Molecular graph generation with Graph Neural Networks

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Abstract

The generation of graph–structured data is an emerging problem in the field of deep learning. Various solutions have been proposed in the last few years, yet the exploration of this branch is still in an early phase. In sequential approaches, the construction of a graph is the result of a sequence of decisions, in which, at each step, a node or a group of nodes is added to the graph, along with its connections. A very relevant application of graph generation methods is the discovery of new drug molecules, which are naturally represented as graphs. In this paper, we introduce a sequential molecular graph generator based on a set of graph neural network modules, which we call MG\textsuperscript{2}N\textsuperscript{2}. Its modular architecture simplifies the training procedure, also allowing an independent retraining of a single module. The use of graph neural networks maximizes the information in input at each generative step, which consists of the subgraph produced during the previous steps. Experiments of unconditional generation on the QM9 dataset show that our model is capable of generalizing molecular patterns seen during the training phase, without overfitting. The results indicate that our method outperforms very competitive baselines, and can be placed among the state of the art approaches for unconditional generation on QM9.

Keywords

Graph Generation, Molecule Generation, Deep Learning, Graph Neural Networks, Drug Discovery.
1 Introduction

Graph generation is a complex problem with several real–world applications, and many different approaches have been devised for solving it. Classical methods resorted to random mathematical models: The Erdős–Rényi model [1] was the first approach in this direction. Since Erdős–Rényi graphs tend to have unrealistically low clustering coefficients, especially with respect to community graphs, two methods were later developed, mainly to obtain more realistic small–world networks: the growth–based Barabási–Albert model [2], and the Watts–Strogatz rewiring model [3]. The recent developments in machine learning have stimulated its application to the field of graph generation. Deep learning techniques, indeed, can capture the characteristics of a given domain from a set of examples, which are then exploited to generate new graphs. Variational Auto–Encoders (VAEs) [4] were the first neural network models to be employed for this purpose [5] [6]. The success of Generative Adversarial Networks (GANs) [7] in image generation has led to replicate the same adversarial approach for graph–structured data [8] [9]. This approach can be improved by adding constraints to the adversarial learning [10]. The different nature of the problem, though, has encouraged the development of alternative solutions as well. While VAEs, by sampling representations from a continuous latent space, can generate graphs as unitary entities, many methods tackle the problem with a sequential approach. The construction of a graph becomes, therefore, a sequence of decisions, in which a node or a group of nodes is added to the graph at each step. On the one hand, many methods make use of Recurrent Neural Networks (RNNs) to handle the decision sequence [11], [12], [13]. On the other hand, Graph Neural Networks (GNNs) [14], with their capability of processing graph–structured data without loss of connectivity information, allow to build very powerful generative models. In particular, at each step, GNNs can exploit all the information contained in the partial graph generated by the previous steps, while recurrent models typically rely only on the sequence of previous decisions. In principle, this holds true for any GNN model, including Graph Nets [15], GraphSAGE [16], Gated Graph Sequence Neural Networks [17], Message Passing Neural Networks [18], and Graph Convolution Networks [19] [20] [21]. The only approach of this type we are aware of is DeepGMG [22], based on Graph Nets [15].

The generation of molecular graphs is a complex task, which can lead to the development of new instruments for drug discovery, potentially cutting the huge costs, in terms of both time and money, of that fundamental research process [23]. The space of molecular graphs is virtually infinite, and even constraining the size of molecules to few atoms, the number of theoretically possible compounds is overwhelming. Efficient automatic generative techniques are required for the exploration of such huge space, and deep generative models represent ideal candidates. Using SMILES notation [24], molecules can be generated as sequential objects. This approach has been carried out with VAE models [25], also exploiting the grammar of the SMILES language [26] [27]. However, SMILES strings do not preserve the full connectivity information, as molecules are more naturally represented as undirected graphs, with finite (and relatively small) sets of vertex and edge types. Graph–based VAEs have been employed in the generation of molecular graphs [25]. Junction–Tree VAEs build graphs by connecting pre–extracted structural motifs [29], an approach which has been recently extended to larger molecules and polymers by making the VAE hierarchical [30]. This approach can also be improved by exploiting the valence histogram of each atom [31]. Recently, statistical flow models, characterized by an invertible encoder/decoder, have been developed [32]. Graph GANs have been employed for the generation of molecules,
handling the decisions with Reinforcement Learning (RL) techniques \[33\] \[34\]. Finally, the above mentioned approach based on Graph Nets have also been applied to this task with promising results \[22\].

In this work, we present a sequential molecular graph generator based on GNNs \[14\], which we call Molecule Generative Graph Neural Network (MG$^2$N$^2$). A single node is added and connected to the graph, at each step. The method focuses on one node at a time, and generates its neighbors before processing the following node, preventing disconnected components from being created. Similarly to GraphRNN \[11\], we follow a Breadth First Search (BFS) ordering to decide which nodes to expand first. Edges are generated in parallel rather than sequentially, making the approach less computationally demanding. The control flow in MG$^2$N$^2$ depends on decisions implemented by three GNN modules. The sequential and modular nature of our method makes it interpretable. As previously mentioned, at each step, GNNs exploit all the information contained in the subgraph generated until that step. Gumbel softmax \[35\] output layers allow the networks to be trained over discrete stochastic distributions. Moreover, the modules are trained independently of each other: This feature simplifies the learning process and allows to retrain each module independently. The GNN model used in this work was derived from the original GNN approach \[14\], which was proved to be a universal approximator on graphs \[36\]. This property ensures that the GNN model is general enough to be able to make the complex decisions that the modules must implement.

The contributions of this work consist in a new sequential generative model for molecular graphs, MG$^2$N$^2$, and its experimental evaluation on a well-known benchmark for molecule generation, the Quantum Machine 9 (QM9) dataset. The results show that the proposed approach outperforms very competitive baselines and reaches the state of the art for unconditional generation. The experiments also clarify the main properties of the method and show that MG$^2$N$^2$ is capable of generating molecules with chemical characteristics similar to those of the dataset.

The rest of this paper is organized as follows. A short description of the GNN model used in this work is provided in Section 2. Section 3 presents and discusses the generative algorithm and its implementation with neural networks. The experiments and their results are described and commented in Section 4. Our conclusions are drawn in Section 5.

2 The GNN Model

The GNN used in this paper is based on the original model \[14\]. The implementation is described in the following. Each graph $G = (V, E)$ is composed of a set of vertices $V$ and a set of edges $E \subseteq V \times V$. Each vertex $v_i \in V$ is labeled with a vector of attributes $l_i$. Each edge $(v_i, v_j) \in E$ is labeled with a vector of attributes $e_{i,j}$. Based on $E$, a neighborhood function, assigning a set of neighbors $Ne(v_i)$ to each vertex $v_i \in V$ is defined as $Ne(v_i) = \{v_j : (v_j, v_i) \in E\}$.

A state $s_i$ is assigned to each vertex $v_i \in V$. It is initialized as $s_i^0 = l_i$. The state of all the vertices is then updated iteratively, and in parallel, until convergence and/or a maximum number of iterations are reached. During this process, each vertex communicates with its neighbors by sending and receiving messages. The state $s_i^k$ of vertex $v_i$, at step $k$, is calculated as in Eq. (1), based on its state at the previous step $s_i^{k-1}$ and the messages coming from its neighbors $v_j \in Ne(v_i)$.

$$s_i^k = F(s_i^{k-1}, A(\{M(v_i, v_j, k) : v_j \in Ne(v_i)\}))$$ (1)
In Eq. (1), some message passing function $M$ defines the message sent from $v_j$ to $v_i$ at step $k$. A neighborhood aggregation function $A$ defines how the messages coming from all the neighbors of $v_i$ are aggregated. Finally, a state updating function $F$ defines how the new state is calculated. In principle, all of these functions could be learnt by neural networks. In earlier models [37], $M$ was implemented by a MultiLayer Perceptron (MLP), while the other two functions were kept fixed. In this work, we opted for learning $F$ with an MLP, while keeping $A$ and $M$ fixed. This choice is also in line with other recent developments, which are broadly summarized in [15][16]. In particular, we defined $M$ as the concatenation of the state of the source node $s_j^{k-1}$ and the label $e_{i,j}$ of the edge $(v_i,v_j)$, as shown in Eq. (2).

$$M(v_i,v_j,k) = (s_j^{k-1}, e_{i,j})$$

Two different neighborhood aggregation functions were employed, defined as either the arithmetic mean or the element–wise sum of the incoming messages, as shown in [9].

$$A_{avg}(\{M(v_i,v_j,k) : v_j \in Ne(v_i) \}) = \frac{1}{|Ne(v_i)|} \sum_{j:v_j \in Ne(v_i)} M(v_i,v_j,k)$$

$$A_{sum}(\{M(v_i,v_j,k) : v_j \in Ne(v_i) \}) = \sum_{j:v_j \in Ne(v_i)} M(v_i,v_j,k)$$

Therefore, we can introduce a hyperparameter coefficient $a$ which allows us to choose between the aggregation functions $A_{avg}$ and $A_{sum}$, by simply assigning it the values $1/|Ne(v_i)|$ or 1, respectively. As a consequence, the final form of (1) can be written as in [4].

$$s_i^k = F(s_i^{k-1}, a \sum_{j:v_j \in Ne(v_i)} (s_j^{k-1}, e_{i,j}))$$

The state updating process is stopped when either state convergence or a maximum number of iterations $k = k_{max}$ are reached. State convergence is assumed to happen when the distance between the current state and the previous state becomes negligible: $\|s_i^k - s_i^{k-1}\| < \epsilon$, and $k_{max}$ are set as hyperparameters. $\| \cdot \|$ is the Euclidean norm. The output $y \in \mathbb{R}^p$ is then determined by an output function $O$ which depends on the type of problem at hand. In graph–based problems it is defined on the whole graph $G$, in node–based problems it is defined on single vertices $v_i \in V_{out} \subseteq V$, while in edge–based problems it is defined on single edges $(v_i,v_j) \in E_{out} \subseteq E$, where $V_{out}$ and $E_{out}$ are respectively the subset of vertices and the subset of edges for which an output is defined.

In this work, the problem of graph generation is divided into three classification subtasks. One of them is node–based while the other two are edge–based. In the former, the output function is defined as in Eq. (5), while in the latter, the output function is defined as in Eq. (6).

$$y_i = O(s_i^k)$$

$$y_{i,j} = O(s_i^k, s_j^k, e_{i,j})$$

In both equations, $k^*$ represents the last iteration of Eq. (1), determined by one or both of the stopping conditions. The function $O$ is implemented by an MLP, which is put in cascade to the state network. Learning is achieved by comparing, through a loss function $E$, the supervision $y$ to the output $\hat{y}$ produced by the output network. The network weights are updated by Backpropagation Through Time [35].
3 Method

Our method consists in a graph generation algorithm tailored to the task of molecule generation, and its implementation with GNNs. Subsection 3.1 describes the algorithm, while the implementation is presented in Subsection 3.2. The details about preprocessing are contained in Subsection 3.3 while Subsection 3.4 deals with node ordering issues.

3.1 Generation Algorithm

The generation of a labeled graph $G = (V, E)$ is handled as a sequential process, starting with an initially empty $E$ and with a single vertex $V = \{v_0\}$. The label $l_0$ of $v_0$ is sampled from a distribution of labels $D_0$, which is learned from the training set. Each step consists in adding a new node to the graph and connecting it to the other nodes. The algorithm focuses on one node $v_i$ at a time, generating all its neighbors before focusing on the following node $i = i + 1$. This process will be referred to as node expansion. Nodes are indexed according to the order in which they have been generated, so that, for instance, the third generated node $v_3$ will always be the fourth node to be expanded ($v_0$ is the first). The process stops when all the nodes have been expanded ($i > |V|$) or when the maximum number of nodes has been reached ($|V| = |V_{\text{max}}|$).

As a new node $v_j$ is generated, first it is connected to the node $v_i$ which is being expanded, then it can be linked to the other vertices $V \setminus \{v_i, v_j\}$. While the set of edges generated in the latter phase can be empty, the $(v_i, v_j)$ edge is always generated. This constraint ensures that the generated graph is always connected, without impairing generality: any graph can still be produced.

We can define three problems that must be solved to carry out a generative step. Each problem corresponds to a function the model will provide: node generation ($P_1$), first edge classification ($P_2$), additional node linking ($P_3$).

- $P_1$ decides whether to expand $v_i$ with a new neighbor node $v_j$ or to stop its expansion. If $v_j$ is generated, $P_1$ also returns its label $l_j$.
- $P_2$ is called after a new node $v_j$ has been generated. It determines the label $e_{i,j}$ of the edge $(v_i, v_j)$.
- $P_3$ is called after a new node $v_j$ has been generated and connected to $v_i$. It determines the existence of any possible edge connecting $v_j$ to any other vertex $v_k \in V \setminus \{v_i, v_j\}$. The labels of all the generated edges are also returned. All the edges are processed in parallel. The main drawback of this approach is that the dependencies between edges are ignored, but it also brings the advantages of avoiding edge ordering biases and of significantly reducing the time cost.

The generation algorithm is summarized in Algorithm 1.

3.2 Implementation with Graph Neural Networks

In the proposed approach, each of the functions $P_1$, $P_2$, $P_3$ described in Subsection 3.1 is implemented by a dedicated GNN module, which will be referred to as M1, M2, M3, respectively. Each of the modules is trained separately, and one step at a time, assuming the other two modules’ decisions to always correspond to the ground truth. This is a strong assumption, which will prevent the model from exploring possible different solutions, but it dramatically simplifies the training procedure. Another
Algorithm 1 Graph generation algorithm.

procedure Generate($G = (V, E)$)
    $V \leftarrow \{v_0\}, l_0 \sim D_0$
    $E \leftarrow \emptyset$
    $i \leftarrow 0$
    $j \leftarrow 1$
    while $(i < |V|) \land (|V| \leq |V|_{\text{max}})$ do
        $gd \leftarrow P1(V, E, i)$
        while $gd \neq \text{stop}$ do
            $V \leftarrow V \cup \{v_j\}, l_j \leftarrow gd$
            $E \leftarrow E \cup \{(v_i, v_j)\}$
            $e_{i,j} \leftarrow P2(V, E, i, j)$
            for $k \in [0, j - 1], k \neq i$ do
                $ld \leftarrow P3(V, E, k, j)$
                if $ld \neq \text{disconnected}$ then
                    $E \leftarrow E \cup \{(v_k, v_j)\}$
                    $e_{k,j} \leftarrow ld$
                end if
            end for
            $j \leftarrow j + 1$
        end while
        $gd \leftarrow \text{GeneratorDecision}(V, E, i)$
    end while
    $i \leftarrow i + 1$
end procedure
advantage of this paradigm is the fact that, each being trained separately from the others, the modules can be recombined to build new versions of the model. If a module needs to be optimized there is no need of re-training the other two.

In order to generate labeled graphs, we need to make some assumptions on the nature of vertex and edge labels. Three main cases can be identified: unlabeled graphs, graphs with continuous vertex and edge labels, graphs with a finite set of vertex and edge types. In this work, we will focus on the third case, which corresponds to the typical setting in molecule generation problems. Thus, in the following, we assume that the label \( l_i \) of any vertex \( v_i \) belongs to a finite set \( \mathbb{T}_v \), the label \( e_{i,j} \) of any edge \( (v_i, v_j) \) belongs to a finite set of types \( \mathbb{T}_e \), and \( \mathbb{T}_v \) and \( \mathbb{T}_e \) are defined by the dataset.

In the proposed approach, the GNN modules generate nodes and edges along with their labels. With reference to Algorithm 1, the following holds.

- **P1** faces a node-based classification problem, as it decides whether to stop the expansion of the current node \( v_i \) or to generate another neighbor \( v_j \), and, in case, which label to assign to \( v_j \). The set of output vertices of M1 consists only of \( v_i \): \( V_{out} = \{ v_i \} \). The output classes correspond to the union of the stop decision to the set of vertex types \( \{ \text{stop} \} \cup \mathbb{T}_v \).

- **P2** deals with an edge-based classification problem, since it generates the label of the edge connecting the vertex being expanded \( v_i \) and its new neighbor \( v_j \). The set of output edges of M2 consists only of this edge \( E_{out} = \{ (v_i, v_j) \} \). The output classes correspond to the set of edge types \( \mathbb{T}_e \).

- **P3** works on an edge-based classification problem, since it predicts the existence, and, in case, the label, of every possible edge connecting the new vertex \( v_j \) to the other nodes in the graph, except the node being expanded \( v_i \). These calls are parallelized and integrated in a single prediction from M3. This idea has the drawback of considering each predictable edge as if it were independent from the other predictable edges, but it also allows to avoid the biases introduced by taking the decisions in sequence and it speeds up the procedure. To do so, the graph \( G \) is extended with a set of provisional edges \( E_p = \{ (v_k, v_j) : v_k \in V \setminus \{ v_i, v_j \} \} \). The module M3 takes in input the new graph \( G' = (V, E') : E' = E \cup E_p \). The set of output edges for M3 is \( E'_{out} = E_p \). The output classes correspond to the union of the disconnected decision to the set of edge types \( \{ \text{disconnected} \} \cup \mathbb{T}_e \).

To learn a stochastic behavior from the supervisions, which are samples from a categorical distribution, we resorted to a Gumbel softmax output layer \(^{35}\), based on the Gumbel–Max method for sampling from discrete distributions \(^{39} \) \(^{40} \).

This approach allows to backpropagate through an arbitrarily close approximation of the categorical distribution. The softmax can be annealed, by decreasing a temperature parameter \( \tau \), from a less accurate approximation, which tends to a uniform distribution for \( \tau \to \infty \), and to the discrete distribution itself for \( \tau \to 0 \). Lower temperatures come at the cost of an increasing gradient variance. The choice of two parameters \( \tau_{max} \) and \( \tau_{min} \), and a curve, will determine the annealing path. Annealing while training has the positive effect of encouraging the exploration of alternatives to the decision with the highest estimated probability in the early phases, to then converge to more canonical decisions in the final training epochs, when the estimation of the class probabilities has gained higher reliability. This is very important to prevent the networks from learning repetitive patterns, and to avoid mode collapse (i.e. generating always vertices of the same type, based on the highest prior probability).
3.3 Graph Preprocessing

To build the training, validation, and test sets for M1, M2, M3, the molecules from the dataset under analysis are pre–processed. For each generative step, we need an input graph, the index of the focus node, and a supervision. Each molecular graph \( G = (V, E) \) is decomposed in a sequence of incomplete graphs, one for each generative step.

For M1, the sequence is composed of \( n = 2|V| - 1 \) graphs. The first graph contains only one node \( G_0 = (V_0 = \{v_0\}, E_0 = \{\}) \), any intermediate graph \( G_i = (V_i, E_i) \) corresponds to an incomplete subgraph of \( G \), \( G_i = (V_i \subset V, E_i \subset E) \), and the last graph is complete \( G_{n-1} = G \). For M2 and M3, the sequences are composed of \( n = |V| - 1 \) graphs, because M2 and M3 are not called after the \(|V| \) stop decisions from M1 (see Algorithm 1). The graphs \( G_i = (V_i \subset V, E_i \subset E) \) acquire nodes and edges as \( i \) grows.

The sets are built so that graphs from the same generative sequence (which correspond to different levels of completion of the same original graph) belong to the same set (and to the same batch). This is particularly important to avoid evaluation biases deriving from testing or validating on examples which have slightly different replicas in the training set.

3.4 Node Ordering

To define the generative sequences of the graphs, a node ordering needs to be established. This will determine the order in which the nodes of each graph must be generated, and, consequently, the sequences of input graphs and supervisions described in Subsection 3.3. The model is expected to learn this generative strategy from the training set, so that, for instance, a training set in which carbon atoms have higher priority will teach the model to generate carbon neighbors first. Theoretically, being \( V \) a set with no given ordering, the model would benefit from being trained on any possible node ordering. Since this is impossible from a computational point of view, some constraints must be imposed to reduce the number of orderings from \( o(|V|!) \) to a computationally feasible level. In this work we chose a Breadth First Search (BFS) strategy, which has the additional benefit of reducing the number of link predictions needed at each step [11]. Among the nodes at the same depth level in the BFS tree, node types with lower average centrality are expanded first. The average centrality of node types is measured on the training set, according to the Freeman Betweenness Centrality [41]. This boosts both positive effects of the BFS strategy. To further reduce the number of possible orderings of a factor \(|V|\), we decided to always start from the same node, which is node 0 of the original node numbering taken from the dataset. The other nodes are then re-numbered according to the previous rules, making a random choice in any case in which multiple permutations are still possible. The latter two assumptions allow us to retain one unique ordering, coming at the cost of a loss of generality. Although this cost would likely be critical for a truly recurrent model, it is sustainable in this learning framework, in which the correlation between two steps is limited to the output of the first shaping the input of the second. The only input to the model, in fact, is represented by the graph itself, regardless to the many possible sequences of steps that may have brought to its current shape.
4 Experimental Methodology

We performed a series of experiments, testing our model on the QM9 dataset [42], a common benchmark in molecular graph generation, which is introduced in Subsection 4.1. Subsection 4.2 describes the model setup, deals with the hyperparameters, and introduces the experiments we performed. The evaluation methods are described in Subsection 4.3 while Subsection 4.4 contains the results, the comparisons with other models, and the considerations on their significance.

4.1 Dataset Description

To evaluate our method, a set of experiments were run on the Quantum Machine 9 (QM9) dataset [42], which is a subset of GDB–17, a chemical universe of 166 billion molecules [43]. QM9 is an ideal benchmark for a new generative model for molecular graphs, as most competitive methods in this area have been tested on this dataset. It is composed of 133,885 compounds, made of up to 9 heavy atoms (C,O,N,F), plus the hydrogens which are bound to them, for a maximum size of 29 atoms. Each molecule is represented as an undirected graph, in which each vertex corresponds to an atom and each edge corresponds to a chemical bond. The label of each vertex represents the corresponding atom type, through one–hot encoding, so that $|T_v| = 5$. The label of each edge represents, through one–hot encoding, the type of chemical bond connecting its two vertices, which can be either single, double or triple, so that $|T_e| = 3$. The output of the modules M1, M2 and M3, defined in Subsection 3.2 have dimensions, respectively $|\{\text{stop}\} \cup T_v| = 6$, $|T_v| = 3$, and $|\{\text{disconnected}\} \cup T_e| = 4$.

A random splitting procedure is applied to the dataset, in order to obtain a training set, a test set, and a validation set, composed of 120,000, 10,000 and 3,885 molecule graphs, respectively. The validation set is used, during the training phase, to evaluate the performance of our models on data that are not provided to them directly. The held–out test set allows us to compare the statistics of our sets of generated graphs to the statistics of ground–truth graphs which have never been seen by our generative model, assessing the capability of the model of reproducing the chemical characteristics of QM9 compounds.

Inside each graph, the nodes are re–numbered according to the procedure described in Subsection 3.4. To determine the order among the neighbors $N(e(v_{i}))$ of a generic $v_{i} \in V$, the average Freeman Betweenness Centrality is measured on the 120,000 training graphs, obtaining the following values: FBC(Hydrogen) = 0.0, FBC(Fluorine) = 0.0, FBC(Oxygen) = 0.115, FBC(Nitrogen) = 0.246, FBC(Carbon) = 0.382.

4.2 Experimental Setup

The code for training the GNNs [44] and generating graphs was implemented using Tensorflow [45]. All the training runs of module M1 were issued on a Nvidia Tesla–V100 GPU, with 32 GB dedicated memory. Training runs of modules M2 and M3 always took place on a Nvidia 2080–Ti GPU. The training set was randomly split in 20 batches of 6,000 graphs each, to reduce the memory requirements. All the experiments used the same split. During the generation of new graphs, even though all the three modules are kept in memory, far less computational resources are needed. The generation sessions were run on the Nvidia 2080–Ti GPU, but required only 0.5 GB of memory.

Table I shows the configurations of the modules M1, M2, M3 used in the experiments,
Module Aggregation Epochs LR k\textsubscript{max} HU\textsubscript{state} HU\textsubscript{out}
M1(I) sum 700 $4 \times 10^{-3}$ 3 30 50
M1(II) sum 1500 $2 \times 10^{-3}$ 6 100 60
M1(III) sum 2000 $1 \times 10^{-5}$ 6 100 60
M2(I) avg 500 $2 \times 10^{-3}$ 3 20 50
M2(II) avg 1000 $1 \times 10^{-3}$ 4 40 60
M3(I) avg 500 $2 \times 10^{-3}$ 6 20 50
M3(II) sum 500 $2 \times 10^{-3}$ 6 20 50
M3(III)* avg 500 $2 \times 10^{-3}$ 6 20 50
M3(IV)* sum 500 $2 \times 10^{-3}$ 6 20 50

Table 1: Different module configurations are identified by the module number M1, M2 or M3 introduced in Subsection 3.2 and by a sequential version number (I, II, ...). Hyperparameters correspond to: neighbor aggregation function (Aggregation), training epochs (Epochs), initial learning rate (LR), maximum state convergence iterations (k\textsubscript{max}), hidden units of the state network (HU\textsubscript{state}), and hidden units of the output network (HU\textsubscript{out}). M3 versions marked with * were trained with class weights to balance the supervisions.

which include the neighbor aggregation function, the training epochs, the initial learning rate, the maximum number of iterations for state convergence, and the number of hidden units of the state network and the output network. Each GNN module is composed of a state and an output network. The former is a two–layered MLP implementing the state updating function described in Eq. (4). The latter is another two–layered MLP, implementing Eq. (5) in M1, and Eq. (6) in M2 and M3. The initial values M1(I), M2(I) and M3(I) in Table 1 were obtained through a preliminary experimentation, with the goal of maximizing the accuracy of the modules M1, M2, M3, each one independently from the others, on the validation set. Just as if the modules had been classifiers, accuracy was calculated as the percentage of correct outputs, according to the single step supervision, and regardless of molecule validity. For the Gumbel softmax annealing path, based on [35], we initially chose a linear descent from $\tau_{\text{max}} = 5.0$ to $\tau_{\text{min}} = 1.0$ during training. Tests on different linear configurations did not bring improvements. In particular, annealing to temperatures $\tau < 1.0$ brought the model to an unwanted repetitive behavior. Therefore, we kept the initial annealing path for all the successive training runs. All the models were trained with an Adam optimizer [46] and cross–entropy loss, which does not require adjustments to work with the Gumbel softmax output layers.

4.3 Evaluation

The evaluation of generation performances is twofold. On the one hand, the metrics for unconditional generation introduced in [28] are used to measure the validity, uniqueness and novelty of the generated graphs. On the other hand, the distributions of the chemical properties of the compounds can be compared to those measured on the test set, assessing the model’s capability of reproducing the characteristics of QM9 compounds. Both evaluations are carried out on batches of 10,000 generated graphs. Let Gen be the set\textsuperscript{1} of generated compounds, Val $\subseteq$ Gen be the subset of chemically valid compounds, and QM9 be the set of molecules in the dataset. Validity is calcu-

\textsuperscript{1}More precisely, here we are using the multiset, an extension of the standard set which can contain multiple copies of the same instances.
lated as the fraction of chemically valid molecules over the total generated molecules: 
\[ \text{Validity} = \frac{|\text{Val}|}{|\text{Gen}|} \].

Uniqueness is the fraction of unique molecules among the valid ones: 
\[ \text{Uniqueness} = \frac{|\text{uniq}(\text{Val})|}{|\text{Val}|} \], where \( \text{uniq} \) is a function that takes in input a multiset and returns the corresponding set, from which the duplicates are removed. Novelty is the fraction of unique molecules which do not match any QM9 compound: 
\[ \text{Novelty} = \frac{|\text{uniq}(\text{Val})| - |\text{uniq}(\text{Val} \cap \text{QM9})|}{|\text{uniq}(\text{Val})|} \].

We also define an additional measure, that combines the three previous metrics and accounts for the fraction of valid, unique and novel molecules over the total generated ones: 
\[ \text{VUN} = \text{Validity} \times \text{Uniqueness} \times \text{Novelty} \].

The chemical properties include the molecular weight of each compound, the logarithmic octanol/water partition coefficient (logP) [47], and the quantitative estimate of drug–likeness (QED) score [48]. The logP coefficient quantifies the solubility of a molecule in polar or non-polar solvents, while the QED score assesses the drug–likeness of a compound, summarizing in a single measure the following chemical descriptors: polar surface area, molecular weight, logP, number of rotatable bonds, numbers of hydrogen bond donors and acceptors, number of aromatic rings, potential structural issues.

The validity as well as the chemical properties of each compound are assessed with the RDKit package [49]. In order to determine the uniqueness and novelty of a molecule, we resorted to the graph isomorphism function of the NetworkX package [50].

4.4 Results and Discussion

The first experiment consisted in a study of the role played in the algorithm by the hyperparameter \( |V|_{\text{max}} \) which controls the maximum number of nodes in a generated graph. In principle, our model, being trained step by step, can extend its generation procedure for an arbitrary number of iterations, until it stops itself on a complete molecule. This feature could be exploited to extend the generation domain to molecules which are larger than those seen during training, while retaining the same generic patterns. Using M1(I), M2(I) and M3(I), defined in Table 1, we explored different thresholds for the maximum number of generated nodes \( |V|_{\text{max}} \). The natural value for this dataset is \( |V|_{\text{max}} = 29 \), which corresponds to the largest graph size in the training set. As described in Section 3.1, the generation procedure stops when the number of vertices reaches \( |V|_{\text{max}} \). This means that any graph still incomplete at that point will not correspond to a valid molecule. Intuitively, raising \( |V|_{\text{max}} \) will increase the amount of valid generated compounds. Even if this is confirmed by the results reported in Table 2, the additional valid molecules, being heavier than average, alter the property distributions of the batch. Moreover, as shown in Fig. 1 their QED is below average. Falling in a region of low to very–low drug–likeness, these compounds are not useful in the scope of generating new potential drugs. These considerations suggested to keep \( |V|_{\text{max}} = 29 \) for the subsequent experiments.

Starting from the baseline configuration C1 (see Table 4), in which the modules are optimized separately and not on the evaluation metrics chosen for our task, we explored the hyperparameter space in search of a better configuration 3. A systematic search on a grid of configurations was computationally infeasible. Moreover, since the generative models are evaluated with antagonist metrics, it is impossible to optimize
Table 2: Higher values of $|V|_{\text{max}}$, on generation batches from the same model setup, produce more valid and unique compounds. The divergence of average QED and molecular weight from the values taken on the validation set (0.478 and 127.5, respectively), however, suggests that the best configuration is $|V|_{\text{max}} = 29$.

| Max size | Validity | Uniqueness | Avg. QED | Avg. Mol. Wt. |
|----------|----------|------------|----------|--------------|
| 29       | 0.491    | 0.814      | 0.448    | 124.6        |
| 40       | 0.593    | 0.845      | 0.438    | 144.7        |
| 80       | 0.688    | 0.866      | 0.408    | 172.9        |
| 1000     | 0.781    | 0.879      | 0.366    | 231.3        |

Figure 1: Logarithm of the molecular weight (left) and QED (right) distributions of generated graphs with different values of $|V|_{\text{max}}$. It can be observed how higher thresholds imply the generation of heavier compounds, with lower QED.

in increasing $k_{\text{max}}$ and the number of hidden units in the first two modules, in order for them to better capture complex molecular patterns during training. Using this new configuration (M1(II) and M2(II)), we explored different setups for M3. In particular, to establish the ideal neighbor aggregation method, M3(I) was compared to M3(II). Then, the same comparison was repeated on M3(III) and M3(IV), which were trained with class weighted supervision. This latter measure was meant to decrease the learning bias from the very unbalanced prior class probability distribution, which could prevent the model from learning the chemical rules (roughly, 97% of the examples belong to the disconnected class, while the other 3% are distributed over the three bond types). The results of these configurations can be observed in Table 3. This balancing strategy for module M3 did not bring advantages, as it is shown in Table 3. This suggests that the GNN can deal with the unbalanced distribution, and efforts to improve the generation performances should focus on other parameters. Besides, the two neighbor aggregation methods appear to be equivalent. A new version of the node generation module, M1(III) was also trained, increasing the number of training epochs and decreasing the initial learning rate (see Table 1), in order to stabilize the configuration for all of them. Thus, we have heuristically selected the most promising solutions and reported those experiments which, in our opinion, are the most interesting.

The error on each pattern is multiplied by the inverse of the prior of its target class. In this way, the GNN will tend to produce a balanced output over all the classes. At test time, the output is re-multiplied by the vector of prior class probabilities, to restore this important piece of information.
Table 3: Alternative setups for M3. Balancing weights bring no advantage on model performances. The two aggregation functions show equivalent results.

| Config. | M1 | M2 | M3 | Validity | Uniqueness | Avg. QED | Avg. Mol. Wt. |
|---------|----|----|----|----------|------------|----------|---------------|
| C1      | M1(I) | M2(I) | M3(I) | 0.491 | 0.813 | 0.448 | 124.6 |
| C2      | M1(II) | M2(II) | M3(I) | 0.511 | 0.888 | 0.461 | 134.8 |
| C3      | M1(III) | M2(II) | M3(II) | 0.668 | 0.340 | 0.404 | 75.3 |

Table 4: Summary of the best configurations determined by preliminary experiments. C3 produces more valid molecules, while the highest QED is obtained by C2. C1 has the closest average molecular weight to the validation set reference (127.5).

The lack of an explicit global reconstruction penalty is one of the reasons for the very high novelty of the material produced by MG\textsuperscript{2}N\textsuperscript{2}: the model is not forced to perfectly reconstruct the molecules on a global basis, but it is forced to correctly reconstruct the local parts of the graph. This approach is expected to preserve a certain degree of validity while encouraging the model to explore more different molecular patterns. Though GraphVAE and MolGAN have higher validity, our model outperforms both of them in terms of uniqueness of the compounds. The aggregated VUN score shows that MG\textsuperscript{2}N\textsuperscript{2} generates the highest percentage of molecules which are valid, unique, and novel at the same time. Notice that, differently to all of the baselines, our method explicitly generates the hydrogen atoms, and all of the hydrogens are required to have been explicitly generated to mark a molecule as valid. This difference is one of the factors determining the lower performance of our approach on the validity metric.

To further assess the chemical similarity between the generated material and the test set of molecules from QM9, we plotted the distributions of the chemical descriptors, which can be observed in Fig. 2. For a qualitative visual comparison, we extracted VAEs learn to reconstruct the training examples as closely as possible. The reconstruction penalty is calculated on a global basis, as the Kullback-Leibler divergence between the example graph and its reconstructed version. As the KL-divergence cannot be directly optimized, due to the presence of intractable terms, VAEs optimize the Evidence Lower BOund (ELBO) of these terms, which provides a valuable method to enforce a good global reconstruction.
| Model       | Valid | Unique | Novel | VUN | Avg. QED  | Avg. logP | Avg. Mol. Wt. |
|-------------|-------|--------|-------|-----|-----------|-----------|---------------|
| GrammarVAE | 0.602 | 0.093  | 0.809 | 0.045| -         | -         | -             |
| GraphVAE   | 0.557 | 0.760  | 0.616 | 0.261| -         | -         | -             |
| MolGAN     | 0.981 | 0.104  | 0.942 | 0.096| -         | -         | -             |
| Ours(C2)   | 0.511 | 0.888  | 1.000 | 0.454| 0.461 (0.116) | 0.272 (1.336) | 134.8 (45.7) |
| Ours(C3)   | 0.668 | 0.340  | 1.000 | 0.227| 0.404 (0.088) | 0.238 (1.093) | 75.3 (52.8)  |
| Test       | -     | -      | -     | -   | 0.482 (0.096) | 0.270 (1.325) | 127.3 (7.6)  |

Table 5: Validity, Uniqueness, and Novelty of generated compounds assessing the quality of our models and the baselines. The average values of chemical descriptors (Molecular Weight, logP, and QED) are compared to the same quantities measured over the test set. Standard deviations are reported between parentheses. Metrics for GrammarVAE and GraphVAE are taken from the GraphVAE article [28]. The performances of MolGAN are taken from the original paper [33].

some valid molecules at random from each set and plotted their structural formulas with RDKit (see Fig. 3 and Fig. 4).

While achieving an acceptable logP distribution, configuration C3 fails to reproduce the QED distribution of the test set. Configuration C2, instead, generates compounds which have very similar logP and QED distributions with respect to those of the test set. This is due to the further optimization carried out on C3: while achieving the goal of building more valid compounds, it actually went in contrast with the other objectives of generating unique, novel molecules with QM9-like properties. The learning parameters proved to have a role in determining the properties of the model, as we can see by comparing C2 and C3. C2 can be considered as our best model configuration.

Figure 2: logP (left) and QED (right) distributions of generated graphs and training/test molecules. It can be observed how well C2 has generalized the chemical characteristics of the compounds seen during training.

5 Conclusions

In this article, we introduced a generative model for molecular graphs: Molecule Generative Graph Neural Network (MG$^2$N$^2$), and a sequential generation algorithm we devised for this purpose. Our method exploits the capabilities of Graph Neural Net-
works to natively process graph-structured data. This allows to exploit the graph output of the previous step as the network input, which represents an advantage with respect to other sequential methods, that mainly rely on the sequence of previous decisions, rather than on the graph they produce. The modularity of our model implies an easier, less resource demanding, learning process. In line with all the other sequential methods, and contrarily to VAEs, the generation process is easily interpretable: the steps in which errors occur, or in which specific atoms and bonds are created, can be readily identified in the generation sequence. This feature is very important as it simplifies any process of improvement or repurposing of the model.

The model was tested on a benchmark generation task over the QM9 dataset. The distributions of the chemical descriptors retraced those measured on the held out test set. The quality of generated graphs proved to be very high, allowing our model to outperform very competitive baselines.
Figure 4: Grid representation of a random sample of 64 valid molecular graphs generated with configuration C3.

Future work will focus on generalizing the model to other molecular graph generation problems, and on extending the present approach to conditional generation. A conditional generation model could be implemented by concatenating a vector of desired properties to the input of each module.

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