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ABSTRACT

Computational prediction of in-hospital mortality in the setting of an intensive care unit can help clinical practitioners to guide care and make early decisions for interventions. As clinical data are complex and varied in their structure and components, continued innovation of modelling strategies is required to identify architectures that can best model outcomes. In this work, we trained a Heterogeneous Graph Model (HGM) on electronic health record (EHR) data and used the resulting embedding vector as additional information added to a Convolutional Neural Network (CNN) model for predicting in-hospital mortality. We show that the additional information provided by including time as a vector in the embedding captured the relationships between medical concepts, lab tests, and diagnoses, which enhanced predictive performance. We found that adding HGM to a CNN model increased the mortality prediction accuracy up to 4%. This framework served as a foundation for future experiments involving different EHR data types on important healthcare prediction tasks.

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1. INTRODUCTION

Timely prediction of in-hospital mortality within intensive care units (ICU) is beneficial [1, 2] for practitioners to tailor care and allow for earlier interventions to prevent deterioration [3, 4]. Electronic health record (EHR) data consist of information relating to patient encounters with a health system, such as disease diagnoses, vital signs, and medications, among others [5, 6] which are often used for machine learning (ML) predictions for different tasks in the biomedical domain including mortality prediction [7, 8, 9]. The inherent complexity of EHR data often require advanced modeling frameworks to gain robust performance for these tasks. A common modeling approach for EHR research is to use a 2-dimensional convolutional neural networks (CNN) with one dimension as time and the other as clinical features [10, 11, 12]. In healthcare-related CNN models, various medical features are normally concatenated to be directly used as inputs and create embeddings [13, 14, 15]. This form of feature representation can be powerful, but disregards the graphical structure and interconnectivity between medical concepts [16, 17] which can affect the CNN performance especially since EHR data are often sparse due to missingness [10].

In this work, we proposed a Heterogeneous Graph Model (HGM) to create a patient embedding vector, which better accounts for missingness in data for training a CNN model. The HGM model captures the relationships between different medical concept types (e.g., diagnoses and lab tests) due to its graphical structure. This relational representation facilitates capturing more complex patient patterns and encoding similarities.

2. METHODOLOGY

2.1 Data Set

We conducted our experiments on de-identified EHR data from MIMIC-III [18]. This data set contains various clinical data relating to patient admission to ICU, such as disease diagnoses in the form of International Classification of Diseases (ICD)-9 codes, and lab test results as detailed in Supplementary Materials. We collected data for 5,956 patients, extracting lab tests every hour from admission. There are a total of 409 unique lab tests and 3,387 unique disease diagnoses observed. The diagnoses were obtained as ICD-9 codes and they were represented using one-hot encoding where one represents patients with disease and zero indicates those without. We binned the lab test events into 6, 12, 24, and 48 hours prior to patient death or discharge from ICU. From these data, we performed mortality predictions that are 10-fold, cross validated.

2.2 Convolutional Neural Network Model

Convolutional neural networks (CNNs) are often used, and perform well, on image processing tasks [19] due to their inherent feature extraction and abstraction ability, which increase the accuracy for classification tasks. There are also studies that have demonstrated encouraging successes in using CNN for EHR analyses. In this work, we used a standard CNN model as the baseline.

Since CNNs typically require two dimensional inputs, we treated time as the horizontal dimension and medical events as the vertical dimension. For the time dimension, we recorded every event with one-hour binned increments with respect to the patient death or discharge time. In this model, the vertical dimension was constructed by concatenating two medical event vectors: lab tests and diagnoses. Every entry of the lab test vector recorded the value of a specific lab test by hour, and we pre-processed the lab test by considering the values between 0.5 and 99.5 percentile to remove any inaccurate measurement. We then normalized the data by calculating the standard score (z-score). We imputed missing lab values with zeros. For the diagnosis vector, the *i*-th entry is 1 if the *i*-th diagnosis is observed; otherwise 0. We treated mortality prediction as a binary classification, for which we used a softmax layer with two dimensions and cross-entropy for loss.

2.3 Heterogeneous Graph Model

The features used in baseline CNN model are essentially raw data concatenated together, which do not consider the relationships between medical concepts. We used an HGM to capture these inherent relationships by creating three different types of nodes: patient, lab test, and diagnosis. These different types of nodes are connected by two relation types: tested and diagnosed. These could be represented with two triples:

Patient lab: {patient, tested, lab} Patient diagnosis: {patient, diagnosed, diagnosis}

The testing relationship shows whether a specific lab test was given to a patient at a specific time, and the diagnosed relationship shows whether a patient was diagnosed with a disease.

To represent the lab test and diagnosis node types, we used multi-hot encoding vector: $X_i \in \{0,1\}^{409}$ and $X_d \in \{0,1\}^{3387}$, and the *i*-th entry with the value of 1 indicating whether a specific lab test was performed or a specific diagnosis was given.

2.3.1 Node Embeddings

For capturing the relations between different medical events related to a patient, we first utilized the TransE model to project different types of nodes into the same latent space, and then classified those nodes that were connected as a similar group and the disconnected nodes as a dissimilar group.

The TransE model uses a set of 1) projection matrices and 2) relation vectors. After initialization, projections and translations are optimized end-to-end. Heterogeneous nodes X_{pr} , X_{lr} and X_d are projected into a shared latent space with trainable projection matrices W_{pr} , W_{lr} and W_d using the nonlinear mappings with Equation (1):

$$c_{p} = \sigma \left(W_{p} \cdot X_{p} \right)$$

$$c_{i}^{*} = \sigma \left(W_{i} \cdot X_{i} \right)$$

$$c_{d}^{*} = \sigma \left(W_{d} \cdot X_{d} \right)$$
(1)

where σ is a non-linear activation function and c_p, c_i^* , and c_d^* are the latent representations of each type of node. Despite the fact that the EHR uses different dimensions for different data types X_p, X_p and X_d , all node types are projected into the same latent space. Then we applied translation operations to link these different types of nodes with Equation (2):

$$c_i = c_i^* - r_{ip}$$

$$c_d = c_d^* - r_{dp}$$
(2)

where r_{ip} and r_{dp} are the relation vectors connecting patients to lab tests and diagnoses, respectively. c_i and c_d are the semantically translated projection representation into the same latent space of patient embedding c_p .

2.3.2 Optimization Model

For training the HGM, we applied a skip-gram optimization model, which increases the proximity between embedding points whose corresponding graph nodes are often connected after the projection and translation operations (Equation (3)):

$$\max \sum_{u \in V} \sum_{t \in T_{V}} logPr(N_{t}(u) \mid u)$$
(3)

where $N_t(u)$ are the neighborhood vertices of center node u, and $t \in T_v$ is the node type. Here, we learned the node embeddings by maximizing the probability of correctly predicting the patient node's associated lab tests and diagnoses. The prediction probability is modeled as a softmax function with Equation (4):

$$Pr(c_t \mid f(u)) = \frac{e^{\bar{c}_t \cdot \bar{u}}}{Z_u}$$
(4)

where \vec{u} is the latent representation of patient u, \vec{c}_t is the latent representation of lab and diagnosis neighbors of node u, and $\vec{c}_t \cdot \vec{u}$ is the inner product of the two embedding vectors representing their similarity. Z_u is the normalization term $Z_u = \sum_{v \in V} e^{\vec{v}_t \cdot \vec{u}}$ that is a sum over all vertices V, each of which is represented as \vec{v}_t including all node types. Therefore, Equation (3) is simplified to Equation (5):

$$\mathcal{L}_{s} = -\sum_{t \in T} \sum_{u \in V} \left[\sum_{c_{t} \in N_{t}(u)} \overrightarrow{c_{t}} \cdot \overrightarrow{u} - log Z_{u} \right]$$
(5)

Numerical computation of Z_u is intractable for large-scale graphs. So we adopted a negative sampling strategy to approximate the normalization factor. We eventually used the following optimization function (Equation (6)):

$$\mathcal{L}_{s} = -\sum_{t \in T} \sum_{u \in V} \left[\sum_{c_{t} \in N_{t}(u)} log\sigma(\vec{c_{t}} \cdot \vec{u}) + \sum_{j=1}^{K} E_{c_{j} - P_{v}(c_{j})} log\sigma(-\vec{c_{j}} \cdot \vec{u}) \right]$$
(6)

where σ (*x*) is the sigmoid function, which operates on the dot product between (c_{i^*u}) and $\sigma(x) = \frac{1}{1 + \exp(-x)}$, and \mathbb{K} is the number of negative samples. $P_v(c_i)$ is the negative sampling distribution.

For training HGM, we performed heterogeneous neighborhood sampling by its one-hop connectivity, and picked *Patient* node as the center node, since it has one-hop connections to both *Diagnoses* and

Lab_test nodes. Specifically, for one training center *Patient* node, we uniformly sampled 10 *Diagnoses* onehop direct connected nodes, and 10 *Lab_test* one-hop direct connected nodes. From these sampled 10 *Diagnoses* nodes, we sampled another 10 *Patient* nodes, each having connections with each of the prior 10 *Diagnoses* nodes. In this way, we connected the center patient node with similar other *Patient* nodes by their common diagnoses. We also sampled the patient node which belongs to the next hour corresponding to the center *Patient* node. For negative sampling, we performed uniform sampling through all *Diagnoses* nodes and *Lab_test* nodes that do not have one-hop connections with the center training patient node. We then projected these different nodes into the same latent space through TransE model. After unifying the embeddings for different node types, each concept is represented as a point in a Euclidean space. In this space, we can measure the similarity between any two vectors using dot product.

2.3.3 HGM Embeddings with CNN Model

The HGM embedding vector encodes not only a patient's information, but also their relation with diagnoses, lab tests, and subsequent lab test results in time. The patient node is represented as a vector $X_p \in \mathbb{R}^{477}$ containing the numerical values measured from lab tests averaged at that time step. We concatenated the resulting embedding vectors to feed into the baseline CNN vertical feature dimension to form a final feature vector within every hour, and used these new features as the CNN input to predict mortality. In addition, since we encoded time as a relation type, we can infer the embedding vector of time steps with missing data based on information from the previous hour. We visualized this procedure in Figure 1.

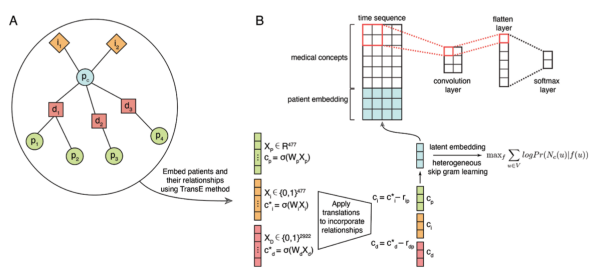


Figure 1. (A) A graphical representation of the HGM for p: patient, i: lab test, and d: diagnosis data. (B) All graph nodes in (A) have a corresponding vector like those shown in (B). The vector representations can be projected into a shared space with the TransE method, and this projection is optimized for retaining relations in the original data in the embedding via skip-gram optimization. Finally, these vectors are concatenated into the CNN model for mortality prediction.

3. EXPERIMENTS

We aim to predict mortality 6, 12, 24, and 48 hours prior to death and/or discharge. The CNN model was used for prediction as introduced in Section 2.2. The CNN model architecture has two convolutional layers, where the first convolution filter is 5x2, the second layer filter is 3x2, and a maxpooling layer between these two layers. Following the convolution layer is a fully connected layer with latent dimension of 100 neurons. The final layer is a sigmoid layer for predicting the output probability. We compared three different scenarios to test the impact of adding HGM embedding vectors as additional features to the framework:

- HGM: Embed patient labs and diagnosis raw data
- CNN: Use raw lab test feature
- HGM+CNN: Concatenate the HGM patient embedding vector, and the raw lab test feature vector

For baselines, we also compared our developed models with three traditional machine learning models: Logistic Regression (LR), Support Vector Machine (SVM), and Random Forest (RF). In this work, we used AUROC and AUPRC scores as the primary performance metric. We tabulated the results in Tables 1 and 2, and we show the evaluation AUROC and AUPRC curves for these tasks in Figure 2.

Model	Hours prior to death				
	6	12	24	48	
LG	0.689±0.01	0.691±0.01	0.672±0.02	0.675±0.02	
SVM	0.654±0.01	0.661±0.02	0.652±0.01	0.653±0.01	
RF	0.667±0.02	0.671±0.01	0.663±0.02	0.654±0.02	
HGM	0.714±0.02	0.715±0.03	0.653±0.03	0.641±0.03	
CNN	0.782±0.01	0.771±0.02	0.775±0.01	0.767±0.01	
HGM+CNN	0.800 ±0.01	0.791 ±0.02	0.796 ±0.01	0.771 ±0.01	

 Table 1. Mortality prediction AUROC evaluation.

Note: Mean values from 10-fold cross validation with standard deviation for confidence intervals.

 Table 2. Mortality prediction AUPRC evaluation.

Model	Hours prior to death					
	6	12	24	48		
LG	0.545±0.01	0.556±0.02	0.542±0.01	0.539±0.01		
SVM	0.487±0.02	0.501±0.01	0.498±0.02	0.487±0.02		
RF	0.512±0.02	0.523±0.02	0.510±0.01	0.503±0.01		
HGM	0.557±0.02	0.559 ± 0.02	0.578±0.02	0.567 ± 0.03		
CNN	0.590±0.01	0.577±0.02	0.589±0.01	0.585 ± 0.02		
HGM+CNN	0.601±0.01	0.600±0.01	0.604±0.01	0.617 ±0.02		

Note: Mean values from 10-fold cross validation with standard deviation for confidence intervals.

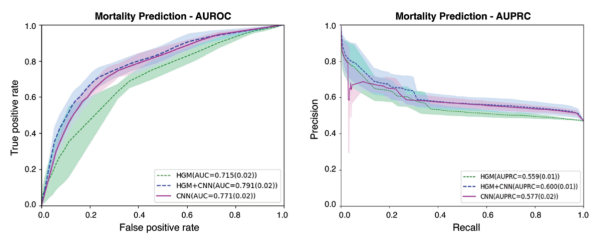


Figure 2. Evaluation of AUROC and AUPRC curves for HGM, CNN, and HGM+CNN models.

The testing results show that the HGM+CNN outperforms all baseline models and both the basic HGM and CNN models, indicating the additional information added from the HGM patient embeddings increases the accuracy of predicting in-patient mortality. The prediction accuracy of using different hours prior to death and/or discharge does not vary by much, indicating that different time windows do not have a major impact on the result for this particular task and modeling strategy. The prediction accuracy in the CNN model drops by 1% in the case of six hours prior to death and/or discharge, but not in the other two models, indicating that using the embedding features from HGM model is slightly more robust than the raw data.

4. DISCUSSION AND CONCLUSION

In this work, we proposed a method to incorporate patient embedding vector from an HGM model into a CNN model in order to provide more information via interconnectivity between different clinical concepts. We assessed the value of this implementation on a task of predicting mortality in EHR data. The results of our experiment show the superior performance of adding the additional patient embedding vector, which is pretrained from the HGM model, compared to pure raw features as the input to CNN model and traditional ML models, too. In one aspect, this is due to the fact that the HGM embedding vector captures additional relational information between different medical concepts, thus providing additional information to the CNN model.

Furthermore, we observed that concatenating the HGM embedding vector with diagnosis feature vectors did not increase the accuracy *versus* using the concatenation between raw lab test and diagnosis feature vectors. This finding indicates that the raw lab test feature vector can provide unique information for CNN to utilize. At the same time, this finding indicates that the embedded patient vector from HGM model could lose some information from the raw lab test feature along the process of projecting these data into a low dimensional latent space. By concatenating all feature vectors, we aim to preserve the information from different data points, which helps to achieve higher mortality prediction accuracy. There are a few limitations

to this study. First, these findings need to be replicated in another data set. Also, exploring more baselines other than the ones shown in this work is beneficial for evaluating the improvements overall. We hope the findings from this work can be expanded in future directions that may add more EHR node types and time components on a variety of other important health-related predictive tasks.

AUTHOR CONTRIBUTIONS

T.Y. Wanyang (tingyi.wanyan@mssm.edu), A. Azad (azad@iu.edu), Y. Ding (ying.ding@austin.utexas.edu), and B.S. Glicksberg (benjamin.glicksberg@mssm.edu) conceived of the project. T.Y. Wanyang and B.S. Glicksberg collected the data. TW and H. Honarvar (hoseinhonarvar@gmail.com) performed the analyses and made the figures. T.Y. Wanyang, HH, and B.S. Glicksberg wrote the manuscript. TW, HH, A. Azad, Y. Ding, and B.S. Glicksberg edited the manuscript and provided revisions. A. Azad, Y. Ding, and B.S. Glicksberg jointly supervised the work.

DATA AVAILABILITY STATEMENT

All the data are available in the Science Data Bank repository, https://doi.org/10.11922/sciencedb. j00104.00094, under an Attribution 4.0 International (CC BY 4.0).

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