Time-Series Event Prediction with Evolutionary State Graph

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ABSTRACT

The accurate and interpretable prediction of future events in time-series data often requires the capturing of representative patterns (or referred to as states) underpinning the observed data. To this end, most existing studies focus on the representation and recognition of states, but ignore the changing transitional relations among them. In this paper, we present evolutionary state graph, a dynamic graph structure designed to systematically represent the evolving relations (edges) among states (nodes) along time. We conduct analysis on the dynamic graphs constructed from the time-series data and show that changes on the graph structures (e.g., edges connecting certain state nodes) can inform the occurrences of events (i.e., time-series fluctuation). Inspired by this, we propose a novel graph neural network model, Evolutionary State Graph Network (EvoNet), to encode the evolutionary state graph for accurate and interpretable time-series event prediction. Specifically, Evolutionary State Graph Network models both the node-level (state-to-state) and graph-level (segment-to-segment) propagation, and captures the node-graph (state-to-segment) interactions over time. Experimental results based on five real-world datasets show that our approach not only achieves clear improvements compared with 11 baselines, but also provides more insights towards explaining the results of event predictions.

KEYWORDS

Time series prediction, evolutionary state graph, graph networks

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1 INTRODUCTION

The prediction of future events (e.g., anomalies) in time-series data has been an important task for temporal data mining [1, 14, 31, 32]. One common approach is latent state machines. For example, HMM [38], RNN [6] and their variants [12, 23] use series of latent representations to encode temporal data. However, such black-box encoding does not directly capture representative patterns (or referred to as “states”) that carry physical meanings in practice, such as walk or run in the observations from fitness-tracking devices. While these methods sometimes can obtain strong results, they are still sensitive to noises [41], provide poor interpretability, and are hard to debug when things go wrong. For this reason, many recent studies focus on discretizing time-series and finding the underlying states, with methods such as sequence clustering [20, 45], dictionaries (e.g. SAX [28, 41], BoP [29]) and shapelets [30, 39]. While effectively handling noises and providing better interpretability, they only recognize the states but ignore the potential effects of relations among them.

To jointly model the states and their relations, recent studies have started to explore the usage of graph structures, such as GCN-LSTM [31] and Time2Graph [10]. However, GCN-LSTM requires an explicit graph as input (e.g., in-app action graph), which is difficult to directly get from general time-series data. Time2Graph uses shapelets to discover states and relations, but it only computes a single static graph over the whole timeline, despite the fact that the...
state relations might change over time (e.g., node-level dynamics and graph-level migration, cf. Section 3.1 for details). To the best of our knowledge, no existing studies have successfully captured and modeled the time-varying relations among the time-series states.

In this work, we observe that time-series are often affected by the joint influence of different states, and in particular, the change of relations among states. For example, in the sequential observations from fitness-tracking devices, stopping exercise from an intense run may cause the fainting event, while the monitoring data will look normal if one stops exercise from jogging; from online shopping records, a sudden interest change from electronics to cosmetics might be more suspicious than a smooth one from cosmetics to fashion.

Motivated by such observations, we propose a novel framework for time-series event prediction, by constructing and modeling a dynamic graph structure as shown in Figure 1. Following existing studies [10, 29, 30, 41], we model time-series based on the underlying states. However, to preserve more information from the original time-series data, we model each time-series segment as belonging to multiple states with different recognition weights, and leverage a directed graph to model the transitional relations among states between adjacent segments. Since the graph evolves along the time-series, we refer to it as an evolutionary state graph. Our empirical observations find that: 1) time-series evolution can be translated into different levels of graph dynamics; 2) when an event occurs, the time-series fluctuation can be expressed as the migration of graph structure, in particular, the dynamics of some edges connecting certain states (Section 3.1).

Despite the insights provided by our empirical observations, there still remains the challenge of how to quantitatively leverage the evolutionary state graph to improve the performance of time-series event prediction. Existing GNN models only consider a static graph or node-level dynamics [10, 27, 31, 35], which cannot be directly used for learning with our evolutionary state graph. In light of this, we propose a novel GNN model, Evolutionary State Graph Network (EvoNet), to further model the graph-level propagation and node-graph interactions with a temporal attention mechanism. The learned representations are then fed into an end-to-end model for time-series event prediction (Section 3.2).

To validate the effectiveness of EvoNet, we conduct experiments on five real-world datasets. Our experimental results demonstrate the superiority of EvoNet over 11 state-of-the-art baselines on time-series event prediction (Section 4.4). We further conduct comprehensive ablation and hyper-parameter studies to validate the effectiveness of our proposed method (Section 4.5). Finally, we demonstrate the insights towards prediction explanation by visualizing EvoNet and its evolutionary state graph (Section 4.6).

The main contributions of this work are summarized as follows:

- Through real-world data analysis, we find the time-varying relations among states important for time-series event prediction.
- We propose the evolutionary state graph to capture the dynamic relations among states, and develop EvoNet to improve the performance of event prediction based on such graphs.
- We conduct extensive experiments on five datasets to demonstrate that our method can both make more accurate predictions, and provide more insight towards explaining them.

2 BACKGROUND AND PROBLEM

Time-series event prediction. We consider the task of predicting future events in a given time-series sequence, following similar definition in previous work [1, 14, 31, 32]. Each time-series sequence with $T$ chronologically paired segments can be represented as $(X_{1:T}, Y_{1:T}) = \{(X_1, Y_1), (X_2, Y_2), \ldots, (X_T, Y_T)\}$, where $X_t \in \mathbb{R}^{\times d}$ and $Y_t \in \mathbb{Z}$ denote a time-series segment [3] and the observed event in the corresponding time (e.g., anomalies), respectively. Each segment $X_t$ is a contiguous subsequence, i.e., $X_t = \{x_1, \ldots, x_r\}$, where $x_i \in \mathbb{R}^d$ is a $d$-dimensional observation at the $i$-th time unit; segment length $r$ is a hyper-parameter which indicates certain physical meanings (e.g. 24 hours). If a time-series sequence can be divided by $T$ segments of equal length $r$, we then have $(X_{1:T}, Y_{1:T}) = \{(x_{r+t-1}, \ldots, x_{r+t+r}), Y_t\}_{0 \leq t < T}$. In this work, we aim to predict the future event $Y_{T+1}$ via discovering time-series states behind $(X_{1:T}, Y_{1:T})$ and modeling their dynamic relations.

State. A state $s$ is a segment that indicates a representative pattern in the time-series sequence, denoted as $\Theta_s \in \mathbb{R}^{\times rd}$. In our study, we adopt existing methods (e.g., Symbolic Aggregate Approximation [28, 41], Bag of Patterns [29], Shapelets [30, 39], sequence clustering [20, 43]) for recognizing interpretable states from time-series data (e.g., symbolic values, shapes or clusters), which are shown to be effective in handling noises and providing good interpretability. As a minor but necessary contribution, we present different implementations of state recognition in the appendix (Section A.2), which act as interchangeable data pre-processors in our framework, and we conduct experiments in Section 4.5 to compare them.

Segment-to-state representation. Once the states have been recognized, one can then models each time-series segment $X_t$ as a composition of states—i.e., quantify the recognition weight of each state to a segment to characterize the segment-state associations. Formally, given a segment $X_t$ and a state $\Theta_s$, the recognition weight $P(\Theta_s|X_t)$ is a measurement of similarity, defined as follows.

$$P(\Theta_s|X_t) = \frac{\max([D(X_t, \Theta_s)]_{\in \mathcal{V}})}{\max([D(X_t, \Theta_s)]_{\in \mathcal{V}}) - \min([D(X_t, \Theta_s)]_{\in \mathcal{V}})},$$

where $D(X_t, \Theta_s)$ can be formalized as the Euclidean Distance or other distances based on different time-series representation and state recognition methods. (cf. Section A.2 for details in the appendix). The smaller this distance, the higher the weight $P(\Theta_s|X_t)$.

3 EVONET FRAMEWORK

In this section, we present a novel framework for time-series event prediction. We name the proposed framework Evolutionary State Graph Network (EvoNet), as it transforms the time-series into a dynamic graph based on the states and recognition weights, and constructs a GNN-based neural network to capture significant correlations and improve the ability of event prediction.

3.1 Evolutionary State Graph

Inspired by existing models introduced in Section 2, we aim to leverage the underlying states for effective and interpretable modeling of time-series. A straightforward approach is to regard a time series as a sequence of the most likely states (for each segment in the sequence), and then model their sequential dependencies [1, 20]. However, one segment may not belong only to a single state; rather it should be recognized as multiple states with different weights.
To this end, one can adopt a multiscale recurrent network (MRNN) [36] to model a multidimensional sequence of state weights, but this method does not highlight the transitions among the states, which may essentially determine whether an event occurs. Therefore, in this work we propose a novel dynamic graph structure to describe the relations among the states and explore how the dynamic shifts of states can reveal time-series evolution.

**Evolutionary state graph.** We define the evolutionary state graph as a sequence of weighted-directed graphs $(G^{(t)})_{t=1}^T$. Specifically, each graph is formulated as $G^{(t)} = (V^{(t)}, E^{(t)}, M^{(t)})$ to represent the transitions from the states of segments $X_{t-1}$ to those of $X_t$. Each node in the graph indicates a state $v$; each edge $e^{(t)}_{(v,v')}$ represents the transition weight (or relation in short) from $v$ to $v'$, along with the transition weight $m^{(t)}_{(v,v')}$, which is the joint weight that $X_{t-1}$ is recognized to the state $v$, while $X_t$ is recognized to the state $v'$.

Compared with existing time-series representations based on states, our evolutionary state graph preserves more information from the original data along the timeline through the modeling of multiple states in each segment and their changing transitional relations. It allows the subsequent model to be more powerful and provide richer interpretations in its predictions, while inheriting from state-based representations the robustness towards noises.

**Real-world example and analysis of evolutionary state graph.** To demonstrate how the evolutionary state graph reveals the evolution of time-series and helps the prediction of events, we conduct an observational study on the NetFlow dataset (cf. Section 4.1 for details of the dataset). As we can see from the case shown in Figure 2, when an anomaly event occurs, the state transitions ($#2\rightarrow#16$) and ($#2\rightarrow#8$) are more frequent at time I; similarly, the state transitions ($#2\rightarrow#8$) and ($#8\rightarrow#23$) are obvious at time II. These transitions reveal that the unbalanced inflow and outflow (state 8 and state 16), or flow drop (state 23), will cause anomalies of network devices. At time III, no anomaly occurs during this period. We can see that states primarily stay in $#2$. There is then an anomaly in the next immediate moment at time IV. Accordingly, we can see a clear increase of the state transition $#2\rightarrow#16$.

In light of the observations, we conduct statistic analysis related to the constructed evolutionary state graph based on the abnormal state transitions. This analysis reveals that the abnormal state transitions ($#2\rightarrow#16$) and ($#2\rightarrow#8$) are more frequent than normal transitions. This finding is consistent with the observations made in the real-world example. Therefore, the constructed evolutionary state graph provides a more comprehensive and accurate model of time-series evolution, which can be used for event prediction and anomaly detection.
samples (an anomaly occurs at time $t$) and normal samples (no anomaly occurs). The distributions of the different graph-level and node-level measurements at different times (before and after anomaly $t$) are visualized in Figure 3. From the figure, we can clearly see that when an anomaly occurs, the abnormal graph (red bar) tends to be denser; i.e., the betweenness scores gets lower, while the closeness scores gets higher. Figure 3b presents three typical states and compares their in-degree before and after anomaly $t$. We can see that the in-degrees of state 8 and 16, indicating the unbalanced inflow and outflow, gradually increase before $t$; this illustrates that the network gradually becomes abnormal. The in-degree of state 23 suddenly increases, indicating that the flow drop is an unexpected event. When no anomaly occurs, we can see that the normal evolutionary state graph (blue lines) generally remains unchanged.

Through the example, we show how the transformation of time-series into evolutionary state graphs allows us to capture the relations between states and their evolution. Meanwhile, we also learn that the graph-level and node-level evolutions can reveal different contextual information related to the time-series events: the node-level evolution reveals the states’ skips when events occur, while the graph-level evolution presents the time-series migration. Intuitively, we shall capture these two levels of information simultaneously when learning with evolutionary state graphs.

### 3.2 Evolutionary State Graph Network

**Overview.** Motivated by Section 3.1, unlike most existing works [1, 30, 41] which model the independent effects of each state, we develop EvoNet to capture the following two types of information through the leverage of our evolutionary state graph:

- **Local structural influence:** the same state $v$ will cause different observations when $v$ is transmitted from different states. In other words, the relations among states matter. For example, *stopping exercise* from an *intense run* may cause fainting, while the monitoring data will look healthier if one *stops exercise* from *jogging*.

- **Temporal influence:** previous transitions of states will influence the current observed data. For example, (*intense run* → *jogging* → *· · · → *stopping exercise*) and (*jogging* → *jogging* → *· · · → *stopping exercise*) lead to different fitness effects.

The above two types of influence can be naturally represented by the evolutionary state graph: the local structural influence is primarily determined by local-pairwise relations among nodes in each graph, while the temporal influence is determined by how relations evolve over different graphs. Inspired by Graph Neural Networks (GNN) [4], we model both the structural and temporal influences of evolutionary state graph by designing two mechanisms: local information aggregation and temporal graph propagation.

Figure 4 illustrates the overall structure of EvoNet. Given the observations $(X_{\cdot|T}, Y_{\cdot|T})$, we first recognize states for each segment $X_t$ and construct the evolutionary state graph $(G^{(1:T)}|T)$. Next, we define a representation vector $h_v^{(t)} \in \mathbb{R}^{|h|}$ for each node $v$ in graph $G^{(t)}$ to encode $v$’s node-level patterns, and define a representation vector $U^{(t)} \in \mathbb{R}^{|U|}$ for $G^{(t)}$ to encode the graph-level information. Based on this, EvoNet aggregates local structural information by means of message passing, and further incorporates temporal information using the recurrent *EvoBlock*. EvoNet then applies the learned representations $(h, U)$ towards the prediction task.

**Local information aggregation.** In order to aggregate the local structural information in each $G^{(t)}$, EvoNet aims to make two linked nodes share similar representations. To achieve this, we let each node representation $h_v^{(t)}$ in $G^{(t)}$ aggregate the messages of its neighbors, and thus compute its new representation vector. Initially, we let $h_v^{(0)} = \Theta_v$. Recall that $\Theta_v$ is obtained from the state recognition on all segments, which records the time-series information of state $v$. Then, following the message-passing neural network (MPNN) [19] directly, we have the following aggregation scheme:

$$
H_v^{(t)} = \sum_{v' \in N(v)} f_{\text{MP}} \left( h_{v'}^{(t-1)} , e_{v,v'}^{(t)} \right)
$$ (3)
where $H^{(t)}_0$ is the intermediate representation of node $v$ following aggregation, which combines the messages from all neighbors $N(v)$ in the graph $G^{(t)}$. The message function $\mathcal{F}_{\text{MP}}(\cdot, \cdot)$ can be implemented by many existing neural networks, such as GGNN [27]:

$$
\mathcal{F}_{\text{MP}}(h^{(t)}_v, \epsilon_{(u,v)}) = W_{\text{MP}} \left[ m_{(u,v)} \times h^{(t-1)}_u \right] + b_{\text{MP}} \tag{4}
$$

where $m_{(u,v)} \times h^{(t-1)}_u$ is the passing message, while $W_{\text{MP}}$ and $b_{\text{MP}}$ are the learnable parameters, indicating the passing weight and bias. We also have other implementations for $\mathcal{F}_{\text{MP}}$, such as pooling, GCN [15], GraphSAGE [22], GAT [42], etc. (cf. Section A.3 for details in the appendix). Herein, we serve $\mathcal{F}_{\text{MP}}$ as interchangeable modules in EvoNet and conduct experiments in Section 4.5 to analyze the effectiveness of different implementations.

**Temporal graph propagation.** In addition to aggregating the local structural information, previous transitions also influence current representations. Moreover, when events occur, the modes of the graph-level and node-level evolution will change (Section 3.1). Intuitively, we should capture these two kinds of temporal information simultaneously. To achieve this, we design a recurrent block, named EvoBlock, to capture the evolving information in the evolutionary state graph. EvoBlock combines the local aggregated representation $H^{(t)}_v$ and the past representation $h^{(t-1)}_v$, formulated as

$$
h^{(t)}_v, U^{(t)} := \mathcal{F}_{\text{recur}} \left( h^{(t-1)}_v, h^{(t-1)}_v, U^{(t-1)} \right) \quad \text{for } \alpha \in \mathcal{V} \tag{5}
$$

where $\mathcal{F}_{\text{recur}}$ indicates a recurrent function that allows us to incorporate information from the previous timestamp in order to update current representations. When there are fewer messages from other nodes, i.e., $m_{(u,v)} \to 0$, $h^{(t)}_v, U^{(t)}$ will be more influenced by the previous $h^{(t-1)}_v, U^{(t-1)}$. Otherwise, the messages will influence current representations more.

As shown in Figure 4a, most existing works implement $\mathcal{F}_{\text{recur}}$ using simple recurrent neural networks on node-level propagation (e.g., GGNN [27] adopts GRU [12], GCN-LSTM [31] adopts LSTM [23], etc.). For the graph-level propagation $U^{(t)}$, these methods simply pool the node-level representations, i.e., $U^{(t)} = \sum_{v \in \mathcal{V}} h^{(t)}_v$. However, in our empirical observations (Section 3.1), both the graph and nodes in the evolutionary state graph will present different temporal information when events occur. In order to improve the ability of event prediction, $\mathcal{F}_{\text{recur}}$ shall consider the contextual information of previous events $Y_{1:T}$ when modeling the graph-level propagation, and then influence the node-level representations via the node-graph interactions. Accordingly, events are generally scattered in the timeline; thus, we propose a temporal attention mechanism for capturing significant temporal information in node-graph interactions. More specifically, as shown in Figure 4b, we have

$$
\mathcal{F}_{\text{recur}} \left( h^{(t)}_v, h^{(t-1)}_v, U^{(t-1)} \right) := h^{(t)}_v = \Phi_h \left( h^{(t-1)}_v, U^{(t-1)} \right) \tag{6}
$$

where “@” indicates the concatenation operator. The current node-level representation is computed using the function $\Phi_h(\cdot, \cdot)$, based on the past representations $\left( h^{(t-1)}_v, U^{(t-1)} \right)$ and current aggregations $h^{(t)}_v$, while the current graph-level representation is computed by $\Phi_U(\cdot, \cdot)$ based on the past $U^{(t-1)}$ and current event $Y_t$, as well as all node representations $h^{(t)}_v$. The attention score $\alpha_t$ re-weights the node-graph interaction of the $t$-th temporal step, which is computed based on the concatenated patterns of $U^{(t-1)}$ and all aggregations $\sum_{v \in \mathcal{V}} h^{(t)}_v$ under the learnable weight $W_a$. We use the softmax function to normalize $\alpha_t$ during different time steps.

Recurrent function $\Phi_h(\cdot, \cdot)$ smooths the two inputted vectors of each temporal step, and can be implemented using many existing approaches. Herein, we provide an example of $\Phi_h(\cdot, \cdot)$ implemented by LSTM. Formally, we have

$$
\Phi_h \left( h^{(t-1)}_v, U^{(t-1)} \right) =
\begin{cases}
\left( \begin{array}{c}
\alpha_t h^{(t)}_v + \alpha_t U^{(t-1)} \\
1 = \sigma(W_f h^{(t)}_v + b_f) + \beta f \\
C^{(t)} = \sigma(W_c h^{(t)}_v + b_c) \circ \tanh(W_c h^{(t)}_v + b_c) \\
O^{(t)} = \sigma(W_o h^{(t)}_v + b_o) \circ \tanh(C^{(t)})
\end{array} \right)
\end{cases}
\tag{7}
$$

where $f, C^{(t)}$ and $O^{(t)}$ are forget gate, input gate and output gate respectively, while $\sigma$ is a sigmoid activation function. The current node vectors are updated by receiving their own previous memory and current memory. In our experiments, we compare the performance of different methods for EvoBlock (Table 2).

**End-to-End Model Learning.** Thus, the representations $h^{(t)}_v$ and $U^{(t)}$ capture both the node-level and graph-level information respectively until the $t$-th temporal step, which can then be applied to predict the next event $Y_{t+1}$. More specifically, we encode the current evolutionary state graph $G^{(t)}$ into representation $h^{(t)}_G$ based on the concatenated features of all $h^{(t)}_v$ and $U^{(t)}$, which can be formulated as

$$
h^{(t)}_G = \mathcal{F}_k \left( U^{(t)} \oplus \sum_{v \in \mathcal{V}} h^{(t)}_v \right) \tag{8}
$$

where $\mathcal{F}_k$ acts as a fully connected layer. We then learn a classifier, such as a neural network or XGBoost [9], which takes $h^{(t)}_G$ as input and estimates the probability of the next event, $P(Y_{t+1} | h^{(t)}_G)$. To learn the parameters $\theta$ of the proposed EvoNet and classifier, we employ an end-to-end framework, based on the Adam optimization algorithm [26] to minimize the cross-entropy loss $L$ as follows:

$$
L = - \sum_{Y_{t+1}} \log P \left( Y_{t+1} | h^{(t)}_G \right) + (1 - Y_{t+1}) \log \left( 1 - P \left( Y_{t+1} | h^{(t)}_G \right) \right) \tag{9}
$$

where $Y_{t+1} \in \{0, 1\}$ is the ground truth that indicates whether a future event will occur. The procedure of state recognition and graph propagation are carried out step by step: we first recognize the states and construct an evolutionary state graph, then conduct the evolutionary state graph propagation to model the time-series. $|\mathcal{V}|$-node graphs are constructed in $T$ segments, such that the time complexity of each iteration is $O(T \times |\mathcal{V}|^2)$.

## 4 EXPERIMENTS

We apply our method to the prediction of upcoming events in time-series data, and aim to answer the following three questions:

- **Q1:** How does EvoNet perform on the time-series prediction task, compared with other baselines from the state-of-the-art?
\begin{itemize}
  \item Q2: How does the proposed EvoBlock effectively bridge the graph-level and node-level information over time?
  \item Q3: How do different configurations, e.g., state number, segmentation length, implementation of state recognition and message passing, influence the performance?
\end{itemize}

### 4.1 Datasets

We employ five real-world datasets to conduct our experiments, including two public ones (DJIA30 and WebTraffic) from Kaggle\(^1\), and another three (NetFlow, ClockErr and AbServe) provided by China Telecom\(^2\), State Grid\(^3\) and Alibaba Cloud\(^4\), respectively. Table 1 presents the overall dataset statistics.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>#samples</th>
<th>Positive Ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DJIA30</td>
<td>15,540</td>
<td>9.5</td>
</tr>
<tr>
<td>WebTraffic</td>
<td>2,992,184</td>
<td>28.2</td>
</tr>
<tr>
<td>NetFlow</td>
<td>238,000</td>
<td>8.6</td>
</tr>
<tr>
<td>ClockErr</td>
<td>6,879,834</td>
<td>14.9</td>
</tr>
<tr>
<td>AbServe</td>
<td>12,224</td>
<td>1.5</td>
</tr>
</tbody>
</table>

#### 4.2 Baseline Methods

We compare our proposed EvoNet with several groups of baselines:

1. An online community of data scientists and machine learners.
2. A major mobile service provider in China.
3. A major electric power company in China.
4. The largest cloud service provider in Asia.

### Feature-based models

Several popular feature-based algorithms have been proposed for time-series analysis. In this paper, we choose some typical algorithms to compare with our model: Bag of Patterns (BoP)\(^{29}\), Vector Space Model using SAX (SAX-VSM)\(^{41}\) and Fast Shapelet (FS)\(^{39}\). These methods capture different state representations, which serve as features for event predictions.

#### Graph-based models

Recently, many GNN-based works are proposed to model the (dynamic) graphs. In this paper, we choose several state-of-the-art algorithms as baselines to model the evolutionary state graph, and conduct the same approaches for event prediction as EvoNet: gated graph neural network (GGGNN)\(^{27}\) initializes the node vector \(h^0\) using a one-hot vector of the corresponding state; it conducts GGGNN\(^{27}\) for local message passing and only adopts a GRU structure\(^{12}\) for node-level propagation. GCN-LSTM\(^{31}\) uses states’ patterns \(\Theta\) to initialize the node vector \(h^0\); it conducts GCN\(^{15}\) for local message passing and LSTM structure\(^{23}\) for node-level propagation. EvoWeGCN\(^{35}\) is a dynamic graph neural network that builds a multi-layer framework to combine RNN and GCN; it also focuses on node-level propagation. ST-MGCN\(^{18}\) is a spatiotemporal multi-graph convolution network, in which \(Y_t\) serve as contextual information for propagation. It directly fuses the contextual information into node-level representations rather than learning graph-level representations and modeling the node-graph interactions. Time2Graph\(^{10}\) adopts shapelet to extract states; it aggregates the graphs at different times as a static graph and conduct DeepWalk\(^{37}\) to learn graph’s representations, which then serve as features for event predictions.

#### EvoNet variants

We also compare EvoNet with its derivatives by modifying some key components to see how they fare: 1) we sample the most possible state sequence (i.e., each segment is recognized with highest state weight) for each time-series, and directly use LSTM to model the new sequence without building and modeling the evolutionary state graph, denoted as EvoNet w/o G; 2) we build evolutionary state graph for time-series but model it without conducting temporal attention mechanism, denoted as EvoNet w/o A; 3) we conduct complete EvoNet for time-series modeling, denoted as EvoNet. Herein, EvoNet uses the state patterns \(\Theta\) to initialize node vector \(h^0\) and conducts graph-level and node-level propagation for \(G^{(1:T)}\). We implement state recognition and local message...
Table 2: Comparison of prediction performance on five real-world datasets (%). The bold text indicates the best performance among all methods, while the underline text indicates the second-best performance.

<table>
<thead>
<tr>
<th>Datasets</th>
<th>Models</th>
<th>Feature-based models</th>
<th>Sequential models</th>
<th>Graphical models</th>
<th>Our models</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>F1-score</td>
<td>AUC</td>
<td>F1-score</td>
<td>AUC</td>
</tr>
<tr>
<td></td>
<td>BoP [29]</td>
<td>24.92±0.40</td>
<td>50.92±0.19</td>
<td>44.31±0.33</td>
<td>66.87±0.09</td>
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<tr>
<td></td>
<td>FS [39]</td>
<td>24.38±0.97</td>
<td>55.50±0.42</td>
<td>48.39±0.76</td>
<td>69.66±0.23</td>
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<tr>
<td></td>
<td>SAX-VSM [41]</td>
<td>26.06±0.45</td>
<td>51.42±0.20</td>
<td>46.66±0.49</td>
<td>67.63±0.15</td>
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<tr>
<td>Features</td>
<td>S-HMM [1]</td>
<td>25.20±0.48</td>
<td>51.14±0.20</td>
<td>43.09±0.41</td>
<td>66.54±0.12</td>
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<td></td>
<td>MRNN [36]</td>
<td>21.20±0.42</td>
<td>49.39±0.19</td>
<td>44.43±0.57</td>
<td>67.51±0.17</td>
</tr>
<tr>
<td></td>
<td>HRNN [11]</td>
<td>26.43±0.87</td>
<td>52.66±0.29</td>
<td>45.79±0.82</td>
<td>68.27±0.26</td>
</tr>
<tr>
<td></td>
<td>ST-MGCN [18]</td>
<td>23.72±0.91</td>
<td>51.56±0.31</td>
<td>43.30±1.25</td>
<td>67.14±0.38</td>
</tr>
<tr>
<td></td>
<td>EvolveGCN [35]</td>
<td>25.76±0.85</td>
<td>52.66±0.30</td>
<td>45.67±0.90</td>
<td>68.15±0.29</td>
</tr>
<tr>
<td></td>
<td>Time2Graph [10]</td>
<td>26.16±1.24</td>
<td>53.01±0.55</td>
<td>45.90±1.58</td>
<td>68.38±0.41</td>
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<tr>
<td></td>
<td>ST-MGCN [18]</td>
<td>26.93±0.97</td>
<td>53.39±0.39</td>
<td>45.96±0.91</td>
<td>68.74±0.27</td>
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<td></td>
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<td>53.28±0.39</td>
<td>46.03±1.12</td>
<td>68.74±0.43</td>
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<td>55.07±0.39</td>
<td>47.02±0.95</td>
<td>69.03±0.27</td>
</tr>
</tbody>
</table>

The bold text indicates the best performance among all methods, while the underline text indicates the second-best performance.
Figure 5: The impact of the different parameters. (a)-(d) present comparisons of state number $|\mathcal{V}|$, segment length $\tau$, implementation of message passing and state recognition, respectively, over three datasets (WebTraffic, NetFlow and AbServe) in Section 4.1.

Figure 6: A case of EvoNet conducted for anomaly analysis of cloud service. (a) visualizes the raw time-series of CPU utilization. The red lines indicates anomaly events during different intervals ($\tau = 5$). We set 10 states for constructing evolutionary state graph and conducting EvoNet. Heat map in (b) presents the attention score $\alpha_t$ of EvoNet at different temporal steps. (c)-(d) visualize the evolutionary state graph in two different intervals (I, II marked in (a)) and the aggregated graph in the whole timeline, respectively. (e) presents the raw segments corresponding to the five different states.

1. Sensitivities of state number $|\mathcal{V}|$. As shown in Figure 5a, prediction performance curves differ depending on the dataset, illustrating that the state number $|\mathcal{V}|$ is sensitive to the data owns patterns. Moreover, the performance is not bound to improve as $|\mathcal{V}|$ increases, suggesting that $|\mathcal{V}|$ is an empirically determined parameter and is unsuitable for large values.

2. Sensitivities of segment length $\tau$. Another sensitive parameter is the segment length $\tau$, the variation of which may change the temporal scale of event $Y_t$, and thus the positive ratio of ground truth. We can see the performances in Figure 5b do not vary significantly, meaning that it can be an empirical parameter that is generally determined by the realistic demand (e.g. an acceptable temporal scale of anomaly detection, etc.)

3. Implementation of message passing. Figure 5c presents the comparisons for different implementations of message passing. We can observe that GAT and GraphSAGE perform poorly and are unstable due to their full attention or sampling operation, which is unsuitable for the small-scale graph. The performances of GGNN and GCN are similar, and both outperform the pooling method.

4. Implementation of state recognition. As shown in Figure 5d, we test different implementations of state recognition, and further compare them with some feature-based baselines (i.e., SAX-VSM [41] and Fast Shapelets [39]). We can see that EvoNet can clearly improve the performance of SAX-VSM and Fast Shapelets when models the relations. Moreover, the implementations of cluster methods and shapelet outperform the SAX word; this is because each SAX word is simply a symbolic value representing state, while other representations are a vector describing state patterns, which provide more information for modeling the evolutionary state graph.

4.6 Case Studies

In this section, we apply our EvoNet method to a real-world anomaly prediction scenario in Alibaba Cloud\(^3\), enabling us to demonstrate how this method can be used to find meaningful relational clues to explain its results. As described in Section 4.1, the minutely time-series of server monitor are segmented by the interval $\tau = 5$ (empirical length). In order to present clearly, we cluster 10 states for constructing evolutionary state graph and conduct EvoNet for anomaly prediction. We visualize the results including several states and the evolutionary state graph at different times. The temporal attention scores learned by EvoNet are also visualized to validate its effectiveness. All results are presented in Figure 6.

1. Effectiveness of temporal attention mechanism. As shown in Figure 6(a)-(b), we adopt heat map to visualize the attention scores $\alpha_t$ learned by Eq 6 at different times. We can see that the attention scores successfully highlight the positions of anomalies in (a) (i.e., the positions near 13:00 and 20:00), which demonstrate that the temporal attention mechanism is useful for EvoNet to capture significant temporal information.

\(^3\)Our method has been deployed by SLS, Alibaba Cloud, the largest log service provider in China, acting as a common function.
2. Interpretability of evolutionary state graph. We then explore how an evolutionary state graph can be used to find meaningful insights that can explain anomaly event. As shown in Figure 6(a), we mark two intervals, I and II, to visualize evolutionary state graph and explore some meaningful insights. The results are shown in Figure 6(c)-(d). We can see that there is a major transition #2→#4→#0 (i.e., thick edges) in the graph of I, while #3→#5 is a major transition in the graph of II. Note that there is an anomaly occurring immediately after interval II. When we aggregate all evolutionary state graphs in the timeline (Figure 6(d)), we can find that the transition #2→#4→#0 is the major path in the graph, while #3→#5 is a rare path (i.e., thin edges). These observations indicate that the transition #3→#5 occurred in interval II is abnormal, which is consistent with the anomaly of cloud service. As shown in Figure 6(e), we present the average curve of segments with different states. We can see that the transition #2→#4→#0 indicates a process of service, i.e., CPU utilization rises from 0.25 to 0.75 and drops after maintaining a period. On the contrary, transition #3→#5 indicates that CPU utilization rises to 0.5 and then drops immediately. These observations demonstrate that this anomaly may be caused by the CPU’s fault.

5 RELATED WORK

Time-series modeling. Time-series modeling aims to capture the representative patterns underpinning observed data. One important trend here is sequential modeling, such as HMM [38], RNN [6] and their variants [12, 23, 24, 45] and fitting auto-regressive models [2]. They define one latent representation to capture all the patterns by modeling the sequential dependencies, rather than distinguishing different states. Another trend is mining discretized sequential patterns, such as switch time-series models [1] and dictionaries [28, 29, 41]. They model the time-series by capturing different states of segments independently, but ignore the influence from their relations. Recently, some works use hierarchical or attention connections to incorporate the above two techniques and get good performance [11, 49]. However, most of them only capture the patterns of states, ignoring their relations. Some works have applied graph structure into the relation modeling of time-series states [10, 20, 31], which aims to represent different segments, rather than capturing the dynamics. To the best of our knowledge, no existing studies have successfully modeled the time-varying relations among states.

Graph neural networks. Models in the graph family [4, 15, 16, 22, 27, 42, 46] have been applied to many real-world scenarios, including learning the dynamics of physical systems [5, 40], predicting the chemical properties of molecules [17], predicting traffic on roads [18] and reasoning about knowledge graphs [21], etc. These studies present the effectiveness of GNNs for modeling structural information. Some works focus on summarizing models and refining formal expressions. The message-passing neural network (MPNN) unified various graph convolutional network and graph neural network approaches by analogy to message-passing in graphical models [19]. The non-local neural network (NLNN) has a similar vein, which unified various "self-attention"-style approaches by analogy to methods from graphical models and computer vision for capturing long range dependencies in signals [44]. Recently, some works have attempted to model dynamic graphs using GNNs [18, 31, 35], although they focus primarily on the explicit graphical structure. To the best of our knowledge, no existing studies have successfully modeled dynamic relations in non-graphical data, such as time-series.

6 CONCLUSIONS

In this paper, we study the problem of how relations among states reflect the evolution of temporal data. We propose a novel representation, the evolutionary state graph, to present the time-varying relations among time-series states. In order to capture these effective patterns for downstream tasks, we further propose a GNN-based model, EvoNet, to conduct dynamic graph modeling. As for the validation of EvoNet’s effectiveness, we conduct extensive experiments on five real-world datasets. Experimental results demonstrate that our model clearly outperforms 11 state-of-the-art benchmark methods. Based on this, we can find some meaningful relations among the states that allow us to understand temporal data.

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REFERENCES

A APPENDIX

A.1 Algorithm Details

In order to outline our proposed model in detail, we present the complete pseudo code of EvoNet to illustrate the learning procedure. Given the observations \( \{X_{1:T}, Y_{1:T}\} \) and parameters \( \{\|V\|, \tau, \mathcal{F}_{\text{state}}, \mathcal{F}_{\text{MP}}\} \), EvoNet first captures different states by means of the recognition function \( \mathcal{F}_{\text{state}} \). It then constructs the evolutionary state graph \( \mathcal{G}^{(1:T)} \) and conducts graph propagation by means of the message function \( \mathcal{F}_{\text{MP}} \) and EvoBlock. Finally, the learned representations are fed into an output model for prediction tasks; we use a backpropagation learning algorithm with cross-entropy loss to train the entire networks. More details can be found in Algorithm 1.

Algorithm 1 The learning procedure of EvoNet

| Input: | Observations \( \{X_{1:T}, Y_{1:T}\} \), parameters \( \{\|V\|, \tau, \mathcal{F}_{\text{state}}, \mathcal{F}_{\text{MP}}\} \) |
| Output: | Train state recognition model |

1. \( \mathcal{F}_{\text{state}} \leftarrow \{X_{1:T}, \|V\|\},\text{ train state recognition model} \)
2. for each segment \( X_{t} \in X_{1:T} \) do
   3. \( \{P(\mathcal{O}_i|X_{t}) \leftarrow \mathcal{F}_{\text{state}}(X_{t}, \mathcal{O}_i)\}_{v \in V}, \text{ get the recognized weights of states as Eq 1} \)
   4. \( G^{(t)} \leftarrow \text{construct the evolutionary state graph as Eq 2} \)
3. end for
4. while the parameters of EvoNet have not converged do
   5. for each \( X_{t} \in X_{1:T}, Y_{t} \in Y_{1:T}, G^{(t)} \in \mathcal{G}^{(1:T)} \) do
      6. \( \mathcal{H}^{(t)}_{v} \leftarrow \mathcal{F}_{\text{MP}} \{G^{(t)}, \mathcal{H}^{(t-1)}\}_{v \in V}, \text{ conduct message passing as Eq 4} \)
      7. \( \alpha_{t} \leftarrow \text{compute attention score as Eq 6.3} \)
      8. \( \mathcal{H}^{(t)}_{v} \leftarrow \text{node-level propagation as Eq 6.1} \)
      9. \( \mathcal{U}^{(t)} \leftarrow \text{graph-level propagation as Eq 6.2} \)
     10. \( \mathcal{H}^{(t)}_{G} \leftarrow \text{compute current feature embedding as Eq 8} \)
     11. \( P(Y_{t+1}^{\tau}, \mathcal{H}^{(t)}_{G}) \leftarrow \text{estimate the probabilities of the next event} \)
   12. end for
13. \( \theta \leftarrow \nabla_{\theta} \left\{ \frac{1}{N} \sum_{n=1}^{N} \mathcal{L} \right\}, \text{ back-propagate the loss and train the whole EvoNet as Eq 9} \)
14. end while

A.2 Implementation of State Recognition

In this section, we present several implementations for state recognition, including sequence clustering [20], SAX words [28] and Shapelets [30], which have been proven to be competitive for capturing the representative patterns (or states), in previous works.

Sequence Clustering. Cluster methods allow us to find the repeated patterns in time-series segments, which can reduce the dimension and allow us to derive insights capable of explaining time-series evolution [1, 207]. Herein, we take Kmeans[25] as example; the aim here is to partition the \( n \) segments \( X_{t} \) into \( |V| \) sets \( \Omega = \{\Omega_1,...,\Omega_{|V|}\} \), so as to minimize the within-cluster sum of squares, i.e., variance. Formally, the objective is to find:

\[
\arg\min_{\Omega} \sum_{v \in \Omega} \sum_{X_{t} \in \Omega_v} \|X_{t} - \Theta_v\|^2 = \arg\min_{\Omega} \sum_{v \in \Omega} \|\Omega_v\| \text{Var} \Omega_v
\]

where \( \Theta_v \) is the mean of all segments in \( \Omega_v \).

In this section, we present several implementations for state recognition, including sequence clustering [20], SAX words [28] and Shapelets [30], which have been proven to be competitive for capturing the representative patterns (or states), in previous works.

SAX word. Symbolic aggregate approximation (SAX) is the first symbolic representation for time series that allows for dimensional reduction and indexing with a lower-bounding distance measure. It transforms the original time-series segments into several average values (PAA representation\(^4\)) and converts them into a string.

Herein, we can consider each SAX word as a state \( v \) and extend the corresponding average value \( a \) as representative patterns of the time series segments, i.e., \( \Theta_v = [a, ...] \). Based on this, we can normalize the distance \( D(X_{t}, \Theta_v) \) as the recognition weight, following the approach outlined in Eq 1. Subsequently, we can construct the evolutionary state graph to represent the relations among SAX representations.

Shapelet. A shapelet \( \Theta_v \) is a segment that is representative of a certain class. More precisely, it can separate segments into two smaller sets, one that is close to \( \Theta_v \) and another that is far from \( \Theta_v \) according to some specific criteria, such that for a given time series classification task, positive and negative samples can be put into different groups. The criteria for these can be formalized as

\[
\mathcal{L}_{\text{shapelet}} = -g \left( \mathcal{D}_{\text{pos}}(X_{t}, \Theta_v), \mathcal{D}_{\text{neg}}(X_{t}, \Theta_v) \right)
\]

where \( \mathcal{L}_{\text{shapelet}} \) measures the dissimilarity between positive and negative samples towards the shapelet \( \Theta_v \). \( \mathcal{D}_{\text{pos}}(X_{t}, \Theta_v) \) denotes the set of distances with respect to a specific group, i.e., positive or negative class; the function \( g \) takes two finite sets as input and returns a scalar value to indicate how far apart these two sets are. This could be information gain or some dissimilarity measurements on sets (i.e., KL divergence). We can then adopt the same approaches as in the above definitions to recognize states’ weights and construct the evolutionary state graph to represent the relations among shapelets.

A.3 Implementation of Message Passing

As for the implementations of message passing in local information aggregation, there are many existing works addressing this issue, such as pooling, GGNN [27], GCN [15], GraphSAGE [22], GAT [42], etc.. Herein, we present their implementation details. Broadly speaking, the aim of message passing is to aggregate the messages

\[^4https://jmotif.github.io/sax-vsm_site/morea/algorithm/PAA.html\]
of node $v$’s neighbors, and thus to compute its new representation vector, the scheme of which is
\begin{equation}
H_{v}^{(t)} = \sum_{\nu' \in N(v)} \mathcal{F}_{MP}(h_{\nu'}^{(t-1)}, e_{(v,\nu')}^{(t)})
\end{equation}
where $H_{v}^{(t)}$ is the intermediate representation of node $v$ after aggregation; moreover, $\mathcal{F}_{MP}(\cdot, \cdot)$ is the specific message function, which combines the messages from all $v$’s neighbors $N(v)$ in graph $G^{(t)}$.

**Pooling.** Pooling is a simple implementation, which receives the neighbors’ messages by computing the production of these neighbors’ representation and current transition weight. This approach can be formulated as
\begin{equation}
\mathcal{F}_{MP}(h_{\nu}^{(t-1)}, e_{(\alpha,\nu')}^{(t)}) = m_{\nu}^{(t)}(\alpha,\nu') \times h_{\nu}^{(t-1)}
\end{equation}
where $\alpha' \in N(v)$ is a neighbor of node $v$ and $h_{\nu}^{(t-1)}$ is its representation of the last temporal point. $m_{\nu}^{(t)}(\alpha,\nu')$ is the current relation weight, which is computed by Eq 2 (see details in Section 3.1).

**GGNN.** Gated Graph Neural Networks [27] implement a message-feedback mechanism: in short, when node $\alpha'$ passes a message to node $v$ via edge $(\alpha' \rightarrow v)$, $v$ will send a feedback message to $\alpha'$. This approach aggregates the in-degree and out-degree messages from its neighbors, which is formulated as
\begin{equation}
\mathcal{F}_{MP}(h_{\nu}^{(t-1)}, e_{(\alpha,\nu')}^{(t)}) = W_{in} \cdot m_{(\alpha,\nu')}^{(t)} \times h_{\nu}^{(t-1)} + b + W_{out} \cdot m_{(\nu,\alpha)}^{(t)} \times h_{v}^{(t-1)}
\end{equation}
where $W, b$ is the learnable weight and bias, which is related to the downstream task. From the perspective of the whole graph (adjacency matrix), we in fact build a new graph with the opposite directed edges. Hence, the above scheme can be reformulated as
\begin{align}
\mathcal{M}^{(t)} &= \begin{bmatrix}
\mathcal{M}_{in}^{(t)} \\
\mathcal{M}_{out}^{(t)}
\end{bmatrix} = \begin{bmatrix}
m_{(\alpha,\nu')}^{(t)} \\
m_{(\nu,\alpha')}^{(t)}
\end{bmatrix}_{\alpha',\nu' \in V} \\
H^{(t)} &= W \cdot \mathcal{M}^{(t)} \cdot h^{(t-1)} + b
\end{align}

where $\mathcal{M}_{in} = \begin{bmatrix}
m_{(\alpha,\nu')}^{(t)} \\
m_{(\nu,\alpha')}^{(t)}
\end{bmatrix}_{\alpha',\nu' \in V}$ is the adjacency matrix in graph $G^{(t)}$, $\top$ indicates the transposition operator, i.e., $\mathcal{M}_{out}$ is actually the transposition matrix of $\mathcal{M}_{in}$.

**GCN.** Graph Convolution Networks [15] adopt spectral approaches to represent the graph. It computes the eigendecomposition of the graph Laplacian, defined as
\begin{equation}
H^{(t)} = \mathbf{U}^{(t)}(g(\Lambda^{(t)})\mathbf{U}^{(t)})^{T} \cdot h^{(t-1)}
\end{equation}
where $\mathbf{U}^{(t)}$ is the matrix of eigenvectors of the normalized graph Laplacian $L^{(t)} = L_{V} - D^{(t)}^{-\frac{1}{2}} \Lambda^{(t)} D^{(t)}^{-\frac{1}{2}} = \mathbf{U}^{(t)} g(\Lambda^{(t)}) \mathbf{U}^{(t)}^{T}$ ($D^{(t)}$ is the degree matrix and $M^{(t)}$ is the adjacency matrix of the graph $G^{(t)}$, with a diagonal matrix of its eigenvalues $\Lambda^{(t)}$, $g(\cdot)$ is the filter function, which can be approximated by a truncated expansion in terms of Chebyshev polynomials [13].

**GraphSAGE.** In order to avoid transductive learning and naturally generalize to unseen nodes, Hamilton et al. [22] proposed the general inductive framework, GraphSAGE, which generates new representation by sampling and aggregating features from a node’s local neighborhood. The difference between this approach and the aforementioned GGNN (Eq 15) is that the former does not utilize the full set of neighbors, but rather fixed-size set of neighbors through uniform sampling.

**GAT.** Graph Attention Networks adopt a self-attention strategy, which involves computing the representations of each node attending to it over its neighbors. The attention coefficients are computed in the node pair $(v, v')$
\begin{equation}
\alpha_{(v,v')}^{(t)} = \frac{\exp \left( \text{LeakyReLU} \left( W \left( h_{v}^{(t-1)} \oplus h_{v'}^{(t-1)} \right) \right) \right)}{\sum_{v'' \in N(v)} \exp \left( \text{LeakyReLU} \left( W \left( h_{v}^{(t-1)} \oplus h_{v''}^{(t-1)} \right) \right) \right)}
\end{equation}
where $\alpha_{(v,v')}^{(t)}$, is the attention coefficient of node $v$ and $v'$ in $G^{(t)}$, which reweights the edge $m_{(v,v')}^{(t)}$. We can then adopt an approach similar to Eq 3 to obtain $H_{v}^{(t)}$ of each node.

### A.4 Hyperparameter Settings
We have discussed several important hyperparameter settings of the proposed model in Section 4.5. We conduct grid search for our proposed model and baselines in order to find the adaptive hyperparameters and compare fairly. The remaining aspects of parameter options are introduced below to facilitate better reproductivity.

**Hyperparameters in EvoNet.** We test EvoNet at the number of states $|V|$, segment length $\tau$, the size of graph-level representation $|U|$ (the size of node-level representation $|h|$ is determined by state recognition, since $h_{0} = 0$, while the search space may differ between different datasets. We test $|V|$ with values from 5 to 100 with interval 10, and further test $\tau$ with different lengths that are smaller or greater than the period length of the corresponding dataset. We test $|U|$ from $2^{4}$ to $2^{10}$ with exponential interval 1. In batch-wise training for EvoNet, the batch size is set to 1000, and we choose the Adam algorithm [26] as the loss optimizer.

**Hyperparameters in baselines.** As for baselines, we use the source code provided on TSLearn\footnote{https://tslearn.readthedocs.io/en/latest} for several feature-based models, and code the sequential models by ourselves. For the graphical models, we conduct the experiments on the provided codes in GitHub. If the parameter interface is open, we adopt the same grid search approach to search the best parameters. Due to the binary event prediction tasks, we use XGBoost [9] with same parameters for all methods in order to improve the overall performance.