# **Appendix**

## A. Grammars for equations and SMILES

The grammar for the single-variable equations includes 3 binary operators, 2 unary operators, 3 constants, and grouping symbols; the start symbol is S. Training data for the VAE came by generating 100,000 different equations with parse tree depth less than 7, corresponding to equations which can be produced using up to 15 production rule applications.

The grammar for SMILES is based on the official OPENSMILES specification (Weininger, 1988), and starts with smiles.

```
smiles \rightarrow chain
atom → bracket_atom | aliphatic_organic | aromatic_organic
BAI \rightarrow isotope symbol BAC \mid symbol BAC \mid isotope symbol \mid symbol
BAC → chiral BAH | BAH | chiral
BAH → hcount BACH | BACH | hcount
BACH → charge class | charge | class
symbol → aliphatic_organic | aromatic_organic
isotope \rightarrow DIGIT | DIGIT DIGIT DIGIT DIGIT DIGIT DIGIT DIGIT DIGIT \rightarrow '1' | '2' | '3' | '4' | '5' | '6' | '7' | chiral \rightarrow '@' | '@@'
                              | '4' | '5' | '6' | '7' | '8'
hcount \rightarrow 'H' | 'H' DIGIT charge \rightarrow '-' | '-' DIGIT | '-' DIGIT DIGIT | '+' | '+' DIGIT | '+' DIGIT DIGIT bond \rightarrow '-' | '=' | '#' | '/' | '\'
ringbond \rightarrow DIGIT + bond DIGIT
branched\_atom \rightarrow atom \mid atom RB \mid atom BB \mid atom RB BB
RB → RB ringbond | ringbond
BB \rightarrow BB branch | branch
branch \rightarrow '(' chain ')' | '(' bond chain ')'
chain → branched_atom | chain branched_atom | chain bond branched_atom
```

#### **B.** Network structure

We briefly overview recent sequence modeling advances which inform our encoder and decoder models. An encoder  $q_{\phi}(\mathbf{z}|\mathbf{X})$  takes a sequence of T timesteps  $\mathbf{X} = [\mathbf{x}_1, \dots, \mathbf{x}_T]$  as input and returns a distribution over real-valued vectors z, whereas the decoder  $p_{\theta}(\mathbf{x}|\mathbf{z})$  takes a real-valued vector  $\mathbf{z}$  as input and generates a distribution over sequences themselves  $\mathbf{X}$ . We use the same encoder and decoder as Gómez-Bombarelli et al. (2016b), inspired by Bowman et al. (2016), with a few modifications to incorporate a grammar as described in Section 3. We begin by encoding the data as sequences of one-hot vectors. We then pass these vectors through our encoder network which consists of 3 layers of one-dimensional convolutions (Kalchbrenner et al., 2014). We then flatten the resulting sequence into a vector and pass it through a fully connected layer. We then pass the resulting vector through two separate fully connected layers to produce the mean and variance of the latent distribution  $q_{\phi}(\mathbf{z}|\mathbf{X})$  over  $\mathbf{z}$ . To decode, we sample a  $\mathbf{z}$  from this distribution and pass it through a fully connected layer. We then repeat the output of this layer  $T_{max}$  times, producing a matrix of dimension  $H \times T_{max}$ , where H is the dimensionality of  $\mathbf{z}$ . This can be thought of as a sequence over  $T_{max}$  timesteps. We then pass this matrix through 3 layers of GRU recurrent units (Cho et al., 2014), and finally through a fully connected layer, whose weights are repeated at each timestep. The result are the logit vectors  $\mathbf{F}$ .

## C. Implementation details

We make two implementation changes to the standard VAE. We begin by reconsidering the ELBO as the sum of a reconstruction loss term and a negative KL-divergence term (Kingma & Welling, 2014),

$$\mathcal{L}(\phi, \theta; \mathbf{X}) = \mathbb{E}_{q_{\phi}(\mathbf{z}|\mathbf{X})} \left[ \log p_{\theta}(\mathbf{X}|\mathbf{z}) \right] - KL(q_{\phi}(\mathbf{z}|\mathbf{X})||p(\mathbf{z})).$$

The first term ensures that the latent representation z serves as an accurate 'encoding' or 'compression' of the original data X. The second term serves to regularize the latent representation so that the latent space is not too complex. Combined together, these terms are aimed at producing a latent space which is the most efficient encoding of the data X. However, when optimizing this ELBO in the usual way, we noticed that the VAE had the tendency to ignore the first term and simply

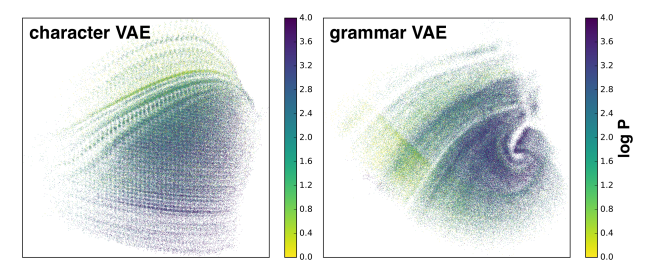


Figure 6. The logP values of a 2-dimensional character and grammar VAE. The grammar VAE leads to a low-dimensional latent space which is visually smother with respect to the property of interest.

try to optimize the second term by trying to have the encoder  $q_{\phi}(\mathbf{z}|\mathbf{X})$  match the prior distribution  $p(\mathbf{z})$ . Thus, the learned latent representation  $\mathbf{z}$  contains essentially no information about  $\mathbf{X}$ . We believe this happened because our decoder is such a powerful recurrent neural network, that it easily produces decodings that 'appear' similar enough to the data  $\mathbf{X}$ . We note that this behavior has been observed in other recent work (Zhao et al., 2017). To fix this we made two small modifications, inspired by the character VAE on which our code is based<sup>4</sup>. First, we scale the standard deviation of the encoding distribution  $q_{\phi}(\mathbf{z}|\mathbf{X})$  by 0.01. We found that this helped the VAE produce more consistent encodings  $\mathbf{z}$  for data  $\mathbf{X}$ , thus making the latent-space better resemble the data. Second, we divide the KL-divergence term by the number of latent dimensions H. This effectively downweights the importance of the KL term so the VAE focusses more of an effort at improving reconstruction.

Table 6. Reconstruction accuracy and sample validity results.

Method	% Reconstruct	% Prior Valid
GVAE	53.7	25.9
CVAE	44.6	0.001

### D. Additional experiments

Molecule reconstruction & validity. We characterize how well the VAE models over molecules are able to reconstruct input sequences from their corresponding latent representations and to also decode valid sequences when sampling from the prior in latent space. Comparisons of full reconstruction accuracy for both the character and grammar VAEs are shown in Table 6. To compute reconstruction error we start with 5000 true molecules from a hold-out set. For each molecule we encode it 10 times, and we decode each encoding 100 times (as encoding and decoding are stochastic). This results in 1000 decoded molecules for each of the 5000 input molecules. We compute the average of these 1000 decodings that are identical to the input molecule. We then average these averages across all 5000 inputs to get the percentage of molecules that reconstruct out of the 5,000,000 attempts. To compute the percentage prior validity we sample 1000 latent points from the prior distribution  $p(\mathbf{z}) = \mathcal{N}(0, \mathbf{I})$ . We decode each of these points 500 times and test which of the decoded SMILES strings correspond to valid molecules. We average across all 1000 points and 500 trials to yield the percentages in Table 6. These results clearly indicate that the proposed GVAE has higher reconstruction accuracy, and produces a higher proportion of valid sequences when sampling from the prior.

<sup>&</sup>lt;sup>4</sup>https://github.com/maxhodak/keras-molecules

#### **Grammar Variational Autoencoder**

**LogP Visualization.** To visualize the latent space of the VAEs on molecules we train a CVAE and GVAE on the ZINC dataset (Gómez-Bombarelli et al., 2016b) with a 2-dimensional latent space. We plot the training set colored by the logP values of the molecules in Figure 6. We note that the CVAE seems to disperse molecules with higher logP values (corresponding to molecules with better drug properties) throughout a large region in the lower half of the latent space. The GVAE on the other hand concentrates molecules with high logP in a small region of latent space. We suspect this makes Bayesian optimization for molecules with high logP much easier in the GVAE.