Scalable Deep Generative Modeling for Sparse Graphs

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Abstract

Learning graph generative models is a challenging task for deep learning and has wide applicability to a range of domains like chemistry, biology and social science. However, current deep neural methods suffer from limited scalability: for a graph with \( n \) nodes and \( m \) edges, existing deep neural methods require \( \Omega(n^2) \) complexity by building up the adjacency matrix. On the other hand, many real world graphs are actually sparse in the sense that \( m \ll n^2 \). Based on this, we develop a novel autoregressive model, named BiGG, that utilizes this sparsity to avoid generating the full adjacency matrix, and importantly reduces the graph generation time complexity to \( O((n+m) \log n) \). Furthermore, during training this autoregressive model can be parallelized with \( O(\log n) \) synchronization stages, which makes it much more efficient than other autoregressive models that require \( \Omega(n) \). Experiments on several benchmarks show that the proposed approach not only scales to orders of magnitude larger graphs than previously possible with deep autoregressive graph generative models, but also yields better graph generation quality.

1. Introduction

Representing a distribution over graphs provides a principled foundation for tackling many important problems in knowledge graph completion (Xiao et al., 2016), de novo drug design (Li et al., 2018; Simonovsky & Komodakis, 2018), architecture search (Xie et al., 2019) and program synthesis (Brockschmidt et al., 2018). The effectiveness of graph generative modeling usually depends on learning the distribution given a collection of relevant training graphs. However, training a generative model over graphs is usually quite difficult due to their discrete and combinatorial nature.

Classical generative models of graphs, based on random graph theory (Erdős & Rényi, 1960; Barabási & Albert, 1999; Watts & Strogatz, 1998), have been long studied but typically only capture a small set of specific graph properties, such as degree distribution. Despite their computational efficiency, these distribution models are usually too inexpressive to yield competitive results in challenging applications. Recently, deep graph generative models that exploit the increased capacity of neural networks to learn more expressive graph distributions have been successfully applied to real-world tasks. Prominent examples include VAE-based methods (Kipf & Welling, 2016; Simonovsky & Komodakis, 2018), GAN-based methods (Bojchevski et al., 2018), flow models (Liu et al., 2019; Shi et al., 2020) and autoregressive models (Li et al., 2018; You et al., 2018; Liao et al., 2019). Despite the success of these approaches in modeling small graphs, e.g. molecules with hundreds of nodes, they are not able to scale to graphs with over 10,000 nodes.

A key shortcoming of current deep graph generative models is that they attempt to generate a full graph adjacency matrix, implying a computational cost of \( \Omega(n^2) \) for a graph with \( n \) nodes and \( m \) edges. For large graphs, it is impractical to sustain such a quadratic time and space complexity, which creates an inherent trade-off between expressiveness and efficiency. To balance this trade-off, most recent work has introduced various conditional independence assumptions (Liao et al., 2019), ranging from the fully auto-regressive but slow GraphRNN (You et al., 2018), to the fast but fully factorial GraphVAE (Simonovsky & Komodakis, 2018).

In this paper, we propose an alternative approach that does not commit to explicit conditional independence assumptions, but instead exploits the fact that most interesting real-world graphs are sparse, in the sense that \( m \ll n^2 \). By leveraging sparsity, we develop a new graph generative model, BiGG (BiG Graph Generation), that streamlines the generative process and avoids explicit consideration of every entry in an adjacency matrix. The approach is based on three key elements: (1) an \( O(\log n) \) process for generating each edge using a binary tree data structure, inspired by R-MAT (Chakrabarti et al., 2004); (2) a tree-structured autoregressive model for generating the set of edges associated with each node; and (3) an autoregressive model defined over the sequence of nodes. By combining these elements, BiGG can generate a sparse graph in \( O((n+m) \log n) \) time.
which is a substantial improvement over $\Omega(n^2)$.

For training, the design of BiGG also allows every context embedding in the autoregressive model to be computed in only $O(\log n)$ sequential steps, which enables significant gains in training speed through parallelization. By comparison, the context embedding in GraphRNN requires $O(n^2)$ sequential steps to compute, while GRAN requires $O(n)$ steps. In addition, we develop a training mechanism that only requires sublinear memory cost, which in principle makes it possible to train models of graphs with millions of nodes on a single GPU.

On several benchmark datasets, including synthetic graphs and real-world graphs of proteins, 3D mesh and SAT instances, BiGG is able to achieve comparable or superior sample quality than the previous state-of-the-art, while being orders of magnitude more scaleable.

To summarize the main contributions of this paper:

- We propose an autoregressive generative model, BiGG, that can generate sparse graphs in $O((n+m) \log n)$ time, successfully modeling graphs with 100k nodes on 1 GPU.
- The training process can be largely parallelized, requiring only $O(\log n)$ steps to synchronize learning updates.
- Memory cost is reduced to $O(\sqrt{m \log n})$ for training and $O(\log n)$ for inference.
- BiGG not only scales to orders of magnitude larger graphs than current deep models, it also yields comparable or better model quality on several benchmark datasets.

Other related work There has been a lot of work (Chakrabarti et al., 2004; Robins et al., 2007; Leskovec et al., 2010; Airoldi et al., 2009) on generating graphs with a set of specific properties like degree distribution, diameter, and eigenvalues. All these classical models are hand-engineered to model a particular family of graphs, and thus not general enough. Besides the general graphs, a lot of work also exploits domain knowledge for better performance in specific domains. Examples of this include (Kusner et al., 2017; Dai et al., 2018; Jin et al., 2018; Liu et al., 2018) for modeling molecule graphs, and (You et al., 2019) for SAT instances. See Appendix A for more discussions.

2. Model

A graph $G = (V, E)$ is defined by a set of nodes $V$ and set of edges $E \subseteq V \times V$, where a tuple $e_i = (u, v) \in E$ is used to represent an edge between node $u$ and $v$. We denote $n = |V|$ and $m = |E|$ as the number of nodes and edges in $G$ respectively. Note that a given graph may have multiple equivalent adjacency matrix representations, with different node orderings. However, given an ordering of the nodes $\pi$, there is a one-to-one mapping between the graph structure $G$ and the adjacency matrix $A^\pi \in \{0, 1\}^{n \times n}$.

Our goal is to learn a generative model, $p(G)$, given a set of training graphs $\mathcal{D}_{\text{train}} = \{G_1, G_2, \ldots, G_{|\mathcal{D}_{\text{train}}|}\}$. In this paper, we assume the graphs are not attributed, and focus on the graph topology only. Such an assumption implies

$$p(G) = p(V)p(E|V) = p(|V| = n) \sum_\pi p(E, \pi|n) = p(|V| = n) \sum_\pi p(A^{\pi}),$$

where $p(E, \pi|n)$ is the probability of generating the set of edges $E$ under a particular ordering $\pi$, which is equivalent to the probability of a particular adjacency matrix $A^\pi$. Here, $p(|V| = n)$ is the distribution of number of nodes in a graph. In this paper, we use a single canonical ordering $\pi(G)$ to model each graph $G$, as in (Li et al., 2018):

$$p(G) \simeq p(|V| = n) p(A^{\pi(G)}),$$

which is clearly a lower bound on $p(G)$ (Liao et al., 2019).

We estimate $p(|V| = n)$ directly using the empirical distribution over graph size, and only learn an expressive model for $p(A)$. In the following presentation, we therefore omit the ordering $\pi$ and assume a default canonical ordering of nodes in the graph when appropriate.

As $A$ will be extremely sparse for large sparse graphs ($m \ll n^2$), generating only the non-zero entries in $A$, i.e., the edge set $E$, will be much more efficient than the full matrix:

$$p(A) = p(e_1)p(e_2|e_1)\ldots p(e_m|e_1, \ldots, e_{m-1}),$$

where each $e_i = (u, v)$ include the indices of the two nodes associated with one edge, resulting in a generation process of $m$ steps. We order the edges following the node ordering, hence this process generates all the edges for the first row in $A$, before generating the second row, etc. A naive approach to generating a single edge will be $O(n)$ if we factorize $p(e_i) = p(u)p(v|u)$ and assume both $p(u)$ and $p(v|u)$ to be simple multinomials over $n$ nodes. This, however, will not give us any benefit over traditional methods.

2.1. Recursive edge generation

The main scalability bottleneck is the large output space. One way to reduce the output space size is to use a hierarchical recursive decomposition, inspired by the classic random graph model R-MAT (Chakrabarti et al., 2004). In R-MAT, each edge is put into the adjacency matrix by dividing the 2D matrix into 4 equally sized quadrants, and recursively descending into one of the quadrants, until reaching a single entry of the matrix. In this generation process, the complexity of generating each edge is only $O(\log n)$. 

Figure 1. Recursive generation of the edge $e = (u, v)$ given $u$. 

We adopt the recursive decomposition design of R-MAT, and further simplify and neutralize it, to make our model efficient and expressive. In our model, we generate edges following the node ordering row by row, so that each edge only needs to be put into the right place in a single row, reducing the process to 1D, as illustrated in Figure 1. For any edge \((u, v)\), the process of picking node \(v\) as one of \(u\)’s neighbors starts by dividing the node index interval \([1, n]\) in half, then recursively descending into one half until reaching a single entry. Each \(v\) corresponds to a unique sequence of decisions \(x_1^t, ..., x_d^t\), where \(x_i^t \in \{\text{left, right}\}\) is the \(i\)-th decision in the sequence, and \(d = \lceil \log_2 n \rceil\) is the maximum number of required decisions to specify \(v\).

The probability of \(p(v|u)\) can then be formulated as
\[
p(v|u) = \prod_{i=1}^{\lceil \log_2 n \rceil} p(x_i = x_i^t),
\]
where each \(p(x_i = x_i^t)\) is the probability of following the decision that leads to \(v\) at step \(i\).

Let us use \(E_u = \{(u, v) \in E\}\) to denote the set of edges incident to node \(u\), and \(N_u = \{v \mid (u, v) \in E_u\}\). Generating only a single edge is similar to hierarchical softmax (Mnih & Hinton, 2009), and applying the above procedure repeatedly can generate all of \(|N_u|\) edges in \(O(|N_u| \log n)\) time. But we can do better than that when generating all these edges.

**Further improvement using binary trees.** As illustrated in the left half of Figure 2, the process of jointly generating all of \(E_u\) is equivalent to building up a binary tree \(T_u\), where each tree node \(t \in T_u\) corresponds to a graph node index interval \([v_l, v_r]\), and for each \(v \in N_u\), the process starts from the root \([1, n]\) and ends in a leaf \([v, v]\). Taking this perspective, we propose a more efficient generation process for \(E_u\), which generates the tree directly instead of repeatedly generating each leaf through a path from the root. We propose a recursive process that builds up the tree following a depth-first or in-order traversal order, where we start at the root, and recursively for each tree node \(t\): (1) decide if \(t\) has a left child denoted as \(lch(t)\), and (2) if so recurse into \(lch(t)\) and generate the left sub-tree, and then (3) decide if \(t\) has a right child denoted as \(rch(t)\), (4) if so recurse into \(rch(t)\) and generate the right sub-tree, and (5) return to \(t\)’s parent. This process is shown in Algorithm 1, which will be elaborated in next section.

Overloading the notation a bit, we use \(p(lch(t))\) to denote the probability that tree node \(t\) has a left child, when \(lch(t) \neq \emptyset\), or does not have a left child, when \(lch(t) = \emptyset\), under our model, and similarly define \(p(rch(t))\). Then the probability of generating \(E_u\) or equivalently tree \(T_u\) is
\[
p(E_u) = p(T_u) = \prod_{t \in T_u} p(lch(t))p(rch(t)).
\]
This new process generates each tree node exactly once, hence the time complexity is proportional to the tree size \(O(|T_u|)\), and it is clear that \(|T_u| \leq |N_u| \log n\), since \(|N_u|

### Algorithm 1 Generating outgoing edges of node \(u\)

1: \textbf{function} recursive\((u, t, h_u^t(op))\)
2: \hspace{1em} if \text{is leaf}(t) \textbf{then}
3: \hspace{2em} \text{Return } \tilde{I}, \{\text{edge index that } t \text{ represents}\}
4: \hspace{1em} \textbf{end if}
5: \hspace{1em} \text{has_left} \sim p(lch_u(t)|h_u^t(op)) \text{ using Eq. (8)}
6: \hspace{1em} \text{if has_left then}
7: \hspace{2em} \text{Create } lch_u(t) \text{, and let } h_u^{lch}(lch_u(t)), N_u^{lch} \leftarrow \text{recursive}(u, lch_u(t), h_u^t(op) (lch_u(t)))
8: \hspace{1em} \text{else}
9: \hspace{2em} \text{has_right} \sim p(rch_u(t)|h_u^t(op)(rch_u(t))) \text{ using Eq. (9)}
10: \hspace{2em} \text{if has_right then}
11: \hspace{3em} \text{Create } rch_u(t), \text{ and let } h_u^{rch}(rch_u(t)), N_u^{rch} \leftarrow \text{recursive}(u, rch_u(t), h_u^t(op)(rch_u(t)))
12: \hspace{2em} \text{else}
13: \hspace{3em} h_u^{bot}(rch_u(t)) \leftarrow \emptyset, N_u^{bot} = \emptyset
14: \hspace{2em} \text{end if}
15: \hspace{1em} \text{end if}
16: \hspace{1em} h_u^{bot}(lch_u(t)) \leftarrow \emptyset, N_u^{bot} = \emptyset
17: \hspace{1em} \text{end if}
18: \hspace{1em} h_u^{bot}(t) = \text{TreeCell}_{lch}(h_u^{lch}(lch_u(t)), h_u^{bot}(rch_u(t)))
19: \hspace{1em} N_u^t = N_u^{lch} \cup N_u^{rch}
20: \hspace{1em} \text{Return } h_u^{bot}(t), N_u^t
21: \textbf{end function}

is the number of leaf nodes in the tree and \(\log n\) is the max depth of the tree. The time saving comes from removing the duplicated effort near the root of the tree. When \(|N_u|\) is large, i.e., as some fraction of \(n\) when \(u\) is one of the “hub” nodes in the graph, the tree \(T_u\) becomes dense and our new generation process will be significantly faster, as the time complexity becomes closer to \(O(n)\) while generating each leaf from the root would require \(\Omega(n \log n)\) time.

In the following, we present our approach to make this model fully autoregressive, i.e. making \(p(lch(t))\) and \(p(rch(t))\) depend on all the decisions made so far in the process of generating the graph, and make this model neutralized so that all the probability values in the model come from expressive deep neural networks.

### 2.2. Autoregressive conditioning for generating \(T_u\)

In this section we consider how to add autoregressive conditioning to \(p(lch(t))\) and \(p(rch(t))\) when generating \(T_u\).

In our generation process, the decision about whether \(lch(t)\) exists for a particular tree node \(t\) is made after \(t\), all its ancestors, and all the left sub-trees of the ancestors are generated. We can use a top-down context vector \(h_u^{top}(t)\) to summarize all these contexts, and modify \(p(lch(t))\) to \(p(lch(t)|h_u^{top}(t))\). Similarly, the decision about \(rch(t)\) is made after generating \(lch(t)\) and its dependencies, and \(t\)’s entire left-subtree (see Figure 2 right half for illustration). We therefore need both the top-down context \(h_u^{top}(t)\), as well as the bottom-up context \(h_u^{bot}(lch(t))\) that summarizes the sub-tree rooted at \(lch(t)\), if any. The autoregressive
model for \( p(T_u) \) therefore becomes
\[
p(T_u) = \prod_{t \in T_u} p(lch(t)|h^\text{top}_u(t)) p(rch(t)|h^\text{top}_u(t), h^\text{bot}_u(lch(t))),
\]
and we can recursively define
\[
\begin{align*}
  h^\text{bot}_u(t) &= \text{TreeCell}^{\text{bot}}(h^\text{bot}_u(lch(t)), h^\text{bot}_u(rch(t))) \\
  h^\text{top}_u(lch(t)) &= \text{LSTMCell}(h^\text{top}_u(t), \text{embed(left)}) \\
  h^\text{top}_u(rch(t)) &= \text{LSTMCell}(h^\text{top}_u(t), \text{embed(right)}),
\end{align*}
\]
where TreeCell^{\text{bot}} and TreeCell^{\text{top}} are two TreeLSTM cells (Tai et al., 2015) that combine information from the incoming nodes into a single node state, and embed(left) and embed(right) represents the embedding vector for the binary values “left” and “right”. We initialize \( h^\text{bot}_u(\emptyset) = 0 \), and discuss \( h^\text{top}_u(\text{root}) \) in the next section.

The distributions can then be parameterized as
\[
\begin{align*}
  p(lch(t)|\cdot) &= \text{Bernoulli}(\sigma(W^l_i h^\text{top}_u(t) + b_i)), \\
  p(rch(t)|\cdot) &= \text{Bernoulli}(\sigma(W^r_i h^\text{top}_u(rch(t)) + b_r)).
\end{align*}
\]

### 2.3. Full autoregressive model

With the efficient recursive edge generation and autoregressive conditioning presented in Section 2.1 and Section 2.2 respectively, we are ready to present the full autoregressive model for generating the entire adjacency matrix \( A \).

The full model will utilize the autoregressive model for \( N_u \) as building blocks. Specifically, we are going to generate the adjacency matrix \( A \) row by row:
\[
p(A) = p(A|N_u) = \prod_{u \in V} p(A|N_u\{N_{u'} : u' < u\}).
\]

Let \( g^0_u = h^\text{bot}_u(t_1) \) be the embedding that summarizes \( T_u \), suppose we have an efficient mechanism to encode \( g_1, g_2, \ldots, g_u \) into \( h^\text{row}_u \), then we can effectively use \( h^\text{row}_u \) to generate \( T_u \) and thus the entire process would become autoregressive. Again, since there are \( n \) rows in total, using a chain structured LSTM would make the history length too long for large graphs. Therefore, we use an approach inspired by the Fenwick tree (Fenwick, 1994) which is a data structure that maintains the prefix sum efficiently. Given an array of numbers, the Fenwick tree allows calculating any prefix sum or updating any single entry in \( O(\log L) \) for a sequence of length \( L \). We build such a tree to maintain and update the prefix embeddings. We denote it as row-binary forest as such data structure is a forest of binary trees.

Figure 3 demonstrates one solution. Before generating the edge-binary tree \( T_u \), the embeddings that summarize each individual edge-binary tree \( R_u = \{T_{u'} : u' < u\} \) will be organized into the row-binary forest \( G_u \). This forest is organized into \( \lceil \log(u-1) \rceil + 1 \) levels, with the bottom 0-th level as edge-binary tree embeddings. Let \( g_j \in G_u \) be the \( j \)-th node in the \( i \)-th level, then
\[
g^i_j = \text{TreeCell}^{\text{row}}(g^{i-1}_{j+2^i-1}, g^{i-1}_{j+2^i}),
\]
where \( 0 \leq i \leq \lceil \log(u-1) \rceil + 1, 1 \leq j \leq \lfloor \frac{R_u}{2^i} \rfloor \).

#### Embedding row-binary forest

One way to embed this row-binary forest is to embed the root of each tree in this forest. As there will be at most one root in each level (otherwise the two trees in the same level will be merged into a larger tree for the next level), the number of tree roots will also be \( O(\log n) \) at most. Thus we calculate \( h^\text{row}_u \) as follows:
\[
h^\text{row}_u = \text{LSTM} \left( \left[ g^i_u \right]^{\left\lfloor \frac{R_u}{2^i} \right\rfloor} \right),
\]
where \( u \& 2^i = 2^i \).

Here, \( \& \) is the bit-level ‘and’ operator. Intuitively as in Fenwick tree, the calculation of any prefix sum of length \( L \) requires the block sums that corresponds to each binary digit in the binary bits representation of integer \( L \). Recall the operator \( h^\text{top}(\cdot) \) defined in Section 2.2, here \( h^\text{top}(\text{root}) = h^\text{row}_{u-1} \), when \( u > 1 \), and equals to zero when \( u = 1 \). With the embedding of \( G_u \) at each state served as ‘glue’, we can connect all the individual row generation modules in an autoregressive way.

#### Updating row-binary forest

It is also efficient to update such forest every time a new \( g^0_u \) is obtained after the generation of \( T_u \). Such an updating procedure is similar to the Fenwick tree update. As each level of this forest has at most one root node, merging in the way defined in (11) will
Algorithm 2: Generating graph using BiGG

1: function update_forest(u, G_{u-1}, g_u^0)
2: \( G_u = G_{u-1} \cup \{ g_u^0 \} \)
3: for \( i \leftarrow 0 \) to \( \lfloor \log(u-1) \rfloor \) do
4: \( j \leftarrow \arg \max_j \| g_j^i \in G_u \|
5: \text{if such } j \text{ exists and } j \text{ is an even number then}
6: \( g_{j+1}^{i+1} \leftarrow \text{TreeCellRow}(g_{j-1}^i, g_j^i) \)
7: \( G_u \leftarrow G_u \cup g_{j+1}^{i+1} \)
8: end if
9: end for
10: Update \( h_{row}^0 \) using Eq (12)
11: Return: \( h_{row}^u, G_u, \mathcal{R}_{u-1} \cup \{ g_0^u \} \)
12: end function

13: Input: The number of nodes \( n \)
14: \( h_{row}^0 \leftarrow 0, \mathcal{R}_0 = \emptyset, \mathcal{G}_0 = \emptyset, E = \emptyset \)
15: for \( u \leftarrow 1 \) to \( n \) do
16: \( t_1 \) be the root of an empty edge-binary tree
17: \( g_0^0, \mathcal{N}_0 \leftarrow \text{recursive}(u, t_1, h_{row}^0) \)
18: \( h_{row}^0, G_u, \mathcal{R}_u \leftarrow \text{update_forest}(u, \mathcal{R}_{u-1}, G_{u-1}, g_0^u) \)
19: end for
20: Return: \( G \) with \( V = \{ 1, \ldots, n \} \) and \( E = \cup_{u=1}^{n} \mathcal{N}_u \)

happen at most once per each level, which effectively makes the updating cost to be \( O(\log n) \).

Algorithm 2 summarizes the entire procedure for sampling a graph from our model in a fully autoregressive manner.

**Theorem 1** BiGG generates a graph with \( n \) nodes and \( m \) edges in \( O((n + m) \log n) \) time. In the extreme case where \( m \approx n^2 \), the overall complexity becomes \( O(n^2) \).

**Proof** The generation of each edge-binary tree in Section 2.2 requires time complexity proportional to the number of nodes in the tree. The Fenwick tree query and update both take \( O(\log n) \) time, hence maintaining the data structure takes \( O(\log n) \). The overall complexity is \( O(n \log n + \sum_{u=1}^{m} |\mathcal{T}_u|) \). For a sparse graph \( |\mathcal{T}_u| = O(|\mathcal{N}_u| \log n) \), hence \( \sum_{u=1}^{m} |\mathcal{T}_u| = \sum_{u=1}^{m} |\mathcal{N}_u| \log n = O(m \log n) \). For a complete graph, where \( m = O(n^2) \), each \( \mathcal{T}_u \) will be a full binary tree with \( n \) leaves, hence \( |\mathcal{T}_u| = 2n - 1 \) and the overall complexity would be \( O(n \log n + n^2) = O(n^2) \). \( \square \)

### 3. Optimization

In this section, we propose several ways to scale up the training of our auto-regressive model. For simplicity, we focus on how to speed up the training with a single graph. Training multiple graphs can be easily extended.

#### 3.1. Training with \( O(\log n) \) synchronizations

A classical autoregressive model like LSTM is fully sequential, which allows no concurrency across steps. Thus training LSTM with a sequence of length \( L \) takes \( \Omega(L) \) of

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**Figure 4.** Parallelizing computation during training. The four stages are executed sequentially. In each stage, the embeddings of nodes with the same color can be computed concurrently.

Given a graph \( G \), estimating the likelihood under the current model can be divided into four steps.

1. **Computing** \( h_{row}^{bot}(t), \forall u \in V, t \in \mathcal{T}_u : \) as the graph is given during training, the corresponding edge-binary trees \{\( \mathcal{T}_u \)\} are also known. From Eq (7) we can see that the embeddings \( h_{row}^{bot}(t) \) of all nodes at the same depth of a tree can be computed concurrently without dependence, and the synchronization only happens between different depths. Thus \( O(\log n) \) steps of synchronization is sufficient.

2. **Computing** \( g_j^i \in \mathcal{G}_u : \) the row-binary forest grows monotonically, thus the forest \( \mathcal{G}_u \) in the end contains all the embeddings needed for computing \( \{ h_{row}^u \} \). Similarly, computing \( \{ g_j^i \} \) synchronizes \( O(\log n) \) steps.

3. **Computing** \( h_{row}^{top}, \forall u \in V : \) as each \( h_{row}^u \) runs an LSTM independently, this stage simply runs LSTM on a batch of \( n \) sequences with length \( O(\log n) \).

4. **Computing** \( h_{row}^{top} \) and likelihood: the last step computes the likelihood using Eq (8) and (9). This is similar to the first step, except that the computation happens in a top-down direction in each \( \mathcal{T}_u \).

Figure 4 demonstrates this process. In summary, the four stages each take \( O(\log n) \) steps of synchronization. This allows us to train large graphs much more efficiently than a simple sequential autoregressive model.

#### 3.2. Model parallelism

It is possible that during training the graph is too large to fit into memory. Thus to train on large graphs, model parallelism is more important than data parallelism.

To split the model, as well as intermediate computations,
into different machines, we divide the adjacency matrix into multiple consecutive chunks of rows, where each machine is responsible for one chunk. It is easy to see that by doing so, Stage 1 and Stage 4 mentioned in Section 3.1 can be executed concurrently on all the machines without any synchronization, as the edge-binary trees can be processed independently once the conditioning states like \( \{ h_u^{row} \} \) are made ready by synchronization.

Figure 5. Model parallelism for training single graph. Red circled nodes are computed on GPU 1 but is required by GPU 2 as well.

Figure 5 illustrates the situation when training a graph with 7 nodes using 2 GPUs. During Stage 1, GPU 1 and 2 work concurrently to compute \( \{ g_u^1 \}_{u=1}^3 \) and \( \{ g_u^2 \}_{u=4}^7 \), respectively. In Stage 2, the embeddings \( g_u^1 \) and \( g_u^2 \) are needed by GPU 2 when computing \( g_u^2 \) and \( g_u^3 \). We denote such embeddings as ‘g-messages’. Note that such ‘g-messages’ will transit in the opposite direction when doing a backward pass in the gradient calculation. Passing ‘g-messages’ introduces serial dependency across GPUs. However as the number of such embeddings is upper bounded by \( O(\log n) \) depth of row-binary forest, the communication cost is manageable.

3.3. Reducing memory consumption

Sublinear memory cost: Another way to handle the memory issue when training large graphs is to recompute certain portions of the hidden layers in the neural network when performing backpropagation, to avoid storing such layers during the forward pass. Chen et al. (2016) introduces a computation scheduling mechanism for sequential structured neural networks that achieves \( O(\sqrt{L}) \) memory growth for an \( L \)-layer neural network.

We divide rows as in Section 3.2. During the forward pass, only the ‘g-messages’ between chunks are kept. The only difference from the previous section is that the edge-binary tree embeddings are recomputed, due to the single GPU limitation. The memory cost will be:

\[
O(\max \{ k \log n, \frac{m}{k} \})
\]

Here \( O(k \log n) \) accounts for the memory holding the ‘g-message’, and \( O(\frac{m}{k}) \) accounts for the memory of \( T_u \) in each chunk. The optimal \( k \) is achieved when \( k \log n = \frac{m}{k} \), hence \( k = O(\sqrt{\frac{m}{\log n}}) \) and the corresponding memory cost is \( O(\sqrt{m \log n}) \). Also note that such sublinear cost requires only one additional feedforward in Stage 1, so this will not hurt much of the training speed.

3.4. Position encoding:

During generation of \( \{ T_u \} \), each tree node \( t \) of the edge-binary tree knows the span \([v_l, v_r] \) which corresponds to the columns it will cover. One way to augment \( h_u^{top} \) with the position encoding as:

\[
h_u^{top}(t) = h_u^{top}(t) + PE(v_r - v_l)
\]

where PE is the position encoding using sine and cosine functions of different frequencies as in Vaswani et al. (2017). Similarly, the \( h_u^{row} \) in Eq (12) can be augmented by PE\((n - u) \) in a similar way. With such augmentation, the model will know more context into the future, and thus help improve the generative quality.

Please refer to our released open source code located at https://github.com/google-research/google-research/tree/master/bigg for more implementation and experimental details.

4. Experiment

4.1. Model Quality Evaluation on Benchmark Datasets

In this part, we compare the quality of our model with previous work on a set of benchmark datasets. We present results on median sized general graphs with number of nodes ranging in 0.1k to 5k in Section 4.1.1, and on large SAT graphs with up to 20k nodes in Section 4.1.2. In Section 4.3 we perform ablation studies of BiGG with different sparsity and node orders.
4.1.1. General graphs

The general graph benchmark is obtained from Liao et al. (2019) and part of it was also used in (You et al., 2018). This benchmark has four different datasets: (1) Grid, 100 2D grid graphs; (2) Protein, 918 protein graphs (Dobson & Doig, 2003); (3) Point cloud, 3D point clouds of 41 household objects (Neumann et al., 2013); (4) Lobster, 100 random Lobster graphs (Golomb, 1996), which are trees where each node is at most 2 hops away from a backbone path. Table 1 contains some statistics about each of these datasets. We use the same protocol as Liao et al. (2019) that splits the graphs into training and test sets.

Baselines: We compare with deep generative models including GraphVAE (Simonovsky & Komodakis, 2018), GraphRNN, GraphRNN-S (You et al., 2018) and GRAN (Liao et al., 2019). We also include the Erdős–Rényi random graph model that only estimates the edge density. Since our setups are exactly the same, the baseline results are directly copied from Liao et al. (2019).

Evaluation: We use exactly the same evaluation metric as Liao et al. (2019), which compares the distance between the distribution of held-out test graphs and the generated graphs. We use maximum mean discrepancy (MMD) with four different test functions, namely the node degree, clustering coefficient, orbit count and the spectra of the graphs from the eigenvalues of the normalized graph Laplacian. Besides the four MMD metrics, we also use the error rate for Lobster dataset. This error rate reflects the fraction of generated graphs that doesn’t have Lobster graph property.

Results: Table 1 reports the results on all the four datasets. We can see the proposed BiGG outperforms all other methods on all the metrics. The gain becomes more significant on the largest dataset, i.e., the 3D point cloud. While GraphVAE and GraphRNN gets out of memory, the orbit metric of BiGG is 2 magnitudes better than GRAN. This dataset reflects the scalability issue of existing deep generative models. Also from the Lobster graphs we can see, although GRAN scales better than GraphRNN, it yields worse quality due to its approximation of edge generation with mixture of conditional independent distributions. Our BiGG improves the scalability while also maintaining the expressiveness.

4.1.2. SAT graphs

In addition to comparing with general graph generative models, in this section we compare against several models that are designated for generating the Boolean Satisfiability (SAT) instances. A SAT instance can be represented using bipartite graph, i.e., the literal-clause graph (LCG). For a SAT instance with \( n_x \) variables and \( n_c \) clauses, it creates \( n_x \) positive and negative literals, respectively. The canonical node ordering assigns 1 to \( 2 \times n_x \) for literals and \( 2 \times n_x + 1 \) to \( 2 \times n_x + n_c \) for clauses. The following experiment largely follows G2SAT (You et al., 2019). We use the train/test split of SAT instances obtained from G2SAT website. This result in 24 and 8 training/test SAT instances, respectively. The size of the SAT graphs ranges from 491 to 21869 nodes. Note that the original paper reports results using 10 small training instances instead. For completeness, we also include such results in Appendix B.3 together with other baselines from You et al. (2019).

Baseline: We mainly compare the learned model with G2SAT, a specialized deep graph generative model for bipartite SAT graphs. Since BiGG is general purposed, to guarantee the generated adjacency matrix \( A \) is bipartite, we let our model to generate the upper off-diagonal block of the adjacency matrix only, i.e., \( A[0 : 2 \times n_x, 2 \times n_x : 2 \times n_x + n_c] \). G2SAT requires additional ‘template graph’ as input when generating the graph. Such template graph is equivalent to specify the node degree of literals in LCG. We can also enforce the degree of each node \( |\mathcal{V}_v| \) in our model.

Evaluation: Following G2SAT, we report the mean and standard deviation of statistics with respect to different test functions. These include the modularity, average clustering coefficient and the scale-free structure parameters for different graph representations of SAT instances. Please refer to Newman (2001; 2006); Ansótegui et al. (2009); Clauset et al. (2009) for more details. In general, the closer the statistical estimation the better it is.

Results: Following You et al. (2019), we compare the statistics of graphs with the training instances in Table 2. To mimic G2SAT which picks the best action among sampled options each step, we perform \( \epsilon \)-sampling variant (which is denoted BiGG-\( \epsilon \)). Such model has \( \epsilon \) probability to sample from Bernoulli distribution (as in Eq (8) (9)) each step, and \( 1 - \epsilon \) to pick best option otherwise. This is used to demonstrate the capacity of the model. We can see that the proposed BiGG can mostly recover the statistics of training graph instances. This implies that despite being general, the full autoregressive model is capable of modeling complicated graph generative process. We additionally report the statistics of generated SAT instances against the test set in Appendix B.3, where G2SAT outperforms BiGG in 4/6 metrics. As G2SAT is specially designed for bipartite graphs, the inductive bias it introduces allows the extrapolation to large graphs. Our BiGG is general purposed and has higher capacity, thus also overfit to the small training set more easily.

4.2. Scalability of BiGG

In this section, we will evaluate the scalability of BiGG regarding the time complexity, memory consumption and the quality of generated graphs with respect to the number of nodes in graphs.
Scalable Deep Generative Modeling for Sparse Graphs

### Table 1
Performance on benchmark datasets. The MMD metrics uses test functions from \{Deg., Clus., Orbit., Spec.\}. For all the metrics, the smaller the better. Baseline results are obtained from Liao et al. (2019), where OOM indicates the out-of-memory issue.

<table>
<thead>
<tr>
<th>Datasets</th>
<th>Methods</th>
<th>Training-24</th>
<th>G2SAT</th>
<th>BigG-0.1</th>
<th>BigG-0.01</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grid</td>
<td>0.79</td>
<td>2.00</td>
<td>0.68</td>
<td>5.64e^{-2}</td>
<td>1.00</td>
</tr>
<tr>
<td>$</td>
<td>V</td>
<td>_{max}=361,</td>
<td>V</td>
<td>_{avg} \approx 210$</td>
<td>0.13</td>
</tr>
<tr>
<td>$</td>
<td>E</td>
<td>_{max}=684,</td>
<td>E</td>
<td>_{avg} \approx 392$</td>
<td>1.12e^{-2}</td>
</tr>
<tr>
<td>Protein</td>
<td>0.21 (8%)</td>
<td>0.36</td>
<td>0.53</td>
<td>9.13e^{-2}</td>
<td>0.33</td>
</tr>
<tr>
<td>$</td>
<td>V</td>
<td>_{max}=500,</td>
<td>V</td>
<td>_{avg} \approx 1575$</td>
<td>0.36</td>
</tr>
<tr>
<td>$</td>
<td>E</td>
<td>_{max}=258,</td>
<td>E</td>
<td>_{avg} \approx 646$</td>
<td>0.68</td>
</tr>
<tr>
<td>3D Point Cloud</td>
<td>0.21</td>
<td>0.00</td>
<td>0.42</td>
<td>9.26e^{-2}</td>
<td>0.14</td>
</tr>
<tr>
<td>$</td>
<td>V</td>
<td>_{max}=5037,</td>
<td>V</td>
<td>_{avg} \approx 1377$</td>
<td>0.21</td>
</tr>
<tr>
<td>$</td>
<td>E</td>
<td>_{max}=10886,</td>
<td>E</td>
<td>_{avg} \approx 3074$</td>
<td>0.21</td>
</tr>
<tr>
<td>Lobster</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.12</td>
<td>0.00</td>
</tr>
<tr>
<td>$</td>
<td>V</td>
<td>_{max}=100,</td>
<td>V</td>
<td>_{avg} \approx 53$</td>
<td>0.12</td>
</tr>
<tr>
<td>$</td>
<td>E</td>
<td>_{max}=99,</td>
<td>E</td>
<td>_{avg} \approx 52$</td>
<td>0.00</td>
</tr>
</tbody>
</table>

Table 2. Training and generated graph statistics with 24 SAT formulas used in You et al. (2019). The neural baselines in Table 1 are not applicable due to scalability issue. We report mean and std of different test statistics, as well as the gap between true SAT instances.

### 4.2.1. Runtime and Memory Cost

Here we empirically verify the time and memory complexity analyzed in Section 2. We run BiGG on grid graphs with different numbers of nodes $n$ that are chosen from \{100, 500, 1k, 5k, 10k, 50k, 100k\}. In this case $m = \Theta(n)$. Additionally, we also plot curves from the theoretical analysis for verification. Specifically, suppose the asymptotic cost function is $f(n, m)$ w.r.t. graph size, then if there exist constants $c_1, c_2, n', m'$ such that $c_1 g(n, m) < f(n, m) < c_2 g(n, m), \forall n > n', m > m'$, then we can claim $f(n, m) = \Theta(g(n, m))$. In Figure 6 to 8, the two constants $c_1, c_2$ are tuned for better visualization.

Figure 6 reports the time needed to sample a single graph from the learned model. We can see the computation cost aligns well with the ideal curve of $O(n + m \log n)$. To evaluate the training time cost, we report the time needed for each round of model update, which consists of forward, backward pass of neural network, together with the update of parameters. As analyzed in Section 3.1, if there is a device with infinite FLOPS, then the time cost would be $O(\log n)$. We can see from Figure 7 that this analysis is consistent when graph size is less than 5,000. However as graph gets larger, the computation time grows linearly on a single GPU due to the limit of FLOPS and RAM.

Finally Figure 8 shows the peak memory cost during training on a single graph. We select the optimal number of chunks $k^* = O(\sqrt{\frac{m}{\log n}})$ as suggested in Section 3.3, and thus the peak memory grows as $O(\sqrt{m \log n})$. We can see such sublinear growth of memory can scale beyond sparse graphs with 100k of nodes.

### 4.2.2. Quality w.r.t. Graph Size

In addition to the time and memory cost, we are also interested in the generated graph quality as it gets larger. To do so, we follow the experiment protocols in Section 4.1.1 on a set of grid graph datasets. The datasets have the average number of nodes ranging in \{0.5k, 1k, 5k, 10k, 50k, 100k\}. We train on 80 of the instances, and evaluate results on
4.3. Ablation study

In this section, we take a deeper look at the performance of BiGG with different node ordering in Section 4.3.1. We also show the effect of edge density to the generative performance in Section 4.3.2.

4.3.1. BiGG with different node ordering

In the previous sections we use DFS or BFS orders. We find these two orders give consistently good performance over a variety of datasets. For the completeness, we also present results with other node orderings.

We use different orders presented in GRAN’s Github implementation. We use the protein dataset with spectral-MMD as evaluation metric against held-out test graphs. In summary: 1) BFS/DFS give our model consistently good performance over all tasks, as it reduces the tree-width for BiGG (similar to Fig5 in GraphRNN) and we suggest to use BFS or DFS by default; 2) BiGG is also flexible enough to take any order, which allows for future research on deciding the best ordering.

Determining the optimal ordering is NP-hard, and learning a good ordering is also difficult, as shown in the prior works. In this paper, we choose a single canonical ordering among graphs, as Li et al. (2018) shows that canonical ordering mostly outperforms variable orders in their Table 2, 3, while Liao et al. (2019) uses single DFS ordering (see their Sec 4.4 or github) for all experiments.

4.3.2. Performance on random graphs with decreasing sparsity

We here present experiments on Erdos-Renyi graphs with on average 500 nodes and different densities. We report spectral MMD metrics for GRAN, GT and BiGG, where GT is the ground truth Erdos-Renyi model for the data.

<table>
<thead>
<tr>
<th>Edge density (%)</th>
<th>GRAN</th>
<th>GT</th>
<th>BiGG</th>
</tr>
</thead>
<tbody>
<tr>
<td>1%</td>
<td>3.50e-4</td>
<td>9.97e-4</td>
<td>9.47e-4</td>
</tr>
<tr>
<td>2%</td>
<td>1.23e-4</td>
<td>4.55e-4</td>
<td>5.08e-4</td>
</tr>
<tr>
<td>5%</td>
<td>7.81e-4</td>
<td>2.82e-4</td>
<td>3.18e-4</td>
</tr>
<tr>
<td>10%</td>
<td>1.31e-3</td>
<td>1.94e-4</td>
<td>8.38e-4</td>
</tr>
</tbody>
</table>

Table 5. Graph generation quality with decreasing sparsity. We use spectral-MMD as evaluation metric against held-out test graphs.

Our main focus is on sparse graphs that are more common in the real world, and for which our approach can gain significant speed ups over the alternatives. Nevertheless, as shown in Table 5, we can see that BiGG is consistently doing much better than GRAN while being close to the ground truth across different edge densities.

5. Conclusion

We presented BiGG, a scalable autoregressive generative model for general graphs. It takes $O((n + m) \log n)$ complexity for sparse graphs, which substantially improves previous $\Omega(n^2)$ algorithms. We also proposed both time and memory efficient parallel training method that enables comparable or better quality on benchmark and large random graphs. Future work include scaling up it further while also modeling attributed graphs.
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References


