Scalable Spatiotemporal Graph Neural Networks

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Abstract

Neural forecasting of spatiotemporal time series drives both research and industrial innovation in several relevant application domains. Graph neural networks (GNNs) are often the core component of the forecasting architecture. However, in most spatiotemporal GNNs, the computational complexity scales up to a quadratic factor with the length of the sequence times the number of links in the graph, hence hindering the application of these models to large graphs and long temporal sequences. While methods to improve scalability have been proposed in the context of static graphs, few research efforts have been devoted to the spatiotemporal case. To fill this gap, we propose a scalable architecture that exploits an efficient encoding of both temporal and spatial dynamics. In particular, we use a randomized recurrent neural network to embed the history of the input time series into high-dimensional state representations encompassing multi-scale temporal dynamics. Such representations are then propagated along the spatial dimension using different powers of the graph adjacency matrix to generate node embeddings characterized by a rich pool of spatiotemporal features. The resulting node embeddings can be efficiently pre-computed in an unsupervised manner, before being fed to a feed-forward decoder that learns to map the multi-scale spatiotemporal representations to predictions.

The training procedure can then be parallelized node-wise by sampling the node embeddings without breaking any dependency, thus enabling scalability to large networks. Empirical results on relevant datasets show that our approach achieves results competitive with the state of the art, while dramatically reducing the computational burden.

1 Introduction

As graph neural networks (GNNs; Scarselli et al. 2008; Bacciu et al. 2020) are gaining more traction in many application fields, the need for architectures scalable to large graphs – such as those associated with large sensor networks – is becoming a pressing issue. While research to improve the scalability of models for static graph signals has been very prolific (Hamilton, Ying, and Leskovec 2017; Chiang et al. 2019; Zeng et al. 2019; Frasca et al. 2020), little attention has been paid to the additional challenges encountered when dealing with discrete-time dynamical graphs, i.e., spatiotemporal time series. Several of the existing scalable training techniques rely on subsampling graphs to reduce the computational requirements of the training procedure, e.g., (Hamilton, Ying, and Leskovec 2017; Zeng et al. 2019). However, sampling the node-level observations as if they were i.i.d. can break relational (spatial) dependencies in static graphs and it is even more problematic in the dynamic case, as dependencies occur also across the temporal dimension. Indeed, complex temporal and spatial dynamics that emerge from the interactions across the whole graph over a long time horizon, can be easily disrupted by perturbing such spatiotemporal structure with subsampling. As an alternative, precomputing aggregated features over the graph allows for factoring out spatial propagation from the training phase in certain architectures (Frasca et al. 2020). However, similarly to the subsampling approach, extending this method to the spatiotemporal case is not trivial as the preprocessing step must account also for the temporal dependencies besides the graph topology.

In this paper, we propose a novel scalable encoder-
decoder architecture for processing spatiotemporal data; Fig. 1 shows high-level schematics of the proposed approach. The spatiotemporal encoding scheme is training-free: first, it exploits a deep randomized recurrent neural network (Jaeger 2001; Gallicchio, Micheli, and Pedrelli 2017) to encode the history of each sequence in a high-dimensional vector embedding; then, it uses powers of the graph adjacency matrix to build informative node representations of the spatiotemporal dynamics at different scales. According to the downstream task at hand, the decoder maps the node representations into the desired output, e.g., the future values of the time series associated with each node. To improve efficiency, we exploit the structure of the extracted embedding to design the decoder to act as a collection of filters localized at different spatiotemporal scales.

Since the spatiotemporal encoder requires neither training nor supervision, the representation of each node and time step can be computed in a preprocessing stage, without the constraints that come from online training on GPUs with limited memory. The decoder is the only component of the architecture with trainable parameters. However, since spatiotemporal relationships are already embedded in the representations, the embeddings can be processed independently from their spatiotemporal context with two consequent advantages. First, training can be done node-wise, allowing for sampling node representations in mini-batches of a size proportional to the hardware capacity. Second, the decoder can be implemented similarly to a standard multilayer perceptron (MLP) readout, which is fast and easy to train.

Let $T$ and $E$ be the number of steps and the number of edges in the input graph, respectively. The cost of training a standard spatiotemporal GNN on a mini-batch of data has a computational and memory cost that scales as $O(TE)$, or $O(T^3E)$ in attention-based architectures (Wu et al. 2022). Conversely, in our approach mini-batches can be sampled disregarding the length of the sequence and size of the graph, thus making scalability in training constant, i.e., $O(1)$, w.r.t. the spatiotemporal dimension of the problem.

Our contributions can be summarized as follows.

- We propose a general scalable deep learning framework for spatiotemporal time series, which exploits a novel encoding method based on randomized recurrent components and scalable GNNs architectures.
- We apply the proposed model to forecast multivariate time series, whose channels are subject to spatial relationships described by a graph.
- We carry out a rigorous and extensive empirical evaluation of the proposed architecture and variations thereof. Notably, we introduce two benchmarks for scalable spatiotemporal forecasting architectures.

Empirical results show that our approach performs on par with the state of the art while being easy to implement, computationally efficient, and extremely scalable. Given these considerations, we refer to our architecture as Scalable Graph Predictor (SGP).

2 Preliminaries and Problem Definition

We consider discrete-time spatiotemporal graphs. In particular, given $N$ interlinked sensors, we indicate with $x^i_t \in \mathbb{R}^{d_x}$ the $d_x$-dimensional multivariate observation associated with the $i$-th sensor at time-step $t$, with $X^i_t \in \mathbb{R}^{N \times d_x}$ the node attribute matrix encompassing measurements graph-wise, and with $X^{i:t+T}_t$ the sequence of $T$ measurements collected in the time interval $[t, t + T]$ at each sensor. Similarly, we indicate with $U^i_t \in \mathbb{R}^{N \times d_u}$ the matrix containing exogenous variables (e.g., weather information related to a monitored area) associated with each sensor at the $t$-th time-step. Then, we indicate additional, optional, static node attributes as $V \in \mathbb{R}^{N \times d_v}$. The relational information is encoded in a, potentially dynamic, weighted adjacency matrix $A^i_t \in \mathbb{R}^{N \times N}$. We indicate with the tuple $G^i_t = (X^i_t, U^i_t, V, A^i_t)$ the graph signal at the $t$-th time-step. Note that the number of sensors in a network is here considered fixed only to ease the presentation; we only request nodes to be distinguishable across time steps. The objective of spatiotemporal forecasting is to predict the next $H$ observations given a window of $W$ past measurements. In particular, we consider the family of forecasting models $F_\theta(\cdot)$ s.t.

$$\hat{X}^{i:t+H}_t = F_\theta(G^i_{t-W:t}),$$

where $\theta$ indicates the learnable parameters of the model and $\hat{X}^{i:t+H}_t$ the $H$-step ahead point forecast.

Echo-State Networks Echo state networks (Jaeger 2001; Lukoševičius and Jaeger 2009) are a class of randomized architectures that consist of recurrent neural networks with random connections that encode the history of input signals into a high-dimensional state representation to be used as input to a (trainable) readout layer. The main idea is to feed an input signal into a high-dimensional, randomized, and non-linear reservoir, whose internal state can be used as an embedding of the input dynamics. An echo state network is governed by the following state update equation:

$$h_t = \sigma(W_x x_t + W_h h_{t-1} + b),$$

where $x_t$ indicates a generic input to the system, $W_x \in \mathbb{R}^{d_x \times d_x}$ and $W_h \in \mathbb{R}^{d_h \times d_h}$ are the random matrices defining the connectivity pattern in the reservoir, $b \in \mathbb{R}^{d_h}$ is a randomly initialized bias, $h_t$ indicates the reservoir state, and $\sigma$ is a nonlinear activation function (usually tanh). If the random matrices are defined properly, the reservoir will extract a rich pool of dynamics characterizing the system underlying the input time series $x_t$, and, thus, the reservoir states become informative embeddings of $x_{t-T:t}$ (Lukoševičius and Jaeger 2009). Thanks to the non-linearity of the reservoir, the embeddings are commonly processed with a linear readout that is optimized with a least squares procedure to perform classification, clustering, or time series forecasting (Bianchi et al. 2020).

3 Scalable Spatiotemporal GNNs

This section presents our approach to building scalable GNN architectures for time series forecasting. Our method is
Figure 2: Overview of the SGP encoder. Input time series are fed into a randomized network with recurrent connections and embedded into a hierarchical vector representation. A graph shift operator is used to propagate and aggregate spatial information of different order which is then concatenated to obtain a final embedding.

based on a hybrid encoder-decoder architecture. The encoder first constructs representations of the time series observed at each node by using a reservoir that accounts for dynamics at different time scales. Representations are further processed to account for spatial dynamics described by the graph structure. In particular, as shown on the right-hand side of Fig. 2, we use incremental powers of the graph adjacency matrix to propagate and aggregate information along the spatial dimension. Each power of the propagation matrix accounts for different scales of spatial dynamics. The final embedding is then built by concatenating representations for different propagation steps, thus resulting in a rich encoding of both spatial and temporal features.

The encoder does not need any training and, once computed, the embeddings can be uniformly sampled over time and space when training a nonlinear readout to perform H-step-ahead predictions. The straightforward choice for the decoder (i.e., readout) is to map the encodings to the outputs (i.e., predictions) by using a linear transformation or a standard MLP. However, to further enhance scalability, our decoder exploits the structure of the embedding to reduce the number of parameters and learn filters that are localized in time and space. As we will discuss in Sec. 3.2, this is done by learning separate weight matrices for each spatiotemporal scale.

The following subsections describe in detail each component of the architecture.

### 3.1 Spatiotemporal Encoder

We consider as temporal encoders deep echo state networks (DeepESN; Gallicchio, Micheli, and Pedrelli 2017) with leaky integrator neurons (Jaeger et al. 2007). In particular, we consider networks where the signal associated with each node is encoded by a stack of L randomized recurrent layers s.t.

\[
\begin{align*}
    h_{i}^{(0)} & = [x_{i}^{0} | u_{i}^{0}], \\
    h_{i}^{(l)} & = \tanh \left( W_{u}^{(l)} h_{i}^{(l-1)} + W_{h}^{(l)} h_{i}^{(l-1)} + b^{(l)} \right), \\
    h_{i}^{(l)} & = (1 - \gamma_{l}) h_{i}^{(l-1)} + \gamma_{l} h_{i}^{(l)}, \quad l = 1, \ldots, L,
\end{align*}
\]

where \( \gamma_{l} \in (0, 1] \) is a discount factor associated with \( l \)-th layer, \( W_{u}^{(l)} \in \mathbb{R}^{d_{hi} \times d_{hi-1}}, W_{h}^{(l)} \in \mathbb{R}^{d_{hi} \times d_{hi}}, b^{(l)} \in \mathbb{R}^{d_{hi}} \) are random weight matrices, \( h_{i}^{(l)} \) indicates the hidden state of the system w.r.t. the \( i \)-th node at the \( l \)-th layer, and \( \parallel \) indicates node-wise concatenation. As Eq. 3 shows, DeepESNs are a hierarchical stack of reservoir layers that, e.g., by changing the discount factor at each layer, extract a rich pool of multi-scale temporal dynamics (Gallicchio, Micheli, and Pedrelli 2017). Given a DeepESN encoder, the input is represented by the concatenation of the states from each layer, i.e., we obtain node-level temporal encodings \( \vec{h}_{i}^{t} \) for each node \( i \) and time-step \( t \) as

\[
\vec{h}_{i}^{t} = (h_{i}^{(0)} || h_{i}^{(1)} || \ldots || h_{i}^{(L)}) .
\]

We indicate as \( \mathbf{H}_{t} \) the encoding for the whole graph at time \( t \). The extraction of the node-level temporal embeddings is depicted on the left-end side of Fig. 2, where, to simplify the drawing, we depict an ESN with a single layer.

The next step is to propagate information along the spatial dimension. As discussed at the beginning of the section, we use powers of a graph shift operator \( \tilde{A} \) to propagate and aggregate node representations at different scales. By using a notation similar to Eq. 4, we obtain spatiotemporal encodings as

\[
\begin{align*}
    S_{t}^{(0)} & = \mathbf{H}_{t} = (H_{t}^{(0)} || H_{t}^{(1)} || \ldots || H_{t}^{(L)}), \\
    S_{t}^{(k)} & = \tilde{A} S_{t}^{(k-1)} = (\tilde{A}^{k} H_{t}^{(0)} || \tilde{A}^{k} H_{t}^{(1)} || \ldots || \tilde{A}^{k} H_{t}^{(L)}), \\
    \bar{S}_{t} & = (S_{t}^{(0)} || S_{t}^{(1)} || \ldots || S_{t}^{(K)}) ,
\end{align*}
\]

where \( \tilde{A} \) indicates a generic graph shift operator matching the sparsity pattern of the graph adjacency matrix. In practice, by indicating with \( D \) the graph degree matrix, we use \( \tilde{A} = D^{-1/2} A D^{-1/2} \) in the case of a directed graph and the symmetrically normalized adjacency \( A = D^{-1/2} A D^{-1/2} \) in the undirected case. Furthermore, for directed graphs we optionally increase the number of representations to \( 2K + 1 \).

\[\text{We refer to (Gallicchio, Micheli, and Pedrelli 2018) for more details on the properties and stability of DeepESNs.}\]
3.2 Multi-Scale Decoder

The role of the decoder is that of selecting and weighing from the pool of (possibly redundant) features extracted by the spatiotemporal encoder and mapping them to the desired output. Representations $\mathbf{S}_t$ can be fed into an MLP that performs node-wise predictions. Since the representations are large vectors, a naive implementation of the MLP results in many parameters that hinder scalability. Therefore, we replace the first MLP layer with a more efficient implementation that exploits the structure of the embeddings.

As described in Sec. 3.1, $\mathbf{S}_t$ is the concatenation of the representations corresponding to different spatial propagation steps which, in turn, are obtained from the concatenation of multi-scale temporal features. To exploit this structure, we design the first layer of the decoder with a sparse connectivity pattern to learn representations

$$Z_t^{(k)} = \sigma \left( \mathbf{A}_k \mathbf{H}_1^{(0)} \Theta_k^{(0)} \| \ldots \| \mathbf{A}_k \mathbf{H}_L^{(L)} \Theta_k^{(L)} \right),$$

where $\Theta_k^{(l)} \in \mathbb{R}^{d_l \times d_k}$ are the learnable parameters and $\sigma$ is an activation function. In practice, representations $\mathbf{Z}_t$ can be efficiently computed by exploiting grouped 1-d convolutions (e.g., see Krizhevsky, Sutskever, and Hinton 2012) to parallelize computation on GPUs. In particular, if we indicate the 1-d grouped convolution operator with $g$ groups and kernel size $r$ as $\star_{r,g}$, and the collection of the decoder parameters $\Theta_k^{(l)}$ as $\Theta$ we can compute $\mathbf{Z}_t$ as

$$\mathbf{Z}_t = \sigma \left( \Theta \star_{1,g} \mathbf{S}_t \right),$$

with $g = L(K + 1)$ in the case of undirected graphs and $g = L(2K + 1)$ for the directed case. Besides reducing the number of parameters by a factor of $L(K + 1)$, this architecture localizes filters $\Theta_k^{(L)}$ w.r.t. the dynamics of spatial order $k$ and temporal scale $l$. In fact, as highlighted in Eq. 6–8, representations $\mathbf{Z}_t$ can be seen as a concatenation of the results of $L(K + 1)$ graph convolutions of different order. Finally, the obtained representations are fed into an MLP that predicts the $H$-step-ahead observations as

$$\tilde{x}_{t:t+H}^i = \text{MLP} \left( \mathbf{z}_t^i, v^i \right),$$

where the static node-level attributes $v^i$ can also be augmented by concatenating a set of learnable parameters (i.e., a learnable positional encoding).

4 Related Works

Spatiotemporal GNNs are essentially based on the idea of integrating message-passing modules in architectures to process sequential data. Notably, Seo et al. (2018) and Li et al. (2018) use message-passing to implement gates of recurrent neural networks. Yu, Yin, and Zhu (2018) and Wu et al. (2019, 2020) proposed architectures alternating temporal and spatial convolutions. Wu et al. (2022) and Marisca, Cini, and Alippi (2022), instead, exploit the attention mechanism to propagate information along both time and space. Modern architectures often combine some type of relational inductive bias, with full Transformer-like attention (Vaswani et al. 2017) along the spatial dimension (Zheng et al. 2020; Oreshkin et al. 2021; Satorras, Rangapuram, and Januschowski 2022), which, however, makes the computation scale quadratically with the number of nodes. SGP falls within the category of time-then-graph models, i.e., models where the temporal information is encoded before being propagated along the spatial dimension. Gao and Ribeiro (2022) showed that such models can be more expressive than architectures that alternate temporal and spatial processing steps.

Research on scalable models for discrete-time dynamic graphs has been relatively limited. Practitioners have mostly relied on methods developed in the context of static graphs which include node-centric, GraphSAGE-like, approaches (Hamilton, Ying, and Leskovec 2017) or subgraph sampling methods, such as ClusterGCN (Chiang et al. 2019) or GraphSAINT (Zeng et al. 2019). Wu et al. (2020); Gandhi et al. (2021); Wu et al. (2021) are examples of such approaches. Among scalable GNNs for static graphs, SIGN (Frasca et al. 2020) is the approach most related to our method. Like in our approach, SIGN performs spatial propagation in a preprocessing step by using different shift operators to aggregate across different graph neighborhoods, which are then fed to an MLP. However, SIGN is limited to static graphs and propagates raw node-level attributes.
Finally, similar to our work, DynGESN (Micheli and Torrerella 2022) processes dynamical graphs with a recurrent randomized architecture. However, the architecture in DynGESN is completely randomized, while ours is hybrid as it combines randomized components in the encoder with trainable parameters in the decoder.

5 Empirical Evaluation

We empirically evaluate our approach in 2 different scenarios. In the first one, we compare the performance of our forecasting architecture against state-of-the-art methods on popular, medium-scale, traffic forecasting benchmarks. In the second one, we evaluate the scalability of the proposed method on large-scale spatiotemporal time series datasets by considering two novel benchmarks for load forecasting and PV production prediction. Further details on datasets, baselines, and experimental settings are detailed in the supplemental material. We provide an efficient open-source implementation of SGP together with the code to reproduce all the experiments.

Datasets

In the first experiment we consider the METR-LA and PEMS-BAY datasets (Li et al. 2018), which are popular medium-sized benchmarks used in the spatiotemporal forecasting literature. In particular, METR-LA consists of traffic speed measurements taken every 5 minutes by 207 detectors in the Los Angeles County Highway, while PEMS-BAY includes analogous observations recorded by 325 sensors in the San Francisco Bay Area. We use the same preprocessing steps of previous works to extract a graph and obtain train, validation and test data splits (Wu et al. 2019). For the second experiment, we introduce two larger-scale datasets derived from energy analytics data. The first dataset contains data coming from the Irish Commission for Energy Regulation Smart Metering Project (CER-E; Commission for Energy Regulation 2016), which has been previously used for benchmarking spatiotemporal imputation methods (Cini, Marisca, and Alippi 2022); however, differently from previous works, we consider the full sensor network consisting of 6345 smart meters measuring energy consumption every 30 minutes at both residential and commercial/industrial premises. The second large-scale dataset is obtained from the synthetic PV-US dataset (Hummon et al. 2012), consisting of simulated energy production by 5016 PV farms scattered over the United States given historic weather data for the year 2006, aggregated in half an hour intervals. Since the model does not have access to weather information, PV production at neighboring farms is instrumental for obtaining good predictions. Notably, CER-E and PV-US datasets are at least an order of magnitude larger than the datasets typically used for benchmarking spatiotemporal time series forecasting models. Note that for both PV-US and CER-En the (weighted) adjacency is obtained by applying a thresholded Gaussian kernel to the similarity matrix obtained by considering the geographic distance among the sensors and the correntropy (Liu, Pokharel, and Principe 2007) among the time series, respectively. We provide further details on the datasets in the supplemental material.

Baselines

We consider the following baselines:

1. LSTM: a single standard gated recurrent neural network (Hochreiter and Schmidhuber 1997) trained by sampling window of observations from each node-level time series by disregarding the spatial information;

2. FC-LSTM: an LSTM processing input sequences as if they were a single high-dimensional multivariate time series;

3. GESN is completely randomized, while ours is hybrid as it combines randomized components in the encoder with trainable parameters in the decoder.

Table 1: Results on benchmark traffic datasets (averaged over 3 independent runs). We report MAE, MSE, and MAPE averaged over a one-hour (12 steps) forecasting horizon. We also show MAE for $H \in \{15, 30, 60\}$ minutes time horizons. Bold numbers are within a standard deviation from the best average.
Table 3: Additional information on the considered datasets.

| Dataset              | # steps | # nodes | # edges | sparsity |
|----------------------|---------|---------|---------|----------|
| METR-LA              | 34272   | 207     | 1515    | 3.54%    |
| PEMS-BAY             | 52116   | 325     | 2369    | 2.24%    |
| PV-US (100mn)        | 8868    | 5016    | 417,199 | 1.66%    |
| CER-En (100mn)       | 8868    | 6435    | 639,369 | 1.54%    |
| PV-US                | 8868    | 5016    | 3,710,008 | 14.75%   |
| CER-En               | 8868    | 6435    | 3,186,369 | 7.69%    |

Table 2: Results on large-scale datasets (averaged over at least 3 independent runs). We report MAE over $H$-step-ahead predictions, $H = \{30\text{m}, 7h30\text{m}, 11\text{h}\}$, together with timings and memory consumption. *indicates that subsampling was needed to comply with the memory constraints. Bold numbers are within a standard deviation from the best average.

|                 | Prediction error (MAE) | Resource utilization | Prediction error (MAE) | Resource utilization |
|-----------------|------------------------|----------------------|------------------------|----------------------|
|                 | 30m  | 7h30m | 11h | Batch/s | Memory | Batch | 30m  | 7h30m | 11h | Batch/s | Memory | Batch |
| DCRRN           | 1.39±0.09 | 3.34±0.22 | 3.54±0.48 | 2.04±0.01 | 9.63 GB | 11.14 GB | 1.50±0.08 | 2.94±0.25 | 3.12±0.14 | 8.41±0.09 | 11.46 GB | 11.70 GB |
| GWNet           | 1.45±0.13 | 5.09±0.63 | 5.26±1.13 | 2.01±0.02 | 11.64 GB | 11.35 GB | 1.52±0.06 | 2.05±0.01 | 2.00±0.01 | 8.83±0.09 | 11.35 GB | 11.64 GB |
| UG-GNN          | 1.33±0.08 | 2.94±0.25 | 3.12±0.14 | 8.41±0.09 | 11.46 GB | 11.70 GB | 1.52±0.06 | 2.05±0.01 | 2.00±0.01 | 8.83±0.09 | 11.35 GB | 11.64 GB |

3. **DCRRN**: a recurrent graph network presented in (Li et al. 2018) – differently from the original model we use a recurrent encoder followed by a linear readout (more details in the appendix);

4. **Graph WaveNet (GWNet)**: a residual network that alternates temporal and graph convolutions over the graph that is given as input and an adjacency matrix that is learned by the model (Wu et al. 2019);

5. **GatedGN (GGN)**: a state-of-the-art time-then-graph (Gao and Ribeiro 2022) model introduced in (Satorras, Rangapuram, and Januschowski 2022) for which we consider two different configurations. The first one (FC) uses attention over the full node set to perform spatial propagation, while the second one (UG) constrains the attention to edges of the underlying graph.

6. **DynGESN**: the echo state network for dynamical graphs proposed in (Micheli and Tortorella 2022).

For all the baselines, we use, whenever possible, the configuration found in the original papers or in their open-source implementation; in all the other cases we tune hyperparameters on the holdout validation set.

**Experimental setup** For the traffic datasets, we replicate the setup used in previous works. In particular, each model is trained to predict the 12-step-ahead observations. In SGP, the input time series are first encoded by the spatiotemporal encoder, and then the decoder is trained by sampling mini-batches along the temporal dimension, i.e., by sampling $B$ sequences $G_{-W:j}$ of observations.

For the large-scale datasets, we focus on assessing the scalability of the different architectures rather than maximizing forecasting accuracy. In particular, for both datasets, we consider the first 6 months of data (4 months for training, 1 month for validation, and 1 month for testing). The models are trained to predict the next $\{00:30, 07:30, 11:00\}$ hours. We repeat the experiment in two different settings to test the scalability of the different architectures w.r.t. the number of edges. In the first setting, we extract the graph by sparsifying the graph adjacency matrix imposing a maximum of 100 neighbors for each node, while, in the second case, we do not constrain the density of the adjacency matrix. Tab. 3 reports some details for the considered benchmarks. To assess the performance in terms of scalability, we fix a maximum GPU memory budget of 12 GB and select the batch size accordingly; if a batch size of 1 does not fit in 12 GB, we uniformly subsample edges of the graph to reduce the memory consumption. Differently from the other baselines, in SGP we first preprocess the data to obtain spatiotemporal embeddings and then train the decoder by uniformly sampling the node representations. We train each model for 1 hour, then restore the weights corresponding to the minimum training error and evaluate the forecasts on the test set. The choice of not running validation at each epoch was dictated by the fact that for some of the baselines running a validation epoch would take a large portion of the 1 hour budget.

The time required to encode the datasets with SGP’s encoder ranges from tens of seconds to $\approx 4$ minutes on an AMD EPYC 7513 processor with 32 parallel processes. To ensure reproducibility, the time constraint is not imposed as a hard time out; conversely, we measure the time required for the update step of each model on an NVIDIA RTX A5000 GPU and fix the maximum number of updates accordingly. For SGP, the time required to compute node embeddings was considered as part of the training time and the number of updates was appropriately reduced to make the comparison fair. For all the baselines, we keep the same architecture.
The bottom of Tab. 1 reports results for the ablation of key elements of the proposed architecture. No-Graph indicates that the embeddings are built without the spatial propagation step; FC-Dec. consider the case where the structure of the embedding is ignored in the readout and the sparse weight matrix in Eq. 7 is replaced by a fully-connected one; G-Dec. indicates that the spatial propagation is limited to the neighbors of order $K = 1$ and, thus, the decoder behaves similarly to a single-layer graph convolutional network. Results clearly show the optimality of the proposed architectural design.

Large-scale experiment Tab. 2 reports the results of the scalability experiment where we considered only the spatiotemporal GNNs trained by gradient descent. We excluded the full-attention baseline (FC-GatedGN) as its $O(N^2)$ complexity prevented scaling to the larger datasets; however, we considered the UG version where attention is restrained to each node’s neighborhood. There are several comments that need to be made here. First of all, batch size has a different meaning for our model and the other baselines. In our case, each sample corresponds to a single spatiotemporal (preprocessed) observation; for the other methods, a sample corresponds to a window of observations $G_t - W: t$ where edges of the graph are eventually subsampled if the memory constraints could not be met otherwise. In both cases, the loss is computed w.r.t. all the observations in the batch. The results clearly show that SGP can be trained efficiently also in resource-constrained settings, with contained GPU memory usage. In particular, the update frequency (batch/s) is up to 2 order of magnitude higher. Notably, resource utilization at training time remains constant (by construction) in the two considered scenarios, while almost all the baselines require edge subsampling in order to meet the resource constraints. Fig. 3 shows learning curves for the PV-US dataset, further highlighting the vastly superior efficiency, scalability, and learning stability of SGP. Finally, results concerning forecasting accuracy show that performance is competitive with the state of the art in all the considered scenarios.

6 Remarks and Conclusion

We proposed SGP, a scalable architecture for graph-based spatiotemporal time series forecasting. Our approach competes with the state of the art in popular medium-sized benchmark datasets, while greatly improving scalability in large sensor networks. While in SGP sampling largely reduces GPU memory usage compared to the other methods, the entire processed sequence can take up a large portion of system memory, depending on the size of the reservoir. Nevertheless, the preprocessing can be distributed, the preprocessed data stored on disk and loaded in batches during training, as customary for large datasets. We believe that SGP constitutes an important stepping stone for future research on scalable spatiotemporal forecasting and has the potential of being widely adopted by practitioners. Future work can explore a tighter integration of the spatial and temporal encoding components and assess performance on even larger benchmarks.
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