Desiderata for the development of next-generation
electronic health record phenotype libraries

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Abstract

Background: High-quality phenotype definitions are desirable to enable the extraction of patient cohorts from large electronic health record repositories and are characterized by properties such as portability, reproducibility, and validity. Phenotype libraries, where definitions are stored, have the potential to contribute significantly to the quality of the definitions they host. In this work, we present a set of desiderata for the design of a next-generation phenotype library that is able to ensure the quality of hosted definitions by combining the functionality currently offered by disparate tooling.

Methods: A group of researchers examined work to date on phenotype models, implementation, and validation, as well as contemporary phenotype libraries developed as a part of their own phenomics communities. Existing phenotype frameworks were also examined. This work was translated and refined by all the authors into a set of best practices.

Results: We present 14 library desiderata that promote high-quality phenotype definitions, in the areas of modelling, logging, validation, and sharing and warehousing. Conclusions: There are a number of choices to be made when constructing phenotype libraries. Our considerations distil the best practices in the field and include pointers towards their further development to support portable, reproducible, and clinically valid phenotype design. The provision of high-quality phenotype definitions enables electronic health record data to be more effectively used in medical domains.

Keywords: electronic health records; EHR-based phenotyping; computable phenotype; phenotype library

Introduction

As a result of the digitization of health systems worldwide, electronic health record (EHR) data repositories have emerged as the main source of data for medical cohort research studies. To extract these cohorts, there is an increasing reliance on EHR-based phenotype definitions (also referred to as phenotyping al-
gories), which identify individuals who exhibit certain phenotypic traits, such as the same diseases, characteristics, or set of comorbidities. These definitions can be represented in many forms, including narrative descriptions, pseudo-code, or, in some cases, may already be directly executable. Conceptually, they may vary from simple code lists, via rule-based algorithms, to more involved machine learning (ML) tasks and high-throughput approaches using natural language processing (NLP).

While traditional big data techniques can successfully address the scale of the EHR data available, the effectiveness of phenotype definitions is affected by a range of other syntactic and semantic issues, including variations in the way data are structured and the coding systems used.

To overcome these issues and enable effective cohort extraction, a phenotype definition must exhibit certain properties. It must be reproducible, allowing for accurate (re)implementation, irrespective of the idiosyncrasies of the dataset against which the definition was originally developed; portable, allowing for straightforward implementation, irrespective of the structure of the target dataset; and valid, effectively capturing the disease or condition modelled. A definition that exhibits all of these properties we refer to as “high quality”.

To ensure high-quality phenotype definitions, support should be provided to the authoring, implementation, validation, and dissemination processes of a phenotype’s lifecycle. While such support is currently available, it is often sporadic and inconsistent because it is delivered via a wide range of different tools. Instead, building on the work of Richesson et al. [1], we propose that the functionality provided by these tools should instead be provided centrally, through the phenotype libraries where definitions are hosted. For example, libraries should enable phenotypes to be developed according to some set of standard models, and track the evolution of definitions under these models, so as to ensure that hosted definitions are clearer to understand and thus have the potential to be more reproducible. Moreover, libraries should assist in the derivation of directly computable phenotype definitions, through the provision of implementation tooling, to improve portability by enabling the execution of phenotypes in local use cases. Similarly, libraries should directly validate the definitions they host, through, for example, automated comparisons with gold standards.

To this end, in this work we contribute a number of desiderata for the development of phenotype libraries, which not only ensure that definitions are accessible but also maximize the quality of the phenotypes they contain by supporting all parts of the definition lifecycle. These desiderata are based on both the lessons learned during the development of contemporary libraries within the authors’ own phenomics communities, as well as a review of the functionality currently offered by phenotype tooling, which represent practices that have led to the development of high-quality phenotype definitions. By providing access to high-quality definitions, phenotype libraries enable both efficient and accurate use of EHR data for activities such as medical research, decision support, and clinical trial recruitment.

Background

Human phenomics is the study of human phenotypes and includes the science and practice of defining observable medical phenomena that indicate phenotypes to advance research and personalized care. The concept of a phenotype originated as a complement to the genotype, and the phenotype was defined as the complete set of an individual’s inheritable characteristics. Rather than describing someone’s genetic information, a phenotype captures all the observable properties (phenotypes) that result from the interaction of genetic make-up and environmental factors, including their demographic information, such as height or eye color, and medical histories.

With the emergence of large-scale EHR data repositories, the term “phenotype” has evolved to denote traits shared by groups of patients, such as a disease or condition that a cohort, or set of individuals, has. This may also include other complex combinations of traits, exposures, or outcomes, including comorbidities, polypharmacy, and demographic data. Defining these phenotypes, and validating them to ensure their accuracy and generalizability, is a process known as “phenotyping”, with “EHR-based phenotyping” relying primarily on data in the EHR. “Computational phenotyping” (also known as “deep phenotyping”) uses either supervised ML techniques to discover new members of a priori defined cohort or unsupervised techniques to discover entirely new phenotypes and investigate their properties.

EHR data repositories bring with them a very specific set of data challenges in terms of managing syntactic and semantic complexity, which act as a barrier to studies that need to use patient information from across multiple data sources and for the needs of different studies. For example, by the nature of healthcare delivery and how EHRs are used to document, a patient who has received a diagnosis of diabetes mellitus may be represented slightly differently in two EHR systems and will almost certainly be represented differently in EHRs for different countries.

Phenotype libraries—where definitions can be uploaded, stored, indexed, retrieved, and downloaded by users—provide a logical place in which to ensure that definitions are of a suitable quality to overcome many of the issues associated with extracting cohorts from complex EHR datasets. This is accentuated by the fact that the development of phenotype libraries is a rapidly growing area. Of particular note is the Observational Health Data Sciences and Informatics (OHDSI) Gold Standard Phenotype Library, which aims to support OHDSI community members in finding, evaluating, and utilizing cohort definitions that are validated by the research community. An initial version of the library is currently available, alongside a wider set of requirements to guide its future development [2]. Other libraries planned for development include the VAPheLib [3], which aims to collect, store, and make available 1,000 curated phenotype definitions for the clinical operations research community by the end of 2021. Phenotype libraries are also being developed as a part of wider phenotype frameworks. Alongside Richesson’s reusable phenotype definition framework sit initiatives such as the Phenotyping Pipeline (PheP), which aims to extract, structure, and normalize phenotypes from EHR data collected across participating sites [4].

Methods

To determine the functionality that should be provided by a next-generation phenotype library, a team of international researchers—comprising Health Data Research UK (HDR UK) Phenomics theme members and US researchers from the Mobilizing Computable Biomedical Knowledge (MCKB) and Phenotype Execution and Modelling Architecture (PhEMA) communities—first examined a range of tools supporting different parts of the definition lifecycle, which were developed within their respective phenomics communities. This was en-
riched with a wider review of the literature via Web of Science (WoS) [5] and the grey literature via Google to identify third-party projects that have developed phenotype tooling, or are planning its development, and future trends. Our decision to include the grey literature was informed by our a priori knowledge of tools under development that do not yet have published peer-reviewed articles. The tools reviewed included those that support authoring (e.g., modelling using the Quality Data Model [QDM] logic [6], the Clinical Quality Language [CQL] [7], and use of the Observational Medical Outcomes Partnership [OMOP] Common Data Model [CDM] [8] and associated tooling such as OHDSI’s Automated PHenotype Routine for Observational Definition (APHRODITE) [9]), implementation (e.g., definition translators [10]), and validation (e.g., electronic phenotyping validation [11]). Common functionality provided by the tools identified—representing opportunities for new phenotype library functionality—was extracted and summarized.

In addition, the authors examined existing libraries from within their own communities—including the Phenotype Knowledge Base (PheKB) [12], CALIBER [13], Phenoflow [14], and the Concept Library [15]—to identify instances of functionality currently supporting the phenotype definition lifecycle. Common functionality provided by these libraries—which has been shown to result in reproducible, portable, and valid phenotype definitions, and thus represent best practice—was also extracted and summarized.

Both of these summaries were translated to a draft set of desiderata via discussion amongst a subset of the authors (M.C., S.M., E.J., S.D., V.C.). All authors participated in an asynchronous iterative review process to critique, consolidate, refine, and define the final set of desiderata. The desiderata were further classified into logical categories.

**Desiderata**

In total, the authors arrived at a finalized collection of 14 desiderata, which are organized across the following sections into 5 categories: modelling, logging, implementation, validation, and sharing and warehousing. Figure 1 shows how the desiderata presented promote the design of a phenotype library that supports all parts of the phenotype definition lifecycle.

**Modelling**

Phenotype models govern the structure and syntax of phenotype definitions. For example, phenotype definitions are traditionally rule-based, meaning that they are composed of individual logical statements that each evaluate to a Boolean value, typically by relating data elements (with associated values)—such as the presence of a particular set of ICD-10 codes or a particular laboratory test result—to each other. The set of operators available to an author when connecting data elements (e.g., logical connectives such as conjunction and disjunction) would be established within a phenotype definition model. A model may dictate that a phenotype be represented in an unstructured, semi-structured, structured, or executable manner [16]. A summary of different phenotype definition formats, governed by phenotype models, is given in Table 1.

Implementing a phenotype definition involves translating the abstract definition (if unstructured or semi-structured) into an executable form that can be directly run against a patient dataset to derive the cohort exhibiting the defined phenotype. Typically this requires the logic of the definition to be realized in a programming language, such as translating abstract conditional clauses into a set of tangible Python conditional statements. We refer to these implementations as “computable phenotypes”. For a definition to be reproducible, it must be realized in a formal structure that can be accurately interpreted and implemented. Given the potential for human error in translating from an unstructured narrative to something computable, formal phenotype models provide such a structure.

Phenotype models are also key in ensuring semantic interoperability between definitions. That is, while the development of phenotype definitions can involve deriving a curated, canonical set of phenotype definitions containing “definitive” versions for each disease or condition being modelled for a particular domain (e.g., a national stroke body may want to maintain their set of stroke phenotyping algorithms), more often than not, it is perfectly valid to have overlapping phenotype definitions for different uses. For example, an eligibility criterion for a clinical trial may differ from a rule that triggers a decision support tool in an EHR system, and both would differ from a definition used in a population health study, even if all three nominally refer to the same disease [17]. Internationally, definitions for the same disease may also differ [18]. While this overlap is permissible, different definitions for the same condition must still be compatible, enabling, for example, their relative functionality to be compared. The adoption of a phenotype model enables such compatibility.

Given these benefits, a phenotype library should adopt a formal phenotype model to control the structure of hosted definitions. To ensure the use of such a model, a library can offer a graphical authoring environment—in the same way that tools such as the Phenotype Execution and Modelling architecture (PhEMA) Authoring Tool (PhAT) do [6]—through which new definitions can be authored. Similarly, existing definitions can be automatically checked for their adherence to the chosen model when uploaded.

Desiderata relating to the adoption of a phenotype model by a library are listed in the following sections. We view these desiderata as complementary to the well-established desiderata for phenotype definition model development put forward by Mo et al. [19].

**Support modelling languages**

The phenotype definition model adopted by a library should be supported by a (non-executable) high-level modelling language that dictates the syntax available to an author when defining the logic of a phenotype. A computable form of the definition can then be realized for execution in a local use case. When selecting or developing a definition model, the temptation may be to select a lower-level, executable programming language, in an attempt to expedite local implementation. For example, one could argue that a language such as Python is sufficient for simultaneously defining phenotypes and realizing them computationally. However, we would argue that using such a language as a means to express the logic of a definition ties the definition to general purpose, low-level language constructs, reducing clarity and thus reproducibility. This conclusion is supported by work such as that of Papez et al. [20], which found openEHR an overly restrictive standard when attempting to express phenotype definitions in a form that can be directly executed. An example of a phenotype definition realized in an executable language (Python) is given in Fig. 2.

In contrast, the syntax of higher-level modelling languages, while still precise, is often clearer, as well as often being domain specific. For example PhEMA’s PhAT allows users to define phe-
Figure 1: The stages of the phenotype definition lifecycle supported by a next-generation phenotype library.

**Table 1: Phenotype definition formats**

| Format                  | Description                                                                 | Example                                                                 | Category          |
|-------------------------|----------------------------------------------------------------------------|------------------------------------------------------------------------|-------------------|
| Code list               | A set of codes that must exist in a patient's health record in order to include them within a phenotype cohort | COVID-19 ICD-10 code "U07.1"                                            | Rule-based        |
| Simple data elements    | Formalizing the relationship between code-based data elements using logical connectives | COVID-19 ICD-10 code "U07.1" AND ICD-11 code "RA01.0"                   | Rule-based        |
| Complex data elements   | Formalizing the relationship between complex data elements, such as those derived via NLP | Patient's blood pressure reading > 140 OR patient notes contain "high BP" | Rule-based        |
| Temporal                | Prefix rules with temporal qualifiers                                      | Albumin levels increased by 25% over 6 hours, high blood pressure reading has to occur during hospitalization | Rule-based        |
| Trained classifier      | Use rule-based definitions as the basis for constructing a classifier for future (or additional) cohorts | A k-fold cross-validated classifier capable of identifying patients with COVID-19 | Probabilistic     |

![Figure 2: Python (executable) vs CQL (modelling) [21] representation of pharyngitis phenotype.](https://academic.oup.com/gigascience/article-10/9/giab059/6368631 by UCL, London user on 29 September 2021)

It is also important to note that the use of a modelling language as the basis for a phenotype model does not preclude the utility or use of higher-level, (more) human-readable representations such as flow charts. In fact, modelling languages typically connect well with such representations. For example, flow charts can be directly generated from Phenoflow’s workflow model. QDM is linked to a graphical HTML layer, and OHDSI cohorts can be viewed graphically using the ATLAS cohort editor.

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1. ```python
   valueset={} 
   2. valueset ['Acute Pharyngitis'] = '2.16.840...1011' 
   3. valueset ['Acute Tonsillitis'] = '2.16.840...1012' 
   4. 
   5. def Pharyngitis(): 
   6.   conditionA = valueset ['Acute Pharyngitis'] 
   7.   conditionB = valueset ['Acute Tonsillitis'] 
   8.   return conditionA + ' ' + conditionB; 
   9. 
   10. valueset ['Acute Pharyngitis'] = '2.16.840...1011' 
   11. valueset ['Acute Tonsillitis'] = '2.16.840...1012' 
   12. 
   13. define Pharyngitis: 
   14.   [Condition: 'Acute Pharyngitis'] union 
   15.   [Condition: 'Acute Tonsillitis']

Figure 2: Python (executable) vs CQL (modelling) [21] representation of pharyngitis phenotype.
Support Natural Language Processing–based and machine learning–based definitions

The modelling language selected to form the basis of a phenotype definition model should also support the representation of a wider range of definition types (Table 1). That is, under a definition model, one should be able to express not only standard rule-based definitions but also more complex definitions based on ML and NLP techniques. These techniques are becoming increasingly prevalent, particularly in those situations where the datasets against which the implemented definition is to be executed are of varying completeness or lack consistent record coding. The use of modelling languages to represent these types of definitions is also important for reproducibility because the use of an abstract representation reduces the potential for references to implemented libraries, commonly used by NLP and ML techniques.

Critically, in order to sufficiently represent both ML- and NLP-based phenotypes, a modelling language must be able to represent not only static information (as in rule-based phenotypes) but also complex processes. For example, in the case of ML, a definition may consist of a static, high-level specification of a trained patient classifier (via the provision of values such as feature coefficients) or may be a more complex description of the workflow used to train a classifier for a given condition, with a view to the classifier being re-implemented in new use cases or training a new model in new use cases, respectively. The workflow used to train a classifier may involve the identification of cases using the presence of certain keywords within an EHR [22] or, as in the case of the PheNorm framework, may involve additional steps, such as normalization (to factor in number of encounters when looking at the significance of a larger number of keywords or codes) and denoising (to look at the wider context of a keyword or code count, e.g., competing diagnoses) [23].

The high-level definition of ML-based, or probabilistic, phenotypes in this way is supported in the OHDSI’s Automated PHenotype Routine for Observational Definition, Identification, Training and Evaluation (APHRODITE) computable phenotype framework, which, although also linked to the OMOP CDM, offers a level of abstraction at which trained classifiers can be represented and ported between sites, or a defined workflow can be used to construct site-specific classifiers, when used in an executable form [24].

In the case of NLP, a definition may consist of a simple list of keywords relating to a given medical concept, or a set of regular expressions (not tied to any specific programming language), with a view to these being used as the basis for identifying conditions from free-text in a medical record, when realized in a computable form. However, like ML, NLP-based phenotype definitions are also often associated with complex processes, especially when used to conduct high-throughput phenotyping. For example, a PheMap phenotype definition consists of a set of linked concepts, the presence of which in a patient’s EHR is used to determine the probability of the patient having the condition represented [25]. The association of a phenotype with different concepts is defined within the PheMap knowledge base, which is constructed on the basis of a process that uses a specific set of NLP tools to derive these associations on the basis of the content of various text-based resources. Therefore, it is also important to represent this process as a part of any definition, especially if the knowledge base needs to be reconstructed within different domains. In instances such as these, it may be important for the definition model used to include guidance on the use of specific tools, but it must do so in a manner that retains clarity and generalizability, thus balancing reproducibility requirements. Modelling languages like CQL have the potential to link to external tooling, for the purposes of effectively capturing NLP and ML processes such as these.

Support multi-dimensional descriptions

A significant hurdle in porting a phenotype definition from one setting (institution or dataset) to another—a process we refer to as localization—is understanding its structure and semantics in order to derive a local computable form or modify an existing one. Complex rules and the use of idiomatic clinical terminology, although often necessary components of a definition, are both barriers to this understanding and thus to reproducibility. To address this issue, a phenotype definition model should allow an author to express the same logic of a phenotype at different levels of technical complexity. This approach aims to communicate supplementary information alongside the provision of the core definition logic. For example, the workflow-based PheFlow model allows an author to use the technical terminology and rules required to express a phenotype definition but then also requires an author to provide longer definitions of this functionality to improve clarity, and to also classify each unit of functionality under a given ontology, enabling a high-level understanding of the functionality to always be accessible. In other modelling languages like CQL, such information can be communicated using constructs such as inline comments.

Logging

The development of a phenotype definition is an incremental process. Capturing and communicating this process is key in ensuring that a definition can be accurately interpreted and is thus reproducible. Moreover, this information strengthens the trustworthiness of a phenotype and thus its potential applications. Therefore, phenotype libraries should provide a mechanism for logging the evolution of a phenotype definition.

Support versioning and data provenance

One way in which a phenotype can evolve is through a series of iterative refinements. SAIL Databank’s Concept Library stores phenotypes as sets of codes, with a view to making these phenotypes available in different studies and use cases [15]. The concept library, as the name suggests, focuses on a model under which phenotypes are collections of grouped medical “concepts” or “working sets”. The Concept Library records and communicates the evolution of a phenotype definition using methods akin to standard version control, logging the state of a phenotype after each revision, and thus provides an overview of the definition’s progression. This versioning process often relies on attributing a universally unique identifier (UUID) to each definition and each subsequent revision of that definition. Such an identifier might simply be incremental or convey some details of the phenotype itself. It should also be independent of other identifiers to maximize clarity [26]. For example, within APHRODITE a UUID is derived by committing (each version of) a generated definition to a GitHub repository and extracting the unique commit hash value, in accordance with the FAIR (Findable, Accessible, Interoperable and Reusable) principles [27].

A more comprehensive way to capture the evolution of a definition—and thus contribute to its reproducibility—is to deploy formal data provenance capture tools to capture richer, real-time information about the evolution of an entity. This might include information about updates to the structure of a definition or details of how that definition was validated. It might also include information about how the definition was de-
Next-generation electronic health record phenotype libraries

expression after exhibiting bipolar disorder) [32]. Establishing this relationship further contributes to the provenance of a phenotype, the precision of its definition, and, consequently, its reproducibility.

Conversely, “sub-phenotypes” may be computationally derived from existing phenotypes by clustering those features (e.g., demographic, diagnosis, medication) identified, by a trained classifier, to be key attributes of those patients exhibiting the parent phenotype [33]. Such a relationship should also be logged by a phenotype library, to establish the evolution of a definition, and track changes and dependencies across phenotype definitions.

Implementation

Our initial desiderata determined that phenotype definitions should not themselves be executable. While important for reproducibility, this raises natural issues around the complexity of realizing a phenotype defined using a modelling language computationally for individual use cases, something that hinders portability. This issue can be addressed by meeting several requirements, which are explored in the following sections.

Communicate implementation information in the model

One way in which implementation can be supported is through the definition itself, by communicating information pertinent to its computable realization. To do this, one might select a phenotype definition model based on a modelling language that allows an author to express additional information at different levels of abstraction. For example, the Phenoflow model frames the traditional (rule-based) logic of a phenotype definition as an “abstract” layer and allows an author to complement this layer with additional layers, each of which gradually communicates more implementation information: a “functional” layer, introducing the concept of data types, and a “computational” layer, expressing details such as target execution environments. The fact that these layers sit alongside the traditional, abstract logic layer allows for more concrete implementation to be expressed without affecting portability.

The abstract layer of the Phenoflow model is split into individual modules, each of which represents a distinct unit of functionality, and which collectively define the process required for deriving a patient cohort from a set of health records. Each module in the abstract layer has an equivalent module in both the functional and computational layers, ensuring a correspondence between each level of representation within the model. However, these modules also provide another means by which implementation information can be communicated through a definition model, in that they provide a clear template for development; each module represents a single unit of functionality that must be implemented by a developer when realizing the computable form. This reduces the implementation burden on developers and thus improves portability. Modelling languages like CQL, which support the definition of individual functions as a part of an abstract layer, offer similar benefits.

Support tooling that provides multiple programming language implementations

Phenotype implementation tooling automatically takes an abstract phenotype definition and translates it into a computable form. This naturally improves portability. Examples of this tooling include the “translators” developed by the PhEMA initiative, which are able to take a modelling language definition of a phenotype—such as definitions expressed in QDM, as produced by the PhAT, or in CQL—and transform them into executable for-
mats (e.g., pipelines [10]). In addition, the OHDSI tools provide ways to take their domain-specific representation and translate it to SQL queries that execute against multiple database systems adopting the OMOP CDM. Although all definitions are SQL, the different dialects used by database vendors are akin to separate programming languages.

Given these benefits, a phenotype library should provide access to implementation tooling. In the simplest form, access should be provided to this tooling by hosting and indexing it in a library, in the same way that the definitions themselves are hosted and indexed. This tooling can then be downloaded, along with a definition, and executed locally to produce a computable form. More advanced integrations will provide the functionality offered by implementation tooling directly through the library, by running it as a service that can be accessed by users via the library in order to download the automatically generated computable form of a phenotype. This is the approach taken by the Phenoflow platform, which allows users to obtain computable copies of a phenotype definition directly, by running a microservice generation architecture.

The tooling indexed should be able to support implementations in a variety of different programming languages. While the programming language used might seem to be of little consequence, in practice, even with the presence of a translator, the researcher generating a computable form for a new use case is likely to still have to modify (localize) that computable phenotype for local use. Such modifications might include optimizations to the structure of the implementation to allow the computable form to operate in low-memory environments or to operate as a part of existing infrastructure (e.g., a clinical trial platform [34]). In this instance, having that definition in a language that the researcher is comfortable with editing is important. For example, the pipeline-based implementation originally produced by the PhEMA translator only supports the KNIME format. As such, a researcher has to be comfortable with this format to make edits. To maximize portability, phenotype libraries should aim to support implementation tooling capable of producing executable definitions in multiple languages. An example of this is seen within the Phenoflow platform, where one can generate a workflow that uses modules from a variety of languages, including Python and Javascript, with containerized environments supporting the straightforward execution of these units locally.

**Support tooling that provides connectivity with multiple data standards**

When a phenotype definition is translated by a piece of tooling into an executable form, it is typical for that definition to be tied to a given data source format, from which the resulting cohort is identified. In certain cases, that data format is always the same. For example, OMOP cohort definitions, when translated into a computable form (SQL), are always tailored for the OMOP CDM. While beneficial in the sense that this provides an automated translation process that works across sites, those sites must all adopt the OMOP CDM, which is not always feasible. Instead, in reality, sites may use a variety of implementation formats, such as i2b2 and FHIR. For these reasons, phenotype libraries should index implementation tooling that not only supports multiple language implementations but also supports the realization of definitions for different data formats. Naturally, the more data source formats supported, the more portable the definitions stored within a library are. For example, the computable forms generated by PhEMA’s translators can be tailored for a variety of local data formats, including FHIR and the OMOP CDM itself. Similarly, in the Phenoflow library, interacting with a data source is considered to be the first step in a phenotype’s definition, and as such different “connectors” are available when generating the computable form of a definition. These connectors support a variety of different standards such as OMOP and i2b2, and plans are in place to support dataset-specific standards, such as the standard used by UK Biobank (via tooling such as Funpack [35]).

The connector approach also provides a natural point at which to conduct any necessary (automatic) translation between the coding system adopted by a target data source and the coding system expected by the implemented definition. For example, if the target datasource adopts Read codes but the computable phenotype relies on sets of ICD codes, a connector might not only ingest data but also perform code mappings accordingly.

Despite these benefits, the requirement to produce a new translator, or new connector, for each new data source format is a natural drawback to each of these approaches. However, the advantages over manual translation are still clear.

**Validation**

Validating a phenotype definition involves confirming its accuracy. To do this, the cohort identified by a computable phenotype is typically compared to a reference standard, such as the cohort identified by manual review of medical records from the same patient population (a “gold standard”). The extent to which the two cohorts overlap determines the validity of the definition. While reference standards are a common means of phenotype validation, other techniques exist and are listed in Table 2. Phenotype definitions that are shown to be accurate are considered to be of a higher quality. Therefore, phenotype libraries should facilitate the validation process.

**Support a defined validation process**

To support the validation of stored definitions, a phenotype library should have a clear and scalable process for the submission of existing validation information by a user, across a variety of the mechanisms listed in Table 2. This information can then be stored and presented alongside each definition. For example, the CALIBER library stores phenotypes as code sets (342, at the time of writing), with a view to providing a framework for the definition of consistent phenotypes, which can then be reused by care service providers for nationwide EHR-based observational research [13]. Each definition in CALIBER appears alongside algorithmic information about the relationship between the code sets and key validation information. Specifically, the CALIBER library offers up to 6 different techniques, which are used to validate a single definition. Similarly, the OHDSI gold standard phenotype library is so-called because there is a well-defined process proposed for the submission of phenotypes based on different user roles. Specifically, the submission of a computable phenotype definition to the library (which can occur using the APHRDITE framework) will require definitions to be submitted by those in the “author” role, vetted by “librarians”, validated by users who act as “validators”, and used by standard “users” [36].

**Automate multiple validation techniques**

When new definitions are submitted without validation information to a library, it should seek to automatically validate these definitions by comparing them, or their outputs, against assets that are hosted alongside the definitions, such as gold standard datasets. For example, in [11], Kukhareva et al. present “electronic phenotyping validation”, a framework for the automated...
Next-generation electronic health record phenotype libraries

Table 2: Phenotype validation mechanisms

| Mechanism               | Description                                                                 | Example                                                                                           |
|------------------------|-----------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|
| Disease registries     | Compare the phenotype cohort with those present in the registry             | Comparison of a diabetes phenotype cohort with those patients present in a diabetes registry (e.g., T1D exchange) |
| Chart review           | Compare the phenotype cohort with the patients identified by manual review of medical records | Comparison with a diabetes gold standard, produced by double manual review of patient medical records |
| Cross-EHR concordance  | Compare percentage of cases identified by a phenotype across different sources, and identify any overlap | Comparison of the percentage of patients identified by a diabetes phenotype in primary and secondary care EHRs, and the identification of any case overlap |
| Risk factors           | Compare the magnitude of the phenotype cohort with standard risk calculations | Comparison with a survival analysis                                                               |
| Prognosis              | Compare the magnitude of the phenotype cohort with external prognosis models | A patient is more likely to be a valid member of a diabetes cohort if they have the HLA-DR3 gene |
| Genetic associations    | Compare whether the presence of a patient in a phenotype cohort is consistent with their genetic profile |                                                                                                |

A comparison of a definition with the results of a manual review of medical records. In the absence of such assets, a portal might host tooling designed to derive these assets automatically. For example, PheValuator trains a linear model based upon cases and controls identified using some of the techniques already discussed, such as the presence (or absence) of a large number of clinical codes relating to a certain condition within a patient’s record [37]. This model is used, in turn, to construct an evaluation cohort, which matches each individual in the cohort to a probability value indicating the likelihood of them having the condition of interest. This cohort can then act as a silver standard against which phenotype definitions can be validated, in this case by using the matched probabilities to construct totals from which sensitivity, specificity, and positive predictive value are calculated.

There is also an argument for the automated combination of different validation approaches to avoid the shortcomings of each individual approach. For example, using a disease registry approach alone as a gold standard for phenotypes related to that disease is not scalable or feasible for patient cohorts focusing on multi-morbidities and complex demographic criteria. Similarly, validating using clinical notes review, where phenotype patient matches are manually reviewed, is not sustainable for large learning health system infrastructures. While the manual text extraction of phenotypes can be effective in smaller scenarios, it is heavily dependent on the human expert and the sample being analysed and is not well suited to cross-site studies with differences in clinical and operational procedures and opinion between sites. As such, phenotype libraries should offer novel hybrid approaches to validation that encompass structured data, free text, and ancillary sources for both structured and unstructured data.

Enable feedback

To facilitate any (informal) user-based validation of stored definitions, a phenotype library should support social interactions between the authors and researchers that use it, with a view to providing authors with feedback and allowing them to address this feedback accordingly. Social functionality is supported by the Phenotype Knowledge Base (PheKB), which currently hosts ∼70 phenotype definitions [12]. For example, within the library, users can post comments or questions against different phenotype definitions. A researcher can also request collaboration on the development of phenotype definitions.

However, those users permitted to interact with a phenotype definition within a portal may be restricted. Within PheKB, only users with certain organizational affiliations (e.g., the eMERGE network or the Phenome-Wide Association Studies [PheWAS] community [38]) are provided with access by default, with other users required to request an account prior to providing feedback on definitions. Other portals may restrict access to different countries or regions.

In many cases, these restrictions are necessary during the development of a phenotype. For example, APHRODITE’s definition repositories are kept private while they are still under development. However, once developed, definitions can be accessed through the repository via any web browser or through an R Shiny app. Based on practices such as these, phenotype libraries should limit the restrictions they place on those who can engage with the definitions in phenotype libraries, once developed. By eliciting comments on the validity of hosted definitions from a wider audience, one is likely to gain a greater understanding as to the quality of a definition.

Sharing and warehousing

Once a phenotype definition is appropriately reproducible, portable, and validated, it should then be accessible for use by others. While the traditional and default role of a phenotype library is to provide such access, this can be optimized, as discussed in the following sections.

Expose a standard API

To maximize accessibility, a phenotype library should facilitate user interactions via multiple interfaces. The definitions in a library are usually available via a single interface: a graphical front end. While this provides a reasonable baseline for accessibility, it does not maximize it. For example, a user cannot instruct a piece of software to interact with the library, to include definitions directly within a piece of code, resulting in potential inconsistencies arising from manual entry. Similarly, existing software systems, such as decision-support systems, cannot autonomously access phenotypic information. Perhaps most importantly, a lack of programmatic accessibility means that one...
library cannot easily access the functionality of another in order to provide complementary functionality.

To address these issues, phenotype libraries should offer API-level web services that (at a minimum) duplicate the functionality available in a user interface. In doing so, several considerations should be addressed. First, the level of API access needs to be considered, including whether to provide access only to trusted partners, and thus provide suitable authentication mechanisms (e.g., OAuth), or whether to make the API publicly accessible. The selection of the type of API-level access provided to the functionality of the web resource should be subject to the policy of the organization developing the library. Second, the protocol used to facilitate communication with the API should be considered, such as Remote Procedure Call (RPC), Service Object Access Protocol (SOAP), and Representation State Transfer (REST). REST is a simple and widely adopted specification model and is thus the technology that is likely to be most attractive when constructing a library API. Next, to support programmatic access and enable definitions to be differentiated automatically, a formal identification system should be established for each definition. The most straightforward way to this is to leverage the UUID attributed to each phenotype version.

The functionality of the API itself also needs to be considered. In [1], Richesson et al. propose that an API service should be used to construct phenotype definitions for the purpose of defining inclusion and exclusion criteria for clinical research trials. Building on this outline, we consider several additional API-level use cases, including searching phenotype definitions, extracting a specific phenotype definition, submitting a new phenotype definition, submitting a new use case for an existing phenotype definition, or validating an existing definition and linking a phenotype definition with a data source, and vice versa. Examples of specific functionality that an API-level phenomics resource should support within each of these use cases are given in Table 3.

The benefits of API functionality are evident in the CALIBER, Phenoflow, and Concept Library libraries, all of which communicate together to collectively form, along with a dataset Gateway, the HDR UK phenomics resource. As shown in Fig. 4, each library operates as a service, and collectively these services are able to deliver the functionality of a single library to a user. The services at the core of this library are the Concept Library and the CALIBER library, each of which stores phenotype definitions. Using provided APIs, the Concept Library is able to import definitions from the CALIBER library, enabling phenotypes to be both formally stored and validated across both services, respectively. Similarly, the Phenoflow service—also capable of automatically importing and representing definitions using a workflow-based model, and generating a corresponding computable form for execution against a local dataset—is able to import definitions from both the Concept Library and CALIBER. Finally, the Gateway service provides access to a comprehensive collection of datasets, which are linked to by services such as CALIBER, when a given phenotype definition is present in one of the hosted datasets. Similarly, the Gateway links back to CALIBER when a phenotype is present in a dataset, to facilitate searches based upon these definitions.

**Offer advanced search capabilities**

The accessibility of existing phenotypes within a library relies on its search capabilities. Searches based on given name or identifier and version should enable simple use cases. For example, PheKB offers comprehensive search functionality, with users able to perform searches not only against the definitions themselves using given keywords but also against supporting definition content, such as articles, implementations, and datasets. Alternatively, the library has the option to list all phenotype definitions—including phenotype definitions under development, if the user is logged in—where a user can instead filter the definitions returned after the fact, based on properties such as the authoring institution.

While the search functionality offered by PheKB is helpful, more advanced search capabilities should be supported to facilitate both more complex cases and improved information retrieval. This includes searches based on specific codes, or groups of codes, or an approximate pattern matching, based on regular patterns or even text similarity. Synonyms (including abbreviations and acronyms) may also be used as a mechanism to improve search results over keyword searches. For example, a search for “diabetes” would likely fail to find a phenotype that refers to “T2DM” throughout, although “T2DM” is a recognized abbreviation that can be semantically linked via the UMLS.

Even more advanced capabilities might include searches using semantic similarity between a given set of concepts and the stored phenotypes supported by phenotype ontologies. This could enable the discovery of semantically identical or closely related concepts within the library. Similarly, similarity metrics between phenotype definitions, facilitated by the adoption of a formal phenotype model, are likely to assist in scalable searches across different repositories, whereby a partial match may indicate a usable cohort definition to investigate.

**Include comprehensive metadata**

The search and browse features described must be supported by appropriate metadata, which can be used to describe both the subject and format of phenotypes in ways that make them findable to users with specific research or clinical needs. Such definitions we might refer to as “FAIR Phenotypes” [41]. To achieve this, each phenotype definition should include structured data that describe the subject (i.e., clinical condition) and intent (screening, etc.) of the definition, as well as the source, date, publisher, and so forth, similar to the tagging of resources in traditional libraries. Additionally, each component of the phenotype model (e.g., underlying data model, data elements, value sets, code lists, coding language) must be specified with an assigned code or value so that users can search on these features or have them displayed when browsing a phenotype library or repository. Examples of existing libraries that look to attribute
Table 3: Suggested library API functions with all requests made in, and responses returned in, YAML+Markdown/JSON/XML formats

| Function                          | User access level | Description                                                                                                                                 |
|----------------------------------|-------------------|---------------------------------------------------------------------------------------------------------------------------------------------|
| Search                           | Simple search     | A free text search, examining the entire contents of the portal and returning a list of phenotypes that match the search criteria.          |
|                                  | Advanced search   | A free text search, examining specified sections of the portal (e.g., main content, just metadata) and returning a list of phenotypes that match the search criteria. |
| Phenotype extraction             | Extracting specific phenotype(s) | Given a phenotype ID supplied by a user (or generated by the platform), the API returns the phenotype definition. Only authorized users should be allowed to submit either a single or group of phenotype definitions. |
|                                  | Extracting all phenotypes | Return a full list of phenotypes. Each aspect of a phenotype definition—including constituent code lists, links to datasets where that phenotype appears, and other metadata—can be updated by passing a phenotype ID and the names of the fields to update and their new values. Each update should mark a version number to keep record of any updates over time. |
| Adding new phenotype(s)          | Updating the contents of a specific phenotype | An administrator of the portal can hide a phenotype definition by providing a phenotype ID. An administrator of the portal can delete a phenotype definition entirely by providing a phenotype ID. |
|                                  | Updating a complete phenotype with multiple features | Update a phenotype's contents by passing a phenotype ID and submitting an updated phenotype definition file to replace the previous version for public view. |
|                                  | Submission of a new validation case study for an existing phenotype | Adding a new use case to validate an existing phenotype (identified by a phenotype ID) by passing a file. |
| Deletion of a phenotype          | Removing a phenotype from public view (soft delete) | An administrator of the portal can hide a phenotype definition by providing a phenotype ID. |
|                                  | Removing a phenotype from the library (hard delete) | An administrator of the portal can delete a phenotype definition entirely by providing a phenotype ID. |

appropriate metadata to stored definitions include CALIBER and PheKB (Fig. 5).

In addition to supporting search, the use of metadata is important for a number of other reasons. First, metadata can make clear characteristics of phenotypes related to their accessibility, interoperability, and reuse. To this end, as part of the Mobilizing Computable Biomedical Knowledge (MCBK) initiative, Alper et al. have proposed 12 categories of metadata that are required to fully represent knowledge objects, including phenotypes, for FAIR principled criteria [42]. In addition, metadata fields that describe the versioning aspects of a definition can be populated to further formalize the provenance of the phenotypes in a collection. Next, as the intent, development, and validation of phenotypes are essential for potential implementers to understand in order to trust the quality and appropriateness of a phenotype for a new purpose, representing aspects of the phenotype development and validation process formally is critical. To do this, the Trust and Policy Work Group of the Patient–Centered Clinical Decision Support Learning Network defined an extensive set of metadata for trust [43]. Finally, metadata can be used to formally represent many aspects of the implementation and tooling described, enabling potential implementers to search on these features, such as language, and possibly support automated translations.

While more and robust metadata are beneficial from a library perspective, populating these metadata accurately and consistently requires resources, and the extent and detail of metadata will depend upon a balance to adequately meet the needs at the expense that the library sponsor will bear. One potential solution to this issue is to automatically generate metadata, which is the approach taken in data management platforms [44]. Overall, time will show how the community of phenotype users can develop consensus on a minimum set of metadata, library, or indexing best practices to complement and formalize the desiderata described here, and also build a compelling value case for their use to support high-quality phenotyping across countries.

Conclusions

While making significant advances, computable phenotyping is still at an early stage where methods and repositories are emerging to meet the needs of a range of medical research domains, with little methodological consensus. As tooling gradually matures beyond the realm of early adopters to become usable for a broad spectrum of researchers and implementers, the focus needs to move away from one-size-fits-all “perfect” phenotype definitions to acknowledging the diversity of phenotype application areas, the resultant explosion in the numbers and varia-
tions of phenotypes to be stored—in particular the arrival of advanced probabilistic and NLP-based phenotypes to sit alongside traditional rule-based definitions—and the challenges of deploying them in the real world, especially in the presence of high-throughput requirements. Portability and reproducibility are essential in addressing this scaling-up, with techniques needed to move phenotype definitions between both data sources and different health settings.

Phenotype libraries offer a natural meeting point of these multiple use cases and domains to support high-quality phenotype definitions. In terms of designing phenotype libraries as technical entities that enable the storage and retrieval of definitions, there is a clear need to track the evolution of phenotype definitions as they are authored, support advanced search techniques that enable these definitions to be located by others, and establish a collaborative process through which the validity of definitions can be critiqued. All of this functionality should be accessible within a library via multiple channels, in particular comprehensive, standards-based API functionality to ensure interoperability. Authoring and storing phenotype definitions according to a standard model is another aspect through which phenotype libraries can contribute to definition reproducibility. The model adopted by a phenotype library should exist at the correct level of abstraction, prioritizing modelling languages over executable programming languages, and offset this, in terms of implementation, by incorporating key implementation information and improving clarity through multi-dimensional descriptions. Finally, a phenotype library should encourage the use of phenotype definitions in new use cases by supporting the validation process, both automatically and through the definition of a structured validation process.

The impact of supporting the development and implementation of high-quality phenotype definitions is significant, particularly because these definitions provide efficient access to accurate cohort data by overcoming many of the complexities associated with patient datasets. Cohort data support not only research studies (e.g., the identification of predictors for a certain condition) but also the provision of decision support (e.g., access to the medical histories of one or more individuals) and clinical trials (e.g., the establishment of trial cohorts). The use of computable phenotypes to determine cohorts from complex datasets for these purposes can be complemented by using traditional big data techniques to manage scale, by an increased focus on multi-morbidities—the complex interactions of diseases in patients—which are a crucial factor in personalized decision support systems, and by N-of-1 clinical trial design.

Overall, running through these desiderata is the awareness that cross-domain sharing of phenotype definitions can only occur through curated libraries that evolve in a controlled manner. Such libraries have to be (i) clinically and scientifically valid, (ii) technically realizable, and (iii) usable by researchers in different domains. Through the use of our desiderata, we believe the current and future phenotype libraries will deliver on these three fronts.

Data Availability
Not applicable.

Abbreviations
APHRODITE: Automated PHenotype Routine for Observational Definition, Identification, Training and Evaluation; API: Application Programming Interface; CQL: Clinical Quality Language; EHR: Electronic Health Record; ICD-10: International Classification of Diseases, Tenth Revision; MCBK: Mobilizing Computable Biomedical Knowledge; ML: machine language; NLP: natural language processing; OHDSI: Observational Health Data Sciences and Informatics; PhAT: PhEMA Authoring Tool; PheKB: Phenotype Knowledge Base; PheP: Phenotyping pipeline; PhEMA: Phenotype Execution and Modelling Architecture; QDM: Quality
Data Model; UMLS: Unified Medical Language System; UUID: universally unique identifier.

Competing Interests
The authors declare that they have no competing interests.

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