Order Agnostic Autoregressive Graph Generation

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Abstract

Graph generation is a fundamental problem in various domains, including chemistry 1 and social networks. Recent work has shown that molecular graph generation 2 using recurrent neural networks (RNNs) is advantageous compared to traditional 3 4 generative approaches which require converting continuous latent representations 5 into graphs. One issue which arises when treating graph generation as sequential 6 generation is the arbitrary order of the sequence which results from a particular choice of graph flattening method. In this work we propose using RNNs, taking into 7 account the non-sequential nature of graphs by adding an Orderless Regularization 8 (OLR) term that encourages the hidden state of the recurrent model to be invariant 9 to different valid orderings present under the training distribution. We demonstrate 10 11 that sequential graph generation models benefit from our proposed regularization scheme, especially when data is scarce. Our findings contribute to the growing body 12 of research on graph generation and provide a valuable tool for various applications 13 requiring the synthesis of realistic and diverse graph structures. 14

15 **1** Introduction

Graphs are powerful representations of complex relationships and structures found in a wide range 16 of domains, including social networks, molecular chemistry, transportation networks, distributed 17 algorithms and many more. A dedicated class of architectures, Graph Neural Networks (GNNs), has 18 been developed to handle the specific properties of graphs. Graphs are naturally versatile objects, 19 but such versatility comes at the cost of lack of structure and no naturally induced order. Most GNN 20 architectures therefore operate by applying a neural architecture at the node level followed by an 21 aggregation step which takes into account the local neighborhood structure of the graph. By stacking 22 multiple such layers, a GNN is able to perform node-level or graph-level tasks that take into account 23 the entire structure of the graph. 24

The ability to generate realistic and structured graphs is essential for various applications ranging from 25 drug design [17, 20, 34, 36, 44, 55, 66, 69] to program synthesis [9, 32, 11, 1, 30, 65, 7, 15]. In recent 26 years a wide variety of generative models have been developed, including generative adversarial 27 networks (GANs), variational autoencoders (VAEs), normalizing flows, and diffusion models. These 28 algorithms devise different strategies to learn continuous mappings from a latent distribution to a space 29 30 of realistic examples. Unfortunately, graphs do not admit a natural representation in a continuous space; consequently, the discrete and unordered nature of graphs make them less amenable to the 31 methods mentioned above for the task of graph generation. A different type of generative model 32 relies on autoregressive architectures which enable processing a sequence and generating the next 33 element; for example, these architectures are commonly used for large language models. Generally, 34 autoregressive models are applicable when the generated objects admit a sequential order. 35

In this work we focus on sequential generation of graphs using autoregressive neural architectures. A strong motivating factor for choosing autoregressive architectures is that we are particularly interested in molecular graph generation; and in this context Flam *et al.* [23] have shown that sequential

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generation is favorable compared to other approaches. The specific representation we consider is depth-first search (DFS) trajectories of graphs. The reasons for this choice of representation is twofold: (a) DFS is a natural way of flattening graphs into sequences; (b) in the chemistry community DFS is used to convert molecules into strings. However, an issue arises when converting graphs into sequences: there are many DFS trajectories for a given graph. Indeed, for many graph flattening methods, there is an arbitrariness in the order of the sequence which results [63, 12].

⁴⁵ In order to alleviate the dependency on a specific trajectory, we add a regularization term dubbed

⁴⁶ Orderless Regularization (OLR) which ensures the learnt model is invariant to different DFS orderings

⁴⁷ of the same graph. For the sake of training with OLR, one needs to generate different DFS trajectories ⁴⁸ with a common end-vertex which is known to be hard [5]. We formalize the notion of graph-level

invariance and devise an efficient algorithm to generate such trajectories under certain constraints.

⁵⁰ Finally, we demonstrate empirically that our regularization term is beneficial when the amount of

⁵¹ training data is limited by considering the use case of small molecule generation.

The reminder of the paper is structured as follows: in Section 2 we provide background and introduce the concepts, definitions, and notations used throughout the paper. Section 3 goes into the details of

54 OLR over DFS trajectories. Section 4 is devoted to related work. In Section 5 we provide empirical

⁵⁵ evidence for the effectiveness of OLR. Section 6 provides concluding remarks.

56 2 Background

In this section we formally define the problem of graph generation, the notations and definitions necessary to present our proposed method. We denote matrices by bold uppercase letters, $\mathbf{M} \in \mathbb{R}^{n \times m}$, vectors by bold lowercase letters, v, and the i^{th} entry of v by v_i . We proceed with a general formulation of recurrent models.

61 2.1 Recurrent Models

⁶² Let \mathcal{X}, \mathcal{H} , and \mathcal{Y} be the spaces of inputs, hidden states, and outputs, respectively. Given an input ⁶³ sequence $\mathbf{x} \equiv (\mathbf{x}_1, \dots, \mathbf{x}_n) \in \mathcal{X}^n$, a recurrent model consists of two functions, the state update ⁶⁴ function $f_u : \mathcal{H} \times \mathcal{X} \to \mathcal{H}$,

$$\boldsymbol{h}_{t+1} = f_u(\boldsymbol{h}_t, \boldsymbol{x}_t), \tag{2.1}$$

and the output function $f_o: \mathcal{H} \times \mathcal{X} \to \mathcal{Y}$,

$$\boldsymbol{y}_t = f_o(\boldsymbol{h}_t, \boldsymbol{x}_t). \tag{2.2}$$

where $h_0 \in \mathcal{H}$. We overload the notation and denote the hidden state and output of a recurrent model over a sequence as $f_u(\mathbf{x})$ and $f_o(\mathbf{x})$ respectively.

The formulation presented of recurrent models is broad and able to capture RNNs as well as more
 complex architectures such as Gated Recurrent Units (GRUs, [13]) and Long-Short Term Memory
 networks (LSTMs, [33]). For example, in the case of a very simple, vanilla RNN,¹

$$\boldsymbol{h}_{t+1} = \sigma_h (\mathbf{A}\boldsymbol{h}_t + \mathbf{B}\boldsymbol{x}_t) \tag{2.3}$$

71 and

$$\boldsymbol{y}_t = \sigma_y (\mathbf{C}\boldsymbol{h}_t + \mathbf{D}\boldsymbol{x}_t) \tag{2.4}$$

where A, B, C, D are matrices with the appropriate dimensions and σ_h, σ_y are standard nonlinearities such as *sigmoid* or *tanh*.

74 2.2 Graph Generation

A graph is given by $G = (\mathcal{V}, \mathcal{E})$ where \mathcal{V} is a set of nodes (or vertices) and $\mathcal{E} \subseteq \mathcal{V} \times \mathcal{V}$ is a set of tuples denoting the nodes connected by an edge in the graph. Additionally, for each $v \in \mathcal{V}$, denote by $x_v \in \mathbb{R}^m$ the features of node v. Similarly, $e_{uv} \in \mathbb{R}^k$ denotes the features of the edge $(u, v) \in \mathcal{E}$. For example, in a molecular graph, nodes are atoms, and their features will contain the element; and edges correspond to bonds, and their features will contain the bond types (single, double, etc.).²

¹Note that the bias term may be encapsulated into the input processing matrices by expanding the input with an additional dimension and assigning a fixed value of 1 on that coordinate.

²Molecular node and edge features may contain other properties as well.

80 Another example is of social networks, where nodes corresponds to users and their features to user

81 profiles; and edges correspond to connections between users and their features contain metadata on 82 this connection.

The topic of designing neural networks to operate specifically on graphs is dominated by Graph Neural Networks (GNNs) which mostly rely on a message-passing scheme to propagate information between nodes. While these architectures are extremely successful in node level and graph level prediction, they are not as prevalent in the context of graph generation, and many such approaches are restricted to small graphs (though [16] is a notable exception).

Formally, the task of graph generation is usually concerned with learning to model distributions: 88 concretely, given a set of N graphs $\{G_i\}_{i=1}^N$ originating from an underlying distribution p, the 89 goal of graph generation is to devise an algorithm that generates new graphs from the underlying 90 distribution p. Prior work has mostly adapted successful generative methods over a continuous space 91 to the domain of graphs [25, 8, 37, 51]. In this work we focus on using recurrent models which 92 can be employed naturally to generate discrete objects. Crucially, Flam et al. [23] have shown that 93 sequential generation is favorable compared to competing approaches in the context of molecular 94 graph generation. 95

96 2.3 Sequential Graph Generation

When applying recurrent models for graph generation, the graph first needs to be "flattened" into 97 a sequence. As there is no natural order for a graph, one must artificially induce such an order; for 98 example, the approach taken by [67] considers generation of breadth-first search (BFS) trajectories. 99 While there are many ways to convert a graph into a sequence, in this work we focus on depth-first 100 search (DFS); a strong motivation for this choice is that this is the method used to convert graph 101 molecules into a linear representation called SMILES strings [64]. By convention, the output of the 102 DFS algorithm is a spanning tree and we consider the induced order of the graph as the order in 103 which the vertices were visited during the DFS run (also known as pre-order traversal). 104

¹⁰⁵ In what follows we formally define the concepts discussed.

Definition 2.1. Given a connected graph $G = (\mathcal{V}, \mathcal{E})$ with $|\mathcal{V}| = n$, we say the permutation $\pi \in \mathbb{S}_n$

107 is a valid ordering of G if it is possible to run DFS over G and visit the vertices in the order induced

by π . Denote the sequence corresponding to a valid ordering π of G by

$$\mathbf{s}(G,\pi) = (v_{\pi(1)}, \dots, v_{\pi(n)}). \tag{2.5}$$

109 Denote the set of all such sequences for a given graph G by

$$\mathcal{S}(G) = \{ \mathbf{s}(G, \pi) : \pi \text{ is a valid ordering of } G \}.$$
(2.6)

Clearly, for a non-trivial graph S(G) will contain many sequences. In this work we have a special interest in sequences that share the same end vertex.

Definition 2.2. Let S(G, v) denote all sequences terminating at node $v \in V$, formally,

$$\mathcal{S}(G,v) = \{ \mathbf{s} \in \mathcal{S}(G) : s_n = v \}$$
(2.7)

In the following section we discuss the desired properties for recurrent models when used for graph generation.

115 **3** Structure Agnostic Recurrent Models

Recurrent models are a natural choice when generating discrete objects such as text. On the other hand, graphs are discrete objects with no naturally induced order. In Section 2 we described a mapping between graphs and sequences; and in particular, the fact that many different sequences correspond to the same graph. In this section we present our method that overcomes the issues described.

121 3.1 Generating Depth-First Search Traversals

¹²² In this work we use recurrent models to generate DFS traversals of graphs. Clearly when generating a DFS traversal, the next node to be generated depends on the nodes generated thus far and in particular



Figure 1: Illustration of two DFS traversals of the same graph starting from node A and terminating at node D, blue lines denote traversal order. (Left) traversal resulting in the sequence A(BEF)(C)D. (Right) traversal resulting in the sequence A(C)(BFE)D. The parentheses denote the opening and closing of branches when traversing the tree; with this syntax it is possible to reconstruct the tree from such sequences. Note that multiple sequences correspond to the same tree, a fact that lies at the heart of this work.

the last generated node. An important observation is that the output of the recurrent model should be

invariant to different valid orderings corresponding to the same subgraph as long as they lead to the

same node. The following definition formalizes this notion,

Definition 3.1. We say a recurrent model is structure invariant with respect to a connected graph G if

$$\forall v \in \mathcal{V}, \ \forall \mathbf{s}_1, \mathbf{s}_2 \in \mathcal{S}(G, v) \ it is the case that \ f_o(\mathbf{s}_1) = f_o(\mathbf{s}_2).$$
(3.1)

¹²⁹ If the above condition is satisfied for all $G \sim D$, we say that the recurrent model is structure ¹³⁰ invariant with respect to a distribution D.

Figure 1 depicts a graph and two different DFS traversals sharing the same root and terminal node. A recurrent model processing the two DFS traversals will ideally generate the same node that will be attached to node *D*.

Definition 3.1 describes the structure invariance property with respect to a graph. Since recurrent models generate the traversal sequentially, we would like this property to hold at any moment during generation, i.e., we want to modify Definition 3.1 to take into account partial DFS traversals.

Definition 3.2. For a connected graph G, we say a connected subgraph $\tilde{G} \subseteq G$ is **induced by DFS over** G if there exists a valid ordering $\pi \in S_n$ of G, and $k \leq n$ such that $(v_{\pi(1)}, \ldots, v_{\pi(k)})$ is a valid ordering of \tilde{G} . Denote the set of all DFS induced subgraphs over G by $\mathcal{G}_{DFS}(G)$.

At this stage, a reader might question the necessity of Definition 3.2 and why $\mathcal{G}_{DFS}(G)$ differs from the set of all connected subgraphs of G. We note that for a general connected graph $\mathcal{G}_{DFS}(G)$ does **not** correspond to the set of all connected subgraphs.

143 **Proposition 3.3.** For a connected graph G,

$$\mathcal{G}_{DFS}(G) \neq \left\{ \tilde{G} \mid \tilde{G} \subseteq G \text{ and } \tilde{G} \text{ is connected} \right\}$$
(3.2)

- Figure 2 depicts a graph and two connected subgraphs, one which is induced by DFS and the other that cannot be obtained by a DFS traversal.
- With the notion of DFS induced subgraphs in hand, we now present the following definition of *total structure invariance*:
- **Definition 3.4.** We say a recurrent model is **totally structure invariant** with respect to a connected graph *G*, if

$$\forall \hat{G} \in \mathcal{G}_{DFS}(G), \ \forall v \in \mathcal{V}(\hat{G}), \ \forall \mathbf{s}_1, \mathbf{s}_2 \in \mathcal{S}(\hat{G}, v) \ it is the case that \ f_o(\mathbf{s}_1) = f_o(\mathbf{s}_2).$$
(3.3)



Figure 2: Illustration of the same graph with two connected subgraphs: (Left) subgraph which is not induced by DFS. (Right) subgraph induced by DFS, arrows depict a traversal resulting in the sequence BA(CF)D.

- 150 If the above condition is satisfied for all $G \sim D$, we say that the recurrent model is **totally structure**
- 151 *invariant with respect to a distribution* \mathcal{D} .
- ¹⁵² In the next section we discuss how to train recurrent models which are totally structure invariant with ¹⁵³ respect to a given training distribution over graphs.

154 3.2 Regularizing Towards Total Structure Invariance

Motivated by the observation discussed in Section 3.1, we propose training recurrent models that are totally structure invariant with respect to the underlying distribution over graphs. It would be appealing to characterize the class of all totally structure invariant functions and optimize over those. Unfortunately, it is difficult to attain a crisp characterization of structure invariance as this property depends on the training distribution.

Instead, we propose encouraging total structure invariance via regularization. Specifically, we would
 like to minimize the following auxiliary loss,

$$\mathbb{E}_{G\sim\mathcal{D}}\mathbb{E}_{\tilde{G}\sim\mathcal{G}_{DFS}(G)}\mathbb{E}_{v\in\mathcal{V}(\tilde{G})}\mathbb{E}_{\mathbf{s}_{1},\mathbf{s}_{2}\in\mathcal{S}(\tilde{G},v)}\left[\left(f_{o}(\mathbf{s}_{1})-f_{o}(\mathbf{s}_{2})\right)^{2}\right]$$
(3.4)

which we refer to as *Orderless Regularization* (OLR). Examining Equation 3.4, we note that sampling from $\mathcal{G}_{DFS}(G)$ is easily done by randomly selecting a root node and running DFS with stochastic decision making. On the other hand, given \tilde{G} and v, sampling from $\mathcal{S}(\tilde{G}, v)$ is hard and has been shown to be NP-complete [5].

166 3.3 Sampling Trajectories with Common End Vertex

The problem of generating all DFS trajectories that terminate at the same vertex is hard and there are no known efficient algorithms for this task. In order to overcome this obstacle we apply a heuristic for computing such trajectories. We highlight that our proposed scheme is not equivalent to a uniform sampling over all possible trajectories; however, in Section 5 we show that the resulting regularization scheme is effective empirically.

- Next, we formally show that for practical graphs there exists efficient algorithms to generate such
 trajectories.
- **Definition 3.5.** Let $G = (\mathcal{V}, \mathcal{E})$ be an arbitrary graph. *G* is said to be *k*-edge-connected if the subgraph $G' = (\mathcal{V}, \mathcal{E} \setminus \tilde{\mathcal{E}})$ is connected for all $\tilde{\mathcal{E}} \subseteq \mathcal{E}$ such that $|\tilde{\mathcal{E}}| < k$.
- **Proposition 3.6.** There is an efficient algorithm to find distinct DFS trajectories with common end vertex for any k-edge connected graph for $k \leq 2$.

We note that in many real world tasks, graph are rarely k-edge connected for k > 2. For example, in the ZINC molecular dataset, more than 99.5% of molecular graphs are 1-edge connected.

Proof Sketch. Find a min-cut: by the definition of 1-edge-connected graphs the min-cut includes a single crossing edge (u, v). By removing (u, v) the graph is partitioned into two connected components, G_1 and G_2 containing u and v respectively. Run a DFS on G_1 with u as the root vertex to result in (u_1, \ldots, u_k) , and similarly for G_2 to result in (v_1, \ldots, v_m) (where $v_1 = v$ and $u_1 = u$).³ We can now construct a DFS traversal on G by 'gluing' together the sequences as,

$$(v_1, u_1, \dots, u_k, v_2, \dots, v_m) \tag{3.5}$$

We can run another (stochastic) DFS on G_1 from u to obtain $(u_{\tilde{\pi}(1)}, \dots, u_{\tilde{\pi}(k)})$ where $\pi \in S_k$ and $\tilde{\pi}(1) = 1$. We can construct a second DFS sequence as in Equation 3.5,

$$(v_1, u_{\tilde{\pi}(1)}, \dots, u_{\tilde{\pi}(k)}, v_2, \dots, v_m)$$
 (3.6)

187 We have created two valid DFS sequences that both terminate at v_m .

188 See Appendix B for the full details and the case of 2-edge connected graphs.

189 4 Related Work

In this section we discuss several relevant topics to graph generation. For a comprehensive review on graph generation see [28, 71].

One Shot Generation Classic generative architectures (e.g. Variational autoencoders (VAEs) [39], 192 Generative adversarial networks (GANs) [26], etc.) work by learning a continuous mapping from 193 a latent distribution to generate new examples with similar properties to the training distribution. 194 These models usually incorporate a neural architecture that maps directly from the latent space to 195 the domain of the training data (e.g. images) and therefore the output space must be predetermined. 196 These properties pose a challenge when applied to the domain of graphs, as the latter are discrete 197 objects with variable size and no naturally induced order. In order to circumvent these caveats, prior 198 work [3, 17, 19, 21, 22, 29, 34, 43, 44, 55, 58, 72] has used a one-shot generation strategy. That is, 199 the output space is limited by design to a specific representation of graphs (i.e. adjacency matrix 200 201 or adjacency list) of specific size and the output is generated in a single forward pass. While the one-shot strategy has its merits, there are a few significant drawbacks such as the inability to generate 202 graphs with arbitrary large number of nodes. 203

Sequential Generation The idea of using autoregressive models for graph generation is not new 204 and there have been several works in this vein. GraphRNN [67] proposes generating BFS trajectories 205 in order to limit the number of possible orderings per graph. Other works take a different approach 206 of generating edges in an autoregressive manner [4, 27]. Additional approaches include Molecular-207 RNN [50] which incorporates a reinforcement learning environment to generate nodes and edges 208 sequentially. Yet another approach includes sequentially generating subgraph structures [36, 41, 47]. 209 Another recent work [10] treats the induced order as a problem of dimensionality reduction and 210 attempts to learn mappings from graphs to sequences. In this work we argue that the most effective 211 inductive bias for the use of autoregressive models to generate graphs is to be invariant to different 212 orderings possible under the training distribution. 213

Molecule Generation One of the most prominent uses of graph generation, which is used for evaluation in this work, is that of molecule generation. Molecular generation is applicable to the development of synthetic materials, drug development and more. Molecules are 3D objects which are naturally represented as point clouds⁴ with corresponding geometric approaches [24, 56, 57, 35] which utilize inherent symmetries in the architectures employed. While 3D representations are richer and carry significant information that does not transfer to 1D and 2D representations, they

 $^{{}^{3}}k$ and m denote the size of partitions and satisfy $k + m = |\mathcal{V}|$.

⁴In a point-cloud representation of a molecule each point represents an atom and bonds are implicit from the distances between atoms.

are costly to obtain and therefore the corresponding amount of data is limited as compared to 1D and 2D representations, which are ubiquitous. Another aspect of molecule generation is when the generation is conditioned to satisfy certain properties. For example, [59, 70, 53] generate molecules that are conditioned to bind to specific ligand structures, [38, 69] generate molecules that fulfill certain chemical properties. In this work we consider the task of de-novo generation [2, 42, 48, 61] where the objective is to generate molecules with similar properties to those in the training data.

Permutation Invariant Recurrent Models Another relevant topic is the use of autoregressive 226 models for problems over sets which, like graphs, lack a natural order. There have been many works 227 focusing on problems over sets. The most prominent of these is DeepSets [68] which applies a 228 deep neural network on each element of the set and then aggregates the result with a permutation 229 invariant operator (e.g. sum or max), finally applying another deep neural network on the aggregated 230 result. There have also been autoregressive works designed for sets: [46] use RNNs on different 231 permutations and output the average. While this requires n! orderings for a set of size n, the authors 232 have presented several approximation techniques and justified them empirically. [14] have shown 233 that while DeepSets are universal, some permutation invariant functions require unbounded width to 234 implement successfully and have proposed using RNNs with a regularization term which enforces 235 permutation invariance. 236

237 5 Experiments

A prominent application of graph generation is that of molecule design. Graph generation tasks range 238 from de novo generation where the objective is to generate molecules with similar properties to a 239 given dataset, to conditional generation for which the task is to generate a graph given a second graph 240 with specific characteristics, i.e. a ligand that binds to a specific target. Our empirical evaluation 241 242 focuses on the former. We evaluate our proposed regularization method on the MOSES benchmark [49] and compare to relevant baselines. Our implementation is based on the work of CharRNN which 243 use three layers of the LSTM architecture each with hidden dimension of 600 (for complete details 244 refer to [54]). We find a consistent improvement when adding OLR to the objective of autoregressive 245 models. 246

The data curated by [49] is refined from the ZINC dataset [60] which contains approximately 4.6Mmolecules. The authors filter the data based on molecular weights, number of rotational bonds, lipophilicity, etc. to result in a total of 2M molecules. The authors provide partitions of the data into train, test and scaffold test to allow fair evaluation.⁵

251 5.1 Computing Trajectories

OLR works by feeding two different trajectories that terminate at the same node. While this calculation is feasible to perform during the forward pass it introduces a computational bottleneck. In order to circumvent this issue we employ the following calculations offline. For each molecule we first index all min-cuts and randomly select one. We then generate multiple (10) traversals terminating at the same node as described in Section 3.3 and write the sequences into a file along with the original molecule from which the trajectories are derived from. When loading the data, two trajectories are selected at random and used as inputs to the OLR objective described in Section 3.2.

259 5.2 Data Filtering

Our offline computation of trajectories in Section 5.1 requires that there are min-cuts that induce sufficient number of different DFS traversals terminating at the same node. While 99.9% of the molecules in MOSES have at least two such trajectories, we filter the data to remain with molecules which have at least 10 different trajectories satisfying the criteria defined. After filtering we are left with approximately 500K molecules for training, and 55K for test and scaffold test partitions. We note that in following sections we show our method is most effective when training data is scarce and therefore the filtering process does not limit the applicability of our proposed regularization scheme.

⁵The scaffold of a molecule is the structure induced by its ring systems along with the connectivity pattern between these systems. The scaffold test partition contains molecules with structures that did not appear in the train and test partitions. The scaffold test allows for the evaluation of how well the model can generate previously unobserved scaffolds.

	Canonical	Randomized	OLR + Randomized
Unique@1K ([†])	1.0	1.0	1.0
Unique@10K ([†])	0.9965	0.9975	0.9981
FCD/Test (\downarrow)	0.6623	0.8568	0.7784
FCD/TestSF (\downarrow)	1.1980	0.4967	0.4936
SNN/Test (†)	0.5012	0.9947	0.9958
SNN/TestSF ([†])	0.4835	0.8246	0.8220
Frag/Test (1)	0.9960	1.4236	1.3089
Frag/TestSF (1)	0.9943	0.4795	0.4769
Scaf/Test (↑)	0.8193	0.9919	0.9926
Scaf/TestSF ([†])	0.0862	0.1185	0.0931
IntDiv (↑)	0.8515	0.8508	0.8537
IntDiv2 (↑)	0.8457	0.8449	0.8479
Filters (1)	0.9720	0.9705	0.9702
Novelty (\uparrow)	0.9749	0.9797	0.9809
Rank Average	2.42	1.85	1.50

Table 1: Generation results at validity threshold of 0.8. Leading result highlighted in bold for each metric. Rank Average is the average position of each method over all metrics considered. As can be seen, OLR outperforms the baselines considered. Refer to the text for further details.

267 5.3 Results

Our results for training with OLR compared to other baselines trained on the same data are shown in 268 Table 1. The most relevant baselines is CharRNN [54] which is an autoregressive model trained on 269 Canonical SMILES. We further compare to a randomized version of CharRNN inspired by the finding 270 of [2] which show that augmenting the data by using randomly generated SMILES representations 271 of the same molecule improves performance. We also attempted to compare our method to other 272 non-autoregressive models such as those based on Variational Autoencoders (VAEs) [8, 25, 37]; 273 however, we found that the models did not produce valid molecules when trained with 1000 examples. 274 so we do not report these results. We use the metrics defined by the MOSES benchmark [49]; see 275 Appendix A for a thorough description of these metrics. 276

In order to demonstrate the effectiveness of OLR we use 1000 randomly sampled data points from the training set and evaluate over the entire test set. When training with small amounts of data there is a trade-off between the validity of the generated molecules and the uniqueness and other metrics. Our evaluation considers the best performing models for each method providing the validity of the generated molecules exceeds 80%.

Results are depicted in Table 1. As can be seen, adding randomized variants of the molecules
outperforms the original work of [54] which train an RNN as a language model using only canonical
SMILES. Furthermore, adding the OLR objective exceeds the performance of randomized SMILES.
In order to clearly depict the performance difference, we calculate the rank of each method on each
metric considered. The average rank of each method is added as the last row of Table 1.

287 6 Conclusions

In this work we highlight the innate gap that every autoregressive model for graph generation must 288 mitigate - the induced order on graphs. We propose a different approach to previous works by 289 introducing a novel regularization scheme that encourages learning hypotheses that are invariant to 290 different DFS orderings. We demonstrate empirically that our proposed method improves performance 291 for autoregressive models and is especially effective when the available datasets are small, as is the 292 case in many real world problems. We believe that our approach can contribute to the applicability of 293 autoregressive models for graph generation and that similar ideas may be incorporated in various 294 generation strategies beyond the scope of this work. 295

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492 A Metric Details

⁴⁹³ In this section we provide details for the metrics reported in Table 1.

A few of the similarity measures (SNN and IntDiv) are based on the *Tanimoto coefficient*. In order to compute the Tanimoto coefficient, the molecules are mapped to a vector of fingerprints where each bit in the vector represents the presence (or absence) of a specific fragment.⁶ For molecules A, B, denote their fingerprints by m_A and m_B respectively, the Tanimoto coefficient is then calculated as the Jaccard index of the two vectors,

$$J(m_A, m_B) = \frac{|m_A \cap m_B|}{|m_A \cup m_B|} = \frac{|m_A \cap m_B|}{|m_A| + |m_B| - |m_A \cap m_B|}.$$
 (A.1)

499 We denote the Tanimoto coefficient of molecules A, B by T(A, B).

Unique@K report the fraction of uniquely generated valid SMILES strings amongst the Kmolecules generated (validity is determined by the RDKit library). We generate 30,000 molecules for each model and report for K = 1,000 and K = 10,000. High uniqueness values ensure the models do not collapse into repeatedly producing the same set of molecules.

Fréchet ChemNet Distance (FCD) is a metric for evaluating generative models in the chemical context, the method is based on the well established *Fréchet Inception Distance* (FID) metric used to evaluate the performance of generative models in computer vision [31].

Fréchet distance measure the Wasserstein-2 distance [62] from the distributions induced by taking the activations of the last layer of a relevant deep neural net. In the case of FCD, molecule activations are probed from ChemNet [45]. Given a set of generated molecules, denote by G the set of vectors as obtained by the activations of ChemNet, one can calculate the mean and covariance μ_G and Σ_G . Similarly, denote μ_R and Σ_R the mean and covariance of the set of molecules in the reference set, the FCD is calculated as follows,

$$FCD(G,R) = \|\mu_G - \mu_R\|^2 + Tr\left(\Sigma_G + \Sigma_R - 2(\Sigma_G \Sigma_R)^{1/2}\right).$$
 (A.2)

where Tr(M) denotes the trace of the matrix M. Low FCD values indicate that the generated molecules distribute similarly to the reference set.

Similarity to Nearest Neighbor (SNN) is the average of the Tanimoto coefficient of the generated molecule set denoted by G and their respective nearest neighbor in a reference set of molecules denote by R. High SNN indicates the generated molecules have similar structures to those in the reference set. This metric is in the range of [0, 1].

Fragment similarity (Frag) is a fragment similarity measure based on the BRICS fragments [18]. Denote the set of BRICS fingerprints vectors of the generated molecules by G and similarly R for the reference molecules. The fragment similarity is defined as the cosine similarity of the sum vectors,

$$Frag(G, R) = cosine\left(\sum_{g \in G} g, \sum_{r \in R} r\right)$$
 (A.3)

⁶The molecular fingerprints are obtained from RDKit [40] and are based on the extended-connectivity fingerprints [52].



Figure 3: Proof illustration - S has a cycle and two different trajectories starting from u and ending with w (urw and uw(r)). Concatenating with the trajectory from z to v we obtain two different DFS trajectories with a shared suffix.

The Frag measure is in the range of [0, 1], values closer to 1 indicate that the generated and reference molecule set have a similar distribution of BRICS fragment.

Scaffold similarity (Scaff) is similar to the fragment similarity, instead of the BRICS fragment, Scaff is based on mapping molecules to their Bemis–Murcko scaffolds [6].⁷ The measure also has a range of [0, 1], values closer to 1 indicate that the generated molecule set has a similar distribution of scaffold to the reference set.

Internal diversity (IntDiv) is a mesure of the chemical diversity within a generated set of molecules G. This metric indicates

$$IntDiv_{p} = 1 - \left(\frac{1}{|G|^{2}} \sum_{A,B \in G} T(A,B)^{p}\right)^{1/p}$$
(A.4)

We report the internal diversity for p = 1, 2. This measure is in the range [0, 1]. Low values indicate a lack of diversity in the generated molecules, i.e. that the model outputs molecules with similar fingerprints.

Filters is the fraction of generated molecules that pass a certain filtering that has been applied to the training data. The metric is in the range of [0, 1], high values indicate that the model has learnt to generate molecules which avoid the structures omitted by the filtering process.

Novelty is the fraction of generated molecules that does not appear in the training set. This measure is in the range of [0, 1] and is an indication of the whether the model overfits the training data.

538 **B** Missing Proofs

In this section we show how to construct distinct DFS trajectories with common end vertex for a 2–edge connected graph conditioned that the graph is not a cycle.

Proof. From our assumption that the graph is not a cycle, there exists at least two nodes with degree 3. Denote by C = (S, T) a minimal cut of size 2 (such a cut exists from our assumption that the graph is 2-connected). Denote the edges of the minimal cut by $e_1 = (u, v)$ and $e_2 = (w, z)$ such that $u, w \in S$ and $v, z \in T$. Next, we claim that at least one of the partitions contains a cycle, otherwise there is a path connecting S and T since there are nodes in the graph which have a degree of 3 in the original graph with a path between them. Assume with out loss of generality that S is the partition with a cycle, therefore there are at least 2 different traversals of S that start with u and end

⁷Bemis–Murcko scaffold is the ring structure of a molecule along with the bonds connecting the rings, i.e. the molecule without the side chains.

with w. There is also a trajectory between z and v. Putting together, there are at least 2 trajectories of the entire graph with a common suffix which is the traversal of T. Figure 3 illustrates the proof concept.