Deep generative models have been enjoying success in modeling continuous data. However, it remains challenging to capture the representations for discrete structures with formal grammars and semantics, e.g., computer programs and molecular structures. How to generate both syntactically and semantically correct data still remains largely an open problem. Inspired by the theory of compiler where syntax and semantics check is done via syntax-directed translation (SDT), we propose a novel syntax-directed variational autoencoder (SD-VAE) by introducing stochastic lazy attributes. This approach converts the offline SDT check into on-the-fly generated guidance for constraining the decoder. Comparing to the state-of-the-art methods, our approach enforces constraints on the output space so that the output will be not only syntactically valid, but also semantically reasonable. We evaluate the proposed model with applications in programming language and molecules, including reconstruction and program/molecule optimization. The results demonstrate the effectiveness in incorporating syntactic and semantic constraints in discrete generative models, which is significantly better than current state-of-the-art approaches.

1 INTRODUCTION

Recent advances in deep representation learning have resulted in powerful probabilistic generative models which have demonstrated their ability on modeling continuous data, e.g., time series signals (Oord et al., 2016; Dai et al., 2017) and images (Radford et al., 2015). Despite the success in these domains, it is still challenging to correctly generate discrete structured data, such as graphs, molecules and computer programs. Since many of the structures have syntax and semantic formalisms, the generative models without explicit constraints often produce invalid ones.

Conceptually, an approach in generative model for structured data can be divided in two parts, one being the formalization of the structure generation and the other one being a (usually deep) generative model producing parameters for stochastic process in that formalization. Often the hope is that with the help of training samples and capacity of deep models, the loss function will prefer the valid patterns and encourage the mass of the distribution of the generative model towards the desired region automatically.

Arguably the simplest structured data are sequences, whose generation with deep model has been well studied under the seq2seq (Sutskever et al., 2014) framework that models the generation of sequence as a series of token choices parameterized by recurrent neural networks (RNNs). Its widespread success has encouraged several pioneer works that consider the serialization of more complex structure data into sequences and apply sequence models to the represented sequences. Gómez-Bombarelli et al. (2016) (CVAE) is a representative work of such paradigm for the chemical molecule generation, using the SMILES line notation (Weininger, 1988) for representing molecules. However, because of the lack of formalization of syntax and semantics serving as the restriction of the particular structured data, underfitted general-purpose string generative models will often lead to invalid outputs. Therefore, to obtain a reasonable model via such training procedure, we need to prepare large amount of valid combinations of the structures, which in general is not practical.

To tackle such a challenge, one approach is to incorporate the structure restrictions explicitly into the generative model. For the considerations of computational cost and model generality, context-free grammars (CFG) have been taken into account in the decoder parametrization. For instance, in
Before introducing our model and the learning algorithm, we first provide some background knowledge which is important for understanding the proposed method.
2.1 Variational Autoencoder

The variational autoencoder (Kingma & Welling, 2013; Rezende et al., 2014) provides a framework for learning the probabilistic generative model as well as its posterior, respectively known as decoder and encoder. We denote the observation as $x$, which is the structured data in our case, and the latent variable as $z$. The decoder is modeling the probabilistic generative processes of $x$ given the continuous representation $z$ through the likelihood $p_{\theta}(x|z)$ and the prior over the latent variables $p(z)$, where $\theta$ denotes the parameters. The encoder approximates the posterior $p_{\theta}(z|x) \propto p_{\psi}(x|z)p(z)$ with a model $q_{\psi}(z|x)$ parametrized by $\psi$. The decoder and encoder are learned simultaneously by maximizing the evidence lower bound (ELBO) of the marginal likelihood, i.e.,

$$
\mathcal{L}(X; \theta, \psi) := \sum_{x \in X} \mathbb{E}_{q(z|x)} [\log p_{\theta}(x|z)p(z) - \log q_{\psi}(z|x)] \leq \sum_{x \in X} \log p_{\theta}(x)p(z)dz,
$$

where $X$ denotes the training datasets containing the observations.

2.2 Context Free Grammar and Attribute Grammar

Context free grammar A context free grammar (CFG) is defined as $G = (\mathcal{V}, \mathcal{X}, \mathcal{R}, s)$, where symbols are divided into $\mathcal{V}$, the set of non-terminal symbols, $\mathcal{X}$, the set of terminal symbols and $s \in \mathcal{V}$, the start symbol. Here $\mathcal{R}$ is the set of production rules. Each production rule $r \in \mathcal{R}$ is denoted as $r = \alpha \rightarrow \beta$, where $\alpha \in \mathcal{V}$ is a nonterminal, and $\beta = u_1u_2 \ldots u_{|\beta|} \in (\mathcal{V} \cup \mathcal{X})^*$ is a sequence of terminals and/or nonterminals.

Attribute grammar To enrich the CFG with “semantic meaning”, Knuth (1968) formalizes attribute grammar that introduces attributes and rules to CFG. The attribute is an attachment to the corresponding nonterminal symbol in CFG, written in the format $\langle v \rangle.a$ where $v \in \mathcal{V}$. There can be two types of attributes assigned to non-terminals in $G$: the inherited attributes and the synthesized attributes. An inherited attribute depends on the attributes from its parent and siblings, while a synthesized attribute is computed based on the attributes of its children. Formally, for a production $u_0 \rightarrow u_1u_2 \ldots u_{|\beta|}$, we denote $I(u_i)$ and $S(u_i)$ be the (disjoint) sets of inherited and synthesized attributes of $u_i, i \in \{0, \ldots, |\beta|\}$.

2.2.1 A Motivational Example

We here exemplify how the above defined attribute grammar enriches CFG with non-context-free semantics. We use the following toy grammar, a subset of SMILES that generates either a chain or a cycle with three carbons.

Production | Semantic Rule
--- | ---
$s \rightarrow \langle atom \rangle_1 \cdot 'C' \langle atom \rangle_2$ | $(s).\text{matched} \leftarrow (atom)_1.\text{set} \cap (atom)_2.\text{set}$
$s \rightarrow (s).\text{ok} \leftarrow (atom)_1.\text{set} = (s).\text{matched} = (atom)_2.\text{set}$
$(atom)_1 \rightarrow 'C' | 'C' \langle bond \rangle \langle digit \rangle$ | $(atom).\text{set} \leftarrow \emptyset | \text{concat}(\langle bond \rangle.\text{val}, \langle digit \rangle.\text{val})$
$(bond) \rightarrow '\cdot' | '\cdot' \langle digit \rangle$ | $(bond).\text{val} \leftarrow '\cdot' | '\cdot' \langle digit \rangle.\text{val}$
$(digit) \rightarrow '1' | '2' | \ldots | '9$ | $(digit).\text{val} \leftarrow '1' | '2' \ldots | '9$

where we show the production rules in CFG with $\rightarrow$ on the left, and the calculation of attributes in attribute grammar with $\leftarrow$ on the left. Here we leverage the attribute grammar to check (with attribute matched) whether the ringbonds come in pairs: a ringbond generated at $(atom)_1$ should match the bond type and bond index that generated at $(atom)_2$, also the semantic constraint expressed by $(s).\text{ok}$ requires that there is no difference between the set attribute of $(atom)_1$ and $(atom)_2$. Actually such constraint in SMILES is known as cross-serial dependencies (CSD) (Bresnan et al., 1982) which is non-context-free (Shieber, 1985). Another example of CSD is a sequence of multiple different types of parentheses where each separately balanced disregarding the others. Figure 2a further illustrates the example. Here all the attributes are synthetic, i.e., calculated in a bottom-up direction.

In the semantic correctness checking procedure, one need to perform (possibly multiple) bottom-up and top-down procedures for calculating the attributes after the parse tree is generated, however, in
the structure generating process, the parse tree is not ready for semantic checking, since the synthesized attributes coming from children are not generated yet. Due to such dilemma, it is nontrivial to use the attribute grammar to guide the top-down generation of the tree-structured data. One straightforward way is using acceptance-rejection sampling scheme, i.e., using the CFG decoder in grammar VAE \cite{kusner2017grammar} as a proposal and the semantic checking as the threshold. It is obvious that since the decoder does not include semantic guidance, the proposal distribution may raise semantically invalid candidate frequently, therefore, wasting the computational cost in vain.

3 SYNTAX-DIRECTED VARIATIONAL AUTOENCODER

As described in Section 2.2.1, directly using attribute grammar to address both syntax and semantics constraints is not efficient. In this section we describe how to bring forward the attribute grammar online and incorporate it into variational autoencoders such that our VAE generates both syntactic and semantic valid outputs by definition. We name our proposed method Syntax-Directed Variational Autoencoder (SD-VAE).

3.1 STOCHASTIC SYNTAX-DIRECTED DECODER

By scrutinizing the tree generation, the major difficulty in incorporating the attributes grammar into the processes is the appearance of the synthesized attributes. For instance, when expanding the start symbol \( s \), the corresponding synthesized attribute \( s \).matched is not ready yet. Since none of its children is generated, their synthesized attributes are also absent at this time, making the \( s \).matched unable to be computed. To enable the on-the-fly computation of the synthesized attributes for semantic validation during tree generation, besides the two types of attributes, we introduce the stochastic lazy attributes to enlarge the existing attribute grammar, so that the synthesized attributes will be transformed to inherited constraints in generating procedure and instantiated once all the dependent attributes are ready (also named as lazy linking in the following content).

We demonstrate how the decoder with stochastic lazy attributes will generate semantic valid output through a pedagogical example with the subset of SMILES grammar in figure 2(b). Following the terminology in compiler theory, we named it as stochastic syntax-directed decoder.

The tree generation procedure is indeed sampling from the decoder \( p_{θ}(x | z) \), which can be decomposed into several steps that elaborated below:

i) **stochastic predetermination**: in figure 2(b), we start from the node \( s \) with the synthesized attributes \( s \).matched determining the index and bond type of the ringbond that will be matched at node \( s \). Since we know nothing about the children nodes right now, the only thing we can do is to ‘guess’ a value. That is to say, we associate a stochastic attribute \( \langle s \rangle.sa \in \{0, 1 \}^{C_0} \sim \prod_{i=1}^{C_0} B(sa_i | z; p) \) as a predetermination for the sake of the absence of synthesized attribute \( s \).matched. \( B(\cdot) \) is the bernoulli distribution. Here \( C_0 \) is the maximum cardinality.
Algorithm 1 Decoding with Stochastic Syntax-Directed Decoder

1: **Global variables:** CFG: \( G = (\mathcal{V}, \Sigma, \mathcal{R}, s) \), decoder network parameters \( \theta \)

2: **procedure** \( \text{GENTree}(\text{node}, T) \)

3: Sample stochastic lazy attribute \( \text{node}.sa \sim B_\theta(sa|\text{node}, T) \) \hspace{1em} \( \triangleright \) when introduced on \( \text{node} \)

4: Sample production rule \( r = (\alpha \rightarrow \beta) \in \mathcal{R} \sim p_\theta(r|ctx, node, T) \). \hspace{1em} \( \triangleright \) The conditioned variables encodes the semantic constraints in tree generation.

5: \( ctx \leftarrow \text{RNN}(ctx, r) \) \hspace{1em} \( \triangleright \) update context vector

6: **for** \( i = 1, \ldots, |\beta| \) **do**

7: \( v_i \leftarrow \text{Node}(u_i, node, \{v_j\}_{j=1}^{i-1}) \) \hspace{1em} \( \triangleright \) node creation with parent and siblings’ attributes

8: \( \text{GenTree}(v_i, T) \) \hspace{1em} \( \triangleright \) recursive generation of children nodes

9: Update synthetic and stochastic attributes of \( \text{node} \) with \( v_i \) \hspace{1em} \( \triangleright \) Lazy linking

10: **end for**

11: **end procedure**

possible\(^1\) for the corresponding attribute \( a \). In above example, the 0 indicates no ringbond and 1 indicates one ringbond at both \( \langle \text{atom}_1 \rangle \) and \( \langle \text{atom}_2 \rangle \), respectively.

**ii) constraints as inherited attributes:** we pass the \( \langle s \rangle.sa \) as inherited constraints to the children of node \( \langle s \rangle \), i.e., \( \langle \text{atom}_1 \rangle \) and \( \langle \text{atom}_2 \rangle \) to ensure the semantic validation in the tree generation.

**iii) sampling under constraints:** assume in the valid order, \( \langle \text{atom}_1 \rangle \) is selected before \( \langle \text{atom}_2 \rangle \), we then sample the rules from \( p_\theta(r|\langle \text{atom}_1 \rangle, \langle s \rangle.z) \) for expanding \( \langle \text{atom}_1 \rangle \), and so on and so forth to generate the subtree recursively. Since we carefully designed sampling distribution that is conditioning on the stochastic property, the inherited constraints will be eventually satisfied. In the example, due to the \( \langle s \rangle.sa = \langle '1' \rangle \), when expanding \( \langle \text{atom}_1 \rangle \), the sampling distribution \( p_\theta(r|\langle \text{atom}_1 \rangle, \langle s \rangle.z) \) only has positive mass on rule \( \langle \text{atom} \rangle \rightarrow \langle 'c' \rangle \langle \text{bond} \rangle \langle 'digit' \rangle \).

**iv) lazy linking:** once we complete the generation of the subtree rooted at \( \langle \text{atom}_1 \rangle \), the synthesized attribute \( \langle \text{atom}_1 \rangle . \text{set} \) is now available. According to the semantic rule for \( \langle s \rangle . \text{matched} \), we can instantiate \( \langle s \rangle . \text{matched} = \langle \text{atom}_1 \rangle . \text{set} = \{ '1' \} \). When expanding \( \langle \text{atom}_2 \rangle \), the \( \langle s \rangle . \text{matched} \) will be passed down as inherited attribute to regulate the generation of \( \langle \text{atom}_2 \rangle \).

In summary, the general syntax tree \( T \in L(G) \) can be constructed step by step, within the languages \( L(G) \) covered by grammar \( G \). In the beginning, \( T^{(0)} = \text{root} \), where \( \text{root}.\text{symbol} = s \) which contains only the start symbol \( s \). At step \( t \), we will choose an nonterminal node in the \( \text{frontier} \) of partially generated tree \( T^{(t)} \) to expand. The generative process in each step \( t = 0, 1, \ldots \) can be described as:

1. Pick node \( v^{(t)} \in F_T(T^{(t)}) \) where its attributes needed are either satisfied, or are stochastic attributes that should be sampled first according to bernoulli distribution \( B(=v^{(t)}, T^{(t)}) \);  
2. Sample rule \( r^{(t)} = \alpha^{(t)} \rightarrow \beta^{(t)} \in \mathcal{R} \) according to distribution \( p_\theta(r^{(t)}|v^{(t)}, T^{(t)}) \), where \( v^{(t)} . \text{symbol} = \alpha^{(t)} \), and \( \beta^{(t)} = u_1^{(t)} u_2^{(t)} \ldots u_{|\beta^{(t)}|}^{(t)} \), i.e., expand the nonterminal with production rules defined in CFG.
3. \( T^{(t+1)} = T^{(t)} \cup \{(v^{(t)}, u_1^{(t)})\}_{i=1}^{|\beta^{(t)}|} \), i.e., grow the tree by attaching \( \beta^{(t)} \) to \( v^{(t)} \). Now the node \( v^{(t)} \) will have children represented by symbols in \( \beta^{(t)} \).

The above process continues until all the nodes in the frontier of \( T^{(T)} \) are all terminals after \( T \) steps. Then, we obtain the algorithm\(^2\) for sampling both syntactic and semantic valid structures.

In fact, in the model training phase, we need to compute the likelihood \( p_\theta(x|z) \) given \( x \) and \( z \). The probability computation procedure is similar to the sampling procedure in the sense that both of them requires tree generation. The only difference is that in the likelihood computation procedure, the tree structure, i.e., the computing path, is fixed since \( x \) is given, while in the sampling procedure, it is sampled following the learned model. Specifically, the generative likelihood can be written as:

\[
p_\theta(x|z) = \prod_{t=0}^T p_\theta(r_t|ctx^{(t)}, node^{(t)}, T^{(t)}) B_\theta(sa_t|node^{(t)}, T^{(t)})
\]  \hspace{1em} \( (2) \)

\(^1\) Note that setting threshold for \( C_s \) assumes a mildly context sensitive grammar (e.g., limited CSD).

\(^2\) Here frontier is the set of all nonterminal leaves in current tree.
where $\text{ctx}^{(0)} = z$ and $\text{ctx}^{(t)} = \text{RNN}(r_t, \text{ctx}^{(t-1)})$. Here RNN can be commonly used LSTM, etc..

### 3.2 Structure-Based Encoder

As we introduced in section 2, the encoder, $q_\psi(z|x)$ approximates the posterior of the latent variable through the model with some parametrized function with parameters $\psi$. Since the structure in the observation $x$ plays an important role, the encoder parametrization should take care of such information. The recently developed deep learning models [Duvenaud et al., 2015; Dai et al., 2016; Lei et al., 2017] provide powerful candidates as encoder. However, to demonstrate the benefits of the proposed syntax-directed decoder in incorporating the attribute grammar for semantic restrictions, we will exploit the same encoder in Kusner et al. [2017] for a fair comparison later.

We provide a brief introduction to the particular encoder model used in Kusner et al. [2017] for a self-contained purpose. Given a program or a SMILES sequence, we obtain the corresponding parse tree using CFG and decompose it into a sequence of productions through a pre-order traversal on the tree. Then, we convert these productions into one-hot indicator vectors, in which each dimension corresponds to one production in the grammar. We will use a deep convolutional neural networks which maps this sequence of one-hot vectors to a continuous vector as the encoder.

### 3.3 Model Learning

Our learning goal is to maximize the evidence lower bound in Eq. During training, each instance $x$ is first parsed into syntax tree with CFG parser. Given the encoder, we can then map the structure input into latent space $z$. The variational posterior $q(z|x)$ is parameterized with Gaussian distribution, where the mean and variance are the output of corresponding neural networks. The prior of latent variable $p(z) = \mathcal{N}(0, I)$. Since both the prior and posterior are Gaussian, we use the closed form of KL-divergence that was proposed in Kingma & Welling [2013]. In the decoding stage, our goal is to maximize $p_\theta(x|z)$. Using the Algorithm 1, we can compute the corresponding conditional likelihood.

For efficient learning, during the training time we divide the calculation in our stochastic decoder into two phases: the first phase generates tree and the second phase only consists of a sequence of updates to the context vector. This decoupling is possible since at training time we know the all decisions in the decoder since the tree’s calculation is deterministic. In doing so, the first phase can be accelerated using multiple CPU cores in parallel and the second one can effectively be computed using any mini-batch batch optimization on GPU. In practice, we observe no significant time penalty measured in wall clock time compared to previous works.

### 4 Related Work

Generative models with discrete structured data have raised increasing interests among researchers in different domains. The classical sequence to sequence model [Sutskever et al., 2014] and its variations have also been applied to molecules [Gómez-Bombarelli et al., 2016]. Since the model is quite flexible, it is hard to generate valid structures with limited data. Techniques including data augmentation [Bjerrum, 2017], active learning [Janz et al., 2017] and reinforcement learning [Guimaraes et al., 2017] have also been proposed to tackle this issue. However, according to the empirical evaluations from Benhenda [2017], the validity is still not satisfactory. Even when the validity is enforced, the models tend to overfit to simple structures while neglect the diversity.

Since the structured data often comes with formal grammars, it is very helpful to generate its parse tree derived from CFG, instead of generating sequence of tokens directly. The Grammar VAE [Kusner et al., 2017] introduced the CFG constrained decoder for simple math expression and SMILES string generation. The rules are used to mask out invalid syntax such that the generated sequence is always from the language defined by its CFG. Parisotto et al. [2016] uses a RecursiveReverse-Recursive Neural Network (R3NN) to capture global context information while expanding with CFG production rules. Although these works follows the syntax via CFG, the context sensitive information can only be captured using variants of sequence/tree RNNs [Alvarez-Melis & Jaakkola, 2016; Dong & Lapata, 2016; Zhang et al., 2015], which may not be time and sample efficient.
In our work, we capture the semantics with proposed stochastic lazy attributes when generating structured outputs. By addressing the most common semantics to harness the deep networks, it can greatly reshape the output domain of decoder (Hu et al., 2016). As a result, we can also get a better generative model for discrete structures.

5 Experiments

We show the soundness and utility of our proposed SD-VAE using two sets of applications: programs and molecules. We compare our method with CVAE (Gómez-Bombarelli et al., 2016) and GVAE (Kusner et al., 2017). CVAE operates on character level and GVAE on the context-free grammar level. To make a fair comparison, we closely follow the experimental protocols that were set up in Kusner et al. (2017). The training details are included in Appendix B.

Our method proves significantly superior results than previous works, with better reconstruction accuracy and prior validity by large margins, while also having comparative diversity as a generative model. More importantly, our model produces a smooth continuous latent space which enables the optimization for finding best programs and molecules.

5.1 Settings

Here we first describe our datasets in detail. The first is for modeling of programs that are represented as a list of statements. Each statement is an atomic arithmetic operation on variables (labeled as v0, v1, ..., v9) and/or immediate numbers (1, 2, ..., 9). Some examples are list in the following:

\[
\begin{align*}
  v3 &= \sin(v0); v8 = \exp(2); v9 = v3 - v8; v5 = v0 + v9; \text{return: v5} \\
  v2 &= \exp(v0); v7 = v2 * v0; v9 = \cos(v7); v8 = \cos(v9); \text{return: v8}
\end{align*}
\]

Here v0 is always the input, and the variable specified by return (respectively v5 and v8 in the examples) is the output, therefore it actually represent univariate functions \( f : \mathbb{R} \to \mathbb{R} \). Note that a correct program should, besides the context-free grammar specified in Appendix A.1, also respect the semantic constraints such as that a reference to variable should not consult the unknown variables. We randomly generate 130,000 program each consisting of 1 to 5 valid statements. We hold 2000 programs out for testing and the rest for training VAE.

The second dataset is for molecules. It contains 250000 SMILES string, prepared by Kusner et al. (2017) using randomly extraction from the ZINC database (Gómez-Bombarelli et al., 2016). We use 5000 SMILES strings as the holdout set for testing and the reset for training, the same split as Kusner et al. (2017). For syntax, our formalization of SMILES follows the grammar specified in Appendix A.2, which is the same as Kusner et al. (2017).

For our SD-VAE, we address some of the most common semantics:

**Program semantics** We address the following: \( a \) variables should be defined before use, \( b \) program must return a variable, \( c \) number of statements should be less than 10.

**Molecule semantics** The SMILES semantics we addressed includes: \( a \) ringbonds should satisfy cross-serial dependencies, \( b \) explicit valence of atoms should not go beyond permitted. For more details about the semantics of SMILES language, please refer to Appendix A.3.

5.2 Reconstruction Accuracy and Prior Validity

<table>
<thead>
<tr>
<th></th>
<th>Program Reconstruction %*</th>
<th>Valid Prior %</th>
<th>Zinc SMILES Reconstruction %</th>
<th>Valid Prior %</th>
</tr>
</thead>
<tbody>
<tr>
<td>SD-VAE</td>
<td>96.46 (99.90, 99.12, 90.37)</td>
<td>100.00</td>
<td>76.2</td>
<td>43.5</td>
</tr>
<tr>
<td>GVAE</td>
<td>71.83 (96.30, 77.28, 41.90)</td>
<td>2.96</td>
<td>53.7</td>
<td>7.2</td>
</tr>
<tr>
<td>CVAE</td>
<td>13.79 (40.46, 0.87, 0.02)</td>
<td>0.02</td>
<td>44.6</td>
<td>0.7</td>
</tr>
</tbody>
</table>

Table 1: Reconstructing Accuracy and Prior Validity estimated using Monte Carlo method. Our proposed method (SD-VAE) performance significantly better than existing works.

* We also report the reconstruction % grouped by number of statements (3, 4, 5) in parentheses.
One metric of the soundness of VAE is to measure the ability of the model to encode data (in our case program/molecule) into a representation in the latent space and reconstruct the input by decoding from that point. Another metric is how often the model can decode into a valid data when the prior is randomly sampled. Since both encoding and decoding are stochastic, we follow the estimation by Monte Carlo method similar to that proposed in [Kusner et al., 2017]: For reconstruction, we select 1, 500 programs or 5, 000 molecules in the hold-out test set, and for each of them we encode it 10 times and decoded (for each encoded latent space representation) 25 times, and report the portion of decoded molecules that are the same as the input one;

For validity of prior, we sample 1000 latent representation \( \mathbf{z} \sim \mathcal{N}(O, I) \), for each of them decode 100 times, and calculate the portion of 100,000 decoded results that corresponds to valid Program or SMILES sequences.

**Program** We show in the left part of Table 1 that our model has near perfect reconstruction rate, and most importantly, a perfect valid decoding program from prior. This huge improvement is due to our model that utilizes the full semantics that previous work ignores, thus in theory guarantees perfect valid prior and in practice enables high reconstruction success rate. For a fair comparison, we run and tune the baselines in 10% of training data and report the best result. In the same place we also report the reconstruction successful rate grouped by number of statements. It is shown that our model keeps high rate even with the size of program growing.

**SMILES** The CVAE and GVAE results are included directly from [Kusner et al., 2017]. We show in the right part of Table 1 that our model produces a much higher rate of successful reconstruction and ratio of valid prior. Figure 1 in Appendix C. 2 also demonstrates some decoded molecules from our method. Note that the results we reported have not included the semantics specific to aromaticity into account. If we use an alternative kekulized form of SMILES to train the model, then the valid portion of prior can go up to 97.3%.

### 5.3 Bayesian Optimization

The variational autoencoder realizes the conversion from data (program/molecule) space to a continuous latent space via the encoder and vice versa via the decoder. This naturally leads to following two important applications: First, we can now train an extra model that predicts the data’s property from the representation in latent space, as suggested in [Gomez-Bombarelli et al., 2016]. Second and more importantly, the continuous nature of latent space makes possible the optimization of finding new data with better properties. Following the protocol used in [Kusner et al., 2017], we use Bayesian Optimization (BO) to search the programs and molecules with desired properties in latent space. Details about BO settings and parameters can be found in Appendix C.1.

![Figure 3: On the left are best programs found by each method using Bayesian Optimization. On the right are top 3 closest programs found by each method along with the distance to ground truth (lower distance is better). Both our SD-VAE and CVAE can find similar curves, but our method aligns better with the ground truth. In contrast the GVAE fails this task by reporting trivial programs representing linear functions.](image)

**Finding program** In this application the model is tasked with finding in the latent space a point corresponding a program close to a ground truth program which the model does not know. Here the closeness is measured by \( \log(1 + \text{MSE}) \) between two programs’ returned value estimated by sampling 1000 value for the input \( v_0 \in [-5, 5] \) in evenly paced sace. In Figure 5 we show that our method finds the best program to the ground truth one compared to CVAE and GVAE. Interestingly, the GVAE fails shot of finding non-trivial program. We hypothesize that this is due to the deep generative model in GVAE being highly relying on the space of production sequences from CFG syntax, which may not be optimization-friendly without constraints such as ones in our SD-VAE.
Molecules  Here we optimize the drug properties of molecules. In this problem, we ask the model to optimize for octanol-water partition coefficients (a.k.a log P), an important measurement of drug-likeness of a given molecule. As Gomez-Bombarelli et al. (2016) suggests, for drug-likeness assessment log P is penalized by other properties including synthetic accessibility score (Ertl & Schuffenhauer, 2009). In Figure 4 we show the the top-3 best molecules found by each method, where our method found molecules with better scores than previous works.

![Figure 4: Best top-3 molecules and the corresponding scores found by each method using Bayesian Optimization.](image)

5.4 PREDICTIVE PERFORMANCE OF LATENT REPRESENTATION

<table>
<thead>
<tr>
<th>Method</th>
<th>LL (Program)</th>
<th>RMSE (Program)</th>
<th>LL (Zinc)</th>
<th>RMSE (Zinc)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CVAE</td>
<td>-4.943 ± 0.058</td>
<td>3.757 ± 0.026</td>
<td>-1.812 ± 0.004</td>
<td>1.504 ± 0.006</td>
</tr>
<tr>
<td>GVAE</td>
<td>-4.140 ± 0.038</td>
<td>3.378 ± 0.020</td>
<td>-1.739 ± 0.004</td>
<td>1.404 ± 0.006</td>
</tr>
<tr>
<td>SD-VAE</td>
<td>-3.754 ± 0.045</td>
<td>3.185 ± 0.025</td>
<td>-1.697 ± 0.015</td>
<td>1.366 ± 0.023</td>
</tr>
</tbody>
</table>

Table 2: Predictive performance using encoded mean latent vector. Test LL and RMSE are reported.

We seek to to know how well our latent space predicts the properties of programs and molecules. We train the same sparse Gaussian Process as in Sec 5.3 with the same target value (namely the error for programs and the drug-likeness for molecules) for regression. We test the performance in the hold-out test dataset. In Table 2 we report the result in Log Likelihood (LL) and Regression Mean Square Error (RMSE), which show that our SD-VAE always produces latent space that are more discriminative than both CVAE and GVAE baselines. This also shows that, with a properly designed decoder, the quality of encoder will also be improved via end2end training.

5.5 DIVERSITY OF GENERATED MOLECULES

<table>
<thead>
<tr>
<th>Similarity Metric</th>
<th>MorganFp</th>
<th>MACCS</th>
<th>PairFp</th>
<th>TopologicalFp</th>
</tr>
</thead>
<tbody>
<tr>
<td>GVAE</td>
<td>0.92 ± 0.10</td>
<td>0.83 ± 0.15</td>
<td>0.94 ± 0.10</td>
<td>0.71 ± 0.14</td>
</tr>
<tr>
<td>SD-VAE</td>
<td>0.92 ± 0.09</td>
<td>0.83 ± 0.13</td>
<td>0.95 ± 0.08</td>
<td>0.75 ± 0.14</td>
</tr>
</tbody>
</table>

Table 3: Diversity as statistics from pair-wise distances measured as $1 - s$, where $s$ is one of the similarity metrics. So higher values indicate better diversity. We show mean ± stddev of $\binom{100}{2}$ pairs among 100 molecules. Note that we report results from GVAE and our SD-VAE, because CVAE has very low valid priors, thus completely only failing this evaluation protocol.

Inspired by Benhenda (2017), here we seek to measure the diversity of generated molecules as an assessment of our methods. The intuition is that a good generative model should be able to generate diverse data and avoid model collapse in the learned space. In detail, we conduct this experiment in SMILES dataset, where we sample 100 points from the prior distribution, and for each point, we associate it as a molecule, which is the most frequent occurring valid SMILES decoded (we use 50 decoding attempts since the decoding is stochastic). We then, with one of several molecular similarity, compute the pair-wise similarity and report the mean and standard deviation in Table 3. We see both methods do not have the model collapse problem, while producing similar diversity scores. It indicates that although our method has more restricted decoding space than baselines, the diversity is not sacrificed. This is because we never rule-out the valid molecules. And a more compact decoding space leads to much higher probability in obtaining valid molecules.
5.6 Visualizing the Latent Space

We seek to visualize the latent space as an assessment of how well our generative model is able to produces a coherent and smooth space of program and molecules.

Program  Following Bowman et al. [2016], we visualize the latent space of program by interpolation between two programs. More specifically, given two programs which are encoded to \(p_a\) and \(p_b\) respectively in the latent space, we pick 9 evenly spaced points between them and for each point as prior generate a program using the decoder. In Table 4 we compare our results with previous works. Our SD-VAE can pass though points in the latent space that can be decoded into valid programs without error and with visually more smooth interpolation than previous works. Meanwhile, CVAE makes both syntactic and semantic errors, and GVAE produces only semantic errors (reference of undefined variables), but still in a considerable amount.

<table>
<thead>
<tr>
<th>CVAE</th>
<th>GVAE</th>
<th>SD-VAE</th>
</tr>
</thead>
<tbody>
<tr>
<td>(v^0=\text{cos}(3), y^0=\text{map}(7), y^0=\text{vol}(5), v^0=\text{return}(v))</td>
<td>(v^0=\text{cos}(3), y^0=\text{map}(7), y^0=\text{vol}(5), v^0=\text{return}(v))</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4: Interpolation between two valid programs (the top and bottom ones in brown) where each program occupies a row. Programs in red are with syntax errors. Statements in blue are with semantic errors such as referring to unknown variables. Rows without coloring are correct programs. Observe that when a model passes points in its latent space, our proposed SD-VAE enforces both syntactic and semantic constraints while making visually more smooth interpolation. In contrast, CVAE makes both kinds of mistakes, GVAE avoids syntactic errors but still produces semantic errors, and both methods produce subjectively less smooth interpolations.

SMILES  For molecules, we visualize the latent space in 2 dimensions. We first embed a random molecule into latent space. Then we randomly generate 2 orthogonal unit vectors \(A\). To get the latent representation of neighborhood, we interpolate the 2-D grid and project back to latent space with pseudo inverse of \(A\). Finally we show decoded molecules. In Figure 5 we present two of such grid visualizations. Subjectively compared with figures in Kusner et al. [2017], our visualization is characterized by having smooth differences between neighboring molecules, and more complicated decoded structures.

Figure 5: Latent Space visualization. We start from the center molecule and decode the neighborhood latent vectors (neighborhood in projected 2D space).
REFERENCES


Appendix

A GRAMMAR

A.1 GRAMMAR FOR PROGRAM SYNTAX

The syntax grammar for program is a generative context-free grammar starting with \langle program \rangle.

\[
\begin{align*}
\langle program \rangle & \rightarrow \langle stat \ list \rangle \\
\langle stat \ list \rangle & \rightarrow \langle stat \rangle \ ' ; ' \ (stat \ list) \ | \ (stat) \\
\langle stat \rangle & \rightarrow \langle assign \rangle \ | \ \langle return \rangle \\
\langle assign \rangle & \rightarrow \langle lhs \rangle \ '=' \ \langle rhs \rangle \\
\langle return \rangle & \rightarrow \ ' return ' \ \langle lhs \rangle \\
\langle lhs \rangle & \rightarrow \langle var \rangle \\
\langle var \rangle & \rightarrow \ ' v ' \ \langle var \ id \rangle \\
\langle digit \rangle & \rightarrow \ ' 1 ' \ | \ ' 2 ' \ | \ ' 3 ' \ | \ ' 4 ' \ | \ ' 5 ' \ | \ ' 6 ' \ | \ ' 7 ' \ | \ ' 8 ' \ | \ ' 9 ' \\
\langle rhs \rangle & \rightarrow \ (expr) \\
\langle expr \rangle & \rightarrow \ (unary \ expr) \ | \ (binary \ expr) \\
\langle unary \ expr \rangle & \rightarrow \ (unary \ op) \ \langle operand \rangle \ | \ (unary \ func) \ \langle \ (operand) \ ' \rangle ' \\
\langle binary \ expr \rangle & \rightarrow \ (operand) \ \langle binary \ op \rangle \ \langle operand \rangle \\
\langle unary \ op \rangle & \rightarrow \ ' + ' \ | \ ' - ' \\
\langle unary \ func \rangle & \rightarrow \ ' \ sin ' \ | \ ' \ cos ' \ | \ ' \ exp ' \\
\langle binary \ op \rangle & \rightarrow \ ' + ' \ | \ ' - ' \ | \ ' * ' \ | \ ' / ' \\
\langle operand \rangle & \rightarrow \ \langle var \rangle \ | \ \langle immediate \ number \rangle \\
\langle immediate \ number \rangle & \rightarrow \ \langle digit \rangle \ \langle . \rangle \ \langle digit \rangle \\
\langle digit \rangle & \rightarrow \ ' 0 ' \ | \ ' 1 ' \ | \ ' 2 ' \ | \ ' 3 ' \ | \ ' 4 ' \ | \ ' 5 ' \ | \ ' 6 ' \ | \ ' 7 ' \ | \ ' 8 ' \ | \ ' 9 ' \\
\end{align*}
\]

A.2 GRAMMAR FOR MODECULE SYNTAX

Our syntax grammar is based on OpenSMILES standard, a generative context-free grammar starting with \langle s \rangle.

\[
\begin{align*}
\langle s \rangle & \rightarrow \ \langle atom \rangle \\
\langle smiles \rangle & \rightarrow \ \langle chain \rangle \\
\langle atom \rangle & \rightarrow \ \langle bracket \ atom \rangle \ | \ \langle aliphatic \ organic \rangle \ | \ \langle aromatic \ organic \rangle \\
\langle aliphatic \ organic \rangle & \rightarrow \ \langle B ' \ | \ ' C ' \ | \ ' N ' \ | \ ' O ' \ | \ ' S ' \ | \ ' F ' \ | \ ' I ' \ | \ ' C l ' \ | \ ' B r ' \\
\langle aromatic \ organic \rangle & \rightarrow \ \langle c ' \ | \ ' n ' \ | \ ' o ' \ | \ ' s ' \\
\langle bracket \ atom \rangle & \rightarrow \ ' [ ' \ \langle baracekt \ atom \ \langle isotope \rangle \ ' ] ' \\
\langle baracekt \ atom \ \langle isotope \rangle \rangle & \rightarrow \ \langle isotope \rangle \ \langle symbol \rangle \ \langle baracekt \ atom \ \langle chiral \rangle \rangle \\
&& \ | \ \langle isotope \rangle \ \langle symbol \rangle \ \langle baracekt \ atom \ \langle chiral \rangle \rangle \\
&& \ | \ \langle isotope \rangle \ \langle symbol \rangle \ | \ \langle symbol \rangle \\
\langle baracekt \ atom \ \langle chiral \rangle \rangle & \rightarrow \ \langle chiral \rangle \ \langle baracekt \ atom \ \langle h \ count \rangle \rangle \\
&& \ | \ \langle baracekt \ atom \ \langle h \ count \rangle \rangle \\
&& \ | \ \langle chiral \rangle \\
\langle baracekt \ atom \ \langle h \ count \rangle \rangle & \rightarrow \ \langle h \ count \rangle \ \langle baracekt \ atom \ \langle charge \rangle \rangle \\
&& \ | \ \langle baracekt \ atom \ \langle charge \rangle \rangle \\
&& \ | \ \langle h \ count \rangle \\
\langle baracekt \ atom \ \langle charge \rangle \rangle & \rightarrow \ \langle charge \rangle \\
\langle symbol \rangle & \rightarrow \ \langle aliphatic \ organic \rangle \ | \ \langle aromatic \ organic \rangle \\
\langle isotope \rangle & \rightarrow \ \langle digit \rangle \ | \ \langle digit \rangle \ \langle digit \rangle \ | \ \langle digit \rangle \ \langle digit \rangle \ \langle digit \rangle \\
\langle digit \rangle & \rightarrow \ ' 1 ' \ | \ ' 2 ' \ | \ ' 3 ' \ | \ ' 4 ' \ | \ ' 5 ' \ | \ ' 6 ' \ | \ ' 7 ' \ | \ ' 8 ' \\
\end{align*}
\]
A.3 EXAMPLES OF SMILES SEMANTICS

Here we provide more explanations of the semantics constraints that contained in SMILES language for molecules.

Specifically, the semantics we addressed here are:

1. **Ringbond matching**: the ringbonds should come in pairs. Each pair of ringbonds has an index and bond-type associated. So the SMILES semantics requires something exactly the same as well-known cross-serial dependencies (CSD). CSD is also appeared in some natural languages, such as Dutch and Swiss-German. See Figure 6 for an illustration.

2. **Explicit valence control**: Intuitively, the semantics says each atom cannot have too many bonds associated with it. For example, a normal Carbon atom has maximum valence of 4. This means associating a Carbon atom with two triple-bonds will violate the semantics.

A.4 DEPENDENCY GRAPH INTRODUCED BY ATTRIBUTE GRAMMAR

Suppose there is a production \( r = u_0 \rightarrow u_1 u_2 \ldots u_{|\beta|} \in R \) and an attribute \( u_i.a \) we denote the dependency set \( D^r(u_i.a) = \{ u_j.b | u_j.b \text{ is required for calculating } u_i.a \} \). The union of all dependency sets \( D_T^{(att)} = \bigcup_{r \in \mathcal{T}, u_i \in \mathcal{R}} D^r(u_i.a) \) induces a dependency graph, where nodes are the attributes and directed edges represents the dependency relationships between those attributes computation. Here \( \mathcal{T} \) is an (partial or full) instantiation of the generated syntax tree of grammar \( G \). Let \( D^r(u_i) = \{ u_j | \exists b : u_j.b \in D^r(u_i.a) \} \) and \( D_T = \bigcup_{r \in \mathcal{T}, u_i \in \mathcal{R}} D^r(u_i), i.e., D_T \) is constructed from \( D_T^{(att)} \) by merging nodes with the same symbol but different attributes, we call \( D_T^{(att)} \) is noncircular if the corresponding \( D_T \) is noncircular.

In our paper, we assume the noncircular property of the dependency graph. Such property will be exploited for top-down generation in our decoder.

B TRAINING DETAILS

Since our proposed SD-VAE differentiate itself from previous works (CVAE, GVAE) on the formalization of syntax and semantics, we therefore use the same deep neural network model architecture
for a fair comparison. In encoder, we use 3-layer 1 dimension convolution neural networks (CNNs) following by a full connected layer, whose output would be feed to two separate affine layers for producing $\mu$ and $\sigma$ respectively in reparameterization trick; and in decoder we use 3 layer RNNs following by an affine layer activated by softmax that gives probability for each production rule. In detail, we use 56 dimensions the latent space and the dimension of layers as the same number as in Kusner et al. (2017). As for implementation, we use Kusner et al. (2017)’s open sourced code for baselines, and implement our model with PyTorch framework.$^1$

In a 10% validation set we turn the following hyper parameters and report the test result from setting with best valid loss. For a fair comparison, all tunings are also conducted in the baselines.

We use $\text{ReconstructLoss} + \alpha \text{KLdivergence}$ as the loss function for training. A natural setting is $\alpha = 1$, but Kusner et al. (2017) suggested in their open-sourced implementation$^2$ that using $\alpha = 1/\text{Latent Dimension}$ would leads to better results. We explore both settings.

### C More experiment details

![Figure 7: Visualization of reconstruction. The first column in each figure presents the target molecules. We first encode the target molecules, then sample the reconstructed molecules from their encoded posterior.](image)

#### C.1 Bayesian optimization

The Bayesian optimization is used for searching latent vectors with desired target property. For example, in symbolic program regression, we are interested in finding programs that can fit the given input-output pairs; in drug discovery, we are aiming at finding molecules with maximum drug likeness. To get a fair comparison with baseline algorithms, we follow the settings used in Kusner et al. (2017).

Specifically, we first train the variational autoencoder in an unsupervised way. After obtaining the generative model, we encode all the structures into latent space. Then these vectors and corresponding property values (i.e., estimated errors for program, or drug likeness for molecule) are used to train a sparse Gaussian process with 500 inducing points. This is used later for predicting properties in latent space. Next, 5 iterations of batch Bayesian optimization with the expected improvement

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$^1$http://pytorch.org/

$^2$https://github.com/mkusner/grammarVAE/issues/2
(EI) heuristic is used for proposing new latent vectors. In each iteration, 50 latent vectors are proposed. After the proposal, the newly found programs/molecules are then added to the batch for next round of iteration.

During the proposal of latent vectors in each iteration, we perform 100 rounds of decoding and pick the most frequent decoded structures. This helps regulates the decoding due to randomness, as well as increasing the chance for baselines algorithms to propose valid ones.

C.2 Reconstrution

We visualize some reconstruction results of SMILES in Figure 7. One can see that, in most cases the decoder successfully recover the exact origin input. Due to the stochasticity of decoder, it may have some small variations.