RESEARCH

A sample article title

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Abstract

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Introduction

Data and Information modeling in the healthcare domain have witnessed significant improvements in the last decade owing to advances in the development of state-of-the-art information and communication technologies (ICT) and formalization of storage and messaging standards. Subsequently, the scope of Healthcare Management Information Systems (HMIS), medical ontologies, and Clinical Decision Support Systems (CDSS) has broadened, beyond the operational capabilities of traditional rule based systems. One of the major reasons behind this limitation is due to the numerous heterogeneities in healthcare at data, knowledge, and process level. Thus, healthcare interoperability which aims to provide a solution to this problem, can be compartmentalized into data interoperability, process interoperability, and knowledge interoperability. Data interoperability resolves the heterogeneity between data artificats, to enable, seamless and interpretable communcation among source and target organizations, while preserving the data's original intention during storage, communication, and usage (as defined by IEEE 610.12 [1], Health Level Seven International HL7, and Healthcare Information and Management Systems Society HIMSS [2]). On the other hand, process interoperability regulates the communication among organizational processes to provide compatability between process artifacts within and seamless transformations across different organizations[3]. Lastly, knowledge interoperability provides a sharing mechanism for reusing interpretable medical knowledge, acquired through expert intervention and other mechanisms, across decision support systems [4]. In more tangible terms, healthcare interoperability at data, process, and knowledge level can be exemplified within the healthcare constraints experienced due to the emergence of Covid 19. The operational capabilities of the current healthcare service delivery infrastructure has gone under tremendous stress due to Covid 19. World over, large primary healthcare units have managed to create separate units for managing patients, suffering from extreme cases of the novel coronavirus. For secondary and tertiary care units, government involvement has become necessary to filter coronavirus patients and adhering to a national pandemic response policy. These complex circumstances have enhanced the need for sharing patient data and state-of-theart medical knowledge in real-time, to provide the medical experts with a tool to Doe and Smith Page 2 of 9

make accurate and timely decisions. Data interoperability can enable the front line medical workers to fetch, understand, and use patient data, especially comorbidities, across organizational and physical boundaries, without suffering from societal taboos that may prevent the patient from sharing their complete and accurate medical histories. Knowledge interoperability can improve the knowledge acquisition and sharing protocols to provide the medical experts such as epidieomologists and vaccinologist, with latest information on affected population trends, disease diagnosis, treatment, and followup procedures, and interpretable decisions leading to positive or negative outcomes. Process interoperability can help reduce and in some cases remove the operational redundancies between health centers. In this way, successive healthcare treatments can take benefit from earlier diagnosis, treatment, and followup procedures, thereby reducing the stress on healthcare experts and systems. Standards such as Health Level Seven (HL7) Fast Healthcare Interoperability Resources (FHIR), and openEHR provide the foundations for storing and communicating medical data, through the use of well defined protocols. While systematized nomenclature of medicine—clinical terms (Snomed-CT) [5] and logical observation identifiers names and codes (LOINC) [6] provide a standard definition for clinical terminologies and laboratory tests, respectively. Similarly Medical Logic Module (MLM) provides a standardized way for expressing medical knowledge. However, the plethora of standards, necessitates the creation of bridging standards, that can resolve the heterogeniety between the medical standards. Substantial effort has gone into this endeavor with the Clinical Information Modeling Initiative (CIMI) [7] taking the lead in bridging the gap between HL7v3 and openEHR. Similarly, SNOMED CT and LOINC are working to resolve the redundancies between the two terminological standards since 2013. This healthcare interoperability solution follows a formal, albiet long process, which is greatly dependent on the human factor. However, the current healthcare scenario, requires a quick solution to create a scaffolding of an interoperable bridge between various healthcare providers. It is also important to ensure that this scaffolding should be able to support the formal standardization processes of the future. In [8] we have presented the Ubiquitous Health Platform (UHP), which provides semantic reconciliation-on-read based data curation for resolving data interoperability between various schema. This methodology is based on the creation and management of schema maps, that can provide the framework for transforming a source schema into a target schema. In the current manuscript, we will present our research work to build and manage the schema map knowledge base. Overall, our methodology has two major portions, firstly we apply a novel schema matching technique to create a transformation function σ for the participating legacy schema, and secondly, we have used the Ripple Down Rules (RDR) to manage our knwoledgebase, which will be presented in some detail, focusing primarily on the search and evolution services. In particular,

- Section 2 contains the details of our methodology, where we aim to present a reproducible theoretical framework.
- Section 3 provides the experimental setup
- Section 4 presents the results
- Section presents the related work
- Section 6 concludes the paper.

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Related Work

Althubait et al. [9] proposed an ontology expansion methodology that identifies and extracts new class from text articles using word embedding and machine learning techniques. The authors identified the similarity of tokens and phrases of the text articles with the exiting classes of the ontology. The target ontology is expanded with classes from text articles having greater similarity with that of already added classes. A similar word embedding technique was also used by Nozaki et al. [10], where the authors used instance based schema matching technique to identify the semantic similarity between two instances. The results of the study showed the possibility of detecting similar string attributes of different schemas. Yousfi et al. [11] also utilized semantic base techniques and proposed xMatcher XML schemas matching approach. xMatcher transforms schemas into a set of words, followed by measuring words context, and relatedness score using WordNet. The terms from different schemas having similarities greater or equal to 0.8 are considered similar. Bylygin et al. [12] devised an ontology and schema matching approach by combining lexical and semantic similarity with machine learning approaches. The authors used lexical and semantic measures as features and trained various machine learning algorithms including Naive Bayes, logistic regression, and gradient boosted tree. The result achieved showed that the combination of algorithms outperformed the single modal.

Martono et al. [13] provided overview of previous studies related to linguistic approaches used for schema matching. Linguistic methods focused on finding strings and evaluate there similarity in different schemas. The string are normally normalized before to align both the strings before similarity comparison. The normalized strings are categories based on the information relatedness and element with similar category are compared using various similarity measure including Jaro-distance, Lavenstein (edit-distance), and many more. Alwan et al. [14] have summarized the techniques used in the literature for schemas and instances based schema matching. The information used for schema matching is categories into schema information, instance and auxiliary information. Most of the searchers have used syntactic techniques (including n-gram, and regular expression), semantic techniques (including Latent Semantic Analysis, WordNet/Thesaurus and Google Similarity) for schema level and instance level matching to achieve the final goal of data/information interoperability. Kersloot et al. [15] performed a comprehensive systematic review to evaluate natural language processing (NLP) algorithms used for clinical text mapping onto ontological concepts. The findings of the studies were evaluated with respect to five categories; use of NLP algorithms, data used, validation and evaluation performed, result presentation, and generalization of results. The authors revealed that over one-fourth of the NLP algorithms used were not evaluated and have no validation. The systems that claimed generalization, were self evaluated and having no external validation.

Methodology

Healthcare interoperability, with a focus on non-standard compliant medical schema, is dependent on the generation and validation of schema maps, as discussed above. To this end, the creation of a cohesive workflow is of utmost importance.

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In our earlier work [8] we used maximum sequence identification and matching using suffix trees for syntactic matching of two distinctly sources data schemas. This was followed by semantic concept enrichment and subsequently concept matching, for creating rules in the form of schema maps. The simplified mapping functions, thereby created, provided a simple methodology for converting semi-structured medical data, into a non-persisted, interpretable, model form. In our current methodology we have utilized state-of-the-art machine learning techniques for converting medical schemas into semi-structured form, which is then used to create embedding vectors. These vectors are then used for modeling and creating interpretable rules for applying semantic reconciliation in the form of schema matching and/or transformation. These rules are managed by the RDR which allows fast inference using the relevant knowledge sub-tree, and can well manage knowledge evolution to scale out (by incorporating new schema) and up (by incorporating changes and additions to existing schema) In essence, the aim here is to define a uniqueness property (\mathbb{P}) , as shown in Equation (1), whereby the semantic and syntactic uniqueness of each attribute as a disjoint union, is used.

$$range(\mathbb{P}) = \tau_{syntatic} \uplus \tau_{semantic} \tag{1}$$

The syntactic uniqueness as represented in Equation (2), is also a disjoint union of the measured attribute type, corresponding to one of integer, float, boolean, or string, and it valid values. The elements of the attribute type set have been determined based on the common primitive types available to the applications at an abstract level. While database schema used by EMR systems, correspond to the SQL datatypes[16], the application of semantic reconciliation-on-read strategy, instead necessitates the usage of common programming primitive types. On the other hand, "validValues" are determined based on what is contained in some sampled portion of the database. It is represented by the set of all possible categorical values (such as values of the attribute with type string which can be used to represent names, address, and others), the minimum and maximum values enclosed within open intervals ("()") or closed intervals ("[]") to represent a range of numerical values (such as values of attributes with type float or integer), and boolean values (such as values represented by 1/0 or True/False).

$$\tau_{syntatic} = \{t \mid t\epsilon \{\{integer, float, boolean, string\} \uplus validValues\}\}$$
 (2)

The semantic type is represented by the Equation (3), which is the set of all concepts corresponding to the leaf nodes of the suffix tree, generated from the name of the attribute. Suffix trees are used to divide a string into components, which is a very useful base for quickly identifying the longest common subsequence between a pair of strings and data compression. In our case, these trees provide a manageable list of words, contained with the name of attribute. For each substring, represented by a path of the tree, we then identify the corresponding concept, if it exists, thereby producing a set of concepts which may represent the attribute.

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$$\tau_{semantic} = \{s \mid se \{concept_{i,j} \mid i \leq width(suffixTree(s)) \land je \{substring_i\}\}\}$$
 (3)

Finally, the set AA of Amplified Attribute (AA), holds all unique attributes of participating schema, as shown in Equation (4). For each element of this set, the attribute is unique if its properties from Equation (1), are not similar to any other element of this set. The similarity function " \sim " is a threshold based loose bound, on the disjoint elements of " \mathbb{P} ".

$$\exists a \epsilon A A \mid (\mathbb{P}(a) \land \forall b \epsilon A A \mid \mathbb{P}(b) \to a \sim b) \tag{4}$$

As a possible realization of this theoretical set notation form, and from a practical point of view, our methodology is presented in the following subsections.

Schema acquisition

In the first step of our semantic reconciliation methodology, we collect medical schema from five distinct EMR storage systems (S). These include patient reports from OpenEMR (s_1) , 100,000 patient records from EMRBOTS (s_2) [17], custom database design by Pan et. $al(s_3)$ for supporting regional clinics and health care centers in China [18], clinical knowledge discovery tool MedTAKMI-CDI (s_4) [19], and our custom implementation (s_5) . These five schema follow relational database design, with logical entities, such as demographics, diagnosis, medicine or others, placed into tables which can be further linked to one or more tables. Each of these schema, fulfills the need of their respective information processing applications, however the lack of interoperability is very much visible here. Here, heterogeneity is caused by various factors, including the lack of standard terminologies and different normalization level. While SNOMED-CT provides a mechanism for identifying the standard codes for clinical terms and LOINC can be used for laboratory related terms, most attribute names are created based on the gut feeling of the database designer. While this behavior is not wrong, it does create syntactic differences in the participating schema. Consider the terms "name" and "patientName", which refer to the same attribute of the patient entity, however there is no standard way to represent it, with both being correct. As a result, s_1 and s_3 use the former representation, while s_2 and s_5 use the latter. Differences in normalization cause semantic differences, due to which some data could be available in one schema but absent in others, such as OpenEMR demographics identifying the patient's residential location using specific attributes like "Address", "City", "State", "Postal Code", "Country", and others. Similarly, "EncounterDate" from s_5 is semantically similar to "BeginDate" of "openemr_MedicalProblems" table in s_1 , "Admission-StartDate" of "LabsCorePopulatedTable" in s_2 , "time" in "Diagnosis" table of s_3 , and "dateOfAdmission" in "Diagnosis" and "CareHistory" tables of s_4 . Finally, s_1 and s_3 have separate tables containing the medicinal prescription details, however the same details are unavailable in s_2 , s_4 , s_5 . Once again, this is not an incorrect behavior since this information, might not be a part of the context or the requirements Doe and Smith Page 6 of 9

for these EMR storage systems. In fact, the change in context of the EMR storage system from the initial time of development to a later stage of collaborative processing systems, is the main cause of heterogeneity. In order to provide an interoperable solution, it is therefore necessary to provide syntactic and semantic mappings, while taking the data instances (or tuples) into account. Data instances are used to enrich the semantic context of an attribute (a_i) by establishing its probable data type and possible value set. In our earlier work [8] s_1 , s_2 , and s_5 were used to generate over 115 million patient records, which are converted into a semi-structured form and stored in Hadoop Distributed File System (HDFS). We extended the same setup to further include

Schema processing

In order to process the EMR schema set S, we use the methodology shown in figAlgorithm. For each data representation s_i , we first serialize its tables into Comma Separated Value (CSV) formatted flat files. Each table has a separate file, which contains the attribute names in the header and the data instances (following the same format and CSV struture), in subsequent lines. Each file is loaded in memory and converted into an algorithmic iterable structure. For each attribute, we then generate the suffix tree, which provides all possible substring representations contained within the attribute name. Then for each node of this suffix tree, if its length is greater than 2 (to avoid lookups for alphabets), we then query UMLS with approximate search strategy. Using Lexical Variant Generation (LVG) this strategy retrieves an expanded list of associated semantic concepts. If at least one semantic concept is found for the substring, we replace the substring with the most relevant concept, based on the highest value of the confidence measure, creating the Suffix Concept Tree (SCT). The SCT is structurally smaller than or equal to the suffix tree, since all suffix which are smaller than 2 and those without a qualifying semantic concept, are discarded. Finally for each node of the SCT, the substring from the suffix tree and its corresponding concept is placed into a sentence, which is converted into an embedding vector using **BioBERT**. This embedding vector represents the domain qualified semantics of the source attribute.

We continue this process, until all substrings have been processed. This is followed by syntactic type check of the attribute, which is based on its data. Here we determine, if the attribute values correspond to one of Integer, Float, Boolean, or String types. Additionally, we collect the values of each corresponding data cell, and determine the set of possible values in each case. This semantic enrichment from UMLS concepts and syntactic enrichment from attribute types, is then saved into persistent storage for further processing.

Meanwhile the process continues for the next attribute, then the next table, and finally the next system, till no further processing is possible. The flat enriched schema, following the design shown in **figEnrichedSchema** provides the input to our knowledge base. The logically amplified structure of the attribute, AA, is represented in **figAttributeRepresentation**. Here, the elements of AA are categorized into three parts. "AttributeContext" contains the metadata conforming to an instance of the attribute's existence, with its name, container table name and schema name, acting as a pointer, and schema version, source, and recorded date providing

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the version control features. In this manner, the attribute's fully qualified reference, along with its existential context is identified. "AttributeType" then encapsulates the data type features of this attribute's instances, with its primitive data type, and values providing a representation of the attribute's instance. Finally, "Attribute Semantics" holds, the semantic information of this attribute, in the form of its suffix tree, SCT and Tree Embedding. A pair of AA's with distinct references, are then used as an input to the schema map generation process, which is explained in the next step.

Schema Map generation

Schema Maps, provide an interoperable bridge between two medical systems $(s_i \wedge s_j)$, by identifying the links between their participating attributes. This identification is based on the schema matching process, shown in Algorithm 1, which operates on a pair of amplified attributes, $(aa_i, aa_j) \mid aa_i \epsilon s_i \wedge aa_j \epsilon s_j$, and calculates the similarity score S.

Algorithm 1 Attributes similarity identifier

```
Input: AmplifiedAttributes aa_i, aa_j
                                           Output: Similarity S
 1: Similarity S = 0;
 2: if aa_i.AttributeContext == aa_i.AttributeContext then:
       S=1;
 4: else
 5:
        Syntactic Similarity SynSim = 0
 6:
        at_i = aa_i.AttributeType
 7:
        at_i = aa_i.AttributeType
       if (at_i.DataType == at_j.DataType)||(at_i.DataType \Leftrightarrow at_j.DataType) then:
 8.
 9:
            SynSim = 0.5
10:
        end if
        Semantic Similarity SemSim = 0
11:
12:
        a\vec{e}_i = aa_i.AttributeSemantics.TreeEmbedding
        a\vec{e}_j = aa_j.AttributeSemantics.TreeEmbedding
13:
        SemSim = \frac{a\vec{e}_i \cdot a\vec{e}_j}{\|a\vec{e}_i\| \|a\vec{e}_j\|}
14:
        S = (0.5*SynSim) + (0.5*SemSim)
15:
16: end if
17: return S
```

Schema Map evolution

Experimental Setup

Results

Results goes here

Acknowledgements

Text for this section...

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Abbreviations

Text for this section...

Availability of data and materials

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Ethics approval and consent to participate

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Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

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Figures

Figure 1 Schemas used for knowledge interoperablity.

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Figure 3 Sample figure title

Table ${\bf 1}$ Sample table title. This is where the description of the table should go

	B1	B2	B3
A1	0.1	0.2	0.3
A2			
A3			

Tables

Additional Files

Additional file 1 — Sample additional file title

Additional file descriptions text (including details of how to view the file, if it is in a non-standard format or the file extension). This might refer to a multi-page table or a figure.

 $\label{eq:Additional} \mbox{ Additional file 2} \mbox{ — Sample additional file title } \mbox{ Additional file descriptions text.}$