{aligned}$	93.3	91.2
Shift-Reduce	[cond "Diabetics" cond] who [obs "smoke" obs]	$\begin{aligned} & [\text{intersect} \\ & \quad [\text{cond } "Diabetics" \text{ cond}] \text{ who} \\ & \quad [\text{obs } "smoke" \text{ obs}] \\ & \quad \text{intersect}] \end{aligned}$	89.8	91.7
Shift-Reduce+ Pointer	[cond @1 cond] who [obs @2 obs]	$\begin{aligned} & [\text{intersect} \\ & \quad [\text{cond } @1 \text{ cond}] \text{ who} \\ & \quad [\text{obs } @2 \text{ obs}] \\ & \quad \text{intersect}] \end{aligned}$	89.4	90.4

Table 4.1: Example inputs and logical form syntax styles with fine-tuning performance results using the T5<sub>base</sub> model.

## Chapter 5

# KNOWLEDGE BASE

### 5.1 Overview

Storing biomedical information such as conceptual mappings, controlled vocabularies and terminologies, synonyms, hyponyms, and hypernyms enables a great variety of useful capabilities for a natural language interface for cohort discovery. For example, enabling users to simply specify "Patients without contraindications to Metformin"—without needing to exhaustively list what such contraindications may be—saves user time and energy, and also possibly includes criteria they may not be aware of.

This chapter describes the development of our KB, which combines a variety of sources using a graph database of Resource Description Framework (RDF) [107] triples. Section 5.2 describes our motivation for creating this resources, while Section 5.3 describes the data sources used and methods for KB population. Section 5.4 summarizes the work described in this chapter.

### 5.2 Motivation

While non-specific eligibility criteria can be categorized in a variety of forms, we noticed a number of frequently used patterns during the development of the LCT and LLF corpora. These categories of non-specific criteria formed the basis of the reasoning capabilities we aimed to enabled:

1. **Treatments for Conditions**, such as "*treated for myocardial infarction*". Determining whether a patient was treated for this requires first determining what possible treatment options exist for said condition.

2. **Contraindications to Treatments**, for example "*Contraindicated for MRI*" or "*With known contraindications to ACE inhibitors*". Contraindicated concepts must therefore be searched for procedures, surgical treatments, and drugs (including possible drug-drug interactions).
3. **Observations indicating a Risk**, such as "*At risk for suicide*" or "*At risk for heart attack*". Possible observations indicating risk thus include those for self-harm (such as depression) as well as underlying disease.
4. **Signs and Symptoms of a Condition**. These may include criteria such as "*Symptoms of depression*" or "*Showing signs of COVID-19*".
5. **Conditions affecting a Physiological Function**, for example "*Conditions affecting respiration*".
6. **Indicated Treatments for a Condition**, such as "*Indicated for anticoagulation therapy*" or "*Indicated for endoscopical drainage*".

Beyond facilitating reasoning on non-specific criteria, KBs also serve a vital role in other useful functions, such as determining relevant ICD-10 or SNOMED codes for a given diagnosis, or traversing a concept hierarchy to find relevant related concepts (e.g., determining a range of possible conditions associated with primary hypertension).

### **5.3 Methods**

#### *5.3.1 Data Sources*

Given its widespread use and breadth of vocabulary and terminology coverage, we chose to use the UMLS as the core of our KB. Despite the broad coverage of biomedical concepts it represents, however, the UMLS lacks certain data elements, specifically relations, needed to enable our goals for non-specific concept reasoning. For example, the UMLS contains

only limited data related to contraindications and symptoms. We thus evaluated informatics literature to find publicly available datasets and ontologies which may help fill these gaps. Importantly, as within the UMLS a Concept Identifier, or "CUI", forms the underlying global identifier linking the various biomedical source systems, we focused on the integration of sources directly (via CUI mappings) or indirectly (via ICD-10 or other coding systems) linkable to the UMLS. We integrated the following ontologies and data sources:

1. **The Disease Ontology (DO)** [20]. The DO is a comprehensive KB of human diseases. The DO features cross references to ICD-9, ICD-10, SNOMED, and other systems. For the purposes of this project, we leverage the DO's listing of symptoms for diseases with mappings to the Symptom Ontology (discussed next). For example, DOID\_0080642 ("Middle East respiratory syndrome") includes SYMP\_0000242 ("cough with bloody sputum") as a symptom. Symptoms related to COVID-19 were added to the DO at our request <sup>1</sup>.
2. **The Symptom Ontology (SO)** [21]. The SO is a large KB of symptoms. Like the DO, it includes external references to other vocabularies, including the UMLS. We leverage the DO and SO jointly by mapping both to UMLS CUIs.
3. **The COVID-19 Ontology** [22]. The COVID-19 ontology includes a variety of data elements and mappings related to COVID-19, such as transmission vectors, signs and symptoms, diagnostic methods, prevention and control, as well as genetic and molecular processes. For our purposes, we leverage risk factors for COVID-19 (COVID\_0000207), which in turn link to DO identifiers.
4. **Potential Drug-Drug Interactions (PDDI)** [23]. The PDDI is a large data set of potential drug-drug interactions. The PDDI was developed by merging 14 different

---

<sup>1</sup><https://github.com/DiseaseOntology/HumanDiseaseOntology/issues/916>

sources, including from clinical information sources, NLP corpora and pharmacovigilance sources. We used the PDDI for reasoning upon contraindications to medications.

5. **The Disease-Symptom Knowledge Base (DSKB)** [25]. The DSKB is a listing of diseases and corresponding symptoms derived using the NLP system MedLEE [108] on 25,074 discharge summaries from New York-Presbyterian Hospital. A random subset of the NLP-derived information was reviewed by a clinician for an overall recall of 90% and precision of 92%. The DSKB contains UMLS CUIs and was thus directly linkable to our KB.
6. **LOINC2HPO** [24]. LOINC2HPO is a data set of approximately 3,000 laboratory tests with corresponding phenotypes depending on their results (e.g., high, low, normal, positive, negative) linked to the Human Phenotype Ontology (which in turn is included in the UMLS). While not directly related to our reasoning use cases, we incorporated LOINC2HPO in order to find patients who may not have diagnosis codes for a given condition but whom a condition could be inferred via laboratory test results.

We further integrated a number of vocabulary mappings which we found potentially useful and not present within the UMLS. For example, an eligibility criteria may specify "Patients mechanically ventilated", which would be normalized to concept C0199470 ("Mechanical Ventilation") using MetaMapLite (discussed in Chapter 7). Within the UMLS however, C0199470 is not associated with any ICD-10 PCS codes, while within our enterprise data warehouse, mechanical ventilation events are recorded using ICD-10 PCS codes such as "5A19352", which relate to a different concept, C2695822 ("Respiratory Ventilation, Less than 24 Consecutive Hours"). As these concepts and codes are not linked within the UMLS, our KB would not be able to appropriately generate a database query to find mechanically ventilated patients. Using the following sources, however, enables our KB to fill this gap (among many others):

1. **ICD-9-CM to and from ICD-10-CM and ICD-10-PCS Crosswalk or General Equivalence Mappings** [26]
2. **ICD-9-CM Procedure Codes to SNOMED CT Map** [28]
3. **SNOMED CT to ICD-10-CM Map** [29]
4. **ICD-9-CM Diagnostic Codes to SNOMED CT Map** [27]

The integration of these additional mappings enable our KB to derive codes for a wide variety of use cases.

### *5.3.2 Graph Database Population*

A KB is an abstract, rather than technical concept, which may be instantiated as a relational or document database, graph, or other data structure. The practical needs and use cases related to eligibility criteria in this project drew us toward the use of a graph database as an optimal solution. An extensive comparison of graph databases versus relational other databases is outside the scope of this discussion. In brief however, both graph and relational databases are capable of storing data encompassing the same linkages and tuples [109], though by different means. Generally speaking, graph databases enable a more succinct query syntax (i.e., the same data extract can be accomplished using graph database queries linking tuples than one using a large number of SQL JOIN operations) and further do not necessitate a predefined schema, making them ideal for linking and exploring heterogeneous data sources.

We used GraphDB [110], a robust commercial graph database platform which is free for research use as our KB. To populate our KB, we modified open-source extraction scripts used for BioPortal [73] which extract UMLS data representations from a SQL database and output into the RDF-compatible format, Terse RDF Triple Language (Turtle) [111]. In the original

BioPortal extraction scripts<sup>2</sup>, UMLS CUIs are represented as strings, such as "C0150840", rather than Internationalized Resource Identifiers (IRIs), such as

*< http://bioportal.bioontology.org/ontologies/umls#C0150840 >.*

IRIs serve as critical structures in RDF for linking data sets together. Storing CUIs as strings results in duplication of data, performance degradation (graph databases optimize retrieval speed by caching IRI-based tuple linkages), and more complicated query syntax which runs somewhat contrary to RDF specifications [107]. We therefore modified<sup>3</sup> the BioPortal scripts to treat CUIs as IRIs. We similarly converted data sources not stored in an RDF-related format (such as CSV, HTML, or SQL) to Turtle files by Python script. All Turtle files were subsequently loaded into GraphDB.

### 5.3.3 SPARQL Query Development

After data were loaded in the KB, we developed SPARQL queries for each reasoning and hierarchy traversal use case. SPARQL is a graph query language developed by the W3C and features a syntax relatively similar to SQL [112]. For SPARQL queries related to reasoning use cases, development generally involved exploration of UMLS relations across various vocabularies, in addition to non-UMLS sources in our KB. In cases where more than one relation type was found, we traversed tuples from multiple vocabularies using *UNION* or *IN(<relation1, relation2, ... >)* syntax.

Below is an example of a SPARQL query for extracting concepts related to a sign or symptom. The script accepts a @cui parameter and traverses the KB, outputting sign and symptom-related CUIs and metadata from the DSKB, DO, and Omaha System ontologies (the latter contained within the UMLS but not updated since 2007).

---

<sup>2</sup><https://github.com/ncbo/umls2rdf>

<sup>3</sup><https://github.com/ndobb/umls2rdf>

---

```

# get_signs_and_symptoms_by_condition_cui.sparql
PREFIX rdfs: <http://www.w3.org/2000/01/rdf-schema#>
PREFIX skos: <http://www.w3.org/2004/02/skos/core#>
PREFIX umls: <http://bioportal.bioontology.org/ontologies/umls#>
PREFIX oms: <http://purl.bioontology.org/ontology/OMS#>
PREFIX dis: <http://purl.obolibrary.org/obo/DOID_>
PREFIX dskb: <http://purl.bioontology.org/ontology/DSKB#>

SELECT ?cui ?label ?sab ?code ?code_label ?tui
WHERE
{
  ?subj_iri skos:notation ?subj_cui .
  ?subj_iri skos:prefLabel ?subj_label .
  ?subj_iri ?rel1 ?obj1_iri .
  ?obj1_iri skos:notation ?cui .
  ?obj1_iri skos:prefLabel ?label .
  ?obj1_iri umls:code ?code_iri .
  OPTIONAL
  {
    ?code_iri skos:notation ?code .
    ?code_iri skos:prefLabel ?code_label .
    ?code_iri rdfs:domain ?ont_iri .
    ?ont_iri rdfs:label ?sab .
    ?obj1_iri umls:sty ?sty_iri .
    ?sty_iri skos:notation ?tui .
  }
  FILTER (?subj_cui = @cui)
  FILTER (?rel1 IN (
    dskb:hasSymptom,
    dis:hasSymptom
    oms:has_sign_or_symptom
  ))
}

```

---

## 5.4 Limitations

Our KB has a number limitations. First, the KB does not have a means of representing severity, probability, significance, or other nuanced relationships between a subject concept and reasoned concepts. For example, the PDDI data source lists certain drug-drug interactions as low in significance or only possible, and thus including those as contraindications may be unhelpful in certain cases. Chapter 8 demonstrates that users have the ability to edit reasoned concepts in our web application, which we believe makes this an acceptable

tradeoff. Second, our KB is capable of providing only UMLS concepts as output, which in certain cases this is not ideal. For example, our KB may return a concept such as "Old age" (C1999167) or "Low Platelet Count" (C5201036), which would be translated downstream into SNOMED or other codes for a structured query. We suspect that in most cases though, determining that a patient is over a certain age or has low platelet counts would be more accurately performed by querying structured demographic or laboratory result data directly (e.g., "*WHERE patient\_age >= 65*"). In a related fashion, more complex reasoning tasks, such as for patients "Indicated for bariatric surgery", require complex Boolean logic to be represented appropriately:

1. *Body mass index (BMI) is 40 or higher, called extreme obesity, OR*
2. *BMI is 35 to 39.9, called obesity, and have a serious weight-related health problem, such as type 2 diabetes, high blood pressure or severe sleep apnea.*
3. *In some cases, BMI is 30 to 34 and have serious weight-related health problems.<sup>4</sup>*

Our KB is currently unable to represent reasoning logic like this, as it can respond with only a list of UMLS concepts. In future work, we may examine adaptation of our KB to respond with logical forms (Chapter 4), which would be capable of representing these more complex cases.

## 5.5 Summary

This chapter describes the motivation and development of our KB, instantiated as a graph database which incorporates the UMLS and numerous other linked sources. Our KB forms the underlying foundation which enables reasoning on non-specific criteria (described in Chapter 7).

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<sup>4</sup>Adapted from <https://www.mayoclinic.org/tests-procedures/bariatric-surgery/about/pac-20394258>

## Chapter 6

# SEMANTIC METADATA MAPPING

### **6.1 Overview**

Any system which aims to generate queries to fulfill an aim or process will be necessarily limited in utility if it operates with only a single database schema. Within the health domain, data structures and vocabularies are increasingly complex and diverse, and systems capable of generalizing to other data models and institutions stand to be more impactful. Even the OMOP common data model, though widely used, is often updated to include custom database tables and data to fulfill various real-world needs [113, 114, 115, 116, 117, 118, 119].

This chapter explores what we call a Semantic Metadata Mapping, or SMM, which we leverage as a data-model agnostic means of generating queries for eligibility criteria. Section 6.2 explores related work. Section 6.3 discusses SMM structure and configuration. Section 6.4 summarizes the work of this chapter.

### **6.2 Related Work**

Many recent efforts at creating machine-readable data and resources for cross-system communication originate with the Semantic Web [120], a set of standards undergirded by ontologies, taxonomies, and inference rules designed for the Internet. As explained by the creators of the Semantic Web:

*"An ontology may express the rule "If a city code is associated with a state code, and an address uses that city code, then that address has the associated state code." A [Semantic Web] program could then readily deduce, for instance, that a Cornell University address, being in Ithaca, must be in New York State, which is in the U.S., and therefore should be formatted to U.S. standards."* [120]

The logic behind data model-agnostic query generation using an SMM is similar. For example, suppose an eligibility criteria specifies that patients identify as Hispanic. If an SMM contains a database column, *IsHispanic*, where a given tuple of value "1" means the tuple represents the UMLS concept of Hispanic (C0086409), a program can deduce then that a SQL query with a WHERE clause of "*WHERE IsHispanic = 1*" should retrieve patients who identify as Hispanic. This relatively simple logic is inspired by concepts of the Semantic Web and forms the basis of SMMs. Where SMMs differ most clearly, however, is the use of the UMLS as a means of managing metadata. To the best of our knowledge, this is the first project to do so to enable database query generation.

Using metadata for the purpose of generating database queries has been explored in previous research. Bizer and Seaborne [121] demonstrated that a simple ontology of database schema elements could be used to automatically generate SQL queries. Sequeda *et al* proposed similar methods using First Order Logic and created a publicly available rule engine for the research community [122, 123]. Knoblock *et al* and Gupta *et al* demonstrated the potential of such method for integrating heterogeneously structured databases of protein, gene, and metabolic pathway data [124, 125].

### 6.3 Methods

An SMM includes a listing of available databases, tables, columns, and so on within a given database schema. These database artifacts are "tagged" using a subset of UMLS concepts, many from the National Cancer Institute (NCI) and Health Level 7 (HL7) vocabularies. Examples of categories and concepts possible within an SMM include:

- **Demographic** - Age (C0001779), Ethnic Group (C00015031), Language (C0023008), Female (C0086287), Male (C0086582), Transgender (C3266856), Hispanic (C0086409).
- **Identifiers** - Patient (C5236161), Encounter (C3865224)
- **Metadata** - Concept Mapping (C3858752), Code (C0805701), Code System (C2347818),

Quantitative Value (C0392762).

- **Polarity** - Normal (C0205307), Abnormal (C0205161), Positive (C1514241), Negative (C0205160), Low (C0205251), High (C0205250).
- **Vital Status** - Living (C4551704), Deceased (C0011065).

Alternatively, a column may also be tagged using a UMLS Source Abbreviation, or SAB, such as ICD-10, SNOMED, LOINC, or RXNORM. Tagging a column as an SAB indicates that the tuple contents of a column represent codes associated with a given source system. For example, a column which contains diagnosis codes such as E11.62, I25.10, and others (ICD-10 codes) would have an SAB tag of "ICD10CM". SABs supported by our SMMs are:

- **Current Procedural Terminology (CPT)**
- **Healthcare Common Procedure Coding System (HCPCS)**
- **ICD-9**
- **ICD-10**
- **ICD-10 Procedure Coding System (ICD-10 PCS)**
- **LOINC**
- **National Drug File (NDC)**
- **NCI**
- **RxNorm**
- **SNOMED**

A column tagged by a UMLS concept or SAB can also be conditional, or in other words, only true given some other logic also being true. For example, Table 6.1 shows a hypothetical database table of diagnosis codes. Column *code* contains codes from various vocabularies, including ICD-9, ICD-10, and SNOMED.

<b>patient_id</b>	<b>coding_system</b>	<b>code</b>	<b>dx_name</b>
1	ICD-10	I50	Heart Failure
2	ICD-10	E87.70	Fluid Overload
3	SNOMED	4270721002	AIDS Associated Disorder
4	ICD-9	493.0	Asthma

Table 6.1: Hypothetical database table for patient diagnoses. Codes from a variety of different vocabularies are shown under the *coding-system* column.

In this case, *code* would be tagged with 3 conditional SABs, ICD-9, ICD-10, and SNOMED, with each SAB tag's condition defined as a SQL WHERE clause in the form of a string. For example, the "ICD10CM" tag would be paired with the condition, "WHERE coding\_system = 'ICD-10'". This is a machine-readable equivalent of, "The *code* column represents an ICD-10 code only if the *coding-system* column in the same record is of value 'ICD-10'".

Table 6.2 shows a more complete configuration example of all 3 SABs for the column.

<b>ColumnName</b>	<b>SAB</b>	<b>SqlWhere</b>
code	ICD9CM	coding_system = 'ICD-9'
code	ICD10CM	coding_system = 'ICD-10'
code	SNOMED	coding_system = 'SNOMED'

Table 6.2: Example SMM configuration for the *code* column.

The use of UMLS concepts, SABs, and optional conditions for each enables query generation across a wide variety of potential data models and database schema. An example

of this can be seen in Figure 6.1, which shows strategies by which a given criterion can be used to generate schema-specific queries by leveraging different SMMs. In cases where the LeafAI query engine finds more than one means of querying a concept (e.g., two SQL tables for diagnosis codes), the queries are combined in a UNION statement.

#### **6.4 Limitations**

Our SMM implementation is limited in a number of ways, most notably in that our tagging schema is not entirely machine readable. Specifically, the use of optional conditions in the form of SQL WHERE clauses implemented as simple strings (e.g., "WHERE IsPrimaryDx = 1") means that our system does not enable "understanding" of actual filtering logic or tuple-level values, but rather simply trusts that the given string will filter the condition as expected. In this sense the SQL WHERE strings are a "black box" to the SMM from a systems perspective. A more robust SMM representation may perhaps annotate tuple-level values semantically at a more granular level, though this would necessarily entail far more work on the part of the annotator in exhaustively describing every possible value of every column in a database.

#### **6.5 Summary**

This chapter examined our system for data model-agnostic query generation using SMMs, an encoding system where metadata on various clinical database schema elements are tagged using UMLS concepts and SABs.



Figure 6.1: The LeafAI query engine’s SQL query generation process using two hypothetical database schema to generate queries for platelet counts (shown in logical form after normalization). This example illustrates the flexibility of LeafAI’s semantic metadata mapping system (represented here in JSON format) in adapting to virtually any data model. On the left, ”Tall Table Structure”, platelet counts must be filtered from within a general purpose ”labs” table. The LeafAI KB recognizes that labs may be stored as LOINC codes, and the corresponding SMM indicates that records in this table can be filtered to LOINC values. On the right, ”Pivoted Table Structure”, platelet counts are stored as a specific column in a ”complete\_blood\_counts” table, and thus can be directly queried without further filtering. Additional metadata, columns, tables, types and so on needed in SMMs are omitted for brevity.

# Chapter 7

## QUERY GENERATION

### 7.1 Overview

This chapter describes our overall system for automatic query generation for clinical trial eligibility criteria using a natural language interface, which we call LeafAI. This system incorporates NER models (trained on the LCT corpus, described in Chapter 3), a Seq2Seq model (trained on the LLF corpus, described in Chapter 4), a KB (Chapter 5), and methods for database schema metadata tagging (Chapter 6). Section 7.2 recapitulates the need and motivation for this system. Section 7.3 summarizes relevant related research described earlier in Chapter 2. Section 7.4 describes methods for query generation leveraging components discussed in preceding chapters. Experimental setup and evaluation methods are described in Section 7.5, with results shown in Section 7.6. A discussion of results, implications, and limitations is presented in Section 7.7. Section 7.8 summarizes the content of this chapter.

### 7.2 Motivation

Identifying groups of patients meeting a given set of eligibility criteria is a critical step for recruitment into RCTs. Often, clinical studies fall short of recruitment goals, leading to time and cost overruns or challenges in ensuring adequate statistical power [34, 36]. Failure to recruit research subjects may result from a variety of factors, but often stems from difficulties in translating complex eligibility criteria into effective queries that can sift through data in the EHR [15]. Despite these difficulties, RCT investigators increasingly rely on EHR data queries to identify research candidates instead of labor-intensive manual chart or case report form review [126]. At the same time, the amount and variety of data contained in EHRs is increasing dramatically, creating both challenges and opportunities for patient recruitment

[127]. While more granular and potentially useful data are captured and stored in EHRs now than in the past, the process of accessing and leveraging that data requires technical expertise and extensive knowledge of biomedical terminologies and data models.

Given these challenges, automated or semi-automated means of identifying eligible patients in reducing time and costs in human labor while leveraging more complex but useful data are appealing. In particular, NLP-based cohort discovery methods could be especially valuable since they can key on eligibility criteria described in natural language, a medium that clinicians, researchers and investigators already use.

### **7.3 Related Work**

In this section we briefly summarize recent research in this domain, described earlier in Chapter 2.

Much research has explored methods for query generation, including encoder-decoder neural architectures for transforming clinical natural language questions into SQL queries [52, 53, 54, 55, 12]. The most prominent recent system and most comparable to our approach is Criteria2Query [6, 8]. Criteria2Query utilizes a combination of rule-based and neural modules to generate queries: a CRF-based NER model trained on a corpora of 230 Alzheimer’s Disease eligibility criteria documents [66], relation extraction using Dijkstra’s algorithm [128], BERT [57] for negation detection, Boolean logic detection using heuristics and dependency parsing, a Lucene-based [58] OMOP mapping tool, Usagi [129] for named entity normalization, and SuTime [82] for temporal expression normalization. For query generation, Criteria2Query composes the resulting query representation into JSON and leverages the Observational Health Data Sciences and Informatics (OHDSI) ATLAS [130] API to generate a SQL query.

#### *7.3.1 Gaps and opportunities*

Most programs capable of generating database queries do so for only a single database schema, such as OMOP or MIMIC-III [56]. This lack of flexibility limits their capability to accommodate real-world project needs [113, 114, 115, 116, 117, 118, 119], such as adding new

database tables to OMOP for cancer staging [113]. Moreover, most methods, particularly those using direct text-to-SQL deep learning approaches, tend to generate relatively simple SQL statements with few JOINs or nested sub-queries and typically no support for UNION operators and so on. This relative simplicity contrasts with the complexity of real-world EHR databases, which may contain dozens or even hundreds of tables using various vocabularies and mappings. Furthermore, direct text-to-SQL methods are bound to SQL syntax, and thus incapable of querying other systems such as Fast Healthcare Interoperability Resources (FHIR) [131]. Additionally, few of the methods described provide support for complex logic such as nested Boolean statements or temporal sequences, and none support reasoning on non-specific criteria (e.g., "diseases that affect respiratory function"), phenomena common to study eligibility criteria [15, 16]. Perhaps most importantly, to the best of our knowledge, only one previous work has been tested in terms of matching patients actually enrolled in clinical trials [9], and none have been directly compared to the capabilities of a human database programmer.

## 7.4 Methods

### 7.4.1 System Architecture

The LeafAI query engine was designed using a modular, micro service-based architecture with a central Application Program Interface (API) which orchestrates end-to-end query generation. Inter-module communication is performed using gRPC [132], a robust open-source remote procedure call framework which enables language-agnostic service integration. This allows individual modules to be implemented (and substituted) in programming languages and using libraries well-suited to a given task. We deploy each module as a Docker [133] container. A diagram of the LeafAI query engine architecture is shown in Figure 7.1.

At a high level, query generation is performed in the following steps:

1. A query request is received by the API in the form of inclusion and exclusion criteria as free-text strings.

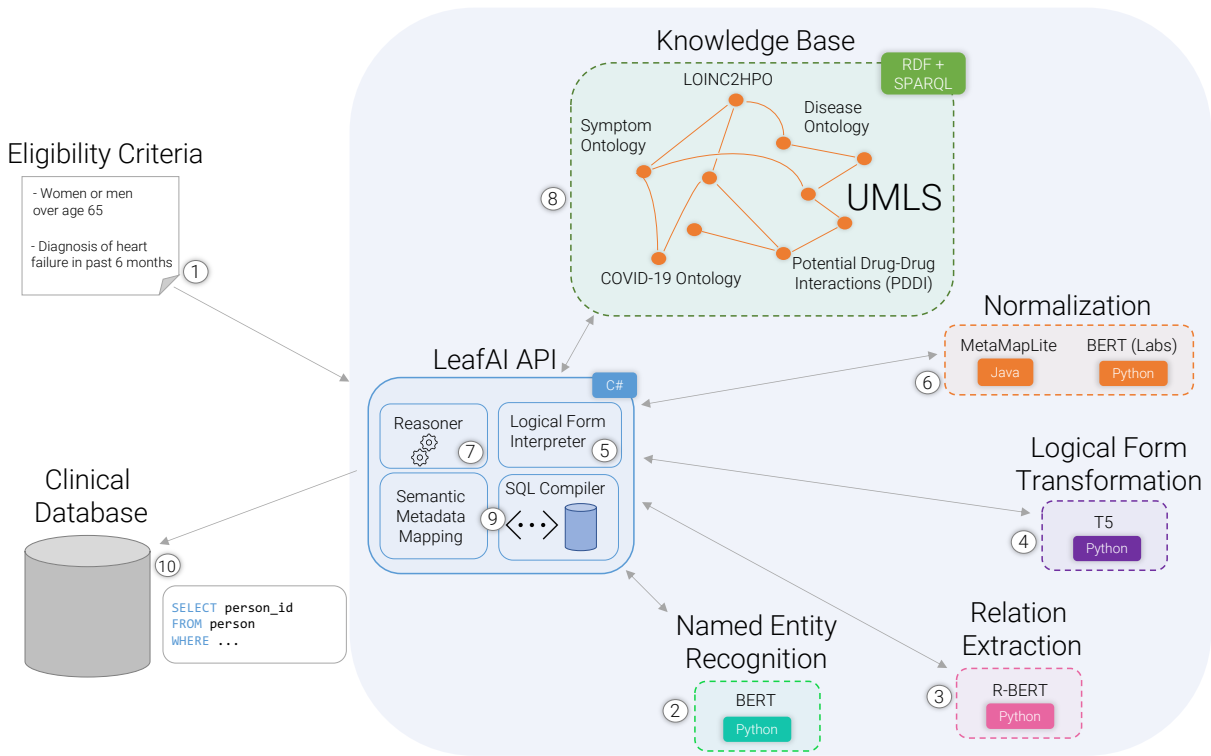


Figure 7.1: LeafAI query architecture. Inter-module communication is performed using the gRPC framework. Individual modules are deployed as Docker containers and communicate solely with the central API, which orchestrates query generation and handles query generation requests.

2. The input texts are tokenized and named entity recognition is performed to determine spans of text representing conditions, procedures, and so on.
3. Relation extraction is performed to determine relations between the entities, such as *Caused-By* or *Numeric-Filter*.
4. The input texts are transformed by replacing spans of "raw" text with logical form names. For example, "Diagnosed with diabetes" would become "Diagnosed with cond("diabetes")." The resulting input texts are in turn transformed into an output

logical representation using a Sequence to Sequence (Seq2Seq) architecture, in the form of a string.

5. A logical form interpreter module implemented as a recursive descent parser [134] reads the logical form string input and instantiates it as an abstract syntax tree (AST) of nested in-memory logical form objects.
6. "Named" logical form objects (i.e., specified with quoted text, such as "cond("diabetes")") are normalized into one or more corresponding UMLS concepts. UMLS child concepts are also added using our KB. For example, "cond("type 2 diabetes")" would also include concepts for type 2 diabetes with kidney complications (C2874072).
7. Working recursively inside-to-outside the AST structure, each logical form object calls a *Reason()* method which executes various rules depending on context.
8. Each reasoning rule is performed as one or more pre-defined SPARQL queries to the KB, concept by concept.
9. The final normalized, reasoned, logical form AST is thus a nested structure of UMLS concepts. Each AST criterion is mapped to zero or more corresponding entries in the semantic metadata mapping (SMM), which in turn lists meanings, roles, and relations of a database schema in the form of UMLS concepts.
10. The final mapped AST object is transformed into a series of database queries, one per line of eligibility criteria text. The output SQL query can either be executed directly on a database or returned to the API caller.

Figure 7.2 illustrates an example of this process. In the following subsections we examine these steps in detail.

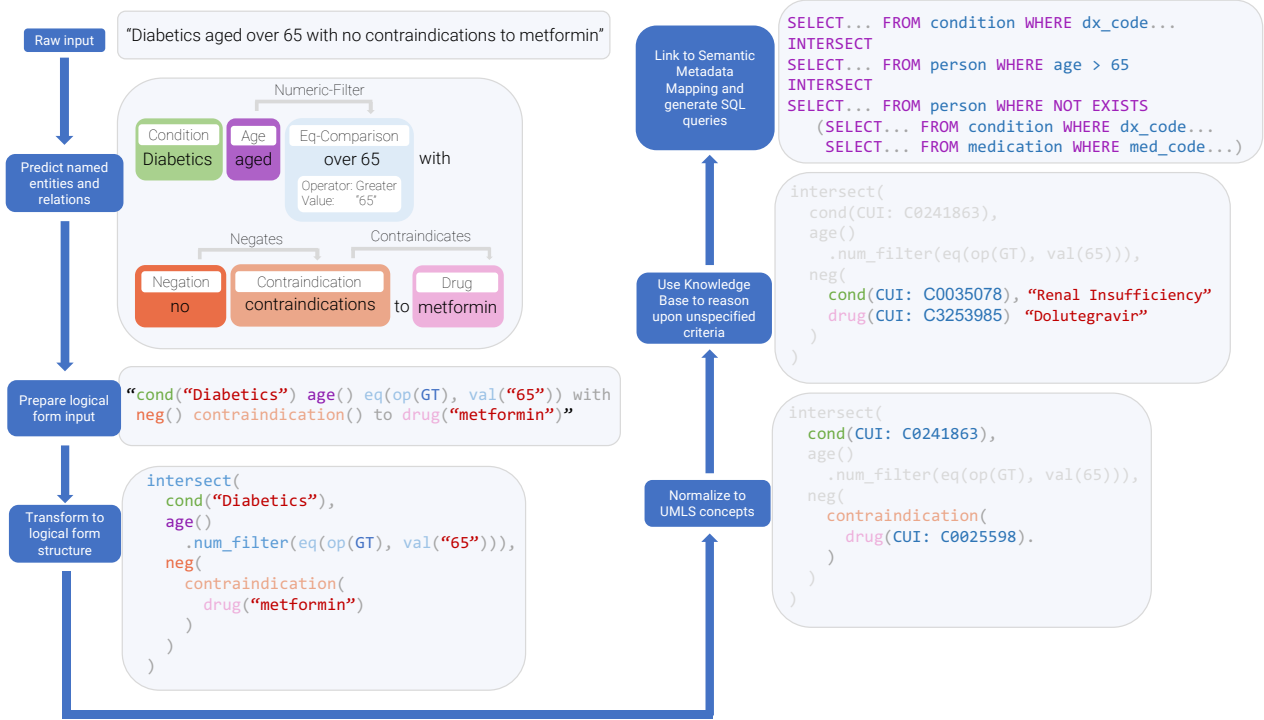


Figure 7.2: LeafAI query generation processes

#### 7.4.2 Named entity recognition and relation extraction

We used the LCT corpus to train two BERT-based NER extractors, one each for LCT general- and fine-grained-entities. Next, we perform relation extraction between named entity pairs similarly using a BERT-based model also trained on the LCT corpus.

#### 7.4.3 Logical form transformation

As discussed in Chapter 4, one of the core challenges of generating queries for eligibility criteria is the problem of logical representation. Generating queries directly based on named entities and relations alone, while practical, performs poorly in cases of nested or particularly complex logic.

After NER and relation extraction are performed, we leverage our best-performing T5

Seq2Seq model fine-tuned for predicting logical forms on the LLF corpus. As inputs to the Seq2Seq model we use the original eligibility criteria with named entity spans replaced by logical form representations, as we found this improved performance compared to training with raw inputs (described in Chapter 4). For example, the criterion

*”Women 55 and older diagnosed with Hepatitis C”*

would be transformed to the augmented criterion

*”female() eq(op(GTEQ), val(“55”)) diagnosed with cond(“Hepatitis C”)”*

and finally returned as the logical form

```
intersect(  
    female(),  
    age().num_filter(eq(op(GTEQ), val("55")))  
    cond("Hepatitis C"),  
)
```

from our Seq2Seq model. The returned logical form string would then be instantiated into an abstract syntax tree (AST) of nested in-memory logical form objects using a recursive descent parser [134] within our API.

#### 7.4.4 Concept normalization

We normalize ”named” logical forms to UMLS concepts using MetaMapLite [135, 136]. We consider a logical form ”named” if it contains a free-text value surrounded by quotes. For example, *cond()* is unnamed and refers to any condition or disease, while *cond(”hypertension”)* is named as it refers to a specific condition.

Normalization using MetaMapLite can often result in high recall but low precision, as MetaMapLite has no NER component and tends to return UMLS concepts which match

a given phrase syntactically but refer to abstract concepts not of interest (e.g., a search for "BMI" may return "body mass index" (C1305855), but also organic chemical "BMI 60" (C0910133)). To improve normalization precision, we employ two strategies. First, our NER component filters predicted UMLS concepts to only those of specific semantic types. For example, we limit condition concepts to only those related to semantic types of diseases or syndrome (dsyn) and so on. Next, using term-frequencies pre-computed across UMLS concept phrases, we compare term frequency-inverse document frequency (tf-idf) on MetaMapLite predictions, removing UMLS concepts whose summed matched spans have a tf-idf score lower than that of unmatched spans in a given named entity. For example, for the string "covid-19 infection", MetaMapLite predicts both "COVID-19" (C5203670) as well as several concepts related to general infections. Our tf-idf strategy removes general infection concepts because "infection" has a lower tf-idf score than the summed scores for "covid" + "-" + "19".

Laboratory values present a particular challenge, as LeafAI expects predicted lab concepts to have directly associated LOINC codes, while MetaMapLite typically normalizes lab test strings to UMLS concepts of semantic type "laboratory test or finding", but which do not have direct mappings to LOINC codes. For example, a search for "platelet count" returns the concept "Platelet Count Measurement" (C0032181), but not the needed concept of "Platelet # Bld Auto" (C0362994) with LOINC code "777-3". Thus similar to Lee and Uzuner with medications [137], we trained a BERT model for sequence classification to normalize lab tests. We trained this model to identify UMLS concepts associated with LOINC codes most frequently used in eligibility criteria [138], with each CUI as a possible class.

#### *7.4.5 Reasoning using an integrated knowledge base*

We leverage our KB (Chapter 5) in order to reason upon non-specific criteria using a combination of relatively simple rules within our API executed as SPARQL queries.

Our KB, nested logical forms, and inside-to-outside normalization methods enable "multi-hop" reasoning on eligibility criteria over several steps. For example, given the non-specific

criterion "Contraindications to drugs for conditions which affect respiratory function", our system successfully reasons that (among other results),

1. **Asthma** causes changes to **respiratory function**
2. **Methylprednisolone** can be used to treat **asthma**
3. **Mycosis** (fungal infection) is a contraindication to **methylprednisolone**

These features allow LeafAI to reason upon fairly complex non-specific criteria.

#### *7.4.6 Query generation using semantic metadata mapping*

To enable data model-agnostic query generation we leverage our SMM system (Chapter 6), implemented as SQL database tables. These include (essentially "meta") records of databases, tables, and columns, with corresponding foreign keys, UMLS CUI annotations, conditions, SAB encodings, and so on. The SMM is loaded from the LeafAI application database upon API startup.

### **7.5 Evaluation**

It is reasonable to expect that an NLP-based system for finding patients based on eligibility criteria would find many patients who actually enrolled in a real clinical trial — assuming that patients enrolled in those trials met the necessary criteria as determined by study investigators. While there are caveats to this approach (for example, certain diagnosis codes may be missing for some patients, etc.), we suggest that tools such as LeafAI be evaluated by their ability to handle real-world eligibility criteria and clinical data.

We compared LeafAI's results to that of a human database programmer experienced in the use of clinical databases and data extraction. Our evaluation was performed as follows:

1. We extracted metadata on 168 clinical trials from our EHR between January 2017 and December 2021 where at least 10 patients were indicated as enrolled and not withdrawn,

and the total number of raw lines within the eligibility criteria (besides the phrases "Inclusion Criteria" and "Exclusion Criteria") was less than or equal to 30.

2. By manual review, we excluded 22 trials with multiple sub-groups, as it would not be possible to know which eligibility criteria applied to which sub-group of enrolled patients.
3. To narrow the scope of our evaluation, we chose to evaluate only trials studying the following 7 disease groups: Cardiology, COVID-19, Crohn's Disease, Multiple Sclerosis (MS), Diabetes Mellitus, Hepatitis C, and Cancer. Using the "condition" field for each trial within metadata from <https://clinicaltrials.gov>, we filtered and grouped the remaining 146 trials into only those studying our diseases of interest.
4. We randomly chose 1 trial from each group, with the exception of Cancer, where given the large number of trials and variety of cancer types, we chose 2 trials. 427 patients were enrolled across the chosen 8 clinical trials.
5. Both the LeafAI query engine and human programmer created queries to find patients for each eligibility criteria, which we executed on an OMOP database derived from our EHR of our institution's entire research-eligible patient population.
6. To ensure results returned would be limited to only data available during the time of each trial, we replaced references to the SQL function for generating a current timestamp (*GETDATE()*) with that of each trial's end date, and similarly replaced OMOP table references with SQL views filtering data to only that existing prior to the end of a trial.
7. To ensure queries would be comparable to LeafAI, the human programmer was instructed to (1) ignore criteria which cannot be computed, (2) make a best effort to reason upon non-specific criteria (e.g., symptoms for a condition), (3) not check whether

patients found by a human query enrolled within a trial, and (4) skip criteria which cause an overall query to find no eligible patients.

## 7.6 Results

Results of the query generation experiment are shown in Table 7.1. Overall, LeafAI matched 212 of 427 (49%) total enrolled patients across 8 clinical trials compared to 180 (42%) found by queries of the human programmer. The mean per-trial percent of patients matched was 43.5% for LeafAI and 27.2% for the human programmer. LeafAI had a greater number of patients deemed eligible across all 8 trials, for a total of 27,225 eligible compared to 14,587 found by the human programmer.

<b>Condition</b>	<b>Enrolled</b>	<b>LeafAI</b>		<b>Human</b>		<b>Time (hrs)</b>
		<b>Matched</b>	<b>Eligible</b>	<b>Matched</b>	<b>Eligible</b>	
CL Lymphoma	83	80 (96%)	3,252	77 (92%)	2,382	1
Hepatitis C	42	33 (78%)	9,529	32 (76%)	9,372	4
Crohn's Disease	16	0 (0%)	113	1 (6%)	9	2
Cardiac Arrest	27	12 (44%)	4,792	0 (0%)	598	5
COVID-19	41	0 (0%)	0	0 (0%)	98	2
Multiple Sclerosis	196	77 (39%)	4,891	69 (35%)	1,016	3
Type 1 Diabetes	11	0 (0%)	1,006	1 (9%)	1,104	4
Ovarian Cancer	11	10 (91%)	1,667	0 (0%)	8	5
<b>Mean</b>		43.5%		27.2%		
<b>Total</b>	427	212 (49%)	27,225	180 (42%)	14,587	26

Table 7.1: Statistics for each clinical trial evaluated by the LeafAI query engine and human programmer. The number of enrolled and matched patients were determined by cross-matching enrollments listed within our EHR. The *Time* column indicates the number of hours the human programmer spent developing queries for each trial.

Table 7.2 shows the number of criteria which were skipped by LeafAI. Of the 103 total criteria across all 8 studies, LeafAI executed queries for 61 (59.3%) and skipped 5 (4.8%) as it found no patients and 42 (40.7%) because no computable concepts were found.

Condition	ID	# Crit.	# No Patients	# Not Computable	# Executed
CL Lymphoma	NCT04852822	4	0 (0%)	0 (0%)	4 (100%)
Hepatitis C	NCT02786537	8	0 (0%)	4 (50%)	4 (50%)
Crohn's Disease	NCT03782376	9	0 (0%)	4 (44.4%)	5 (55.5%)
Cardiac Arrest	NCT04217551	12	0 (0%)	8 (66.6%)	4 (33.3%)
COVID-19	NCT04501952	13	0 (0%)	6 (46.1%)	7 (53.8%)
Multiple Sclerosis	NCT03621761	14	1 (7.1%)	3 (21.4%)	10 (71.4%)
Type 1 Diabetes	NCT03335371	18	2 (11.1%)	8 (44.4%)	8 (44.4%)
Ovarian Cancer	NCT03029611	25	2 (8%)	9 (36%)	14 (56%)
<b>Total</b>		103	5 (4.8%)	42 (40.7%)	61 (59.3%)

Table 7.2: The LeafAI query engine's handling of eligibility criteria for each trial. The column *No Patients* indicates the count of criteria which would, if executed, cause no patients to be eligible. The column *Not Computable* indicates the count of criteria which LeafAI could not generate a query for, for various reasons. Both of these types of criteria were ignored by the system.

Figure 7.3 shows differences in query strategies for 4 trials between LeafAI and the human programmer.

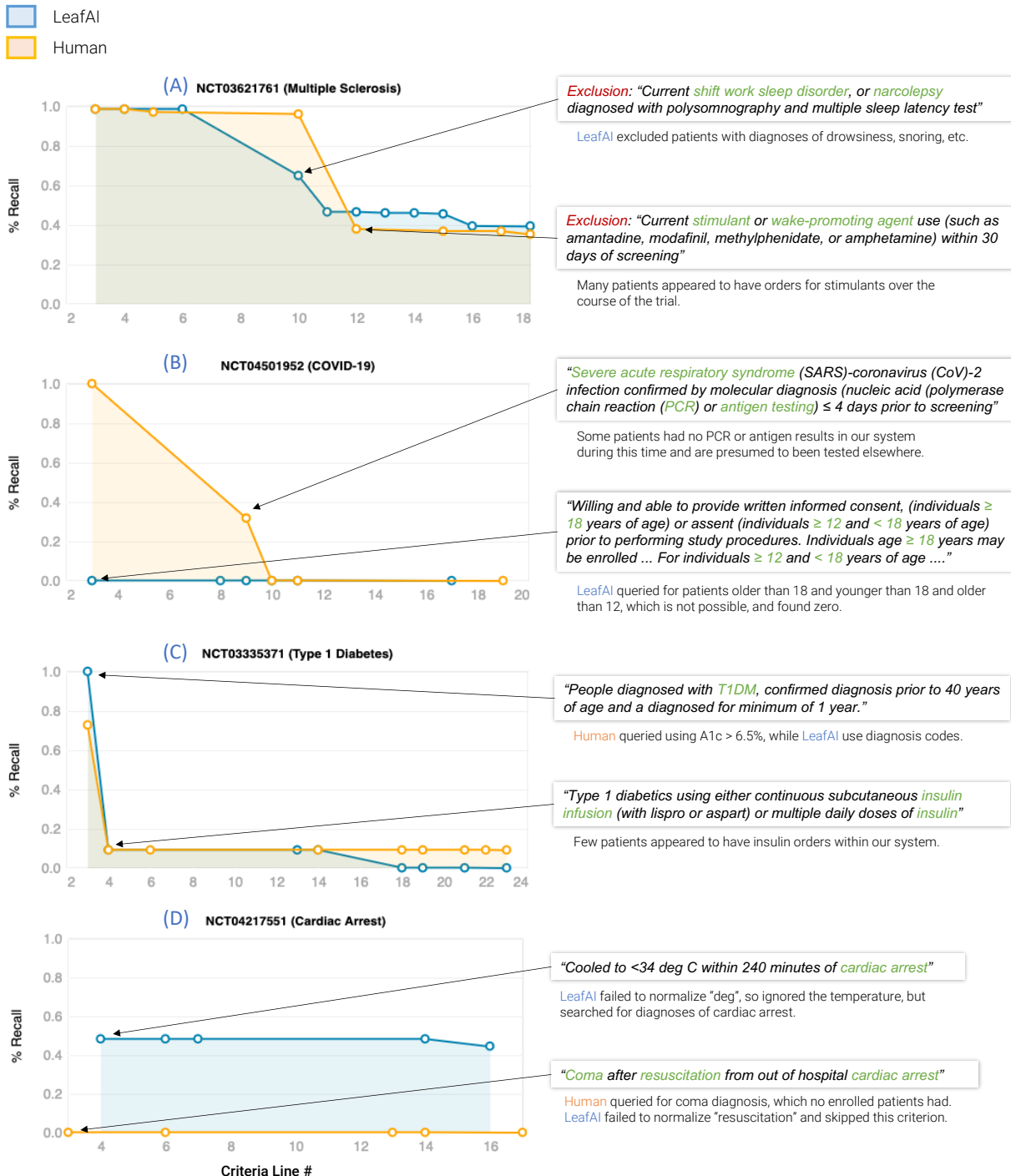


Figure 7.3: Longitudinal results for four trials. The blue line indicates recall for LeafAI and orange the human programmer. The X axis represents the line number for each eligibility criterion. Dots indicate that a query was executed for a given line. On the right, boxes represent the text of a criterion, with comments below discussing strategies and findings.

## 7.7 Discussion

Our results demonstrate that LeafAI is capable of rivaling the ability of a human programmer in identifying patients who are potentially eligible for clinical trials. Indeed, in numerous cases we found LeafAI and the human programmer executing similar queries, such as for Hepatitis C (NCT04852822), Chronic Lymphocytic Leukemia (NCT04852822), MS (NCT03621761), and Diabetes Mellitus (NCT03029611), where both ultimately matched a similar number of patients.

One notable pattern we found is that LeafAI consistently finds a higher number of potentially eligible patients. We hypothesize that in many cases, LeafAI’s KB played a key role both in finding additional eligible patients. For example, in the MS trial, LeafAI searched for 11 different SNOMED codes related to MS (including MS of the spinal cord, MS of the brain stem, acute relapsing MS, etc.), while the human programmer searched for only one, and ultimately LeafAI found nearly 5 times the number of potentially eligible patients (4,891 versus 1,016). It is possible that the human programmer had a lower rate of false positives (and thus higher precision); this will be explored in a future analysis. On the other hand, in the same trial, as can be seen in Figure 7.3 (A), given the exclusion criteria: ”Current shift work sleep disorder, or narcolepsy diagnosed with polysomnography and multiple sleep latency”, LeafAI’s KB unnecessarily excluded otherwise eligible patients by removing those with diagnosis codes for drowsiness, snoring, etc., as within the UMLS those are child concepts of sleep disorder (C0851578). The exclusion of these patients likely resulted in an approximately 40% drop in recall at that stage compared to the human programmer, though ultimately both achieved similar recall (LeafAI: 39% versus Human: 35%).

Beyond performance as measured by recall, it is notable that the human programmer spent approximately 26 hours crafting queries for the 8 trials while LeafAI took only several minutes running on a single laptop. The time saved by using automated means such as LeafAI for cohort discovery may save health organizations significant time and resources.

### *Limitations*

The LeafAI query engine and our evaluation have a number of limitations. First, while the 8 clinical trials we evaluated were randomly selected, we specifically restricted the categories of diseases from which trials were chosen and limited to trials with 30 or less lines of eligibility criteria, and thus our results may not generalize to other kinds of trials. Next, we evaluated our queries using an OMOP-based extract which did not contain the full breadth of data within our EHR. Had our experiments instead been conducted using our enterprise data warehouse (populated by our EHR), it is possible the human programmer would have achieved better results than LeafAI due to knowledge and experience in utilizing granular source data. For example, in the Cardiac Arrest clinical trial, the human programmer noted that data for use of cooling blankets is available in our EHR, but not in OMOP. It is not clear how LeafAI would perform were such data available. We further did not directly compare LeafAI to other NLP systems. While we considered evaluating Criteria2Query as part of our baseline, we ultimately determined that it was inappropriate for our analysis as we aimed to review results longitudinally (i.e., line by line of criteria), a function which Criteria2Query does not perform. Last, the number of truly eligible patients within our institution for each trial is unknown, which hampers our ability to measure system performance. We used each trial’s known enrolled participants as our gold standard, but assume they are only a (possibly small) subset of those actually eligible.

### **7.8 Conclusions**

This chapter introduced LeafAI, a NLP-based system leveraging deep learning and an integrated KB which can automatically generate queries for cohort discovery on virtually any clinical data model. Using an OMOP database representing the entire patient population of our institution, we demonstrated that LeafAI rivals the performance of a human programmer in identifying eligible research candidates.

## Chapter 8

# WEB APPLICATION

### 8.1 Overview

This chapter presents the goals and ongoing development of the LeafAI web application, an interactive user-facing application which integrates with the LeafAI API described in Chapter 7. As development of the web application is in progress, we leave evaluation of this to future work. Section 8.2 describes related work. Section 8.3 describes methods. Section 8.4 discusses future work, including user testing, and Section 8.5 concludes this chapter.

### 8.2 Related Work

Experimental systems enabling the querying of databases using natural language interfaces have been in development since the 1960s. Using relatively simple rule-based parsing systems, Woods [139] created a system for asking natural language questions of a moon rock database, while Epstein and Walker [140] similarly designed a natural language interface for a melanoma database. Decades later, Katz *et al* created START [141], a system capable of basic question-answering using data extracted and parsed from the internet. In the biomedical informatics domain, Cao *et al* developed AskHERMES [142], question-answering software capable of answering medical questions related to drugs, contraindications, and so on, using support vector machines (SVMs) and an internal knowledge base derived from the UMLS.

Cohort discovery systems using natural language are in many ways a subset of systems for question-answering which answer only one (unstated) question, "How many patients meet these criteria?". As discussed, in the biomedical informatics and clinical trials domain, the most well-known and cited system for matching patients to clinical trials using free-text

eligibility criteria is Criteria2Query [6, 8]. Criteria2Query offers a web-based simple and friendly user interface for inputting free-text eligibility criteria and enables users to correct erroneously normalized named entities.

### 8.3 Methods

The web application forms one component of the LeafAI application, which is a 3-tier architecture similar to Leaf [2]: a back-end with clinical and application databases, and server hosting an API (Chapter 7), and a user-facing web application. The web application features a chat-like design.

A chat-like interface for cohort discovery is both novel and a logical design choice given the natural language interface of LeafAI used for query generation. An example of LeafAI’s user interface is shown in Figure 8.1. As can be seen, we assume that the order of user-provided criteria is intentional and important, and leverage that assumption to both structure queries incrementally to report results line by line, with each reported result (e.g., ”421 are aged between 18 and 65”) effectively a subset of the preceding result.

The web application responds to user input in a visual form that is *chat-like*, rather than the result of a general programmatic conversation agent. While the user interface and query generation methods lay a potential foundation for general-purpose question-answering more akin to a conversation agent, we limit our scope to cohort discovery. The user interface is being designed with the following goals:

1. **Accessible history.** Users will be able to immediately scroll to view previous findings.
2. **Rapid feedback and explainability.** LeafAI will perform NER and normalization as users are typing, similar to prefetching search engine responses. This will allow users to preemptively detect concepts and queries which may return unexpected results. LeafAI will also display incremental query results in real-time, providing users faster feedback so users may avoid waiting until all query results are complete.

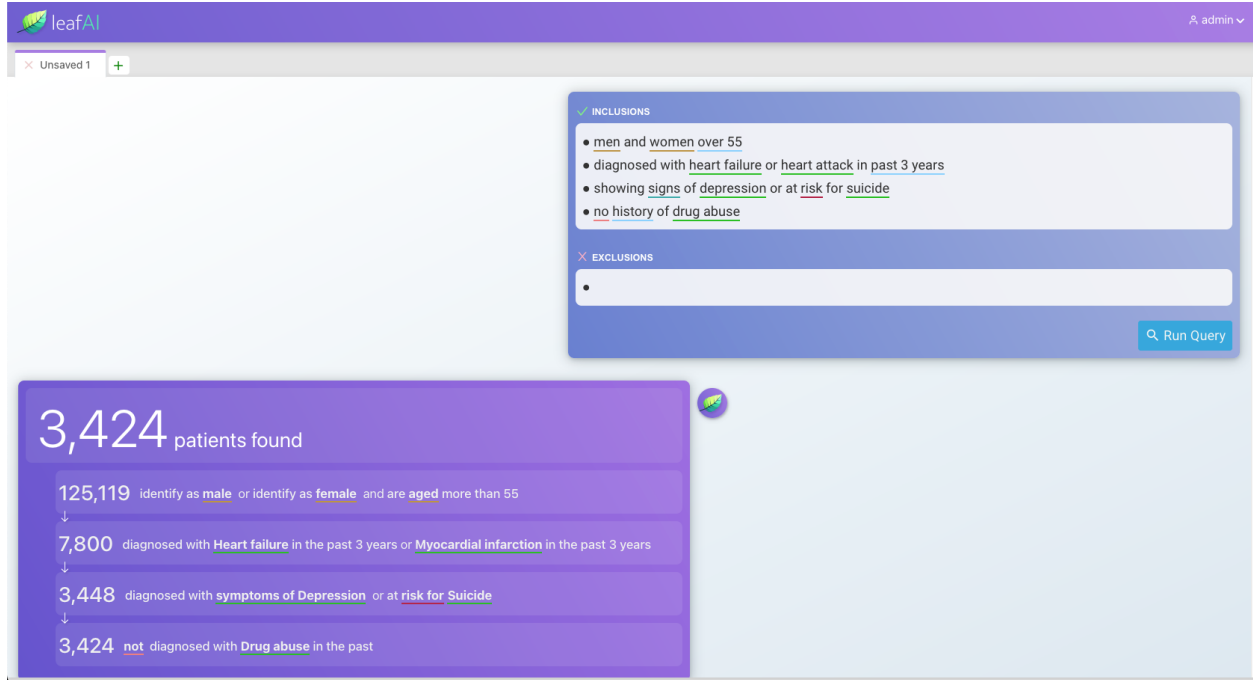


Figure 8.1: The web application user interface. User-entered criteria are shown in the above right, while system responses are shown in the lower left. User criteria are displayed and executed in order, with each count of patients representing a subset of the preceding count.

**3. Direct editing of responses enabling iteration.** As LeafAI returns results of patients found, the eligibility criteria used in a query will be directly editable, saving users’ time and facilitating quick iteration to find intended patients.

We next describe each goal in detail.

#### 8.3.1 Accessible History

Cohort discovery is a form of data exploration. As Derthick and Roth write, ”...the data exploration process is not characterized by monotonic progress towards a goal, but rather involves much backtracking and opportunistic goal revision” [143]. Put another way, user goals and perceptions may change over the course of exploration. With ubiquitous vertical

scrolling - where more recent actions and utterances are inserted downward while history is preserved upward - chat-like user interfaces facilitate user understanding of past utterances and actions. Persisted, easily viewable history of user utterances and actions enable what Gergle *et al* call "conversational grounding" [144], that is, accessible data to guide users to previously acquired findings and information. This history of previous actions can improve the pace of discovery and alleviate the need for users to (often imperfectly) attempt to recall their earlier findings and paths taken [145, 144].

### *8.3.2 Rapid Feedback and explainability*

Users' sense of system latency and responsiveness can significantly affect their satisfaction in using a tool [146, 147, 148]. Faster *preemptive* system responses (i.e., informing users' of a possible consequence before they complete an action) can also both save users' time and reduce loads placed upon systems by preventing unnecessary actions [149, 150]. The web application employs two general strategies for providing rapid feedback to users, one before queries are executed, and the second while results are being reported during query execution.

Consider a hypothetical case where a user begins to type a criteria but misspells "Diabetes Mellitus". Given no notification of the error, the user may take seconds or even minutes of additional typing while adding new criteria without realizing the initial spelling mistake. After the user awaits LeafAI's response, she finds that the query found no patients and is frustrated and confused at the counter-intuitive result, only to finally notice the misspelling, having wasted several minutes. The web application aims to avoid this scenario by preemptively performing NER and normalization while users are typing. Examples of this are shown in Figure 8.2. Named entities found within user criteria are underlined and interactive, enabling users to better estimate whether their queries will succeed or not.

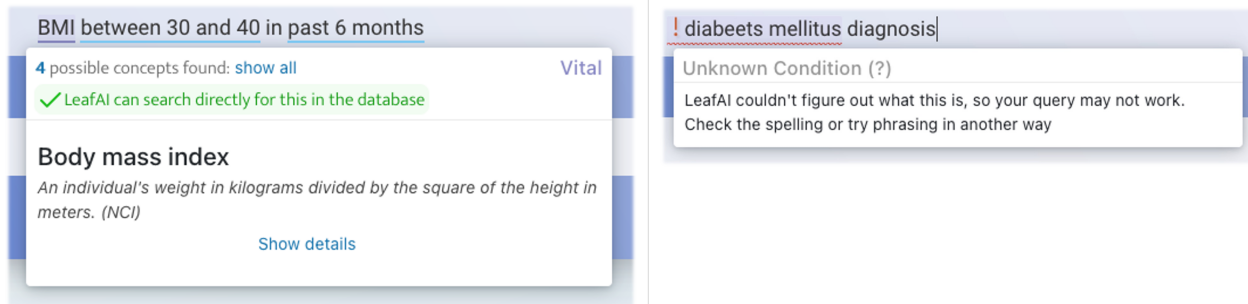


Figure 8.2: Normalized named entities identified in user input before query execution. On the left, the system correctly identified "BMI". On the right, "Diabetes Mellitus" is misspelled, and the user is notified.

Second, as LeafAI generates incremental queries and returns results line by line asynchronously, the user interface shows results as they are reported using a streaming interface. As a result, users will not need to wait until all queries are complete (as in tools such as Leaf and i2b2). An example of this is shown in Figure 8.3.

Taken together, these features for rapid feedback and transparency also enable *explainability* of system actions. That is, rather than simply returning a final count of patients meeting criteria, The web application provides information both before and during query execution of how the system interpreted user intent and how it has executed a query (e.g. what concepts it found or did not, misinterpreted, etc.). The system responses are generated using templated English expressions mapped to logical form types "slot-filled" using normalized UMLS concept names. By avoiding being a "black box", we aim for the system to gain user trust and understanding of how the system operates.

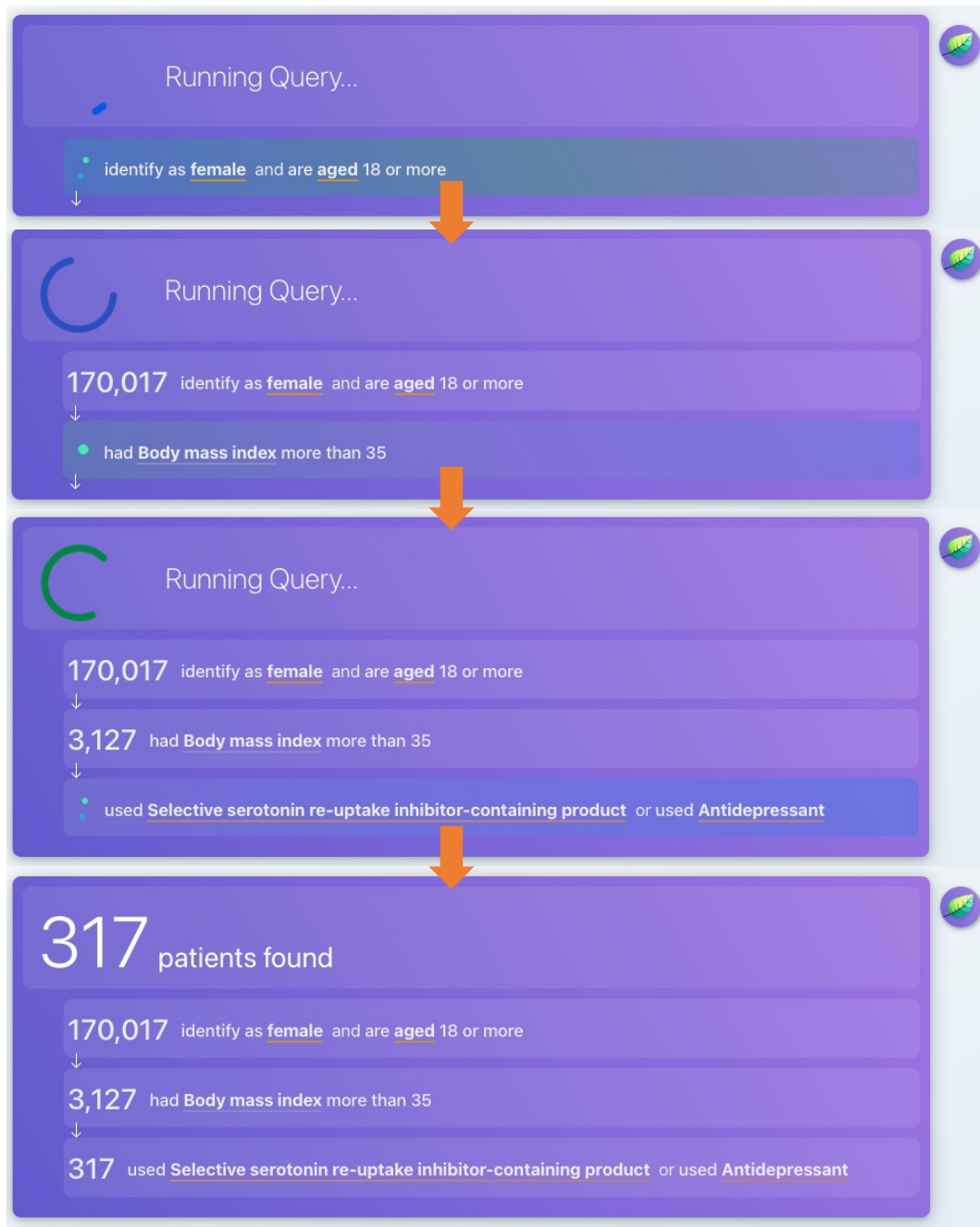


Figure 8.3: A time-lapse representation asynchronously reporting of incremental query results, from top to bottom. This is performed using two streaming interfaces, one from the clinical database to the API, and a second from the API to the web application.

### 8.3.3 Direct editing of responses enabling iteration

Data exploration is iterative: users explore, try, and learn over the course of multiple attempts. As discussed, faster, meaningful results also directly affect user satisfaction. To this end, the web application enables users to immediately and directly edit the returned system responses. This workflow is depicted in Figure 8.4.

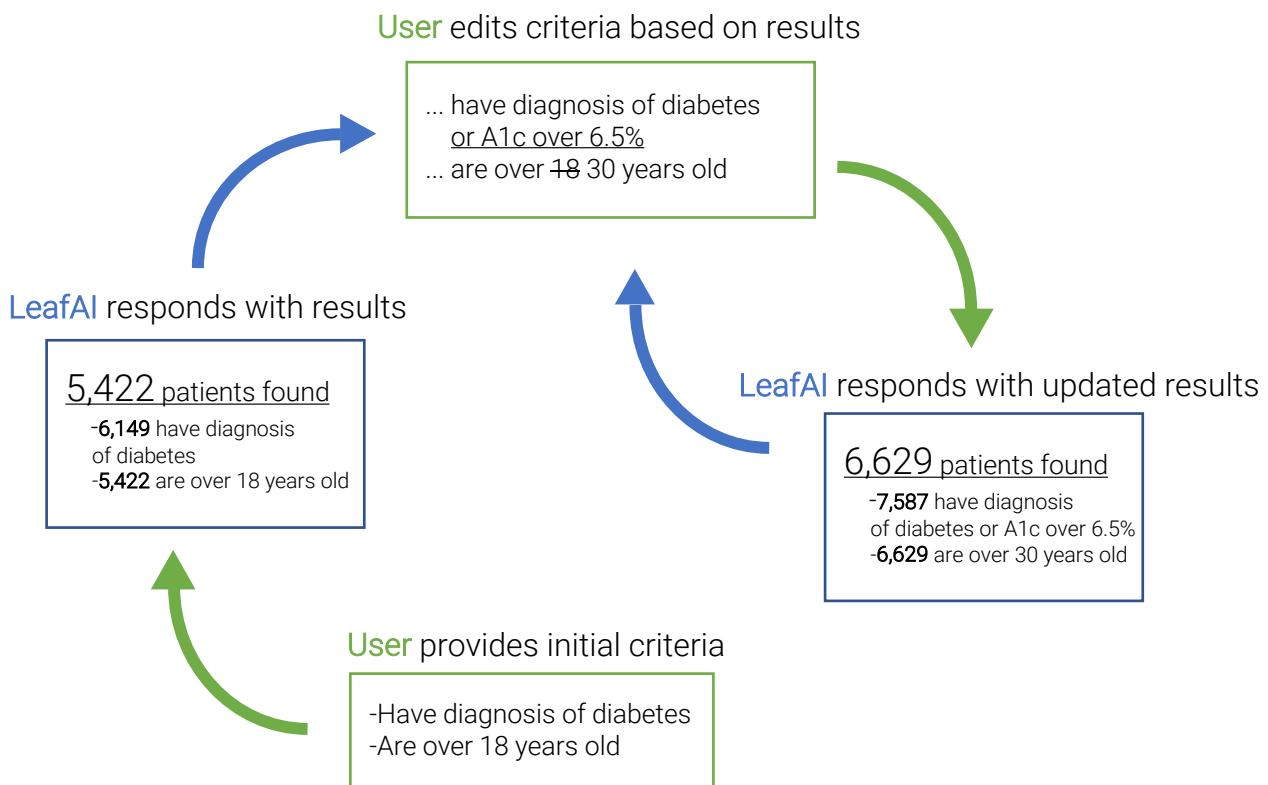


Figure 8.4: An iterative workflow using an example case for adults with Diabetes Mellitus. Users are able to directly edit results and re-execute their queries while preserving query history, saving user time and preserving previous user actions and findings.

For example, after seeing that LeafAI found less patients than expected, a user may realize that she should slightly alter her original query to expand her search. Rather than needing to copy and paste her earlier criteria, instead she will be able to simply click and

modify the earlier results.

Additionally, as discussed, the LeafAI query engine is capable of reasoning upon non-specific criteria. In certain cases, however, the system's findings may be incomplete, incorrect, or undesired for various reasons. Rather than forcing users to accept imperfect reasoning, however, the web application allows users to directly enable or disable reasoned concepts. An example of this is shown in Figure 8.5. In addition to reasoned concepts, the web application enables users to enable and disable named entities from system responses and immediately re-execute queries with a single click, shown in Figure 8.6. This allows for fast iteration and saves user time, obviating the need to alter the original inputs.

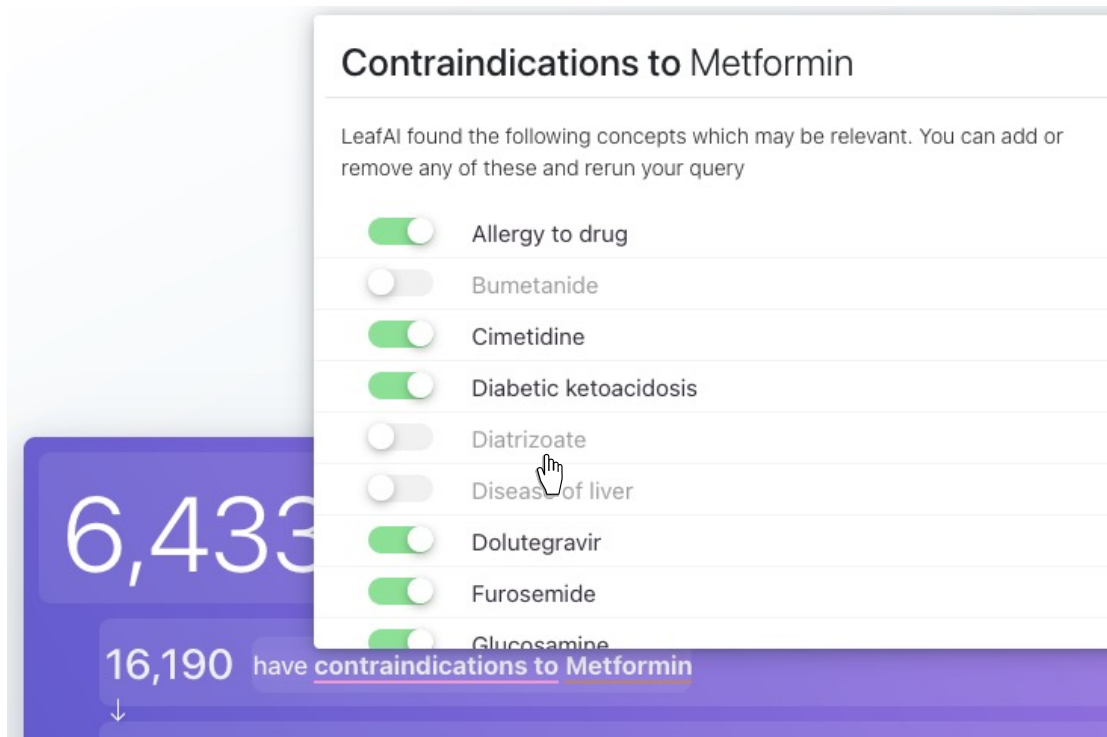


Figure 8.5: An example of a user editing concepts discovered using LeafAI's reasoning system. Users are able to enable or disable reasoned concepts.

The system's reasoning can thus be described as an optional means of helpfully saving users' time, which can be accepted, edited, or discarded as needed.

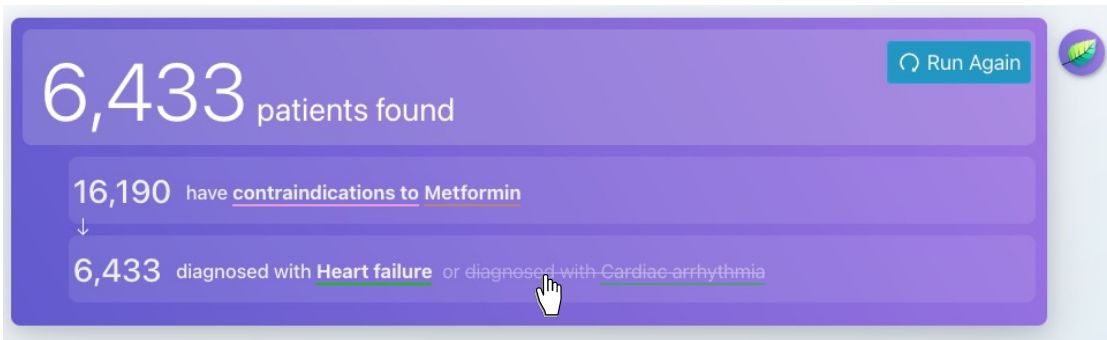


Figure 8.6: An example of a user disabling a named entity from a system response. Named entities can be enabled and disabled by clicking directly on them. Disabled entities are shown with gray colored text and struck through by a line. Upon re-execution, logic for disabled entities is removed from the generated query.

### 8.3.4 Model Inference Speed

LeafAI’s modules for NER, logical form transformation, lab normalization and so on use large Transformer-based neural models [151] with millions of parameters. Using these models as-is for inference can be slow, particularly when performed using CPUs rather than more costly but often far faster GPUs (Graphics Processing Units). Delays in inference time in turn can result in poor system latency, affecting user satisfaction. We further assume that institutions or individuals deploying LeafAI may not necessarily have access to GPUs. To improve inference speed on CPUs, we intend to quantize our models, a process for converting 32-bit floating point values within model weights and biases to 8-bit integers. Quantization has been shown to dramatically reduce model storage size and memory usage and improve inference speed while typically showing limited decreases in performance [152].

## 8.4 Future Work

In future work we will employ a usability and user satisfaction and acceptability study. We hypothesize that the innovative design of the web application user interface will represent a

meaningful advance in user interface design for cohort discovery.

Possible future versions of the application may allow for general question-answering, data visualization, and data analysis as well.

### **8.5 Conclusion**

This chapter described the ongoing development of the LeafAI web application, which incorporates a number of innovative user-facing features such as streaming interface updates during query execution, real-time named entity recognition and normalization, as well as mixed-modal interaction allowing users to manipulate system responses and re-execute generated queries. In future work we will evaluate the web application in a usability and user satisfaction and acceptability study.

## Chapter 9

# CONCLUSIONS

This dissertation describes the development of a novel database query generation system to identify patients eligible for clinical trials using a natural language interface. We conclude with a summary of the primary contributions of this work, a discussion of limitations, and finally remarks on potential directions for future research.

### **9.1 Contributions**

Recruitment of eligible patients for clinical trials remains a costly and time-consuming challenge, and contributes in many cases to slowing the process of bringing new drugs and treatment options to patients who may benefit from them. The process of identifying patients who may qualify for clinical trial is also challenging, as manual chart review is highly labor intensive, interactive tools such as Leaf are not always able to represent appropriate queries to find patients in EHRs, and the technical expertise needed to run database queries to find patients directly from an EHR database is often a limited resource. Applications leveraging natural language descriptions of eligibility criteria to automatically sift through data and find patients therefore hold great potential to simplify and expedite this process.

It is our hope that the research and software applications described in this dissertation are impactful for real-world clinical trials. We summarize our primary contributions here:

1. *Annotated Corpora*: we introduced two new human-annotated corpora, the Leaf Clinical Trials (LCT) corpus and Leaf Logical Forms (LLF) corpus. The LCT corpus was developed using a highly granular named entity- and relation-based schema to enable robust query generation for real-world clinical trials and eligibility criteria. The corpus consists of over 1,000 documents of eligibility criteria from clinical trials. Our best

performing NER prediction baseline model trained on the corpus achieved an  $F_1$  score of 81.3%, while our best performing relation extraction model achieved an  $F_1$  score of 85.2%. The LLF corpus builds upon the LCT corpus by representing 2,000 eligibility criteria lines with corresponding logical intermediate representations, or logical forms. We developed the LLF corpus annotation schema with the aim of creating a data model-agnostic, flexible, robust, and machine-readable representation of the underlying logic of a given criteria. Our best performing model for predicting a logical form from a free-text input achieved a BLEU score of 93.5%.

2. *Model-agnostic query generation*: academic medical centers and research institutions use a variety of clinical database systems and schema. While research-oriented common data models such as OMOP are increasingly used to enable data and tool sharing and cross-institutional analysis, the research database landscape remains heterogeneous. Moreover, many institutions and projects alter common data models in order to accommodate specific project and analysis needs. Because of this, we developed a novel data model-agnostic query generation method enabled by "tagging" clinical database schema metadata using a UMLS concepts in what we call a Semantic Metadata Mapping, or SMM. Our SMM system enables great flexibility in representing a variety of data models, including relatively complex common data models such as OMOP.
3. *Knowledge Base*: a natural language interface for identifying cohorts using eligibility criteria requires an external knowledge representation system because humans often use non-specific phrasing and references for brevity and convenience. We developed a graph-based Knowledge Baes, or KB, by integrating a wide variety of publicly available data sets, ontologies, and vocabulary mapping systems, with the Unified Medical Language System, or UMLS, at its core.
4. *Reasoning upon non-specific criteria*: Indirect criterion which frequently appear in eligibility criteria, such as contraindications to a drug, symptoms of a condition, or

risks of an outcome, must be reasoned upon in order to determine patients meeting a given criterion. Leveraging our KB and logical forms, we demonstrate a novel inside-to-outside sequential reasoning method which allows our system to generate queries meeting non-specific criteria.

5. *Evaluation against a human programmer and actual trial enrollments:* of the natural language interfaces for eligibility criteria developed to date, to our knowledge only one used actual participant enrollments as part of a benchmark, and none compared system performance to that of an experienced human clinical database programmer. In contrast, we evaluated our system’s generated queries to that of a human programmer, setting a new and higher benchmark for this task. We demonstrate that our system is able to rival—and by some measures surpass—a human programmer in matching patients found by generated queries to those who actually enrolled, with our system identifying 212 of 427 (49%) total enrolled patients across 8 clinical trials compared to 180 (42%) found by queries of the human programmer.

## **9.2 Limitations**

The research described in this dissertation has a number of limitations, many of which have been described in preceding chapters. The corpora we introduce, the LCT and LLF corpora, were analyzed in great detail to ensure quality, but were nevertheless annotated by non-clinicians, and in any case, likely any human-annotated corpora many contain errors. Models trained on these corpora may also return erroneous output and thus inaccurate queries downstream. Our KB, while containing a wide variety of sources and thus capable of enabling reasoning across many demonstrated use cases, nevertheless lacks functionality to embed probabilities, significance, or caveats to its output, and thus may return erroneous results. We believe these shortcomings are acceptable given that users of our web application may edit any reasoned concepts.

In our query generation task, it is not clear how well (or poorly) our system may generalize

to other kinds of clinical trials outside of the 8 we tested on. Perhaps most importantly, our evaluation underscores the limits of measures system performance of tools such as LeafAI when there exists no true gold standard indicating patients across an entire medical system who would be eligible for a clinical trial. We use known participant enrollments in actual clinical trials but recognize the shortcomings of this approach.

For the web application, as discussed, we have not yet evaluated whether our system meets or exceeds usability and user acceptance standards, nor how well our system compares to previous user-facing systems.

### **9.3 Future Work**

There are a number of promising avenues for future research which may build upon our findings and systems. We first look forward to testing and improving our web application and query generation methods with robust user testing. Like other self-service tools, it is in the hands of users that the impact of such systems will be made.

Another exciting possible direction is the adaption of our system for general-purpose question-answering or dynamic data visualization. After identifying a cohort of interest, it is natural for inquisitive users to wonder, Of these patients, what were their outcomes post-surgery?, or, Did any show signs of  $X$  in the following year?, or, Show a comparison of their demographics alongside a control cohort. One can imagine such capabilities being of great potential value in expanding the analytic capabilities of tools such as ours.

One additional potential future direction is the creation of a de-identified data set of structured clinical data (and possibly unstructured notes as well) of patients alongside human expert-annotated indications of whether they were eligible or not for a given set of clinical trials. While the manual review required to determine whether patients were eligible or not would not be trivial, the practical value to the research community in improving tools such as our own and objectively comparing approaches would be significant.

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