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ZiMM: a deep learning model for long term and blurry relapses with non-clinical claims data

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Abstract

This paper considers the problems of modeling and predicting a long-term and “blurry” relapse that occurs after a medical act, such as a surgery. We do not consider a short-term complication related to the act itself, but a long-term relapse that clinicians cannot explain easily, since it depends on unknown sets or sequences of past events that occurred before the act. The relapse is observed only indirectly, in a “blurry” fashion, through longitudinal prescriptions of drugs over a long period of time after the medical act. We introduce a new model, called ZiMM (Zero-inflated Mixture of Multinomial distributions) in order to capture long-term and blurry relapses. On top of it, we build an end-to-end deep-learning architecture called ZiMM Encoder-Decoder (ZiMM ED) that can learn from the complex, irregular, highly heterogeneous and sparse patterns of health events that are observed through a claims-only database. ZiMM ED is applied on a “non-clinical” claims database, that contains only timestamped reimbursement codes for drug purchases, medical procedures and hospital diagnoses, the only available clinical feature being the age of the patient. This setting is more challenging than a setting where bedside clinical signals are available. Our motivation for using such a non-clinical claims database is its exhaustivity population-wise, compared to clinical electronic health records coming from a single or a small set of hospitals. Indeed, we consider a dataset containing the claims of almost \textit{all French citizens} who had surgery for prostatic problems, with a history between 1.5 and 5 years. We consider a long-term (18 months) relapse (urination problems still occur despite surgery), which is blurry since it is observed only through the reimbursement of a specific set of drugs for urination problems. Our experiments show that ZiMM ED improves several baselines, including non-deep learning and deep-learning approaches, and that it allows working on such a dataset with minimal preprocessing work.

Equal contributions from A. Kabeshova and Y. Yu
1 Introduction

The increasing volume of Electronic Health Records (EHR) systems of national medical organizations in recent years allows to capture data from millions of individuals over many years. Each individual’s EHR can link data from many sources and hence contain “concepts” such as diagnoses, interventions, lab tests, clinical narratives, and more. This provides great opportunities for data scientists to collaborate on different aspects of healthcare research by applying advanced analytics to these EHR clinical data [60]. There are several challenges in processing EHR data [14]: data quality, high-dimensionality, temporality which refers to the sequential nature of clinical events, sparsity in both medical codes representation and in timestamp representation, irregularly timed observations, biases such as systematic errors in data collection, and mixed data types with missing data. Representation learning can overcome these challenges [60] and the choice of data representation or feature representation plays a significant role in the success of this approach.

1.1 Related works

Recently, much work has been done on developing compact and functional representations of medical records, including the use of deep learning over EHR data [5, 63]. A notable attempt is stacked denoising autoencoders [53]. Denoising autoencoders are also used in [9] to develop patient representation from various binary clinical descriptors on synthetic data. Sums of word-level skip-gram embedded vectors of clinical codes are used in [21] to create full-record representations. Word-level semantic embeddings for diagnosis and intervention codes are constructed in [59], using pooling and concatenation to aggregate embeddings into a vector representing a single admission, while [56] uses word-level embedding as preprocessing for a CNN (Convolutional Neural Networks) architecture.

Only a few studies were made based on claims data due to its complexity and rare availability. Choi et al. learned distributed representations of medical codes (e.g. diagnoses, medications, procedures) from electronic health records (EHRs) and claims data using Skip-gram and applied them to predict future clinical codes and risk groups [17]. Cui2vec is a recent study in learning clinical concept embeddings [8], which applied word2vec [52] and Glove [58] on multiple medical resources such as structured claims data, biomedical journal articles and unstructured clinical notes. Xiang et al. [71] proposed learning a distributional representation of clinical concepts considering temporal dependencies along the longitudinal sequence of a patient’s records based on claims data.

Other efforts have aimed at encoding temporal aspects of EHR data for predictive tasks. In [16], time-stamped events are used as inputs to a particular type of RNNs (Recurrent Neural Networks) to predict future disease diagnosis, while a graph-based attention model is used in [20] to learn concept representations. Another interesting recent effort is [60], which maps raw EHR data to the FHIR format (Fast Healthcare Interoperability Resources [10]) to encode EHR information for several different sequence-oriented models.

RETAIN [18] and GRAM [19] are two state-of-the-art models using RNNs for predicting future diagnoses. However, they cannot handle long sequences effectively. LSTM (Long Short-Term Memory) or bidirectional RNNs [46] can be trained using all available input information in the past and future,
and have been used to alleviate the effect of the long sequence problem and improve the predictive performance. Lipton et al. [43] have been the first to successfully use LSTM to predict patients diagnosis, for multi-label classification. BEHRT [40] is one of the most recent researches that provides the field with an accurate predictive model for the prediction of next diseases based on the transformer-based architecture in Natural Language Processing [28].

Although fixed timesteps are perfectly suitable for many RNN applications, EHRs often contain event-driven and asynchronously sampled samples, in particular for the application considered in the present paper. Recent works try to capture time irregularity using recurrent neural networks. Phased LSTM [55] tries to model the time information by adding a time gate to LSTM, which controls the update of the cell state, the hidden state and thus the final output. Another attempt is [73] where time gates are added to the LSTM in order to model time intervals and specifically better capture both of short-term and long-term sequential events. Time-aware LSTM [7] addresses time irregularity between two events in an LSTM architecture by decomposing the memory of the previous timesteps into short-term memory and long-term memory. Let us point out that, however, most of the works on EHR data cited above either ignores subsequence-level irregularity by partitioning the data into regular time windows and by aggregating the data within each window or handles this irregularity by simply adding the time-span as one coordinate of the features vector, see [16] for instance.

1.2 Aim of this paper and contributions

This paper aims at the construction of a predictive model for long-term and “blurry” relapses that occur after a medical act. We do not consider a short-term complication related to the act itself, but a long-term relapse which is observed only indirectly. In the example considered here, the medical act is a precise surgery, and the relapse is observed through longitudinal prescriptions of a specific set of drugs for urinary problems, over a period of 18 months after surgery. We use all the data available in SNIIRAM (French national health insurance information system, a huge database containing health reimbursements claims of almost all French citizens since 2015, see Section 3.1 for details) for patients on which such a surgery has been performed (TURP surgery, see Section 3.2).

The ZiMM model. The first natural idea is to cast this problem as a classification problem for “relapse” versus “no relapse” after the medical act. However, such a naive approach cannot work here. Indeed, the definition of a binary label would require threshold choices, both about time and dosage: after how many time and amount of drugs prescribed do we consider that there is a relapse? Such thresholds might depend on various clinical practices, habits of the patient, and many other exogenous factors. In this work, we propose to work directly on the data observed, by using all the “blurry” observations at once to train an end-to-end architecture. For this purpose, we introduce a new methodological contribution, namely the ZiMM model (Zero-inflated Multinomial Mixture) which is described in Section 2.1 below.

The ZiMM Encoder-Decoder (ZiMM ED). We introduce an end-to-end Encoder-Decoder architecture trained with an objective, based on the negative log-likelihood of the ZiMM model. All available patients’ claims before the medical act are first encoded into a single embedding vector by the Encoder.
This Encoder combines embedding layers, self-attention layers and recurrent layers to output an embedding vector of the full patient pathway prior to the medical act. This embedding is used as input to a Decoder which is dedicated to the learning of the parameters of the ZiMM model. This architecture is described in Sections 2.3 to 2.5 below.

**Specifics of this work.** The specificity of our approach lies in several points. First of all, it is performed on non-clinical EHR data, and we consider long-term predictions (18 months ahead). Our end-to-end architecture addresses several challenges, such as variable-size inputs, confounding interactions between medical codes, long-term predictions. Finally, we do not inject any prior knowledge: the embeddings of patient pathways are fully trained in an end-to-end fashion, we do not use any strong preprocessing or simplifying aggregations, the data is used almost in its raw form. We keep the smallest granularity possible both on the codes (we use the actual medical codes instead of encompassing categories) and on time, namely we work on the original 1-day time scale.

**Organization of the paper.** The ZiMM model is described in Section 2.1 while the ZiMM ED architecture is described in Sections 2.3 to 2.5. Some details concerning data preprocessing are provided in Section 2.2. Section 3 provides results and a comparison with strong baselines, together with some variations of the architecture to fine-tune the final performance and we conclude in Section 4.

## 2 Proposed architecture

The ZiMM ED architecture has three main building blocks: the ZiMM (Zero-inflated Multinomial Mixture) model (see Section 2.1), an Encoder (see Section 2.3) and a Decoder (see Section 2.4), and is described in Figure 1 below. Given a patient \( i \in \{1, \ldots, n\} \) (among \( n \) patients), with medical act of interest occurring at time \( T^i \), the steps through this architecture are roughly as follows:

- All the medical codes (drugs, diagnosis, medical procedures) observed longitudinally in the life of patient \( i \) before \( T^i \) are embedded, then aggregated in a time window using a self-attention mechanism. The time distance (in days) to \( T^i \) of each day with a non-empty set of codes is embedded as well, and we embed as well hospital stay durations. This sequence of vectors is then fed to a (or several) recurrent layers. This corresponds to Steps 1,2 and 3 in Figure 1. This leads to an embedding vector \( x_i \in \mathbb{R}^d \) of the full pathway of patient \( i \) prior to \( T^i \).

- The vector \( x_i \in \mathbb{R}^d \) is concatenated with other static features of the patient, and used as the input of the Decoder, which is based on several recurrent layers, in order to produce the parameters of the ZiMM model. This corresponds to Step 4 in Figure 1.

Precise descriptions of all steps are provided below, starting with the ZiMM model, followed by some details concerning data preprocessing, and the Encoder-Decoder architecture.

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4
Assume for now that we have an embedding vector $x_i \in \mathbb{R}^d$ that encodes the full health pathway of patient $i \in \{1, \ldots, n\}$ prior to $T_i$. This vector is the output of the ZiMM-Encoder described in Section 2.3 below. Moreover, we observe a vector of labels $y_i = [y_{i,1}, \ldots, y_{i,B}] \in \mathbb{N}^B$, which corresponds to the blurry observation of the relapse after $T_i$. Here, $B$ corresponds to the number of time intervals considered after $T_i$ and $y_{i,b} \geq 1$ is the number of blurry relapses observed in time bucket $b$, so that $y_{i,b} = 0$ means that no blurry relapse is observed in time bucket $b$. In the example considered Section 3 below, $B = 18$ corresponds to a 18-months period and $y_{i,b}$ is the number of drugs (among a set of drugs for urinary problems, see Section 3.3), purchased by patient $i$ in time bucket $b$.

We introduce $n_i = \sum_{b=1}^B y_{i,b} \in \mathbb{N}$, the overall number of blurry relapses of patient $i$. Let us point out that a binary classification problem ($n_i > 0$ versus $n_i = 0$) may wrongly classify very different situations corresponding to the same $n_i$. Consider for instance two extreme situations where first, we have $n_i = 1$ with $y_{i,1} = 1$ (and consequently $y_{i,b} = 0$ for $b = 2, \ldots, B$), and second, we have $n_i = 1$ with $y_{i,B} = 1$ (and consequently $n_{i,b} = 0$ for $b = 1, \ldots, B - 1$). The first case can be due to exogenous factors, such as the patient simply kept buying the drug after the surgery just out of a pure habit (prescriptions can run for a long period), while in the second example, there is no doubt that a relapse is occurring.

Therefore, we need to find a way to model the whole vector $y_i$, so that $n_i$ is not fixed and includes...
zero-inflation, namely a parametrized likelihood for \( n_i = 0 \). For that purpose, we introduce the following ZiMM model, where we suppose that \( n_i \in \{0, 1, \ldots, B\} \) is distributed (conditionally to \( x_i \)) as

\[
P(n_i = k|x_i) = \pi_k(x_i) \quad \text{for} \quad k \in \{0, \ldots, B\},
\]

where \( \pi_k(x_i) \) are such that \( \sum_{k=0}^{B} \pi_b(x_i) = 1 \) and \( \pi_b(x_i) \geq 0 \) (coming out of a softmax activation for instance). These parameters correspond to the categorical distribution specific to patient \( i \), and are constructed by the ZiMM-Decoder (see Section 2.4 below) from \( x_i \). Then, we assume that the distribution of \( y_i \) conditional to \( n_i = b \) and \( x_i \) follows either a Dirac distribution on vector (of size \( B \)) \([0, \ldots, 0]\) whenever \( b = 0 \), or a multinomial distribution of parameters \( b \) and \( p_{b,1}(x_i), \ldots, p_{b,B}(x_i) \), namely

\[
y_i|(x_i, n_i = b) \sim \begin{cases} \delta_{[0,\ldots,0]} & \text{if } b = 0, \\ \text{Multinomial}(b, p_{b,1}(x_i), \ldots, p_{b,B}(x_i)) & \text{otherwise}, \end{cases}
\]

where \( p_{b,1}(x_i), \ldots, p_{b,B}(x_i) \) are the parameters of a multinomial distribution whenever \( n_i = b \), for each \( b = 1, \ldots, B \). Once again, these parameters are specific to patient \( i \) thanks to the embedding vector \( x_i \in \mathbb{R}^d \), and are constructed by the ZiMM-Decoder. These parameters must satisfy \( \sum_{b'=1}^{B} p_{b,b'}(x_i) = 1 \) for \( b = 1, \ldots, B \), and \( p_{b,b'}(x_i) \geq 0 \) for any \( b, b' = 1, \ldots, B \), which is easily achieved with a softmax activation applied row-wise. By combining (1) and (2), with end up with the following mixture distribution for \( y_i \) conditionally on \( x_i \):

\[
y_i|x_i \sim \pi_0(x_i)\delta_{[0,\ldots,0]} + \sum_{b=1}^{B} \pi_b(x_i)\text{Multinomial}(b, p_{b,1}(x_i), \ldots, p_{b,B}(x_i)).
\]

Zero-inflation corresponds to the case where \( n_i = 0 \) and has likelihood measured by \( \pi_0(x_i) \). The ZiMM model has two sets of parameters that are functions of the feature vector \( x \), namely \( \{\pi_b(x)\}_{b \in \{1,\ldots,B\}} \) that parametrizes the distribution of the total number of blurry relapses and \( \{p_{b,b'}(x)\}_{b,b' \in \{1,\ldots,B\}^2} \) that parametrizes the dynamics of these relapses. As explained in Section 2.4 below, the dynamics of \( \{p_{b,b'}(x)\}_{b,b' \in \{1,\ldots,B\}^2} \) for each \( b \) are learned by dedicated recurrent layers in ZiMM-Decoder.

### 2.2 Preprocessing of the data

After a preliminary preprocessing based on our SCALPEL3 library described in [4], the data considered in this paper comes in the form of a table, where each row represents an event with the patient ID, the type of event (drug, diagnosis or medical procedure), its corresponding code, the start date and the end date of the event. Figure 2 provides an illustration of the sequence of claims observed for a single patient. If an event is related to a hospital stay, the start date is the first day of hospitalization while the end is the exit date from the hospital. Otherwise, the start and end dates are the same. The time delta is a day-long, and no aggregation is performed to keep the data as raw as possible.

Multiple events can occur within the same day, and the precise ordering of such events does not carry any information. Thus, we shuffle at random within-day events, to avoid any bias that could occur from the data collection mechanism (a similar strategy can be found in [21]).

All the events observed through claims before \( T_i^s \) are used as inputs to the Encoder described in Section 2.3 below. The output of the Encoder is the embedding vector \( x_i \in \mathbb{R}^d \) of patient \( i \). The blurry relapses observed after \( T_i^s \) are used to build the label vector \( y_i \in \mathbb{N}^B \) of patient \( i \).
Figure 2: Illustration of the sequence of claims observed for a patient. This corresponds to the output of Step 1 from Figure 1. Through claims, we observe three types of events: drug purchases (blue), medical procedures (red) and diagnosis (yellow) before the medical act (prostate surgery in the example) that happens at time $T^i$ for patient $i$. All events are timestamped, and the time delta is a day long. Several events can occur the same day and some days have no event: events are typically very irregularly sampled. All these events, observed before $T^i$, are used to learn the embedding vector $x_i \in \mathbb{R}^d$ of patient $i$. After $T^i$, we only keep the events corresponding to the blurry relapses considered (drug purchases among a set of drugs for urinary problems). These blurry relapses are used to build the label vector $y_i \in \mathbb{N}_B$ of patient $i$.

### 2.3 The ZiMM Encoder

The ZiMM Encoder corresponds to Steps (2) to (4) in Figure 1, a more detailed illustration is shown in Figure 3. Let us provide now details about each step, following the flow of the data.

#### Medical codes and timestamps embeddings.

The following is applied separately for each type of event code (drugs, medical procedures and diagnoses). Codes are first tokenized, and each unique token is individually mapped to an embedding vector in $\mathbb{R}^{d_E}$ which is learned during training, where $d_E$ is a hyper-parameter to be tuned (see Section 3.7). We consider only tokens that occur at least 50 times in the training dataset. To address time irregularity between two asynchronously timestamped events, we do the following: for any event occurring at time $t \leq T^i$ in the observation period of patient $i$, we compute the "time horizon" as $T^i - t$, namely the distance (in days) between the event and medical act. Moreover, whenever it makes sense, we compute end — start, the duration of the event, which corresponds to the duration of a hospital stay defined as the time between hospital admission and discharge. These two integers (in number of days) are also tokenized and replaced by a learned embedding vector. This allows the encoder to learn to put more or less emphasis on events that are close or far from $T^i$, and to exploit the duration of events as a proxy for severity of medical procedures and diagnoses. The patient age in years at $T^i$ is also similarly bucketized and embedded. The embedding of medical codes corresponds to Step (2) of both Figures 1 and 3. In our experiments, the default model uses $d_E = 64$ for the three types of codes and several regularization strategies are applied to avoid overfitting: weights decay ($\ell_2$ regularization with strength $10^{-2}$), multiplicative Gaussian noise, and layer normalization [3]. Since we observed empirically that these embeddings are prone to overfitting for the application considered in the paper, a comparison of the different regularization strategies is provided in Section 3.7.

#### Embeddings aggregation through self-attention.

Several events can occur at the same time (within the same day), so that the number of codes observed within a day is highly heterogeneous. Moreover, such codes are not likely to contribute equally to the vector representation of the day, and their order is not informative. Therefore, we need to perform a trainable aggregation of these codes in order to produce
Figure 3: A graphical illustration of representation learning of patient pathway in the ZiMM Encoder: (2) embedding tensors for each type of medical code; (3) aggregation of embedding vectors through multi-head self-attention inside a time window, where each gray block corresponds to one day \((t_0, t_1, \ldots)\), and \(T^i\) is the day of the medical act of interest; (4) concatenation of aggregated embedding vectors with embeddings for time horizon and hospital length stay.

a representation of the patient history at each timestamp. An approach can consist of using a hierarchy relationship between different diagnosis and treatment inside each patient visit [20]. However, in the data considered here, diagnosis codes are not explicit each time there is treatment; in particular, drugs purchases are almost never related to a diagnosis code (unless they are related to a hospitalization). Hence, we use a self-attention mechanism [68, 42] using a bag-of-features approach to learn how to combine embedding vectors within the same day, following previous successful applications of self-attention for fusing disease embeddings [45].

Let \( C \) be a set of codes with cardinality \(|E|\) (for drugs, medical procedures or diagnoses) and let \( E_C \in \mathbb{R}^{d_E \times |C|} \) be the corresponding matrix of concatenated embedding vectors. The multi-headed self-attention aggregation mechanism contains two layers, the first with \( K \) heads, each of which is a self-attention function that generates a probability vector of size \(|E|\) from \( E_C \):

\[
w_k = \text{softmax}(\alpha_k^\top \tanh(A_k E_C)),
\]

for \( k = 1, \ldots, K \), where \( \alpha_k \in \mathbb{R}^{1 \times d_E} \) and \( A_k \in \mathbb{R}^{d_E \times d_E} \) are to be trained and the second layer outputs a \( d_E \)-dimensional vector given by

\[
E_C \mu_C \quad \text{where} \quad \mu_C = \text{softmax}(b^\top \tanh(BW)),
\]

with \( W = \text{concatenate}(w_1, \ldots, w_K) \in \mathbb{R}^{K \times |C|} \) and \( b \) and \( B \) are trainable parameters. We use the \( \tanh \) squashing activation function following [42] and [45], since in these layers it is considered good
practice [39] to produce centered activations that are bounded in $[-1, 1]$, to help the back-propagation process involved in the training of the model. This self-attention mechanism is trained and performed separately for each type of code, as displayed in Step (3) of Figure 3. Note that this aggregation approach is permutation-invariant for codes that occur within the same day. We use dropout, drop-connect [69], Frobenius-norm weight penalization from [42] and weight-decay to regularize this self-attention mechanism.

**Health pathway encoder.** The inputs of the health pathway encoder, at each timestamp, contains the following embedding vectors, that are concatenated together:

- Medical code embedding computed by Equations (4) and (5);
- Time-horizon embedding (see above);
- Duration of event embedding (see above).

For each patient, this leads to a sequence of fixed-sized embedding vectors that encode both medical and time information at each (non-empty) timestamp $t \leq T^i$, as displayed in Step (4) of Figures 1 and 3. Now, this sequence of vectors is used as the input of a stack of layers, including recurrent layers (LSTM [33], bi-directional LSTM (bi-LSTM) [62], GRU [15]) or convolutional layers, in a sequence-to-one network architecture, since we want to output a single vector $x_i \in \mathbb{R}^d$ that encodes the full pathway of a patient before $T^i$. The results obtained with different types of layers are shown in Section 3.7 below. We use again dropout on the inputs and on the recurrent units to prevent overfitting. Finally, the output vector of the encoder is concatenated with an embedding vector of the age of the patient, leading to the final vector $x_i \in \mathbb{R}^d$ that encodes the full pathway of patient $i$ before $T^i$. This vector is the input of the ZiMM Decoder described in the next Section.

### 2.4 The ZiMM Decoder

The decoder uses the input vector $x_i \in \mathbb{R}^d$ of patient $i$ to construct the parameters of the ZiMM model, and the whole architecture is trained against the negative log-likelihood of the ZiMM model from Section 2.1 computed at the label vector $y_i = [y_{i,1}, \ldots, y_{i,B}]$ containing the blurry relapses. The parameters $\{\pi_b(x_i)\}_{b=1}^B$ and $\{p_{b,b'}(x_i)\}_{b,b' \in \{1, \ldots, B\}^2}$ are highly dependent and are time-ordered, so that a specific architecture is used to model these dependencies. The mixture probabilities are learned through a fully connected feed-forward layer (FFN) that use as input $x_i$, namely

$$\pi_0(x_i), \ldots, \pi_B(x_i) = \text{softmax}(\text{FFN}(x_i)), \quad (6)$$

and another recurrent layer (RNN) uses as well $x_i$ in order to produce hidden states given by

$$h_t = \text{RNN}(h_{t-1}, x_i) \quad (7)$$

for $t = 1, \ldots, B$. Then, we use different recurrent layers (RNN$_b$) in parallel for each value $n_i = b \in \{1, \ldots, B\}$ using

$$h^b_t = \text{RNN}_b(h^b_{t-1}, h_t) \quad (8)$$
for \( t = 1, \ldots, B \), since the multinomial distributions vary strongly conditionally on \( n_i = b \). Finally, a softmax activation is applied on \( h^b_t \) along \( t = 1, \ldots, B \) to produce the parameters \( p_{b,1}(x_i), \ldots, p_{b,B}(x_i) \). Details on the type of layers used and the tuning of hyperparameters is provided in Section 3 below.

### 2.5 Training

The full ZiMM Encoder-Decoder architecture is trained in an end-to-end fashion by minimizing the average negative-log likelihood over all patients \( i = 1, \ldots, n \):

\[
\ell_i(\Theta) = \log(\pi_0(x_i))1_{n_i=0} + \sum_{b=1}^{B} \left[ \log(\pi_b(x_i)) + \log\left( \frac{b!}{\Pi_{y=1}^{B} y_i,b'} \right) + \sum_{b'=1}^{B} y_{i,b'} \log(p_{b,b'}(x_i)) \right] 1_{n_i=b},
\]

where \( \Theta \) stands for the concatenation of all the trainable parameters involved in the layers described in Sections 2.3 and 2.4. The choices of optimizer, learning rate schedule and other hyperparameters are described in Section 3.

### 3 Application: prediction of post-surgical relapse of urinary problems

In this section, we apply the ZiMM ED architecture to predict the blurry relapse of urinary problems after a TURP surgery for patients with benign prostatic hyperplasia (see Section 3.2 below) using a cohort constructed from the French SNIIRAM database (see Section 3.1 below). Then, we explain in details the way the labels are built (Section 3.3), the different steps involved in the cohort construction (Section 3.4), the evaluation metric and implementation details (Section 3.5). Then, the remaining of this section describes the baselines and a comparison with ZiMM ED, followed by an ablation study.

#### 3.1 SNIIRAM: A non-clinical claims dataset

SNIIRAM (French national health insurance information system) contains health reimbursements claims of almost all French citizens since 2015 (more than 65 million) and has been used for health research on many topics, to cite but a few [61, 2, 31]. Since, in France, most health-cares are at least partially reimbursed by the administration, this database contains records corresponding to very various information going from hospital stays to drug purchases in city pharmacies, all coded with different systems [67]. Absence of clinical information and “forced” normalization makes this data highly complex and heterogeneous. A consequence is that it requires a lot of domain expertise about the way healthcare is reimbursed in France in order to prepare it as training data for machine learning algorithms. In this paper, we exploit medication, procedure and diagnosis codes only. Diagnoses are primarily coded with ICD-10, while procedures are coded with CCAM, a French medical classification of clinical procedures. Medications codes use the French pharmaceutical categories CIP13 [11] that we map to the international ATC (Anatomical Therapeutic Chemical Classification) classification system.

Let us stress that while this data is extremely rich and almost population-wide over France, it is not clinical data, but only claims data, clinical information being only latent in the codes appearing in reimbursement information. No vital bedside information, lab tests results or natural language notes from clinicians is available. An important amount of data preprocessing and domain expertise is required for
data extraction, in particular in order to clean existing inaccuracies in codes and timestamps. This is
achieved through our SCALPEL3 library [4], which is a separate topic of research that this paper builds
upon. Furthermore, the identification of a disease state (indication, concomitant disease or outcome) does
not rely only on a single source of information but on the convergence of information from different
sources. This might include, for instance, the presence of a chronic disease registration, hospital diagnoses,
tracer drugs, and procedures or lab tests [11], that are all observed in the data through claims.

We believe that these downsides are counter-balanced by the exhaustive nature of this data. It is
exhaustive both in terms of population (it includes nearly all the population living in France) and in terms
of healthcare events. Indeed, in France, almost all healthcare spendings are at least partially reimbursed
by the government, so almost each healthcare of each individual is associated to at least one event in the
database. This makes it a very rich and powerful database, with a relatively small bias, and many works
have already proved successful in improving population-level health [54, 61, 2, 31]. Though most EHR
studies are dealing with clinical EHR’s [23], it is clearly crucial to exploit such claims-only EHR, and
we hope that our work is a first step towards broader use, in healthcare and machine learning research
community, of non-clinical claims datasets.

Let us note that, since 2016, the access to this database, for public interest research, is possible
through the SNDS access pipeline [65]. It mainly offers access through the SAS software. An access
using classical open source big data and AI frameworks (e.g. R, Python, Spark) including the SCALPEL3
library will be possible, by the end of 2020, through the Health Data Hub [34], a 80m$-funded national
project which aims to be the national unique gateway to most French health data for operating public
interest research (operated by both public or private entities) on modern infrastructures.

3.2 Benign prostatic hyperplasia (BPH)

BPH is a common urological disease that affects aging men all over the world [64, 26]. It causes
urinary tract obstruction due to the unregulated growth of the prostate gland, causing lower urinary tract
symptoms (LUTS) [35]. The options for management of BPH include watchful waiting, pharmacotherapy,
transurethral resection of the prostate (TURP), and other minimally invasive surgical treatments (MISTs)
and open prostatectomy. Most studies consider TURP as the gold standard for surgical management
of BPH [32]. Patients suffering from prostatic hyperplasia regularly take drugs for urination problems.
Successful surgery should cure such problems so that patients should not need to continue their drugs for
urination problems. However, it is often observed that patients retake such drugs after surgery. In some
cases, it merely comes from a habit of taking routine drugs, but in many cases, it is related to a persisting
urination problem [47, 22].

An important problem, from a clinical point of view, is, therefore, to predict the outcome of such
a surgery, on a long-time period following it (18 months is considered here), to improve the decision
making of the clinicians, in particular, to help decide when surgery should be performed. Use of long-term
data after TURP surgery is very scarce in the literature, with only a few studies available [24]. A recent
analysis of 20 contemporary randomized clinical trials with a maximum follow-up of five years reports
that TURP resulted in a significant improvement of maximum flow-rate and quality of life [25, 44].
A second prostatic operation (re-TURP) has been reported at a constant annual rate of approximately
1-2%. A review analyzing 29 RCTs found a re-treatment rate of 2.6% after a mean follow-up of sixteen months [25]. In a large-scale study of 20,671 men, the overall re-treatment rates (i.e., either re-TURP, urethrotomy or bladder neck incision) were 5.8%, 12.3%, and 14.7%, respectively at one, five, and eight years follow-up. More specifically, the respective incidence of re-TURP was 2.9%, 5.8% and 7.4% [48]. However, urology guidelines highlight the lack of extended follow-up after the surgery and no clear evidence explaining re-treatment.

Building a model with the ability to predict the outcome of this surgery is of primary importance for improving population-level health, especially since prostatic problems are a common condition for aging men. In this section, we use ZiMM ED to train such a model using a cohort based on SNIIRAM. The outcome of this surgery is evaluated by the distribution of the blurry relapses, observed through the use of specific drugs related to urination problems.

3.3 Surgery identification and labels construction

The identification of a TURP surgery in SNIIRAM is made through some specific CCAM codes provided by clinicians. However, it happens that two TURP surgeries can occur in a pretty small amount of time. This is likely to correspond to a case where a surgery has clearly failed and that a second surgery is required quickly, which is not the type of complication we are interested in. If the amount of time between the two surgeries is small, it would be improper to say that it corresponds to a relapse. We thus define, following clinicians recommendations, a six week period after the first surgery, as a single surgery block. If another TURP surgery occurs inside the same block, we consider that it is part of the same medical act. The timestamp of the event corresponds to the last surgery within the block, and provides the value of $T_i$ for any patient $i$.

As far as the labels are concerned, a simple but efficient way to identify whether the urinary problems have not ceased or reappeared after a while is to see if the patient, at some point after the surgery, needs to take medications for these urination problems again. For that purpose, we use a list (provided by clinicians) of 136 CIP-13 drugs that are mainly related to urination problems. We choose to drive the prediction on a 18-months period after $T_i$. So, using the notation of Section 2.1, we chose the number of buckets $B = 18$ and bucket size of 30 days (540 days total).

3.4 Cohort construction and exploration

The construction of the cohort follows the flowchart illustrated in Figure 4. We start by selecting all SNIIRAM patients alive, with at least one medical event related to urination problem (observed through medication or surgery codes) between 2010/01/01 and 2015/12/31. This selection gathers a little bit more than 5 million patients. Then, we keep only male patients over 18 years old on 2014/12/31. This reduces the size of the cohort by roughly 1 million patients. Keeping only patients with a TURP surgery between

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1 We considered that a TURP surgery corresponds to the CCAM codes JGFA005, JGFA009, JGFA015, JDPE002 or JGNE003.
2 Following clinicians recommendations, we do not consider TURP surgeries that occur after the first surgery block, and consider only drugs prescriptions as blurry relapses. Consequently, we remove patients with repeated surgeries that are not part of the same block from the cohort construction, as explained in Section 3.4.
2010/01/01 and 2013/06/30 and with at least 2 events (whatever the types) occurring in the follow-up 18-months period leads to a cohort containing 231 747 patients.

As explained in Section 3.3, we compute surgery blocks (any surgery that occurs within a six week period after the first surgery is in the same block), that are considered as a single event of surgery. We remove all the patients that have another surgery out of this first block. We choose a follow-up period of 540 days, corresponding to 18 buckets of 30 days, so that the follow-up period is of the order of 18 months. Indeed, clinicians expect that such a relapse should occur not long after the first year following surgery. As shown in Figure 4, the final cohort has 138 976 patients.

Patients covered by French universal health insurance with at least one urination problem event (surgery or medication) between 2010/01/01 and 2014/12/31 ($n = 5\,136\,308$)

Exported cohort ($n = 4\,091\,074$)

Patients with surgery event between 2010/01/01 and 2013/06/30 ($n = 231\,747$)

Analysed cohort ($n = 138\,976$)

Excluded ($n = 1\,045\,234$):
1. Age < 18; and > 110 if death date is missing ($n = 39\,300$)
2. Women ($n = 1\,005\,933$)
3. Wrong hospital stays dates ($n = 1$)

Excluded ($n = 92\,771$):
1. More than 1 surgery bloc ($n = 9\,973$)
2. Death before $T + 18$ months ($n = 14\,617$)
3. Less than 500 observation days in the 18-months follow-up period ($n = 68\,181$)

Figure 4: Flowchart leading to the final cohort considered in the experiments. The number $n$ stands for the number of patients remaining at each stage of preprocessing.

The numbers of medical codes, namely medications, procedures and diagnoses observed in the final cohort, are displayed in Table 1. We count the number of unique codes, the number of codes observed at least 50 times and the number of codes remaining when keeping only the ones prior to $T^i$ for each patient $i$. The statistics reported in Table 1 below and in all figures from this section use the raw coding scheme (that use CIP-13 for drugs), in order to better describe the statistics of the data, while for training ZiMM ED we replace CIP-13 by ATC (for drugs) since, as illustrated in Section 3.7 below, it allows to reduce the vocabulary size of drugs while keeping the same predictive performance. As for subsequent encoding, we consider only tokens that occur at least 50 times in the training dataset, so that the encoder will see a total of 9 271 medical codes.

On the left-hand side of Figure 5, we show, for this final cohort, the distribution of $n_{i_{12}}$, namely the number of patients with a given total number of drugs prescriptions (related to urinary problems) during
|          | Medications | Medical procedures | Diagnoses | Total |
|----------|-------------|--------------------|-----------|-------|
| #Unique  | 12 785      | 9 578              | 5 885     | 28 248|
| #Unique before $T^i$ | 10 664      | 7 460              | 4 563     | 22 687|
| #Unique seen $\geq$ 50 times before $T^i$ | 6 713        | 1 155              | 1 403     | 9 271 |

Table 1: Number of medical codes observed and remaining when applying the filters used in the cohort construction. Bold values correspond to the vocabulary sizes of the medical codes used when training the ZiMM ED architecture.

the follow-up 18 months period. Namely, the $y$-axis of Figure 5 (left-hand side) displays $\#\{i : n_i = b\}$ where $b = 1, \ldots, B$ is given by the $x$-axis. We observe a sharp decrease for $b = 1, \ldots, 5$ from the mode $b = 1$, which corresponds to patients who continue to buy their drugs and just keep doing it (for several months) after their surgery. However, we observe a second mode around $b = 15$ which certainly corresponds to the relapse we are interested in. This plot shows that the considered problem is in a “weak signal” setting, since the second mode is very small compared to the first (at $b = 1$), and that the prediction of this relapse requires a dedicated methodology indeed. This observation is corroborated by the right-hand side of Figure 5, which displays the number of patients having at least one drug prescription (related to urinary problems) on a specified bucket, namely the $y$-axis displays $\#\{i, y_{i,b} \geq 1\}$ as a function of $b$ given by the $x$-axis. Once again, the function is decreasing from the mode $b = 1$ and plateaus between $b = 7$ and $b = 18$, because of patients who continue to purchase the drug after the surgery. We do not observe a second mode as we did in the left-hand side, since the relapse we are interested in is blurry: the drug purchases that define this blurry relapse are heterogeneously distributed among patients and end up flattened out when aggregated in the display of the right-hand side. This motivates, once again, the dedicated methodology proposed by the ZiMM model.

![Figure 5: Left-hand side. Number of patients as a function of the overall number of drugs prescriptions related to urinary problems during the follow-up 18 months period. The $y$-axis displays $\#\{i : n_i = b\}$ where $b = 1, \ldots, B$ is given by the $x$-axis. We observe a sharp decrease from the mode $b = 1$ followed by a second weak mode around $b = 15$. The number of patients with $n_i = 0$ is equal to 84 328 and is not displayed for readability of the plot. Right-hand side. Number of patients having at least one drug prescription (related to urinary problems) in each time bucket: the $y$-axis displays $\#\{i, y_{i,b} \geq 1\}$ as a function of $b$ given by the $x$-axis. We observe a sharp decrease from the mode $b = 1$ and a plateau between $b = 7$ and $b = 18$, since patients often continue to purchase the drug after the surgery.](image-url)
for a long time before the surgery, while other patients have surgery sooner. On the right-hand side of Figure 6, we show the distribution of the durations of all hospitalizations before $T^i$. We observe that most hospitalizations are one-day short, and that the distribution is heavy-tailed, since these hospitalizations can be related to a large set of possible health problems, leading to very heterogeneous durations. Let us stress that we are considering all hospitalizations that occur before $T^i$, and not only the ones related to a surgery or a urological problem, since all the health events of a patient $i$ are kept before $T^i$.

Figure 6: Left-hand side. Distribution of the observation period of each patient before $T^i$. The strong variability of the observation period displayed here is related to the variability of the medical practice itself: some patients are treated for urological problems with drugs for a long time before the surgery, while other patients have surgery sooner. Right-hand side. Distribution of the durations of all hospitalizations before $T^i$. Most hospitalizations are one-day short, and the distribution is heavy-tailed, since these hospitalizations can be related to a large set of possible health problems.

In the left-hand side of Figure 7, we display the distribution of the number of unique events per patient. We observe that the bulk of the distribution is between 5 and 200 unique events, with an average of 10 unique events. The right-hand side of Figure 7 shows the number of days with at least one medical code in the history of patients before $T^i$. We observe that most patients have more than 10 days in their history with medical codes. Both displays from Figure 7 show that the health pathways of most patients before $T^i$ contain enough variability when assessed by the number of distinct codes, and the number of distinct days with at least one medical code, corresponding to a sufficient amount of variability to carry information for the prediction of the blurry relapse after $T^i$.

Figure 7: Left-hand side. Distribution of the number of events per patient before $T^i$. The bulk of the distribution is between 5 and 200 unique events, with an average of 117 events per patient. Right-hand side. Distribution of the number of days with at least one medical code in the history of patients before $T^i$. We observe that most patients have more than 10 days in their history with medical codes. Both figures show that most patients have a significant amount of medical code variability and time variability before $T^i$. 

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3.5 Implementation and evaluation metrics

Our models are implemented with TensorFlow 2 [1]. The ZiMM ED architecture is open-source in GitHub\(^3\). All the models are trained on a machine equipped with 4 GeForce GTX 1080Ti GPUs and another machine with 3 Tesla V100 GPUs. All the models are trained with the Nadam optimizer [36, 30] with learning rate 0.001. The hyper-parameters of all the models are selected through an extensive random grid search, whereas in order to reduce computation time, some hyper-parameters are fixed “by hand”. An ablation study showing the sensitivity to hyper-parameters is provided in Section 3.7 below.

**Metrics: mean-AP, AUC-PR and AUC-ROC.** In order to evaluate the quality of the prediction of \(y_i = [y_{i,1}, \ldots, y_{i,B}]\) (when using ZiMM ED architecture or any other benchmark models) we use *mean-AP*, defined as the average of the area under the precision-recall curve (AUC-PR), namely we compute the average over the buckets \(b = 1, \ldots, B\) of the AUC-PR for each bucket \(b\), i.e., the AUC-PR of the prediction of \(y_{i,b}\). Moreover, we report also the AUC-PR and the AUC-ROC (area under the ROC curve) for the binary classification problem \(n_i > 0\) against \(n_i = 0\).

**Train, validation and test sets.** All our experiments use the same random data splitting into 70% of patients for training, 15% of patients for validation and 15% for testing. We checked the stationarity across the three splits of labels distribution and the main drugs, diagnoses and medical procedures. All the models are trained on a train set and we tune hyper-parameters using the validation set, while the test set is only used for the final evaluation. We report performances on both the validation and the test sets.

3.6 Baselines

The prediction performances of ZiMM-ED is compared with several baselines, involving several featuring strategies and different predictive models, both for binary prediction (\(n_i > 0\) versus \(n_i = 0\)) using the AUC-PR and AUC-ROC metrics and for the prediction of \(y_i = [y_{i,1}, \ldots, y_{i,B}]\) (the mean-AP score defined in Section 3.5). Results are reported in Table 2 below, where we observe that ZiMM ED improves all the considered baselines, in particular for the prediction of \(y_i\), since the ZiMM model is dedicated to this task. We consider the following baseline models, more details are provided below:

- **LR\(l_2\):** Logistic regression with \(\ell_2\) penalization using the scikit-learn library [57];
- **GBDT:** Gradient boosting using XGBoost [13];
- **MLP:** Multilayer Perceptron model with 128 hidden units;
- **Word2vec:** we first train embeddings of all medical codes following [52], and use these pre-trained embeddings in a MLP with 128 units;
- **LSTM:** a single embedding layer and one forward LSTM layer with 128 hidden units;
- **Patient2Vec** [72].

We evaluated them using 3 types of input features described below.

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\(^3\)https://github.com/stephanegiiffas/zimm.git
**Static features (SF).** This featuring is used for LR12, GBDT and MLP models. Inputs correspond to aggregated counts of grouped medical codes over the entire observation period of a patient. Number of occurrences of each code is then multiplied by the corresponding one-hot encoding. Hence, the input is a \(N\)-dimensional vector representing a patient’s medical history. One logistic regression (LR12) and one gradient boosting decision tree (GBDT) is trained for each bucket \(b\). This input is also used for the multi-layer perceptron (MLP).

**Dynamic features (DF).** This featuring is used for LR12, GBDT and MLP models. We split the sequence into subsequences of 60 days, on which we compute the same features as with SF, but within each interval, in order to incorporate longitudinal information into LR12, GBDT and MLP.

**Irregularly-spaced sequence (ISS).** This featuring is used for Word2vec, LSTM and Patient2Vec models. In this case, we consider the original patient sequence. For both the Word2vec model and the LSTM one, the events are just gathered in a sequence in which the time-stamps or duration of the events are not used. In the Word2vec case, the codes embeddings are trained using Word2vec [52] and prediction is performed with a MLP with 128 hidden nodes. In the LSTM case, an embedding layer and one forward LSTM layer with 128 hidden units are trained in an end-to-end fashion and is applied to the sequence. The Patient2vec uses the same input features as the ZIMM ED architecture (i.e., sequence, including time-stamps and duration of events). It has been only used for the binary classification problem (i.e., \(n_i > 0\) versus \(n_i = 0\)).

The predictive performance of all benchmarks and of ZiMM ED are presented in Table 2, where we report mean-AP scores on the test set, as well as AUC-ROC and AUC-PR scores for the binary classification problem (see Section 3.5 for definition of these scores). According to this table, GBDT-based models on dynamic features (GBDT-DF) performs the best among all benchmark models, however the ZIMM ED architecture outperforms all the benchmark models both for the multi-output \(y_i\) prediction and for the binary prediction.

| Model         | mean-AP | AUC-ROC | AUC-PR |
|---------------|---------|---------|--------|
| LR12-SF       | 0.19    | 0.64    | 0.50   |
| GBDT-SF       | 0.24    | 0.67    | 0.56   |
| MLP-SF        | 0.18    | 0.64    | 0.49   |
| LR12-DF       | 0.21    | 0.65    | 0.53   |
| GBDT-DF       | 0.25    | 0.68    | 0.57   |
| MLP-DF        | 0.19    | 0.65    | 0.50   |
| Word2vec-ISS  | 0.20    | 0.65    | 0.53   |
| LSTM-ISS      | 0.21    | 0.67    | 0.54   |
| Patient2Vec   | -       | 0.68    | 0.55   |
| ZiMM ED (best model) | **0.306** | **0.701** | **0.619** |

Table 2: Predictive performances (on test data) of benchmark models and ZiMM ED architecture. ZiMM ED appears to perform the best among all models both for multi-output and binary prediction. mean-AP stands for average of the area under the precision-recall curve (AUC-PR) over buckets \(b = 1, \ldots, B\).
Table 3: $p$-values of pairwise Mann-Whitney U statistical tests for the null assumptions that the mean-AP scores of ZiMM ED and each baseline are equal. All tests strongly reject this hypothesis. Note that zero $p$-values correspond to situations where the mean-AP score of the baseline was much smaller than the one of ZiMM ED for each bootstrap sample.

Additionally, following suggestions in [49, 6], we also report, in Figure 8 below, boxplots for the mean-AP score computed on 1000 bootstrapped samples based on the test set, together with, in Table 3, the $p$-values of pairwise Mann-Whitney U statistical tests for the null assumption that the mean-AP score of ZiMM ED and each baseline are equal. We observe, both in Figure 8 and Table 3, that ZiMM ED significantly improve the mean-AP scores of all the considered baselines: all the tests strongly reject the null hypothesis (the largest $p$-value is $< 10^{-6}$) and the boxplot of ZiMM ED exhibits a distribution which is significantly shifted towards higher mean-AP scores.

Figure 8: Boxplots of the mean-AP scores computed on 1000 bootstrapped samples based on the test set of all baseline models and ZiMM ED. The boxplot of ZiMM ED exhibits a distribution which is significantly shifted towards higher mean-AP scores.

3.7 Ablation study: performances of our model and variations around it

The performance reported in Table 2 for the ZiMM ED architecture relies on a careful tuning of several hyper-parameters. This corresponds to the so-called ZiMM ED default architecture, where the hyperparameters used are described in Table 4 below.

The hyper-parameters described in Table 4 have been selected using an extensive random search for the best mean-AP metric on the validation set. In Figure 9 below, we illustrate the value of this metric ($y$-axis) for some models that were evaluated during the random search. Each point corresponds to a single model with fixed hyper-parameters, and the set of models is the same on all four displays of Figure 9. In each of these displays we “align” all the models that share a specific hyper-parameter, namely, respectively from left to right: the number of heads used in the self-attention layer, the number of hidden units used in the recurrent layer of the ZiMM Encoder, the number of stacked recurrent layers in ZiMM Encoder and finally the number of hidden units used in the ZiMM Decoder. The red point on
| Hyper-parameter                                                                 | Value   |
|--------------------------------------------------------------------------------|---------|
| Maximum #days observed in patients’ sequence before $T_i$                      | 50      |
| Maximum #medical events within the same day                                    | 24      |
| Vocabulary size for medications coded with ATC                                | 1036    |
| Vocabulary size for diagnoses coded with ICD-10                                | 1391    |
| Vocabulary size for medical procedures coded with CCAM                         | 1146    |
| Embedding dimension of medical codes                                           | 64      |
| L2 penalization rate for medical codes embedding                               | 0.005   |
| Gaussian dropout rate                                                          | 0.3     |
| Embedding dimension for time horizon and hospitalization duration              | 4       |
| L2 penalization rate for time embedding                                         | 0.01    |
| Batch normalization epsilon                                                    | 1e-06   |
| Aggregation mode                                                               | self-attention |
| Number of heads                                                                | 3       |
| Weight drop-connect rate                                                        | 0.3     |
| Dropout rate                                                                   | 0.2     |
| L2 penalization rate for time embedding                                         | 0.01    |
| Recurrent layer type                                                            | LSTM    |
| #hidden units per layer                                                         | 64      |
| #layers                                                                        | 1       |
| Dropout rate                                                                   | 0.3     |
| Recurrent dropout rate                                                          | 0.2     |
| Recurrent layer type                                                            | GRU     |
| #hidden units per layer                                                         | 32      |
| #RNN layers used in parallel for each value $n_i = b$                          | 1       |
| #common RNN layers                                                             | 1       |
| Gaussian dropout rate                                                           | 0.3     |
| Recurrent dropout rate                                                          | 0.2     |
| Optimizer type                                                                 | Nadam   |
| Learning rate                                                                  | 0.001   |
| Batch size                                                                     | 256     |

Table 4: ZiMM-ED default parameters.

all four Figures corresponds to the best model overall, that leads to the ZiMM ED default architecture described in Table 4. Finally, note that the random search included many other hyper-parameters, such as several learning-rate scheduling strategies, dropout rates, types of dropout regularization including input, output, embedding, cell-state and multiplicative Gaussian, that we do not report for the sake of conciseness.

The remaining of this section proposes an ablation study: we modify some hyper-parameters or change some components of ZiMM ED default, and report the impacts on performances in Table 5 below, where the first line reports the performances of ZiMM ED default. A discussion around these ablations is given below.
Figure 9: Validation mean-AP ($y$-axis) of some models evaluated during random search, where each point corresponds to a single model. The set of models is the same on all four displays and the best one (ZiMM ED default) is indicated by a red point. Each display aligns all the models that share a specific hyper-parameter, being, respectively from left to right, the number of heads used in the self-attention layer, the number of hidden units used in the recurrent layer of the ZiMM Encoder, the number of stacked recurrent layers in ZiMM Encoder and finally the number of hidden units used in the ZiMM Decoder.

Data preprocessing. The first part (PP) of Table 5 shows results obtained with variations of the maximum sequence lengths and the classification system used for drugs encoding. The ZiMM ED default uses ATC encoding instead of the raw CIP-13 encoding, which allows to reduce the vocabulary size from 10 664 for CIP-13 to 1 105 for ATC. As observed in Table 5, using CIP-13 (which is a much larger vocabulary for drugs) actually hurts strongly the performances. We observe also that considering longer sequences (100 days) instead of what we do in ZiMM ED default (50 days) deteriorates the performances as well.

Embedding of medical codes and time. Variations around the way embeddings are produced for medical codes and time are reported in the second part (Embedding) of Table 5. We observe that increasing the embedding dimensions does not offer performance increase (using $E_{\text{dim}} = 128$ instead of the default $E_{\text{dim}} = 64$), as well as using the same embedding space for all medical codes. In the ZiMM ED default model, tokenized durations and distance to $T_i$ (see Section 2.3 above) are embedded with trainable embedding vectors. We observe in Table 5 that without these time embeddings, the model performance deteriorates. Also, we find that using trigonometric functions for “positional encoding” as
proposed in [68] instead of learned embeddings does not help in our setting.

**Embeddings aggregation.** Results obtained through different embeddings aggregation techniques are presented in the third part (Aggregation) of Table 5. We applied penalization and dropout on attention weights as proposed in [42], as well as DropConnect [69] for regularizing weights in large fully-connected layers. As explained in [68], multi-head attention allows the model to jointly attend information from different representation subspaces at different positions, however in our setting, increasing SA $N_{\text{heads}}$ (self-attention number of heads) in the self-attention layer does not clearly improve the performance. We tested $N_{\text{heads}} = 8$, as reported in [45], but the best performance in our case was achieved with $N_{\text{heads}} = 3$.

**ZiMM Encoder.** We report in the fourth part (Encoder) of Table 5 the results of extensive experiments performed in order to identify the best combination of hyper-parameters for both recurrent and convolutional layers (CNN): number of hidden units, types of recurrent layers and different dropout rates used in the recurrent layers. Only the best results for each type of layer are reported in Table 5. We observe that GRU layers perform generally as well as LSTM layers. Increasing the number of hidden units in recurrent layers to $N_{\text{units}} = 128$ instead of the default choice $N_{\text{units}} = 64$ does not significantly improve performance, as well as using two stacked LSTM layers $N_{\text{layers}} = 2$ instead of the default $N_{\text{layers}} = 1$. We observed also that in our setting, CNN layers badly under-perform compared to the other types of layers. Finally, we tested “Multi-head transformer” which corresponds to the transformer encoder [68] which takes as input all patient sequence of events. Clinical codes are embedded in the same way as for ZiMM model, which is then passed to the encoder block with a 4-head self-attention mechanism through the entire patient sequence.

**ZiMM Decoder.** Finally, we consider some variations around the ZiMM Decoder, and report the results in the fifth part (Decoder) of Table 5. While the ZiMM Decoder described in Section 2.4 uses a single feed-forward layer to predict the mixture probabilities, a GRU layer to learn hidden states and GRU layers to predict the parameters of each multinomial distributions, one could use instead fully-connected layers to predict all the parameters of the ZiMM distribution, or alternatively simple RNN layers or LSTM layers. The fully-connected layers deteriorates the most the performance, while replacing the GRU layers by RNN or LSTM layers only deteriorates it mildly. We also report the performance obtained with two stacked GRU $N_{\text{layers}} = 2$ instead of the default one $N_{\text{layers}} = 1$ to produce the hidden states (see Equation (7)) and with a smaller hidden size $N_{\text{units}} = 18$ instead of $N_{\text{units}} = 32$.

### 3.8 Visualization of the embeddings produced for diagnoses and drugs codes

We explore the embeddings produced for diagnoses and drugs as a by-product of the ZiMM ED architecture (see Section 2.3 about the embeddings of medical codes). We use UMAP [50] in order to reduce the dimension from 64 to 2. The resulted projections are shown in Figure 10. The UMAP algorithm requires four hyper-parameters: the number of neighbors to consider when approximating the local metric (n-neighbors), the desired separation between close points in the embedding space (min-dist), the number of training epochs (n-epochs) and finally the projection dimension ($d$). We fixed $d=2$ and
| Architecture modification | # params $\times 10^5$ | val set mean-AP | mean-AP AUC-ROC AUC-PR |
|---------------------------|------------------------|----------------|------------------------|
| **ZiMM ED default**      | 3.5                    | 0.292          | 0.304 0.704 0.619      |
| PPs                      |                        |                |                        |
| CIP-13 drugs encoding    | 7.1                    | 0.279          | 0.286 0.690 0.604      |
| 100-days sequence        | 3.5                    | 0.291          | 0.300 0.701 0.616      |
| **Embedding**            |                        |                |                        |
| Common embedding space   | 13.9                   | 0.292          | 0.298 0.698 0.613      |
| Without $\Delta t$ embedding | 3.5                | 0.293          | 0.300 0.702 0.617      |
| Positional encoding      | 3.5                    | 0.292          | 0.302 0.702 0.620      |
| $E_{dim} = 128$          | 7.4                    | 0.291          | 0.300 0.701 0.615      |
| **Aggregation**          |                        |                |                        |
| SA $N_{heads} = 1$       | 3.2                    | 0.293          | 0.303 0.701 0.619      |
| SA $N_{heads} = 5$       | 3.7                    | 0.290          | 0.300 0.704 0.620      |
| SA $N_{heads} = 8$       | 4.1                    | 0.292          | 0.300 0.701 0.616      |
| mean                     | 3.1                    | 0.287          | 0.297 0.696 0.612      |
| **Encoder**              |                        |                |                        |
| $N_{units} = 128$        | 4.6                    | 0.293          | 0.301 0.701 0.618      |
| $N_{layers} = 2$         | 3.8                    | 0.290          | 0.306 0.701 0.619      |
| bi-LSTM                  | 4.2                    | 0.290          | 0.300 0.700 0.617      |
| GRU                      | 3.3                    | 0.294          | 0.300 0.702 0.614      |
| Conv1D                   | 3.2                    | 0.231          | 0.234 0.649 0.536      |
| Multi-head transformer   | 6.2                    | 0.279          | 0.287 0.694 0.607      |
| **Decoder**              |                        |                |                        |
| FC layer                 | 3.3                    | 0.290          | 0.300 0.691 0.617      |
| LSTM                     | 3.5                    | 0.291          | 0.301 0.703 0.619      |
| basic RNN                | 3.4                    | 0.292          | 0.301 0.701 0.619      |
| $N_{units} = 18$         | 3.4                    | 0.292          | 0.298 0.702 0.619      |
| $N_{layers} = 2$         | 3.5                    | 0.288          | 0.297 0.699 0.615      |

Table 5: Performances of some variations around the ZiMM ED default architecture, for which all hyper-parameters are given in Table 4 above. *PP stands for “preprocessing”. SA $N_{heads}$ stands for number of self-attention heads; bi-LSTM stands for bidirectional LSTM; mean-AP stands for average of the area under the precision-recall curve (AUC-PR) over the buckets $b = 1, \ldots, B$; AUC-ROC stands for area under the Receiver operating characteristics curve.

$n$-epochs = 500. ICD10 disease codes are mapped in PheWAS phenotypes [29] and drugs codes are mapped to the first level of the ATC classification (main anatomical group consisting of a single letter), leading to the color scheme used in Figure 10.

On the left-hand side of Figure 10, we can observe the UMAP projections of the embedding vectors of each ICD10 code, colored by the corresponding PheWAS phenotype. The hyper-parameters used here for UMAP are $n$-neighbors=20 and min-dist=0.1. Many points share the same color since many ICD10 codes have the same PheWAS. We observe here that diagnoses that are known to co-occur and belong to the same clinical groups are projected close to each other. The red-orange-highlighted points are the phenotypes the most related to the urogenital system such as **Hyperplasia of prostate**, **Retention of urine**, **Benign neoplasm of male genital organs**, and **Cancer of prostate**. We can also observe a visually strong association of these phenotypes with **Essential hypertension**. This is confirmed by [51], where the association of hypertension and symptoms of benign prostatic hyperplasia is well-studied and understood. Other clinical concepts that are projected close to each other are **Hearing loss** and **Chemotherapy**, **Urinary**...
Hyperplasia of prostate
Essential hypertension
Retention of urine
Benign neoplasm of male genital organs
Cancer of prostate
Atrial fibrillation and flutter
Hypercholesterolemia
Hematuria
Effects radiation NOS
Type 2 diabetes
Urinary tract infection
Hearing loss
Chemotherapy
Renal dialysis

Figure 10: UMAP projections of the embeddings of medical codes. **Left-hand side.** Each point is the projection of the embedding vector of an ICD code, colored with its corresponding PheWAS phenotypes. Note that many points share the same color since many ICD10 codes have the same PheWAS. The legend provides only the most common diagnoses. **Right-hand side.** UMAP projections of the drugs embeddings, colored by the first level of the ATC classification (one letter).

tract infection and Hematuria (blue-highlighted points), which is also confirmed by [66, 41], where it is shown that renal dialysis is associated with a higher risk of cancer.

On the right-hand side of Figure 10, we observe the UMAP projections of drugs embeddings, colored by the first level of the ATC classification (first letter). The UMAP hyper-parameters used here are $n$-neighbors=50 and min-dist=0.2. The dark blue points correspond to Various ATC class (V), that mainly correspond to contrast agents for MRI. We observe that these points do not form any cluster but are dispersed over the whole space. This can be explained by the fact that the considered cohort contain only men with BPH (see Section 3.4), that often have an MRI of the prostate. Furthermore, we observe that the drugs from the class Genito-urinary system and sex hormones (G), that include drugs for urination problems, form one cluster with other age-related medications from the class Nervous system (N), namely analgesics, anti-thrombotics, psycholeptics, psychotropic, and anti-parkinsonian drugs.

4 Conclusion and future works

In this work, we propose ZiMM Encoder Decoder (ZiMM ED), an end-to-end deep learning model trained against the negative log-likelihood of the new ZiMM (Zero-inflated Mixture of Multinomial) model, for the modeling of long-term and blurry relapses. This deep learning model is trained on a cohort based on a large electronic health record database, that contains only claims and no clinical data, from the whole French population.

We show that our model improves the performances of a large number of baselines, including the state-of-the-art, for the considered predictive task. ZiMM ED allows to represent the full health pathways of patients, using all available information, with minimal preprocessing. Therefore, this end-to-end architecture can be used for other tasks as well, through transfer learning or fine tuning of the model. Future works will consider a multi-task version of ZiMM ED (several types of relapses, or other types of events), and other improvements using alternative architectures for attention modeling [37, 27, 38, 70].
The considered dataset and the introduced methodology was developed for non-clinical claims data, which is fundamentally different from clinical datasets. A future effort will be done with the aim of adding a clinical and genomic data to the cohort. In this case a lot of additions analysis can be done ([12]).

This work addresses the problem of predicting the blurry relapses of the TURP surgery, which is the first step towards an evidence-based approach using machine-learning to help the clinical decision. The next step is to exploit these predictions to help to decide the timing of the surgery: given the current health pathway of the patient, what is his probability of a relapse, so that it can help the clinician to decide when to perform the surgery, or to consider alternative treatments.

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