Synergizing Data Imputation and Electronic Health Records for Advancing Prostate Cancer Research: Challenges, and Practical Applications

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Abstract: The presence of detailed clinical information in electronic health record (EHR) systems presents promising prospects for enhancing patient care through automated retrieval techniques. Nevertheless, it is widely acknowledged that accessing data within EHRs is hindered by various methodological challenges. Specifically, the clinical notes stored in EHRs are composed in a narrative form, making them prone to ambiguous formulations and highly unstructured data presentations, while structured reports commonly suffer from missing and/or erroneous data entries. This inherent complexity poses significant challenges when attempting automated large-scale medical knowledge extraction tasks, necessitating the application of advanced tools, such as natural language processing (NLP), as well as data audit techniques. This work aims to address these obstacles by creating and validating a novel pipeline designed to extract relevant data pertaining to prostate cancer patients. The objective is to exploit the inherent redundancies available within the integrated structured and unstructured data entries within EHRs in order to generate comprehensive and reliable medical databases, ready to be used in advanced research studies. Additionally, the study explores potential opportunities arising from these data, offering valuable prospects for advancing research in prostate cancer.

1 INTRODUCTION

Prostate cancer (PCa) is a prevalent disease known for its indolent nature, often characterised by slow development and protracted progression over time (National Cancer Institute, 2023; The American Cancer Society medical and editorial content team, 2019). As such, one specific challenge in performing medical research on PCa is dealing with incomplete medical records and missing data, e.g., as a result of city relocation or disease follow-up across different health providers. This, in turn, can hinder the results of ongoing research studies analysing the effectiveness of diagnoses and various treatment planning approaches (Holmes et al., 2021). Ultimately, this can affect clinical decision-making and the patient’s well-being.

To overcome the limitations of incomplete and/or erroneous data, Electronic Health Records (EHRs) mining has emerged as a crucial approach in medical research as well as within clinical practice (Yadav et al., 2018). EHRs mining leverages advanced data analytic and artificial intelligence (AI) approaches to extract valuable insights from vast amounts of patient data (Ajmal et al., 2023; Javaid et al., 2022). By identifying patterns, trends, and risk factors associated with prostate cancer, EHRs mining facilitates the early detection of advanced diseases and the personal-
isation of treatment strategies (Knighton et al., 2016; Seneviratne et al., 2018; Henkel et al., 2022). However, challenges such as missing data and data security must be addressed to ensure patient information remains complete, confidential, and secure. Additionally, the lack of interoperability between different EHR systems poses hurdles in data sharing and aggregation, limiting the full potential of mining for both prostate cancer research, as well as for general improvement of patient care (De La Torre-Diez et al., 2013). Overcoming these issues and promoting standardised data collection practices and protocols will be pivotal in advancing the field of PCa treatment through EHRs mining (Herp et al., 2023), as well as the overall medical research in general.

In our work, we have designed and developed a data preprocessing pipeline that can leverage routinely collected information from our EHRs (HUS Datalake (Bruck, 2023; Pylkäs, 2023; Misukka, 2022)) to efficiently and accurately retrieve and consolidate clinicians’ work on PCa treatment analysis. Using Microsoft Azure machine learning studio and batches from HUS datalake that are available at the HUS Acamedic environment (a secure scalable data analytics platform developed for medical research (kuorttinen, 2023)), we developed an EHR mining pipeline using Python libraries to read, process, and provide curated data for further research applications.

One of the key clinical inputs exhibiting missing entries within the EHR of a significant number of PCa patients is the occurrence of curative treatment, i.e., radical prostatectomy (RP) or radiation treatment (RT). Since the imputation of such missing data is inevitable, we had to use a different approach to uncover these lost data entries. Using routinely collected values of the prostate-specific antigen (PSA) lab measurements, we were able to successfully identify and even classify curative PCa treatments. To our knowledge, this is the first attempt to approach the inference of EHR missing treatment records through PSA time series data.

Our approach enabled us to enhance our EHR by incorporating approximately 2.8 thousand new curative treatment events, marking a notable 27% growth compared to the treatment events available before-hand. The explanation for this relatively large increase is multi-folded. Some patients might have been treated outside the (Helsinki and Uusimaa) district unit whose database our study is based upon. Others might have been treated within private practice units, which again are not covered by our database. Finally, we can assume that a proportion of these missing treatment events are due to human error in correctly recording them within the EHR.

Another key clinical information (as well as key surrogate measurement within medical research analysis) which is most of the times not directly recorded within EHRs, either in structured or non-structured format, is the time instant when PCa patients are classified as having a biochemical recurrence (BCR). After primary cancer treatment, BCR is achieved when the PSA level in the blood surpasses a certain threshold, thus indicating that the disease may be returning or progressing. Thus, BCR status is an important indicator both clinically, as it signs that further monitoring or treatment may be needed to manage the condition (Stephenson et al., 2006; Artibani et al., 2018), and from a (medical) data analysis perspective, as it is a surrogate for PCa mortality (Zhao et al., 2022; Artibani et al., 2018). By following the PSA measurements as well as all EHR-available PCa treatment records we were able to effectively determine (and report) the status and timing of BCR for all PCa patients.

2 METHODS

2.1 Data Source

Our pipeline starts by identifying patients of interest within a large academic EHR system (Figure 1). We used the Finnprostate dataset, which is a large patient registry study combining Finnish national healthcare data with local hospital data (n=700,000) of men suspected of having PCa (PSA measured) or diagnosed with PCa. From Finnprostate, we gathered a HUS (Hospital District of Helsinki and Uusimaa) sub-cohort of men (n=326,796) having comprehensive patient information regarding out-patient clinic and hospital visits as well as data regarding laboratory tests, medication prescriptions, radiological, pathological, and surgical reports, as well as comorbidities covering the years 1993 to 2019. The above data is embedded within the regional HUS Acamedic datalake.

Medical research commonly encounters missing data. Despite this prevalence, it is nowadays generally accepted to perform various data analysis tasks on partially incomplete records, as long as the missing values are not substantial, and the analysis methods themselves can cope with specific uncertainties. Moreover, the use of advanced imputation techniques such as maximum likelihood (Wald, 1949), multiple imputation (Schafer, 1999), or Bayesian methods (Kong et al., 1994) have a good track record in addressing many of the missing data entries. However, certain complex missing data records, such as the moment and type of a deployed treatment, or the first di-
agnostic biopsy of a tumour and its aggressiveness, are very hard to be addressed by any of the available computational imputation methods.

In our data processing work (yellow box, Figure 1), imputation was reinforced with customised algorithms that rely on clinical guidelines, experts’ interpretations, as well as the intrinsic information redundancy available within EHR, in order to retrieve the missing data. All created algorithms are described in Table 1.

2.2Missing Curative Treatment Detection

The Treatment Detection Algorithm (DTX_algo) plays a pivotal role in enhancing our data quality by identifying and incorporating missing curative treatment records (in Algorithm 4). The algorithm takes all patient’s data as input and returns a list of missing curative treatments.

The Significant PSA Drop Algorithm (SIGDROP) constitutes the initial phase of DTX, meticulously tracking a patient’s PSA values subsequent to their diagnostic biopsy (Algorithm 1). The algorithm takes PSA measurements of patient i, and returns, if any:

- \( \text{drop date} \): The date of the PSA drop, which is the highest (maximum) point from where a significant PSA drop starts; is subsequently considered as a treatment date.
- \( \text{nadir date} \): The date of the PSA nadir, which is the lowest (minimum) point to where the significant drop reached.
- \( \text{PSA}_{\text{min}} \): The minimum values (at the time \( \text{nadir date} \)); this value is used to classify the drop into radical prostatectomy or radiation therapy.

The algorithm’s operation commences with the pursuit of the maximum PSA value (\( PSA_{\text{max}} \), lines 3-4), followed by an endeavour to identify the minimum value within the ensuing \( \delta \leq 12\)-month period (lines 5-32). Upon successful identification of a decreasing value, at lines 15-16, the algorithm calculates \( \alpha \), which is the drop percentage that undergoes rigorous testing to ascertain its adherence to predetermined significance conditions (line 17). This process is indispensable in establishing the genuineness of the observed drop and confirming its clinical significance.

Having validated the drop as significant, and (line 6) with no EHR-recorded curative treatment between the date of drop (\( d_{\text{max}} \)) and the date of the nadir (\( d_{\text{min}} \)), DTX proceeds to collate all such identified drops, systematically categorising them into two distinct treatment modalities: radiation therapy (RT) and radical prostatectomy (RP) (Algorithm 2 line 7-10). This classification not only facilitates comprehensive treatment record augmentation but also provides valuable missing insights into the patient’s therapeutic journey.

2.3Biochemical Recurrence Detection

Biochemical recurrence (BCR) serves as a crucial indicator for PCa mortality. However, its availability in EHRs is not always guaranteed. In such cases, various methods can be employed to retrieve and impute this information. Our Detect Biochemical Recurrence (DBCR_algo) Algorithm is specifically designed to analyse data from treated patients, identifying potential relapses and categorising patients as either having experienced a BCR or not (Algorithm 7). To achieve this outcome, DBCR utilises four (04) distinct functions, each tailored to a specific task.

Clinical guidelines governing PSA relapse are stringent and clearly defined (Van Den Broeck et al., 2020), and these guidelines are meticulously integrated into the PRP and PRT functions (Algorithms 5-6).

- PSA-based relapse after radical prostatectomy \( PRP(p_i) \): this function uses the European Association of Urology (EAU) guidelines (Van Den Broeck et al., 2020) to detect whether a
Table 1: Summary of Algorithms.

<table>
<thead>
<tr>
<th>Algorithm Name</th>
<th>Input</th>
<th>Output</th>
<th>Complexity</th>
<th>Short Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SIGDROP</td>
<td>PSA</td>
<td>drop_date, nadir_date, PSA_{max}</td>
<td>O(M)</td>
<td>Detects significant PSA drop and related dates.</td>
</tr>
<tr>
<td>DTX</td>
<td>PATIENTS_LIST</td>
<td>L</td>
<td>O(M + N)</td>
<td>Detects missing treatments based on PSA data.</td>
</tr>
<tr>
<td>CRT</td>
<td>p_i</td>
<td>d_{zi}</td>
<td>O(1)</td>
<td>Detects Clinical Relapse after RT.</td>
</tr>
<tr>
<td>CRP</td>
<td>p_i</td>
<td>d_{zi}</td>
<td>O(1)</td>
<td>Detects Clinical Relapse after PT.</td>
</tr>
<tr>
<td>PRP</td>
<td>p_i</td>
<td>d_{zi}</td>
<td>O(N)</td>
<td>Detects PSA Relapse after PT.</td>
</tr>
<tr>
<td>PRT</td>
<td>p_i</td>
<td>d_{zi}</td>
<td>O(N)</td>
<td>Detects PSA Relapse after RT.</td>
</tr>
<tr>
<td>DBCR</td>
<td>TREATED PATIENTS</td>
<td>L_{acr}</td>
<td>O(M + N)</td>
<td>Main algorithm for BCR detection.</td>
</tr>
</tbody>
</table>

- RF=Radical prostatectomy, RT=Radiation therapy, BCR=Biochemical recurrence.
- In M*N, M is the number of PSA measurements and N is the number of patients.

Algorithm 1: SIGDROP - Significant PSA drop detection.

Input: PSA

Output: drop_date, nadir_date, PSA_{max}

1: M ← size(PSA)
2: if M ≥ 0 then
3: PSA_{max} ← PSA_{i}[1]
4: date_PSA_{max} ← getDate(PSA_{max})
5: for j = 1 to M − 1 do
6: i ← PSA_{j}[1]
7: if i ≥ 0 then
8: date_PSA_{max} ← date_PSA_{max} − date_PSA_{max}
9: if i ≥ 0 then
10: if (PSA_{max} < PSA_{j}[1] or δ ≥ 12m then
11: PSA_{max} ← PSA_{j}[1]
12: date_PSA_{max} ← getDate(PSA_{max})
13: end if
14: else
15: [i ← PSA_{max} − PSA_{j}[1]
16: α ← PSA_{max}
17: if (α ≥ 0.75 and β ≥ 3) or (α ≥ 0.5 and β ≥ 4) then
18: PSA_{max} ← PSA_{j}[1]
19: else
20: if δ ≥ 12m then
21: PSA_{max} ← PSA_{j}[1]
22: date_PSA_{max} ← getDate(PSA_{max})
23: else
24: γ ← date_PSA_{j}[2] − date_PSA_{max}
25: if j + 2 ≤ M and γ ≥ 12 then
26: PSA_{max} ← PSA_{j}[1]
27: date_PSA_{max} ← getDate(PSA_{max})
28: end if
29: end if
30: end if
31: end if
32: end for
33: end if
34: if PSA_{max} exists then
35: drop_date ← getDate(PSA_{max})
36: nadir_date ← getDate(PSA_{max})
37: return drop_date, nadir_date, PSA_{max}
38: end if
39: return NULL

PSA-based relapse occurred after radical prostatectomy. If an ultrasensitive PSA (Shen et al., 2005) measurement p_{si} was taken for patient p_i then we take this into consideration to define the maximum threshold (lines 3-7).

- PSA-based relapse after radiation therapy \textit{PRT} (p_i): this function also uses the EAU guidelines (Van Den Broeck et al., 2020) to detect whether a PSA-based relapse occurred after radiation therapy. The algorithm searches for the first increase of 2 PSA units from a nadir value.

Going beyond this, our novel BCR detection method is not solely reliant on PSA relapse; instead, it incorporates expert knowledge and translates it into a new tool for detecting BCR based on secondary treatments (Figures 2 and 3). The CRP and CRT functions (Algorithms 2-3) have been developed to identify possible relapses that may have been missed (after an RP or an RT primary treatment, respectively) either due to the absence of PSA tests or because the curating doctor decided on a secondary treatment before the PSA value has crossed the EAU-guideline threshold.

The exact approaches used to define clinical relapse after RP and RT primary treatments are described in Figure 2 and Figure 3, respectively.

Algorithm 2: CRP - Clinical Relapse after RP.

Input: p_i

Output: d_{zi}

1: L ← [1]
2: if lastRPDate(p_i) > firstRPDate(p_i) then
3: if lastRPDate(p_i) − firstRPDate(p_i) > 1yr then
4: L ← L + max(firstRPDateAfterOneYear(p_i))
5: end if
6: if hasHTCT(p_i) and lastHTCTDate(p_i) > firstRPDate(p_i) then
7: if lastHTCTDate(p_i) − firstRPDate(p_i) ≥ 2yr then
8: L ← L + max(firstHTCTDateAfterOneYear(p_i))
9: end if
10: end if
11: else
12: if hasHTCT(p_i) and lastHTCTDate(p_i) > firstRPDate(p_i) then
13: L ← L + max(firstHTCTDateAfterRp(p_i))
14: end if
15: end if
16: d_{zi} ← getMin(L)
17: return d_{zi}
Algorithm 3: CRT - Clinical Relapse after RT.

Input: $p_i$  
Output: $d_{m1}$  
1: $L ← []$  
2: if hasRP($p_i$) and $\text{lastRPDate}(p_i) > \text{firstRTDate}(p_i)$ then  
3: $L ← L + \text{firstRpDateAfterRt}(p_i)$  
4: end if  
5: if hasSecondRT($p_i$) and $\text{secondRTDate}(p_i) - \text{firstRTDate}(p_i) > 1$yr then  
6: $L ← L + \text{secondRTDate}(p_i)$  
7: end if  
8: if hasHTCT($p_i$) and $\text{firstHTCTDate}(p_i) - \text{firstRTDate}(p_i) ≥ 6$m then  
9: $L ← L + \text{firstHTCTDate}(p_i)$  
10: end if  
11: if hasHTCT($p_i$) and $\text{firstHTCTDate}(p_i) - \text{firstRTDate}(p_i) > 3$yr then  
12: $L ← L + \text{firstHTCTDate}(p_i)$  
13: end if  
14: $d_{m1} ← \text{getMin}(L)$  
15: return $d_{m1}$

The DBCR Algorithm then uses all the outputs of the above functions, namely the dates ($d_1, d_2, d_3, d_4$) of possible BCR occurrences, and selects the earliest date (if it exists) as the date of biochemical recurrence for patient $p_i$ (Algorithm 7 lines 7-10).

2.4 Evaluation

Retrieving missing data is of utmost importance in the pre-processing of EHR data for critical and sensitive applications. Additionally, assessing the quality of imputed data holds significant value as it provides insights into the effectiveness of the methods and algorithms employed. In our study, data evaluation involves a two-tier validation process.

The first level (a.k.a. 'step-1’ evaluation) employs automated tests, where we verify the accuracy of our algorithms by taking records without missing treatment data, applying the imputation algorithm, and subsequently scrutinising the outcomes.

The second level (a.k.a. the ‘step-2’ evaluation) entails expert validation, wherein a random selection of imputed data is manually inspected by domain experts, ensuring its correctness.

Due to the absence of biochemical recurrence data in our EHR, we assessed the effectiveness of the DBCR algorithms by manually evaluating patient outcomes and employing descriptive statistics.

3 RESULTS

3.1 Curated Database

The initial phase of this work was to explore the HUS datalake (Bruck, 2023; Pylkäs, 2023) and extract the most accurate and comprehensive prostate cancer data suitable for subsequent medical research applications. As a result, we successfully created a structured and curated database that contains crucial patient information, as defined in Table 3.
Table 2: Evaluation of DTX algorithm performance.

<table>
<thead>
<tr>
<th>Available CTx</th>
<th>Estimated CTx</th>
<th>Correct estimated CTx</th>
<th>New estimated CTx</th>
</tr>
</thead>
<tbody>
<tr>
<td>PID 7563</td>
<td>9723</td>
<td>6962 (92%)</td>
<td>6294 (90%)</td>
</tr>
<tr>
<td>PID-RP 2495</td>
<td>2722</td>
<td>2233 (90%)</td>
<td>1929 (86%)</td>
</tr>
<tr>
<td>PID-RT 5068</td>
<td>7003</td>
<td>4729 (93%)</td>
<td>4365 (92%)</td>
</tr>
</tbody>
</table>

Table 3: The curated data tables.

<table>
<thead>
<tr>
<th>Data</th>
<th>Number of rows (%)</th>
<th>Number of Patients (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1: Pathology 23,393</td>
<td>12,277</td>
<td></td>
</tr>
<tr>
<td>GG1 6618 (28)</td>
<td>3652 (30)</td>
<td></td>
</tr>
<tr>
<td>GG2 6383 (27)</td>
<td>3313 (27)</td>
<td></td>
</tr>
<tr>
<td>GG3 4747 (20)</td>
<td>2503 (20)</td>
<td></td>
</tr>
<tr>
<td>GG4 2310 (10)</td>
<td>1195 (10)</td>
<td></td>
</tr>
<tr>
<td>GG5 335 (14)</td>
<td>164 (13)</td>
<td></td>
</tr>
<tr>
<td>T2: Treatment 40,369</td>
<td>9800</td>
<td></td>
</tr>
<tr>
<td>RP 2743 (7)</td>
<td>2742</td>
<td></td>
</tr>
<tr>
<td>RT 18,254 (45)</td>
<td>7248</td>
<td></td>
</tr>
<tr>
<td>HT 15,804 (39)</td>
<td>4088</td>
<td></td>
</tr>
<tr>
<td>CT 3568 (9)</td>
<td>514</td>
<td></td>
</tr>
<tr>
<td>T3: PSA 1,424,440</td>
<td>238,399</td>
<td></td>
</tr>
<tr>
<td>T4: MRI 20,103</td>
<td>15,807</td>
<td></td>
</tr>
<tr>
<td>T5: Medications 13,837,600</td>
<td>290,055</td>
<td></td>
</tr>
</tbody>
</table>

GG1-GG5 = Gleason grade group 1-5 (associated to each pathological entry)
RP=Radical prostatectomy, RT=Radiation therapy, HT=Hormonal therapy, CT=Chemotherapy.

3.2 Treatments Data

Following the data curation and structuring, we have implemented the DTX algorithm in order to detect and impute the missing curative treatment data. As a result, our database now incorporates n=2763 new PCA-related treatment records, representing a 27% increase compared to the original data found in the HUS data lake. The number of patients with RP has increased by 16% (n=489), while the number of those with RT has increased by 31% (n=2274).

In Table 2 we present the results of 'step-1' DTX performance evaluation, i.e., estimated vs. known (EHR-available) treatment records. We record an imputation performance of 92% (n=6962) correct estimated curative treatments, i.e., treatments estimated using the DTX algorithm that are also found in the existing database. Out of these, 90% (n=6294) are correctly classified as RP or RT, whereas 10% (n=669) are wrongly classified. RP classification was 86% correct, whereas RT classification reached 92%.

The 'step-2' evaluation of the DTX algorithm was performed vs. manual validation by domain experts, where the experts were using the entire collection of unstructured reports associated with the test subjects in order to uncover their treatment history. The 'step-2' evaluation started by sampling 40 random patients, i.e., 20 random RP + 20 random RT, that were detected by the algorithm as having curative treatments (CTx), however this treatment did not appear within the EHR (DTX ∩ DB in Table 2). The results of this manual validation are summarised in Table 4. Only one patient from the RP group was unverifiable (no data = treatment cannot be confirmed), while five RT patients had the same situation. In addition, 95% of RP patients were confirmed to have a curative PCa treatment, and 60% of RT patients were confirmed. In total 79% of the sampled patients (whose treatments were not recorded within EHR) were confirmed to have PCa curative treatment.

Table 4: Manual validation for DTX algorithm performance.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Unverifiable</th>
<th>Verifiable</th>
<th>True CTx</th>
<th>All True CTx</th>
</tr>
</thead>
<tbody>
<tr>
<td>PID-RP</td>
<td>20</td>
<td>01</td>
<td>19 (95%)</td>
<td>18 (95%)</td>
</tr>
<tr>
<td>PID-RT</td>
<td>20</td>
<td>05</td>
<td>15 (75%)</td>
<td>09 (60%)</td>
</tr>
</tbody>
</table>

3.3 BCR Data

Our DBCR algorithm successfully identified 2851 patients (Figures 4 and 5) who developed a biochemical recurrence after a PCa curative treatment. These patients represent 27% of the treated patients.

Among the identified BCR patients, 70% (n=2007) were detected using the PRP and PRT algorithms, which are in accordance with the EAU guidelines (Van Den Broeck et al., 2020). However, around 30% (n=844) were identified using our new algorithms, CRP and CRT, formulated based on the expertise of our clinicians’ team and other contributors to this work. Notably, without applying these new algorithms, these cases might have otherwise gone unnoticed.

Figure 4: BCR detected data: Bar plots representing the distribution of the time from curative treatment to relapse.
3.4 Applications

After cleaning the data and improving its quality, we have successfully carried out multiple applications. The primary application involved developing a data visualisation tool, enabling clinicians and researchers to visualise the trajectory of PCa patients, including their PSA values, treatments, pathological results, medical prescriptions, and others (Figure 6).

Additionally, we have investigated the potential grade inflation in PCa Gleason grade groups during the MRI era; the research focused on patients in Gleason grade groups 1 and 2. This work hypothesised that some patients in grade group 1 in the pre-MRI era are nowadays, in the MRI era, classified and treated as grade group 2 patients. With enough evidence, the work proved the hypothesis which will open serious discussions to reassess current risk stratification tools and clinical decision-making. Updating guidelines on cancer grading and treatments is crucial to be aligned with the precision of modern MRI technology.

Furthermore, we are utilising the curated EHR data to train machine learning models to predict biochemical recurrence within the following 3-to-10 years from initial curative treatment. Knowing that prostate cancer is a slow-developing cancer, BCR is one of the most important and accurate surrogates for prostate cancer mortality. Therefore, predicting BCR would have a significant impact on treatment decisions and treatment planning. Our (preliminary) trained models achieved good performance (Accuracy=0.93, AUC=0.93, Precision=0.88) on an internal validation. The models are trained on n=5262 patients who have had PCa curative treatment.

4 DISCUSSION

Ensuring high data quality is essential when building effective AI models and conducting significant statistical analyses (Gudivada et al., 2017). This importance is particularly heightened in clinical research and applications where decisions may directly impact patients’ lives. Electronic Health Records (EHR), such as the one available at HUS Acamedic, play a critical role in this process, making it imperative to develop robust exploration methods to harness the available data.

In our work, we explored, curated, and augmented bio-medical data from within Finnish healthcare records, with a specific focus on prostate cancer patients. By establishing a new mining framework and developing novel analysis algorithms, we successfully consolidated our data, enabling us to conduct meaningful and impactful medical research.

One of our approaches was to use the time series data on patients’ PSA levels, a subset of medical data which is typically well collected and curated within EHR, in order to infer the existence, and the type, of EHR-missing curative treatment events. To our knowledge, this is the first time PSA time series data were used in this way, although, in (Bettencourt-Silva et al., 2015), the authors employed a similar approach to generate a completeness score for the overall data quality of the cohort. Based on this approach, we were able to consolidate our EHR by adding approx. 2.8k new curative treatment events, representing a 27% increase from the EHR-available treatment events.

Another important outcome of our mining framework was documenting the status and timing of our PCa patients’ BCR. Differently than in previous EHR mining frameworks for PCa medical data, see e.g., (Park et al., 2021b; Park et al., 2021a), we define BCR-status based on both PSA-level measurements (after primary curative treatment, i.e., radiation therapy –RT– or radical prostatectomy –RP–) as well
as based on secondary curative and adjuvant therapies, i.e., PCa related hormonal- and/or chemotherapy. This approach takes into consideration the clinical reality that sometimes, curating doctors decide on secondary therapies before the PSA level crosses the threshold established by current EAU guidelines as the BCR level. Using this approach, we accurately captured an additional 844 BCR events (representing a 42% increase from PSA-only detected BCR events), which otherwise would either not have been found at all or would have been given a significant later timestamp.

One important observation from our EHR data curation and analysis work is that there exists a large amount of redundancy in these data sources. This is particularly observable within the free text input written by doctors during their medical checkups and/or lab, pathological, or imaging reports. On the other hand, due to a multitude of factors, including human error, focusing on only one particular type of data source at a time, such as lab results, pathological reports, or even surgery records, one encounters a significant amount of missing data entries.

Therefore, leveraging the data redundancy feature in EHR not only makes it possible and highly advantageous to recover these missing data entries but also validates and assesses the outcomes of our algorithms. This is why, a “data investigation” approach, such as the one described in this manuscript, is more relevant than classical “data imputation” methods. Indeed, these latter approaches provide only average-like behaviours and also are completely inefficient in detecting missing events, such as a radiation treatment event altogether missing from within the EHR.

Strongly connected to the above reasoning, one could not overlook the potential impact the use of Large Language Models (LLM) could have in detecting and augmenting the existing EHR data (Thirunavukarasu et al., 2023). Such models could be employed to extract (from the free text provided by doctors) relevant information such as missing events, e.g. treatments performed in different clinics, cities, or even countries, or information that is usually not structurally recorded within EHR, e.g., family history, use of alcohol and tobacco products, general health status of the patient, etc. During the current EHR data analysis no LLM was employed; however, the approach is currently actively analysed for future usage within our models.

5 CONCLUSION

This work demonstrates the challenges of mining Finnish electronic health records for prostate cancer (PCa) research, as well as the opportunities it offers in gaining valuable insights. Our methodology, when applied to the HUS dalatke, enabled the detection of missing treatments and biochemical recurrences (BCR), which led to a range of clinically relevant findings, including patients’ timeline histories, the Gleason grade group inflation finding, and the BCR classification models. The results of our framework highlight the potential of EHR data mining to advance PCa research and guide personalised patient care.

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REFERENCES


Algorithm 4: DTX - Missing treatments detection.
Input: PATIENTS_LIST
Output: L
1: L ← |
2: for all pi in PATIENTS_LIST do
3: PSAi ← getPsa(pi)
4: Tx_i ← getTreatments(pi)
5: (d_max, d_min, PSA_min) ← SIGDROP(PSAi)
6: if TXExists(d_max, d_min, PSA_min, Tx_i) = False then
7: if PSA_min < 0.1 then
8: tx_type ← 'RP'
9: else
10: tx_type ← 'RT'
11: end if
12: L ← L + (pi, tx_type, drop_date)
13: end if
14: end for
15: return L

Algorithm 5: PRP - PSA Relapse after RP.
Input: pi
Output: d_{a2}
1: PSA ← getPsaAfterRp(pi)
2: for psa in PSA do
3: if isP(pi(psa)) = TRUE then
4: th ← 0.2
5: else
6: th ← 0.4
7: end if
8: if psa > th then
9: d_{a2} ← getDate(psa)
10: return d_{a2}
11: end if
12: end for
13: return NULL

Algorithm 6: PRT - PSA Relapse after RT.
Input: pi
Output: d_{a2}
1: PSA ← getPsaAfterRt(pi)
2: nadir ← getMax(PSA)
3: for psa in PSA do
4: if nadir > psa then
5: nadir ← psa
6: end if
7: inc ← psa - nadir
8: if inc > 2 then
9: d_{a2} ← getDate(psa)
10: return d_{a2}
11: end if
12: end for
13: return NULL