STUDY PROTOCOL

Exploring polypharmacy with artificial intelligence: data analysis protocol

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Abstract

Background: Polypharmacy is common among older adults and it represents a public health concern, due to the negative health impacts potentially associated with the use of several medications. However, the large number of medication combinations and sequences of use makes it complicated for traditional statistical methods to predict which therapy is genuinely associated with health outcomes. The project aims to use artificial intelligence (AI) to determine the quality of polypharmacy among older adults with chronic diseases in the province of Quebec, Canada.

Methods: We will use data from the Quebec Integrated Chronic Disease Surveillance System (QICDSS). QICDSS contains information about prescribed medications in older adults in Quebec collected over 20 years. It also includes diagnostic codes and procedures, and sociodemographic data linked through a unique identification number for each individual. Our research will be structured around three interconnected research axes: AI, Health, and Law&Ethics. The AI research axis will develop algorithms for finding frequent patterns of medication use that correlate with health events, considering data locality and temporality (explainable AI or XAI). The Health research axis will translate these patterns into polypharmacy indicators relevant to public health surveillance and clinicians. The Law&Ethics axis will assess the social acceptability of the algorithms developed using AI tools and the indicators developed by the Heath axis and will ensure that the developed indicators neither discriminate against any population group nor increase the disparities already present in the use of medications.

Discussion: The multi-disciplinary research team consists of specialists in AI, health data, statistics, pharmacy, public health, law, and ethics, which will allow investigation of polypharmacy from different points of view and will contribute to a deeper understanding of the clinical, social, and ethical issues surrounding polypharmacy and its surveillance, as well as the use of AI for health record data. The project results will be disseminated to the scientific community, healthcare professionals, and public health decision-makers in peer-reviewed publications, scientific meetings, and reports. The diffusion of the results will ensure the confidentiality of individual data.

Keywords: Polypharmacy, Indicators, Medications, Artificial intelligence, Ethics, Social acceptability

Background

Medications represent a large proportion of health care spendings [1–7]. Their use is essential for older adults who suffer from multiple chronic diseases. Canadians aged 65 years and over receive on average 6.9 different classes of medications annually [8]. Polypharmacy, which is the simultaneous use of multiple medications by the same individual, has been associated with a plethora of
harmful health consequences, such as frailty, falls, cognitive problems, hospitalizations, and mortality [9–11]. It thus represents a potential harm for the patient and a financial burden for the health care system [7, 12, 13].

Nonetheless, there are circumstances in which the prescription of multiple medications is appropriate and leads to improved health outcomes. Distinguishing appropriate and inappropriate polypharmacy is an important and complex issue that is difficult to tackle [14]. First, there is a plurality of definitions of polypharmacy [15, 16]. Some of them are based solely on quantitative aspects (for example, more than 4, 5 or 10 simultaneously used medications), while others are based on qualitative characteristics (for example, the presence of inappropriate medications), and some use mixed approaches [15, 16]. Second, each combination of different medications has a different risk/benefit ratio due to the variety of potential drug-drug and drug-disease interactions [17]. The individual characteristics and clinical manifestations are also important drivers of the polypharmacy type and consequences [18–20]. Finally, the concept of polypharmacy is often studied as a static exposure to medications without considering past medication use or subsequent changes, which may limit the conclusions about the consequences of polypharmacy [17].

Identifying what combinations of medications or trajectories of treatment are associated with health outcomes would allow the development of specific polypharmacy indicators for public health surveillance [21, 22]. Such indicators would be useful to guide clinical practice, to implement interventions or policies and would allow for their evaluation thereafter.

Determining health outcomes associated with polypharmacy requires considering concurrent and sequential use of multiple medications, duration of treatment, medical and sociodemographic characteristics of individuals. This type of analysis of large amount of complex data is difficult to perform with traditional statistics, but it can be accomplished with artificial intelligence (AI) methods.

However, using AI in large health administrative data may pose ethical challenges, such as those related to re-identifying individuals or using the data sparingly, that is to restrict attention only to the variables necessary to answer the research question. Moreover, the polypharmacy indicators that are created must neither marginalize nor discriminate against any specific population. Taking into consideration the ethical and social aspects involved will be beneficial for researchers, clinicians, decision-makers, and for the population in general.

**Aim**
The general objective of the research program is to use AI within an ethical framework to develop polypharmacy indicators in older adults for surveillance and clinical practice in the province of Quebec, Canada.

Three interdependent research axes (Fig. 1) will be involved to reach this aim:

- **AI axis**: Define and detect polypharmacy in administrative databases to identify frequent combinations that correlate with health outcomes.
- **Health axis**: Develop polypharmacy indicators in older adults to carry out polypharmacy surveillance in public health and to guide clinicians.
- **Law&Ethics axis**: Explore the ethical and social acceptability aspects related to the use of AI for the development of polypharmacy indicators.

**Methods/design**

**Data source**
We will use the data from the Quebec Integrated Chronic Disease Surveillance System (QICDSS). This database was developed and is managed by the Quebec National Institute of Public Health (Institut national de santé publique du Québec, INS PQ) for the surveillance of chronic diseases [23]. It contains information about medication claims (name of prescribed medication, dose, dates of dispensation, duration of treatment), physician claims (dates, diagnostic codes based on the 9th version of the International Statistical Classification of Diseases and Related Health Problems [ICD-9]), hospitalizations (dates, diagnostic codes based on ICD-9 or ICD-10, provided services and interventions), deaths (date and up to 10 causes), and sociodemographic data (age, sex, region of residence). A unique identification number allows linking information from each file. The database also includes the material and social deprivation index, that is a validated substitute of socio-economic status [24], and a comorbidity index derived from Charlson and Elixhauser indices [25] that allows to quantify the burden of diseases of an individual. The QICDSS medication data cover more than 90% of the older population (65 years and older) since 1996.

**Methodological approach**
The three research axes described herein will closely collaborate to achieve the objectives of the project. As shown in Fig. 1, the results generated by each of the axes will serve as input for the other two axes and will thus make it possible to adjust their work.

**AI axis: Define and detect polypharmacy with AI**

**Context**
The aim of the AI axis is to define and detect polypharmacy associated with health outcomes. The definition of
polypharmacy is expected to incorporate several medications, their type, dosage and duration of use, or specific combinations of medications. For example, based on the QICDSS database, the AI axis will discover medication combinations that are associated with an increased risk of death, hospitalizations or gradual deterioration of the patient state (e.g., worsening of the fragility index and other outcomes that are not binary, which will be identified by the Health axis, objective 2.1). The main challenge will be to identify clinically relevant correlations between the addition of the new medications and health events among the millions of observations, excluding multiple accidental correlations and coincidences that analysis of such a dataset will inevitably reveal.

**Objectives and methodology**

Objective 1.1 aims to define inappropriate polypharmacy using frequent pattern detection algorithms from big datasets. It involves discovering combinations of medications that will allow prediction of subsequent health events such as hospitalization or death. As benchmarks, we will evaluate algorithms such as Apriori [26], FP-Growth [27], and Pattern Fusion [28], which we will compare to the patterns discovered by more novel data mining algorithms based on genetic algorithms [29, 30] and reinforcement learning [31–33]. The obtained list of medication combinations will be refined by taking into consideration such attributes as the dosage of each medication, the duration of treatment, and the order in which the medications were prescribed, and all that in the context of up to 20 years of follow-up observations. The collaboration between the AI axis and the Health axis will be necessary to determine which medications and treatment attributes should be prioritized.

Objective 1.2 will explore the social dimension of the data. Some subgroups of the population may be more sensitive to specific medication combinations and their outcomes, for example based on their age or medical conditions. The challenge will consist in finding an appropriate representation of individuals allowing them to be partitioned depending on demographic or geographic characteristics. This entails integrating heterogeneous individual-level data that contain attributes that are numeric (e.g., postal code that can be used to derive
Euclidean distance between two individuals), binary (e.g., sex), or that describe temporal dimension (e.g., time when a medication was taken and duration of its use). Deep neural networks have previously been proven to learn patient representation from electronic health records and will be considered for this task of clustering [34]. The frequent pattern detection algorithms developed in objective 1.1 will be applied independently to each subpopulation retrieved. They will help to determine which polypharmacy problems are present in these specific subpopulations and their differences from the general population studied in objective 1.1. We will also investigate the possibility of predicting the effect of treatment over time, considering the individual characteristics of a patient. Specifically, we will use recurrent neural networks that have been effective in predicting the series of events from electronic health records [35, 36]. Potential biases that can arise both from the composition of the data and their temporal nature will be of particular interest. For example, the older population in Quebec is mainly white and well-off. Therefore, the neural network is at risk of learning from decisions that benefit this group to the detriment of others. Close collaboration with the Health axis will make it possible to decide which attributes should be prioritized at the beginning of the research, and collaboration with the Law&Ethics axis will prevent the inclusion of an accidental discriminatory bias.

Objective 1.3 will focus on the challenges related to the use of AI strategies in the healthcare, more specifically on the problem of explainability. While demonstrating better predictive performance, the results obtained with neural networks are difficult to explain, which is a limiting factor when one needs to justify why a treatment is being recommended to a patient. Moreover, the complexity of the patterns discovered in the previous objectives and the potential presence of confounding factors [37] can make them very difficult for users to interpret [38]. This is what makes these systems “black boxes”; they give highly predictive solutions, but it is difficult or impossible for the users to understand how they arrive at them, which is a clear problem when patients are asked to trust their health and potentially their lives to these solutions. We will explore different strategies to make the models more explainable and interpretable. The system will notably be characterized in terms of explicitness, faithfulness, and stability [39].

Data preprocessing: The data are currently in the form of relational database tables that contain institutional codes. It will be necessary to transform them into groups of items with different levels of granularity of detail. The institutional codes will be arranged in a hierarchical or in ontological manner to allow algorithms to find similar health conditions and medications with the same mechanisms of action or active ingredients. The data preprocessing is a prerequisite for the proper functioning and success of the algorithms of the three objectives. This work will be carried out in accordance with FAIR principles (Findable, Accessible, Interoperable, Reusable) [40] and international standards (SNOMED CT, LOINC, FHIR) [41, 42]. By documenting the actions and activities carried out, we will promote continuous data valuation, reuse, and development.

Health axis: Development of polypharmacy indicators

Context
The Health axis aims to develop polypharmacy indicators that will be useful for population-level surveillance in order to guide health promotion and disease prevention interventions, and to support clinical practice. Thus, the Health axis will be responsible for translating the information generated by the AI axis into relevant and understandable indicators that will be socially acceptable through recommendations of the Law&Ethics axis (Fig. 1).

Objectives and methodology
Objective 2.1 will define health outcomes that reflect a continuum of changes in health status. Since deaths, and to a lesser extent, hospitalizations, are relatively rare events, only a small portion of the population will be affected. We will identify several types of metrics, such as frailty indices or composite health outcomes (e.g., frequency of medical and emergency room visits due to decompensation of a health condition) that will be useful to define various intermediate stages between states of good health, illness, and death. This approach will enable evaluation of the health outcomes for all individuals. In addition, we will consider institutionalization in the long-term care facilities as one of potentially “negative” health outcomes. The defined continuum of health outcomes will be transmitted to the AI axis and will be used to validate previous algorithms or create new ones, in order to identify polypharmacy patterns and characteristics associated with said outcomes.

Objective 2.2 will interpret the frequent patterns and deep representations discovered in the AI axis and transform them into polypharmacy indicators. The Law&Ethics axis will ensure that these indicators are socially acceptable, fair, and do not ignore potential harmful impacts on small subpopulations. Thereafter, each indicator will be validated for clinical and organizational utility on a test set: the data collected between 2016 and 2019. Predictive models will measure the indicator’s effectiveness to predict health events, such as
mortality and hospitalizations (multivariable Cox model), and continuum of health outcomes (regression analysis).

Cox models will use age as a time scale [43, 44]; all the models will be adjusted for the initial material and social deprivation indices [24], the initial comorbidity index [25] and any other variable deemed relevant by the collaborating experts. Particular attention will be paid to the effect of sex, assessed with stratified analysis. The indicators will be compared according to their discriminating power, i.e., the ability of the model to adequately distinguish between subjects with and without the specific health outcome, using the C-index and the Nagelkerke R², their calibration (slope and calibration curve), and their overall predictive power (observed probabilities compared to predicted probabilities) [45]. Statistics corrected for overfitting will be obtained using bootstrap [46]. Finally, we will evaluate the temporal stability of polypharmacy indicators by comparing their predictive capacities for cohorts from previous years (e.g., 2005–2008, 2010–2013).

An explanatory model will guide the estimation of causal links between indicators and health events. Given the variable nature of medication exposure and potentially confounding covariates, such as comorbidities, the relationship between polypharmacy indicators and clinical outcomes could be subject to time-dependent confounding [47]. We will use causal inference methods such as marginal structural models to estimate the effect of our polypharmacy indicators on hospitalizations, deaths and other health outcomes identified in objective 2.1 [48]. The fraction of events attributable to inappropriate polypharmacy, as defined by each polypharmacy indicator, will be calculated, making it possible to quantify the impact in terms of public health [49].

Finally, the robustness of the results to unmeasured confounding factors will be assessed using the E-value [50, 51]. The E-value indicates the strength of association between an unmeasured confounding variable and (1) the exposure and (2) the event that would be necessary to cancel the association between the exposure and the event.

The indicators will be used for the surveillance that is carried out by the INSPQ. These indicators should be useful and understandable for public health workers, decision-makers, clinicians, and the patients. Thus, the choice of indicators will be made jointly with these potential users with the help of consensus procedures, notably the RAND/Delphi approach [52, 53]. We will recruit approximately twenty participants: patients, clinicians (family doctors, geriatricians, pharmacists, nurses), researchers, public health decision-makers and surveillance specialists to evaluate the proposed indicators. We will assess the basic qualities of the suggested indicators, including validity and reliability, ease of use and ease of understanding [54, 55].

### Law&Ethics axis: Law, ethics, and social acceptability

#### Context

The collection, use, and communication of personal health information to train AI algorithms raise unique legal and ethical concerns and require a responsible approach [56]. Therefore, the Law&Ethics axis of the research project is necessary to ensure the legal conformity of the research process, to validate the usefulness of the results, and the acceptability and legitimacy of their application. This axis will involve researchers, practitioners, and patients.

#### Objectives and methodology

Objective 3.1 will aim to verify that the use of new correlations or additional emerging attributes in the AI Axis is carried out in an ethical manner. Particular attention will be paid to a set of risks associated with stigmatization, re-identification, and fairness. Stigmatization occurs when data reinforces the labelling of population subgroups, that are often already disadvantaged or taken responsible for their health problems. Data linkage operations may increase the risk of identifying patients from data sets that were previously anonymized. It may happen when few of them share specific characteristics, for example, due to the frequency of the studied phenomena, their nature, or the size of the territory where these phenomena are observed. Finally, the issue of fairness can arise when certain population subgroups are unduly excluded from analysis or under-represented in the dataset. The Law&Ethics axis will act as an internal ethics committee for the project, allowing researchers involved in the AI axis to handle these risky situations.

Objective 3.2 will consist of the ethical and social validation of the indicators developed in the Health axis. The approach will be based on grids developed by the INSPQ [57, 58]. Two data collection methods are planned: semi-structured interviews and focus groups. The interviews will be conducted with 30 people. There will be 5 focus groups of 4 to 6 participants. Using these results, we will build a specific ethic assessment grid that will allow weighting individual and collective interests related to the use of polypharmacy indicators and assessing social and ethical validity in the context of surveillance.

Objective 3.3 will focus on the development of markers and guidelines that could frame the ethical reasoning in the context of polypharmacy and AI. We will adopt an approach inspired by narrative ethics, a method primarily developed in medicine, which allows researchers and patients to narrate the challenges and ethical questions
they were confronted with during a particular project [59]. This method is mainly used to investigate emerging and unique ethical issues. The discussions of the experiences lived by researchers and other participants will shed light on many nuances and ethical subtleties of this complex and new practical case [60, 61]. The publication of the ethical considerations will provide the opportunity to highlight certain pitfalls to be avoided or certain attitudes to be adopted.

**Patient and public involvement**

Patients will be involved in the validation process of polypharmacy indicators and in discussions within the Law&Ethics axis. Among other things, they will ensure that the developed indicators meet their priorities and preferences.

**Discussion**

The impact of use of various combinations of multiple medications on individual health and health care system is largely unknown. This research program will be performed by a multi-disciplinary team consisting of specialists in AI, health data, statistics, pharmacy, public health, law, and ethics. Such diversity of expertise will allow investigation of polypharmacy from different points of view and will contribute to a deeper understanding of the clinical, social, and ethical issues surrounding polypharmacy and its surveillance, as well as the use of AI for health record data.

In addition, the polypharmacy indicators will be developed in partnership with patients, clinicians, researchers, and public health decision-makers, ensuring the proposed indicators to be valid, reliable, understandable, and simple for their use in surveillance and clinics.

One of the strengths of the research program is that it will be conducted on an extensive database that contains time-stamped medical and sociodemographic information covering more than 90% of older adults in the province of Quebec, Canada, over two decades.

The main limitation of the research is that the data do not contain variables related to race, ethnicity, gender, health habits, and specific clinical data that could influence both medication use and polypharmacy, and their associated outcomes.

**Anticipated challenges and mitigation strategies**

Each of the axes will meet challenges specific to their objectives. The complexity of the data and their interpretation will make it necessary to test different methodological approaches and will involve a lot of feedback between the axes. The older population is heterogeneous in terms of physiological, psychological, social, and cultural peculiarities. It will not be possible to directly investigate the role of individual factors such as life habits and genetics because they are not contained in the databases. Thus, since many factors other than medications may influence the health outcomes, the interpretation will have to be made with caution.

Similarly, the data mining process may generate a very large number of medication use patterns. Among those latter, some may be ambiguous, inconsistent, or even absurd. Therefore, the interpretation of the results will be of particular importance and will be jointly conducted by researchers and clinicians that will use their technical expertise and clinical experience.

Our team is committed to popularizing the procedures carried out so that all stakeholders can understand the processing that was done with the data. We are also aware that the advent of new techniques can open the path towards inconsistent approaches, and this is the reason why the Law&Ethics team will produce new knowledge, markers, and guidelines for future research in this area.

Our results will be published in peer reviewed scientific journals, presented at local and international scientific meetings, and shared with clinicians and healthcare professionals via the INSPQ leaflets, website, and public health information center.

**Abbreviations**

AI: Artificial intelligence; INSPQ: National Public Health Institute of Quebec; QICDSS: Quebec Integrated Chronic Disease Surveillance System; RAMQ: Régie de l’assurance maladie du Québec; MSSS: Ministry of Health and Social Services.

**Acknowledgements**

Not applicable

**Authors’ contributions**

CS, RK, AD, PLD—conception and design of the research, CS, RK, AD, PLD, OB, YC, DT, AB, PD, CG, FL, AMS, JC, TB, MS, SJ, VE—drafting and revising the paper. All authors have read and approved the manuscript.

**Funding**

This work is supported by the Canadian Institute of Health Research and the Natural Sciences and Engineering Research Council of Canada, grant number [CPRG—170621]. The funders had no role in the study design, and will have no role in data collection and analysis, decision to publish, or preparation of the manuscripts.

**Availability of data and materials**

Data from the QICDSS is not available publicly.

**Declarations**

**Ethics approval and consent to participate**

The use of INSPQ surveillance data is approved by the Quebec Commission d’accès à l’information and government agencies that own the files. The data are stored at the INSPQ and cannot be accessed elsewhere to meet the high security criteria and protection of personal information. All identifiers are encrypted, and several mechanisms ensure data confidentiality. Access to data is logged and limited to authorized personnel according to different levels of access. All researchers and students will follow the mandatory training of the
INSPQ to ensure secure data management. The Ethics Board of the Centre intégré universitaire de santé et de services sociaux de la Capitale Nationale has approved the conduct of this study. All consultations in the Health and Law&Ethics axes will be carried out according to ethical standards and will be approved by the Ethics Committees of our institutions (Université Laval and affiliated research centers). All participants recruited for interviews and focus groups will provide written consent.

Consent for publication
Not applicable.

Competing interests
Authors declare no competing interests.

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Received: 19 March 2021   Accepted: 12 July 2021

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