# Saturn: Sample-efficient Generative Molecular Design using Memory Manipulation

#### **Anonymous Author(s)**

Affiliation Address email

#### **Abstract**

Generative molecular design for drug discovery has very recently achieved a wave of experimental validation, with language-based backbones being the most common architectures employed. The most important factor for downstream success is whether an *in silico* oracle is well correlated with the desired end-point. To this end, current methods use cheaper proxy oracles with higher throughput before evaluating the most promising subset with high-fidelity oracles. The ability to directly optimize high-fidelity oracles would greatly enhance generative design and be expected to improve hit rates. However, current models are not efficient enough to consider such a prospect, exemplifying the sample efficiency problem. In this work, we introduce **Saturn**, which leverages the Augmented Memory algorithm and demonstrates the first application of the Mamba architecture for generative molecular design. We elucidate how experience replay with data augmentation improves sample efficiency and how Mamba synergistically exploits this mechanism. Saturn outperforms 22 models on multi-parameter optimization tasks relevant to drug discovery and may possess sufficient sample efficiency to consider the prospect of directly optimizing high-fidelity oracles. The code is available at https://figshare.com/s/6040d65bfbfc29d6fedf.

# 1 Introduction

2

3

5

6

7

8

9

10

11

12

13

14

15

16

17

19

20

21

23

24

25

27

28

29

30

31

32

33

34

35

Within the last year, there has been a surge of works reporting experimental validation of generative molecular design for drug discovery <sup>1–7</sup>. The fundamental task of generative molecular design is to simulate (from a distribution) molecules with tailored property profiles. All generative models achieve this in one of two ways: distribution learning, where a base model is subjected to transfer learning on a set of known positives, and goal-directed generation, which encompasses both conditional generation and using an optimization algorithm to shift the distribution. Experimental validation has been demonstrated for all methods, but with a notable over-representation from optimization algorithms (as of the last 6 months), and particularly reinforcement learning (RL)<sup>2-7</sup>. Algorithmic molecular optimization always proceeds via the following workflow: generate molecules, assess desirability (using an in silico oracle), update the model, and repeat. When assessing the suitability of molecules absent experimental validation, the crucial indicator to success is correlation of an in silico oracle to the actual end-point. All protocols that *directly* optimize for an oracle (without the use of a surrogate predictor) follow a funnel workflow where less resource-intensive oracles are initially used to prioritize the most promising subset for evaluation with computationally expensive high-fidelity oracles. A concrete and ubiquitous example is designing molecules with high binding affinity to a protein target. By far the most common oracle used to estimate binding affinity is molecular docking, and many works<sup>8–14</sup> have demonstrated the ability to generate molecules with improved docking scores. However, docking scores are often poorly correlated with binding affinity, especially when applied out-of-the-box <sup>8,15</sup>. Correspondingly, the most promising candidates from docking are subjected to higher-fidelity oracles, particularly molecular dynamics (MD) simulations, which offer a much more accurate estimation of binding affinity <sup>15–18</sup>. *Directly* optimizing high-fidelity oracles offers the prospect of learning the distribution and can greatly improve the quality of the generated set <sup>19</sup>. However, doing so is infeasible due to computational cost, exemplifying the sample efficiency problem. Either simulation protocols become much faster without sacrificing accuracy, or generative models become *sufficiently efficient* to optimize under an acceptable oracle budget.

Recently, the proposed Practical Molecular Optimization (PMO)<sup>20</sup> benchmark assessed 25 models across 23 optimization tasks under a 10,000 oracle budget. Since then, other works have explicitly 45 constrained the oracle budget on various drug discovery optimization tasks <sup>10–14,21,22</sup>. Results from the 46 PMO benchmark show that language-based models are, on average, the most sample-efficient models. 47 More recently, Guo et al. 21 proposed Augmented Memory which is built on REINVENT 23,24. It 48 combines experience replay with SMILES augmentation 25 and achieves the new state-of-the-art on 49 the PMO benchmark. In this work, we push towards the prospect of direct optimization of high-fidelity 50 oracles and release **Saturn**. First, we elucidate the mechanism of Augmented Memory<sup>21</sup>, which uses an LSTM <sup>26</sup> recurrent neural network (RNN) as the language model backbone, and characterize how data augmentation and experience replay improve sample efficiency. Next, we systematically 53 assess more advanced generative architectures from just RNNs<sup>26</sup> to decoder transformers<sup>27,28</sup>, and 54 the recent Mamba<sup>29</sup> state space model (SSM). Our results show that the Mamba architecture, in 55 conjunction with data augmentation and experience replay, displays synergistic behavior to improve 56 sample efficiency. Our contribution is as follows: 57

- 1. We show the first application of Mamba<sup>29</sup> for molecular generative design and specifically for goal-directed generation with reinforcement learning.
- 2. We elucidate the mechanism into *how* Augmented Memory<sup>21</sup> improves sample efficiency, as the original work only showed its empirical benefits.
- 3. We comprehensively evaluate language model backbones (> 5,000 experiments) including RNN, decoder transformer<sup>27,28</sup>, and Mamba<sup>29</sup>, which enables us to characterize model-intrinsic and scaling properties that lead to improved sample efficiency.
- 4. We propose **Saturn**, which leverages Mamba<sup>29</sup> and outperforms 22 models on multiparameter optimization drug discovery tasks with fixed oracle budgets.

# 2 Related Work

58

59

60

61

62

63

65

68

71 72

73

75

78

79

80

81

82

**Sample Efficiency in Goal-directed Molecular Design.** The goal of inverse design is to achieve *tailored* molecular generation. Existing works have tackled this problem using a variety of architectures, including SMILES <sup>30</sup>-based RNNs <sup>9,23,24,31-35</sup>, transformers <sup>9,27,36-42</sup>, variational autoencoders (VAEs) <sup>43-46</sup>, adversarial approaches <sup>47-53</sup>, graph-based models <sup>11,54-59</sup>, GFlowNets <sup>10,60,61</sup>, genetic algorithms (GAs) <sup>13,14,62,63</sup>, and diffusion models <sup>12,64,65</sup>. However, many works do not explicitly consider an oracle budget (or use a very lenient budget) and focus mostly on showing that goal-directed generation is possible. The release of the PMO benchmark <sup>20</sup> highlighted that improvements in sample efficiency are vital to even consider the prospect of directly optimizing high-fidelity oracles. Since then, more recent works <sup>10-14,21,22</sup> have enforced fixed oracle budgets when comparing performance with other methods. In this work, we consider fixed oracle budgets in all experiments and, importantly, investigate optimization under small batch sizes, which becomes pertinent when considering high-fidelity oracles that require *at least* one GPU per molecule, which quickly imposes a practical constraint.

Language-based Molecular Generative Models. Text is one of the most widely used molecular representations, with common ones being simplified molecular-input line-entry systems (SMILES)<sup>30</sup> and self-referencing embedded strings (SELFIES)<sup>66,67</sup>. Recent work has shown that the former is generally more performant, despite not enforcing 100% validity<sup>20,68</sup>. Leveraging advances in natural language processing (NLP), language-based molecular generative models are amongst the first and still widely used models, encompassing RNNs<sup>9,23,24,31–35</sup>, transformers<sup>9,27,28,36–42</sup>, and recently SSM S4<sup>69</sup>. In early benchmarks (GuacaMol<sup>70</sup> and MOSES<sup>71</sup>), language-based models have been shown to essentially solve the validity, uniqueness, and novelty metrics. Subsequently, the non-injective syntax of SMILES confers advantageous properties for generative design. Specifically, a single molecule can be expressed as at least *N* (number of heavy atoms) SMILES, in a process known as SMILES augmentation, enumeration, or randomization<sup>25</sup>. This mechanism can be exploited to pre-train

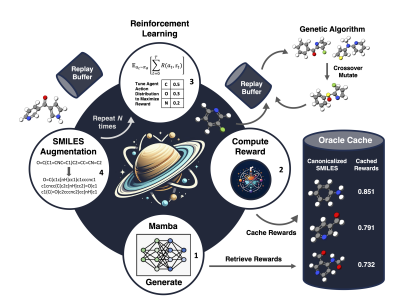


Figure 1: Saturn generative workflow. All generated SMILES and their rewards are stored in the Oracle Cache after canonicalization. A genetic algorithm can be optionally applied using the replay buffer as the parent population. Augmented Memory is used to update the agent numerous times.

models under low data regimes to generalize in chemical space <sup>72–74</sup>, improve sample efficiency <sup>21,35</sup>, and perform transfer learning with a single positive example <sup>75</sup>. Despite the recent trend towards 3D molecular generation <sup>64,65</sup>, language-based models have demonstrated the ability to generate molecules that satisfy 3D-dependent objectives, such as docking <sup>8</sup> in a sample-efficient manner <sup>21,22</sup>. This suggests that language-based models are not entirely 3D-naive and can effectively explore relevant regions of the 3D chemical space. Finally, language models are amongst the most sample-efficient models in the PMO benchmark <sup>20,21</sup> and most works achieving experimental validation of a generated molecule incorporate SMILES-based models <sup>2–7</sup>.

# 3 Method

100

104

105

106

107

108

109

In this section, each component of Saturn (Fig. 1) is described: the language model backbone for molecular generation, the Augmented Memory<sup>21</sup> RL algorithm, the GA, and specific details into key components responsible for sample efficiency and mitigating mode collapse.

Autoregressive Language Model Backbone for Molecular Generation. Molecules are represented as SMILES  $^{30}$  and the task of goal-directed generation is cast as an RL problem. Let  $S_t$  denote the state space representing all intermediate token sequences during molecular generation. The action space,  $A_t(s_t)$ , is defined as the conditional token distribution induced by the policy,  $\pi_\theta$ , and parameterized by a language model backbone. Generation follows a Markov process, and thus, sampling a SMILES, x, is given by the product of conditional token probabilities (Eq. 1):

$$P(x) = \prod_{t=1}^{T} \pi_{\theta_{\text{Agent}}}(a_t \mid s_t)$$
 (1)

The general objective in RL is to maximize the expected reward (Eq. 2):

$$J(\theta) = \mathbb{E}_{a_t \sim \pi_{\theta_{Agent}}} \left[ \sum_{t=1}^{T} R(a_t, s_t) \right]$$
 (2)

R is the reward function and can represent any arbitrary multiparameter optimization (MPO) objective and  $\sigma$  is a scalar factor modulating its effect. Next, the Augmented Likelihood (Eq. 3) is defined, where the prior is the pre-trained model with *frozen* weights:

$$\log \pi_{\text{Augmented}}(x) = \log \pi_{\text{prior}}(x) + \sigma R(x)$$
 (3)

The reward is defined as  $\log \pi_{\text{Augmented}}$  -  $\log \pi_{\theta_{\text{agent}}}$ . Following previous works <sup>21,23,76</sup>, maximizing Eq. 2 is equivalent (up to a factor) to minimizing the squared difference between the Augmented Likelihood and the Agent Likelihood (Eq. 4):

$$L(\theta) = \frac{1}{|B|} \left[ \sum_{a \in A^*} (\log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{agent}}}) \right]^2 \tag{4}$$

 $A^*$  is defined as the actions taken across all time-steps in a given batch. During optimization, the expected reward (Eq. 2) is approximated by sampling a batch, B, of SMILES. The batch size controls for variance as approximating the expectation with fewer samples is necessarily more noisy. See Appendix A.4 for full details on the algorithm and pseudo-code.

Augmented Memory. In Saturn, Augmented Memory maintains a replay buffer of the top 100 SMILES ranked by their rewards. At each generation epoch, the SMILES in the buffer are augmented (randomized)<sup>25</sup> and the agent is updated N augmentation rounds following Eq. 4. Following Blaschke et al.<sup>77</sup>, a Diversity Filter (DF) stores the Bemis-Murcko<sup>78</sup> scaffolds of every SMILES generated. If a scaffold is generated more than a permitted threshold (M = 10 in this work), its reward is truncated to 0. Before executing Augmented Memory, scaffolds associated with penalized rewards are purged from the buffer, preventing mode collapse.

Genetic Algorithm. Saturn adapts the GraphGA<sup>63</sup> algorithm where the replay buffer is treated as the parent population. The motivation is to generate more high reward SMILES to *replace* the buffer SMILES, under the hypothesis that on average, these too, will be high reward (Appendix B.5).

Oracle Caching. In this work, we make the assumption that oracle evaluations are *near deterministic* and store every SMILES generated and its associated reward in a cache. If the same SMILES is generated at a later epoch, the reward is retrieved from the cache and does not impose an oracle call.

#### 4 Results and Discussion

134

139

152

153

155

The results section is comprised of three parts: formulating Saturn, demonstrating sample efficiency in an MPO docking task, and another MPO docking task with comparison to 22 models (including two dataset screening baselines). Every experiment was run across 10 seeds (0-9 inclusive), comprising 4,840 and 200 total runs on test and molecular docking experiments, respectively.

# 4.1 Part 1: Elucidating the Optimization Dynamics of Saturn

We begin by identifying the optimal architecture and hyperparameters for Saturn. First, we experiment with varying the batch size and augmentation rounds of Augmented Memory algorithm<sup>21</sup>, and explicitly demonstrate the trade-off between sample efficiency and diversity. Unlike the original Augmented Memory work, which used an RNN backbone, we investigate more advanced architectures: decoder transformer <sup>27,28</sup> and Mamba <sup>29</sup>. Our analysis elucidates how SMILES augmentation, combined with these architectures, synergistically improves sample efficiency in Saturn.

Experimental Details. Similar to Guo et al.  $^{22}$ , we define a test experiment with the following MPO objective: molecular weight (MW) < 350 Da, number of rings  $\geq$  2, and maximize topological polar surface area (tPSA). Optimizing this objective *requires* generating molecules with rings saturated with heteroatoms, which are dissimilar from the training data. Hence, it is also testing out-of-distribution optimization. All experiments in this section were run across 10 seeds (0-9 inclusive) with an oracle budget of 1,000, and the models were pre-trained with ChEMBL 33  $^{79}$  (Appendix B.1).

**Metrics.** The sample efficiency metrics are **Yield** and **Oracle Burden** (OB). Yield is the number of *unique* generated molecules above a reward threshold, and OB is the number of oracle calls required to generate *N unique* molecules above a reward threshold. The reward threshold in this experiment is 0.7 as molecules start to possess saturated heteroatom rings<sup>22</sup>. Most configurations successfully generate at least *some* molecules passing this threshold within the budget, enabling us to report statistics.

Table 1: Sample efficiency across architectures (batch size 16). 1,000 oracle budget. All metrics are computed at the 0.7 reward threshold. IntDiv1<sup>71</sup> is the internal diversity, Scaffolds is the number of unique Bemis-Murcko<sup>78</sup> scaffolds, OB is Oracle Burden (oracle calls required to generate *N* unique molecules). The number in parentheses in the OB statistics represents how many runs out of 10 were successful. Repeats are the number of times an identical SMILES was generated during the run. The mean and standard deviation across 10 seeds (0-9 inclusive) is reported.

Model	Aug. Rounds	Yield (↑)	IntDiv1 (↑)	Scaffolds $(\uparrow)$	OB 1 (↓)	OB 10 (↓)	OB 100 (↓)	Repeats
RNN	5	107±58	0.814±0.036	101±54	480±118 (10)	721±109 (10)	916±53 (4)	7±7
	6	121±80	0.791±0.040	107±68	493±214 (10)	713±15 (10)6	895±107 (5)	12±11
	7	144±107	0.776±0.026	117±86	467±186 (10)	684±136 (10)	871±116 (6)	38±82
	8	120±95	$0.734\pm0.128$	104±85	481±288 (10)	653±145 (8)	854±54 (5)	18±28
	9	141±104	0.783±0.048	112±72	453±211 (10)	654±154 (9)	871±104 (6)	59±95
	10	106±76	0.76±0.056	84±63	510±201 (10)	733±122 (9)	913±64 (5)	43±47
Decoder	5	154±93	0.748±0.052	122±70	439±151 (10)	679±128 (10)	907±92 (8)	90±90
Transformer	6	116±94	0.748±0.039	86±64	517±165 (10)	728±158 (10)	904±126 (5)	73±42
	7	108±85	0.747±0.051	71±50	510±222 (10)	740±127 (9)	868±48 (4)	126±63
	8	108±94	0.708±0.109	72±57	538±164 (10)	742±116 (9)	887±87 (4)	150±72
	9	78±83	0.687±0.116	51±55	614±244 (10)	790±150 (8)	890±62 (3)	242±139
	10	120±128	0.691±0.042	74±73	663±170 (9)	768±169 (8)	805±65 (4)	344±218
Mamba	5	69±38	0.764±0.052	54±28	542±93 (10)	807±76 (10)	988±17 (3)	178±90
	6	138±46	0.759±0.039	110±42	456±89 (10)	693±75 (10)	919±36 (7)	286±137
	7	174±95	0.737±0.059	127±83	427±177 (10)	643±102 (10)	858±77 (7)	395±147
	8	209±95	0.751±0.030	137±60	461±151 (10)	617±135 (10)	817±71 (8)	482±214
	9	202±98	0.735±0.032	137±80	389±112 (10)	631±102 (10)	841±92 (8)	518±23
	10	306±57	0.714±0.035	206±34	387±148 (10)	555±66 (10)	761±58 (10)	1110±63

Understanding the Limits of Augmented Memory. Augmented Memory<sup>21</sup> improves sample efficiency by repeated learning from high reward SMILES. With decreasing batch size, performance variance increases, as the approximation to the expected reward (Eq. 2) becomes more noisy. In return, fewer oracle calls are imposed, and the agent learns from an increasingly smaller set of unique SMILES. Our hypothesis is that as long as unique high reward SMILES are still generated, sample efficiency can improve with decreasing batch size, at the expense of diversity. We perform a grid search and vary the batch size (64, 32, 16, 8) and augmentation rounds (0-20 inclusive) using the default RNN architecture (Appendix 5). We make the following key observations: with increasing augmentation rounds and decreasing batch size, sample efficiency improves, diversity decreases, and generating repeated SMILES becomes increasingly prevalent but is tolerable with oracle caching. The optimal augmentation rounds and batch size are 5-10 and 16, respectively, as pushing further introduces too much variance, such that apparent improvements are not statistically significant (at the 95% confidence level). In Appendix B.4, we explored the addition of Beam Enumeration 22 but improvements were not consistently statistically significant. In Appendix B.5, we explored allocating a portion of the oracle budget to a GA, which decreases sample efficiency, but recovers diversity, in agreement with previous works 13,80.

159

160

164

165

166

167

168

169

170

171

172

173

174

175

176

179

180

181

182

183 184

186

187

188

189

190

**Small Molecule Goal-directed Generation: Beyond RNNs.** In this section, we move beyond **RNN** (5.8M) to **Decoder** transformer <sup>27,28</sup> (6.3M) and **Mamba** <sup>29</sup> (5.2M), and empirically show that varying the architecture can improve sample efficiency. Complete grid search results are presented in Appendix B.3. Cross-referencing Table 1, we make the following observations: Increasing augmentation rounds decreases diversity and *inconsistently* improves Yield and OB for RNN and transformer. Mamba *more consistently* benefits from increasing augmentation rounds to generate more high reward molecules and also faster. Across the Yield and OB metrics, Mamba consistently outperforms both the RNN and transformer backbones. In particular, Mamba with 10 augmentation rounds successfully generates 100 molecules above the reward threshold (OB 100 metric) in 10/10 replicates, compared to only 5/10 and 4/10 successful replicates for RNN and transformer, respectively (Table 1). Given Mamba's superior sample efficiency, we focus our analysis on comparing it to the RNN baseline in the remainder of this section (transformer results are provided in Appendix B.3).

Mamba: Enhanced Maximum Likelihood. Table 1 shows that the Mamba architecture notably generates repeated SMILES, which can be rationalized with the maximum likelihood objective. Mamba (5.2M) and RNN (5.8M) have similar parameter counts but during pre-training, the former converges to a lower loss during pre-training (Appendix B.1), indicating a better match to the data distribution. Accordingly, and during RL, Eq. 4 aims to make generating high reward SMILES more likely. Mamba generates repeated SMILES suggesting it overfits the data distribution. We demonstrate this by cross-referencing Fig. 2a, which shows that with high augmentation rounds, the

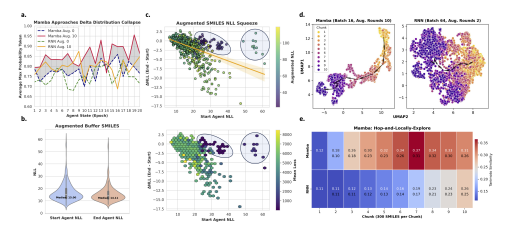


Figure 2: **a.** Average maximum token probability across agent states. Augmentation pushes the agent action distribution towards a delta distribution. **b.** Augmented Memory (10 augmentation rounds) makes the likelihood of generating SMILES in the buffer more likely. **c.** Top: On average, augmented forms of the buffer SMILES become more likely. Bottom: Similar loss magnitudes impose larger changes on improbable sequences and the agent is driven towards generating these specific sequences. When the Augmented Likelihood is equal to the agent likelihood, the loss approaches 0 (circles). **d.** 3,000 oracle budget test experiment chunked into 300 SMILES. UMAP embedding of the agent chemical space traversal (arrows are the centroid of each chunk). Mamba exhibits a directional traversal while RNN (baseline Augmented Memory) continues to sample globally. **e.** Mamba exhibits a "hop-and-locally-explore" behavior where the intra-chunk Tanimoto similarity (top values) are higher than RNN. The bottom value is the inter-chunk similarity.

average max conditional token probability (during generation) approaches 1, and near collapses to a Dirac delta function (less so for RNN). This makes it likely, but *not* deterministic, to generate the same SMILES repeatedly.

194

195

196

197

198

199

200

202

203

204

205

206

207

209

210

211

212

213

214

215

216

217

218

219

220

221

**Squeezing the Likelihood of Augmented SMILES.** While the original Augmented Memory work<sup>21</sup> demonstrated its empirical benefits, we elucidate the underlying mechanism. To isolate its effect, we design a sub-experiment as follows: generate molecules until the buffer is full (100) and then save the agent state before and after executing Augmented Memory (10 augmentation rounds) and save every augmented SMILES form. After execution, the (End) agent becomes more likely to generate the set of augmented SMILES (Fig. 2b). The more *improbable* the SMILES (high NLL), the larger the  $\Delta$ NLL shift (Fig. 2c). According to the loss function (Eq. 4), a larger difference between the Augmented Likelihood (Eq. 3) and Agent Likelihood results in a higher loss. When these terms are near equal, the loss approaches 0 (Fig. 2c circles). The purpose of the Augmented Likelihood is to regularize the agent, preventing it from deviating too far from the prior <sup>23</sup>. Improbable SMILES, which impose a large gradient update, adjust the agent towards a higher probability of generating such sequences. However, already probable (low NLL) SMILES can also impose large loss magnitudes (Fig. 2c), but the  $\Delta$ NLL shift is small because the softmax function saturates, causing minimal changes to the softmax output when the logits are tuned. Taking these observations together, Augmented Memory squeezes the likelihood of augmented SMILES, making the agent more likely to generate any SMILES representation of the same molecular graph. We next demonstrate how the Mamba architecture synergistically leverages this mechanism to enhance sample efficiency.

Mamba: Hop-and-Locally-Explore. Mamba approaches Dirac delta function collapse (Fig. 2a) when learning from repeated augmented SMILES and in the previous section, we have shown that the agent becomes increasingly likely to generate the buffer *molecules*. We hypothesized that Mamba exhibits a "hop-and-locally-explore" behavior: because it is likely to generate *some* SMILES representation of these molecules, small changes to any tokens in these set of augmented sequences equates to small changes to the *same* molecular graph, essentially performing a local exploration (similar molecules, on average, exhibit similar properties, provided the property landscape is not too rough <sup>81,82</sup>). We verify our hypothesis with the following experiment: generate molecules (3,000 oracle budget) and separate the generated set into 10 chunks (each 300 SMILES). We trace the generation trajectory using UMAP<sup>83</sup> and plot the chunk centroids, comparing Mamba and the

baseline (vanilla Augmented Memory<sup>21</sup>) (Fig. 2d). Mamba traverses chemical space in an increased directional manner and the chunks are more locally confined. Further analysis into the intra- and 224 inter-chunk Tanimoto similarity reveals that within chunks, Mamba exhibits much greater similarity 225 than the baseline, and similarity is always lower between chunks (Fig. 2e). Taking these observations 226 together, Mamba (batch size 16) with Augmented Memory (10 augmentation rounds) and oracle 227 caching synergistically improves sample efficiency via "hop-and-locally-explore" behavior (see 228 Appendix C for further quantitative and qualitative analyses). From here on, this model configuration 229 will be referred to as **Saturn** and hyperparameters are *fixed* such that all performance metrics in the 230 following sections are out-of-the-box. 231

#### 4.2 Part 2: Transferability of Sample Efficiency to Physics-based Oracles

In this section, we demonstrate that Saturn's sample efficiency transfers to an MPO objective involving docking against targets related to neurodegeneration (DRD2 <sup>84</sup> and AChE <sup>85</sup>) and inflammation (MK2 kinase <sup>86</sup>). The optimization objective is to constrain MW < 500 Da, maximize the quantitative estimate of drug-likeness (QED) <sup>87</sup>, and minimize AutoDock Vina <sup>88</sup> docking score (see Appendix D.1 for details on the docking protocol). All experiments were run across 10 seeds (0-9 inclusive) and with a 1,000 oracle budget. We compare Saturn (with and without GA) to baseline Augmented Memory <sup>21</sup> using the Yield and OB metrics. Saturn generates more high reward molecules and faster, given the fixed oracle budget (Table 2). This holds even for the more challenging MK2 kinase target where the pre-training data (ChEMBL 33 <sup>79</sup>) is less suited. Furthermore, in agreement with the results from the test experiments, adding a GA on the buffer does not improve sample efficiency but recovers diversity, which can be useful in certain cases.

Table 2: Docking MPO with 1,000 oracle budget. Baseline is vanilla Augmented Memory<sup>21</sup>. IntDiv1<sup>71</sup> is the internal diversity, Scaffolds is the number of unique Bemis-Murcko<sup>78</sup> scaffolds, OB is Oracle Burden (oracle calls required to generate *N* unique molecules). All metrics are computed at the 0.8 reward threshold. The number in parentheses in the OB statistics represents how many runs out of 10 were successful. The mean and standard deviation across 10 seeds (0-9 inclusive) is reported. Best models (statistically significant at the 95% confidence level) are bolded.

Target	Model	Yield (†)	IntDiv1 (†)	Scaffolds (†)	OB 1 (↓)	OB 10 (↓)	OB 100 (↓)
DRD2	Augmented Memory Saturn Saturn-GA	$22 \pm 7$ $369 \pm 62$ $209 \pm 55$	$\begin{array}{c} 0.774 \pm 0.019 \\ 0.671 \pm 0.050 \\ 0.745 \pm 0.041 \end{array}$	$22 \pm 7$ $310 \pm 70$ $189 \pm 57$	$143 \pm 75(10)$ $93 \pm 53(10)$ $96 \pm 56(10)$	$733 \pm 120(10)$ $391 \pm 56(10)$ $403 \pm 75(10)$	Failed $663 \pm 55(10)$ $806 \pm 84(10)$
AChE	Augmented Memory Saturn Saturn-GA	$173 \pm 19$ $480 \pm 79$ $343 \pm 57$	$0.843 \pm 0.009$ $0.757 \pm 0.020$ $0.809 \pm 0.013$	$170 \pm 18$ $400 \pm 96$ $287 \pm 50$	$57 \pm 2(10)$ $32 \pm 24(10)$ $32 \pm 25(10)$	$189 \pm 52(10)$ $185 \pm 82(10)$ $187 \pm 80(10)$	$776 \pm 58(10)$ $508 \pm 80(10)$ $565 \pm 80(10)$
MK2	Augmented Memory Saturn Saturn-GA	$0.2 \pm 0.4$ $14.9 \pm 14.1$ $6.1 \pm 6.5$	$-0.454 \pm 0.212 \\ 0.415 \pm 0.202$	$0.2 \pm 0.4$ $14.1 \pm 13.2$ $5.5 \pm 5.5$	$836 \pm 186(2)$ $677 \pm 186(9)$ $678 \pm 140(9)$	Failed $861 \pm 108(6)$ $911 \pm 11(2)$	Failed Failed Failed

# 4.3 Part 3: Benchmarking Saturn

232

233

236

237

238

239

240

244

245

246

247

248

249

250

252

In this section, we compare Saturn's performance to previous works, including the state-of-the-art Goal-aware fragment Extraction, Assembly, and Modification (GEAM) proposed by Lee et al. <sup>13</sup>, which recently reported impressive results on a docking MPO task, outperforming baselines by a large margin.

**Experimental Details.** To facilitate an exact comparison with GEAM<sup>13</sup>, we used the code from https://anonymous.4open.science/r/GEAM-45EF to reproduce the GEAM results, extract oracle code for our experiments, pre-train on the provided ZINC 250k<sup>89</sup> data (Appendix E,) and used their MPO objective function (Eq. 5),

$$R(x) = \widehat{DS}(x) \times QED(x) \times \widehat{SA}(x) \in [0, 1], \tag{5}$$

where  $\widehat{DS}$  is the normalized QuickVina  $2^{90}$  docking score and  $\widehat{SA}$  is the normalized synthetic accessibility score <sup>91</sup> (see Appendix E for normalization details). Following Lee et al. <sup>13</sup>, docking was performed against 5 targets: **parp1**, **fa7**, **5ht1b**, **braf**, and **jak2**. We ran GEAM and Saturn across 10 seeds (0-9 inclusive) with an oracle budget of 3,000. We emphasize that we do not tune Saturn's hyperparameters for this task and the results in this section are out-of-the-box.

**Metrics.** Following Lee et al.  $^{12,13}$ , we assess the **Hit Ratio** (%) (molecules with a better docking score than the median of known actives, QED > 0.5, SA < 5) and **Novel Hit Ratio** (%) (with the additional constraint of maximum Tanimoto similarity of 0.4 to the training data). We further propose **Strict Hit Ratio** (%) and **Strict Novel Hit Ratio** (%) which filter for the more stringent criteria of QED > 0.7 (based on DrugStore dataset of marketed drugs  $^{87}$ ) and SA < 3 (based on off-the-shelf catalog molecules  $^{91}$ ). While drug candidates need not necessarily meet these stricter thresholds, this metric assesses *optimization capability*, which becomes pertinent when jointly optimizing all components is especially crucial. From an optimization perspective, the objective function (Eq. 5) aims to maximize QED and minimize SA and docking score simultaneously. Therefore, achieving high QED and low SA is part of the goal itself. We additionally measure molecular diversity using **IntDiv1** and #**Circles** with distance threshold 0.75.

Table 3: Novel Hit Ratio (%). Results are from Lee et al. <sup>13</sup> except GEAM and Saturn which we ran across 10 seeds (0-9 inclusive). The mean and standard deviation are reported. Best results (statistically significant at the 95% confidence level) are bolded.

Method			Target Protein		
	parp1	fa7	5ht1b	braf	jak2
REINVENT 23	$0.480 \pm 0.344$	$0.213 \pm 0.081$	$2.453 \pm 0.561$	$0.127 \pm 0.088$	$0.613 \pm 0.167$
GCPN 54	$0.056 \pm 0.016$	$0.444 \pm 0.333$	$0.444 \pm 0.150$	$0.033 \pm 0.027$	$0.256 \pm 0.087$
JT-VAE 45	$0.856 \pm 0.211$	$0.289 \pm 0.016$	$4.656 \pm 1.406$	$0.144 \pm 0.068$	$0.815 \pm 0.044$
GraphAF 93	$0.689 \pm 0.166$	$0.011 \pm 0.016$	$3.178 \pm 0.393$	$0.956 \pm 0.319$	$0.767 \pm 0.098$
GraphGA 63	$4.811 \pm 1.661$	$0.422 \pm 0.193$	$7.011 \pm 2.732$	$3.767 \pm 1.498$	$5.311 \pm 1.667$
MORLD 94	$0.047 \pm 0.050$	$0.007 \pm 0.013$	$0.880 \pm 0.735$	$0.047 \pm 0.040$	$0.227 \pm 0.118$
HierVAE 95	$0.553 \pm 0.214$	$0.007 \pm 0.013$	$0.507 \pm 0.278$	$0.207 \pm 0.220$	$0.227 \pm 0.127$
RationaleRL 55	$4.267 \pm 0.450$	$0.900 \pm 0.098$	$2.967 \pm 0.307$	$0.000 \pm 0.000$	$2.967 \pm 0.196$
GA+D 96	$0.044 \pm 0.042$	$0.011 \pm 0.016$	$1.544 \pm 0.273$	$0.800 \pm 0.864$	$0.756 \pm 0.204$
MARS 97	$1.178 \pm 0.299$	$0.367 \pm 0.072$	$6.833 \pm 0.706$	$0.478 \pm 0.083$	$2.178 \pm 0.545$
GEGL <sup>98</sup>	$0.789 \pm 0.150$	$0.256 \pm 0.083$	$3.167 \pm 0.260$	$0.244 \pm 0.016$	$0.933 \pm 0.072$
GraphDF 99	$0.044 \pm 0.031$	$0.000 \pm 0.000$	$0.000 \pm 0.000$	$0.011 \pm 0.016$	$0.011 \pm 0.016$
FREED 11	$4.627 \pm 0.727$	$1.332 \pm 0.113$	$16.767 \pm 0.897$	$2.940 \pm 0.359$	$5.800 \pm 0.295$
LIMO 100	$0.455 \pm 0.057$	$0.044 \pm 0.016$	$1.189 \pm 0.181$	$0.278 \pm 0.134$	$0.689 \pm 0.319$
GDSS 101	$1.933 \pm 0.208$	$0.368 \pm 0.103$	$4.667 \pm 0.306$	$0.167 \pm 0.134$	$1.167 \pm 0.281$
PS-VAE 102	$1.644 \pm 0.389$	$0.478 \pm 0.140$	$12.622 \pm 1.437$	$0.367 \pm 0.047$	$4.178 \pm 0.933$
MOOD 12	$7.017 \pm 0.428$	$0.733 \pm 0.141$	$18.673 \pm 0.423$	$5.240 \pm 0.285$	$9.200 \pm 0.524$
GEAM 13	$39.159 \pm 2.790$	$19.540 \pm 2.347$	$40.123 \pm 1.611$	$27.467 \pm 1.374$	$\bf 41.765 \pm 3.412$
Saturn (ours) Saturn-Jaccard (ours)	$3.839 \pm 3.316$ $\mathbf{50.552 \pm 9.530}$	$0.470 \pm 0.272$ <b>20.181</b> $\pm$ <b>5.598</b>	$5.731 \pm 6.166$ $54.260 \pm 6.722$	$3.652 \pm 3.777$ $19.820 \pm 10.120$	$6.129 \pm 5.449$ $47.785 \pm 14.041$

**Saturn and GEAM Outperform all Baselines.** We evaluate the Hit Ratio and include random sampling of 3,000 molecules from the ZINC 250k<sup>89</sup> and ChEMBL 33<sup>79</sup> datasets as baselines (Appendix Table 27). The results show that only GEAM<sup>13</sup> and Saturn outperform these baselines, with both methods displaying similar performance. However, Saturn exhibits higher variance, likely due to the small batch size (16) used to approximate the expected reward (Eq. 2). For the Novel Hit Ratio (Table 3), Saturn performs much worse than GEAM, but we rationalize this by cross-referencing Fig. 2. The Mamba backbone excels at maximum likelihood estimation and fits the ZINC 250k<sup>89</sup> training distribution well. It is then unsurprising that generated molecules are not particularly dissimilar to ZINC. We highlight that enforcing molecules to have less than 0.4 Tanimoto similarity to all molecules in the training data is somewhat arbitrary. However, to demonstrate how to solve this problem, we apply curriculum learning <sup>81</sup> to Saturn and further "pre-train" the model to generate molecules with high Jaccard distance (Tanimoto dissimilarity) to the training data (see Appendix E.4). We believe this is still a fair assessment as computing Tanimoto similarity is cheap and this process took minutes and also shows the flexibility of Saturn. We then use this model for the MPO task and show that performance immediately recovers and matches GEAM (Table 3).

**Saturn:** Enhanced MPO. Based on the results so far, it may be desirable to use GEAM over Saturn as it has much lower variance. To investigate this further, we assess the optimization capability of both models by applying a strict filter for QED > 0.7 and SA < 3 (Table 4). The results show that GEAM's Hit Ratios drop drastically while Saturn's remain relatively unchanged, which demonstrates that Saturn optimizes the MPO objective to a much greater degree (see Appendix E for *Novel* Strict Filter results). Importantly, Saturn finds molecules passing this strict filter with much fewer oracle calls (OB metrics in Table 4), trading off diversity to do so. Moreover, for **fa7** and **braf**, GEAM does not find 100 molecules passing the strict filter in 9/10 and 4/10 replicates, respectively, while Saturn is successful in 10/10 for both (Table 4). Finding desirable molecules with fewer oracle calls is of

Table 4: Strict Hit Ratio (%). GEAM and Saturn results are across 10 seeds (0-9 inclusive). OB is Oracle Burden (oracle calls required to generate *N* unique molecules). The number in parentheses in the OB statistics represents how many runs out of 10 were successful. The mean and standard deviation are reported. Best results (statistically significant at the 95% confidence level) are bolded.

Method			Target Protein		
	parp1	fa7	5ht1b	braf	jak2
GEAM 13					
Strict Hit Ratio (†)	$6.510 \pm 1.087$	$2.106 \pm 0.958$	$8.719 \pm 0.903$	$3.685 \pm 0.524$	$7.944 \pm 1.157$
OB (1) (\( \psi \)	$250 \pm 157(10)$	$433 \pm 209(10)$	$114 \pm 112(10)$	$355 \pm 96(10)$	$230 \pm 117(10)$
OB (10) (\( \psi \))	$743 \pm 52(10)$	$1446 \pm 404(10)$	$531 \pm 38(10)$	$892 \pm 144(10)$	$537 \pm 70(10)$
OB (100) (\( \psi\))	$2106 \pm 202(10)$	$2927 \pm 0(1)$	$1527 \pm 110(10)$	$2674 \pm 163(6)$	$1606 \pm 218(10)$
IntDiv1 (†)	$0.766 \pm 0.017$	$0.709 \pm 0.043$	$0.799 \pm 0.017$	$0.751 \pm 0.023$	$0.763 \pm 0.021$
#Circles (↑)	$14 \pm 3$	$7\pm2$	$25 \pm 3$	$11 \pm 2$	$18 \pm 2$
Saturn (ours)					
Strict Hit Ratio	$55.102 \pm 18.027$	$13.887 \pm 9.723$	$64.730 \pm 3.717$	$37.250 \pm 9.615$	$55.903 \pm 13.613$
OB (1) (\( \psi \))	$139 \pm 96(10)$	$352 \pm 206(10)$	$21 \pm 7(10)$	$291 \pm 143(10)$	$88 \pm 56(10)$
OB (10) (\( \psi \))	$518 \pm 92(10)$	$924 \pm 247(10)$	$105 \pm 23(10)$	$581 \pm 123(10)$	$348 \pm 96(10)$
OB (100) (\( \psi \)	$956 \pm 259(10)$	$1776 \pm 551(10)$	$441 \pm 44(10)$	$1057 \pm 187(10)$	$785 \pm 191(10)$
IntDiv1 (†)	$0.596 \pm 0.049$	$0.592 \pm 0.066$	$0.685 \pm 0.021$	$0.597 \pm 0.042$	$0.638 \pm 0.034$
#Circles (↑)	$5 \pm 0$	$3 \pm 1$	$17 \pm 3$	$4 \pm 0$	$7\pm1$

high practical relevance when moving to high-fidelity oracles so as to identify a small set of *excellent* candidates satisfying the MPO objective.

#### 5 Conclusion

293

294

295

296

297

298

299

300

301

304

305

306

307

308

309

311

312

313

314

315

316

317

319

320

321

322

323

326

327

328

In this work, we present **Saturn**, a framework for sample-efficient *de novo* molecular design using memory manipulation. We demonstrate the first application of the Mamba<sup>29</sup> architecture for generative molecular design with reinforcement learning and show how it synergistically leverages SMILES augmentation and experience replay for enhanced sample efficiency. Through systematic study, we elucidate the mechanism of Augmented Memory (original work only showed its empirical benefits) and show it squeezes sequence generation likelihoods such that it becomes increasingly likely to generate some SMILES representation of the replay buffer molecular graphs. Next, we show how Mamba leverages this mechanism to improve sample efficiency through "hop-and-locally-explore" behavior. With the optimal architecture and hyperparameters identified for sample efficiency in a test experiment, we apply Saturn on two MPO tasks relevant to drug discovery, outperforming all baseline models, and matching the recent GEAM 13 model which, when released, outperformed all baselines by a large margin. Compared to GEAM, we further show that Saturn achieves superior MPO, finding desirable molecules faster with fewer oracle calls, albeit with a trade-off in diversity. Our work opens up the prospect of *directly* optimizing expensive high-fidelity oracles (beyond docking), which are more correlated with relevant drug discovery end-points. Recent work has applied multi-fidelity learning <sup>19</sup> or active learning <sup>103,104</sup> to enable on-the-fly update of a surrogate model to predict such oracle evaluations for generative design. These workflows can be applied directly with Saturn, but importantly, we may be *sufficiently efficient* to directly optimize these oracles, mitigating surrogate out-of-domain concerns. Moreover, it is straightforward to augment Saturn with known strategies to improve sample efficiency, such as curriculum learning 81 as we have shown in Part 3. Correspondingly, future work will stress-test Saturn on high-fidelity oracles and interrogate the prospect of directly optimizing QM/MM and free energy 15-18 protocols with modest computational resources.

**Limitations.** While we demonstrate Saturn's broad applicability, it remains to be seen whether performance will carry over to high-fidelity oracles with rougher optimization landscapes <sup>82</sup>, where the "hop-and-locally-explore" behavior may be disadvantageous. However, as we have identified *why* this behavior manifests, we can tailor the sampling behavior for the optimization landscape, if required. For example, activating the genetic algorithm and lowering augmentation rounds loosens the local sampling behavior, as shown in Appendix C.2.

**Broader Impact**. We present a method that enhances sample efficiency in molecular generative models that could impact fields such as drug discovery and functional materials design. There is potential misuse if the generation is steered towards a malicious objective function <sup>105</sup>. As generative design becomes increasingly adopted (in general), measures to ensure safe deployment will be paramount, while maximizing potential societal benefits.

# References

- 1. Frank W Pun, Ivan V Ozerov, and Alex Zhavoronkov. Ai-powered therapeutic target discovery. *Trends in Pharmacological Sciences*, 2023.
- 2. Feng Ren, Xiao Ding, Min Zheng, Mikhail Korzinkin, Xin Cai, Wei Zhu, Alexey Mantsyzov, Alex Aliper, Vladimir Aladinskiy, Zhongying Cao, et al. AlphaFold accelerates artificial intelligence powered drug discovery: efficient discovery of a novel CDK20 small molecule inhibitor. *Chem. Sci.*, 14(6):1443–1452, 2023.
- 33. Wei Zhu, Xiaosong Liu, Qi Li, Feng Gao, Tingting Liu, Xiaojing Chen, Man Zhang, Alex
  Aliper, Feng Ren, Xiao Ding, et al. Discovery of novel and selective SIK2 inhibitors by the
  application of AlphaFold structures and generative models. *Bioorg. Med. Chem.*, 91:117414,
  2023.
- 4. Yangguang Li, Yingtao Liu, Jianping Wu, Xiaosong Liu, Lin Wang, Ju Wang, Jiaojiao Yu, Hongyun Qi, Luoheng Qin, Xiao Ding, et al. Discovery of potent, selective, and orally bioavailable small-molecule inhibitors of CDK8 for the treatment of cancer. *J. Med. Chem.*, 2023.
- Yazhou Wang, Chao Wang, Jinxin Liu, Deheng Sun, Fanye Meng, Man Zhang, Alex Aliper,
   Feng Ren, Alex Zhavoronkov, and Xiao Ding. Discovery of 3-hydroxymethyl-azetidine
   derivatives as potent polymerase theta inhibitors. *Bioorg. Med. Chem.*, page 117662, 2024.
- Feng Ren, Alex Aliper, Jian Chen, Heng Zhao, Sujata Rao, Christoph Kuppe, Ivan V. Ozerov, Man Zhang, Klaus Witte, Chris Kruse, Vladimir Aladinskiy, Yan Ivanenkov, Daniil Polykovskiy, Yanyun Fu, Eugene Babin, Junwen Qiao, Xing Liang, Zhenzhen Mou, Hui Wang, Frank W. Pun, Pedro Torres Ayuso, Alexander Veviorskiy, Dandan Song, Sang Liu, Bei Zhang, Vladimir Naumov, Xiaoqiang Ding, Andrey Kukharenko, Evgeny Izumchenko, and Alex Zhavoronkov. A small-molecule TNIK inhibitor targets fibrosis in preclinical and clinical models. *Nat. Biotechnol.*, March 2024. ISSN 1546-1696. doi: 10.1038/s41587-024-02143-0.
- 7. Jie Zhang, Feng Gao, Wei Zhu, Chenxi Xu, Xiaoyu Ding, Jing Shang, Junwen Qiao, Shan Chen, Xin Cai, Xiao Ding, et al. Ism9682a, a novel and potent kif18a inhibitor, shows robust antitumor effects against chromosomally unstable cancers. *Cancer Research*, 84(6\_Supplement): 5727–5727, 2024.
- 8. Jeff Guo, Jon Paul Janet, Matthias R Bauer, Eva Nittinger, Kathryn A Giblin, Kostas Papadopoulos, Alexey Voronov, Atanas Patronov, Ola Engkvist, and Christian Margreitter. Dockstream: a docking wrapper to enhance de novo molecular design. *Journal of cheminformatics*, 13:1–21, 2021.
- Morgan Thomas, Noel M O'Boyle, Andreas Bender, and Chris De Graaf. Augmented hill-climb
   increases reinforcement learning efficiency for language-based de novo molecule generation.
   Journal of cheminformatics, 14(1):68, 2022.
- Tony Shen, Mohit Pandey, and Martin Ester. Tacogfn: Target conditioned gflownet for drug design. In *NeurIPS 2023 Generative AI and Biology (GenBio) Workshop*, 2023.
- Soojung Yang, Doyeong Hwang, Seul Lee, Seongok Ryu, and Sung Ju Hwang. Hit and lead discovery with explorative rl and fragment-based molecule generation. *Advances in Neural Information Processing Systems*, 34:7924–7936, 2021.
- Seul Lee, Jaehyeong Jo, and Sung Ju Hwang. Exploring chemical space with score-based out-of-distribution generation. In *International Conference on Machine Learning*, pages 18872–18892.
   PMLR, 2023.
- 13. Seul Lee, Seanie Lee, and Sung Ju Hwang. Drug discovery with dynamic goal-aware fragments. arXiv preprint arXiv:2310.00841, 2023.
- 14. Tianfan Fu, Wenhao Gao, Connor Coley, and Jimeng Sun. Reinforced genetic algorithm for structure-based drug design. *Advances in Neural Information Processing Systems*, 35: 12325–12338, 2022.

- 15. Jordan E Crivelli-Decker, Zane Beckwith, Gary Tom, Ly Le, Sheenam Khuttan, Romelia Salomon-Ferrer, Jackson Beall, Rafael Gómez-Bombarelli, and Andrea Bortolato. Machine learning guided agfep: A fast & efficient absolute free energy perturbation solution for virtual screening. 2023.
- 16. Lingle Wang, Jennifer Chambers, and Robert Abel. Protein–ligand binding free energy calculations with fep+. *Biomolecular simulations: methods and protocols*, pages 201–232, 2019.
- J Harry Moore, Matthias R Bauer, Jeff Guo, Atanas Patronov, Ola Engkvist, and Christian
   Margreitter. Icolos: a workflow manager for structure-based post-processing of de novo
   generated small molecules. *Bioinformatics*, 38(21):4951–4952, 2022.
- 18. J Harry Moore, Christian Margreitter, Jon Paul Janet, Ola Engkvist, Bert L de Groot, and Vytautas Gapsys. Automated relative binding free energy calculations from smiles to  $\delta\delta g$ .

  Communications Chemistry, 6(1):82, 2023.
- 19. Peter Eckmann, Dongxia Wu, Germano Heinzelmann, Michael K Gilson, and Rose Yu. Mfbind:
   a multi-fidelity approach for evaluating drug compounds in practical generative modeling. arXiv preprint arXiv:2402.10387, 2024.
- 20. Wenhao Gao, Tianfan Fu, Jimeng Sun, and Connor Coley. Sample efficiency matters: a benchmark for practical molecular optimization. *Advances in neural information processing systems*, 35:21342–21357, 2022.
- 21. Jeff Guo and Philippe Schwaller. Augmented memory: Sample-efficient generative molecular design with reinforcement learning. *JACS Au*, 2024.
- 22. Jeff Guo and Philippe Schwaller. Beam enumeration: Probabilistic explainability for sample efficient self-conditioned molecular design. In *Proc. 12th International Conference on Learning Representations*, 2024.
- 402 23. Marcus Olivecrona, Thomas Blaschke, Ola Engkvist, and Hongming Chen. Molecular de-novo design through deep reinforcement learning. *Journal of cheminformatics*, 9:1–14, 2017.
- Thomas Blaschke, Josep Arús-Pous, Hongming Chen, Christian Margreitter, Christian Tyrchan,
   Ola Engkvist, Kostas Papadopoulos, and Atanas Patronov. Reinvent 2.0: an ai tool for de novo drug design. *Journal of chemical information and modeling*, 60(12):5918–5922, 2020.
- 25. Esben Jannik Bjerrum. Smiles enumeration as data augmentation for neural network modeling of molecules. *arXiv preprint arXiv:1703.07076*, 2017.
- 26. Sepp Hochreiter and Jürgen Schmidhuber. Long short-term memory. *Neural computation*, 9(8): 1735–1780, 1997.
- Ashish Vaswani, Noam Shazeer, Niki Parmar, Jakob Uszkoreit, Llion Jones, Aidan N Gomez,
   Łukasz Kaiser, and Illia Polosukhin. Attention is all you need. Advances in neural information
   processing systems, 30, 2017.
- 28. Alec Radford, Jeffrey Wu, Rewon Child, David Luan, Dario Amodei, Ilya Sutskever, et al. Language models are unsupervised multitask learners. *OpenAI blog*, 1(8):9, 2019.
- 416 29. Albert Gu and Tri Dao. Mamba: Linear-time sequence modeling with selective state spaces.
   417 arXiv preprint arXiv:2312.00752, 2023.
- 30. David Weininger. Smiles, a chemical language and information system. 1. introduction to methodology and encoding rules. *Journal of chemical information and computer sciences*, 28 (1):31–36, 1988.
- 31. Hannes H Loeffler, Jiazhen He, Alessandro Tibo, Jon Paul Janet, Alexey Voronov, Lewis H Mervin, and Ola Engkvist. Reinvent 4: Modern ai–driven generative molecule design. *Journal of Cheminformatics*, 16(1):20, 2024.
- 424 32. Marwin HS Segler, Thierry Kogej, Christian Tyrchan, and Mark P Waller. Generating focused 425 molecule libraries for drug discovery with recurrent neural networks. *ACS central science*, 4(1): 426 120–131, 2018.

- 33. Daniel Neil, Marwin Segler, Laura Guasch, Mohamed Ahmed, Dean Plumbley, Matthew
   Sellwood, and Nathan Brown. Exploring deep recurrent models with reinforcement learning
   for molecule design. In *Proc. 6th International Conference on Learning Representations*, 2018.
- 430 34. Mariya Popova, Olexandr Isayev, and Alexander Tropsha. Deep reinforcement learning for de novo drug design. *Science advances*, 4(7):eaap7885, 2018.
- 432 35. Esben Jannik Bjerrum, Christian Margreitter, Thomas Blaschke, Simona Kolarova, and Raquel
   433 López-Ríos de Castro. Faster and more diverse de novo molecular optimization with double 434 loop reinforcement learning using augmented smiles. *Journal of Computer-Aided Molecular Design*, 37(8):373–394, 2023.
- 36. Viraj Bagal, Rishal Aggarwal, PK Vinod, and U Deva Priyakumar. Molgpt: molecular generation using a transformer-decoder model. *Journal of Chemical Information and Modeling*, 62(9): 2064–2076, 2021.
- 439 37. Ye Wang, Honggang Zhao, Simone Sciabola, and Wenlu Wang. cmolgpt: A conditional
   440 generative pre-trained transformer for target-specific de novo molecular generation. *Molecules*,
   441 28(11):4430, 2023.
- 38. Tao Feng, Pengcheng Xu, Tianfan Fu, Siddhartha Laghuvarapu, and Jimeng Sun. Molecular de novo design through transformer-based reinforcement learning. *arXiv preprint arXiv:2310.05365*, 2023.
- 445 39. Eyal Mazuz, Guy Shtar, Bracha Shapira, and Lior Rokach. Molecule generation using trans-446 formers and policy gradient reinforcement learning. *Scientific Reports*, 13(1):8799, 2023.
- 447 40. Xiuyuan Hu, Guoqing Liu, Yang Zhao, and Hao Zhang. De novo drug design using reinforce-448 ment learning with multiple gpt agents. *Advances in Neural Information Processing Systems*, 449 36, 2024.
- 41. Jiazhen He, Alessandro Tibo, Jon Paul Janet, Eva Nittinger, Christian Tyrchan, Werngard Czechtizky, and Engkvist Ola. Evaluation of reinforcement learning in transformer-based molecular design. 2024.
- 42. Yuyao Yang, Shuangjia Zheng, Shimin Su, Chao Zhao, Jun Xu, and Hongming Chen. Syntalinker: automatic fragment linking with deep conditional transformer neural networks. *Chemical science*, 11(31):8312–8322, 2020.
- 43. Diederik P Kingma and Max Welling. Auto-encoding variational bayes. *arXiv preprint* 457 *arXiv:1312.6114*, 2013.
- 44. Rafael Gómez-Bombarelli, Jennifer N Wei, David Duvenaud, José Miguel Hernández-Lobato,
   Benjamín Sánchez-Lengeling, Dennis Sheberla, Jorge Aguilera-Iparraguirre, Timothy D Hirzel,
   Ryan P Adams, and Alán Aspuru-Guzik. Automatic chemical design using a data-driven
   continuous representation of molecules. ACS central science, 4(2):268–276, 2018.
- 462 45. Wengong Jin, Regina Barzilay, and Tommi Jaakkola. Junction tree variational autoencoder for
   463 molecular graph generation. In *International conference on machine learning*, pages 2323–2332.
   464 PMLR, 2018.
- 46. Alex Zhavoronkov, Yan A Ivanenkov, Alex Aliper, Mark S Veselov, Vladimir A Aladinskiy,
   466 Anastasiya V Aladinskaya, Victor A Terentiev, Daniil A Polykovskiy, Maksim D Kuznetsov,
   467 Arip Asadulaev, et al. Deep learning enables rapid identification of potent ddr1 kinase inhibitors.
   468 Nature biotechnology, 37(9):1038–1040, 2019.
- 47. Ian Goodfellow, Jean Pouget-Abadie, Mehdi Mirza, Bing Xu, David Warde-Farley, Sherjil
   Ozair, Aaron Courville, and Yoshua Bengio. Generative adversarial nets. *Advances in neural* information processing systems, 27, 2014.
- 48. Artur Kadurin, Alexander Aliper, Andrey Kazennov, Polina Mamoshina, Quentin Vanhaelen, Kuzma Khrabrov, and Alex Zhavoronkov. The cornucopia of meaningful leads: Applying deep adversarial autoencoders for new molecule development in oncology. *Oncotarget*, 8(7):10883, 2017.

- 49. Gabriel Lima Guimaraes, Benjamin Sanchez-Lengeling, Carlos Outeiral, Pedro Luis Cunha
   Farias, and Alán Aspuru-Guzik. Objective-reinforced generative adversarial networks (organ)
   for sequence generation models. arXiv preprint arXiv:1705.10843, 2017.
- 50. Benjamin Sanchez-Lengeling, Carlos Outeiral, Gabriel L Guimaraes, and Alan Aspuru-Guzik.
  Optimizing distributions over molecular space. an objective-reinforced generative adversarial network for inverse-design chemistry (organic). 2017.
- 51. Evgeny Putin, Arip Asadulaev, Yan Ivanenkov, Vladimir Aladinskiy, Benjamin Sanchez Lengeling, Alán Aspuru-Guzik, and Alex Zhavoronkov. Reinforced adversarial neural computer
   for de novo molecular design. *Journal of chemical information and modeling*, 58(6):1194–1204,
   2018.
- 52. Nicola De Cao and Thomas Kipf. Molgan: An implicit generative model for small molecular graphs. *arXiv preprint arXiv:1805.11973*, 2018.
- 488 53. Yan A Ivanenkov, Daniil Polykovskiy, Dmitry Bezrukov, Bogdan Zagribelnyy, Vladimir Al489 adinskiy, Petrina Kamya, Alex Aliper, Feng Ren, and Alex Zhavoronkov. Chemistry42: an
  490 ai-driven platform for molecular design and optimization. *Journal of Chemical Information*491 *and Modeling*, 63(3):695–701, 2023.
- Jiaxuan You, Bowen Liu, Zhitao Ying, Vijay Pande, and Jure Leskovec. Graph convolutional
   policy network for goal-directed molecular graph generation. In *Advances in neural information* processing systems. NeurIPS, 2018.
- Wengong Jin, Regina Barzilay, and Tommi Jaakkola. Multi-objective molecule generation using interpretable substructures. In *International conference on machine learning*, pages 4849–4859.
   PMLR, 2020.
- 56. Rocío Mercado, Tobias Rastemo, Edvard Lindelöf, Günter Klambauer, Ola Engkvist, Hongming Chen, and Esben Jannik Bjerrum. Graph networks for molecular design. *Machine Learning:* Science and Technology, 2(2):025023, 2021.
- 57. Sara Romeo Atance, Juan Viguera Diez, Ola Engkvist, Simon Olsson, and Rocío Mercado.
   De novo drug design using reinforcement learning with graph-based deep generative models.
   Journal of Chemical Information and Modeling, 62(20):4863–4872, 2022.
- 58. Krzysztof Maziarz, Henry Jackson-Flux, Pashmina Cameron, Finton Sirockin, Nadine Schneider, Nikolaus Stiefl, Marwin Segler, and Marc Brockschmidt. Learning to extend molecular scaffolds with structural motifs. In *Proc. 10th International Conference on Learning Representations*, 2022.
- 508
   59. Clement Vignac, Igor Krawczuk, Antoine Siraudin, Bohan Wang, Volkan Cevher, and Pas 509 cal Frossard. DiGress: Discrete denoising diffusion for graph generation. In *Proc. 11th* 510 International Conference on Learning Representations, 2023.
- 511 60. Yoshua Bengio, Salem Lahlou, Tristan Deleu, Edward J Hu, Mo Tiwari, and Emmanuel Bengio. Gflownet foundations. *Journal of Machine Learning Research*, 24(210):1–55, 2023.
- 61. Emmanuel Bengio, Moksh Jain, Maksym Korablyov, Doina Precup, and Yoshua Bengio. Flow
   network based generative models for non-iterative diverse candidate generation. Advances in
   Neural Information Processing Systems, 34:27381–27394, 2021.
- 62. Melanie Mitchell. An introduction to genetic algorithms. MIT press, 1998.
- 63. Jan H Jensen. A graph-based genetic algorithm and generative model/monte carlo tree search for the exploration of chemical space. *Chemical science*, 10(12):3567–3572, 2019.
- 64. Ilia Igashov, Hannes Stärk, Clément Vignac, Arne Schneuing, Victor Garcia Satorras, Pascal
   Frossard, Max Welling, Michael Bronstein, and Bruno Correia. Equivariant 3d-conditional
   diffusion model for molecular linker design. *Nature Machine Intelligence*, pages 1–11, 2024.

- Arne Schneuing, Yuanqi Du, Charles Harris, Kieran Didi, Arian Jamasb, Ilia Igashov, Weitao Du, Carla Gomes, Max Welling, Tom Blundell, et al. Flexible structure-based design of small molecules with equivariant diffusion models. In *PROTEIN SCIENCE*, volume 32. WILEY 111 RIVER ST, HOBOKEN 07030-5774, NJ USA, 2023.
- 66. Mario Krenn, Florian Häse, AkshatKumar Nigam, Pascal Friederich, and Alan Aspuru-Guzik.
   Self-referencing embedded strings (selfies): A 100% robust molecular string representation.
   Machine Learning: Science and Technology, 1(4):045024, 2020.
- 67. Mario Krenn, Qianxiang Ai, Senja Barthel, Nessa Carson, Angelo Frei, Nathan C Frey, Pascal
   Friederich, Théophile Gaudin, Alberto Alexander Gayle, Kevin Maik Jablonka, et al. Selfies
   and the future of molecular string representations. *Patterns*, 3(10), 2022.
- 68. Michael A Skinnider. Invalid smiles are beneficial rather than detrimental to chemical language models. *Nature Machine Intelligence*, pages 1–12, 2024.
- 69. Rıza Özçelik, Sarah de Ruiter, Emanuele Criscuolo, and Francesca Grisoni. Chemical language modeling with structured state spaces. 2024.
- 70. Nathan Brown, Marco Fiscato, Marwin HS Segler, and Alain C Vaucher. Guacamol: benchmarking models for de novo molecular design. *Journal of chemical information and modeling*, 59(3):1096–1108, 2019.
- 71. Daniil Polykovskiy, Alexander Zhebrak, Benjamin Sanchez-Lengeling, Sergey Golovanov,
   Oktai Tatanov, Stanislav Belyaev, Rauf Kurbanov, Aleksey Artamonov, Vladimir Aladinskiy,
   Mark Veselov, et al. Molecular sets (moses): a benchmarking platform for molecular generation
   models. Frontiers in pharmacology, 11:565644, 2020.
- 72. Josep Arús-Pous, Simon Viet Johansson, Oleksii Prykhodko, Esben Jannik Bjerrum, Christian
   Tyrchan, Jean-Louis Reymond, Hongming Chen, and Ola Engkvist. Randomized smiles strings
   improve the quality of molecular generative models. *Journal of cheminformatics*, 11:1–13,
   2019.
- 73. Michael Moret, Lukas Friedrich, Francesca Grisoni, Daniel Merk, and Gisbert Schneider.
  Generative molecular design in low data regimes. *Nature Machine Intelligence*, 2(3):171–180,
  2020.
- Michael A Skinnider, R Greg Stacey, David S Wishart, and Leonard J Foster. Chemical language
   models enable navigation in sparsely populated chemical space. *Nature Machine Intelligence*, 3
   (9):759–770, 2021.
- 75. Marco Ballarotto, Sabine Willems, Tanja Stiller, Felix Nawa, Julian A Marschner, Francesca
   Grisoni, and Daniel Merk. De novo design of nurr1 agonists via fragment-augmented generative
   deep learning in low-data regime. *Journal of Medicinal Chemistry*, 66(12):8170–8177, 2023.
- Vendy Fialková, Jiaxi Zhao, Kostas Papadopoulos, Ola Engkvist, Esben Jannik Bjerrum, Thierry
   Kogej, and Atanas Patronov. Libinvent: reaction-based generative scaffold decoration for in
   silico library design. *Journal of Chemical Information and Modeling*, 62(9):2046–2063, 2021.
- 77. Thomas Blaschke, Ola Engkvist, Jürgen Bajorath, and Hongming Chen. Memory-assisted
   reinforcement learning for diverse molecular de novo design. *Journal of cheminformatics*, 12
   (1):68, 2020.
- 78. Guy W Bemis and Mark A Murcko. The properties of known drugs. 1. molecular frameworks.
   Journal of medicinal chemistry, 39(15):2887–2893, 1996.
- Anna Gaulton, Louisa J Bellis, A Patricia Bento, Jon Chambers, Mark Davies, Anne Hersey,
   Yvonne Light, Shaun McGlinchey, David Michalovich, Bissan Al-Lazikani, et al. Chembl: a
   large-scale bioactivity database for drug discovery. *Nucleic acids research*, 40(D1):D1100–D1107, 2012.
- 568 80. Xuhan Liu, Kai Ye, Herman WT van Vlijmen, Michael TM Emmerich, Adriaan P IJzerman, 569 and Gerard JP van Westen. Drugex v2: de novo design of drug molecules by pareto-based 570 multi-objective reinforcement learning in polypharmacology. *Journal of cheminformatics*, 13 571 (1):85, 2021.

- 572 81. Jeff Guo, Vendy Fialková, Juan Diego Arango, Christian Margreitter, Jon Paul Janet, Kostas 573 Papadopoulos, Ola Engkvist, and Atanas Patronov. Improving de novo molecular design with 574 curriculum learning. *Nature Machine Intelligence*, 4(6):555–563, 2022.
- 82. Matteo Aldeghi, David E Graff, Nathan Frey, Joseph A Morrone, Edward O Pyzer-Knapp, Kirk E Jordan, and Connor W Coley. Roughness of molecular property landscapes and its impact on modellability. *Journal of Chemical Information and Modeling*, 62(19):4660–4671, 2022.
- 83. Leland McInnes, John Healy, and James Melville. Umap: Uniform manifold approximation and projection for dimension reduction. *arXiv preprint arXiv:1802.03426*, 2018.
- 84. Sheng Wang, Tao Che, Anat Levit, Brian K Shoichet, Daniel Wacker, and Bryan L Roth.
   Structure of the d2 dopamine receptor bound to the atypical antipsychotic drug risperidone.
   Nature, 555(7695):269–273, 2018.
- 85. Gitay Kryger, Israel Silman, and Joel L Sussman. Structure of acetylcholinesterase complexed
   with e2020 (aricept®): implications for the design of new anti-alzheimer drugs. *Structure*, 7(3):
   297–307, 1999.
- 86. Maria A Argiriadi, Anna M Ericsson, Christopher M Harris, David L Banach, David W Borhani, David J Calderwood, Megan D Demers, Jennifer DiMauro, Richard W Dixon, Jennifer Hardman, et al. 2, 4-diaminopyrimidine mk2 inhibitors. part i: observation of an unexpected inhibitor binding mode. *Bioorganic & medicinal chemistry letters*, 20(1):330–333, 2010.
- 87. G Richard Bickerton, Gaia V Paolini, Jérémy Besnard, Sorel Muresan, and Andrew L Hopkins.
  Quantifying the chemical beauty of drugs. *Nature chemistry*, 4(2):90–98, 2012.
- 593 88. Oleg Trott and Arthur J Olson. Autodock vina: improving the speed and accuracy of docking 594 with a new scoring function, efficient optimization, and multithreading. *Journal of computa-*595 *tional chemistry*, 31(2):455–461, 2010.
- 89. Teague Sterling and John J Irwin. Zinc 15–ligand discovery for everyone. *Journal of chemical information and modeling*, 55(11):2324–2337, 2015.
- 598 90. Amr Alhossary, Stephanus Daniel Handoko, Yuguang Mu, and Chee-Keong Kwoh. Fast,
   599 accurate, and reliable molecular docking with quickvina 2. *Bioinformatics*, 31(13):2214–2216,
   600 2015.
- 91. Peter Ertl and Ansgar Schuffenhauer. Estimation of synthetic accessibility score of drug-like
   molecules based on molecular complexity and fragment contributions. *Journal of cheminfor-matics*, 1:1–11, 2009.
- Yutong Xie, Ziqiao Xu, Jiaqi Ma, and Qiaozhu Mei. How much space has been explored?
   measuring the chemical space covered by databases and machine-generated molecules. In *Proc.* 11th International Conference on Learning Representations, 2023.
- 93. Chence Shi, Minkai Xu, Zhaocheng Zhu, Weinan Zhang, Ming Zhang, and Jian Tang. Graphaf:
   a flow-based autoregressive model for molecular graph generation. In *Proc. 8th International Conference on Learning Representations*, 2020.
- 94. Woosung Jeon and Dongsup Kim. Autonomous molecule generation using reinforcement learning and docking to develop potential novel inhibitors. *Scientific reports*, 10(1):22104, 2020.
- 95. Wengong Jin, Regina Barzilay, and Tommi Jaakkola. Hierarchical generation of molecular
   graphs using structural motifs. In *International conference on machine learning*, pages 4839–4848. PMLR, 2020.
- 96. AkshatKumar Nigam, Pascal Friederich, Mario Krenn, and Alán Aspuru-Guzik. Augmenting
   genetic algorithms with deep neural networks for exploring the chemical space. In *Proc. 8th* International Conference on Learning Representations, 2020.

- 97. Yutong Xie, Chence Shi, Hao Zhou, Yuwei Yang, Weinan Zhang, Yong Yu, and Lei Li. Mars:
   Markov molecular sampling for multi-objective drug discovery. In *Proc. 9th International Conference on Learning Representations*, 2021.
- 98. Sungsoo Ahn, Junsu Kim, Hankook Lee, and Jinwoo Shin. Guiding deep molecular optimization with genetic exploration. volume 33, pages 12008–12021, 2020.
- 99. Youzhi Luo, Keqiang Yan, and Shuiwang Ji. Graphdf: A discrete flow model for molecular
   graph generation. In *International conference on machine learning*, pages 7192–7203. PMLR,
   2021.
- 100. Peter Eckmann, Kunyang Sun, Bo Zhao, Mudong Feng, Michael K Gilson, and Rose Yu. Limo: Latent inceptionism for targeted molecule generation. In *International conference on machine* learning. PMLR, 2022.
- 101. Jaehyeong Jo, Seul Lee, and Sung Ju Hwang. Score-based generative modeling of graphs
   via the system of stochastic differential equations. In *International Conference on Machine Learning*, pages 10362–10383. PMLR, 2022.
- Kiangzhe Kong, Wenbing Huang, Zhixing Tan, and Yang Liu. Molecule generation by principal subgraph mining and assembling. *Advances in Neural Information Processing Systems*, 35: 2550–2563, 2022.
- Hannes Loeffler, Shunzhou Wan, Marco Klähn, Agastya Bhati, and Peter Coveney. Optimal
   molecular design: Generative active learning combining reinvent with absolute binding free
   energy simulations. 2024.
- Michael Dodds, Jeff Guo, Thomas Löhr, Alessandro Tibo, Ola Engkvist, and Jon Paul Janet.
   Sample efficient reinforcement learning with active learning for molecular design. *Chemical Science*, 15(11):4146–4160, 2024.
- 642 105. Fabio Urbina, Filippa Lentzos, Cédric Invernizzi, and Sean Ekins. Dual use of artificial-643 intelligence-powered drug discovery. *Nature Machine Intelligence*, 4(3):189–191, 2022.
- 106. Jeff Guo, Franziska Knuth, Christian Margreitter, Jon Paul Janet, Kostas Papadopoulos, Ola
   Engkvist, and Atanas Patronov. Link-invent: generative linker design with reinforcement
   learning. Digital Discovery, 2(2):392–408, 2023.
- Jiazhen He, Huifang You, Emil Sandström, Eva Nittinger, Esben Jannik Bjerrum, Christian Tyrchan, Werngard Czechtizky, and Ola Engkvist. Molecular optimization by capturing chemist's intuition using deep neural networks. *Journal of cheminformatics*, 13:1–17, 2021.
- Alessandro Tibo, Jiazhen He, Jon Paul Janet, Eva Nittinger, and Ola Engkvist. Exhaustive local
   chemical space exploration using a transformer model. 2023.
- Daniel Flam-Shepherd, Kevin Zhu, and Alán Aspuru-Guzik. Language models can learn complex molecular distributions. *Nature Communications*, 13(1):3293, 2022.
- 654 110. Austin Tripp and José Miguel Hernández-Lobato. Genetic algorithms are strong baselines for molecule generation. *arXiv preprint arXiv:2310.09267*, 2023.
- Matthew Schlegel, Wesley Chung, Daniel Graves, Jian Qian, and Martha White. Importance
   resampling for off-policy prediction. *Advances in Neural Information Processing Systems*, 32,
   2019.
- 659 112. Ronald J Williams. Simple statistical gradient-following algorithms for connectionist reinforce-660 ment learning. *Machine learning*, 8:229–256, 1992.
- 661 113. G Madhavi Sastry, Matvey Adzhigirey, Tyler Day, Ramakrishna Annabhimoju, and Woody Sherman. Protein and ligand preparation: parameters, protocols, and influence on virtual screening enrichments. *Journal of computer-aided molecular design*, 27:221–234, 2013.
- 664 114. Schrödinger release 2019-4: Protein preparation wizard; epik, schrödinger, llc, new york, ny, 2019; impact, schrödinger, llc, new york, ny; prime, schrödinger, llc, new york, ny, 2019.

- Katarina Roos, Chuanjie Wu, Wolfgang Damm, Mark Reboul, James M Stevenson, Chao Lu,
   Markus K Dahlgren, Sayan Mondal, Wei Chen, Lingle Wang, et al. Opls3e: Extending force
   field coverage for drug-like small molecules. *Journal of chemical theory and computation*, 15
   (3):1863–1874, 2019.
- 670 116. Anthony K Rappé, Carla J Casewit, KS Colwell, William A Goddard III, and W Mason Skiff. Uff, a full periodic table force field for molecular mechanics and molecular dynamics simulations. *Journal of the American chemical society*, 114(25):10024–10035, 1992.
- Jason Wei, Yi Tay, Rishi Bommasani, Colin Raffel, Barret Zoph, Sebastian Borgeaud, Dani
   Yogatama, Maarten Bosma, Denny Zhou, Donald Metzler, et al. Emergent abilities of large
   language models. arXiv preprint arXiv:2206.07682, 2022.

# 676 Appendix

The Appendix contains full details on Saturn, grid-search results, algorithmic details, and supplementary results. The code is available at https://figshare.com/s/6040d65bfbfc29d6fedf.

# 679 A What is Saturn?

Saturn is a language-based generative molecular design framework which features minimal implementations of Augmented Memory <sup>21</sup> and Beam Enumeration <sup>22</sup>. These two methods were first implemented here: https://github.com/schwallergroup/augmented\_memory, which in turn was built on REINVENT version 3.2 <sup>23,24</sup>: https://github.com/MolecularAI/Reinvent. REINVENT is still under active development and version 4 <sup>31</sup> was recently released, supporting a wide range of generative tasks including small molecule design <sup>23,24</sup>, library design <sup>76</sup>, linker design <sup>106</sup>, proposing small modifications <sup>107</sup>, and sampling nearest neighbors <sup>108</sup>.

Saturn (at the moment) focuses only on generative small molecule design and research development 687 is on sample efficiency. It is a much smaller code-base than REINVENT 4 and with focus on 688 minimal implementation. That being said, the key new additions to Saturn include: extending small 689 molecule generative architecture from just RNN in REINVENT to decoder transformer 27,28 and 690 Mamba<sup>29</sup>. Secondly, allowing oracle caching to track repeated generations and allow pre-screening 691 specified oracles (in an MPO objective, some oracle components may be computationally inexpensive 692 and it would be practical to first screen a molecules through these oracles before any expensive 693 components). Thirdly, implementation of a genetic algorithm which couples GraphGA<sup>63</sup> on the 694 replay buffer such that new molecules can be generated from the replay buffer parent sequences. In 695 the ensuing subsections, we describe in detail these key new additions. 696

#### 697 A.1 Generative Architecture

Many initial language-based molecular generative models were RNN-based <sup>23,32,34</sup>. Early benchmarks 698 (GuacaMol<sup>70</sup> and MOSES<sup>71</sup>) assessed whether generated molecules were valid (RDKit parsable), 699 unique, and novel (not in the training data). RNNs satisfy these metrics and can learn distributions 700 well <sup>109</sup>. More recently, with the prevalence of the transformer <sup>27,28</sup> architecture, many works <sup>9,36–42</sup> 701 have suggested a replacement of RNNs for generative design. However, many performance assess-702 ments only focus on validity, uniqueness, novelty, and optimizing for permissive oracles such as logP, QED<sup>87</sup> ("drug-likeness"), and the SA score<sup>91</sup>. Some works show that transformers can learn 703 704 longer SMILES sequences better than RNNs 38 (such as natural products). However, often, one actually wants to limit sequence length to constrain design to small molecules. Furthermore, recent 706 works have coupled transformers with reinforcement learning (RL)<sup>9,38-41</sup> but the performance is not 707 necessarily better than RNNs. Consequently, it is unclear whether the benefits of transformers are 708 strictly advantageous for small molecule generation. 709

In this work, we extend Augmented Memory<sup>21</sup> to decoder transformer<sup>27,28</sup> and Mamba<sup>29</sup>. Our results show that transformers display similar performance to RNNs for small molecule generation, in agreement with previous literature findings<sup>9</sup>. We further demonstrate the first application of Mamba<sup>29</sup> for goal-directed generation, supplementing recent work investigating S4 models for transfer learning<sup>69</sup>.

# 715 A.2 Oracle Caching

In many reinforcement learning (RL) set-ups, the reward is assumed to be *stationary*, i.e., it does not change on repeat evaluation. This is an assumption that is not always true for physics-based oracles relevant in drug discovery. For example, docking depends on the initial conformer generated, and even more so for molecular dynamics simulations. However, it is reasonable to assume that the reward is *near deterministic* given a reasonably well behaved protein system (in which preliminary studies were made to verify the oracle stability). In effect, the reward for repeat molecules can be retrieved from a cache, thus not imposing additional oracle evaluations. In this work, we show that under this assumption, Saturn can leverage the Mamba <sup>29</sup> architecture for enhanced sample efficiency. In particular, Mamba displays low uniqueness, but we show this is not detrimental.

As any given molecule can have numerous SMILES representations (via augmentation <sup>25</sup>), it is important to store the *canonical* SMILES in the cache, and also to canonicalize sampled batches when querying the cache. Canonicalization is simply a pre-defined traversal and can differ depending on the method used. As long as all canonicalization operations are performed with the same method, consistency can be guaranteed. In this work, we use RDKit.

#### A.3 Genetic Algorithm

Genetic algorithms (GAs) by themselves can be sample-efficient molecular optimizers <sup>20,63,110</sup>. Previ-731 ous work has shown that GAs can improve diversity of the generated set 80. Recently, Lee et al. 13 732 proposed Goal-aware fragment Extraction, Assembly, and Modification (GEAM) which combines RL with GraphGA<sup>63</sup> and achieves impressive results on generating diverse hits. In Saturn, we implement GraphGA on the replay buffer itself, treating the highest rewarding molecules generated in the entire run so far, as the parent population. Following GEAM <sup>13</sup>, sampling the parents is done with probability proportion to their corresponding rewards. New molecules from crossover and mutation operations are deposited into the Buffer if they are also high rewarding, essentially refreshing the 738 buffer, such that Augmented Memory<sup>21</sup> can learn from these new SMILES. The motivation was 739 to leverage the GA to counteract decreases in diversity and potentially improve sample efficiency. 740 In the results in the main text and in the following sections, we show that applying the GA does 741 not lead to improved sample efficiency but does indeed recover diversity. We believe that this can be a useful modification to the optimization algorithm in cases where relatively expensive oracles 743 are used and diversity is important due to prevalence of false positives. Concretely, higher-fidelity 744 oracles should in principle model physical behavior more accurately, such that true positives are 745 more common. This can be shown in previous works where using free energy simulations provide 746 better correlations with binding affinity 15,19. In such a case, sample efficiency becomes increasingly 747 important, as the goal is to simply generate molecules satisfying this simulation and lower diversity 748 is not detrimental. However, when using lower-fidelity oracles, more false positives means it is 749 beneficial to have more diverse ideas for downstream triaging. Finally, we note that applying the GA and generating new molecules strictly means they were generated off-policy (in the RL context). Therefore, more meaningful updates to the agent may be achieved with importance sampling  $^{111}$ , which we did not explore in the current work. 753

# A.4 Full Algorithm Details and Pseudo-code

754

756

757

758

760

763

764

765

766

767

768

In this section, we derive Saturn's loss function with particular focus on showing its equivalency to maximizing the expected reward. The derivation follows previous works <sup>21,23,76</sup> but with added discussion around implications of the loss function. Specifically, Saturn adapts the Augmented Memory <sup>21</sup> algorithm which is in turn based on REINVENT <sup>23,24,31</sup>. The algorithm itself is reinforcement learning based and can be seen as a modified REINFORCE <sup>112</sup> algorithm. However, while **Saturn** (using Mamba with batch size 16 and 10 augmentation rounds) adapts Augmented Memory, the optimization trajectory is quite different from the original Augmented Memory work due to the "hop-and-locally-explore" sampling behavior. We will focus on highlighting specific points related to this

Saturn's Loss Function. We begin by presenting how Saturn generates SMILES  $^{30}$ , which is the data representation used. SMILES are sequences of alphanumeric characters that can be parsed and mapped to a molecular graph, i.e., a molecule. As SMILES are text-based, it is straightforward to tokenize them, and pre-training Saturn follows next-token prediction. Saturn generates SMILES in an autoregressive manner and thus, SMILES are generated token-by-token from time-step, t to T. This can be viewed from a reinforcement learning perspective by defining  $S_t$  as the state space representing all intermediate token sequences during molecular generation.  $A_t(s_t)$  is the action space which involves sampling a token from a conditional probability distribution, given a token sequence so far, i.e., the current state. Mathematically, the probability of sampling a SMILES, x is given by:

$$P(x) = \prod_{t=1}^{T} \pi_{\theta_{\text{Agent}}}(a_t \mid s_t)$$
 (6)

Just generating SMILES is often not useful because they should satisfy the target objective. Thus, the base pre-trained model needs to be tuned somehow to achieve this. The end goal is to find a **Policy** 

(in the reinforcement learning perspective) which dictates with what probability SMILES should be generated to optimize an objective function. To this end, we define the **Prior** and the **Agent** which 776 share the same architecture (Mamba) and whose weights are exactly the same at the beginning of a 777 generative experiment. The Prior and Agent are general terms to describe the model states but they 778 both are policies as they both induce a probability of sampling SMILES. However, what is different 779 is that the Prior's weights are frozen so it is never updated. By contrast, the Agent is updated and is 780 the model that is learning how to generate "good" SMILES. We now discuss how this is achieved. 781 We define the Augmented Likelihood  $^{23}$  of a SMILES, x, which is a linear combination between the 782 Prior and a reward term: 783

$$\log \pi_{\text{Augmented}}(x) = \log \pi_{\text{Prior}}(x) + \sigma R(x) \tag{7}$$

 $\log \pi_{\mathrm{Prior}}(x)$  is the log-probability of generating a given SMILES, x, under the Prior. Since the Prior's weights are fixed, the probability of sampling a given SMILES never changes. Models are typically parameterized by its weights,  $\theta$ . We take care here and omit  $\theta$  because the Prior, as stated previously, is not updated. Next, R is the reward function which defines the target objective, e.g., minimize docking score. Note that the reward function can contain multiple objectives, in which case, constituting a multi-parameter optimization objective. For example, in Experiment 3 of the main text, R is comprised of minimizing docking score, maximizing QED score  $^{87}$ , and minimizing SA score  $^{91}$ . R takes as input a SMILES, x, and returns a scalar reward  $\in$  [0, 1].  $\sigma$  is a hyperparameter that scales the contribution of the reward function. Importantly, given a SMILES, x, a low  $\sigma$  means the Augmented Likelihood converges to the Prior likelihood while a high  $\sigma$  means the Augmented Likelihood is dominated by the reward. In this work,  $\sigma$  is never changed and is 128 as this was found to work well in the original REINVENT work  $^{23}$ .

784 785

786

787

788

789

790

791

792

793

794

795

798

799

800

801

802

803

804

805

806

807

808

809

812

813

814

815

816

817

819

820

821

822

823

The loss function is defined as the squared difference between the Augmented Likelihood and the Agent Likelihood:

$$L(\theta) = (\log \pi_{\text{Augmented}}(x) - \log \pi_{\theta_{\text{Agent}}}(x))^2$$
 (8)

 $\log \pi_{\text{Agent}}(x)$  is the log-probability of generating a given SMILES, x, under the Agent. Importantly, we explicitly include  $\theta$  here because the Agent is updated. We stop here for a moment to discuss the implications of the loss function. The loss function tries to minimize the distance between the Augmented Likelihood and the Agent likelihood. Since the Augmented Likelihood (Eq. 7 is a linear combination of the Prior likelihood and the reward function, if the Agent generates "bad" SMILES, then the reward goes to 0 and the Augmented Likelihood converges to the Prior Likelihood. In this event, the Agent's weights actually regress back towards the Prior. This is because the Prior is pre-trained on a general dataset containing bio-active molecules (such as ChEMBL<sup>79</sup> and ZINC 250k<sup>89</sup>. The implicit assumption during pre-training is that these general datasets might actually already contain "good" molecules. Therefore, in the event that "bad" molecules are generated, the Prior acts as a "fall-back". On the other hand, when the reward is not 0, the Prior still "anchors" the Agent and does not let its weights deviate *too far* from the Prior (this is controlled by  $\sigma$ ). The reason for this is also because the Prior is assumed to potentially already contain "good" molecules. In practice, the Agent can deviate quite far from the Prior<sup>31</sup>. We now discuss an important implication of this loss function in Saturn. Saturn heavily leverages SMILES augmentation<sup>25</sup> as a data augmentation method to learn from the same molecular graph multiple times. Alternative SMILES sequences, while mapping to the same molecular graph, can have drastically different likelihoods. This is shown in Figure 2 in the main text where Saturn is trained to make it likely to generate all of these alternative SMILES forms. However, this does not always work. Because alternative SMILES forms have different likelihoods, there is the possibility that with the right combination of terms in the Augmented Likelihood, that it equals the Agent likelihood. In this case, the loss contribution is 0 so the Agent actually is not tuned to generate that particular SMILES form with higher likelihood. This is a contributing factor to Saturn's "hop-and-locally-explore" behavior. Given a set of augmented SMILES, if some of these SMILES cancel out in the loss function, then there is a smaller set of augmented SMILES that contribute to the loss function. With a smaller set, overfitting becomes more prone but we show that this mechanism actually benefits sample efficiency.

Finally, Saturn does not generate individual SMILES but rather, batches of SMILES. Therefore, the loss function is a batched loss:

$$L(\theta) = \frac{1}{|B|} \left[ \sum_{a \in A^*} (\log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}}) \right]^2$$
 (9)

The loss magnitude is the mean loss for a given batch, B, of sampled SMILES constructed following the actions,  $a \in A^*$ .

Minimizing the loss function is equivalent to maximizing the expected reward. In reinforcement learning, the general objective is to maximize the expected reward. In this section, we show how maximizing the expected reward is equivalent to minimizing the loss function. We first further define some preliminaries: sampling trajectories means sampling SMILES in our context. While there are often *intermediate* rewards during trajectory sampling, e.g., a drone tasked to fly to a target location might receive various rewards for how balanced it is during the flight, we set all intermediate rewards to 0. This is because rewards are only meaningful if the SMILES is a valid molecule. Technically, since the reward is directly the reward from the full trajectory, it is actually the **Return** in reinforcement learning terminology, but we use the term reward to match existing literature. Mathematically, the cost function (in reinforcement learning, J is used and we follow this convention) describes the expected reward when taking actions from a policy that is parameterized by a neural network (Mamba in our case):

$$J(\theta) = \mathbb{E}_{a_t \sim \pi_{\theta_{Agent}}} \left[ \sum_{t=1}^{T} R(a_t, s_t) \right]$$
 (10)

Since the expectation is in discrete space (sampling tokens is a discrete action), the cost function can be rewritten by transforming the expectation to a sum:

$$J(\theta) = \sum_{t=1}^{T} \sum_{a \in A_t} R(a_t, s_t) \pi_{\theta_{\text{Agent}}}(a_t | s_t)$$
(11)

The double summation is over all time-steps and actions (which token sampled) following the policy,  $\pi_{\theta}$ . Since we want to maximize the cost function, we take the derivative:

$$\nabla_{\theta} J(\theta) = \sum_{t=1}^{T} \sum_{a \in A_t} R(a_t, s_t) \nabla_{\theta} \pi_{\theta_{Agent}}(a_t | s_t)$$
 (12)

Next, the log-derivative trick:

$$\nabla_{\theta} J(\theta) = \sum_{t=1}^{T} \sum_{a \in A_t} R(a_t, s_t) \pi_{\theta_{Agent}}(a_t | s_t) \nabla_{\theta} \log \pi_{\theta}(a_t | s_t)$$
(13)

Using the definition of expectation for discrete space again, the cost function is rewritten:

$$\nabla_{\theta} J(\theta) = \mathbb{E}_{a_t \sim \pi_{\theta_{\text{Agent}}}} \left[ \sum_{t=1}^{T} R(a_t, s_t) \nabla_{\theta} \log \pi_{\theta_{\text{Agent}}}(a_t | s_t) \right]$$
 (14)

Computing the expectation exactly is intractable. This would involve sampling every single SMILES and computing their rewards. Therefore, the expectation is approximated by sampling a batch, B, of SMILES. Next, the set of actions taken in a batch at every time-step, is denoted  $A^*$ , which yield the specific SMILES generated:

$$\nabla_{\theta} J(\theta) = \frac{1}{|B|} \left[ \sum_{a \in A^*} R(a_t, s_t) \nabla_{\theta} \log \pi_{\theta_{Agent}}(a_t | s_t) \right]$$
 (15)

The reward, R is defined according to previous works  $^{21,23,76}$ :

$$R(a_t, s_t) = \log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}}$$
 (16)

Substituting the reward function:

$$\nabla_{\theta} J(\theta) = \frac{1}{|B|} \left[ \sum_{a \in A^*} \log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}} \right] \sum_{a \in A^*} \nabla_{\theta} \log \pi_{\theta_{\text{Agent}}}(a_t | s_t)$$
 (17)

852 Recalling the loss function:

$$L(\theta) = \frac{1}{|B|} \left[ \sum_{a \in A^*} (\log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}}) \right]^2$$
 (18)

Minimizing the loss function requires taking the derivative with respect to  $\theta$ :

$$\nabla_{\theta} L(\theta) = -2 \frac{1}{|B|} \left[ \sum_{a \in A^*} \log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}} \right] \sum_{a \in A^*} \nabla_{\theta} \log \pi_{\theta_{\text{Agent}}}$$
(19)

- The cost function (Eq. 17) is equivalent to the loss function (Eq. 19) up to a factor.
- 855 **Saturn Pseudo-code.** The pseudo-code for Saturn is presented here and the code is available at https://figshare.com/s/6040d65bfbfc29d6fedf.

```
Algorithm 1: Saturn Goal-directed Generation
```

**Input:** Oracle Budget Budget, Prior  $\pi_{Prior}$ , Augmentation Rounds A, Reward Function R, Sigma  $\sigma$ , Replay Buffer Size K, Genetic Algorithm GA

**Output:** Fine-tuned Agent Policy  $\pi_{\theta_{\text{Agent}}}$ , Generated Set G

# **Initialization:**

- 1. Generative Agent  $\pi_{\theta_{Agent}} = \pi_{Prior}$
- 2. Diversity Filter DF
- 3. Replay Buffer  $RB = \{\}$
- 4. Oracle Calls Calls = 0
- 5. Oracle Cache  $Cache = \{\}$
- 6. Generated Set  $G = \{\}$

# while C < Budget do

Sample batch of SMILES 
$$X = \{x_1, \dots, x_b\}$$
 with  $x_i \sim \pi_{\theta_{\Delta_{\text{gent}}}}$ ;

(Optionally) Generate SMILES using the Genetic Algorithm  $X_{\mathrm{GA}} = GA(RB)$ ;

$$X = X \cup X_{GA};$$

#### if X in Cache then

Retrieve rewards  $R_{\text{Cached}}$ 

Compute reward for *new* SMILES  $R(X_{New})$ ;

Update Generated Set tracking  $G = G \cup (X_{New}, R(X_{New}));$ 

Update Oracle Cache  $Cache = ((X_{New}, R_{New}) \cup Cache);$ 

Update Oracle Calls  $C = C + |X_{\text{New}}|$ ;

$$R(X) = R_{\text{Cached}} \cup R(X_{\text{New}});$$

Modify rewards based on the Diversity Filter R(X) = DF(X, R(X));

Update Replay Buffer  $RB = TopK(X \cup RB)$ ;

Compute Augmented Likelihood  $\log \pi_{\text{Augmented}}(X) = \log \pi_{\text{Prior}}(X) + \sigma R(X);$ 

Compute loss 
$$J(\theta) = (\log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}}(X))^2;$$

Update the Agent  $\pi_{\theta_{Agent}}$ ;

Purge Replay Buffer;

# $\mathbf{for}\ i \leftarrow 1\ \mathbf{to}\ A\ \mathbf{do}$

Augment sampled and Replay Buffer SMILES  $X_{\text{Augmented}}$ ;

Compute Augmented Likelihood of augmented SMILES (reward is unchanged)  $\log \pi_{\text{Augmented}} = \log \pi_{\text{Prior}}(X_{\text{Augmented}}) + \sigma R(X_{\text{Augmented}});$ 

 $\text{Compute loss } J(\theta)_{\text{Augmented}} = (\log \pi_{\text{Augmented}} - \log \pi_{\theta_{\text{Agent}}}(X_{\text{Augmented}}))^2;$ 

Update the Agent  $\pi_{\theta_{\operatorname{Agent}}}$ 

return  $\pi_{\theta_{Agent}}$ , G

# 857 B Saturn: Identifying Optimal Hyperparameters and Architecture

In this section, we present results from all hyperparameter investigations for Saturn. In particular, we formulated four questions (each devoted to one subsection) which we answer with empirical results and discussion on the test experiment which has the following multi-parameter optimization (MPO) objective: molecular weight (MW) < 350 Da, number of rings  $\geq$  2, and maximize topological polar surface area (tPSA).

Metrics. Following Guo et al.  $^{22}$ , the sample efficiency metrics are Yield and Oracle Burden (OB). Yield (Eq. 20) is the number of *unique* generated molecules above a reward threshold, T.

$$Yield = \sum_{g=1}^{G} \mathbb{I}[R(g) > T]$$
(20)

Oracle Burden (Eq. 21) is the number of oracle calls (c) required to generate N unique molecules above a reward threshold, T.

Oracle Burden = 
$$c \mid \sum_{g=1}^{G} \mathbb{I}[R(g) > T] = N$$
 (21)

The Yield and OB metrics are used to assess sample efficiency at the 0.7 reward threshold. In all tables, the number after OB parentheses is the number of successful replicates out of 10. All metrics other than IntDiv1<sup>71</sup> are rounded to the nearest integer. All individual experiments were run across 10 seeds (0-9 inclusive) and with a 1,000 oracle budget. All experiments were run sequentially on a workstation equipped with an NVIDIA RTX 3090 GPU and AMD Ryzen 9 5900X 12-Core CPU.

#### B.1 Data Pre-processing and Pre-training

873

882

883

884

885

886

887

888

889

890

891

893

894

Before presenting grid-search results, we first describe the full data pre-processing pipeline and design decisions made. The pre-training data for all experiments except **Part 3: Benchmarking Physics-based MPO Objective** in the main text (ZINC 250k <sup>89</sup> instead), was ChEMBL 33 <sup>79</sup>. We first downloaded the raw ChEMBL 33 from: https://ftp.ebi.ac.uk/pub/databases/chembl/ChEMBLdb/releases/chembl\_33/. There was no particular reason version 33 was chosen, other than it was the latest version at the time of experiments. We note that very recently (March 2024), version 34 was released.

The exact pre-processing steps along with the SMILES remaining after each step are:

- 1. Raw ChEMBL 33 2,372,674
- Standardization (charge and isotope handling) based on https://github.com/ MolecularAI/ReinventCommunity/blob/master/notebooks/Data\_Preparation. ipynb. All SMILES that could not be parsed by RDKit were removed - 2,312,459
- 3. Kept only the unique SMILES 2,203,884
- 4. Tokenize all SMILES based on REINVENT's tokenizer: https://github.com/ MolecularAI/reinvent-models/blob/main/reinvent\_models/reinvent\_core/ models/vocabulary.py
- 5. Keep SMILES  $\leq 80$  tokens 2,065,099
- 6.  $150 \le \text{molecular weight} \le 600 2,016,970$
- 7. Number of heavy atoms  $\leq 40 1,975,282$ 
  - 8. Number of rings  $\leq 8 1,974,522$
  - 9. Size of largest ring  $\leq 8 1,961,690$
- 10. Longest aliphatic carbon chain  $\leq 5 1,950,213$

- 11. Removed SMILES containing the following tokens (due to undesired chemistry and low token frequency): [S+], [C-], [s+], [O], [S@+], [S@@+], [S-], [o+], [NH+], [n-], [N@], [N@@], [N@#], [N@#+], [S@@], [C+], [S@], [c+], [NH2+], [SH], [NH-], [CH-], [O+], [C-], [CH], [SH+], [CH2-], [OH+], [nH+], [SH2] **1,942,081**
- The final vocabulary contained 37 tokens (2 extra tokens were added, indicating <START> and <END>). We note that stereochemistry tokens were kept (this is not the case for REINVENT<sup>24</sup>).
- In this work, we investigated LSTM<sup>26</sup> RNN, decoder transformer<sup>27,28</sup>, and Mamba<sup>29</sup>. Given a vocabulary of 37, the model parameters were as follows:
- 904 1. RNN: 5,807,909 (based on REINVENT<sup>24</sup>)
  - 2. Decoder Transformer 6,337,061 (based on recent work <sup>40</sup> that applied this model size and used a similar loss function to REINVENT)
  - 3. Mamba: 5,265,920 (based on similar size to RNN)

The exact hyperparameters of each architecture are the default arguments in the codebase. Each training step consisted of a full pass through the dataset. The key pre-training parameters were:

- 910 1. Max training steps = 20
- 911 2. Seed = 0

905

906

907

- 912 3. Batch size = 512
- 913 4. Learning rate = 0.0001
- 5. Randomize<sup>25</sup> every batch of SMILES
- 915 The following model checkpoints were used:
- 916 1. RNN: Epoch 18, NLL = 34.61, Validity (10k) = 94.48%
- 2. Decoder Transformer Epoch 20, NLL = 33.38, Validity (10k) = 96.04%
- 918 3. Mamba: Epoch 18, NLL = 32.21, Validity (10k) = 95.60%

# 919 B.2 Understanding the Limits of Augmented Memory

Augmented Memory<sup>21</sup> improves sample efficiency by repeated learning on the high reward SMILES 920 stored in the replay buffer (referred to as Buffer from here on). In the original work, ablation 921 experiments showed that updating the agent with only the Buffer resulted in minimal difference. This 922 suggests that a viable way to exploiting the gains from Augmented Memory is to simply have new 923 examples of high reward SMILES being added to the Buffer. In the original work, the number of 924 augmentation rounds was capped at two to mitigate mode collapse. In this work, we assume near deterministic rewards and use caching to handle repeated generations. Under this assumption, our hypothesis in this subsection is: as long as unique high reward SMILES are generated, increasing 927 augmentation rounds can further improve sample efficiency. Correspondingly, we perform a grid 928 search using Augmented Memory's default generator architecture (LSTM<sup>26</sup> RNN) and vary the batch 929 size (64, 32, 16, 8) and augmentation rounds (0-20 inclusive except 1) where 0 augmentation rounds 930 is equivalent to REINVENT<sup>23,24</sup>. The results are shown in Tables 5, 6, 7, and 8. 931

#### **Increasing augmentation rounds:**

- 1. Decreases diversity, as expected.
- 2. Increases the number of repeated SMILES.

# Decreasing batch size:

932

933

934

935

936

937

938

- 1. Monotonically improves sample efficiency (though not always significant at the 95% confidence level).
- 2. Benefits Augmented memory more than REINVENT (0 augmentation rounds).
- 3. Increases the number of repeated SMILES.

Table 5: RNN batch size 64.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
RNN	0	0±0	_	0±0	584±251 (5)	Failed (0)	Failed (0)	1±1
RNN	2	15±9	0.775±0.073	15±9	644±173 (10)	941±58 (8)	Failed (0)	0±0
RNN	3	33±42	$0.788 \pm 0.043$	32±40	613±96 (10)	927±128 (9)	993±0(1)	0±0
RNN	4	32±16	0.813±0.024	31±16	527±198 (10)	880±90 (10)	Failed (0)	0±0
RNN	5	40±14	0.812±0.023	39±13	459±177 (10)	862±68 (10)	Failed (0)	0±0
RNN	6	41±32	0.805±0.032	39±28	492±184 (10)	852±99 (9)	1041±0(1)	0±0
RNN	7	47±25	0.814±0.019	46±24	543±188 (10)	842±93 (10)	1055±0(1)	0±0
RNN	8	28±16	0.801±0.032	27±16	557±173 (10)	912±82 (9)	Failed (0)	0±0
RNN	9	21±13	$0.742 \pm 0.124$	21±13	596±215 (10)	918±61 (8)	Failed (0)	1±2
RNN	10	27±18	0.796±0.046	27±18	511±266 (10)	859±65 (8)	Failed (0)	0±0
RNN	11	20±14	0.749±0.115	20±14	611±235 (10)	938±85 (8)	Failed (0)	1±2
RNN	12	48±18	0.813±0.022	46±18	468±206 (10)	851±55 (10)	Failed (0)	1±1
RNN	13	57±43	0.808±0.027	54±39	446±213 (10)	822±144 (10)	952±0(1)	1±2
RNN	14	33±13	0.801±0.024	32±13	587±175 (10)	884±79 (10)	Failed (0)	1±1
RNN	15	47±32	0.797±0.037	46±32	532±196 (10)	836±122 (10)	1052±0(1)	2±2
RNN	16	$34 \pm 32$	0.783±0.026	33±30	647±208 (10)	918±97 (10)	1034±0 (1)	3±4
RNN	17	31±29	0.769±0.06	30±29	645±176 (10)	870±99 (7)	Failed (0)	3±4
RNN	18	35±28	0.774±0.035	$32\pm24$	673±125 (10)	898±88 (8)	1053±0(1)	7±5
RNN	19	43±41	0.781±0.034	40±36	659±183 (10)	875±111 (8)	949±0(1)	7±9
RNN	20	51±29	0.792±0.03	48±28	583±187 (10)	837±133 (10)	1056±0 (1)	3±2

Table 6: RNN batch size 32.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
RNN	0	0±0	_	0±0	798±101 (5)	Failed (0)	Failed (0)	1±1
RNN	2	43±25	0.825±0.029	42±24	608±151 (10)	844±90 (9)	Failed (0)	0±0
RNN	3	52±34	0.810±0.059	51±32	522±141 (10)	789±100 (9)	1018±0(2)	0±1
RNN	4	87±33	0.820±0.018	83±31	466±120 (10)	740±77 (10)	987±30 (4)	1±3
RNN	5	98±57	0.817±0.027	89±50	408±184 (10)	714±136 (10)	915±20 (4)	1±2
RNN	6	76±50	$0.808\pm0.028$	71±43	476±159 (10)	783±99 (10)	927±30(2)	1±3
RNN	7	78±40	0.805±0.027	72±40	478±90 (10)	760±70 (10)	942±26 (2)	3±7
RNN	8	89±72	0.798±0.036	78±58	529±165 (10)	767±146 (10)	899±48 (3)	9±13
RNN	9	57±52	0.781±0.046	50±42	608±186 (10)	811±143 (9)	977±36 (3)	5±4
RNN	10	90±65	0.788±0.031	82±55	549±158 (10)	769±142 (10)	977±66 (5)	9±14
RNN	11	60±43	0.755±0.105	57±43	593±207 (10)	781±83 (8)	969±52 (2)	2±2
RNN	12	103±83	0.790±0.021	90±72	534±168 (10)	763±158 (10)	930±105 (4)	10±23
RNN	13	72±57	0.749±0.065	62±52	578±155 (10)	765±134 (8)	958±54 (3)	12±9
RNN	14	95±55	0.779±0.027	83±47	463±173 (10)	758±110 (10)	964±28 (5)	16±15
RNN	15	74±60	0.784±0.036	66±52	554±92 (10)	820±124 (10)	963±54 (4)	22±20
RNN	16	84±60	0.758±0.07	70±44	544±209 (10)	768±105 (9)	957±42 (5)	17±19
RNN	17	112±74	0.765±0.067	96±56	474±131 (10)	729±105 (10)	908±96 (4)	21±21
RNN	18	77±49	0.774±0.039	67±43	533±100 (10)	779±102 (10)	927±12 (2)	35±32
RNN	19	84±56	0.749±0.037	68±50	535±181 (10)	788±127 (10)	951±61 (3)	33±44
RNN	20	76±77	0.717±0.094	64±61	653±200 (10)	810±121 (9)	919±76 (3)	56±64

- 4. Increases variance, as expected (since the expected reward is being approximated with a smaller batch size so it is more noisy).
- 5. Decreases diversity.

Taking these observations together, increasing augmentation rounds and decreasing batch size can trade-off diversity for sample efficiency (inconsistently and with higher variance).

# **B.3** Do Architectures Differ in Behavior?

RNNs essentially solve the validity, uniqueness, and novelty metrics <sup>70,71</sup> and can learn molecular distributions well <sup>109</sup> for small molecule design. In this subsection, we extend Augmented Memory to decoder transformer <sup>27,28</sup> and Mamba <sup>29</sup> to investigate the RL dynamics and empirically investigate potential benefits. Our hypothesis is that since self-attention <sup>27</sup> and selective scanning <sup>29</sup> *can* capture different structural elements <sup>69</sup> (via focusing on different aspects of the sequence), benefits *may* arise from capturing and focusing on favorable moieties. Our analysis is focused solely on sample efficiency metrics and not validity, uniqueness, and novelty.

Similar to the previous subsection, we perform a grid-search over batch size (64, 32, 16, 8) and augmentation rounds (0-20 inclusive except 1). As the results for RNN were presented in the previous subsection, this subsection only shows Decoder and Mamba results (Tables 9, 10, 11, 12, 13, 14, 15, and 16).

Table 7: RNN batch size 16.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
RNN	0	8±9	0.700±0.126	8±9	546±263 (8)	837±144 (3)	Failed (0)	1±1
RNN	2	86±40	0.819±0.026	82±38	409±158 (10)	709±86 (10)	907±14 (2)	2±4
RNN	3	103±47	0.831±0.027	100±44	406±157 (10)	706±98 (10)	942±45 (5)	2±3
RNN	4	90±62	0.828±0.017	83±53	440±152 (10)	741±102 (10)	916±76 (3)	1±1
RNN	5	107±58	0.814±0.036	101±54	480±118 (10)	721±109 (10)	916±53 (4)	7±7
RNN	6	121±80	0.791±0.040	107±68	493±214 (10)	713±156 (10)	895±107 (5)	12±11
RNN	7	144±107	$0.776 \pm 0.026$	117±86	467±186 (10)	684±136 (10)	871±116 (6)	38±82
RNN	8	120±95	0.734±0.128	104±85	481±288 (10)	653±145 (8)	854±54 (5)	18±28
RNN	9	141±104	0.783±0.048	112±72	453±211 (10)	654±154 (9)	871±104 (6)	59±95
RNN	10	106±76	0.760±0.0560	84±63	510±201 (10)	733±122 (9)	913±64 (5)	43±47
RNN	11	120±105	$0.764 \pm 0.032$	95±81	500±220 (10)	741±199 (10)	829±99 (4)	42±37
RNN	12	171±140	0.769±0.028	124±109	389±209 (10)	662±186 (10)	774±128 (5)	39±30
RNN	13	133±106	0.767±0.038	106±93	510±186 (10)	690±162 (10)	826±131 (4)	83±88
RNN	14	166±130	0.769±0.045	129±93	413±237 (10)	659±195 (10)	777±94 (5)	93±69
RNN	15	154±89	$0.732\pm0.064$	127±78	504±162 (10)	647±124 (9)	861±59 (7)	94±75
RNN	16	156±155	0.716±0.094	109±109	517±196 (10)	682±202 (9)	838±182 (6)	143±120
RNN	17	141±82	0.737±0.059	98±49	444±181 (10)	696±128 (10)	894±71 (7)	198±163
RNN	18	189±136	0.727±0.044	152±119	469±212 (10)	657±174 (10)	832±141 (7)	247±210
RNN	19	162±121	$0.654\pm0.165$	119±98	507±257 (10)	625±137 (8)	836±109 (7)	210±128
RNN	20	139±110	0.732±0.045	91±67	492±188 (10)	720±157 (10)	847±110 (5)	262±179

Table 8: RNN batch size 8.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
RNN	0	21±21	0.645±0.133	17±18	481±291 (10)	826±95 (6)	Failed (0)	16±15
RNN	2	136±100	0.807±0.028	113±73	428±169 (10)	665±159 (10)	849±113 (5)	8±9
RNN	3	143±97	0.793±0.037	131±85	395±169 (10)	667±126 (10)	863±109 (6)	27±33
RNN	4	152±115	0.785±0.022	129±96	379±212 (10)	680±179 (10)	865±124 (7)	44±47
RNN	5	164±84	0.786±0.038	123±56	350±158 (10)	643±121 (10)	876±81 (8)	40±41
RNN	6	224±104	0.790±0.041	181±79	352±176 (10)	584±159 (10)	782±56 (8)	49±40
RNN	7	185±111	0.751±0.070	151±96	435±224 (10)	608±127 (9)	814±86 (7)	116±119
RNN	8	159±128	0.775±0.050	128±114	460±195 (10)	646±145 (9)	858±140 (7)	105±77
RNN	9	198±164	$0.732\pm0.072$	151±121	451±227 (10)	641±158 (9)	782±168 (6)	285±396
RNN	10	139±127	0.728±0.078	100±73	512±212 (8)	702±124 (7)	867±145 (4)	112±61
RNN	11	205±173	0.753±0.062	151±120	444±267 (10)	652±234 (10)	737±167 (6)	254±320
RNN	12	261±165	0.762±0.057	211±135	320±246 (10)	579±210 (10)	775±168 (9)	518±760
RNN	13	231±198	0.753±0.061	155±101	444±184 (9)	601±235 (9)	790±214 (8)	351±289
RNN	14	158±103	0.718±0.091	108±60	526±208 (10)	681±127 (9)	845±80 (6)	374±308
RNN	15	221±128	$0.731 \pm 0.043$	150±129	439±196 (10)	618±168 (10)	826±153 (9)	461±292
RNN	16	196±145	0.725±0.043	136±101	470±228 (10)	683±198 (10)	813±141 (7)	694±495
RNN	17	258±130	0.689±0.119	193±94	467±210 (10)	576±139 (9)	787±115 (9)	796±600
RNN	18	253±114	0.727±0.047	195±98	394±175 (10)	605±124 (10)	764±82 (8)	1112±974
RNN	19	268±159	0.714±0.052	204±132	418±161 (10)	579±167 (10)	745±153 (8)	817±811
RNN	20	292±153	0.713±0.039	220±121	397±205 (10)	574±188 (10)	776±173 (10)	1406±1391

# The following observations are similar to RNN. Increasing augmentation rounds:

- 1. Decreases diversity, as expected.
  - 2. Increases the number of repeated SMILES.

# 960 Decreasing batch size:

958

959

961

962

963

964

965

966

967

969

970

971

972

- Monotonically improves sample efficiency (though not always significant at the 95% confidence level).
- 2. Benefits Augmented memory more than REINVENT (0 augmentation rounds).
- 3. Increases the number of repeated SMILES.
- 4. Increases variance, as expected (since the expected reward is being approximated with a smaller batch size so it is more noisy).
- 5. Decreases diversity.

# 968 The following observations contrast RNN with Decoder and Mamba:

- 1. Mamba > Decoder > RNN in terms of NLL convergence (end of Appendix B.1).
- 2. Propensity to generate repeated SMILES follows the same trend and is further supported with the IntDiv1 generally being lower than RNN for the same number of augmentation rounds across all batch sizes.

Table 9: Decoder batch size 64.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Decoder	0	1±1	0.548±0.129	1±1	691±266 (6)	Failed (0)	Failed (0)	2±1
Decoder	2	26±19	0.800±0.061	26±18	524±128 (10)	868±76 (8)	Failed (0)	0±0
Decoder	3	37±24	0.801±0.031	36±23	629±154 (10)	849±85 (9)	Failed (0)	0±0
Decoder	4	49±38	0.797±0.055	48±37	590±142 (10)	851±89 (9)	984±0(1)	0±0
Decoder	5	63±35	0.821±0.014	62±35	545±136 (10)	814±84 (10)	997±21 (2)	1±1
Decoder	6	43±34	0.794±0.033	40±32	649±155 (10)	881±127 (10)	1045±0(1)	2±4
Decoder	7	42±29	0.800±0.039	41±29	585±175 (10)	859±116 (9)	1042±0(1)	4±3
Decoder	8	22±28	0.719±0.119	21±28	717±157 (10)	939±104 (7)	1051±0(1)	6±6
Decoder	9	23±22	0.704±0.156	19±16	618±233 (10)	889±92 (7)	Failed (0)	10±5
Decoder	10	43±48	0.768±0.056	41±47	643±110 (10)	788±104 (6)	980±0 (1)	10±7
Decoder	11	36±45	0.756±0.068	34±44	698±116 (10)	881±108 (8)	891±0(1)	9±7
Decoder	12	47±28	0.795±0.02	43±27	609±101 (9)	862±74 (9)	1046±0(1)	16±9
Decoder	13	66±66	0.727±0.109	56±54	641±216 (10)	788±148 (8)	975±75 (2)	37±25
Decoder	14	38±37	0.696±0.139	33±34	679±169 (10)	868±104 (7)	1004±0(1)	46±28
Decoder	15	38±56	0.671±0.100	25±32	668±241 (9)	809±159 (5)	977±9 (2)	56±28
Decoder	16	33±41	0.716±0.084	25±29	572±221 (10)	900±122 (8)	984±0(1)	78±38
Decoder	17	50±48	0.707±0.091	37±30	595±250 (10)	797±86 (7)	1007±34 (2)	91±42
Decoder	18	30±36	0.732±0.049	26±32	701±135 (8)	886±101 (6)	1025±0(1)	124±41
Decoder	19	$35\pm31$	0.715±0.056	28±21	640±240 (10)	852±155 (8)	1031±0(1)	159±64
Decoder	20	51±51	0.733±0.047	39±38	585±277 (9)	862±136 (8)	984±49 (2)	172±69

- Mamba notably generates many repeated SMILES but sample efficiency improves, thus it is not detrimental under the assumption that the reward is *near deterministic* and oracle evaluations are cached.
- 4. In general, Decoder does not outperform RNN

973

974

975

976

977

978

979

980

981

982

983

984

988

989

990

991

992

993

994

995

996

997

998

999

1000

1001

1002

1003

1004

1005

1006

Taking these observations together and exactly like RNN results, increasing augmentation rounds and decreasing batch size *can* trade-off diversity for sample efficiency (inconsistently and with higher variance). However, of difference, is that Mamba at lower batch sizes (particularly 16) and relatively high augmentation rounds (10) improves sample efficiency in a statistically significant way (at the 95% confidence level).

We further note that we have observed that with low batch size and high augmentation rounds, Mamba can temporarily lose generative ability. Specifically, the validity of the generated batch can be 0. Sampling a new batch can recover this validity but we have observed in extremely rare cases, that validity can be 0 for over 10 successive epochs. We observed this scenario twice in over 5,000 experiments, occurring with a batch size of 8 and augmentation rounds 19 and 20. We speculate the reason is extreme mode collapse to a chemical space where syntax is sensitive. Consequently, once the Selective Memory Purge starts penalizing the reward and the agent is brought back towards the prior, large gradient updates coupled with sensitive syntax may cause invalid SMILES. This process often recovers but in practice, with high-fidelity oracles, one would checkpoint models frequently (even every epoch), as each batch of oracle evaluation would be costly. Alternatively, as all high reward SMILES (so far) generated can be pre-emptively saved. It would be feasible to even start a new run with these SMILES seeded in the replay buffer, akin to inception in REINVENT<sup>24</sup> (transfer learning would work too). This would kick-start the optimization and already guide the agent to this chemical space, preventing optimization progress from completely "lost". Moreover, we also do not recommend a batch size of 8 and augmentation rounds above 10 as the performance variance becomes high. This behavior is likely also highly dependent on the objective function which affects the optimization landscape. Finally, in the rare cases this occurs, and when validity recovers, the effect is minimal as sampling is cheap compared to oracle evaluations. We write this note for full transparency into all the behavior we have observed in our grid-search.

#### **B.4** Are Increased Augmentation Rounds still Synergistic with Beam Enumeration?

Beam Enumeration <sup>22</sup> extracts the most probable substructures for self-conditioned generation and has been shown to be synergistic with Augmented Memory <sup>21</sup> such that the Yield and OB improve. In the original work, the oracle budget in the experiments was 5,000. In this work, we are interested in minimizing the oracle budget and all experiments thus far use a 1,000 oracle budget. Beam Enumeration has a *Patience* criterion which controls when substructures are extracted: only when the average reward improves for *Patience* number of successive epochs. Since we are operating at a much lower oracle budget, it is especially unclear whether Beam Enumeration can still benefit

Table 10: Decoder batch size 32.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Decoder	0	4±4	0.710±0.023	4±4	647±232 (6)	982±39 (2)	Failed (0)	10±13
Decoder	2	45±23	0.813±0.021	43±22	557±174 (10)	844±91 (10)	Failed (0)	1±1
Decoder	3	66±44	0.801±0.033	63±43	515±146 (10)	779±70 (9)	918±0(1)	1±1
Decoder	4	111±88	0.791±0.017	100±80	476±131 (10)	726±133 (10)	908±81 (5)	3±3
Decoder	5	94±70	0.791±0.043	81±53	489±155 (10)	753±112 (9)	897±63 (3)	3±2
Decoder	6	94±66	0.770±0.075	82±60	476±204 (10)	696±126 (9)	921±52 (4)	11±6
Decoder	7	117±87	$0.730\pm0.084$	105±84	473±270 (10)	659±99 (8)	936±93 (6)	54±84
Decoder	8	78±69	0.776±0.032	67±52	519±204 (10)	797±147 (10)	926±94 (3)	35±13
Decoder	9	59±35	0.767±0.032	51±32	575±76 (10)	856±83 (10)	968±0(1)	44±33
Decoder	10	91±75	0.742±0.065	68±52	492±176 (9)	769±121 (9)	879±66 (2)	77±56
Decoder	11	70±46	0.739±0.059	57±36	559±128 (10)	811±96 (10)	974±6 (3)	84±45
Decoder	12	114±58	0.730±0.041	82±45	559±177 (10)	715±59 (9)	942±48 (6)	124±81
Decoder	13	93±83	0.741±0.064	77±68	598±114 (10)	788±129 (9)	874±34 (3)	146±76
Decoder	14	147±112	0.752±0.064	109±84	486±147 (9)	694±152 (9)	791±37 (4)	257±269
Decoder	15	140±100	0.718±0.085	111±78	516±256 (10)	676±143 (9)	916±106 (7)	222±128
Decoder	16	130±142	0.709±0.045	82±66	552±177 (10)	772±164 (10)	851±173 (4)	405±272
Decoder	17	130±125	0.720±0.075	95±89	624±209 (10)	771±186 (10)	841±137 (4)	444±265
Decoder	18	153±165	0.718±0.055	110±130	565±191 (10)	718±197 (9)	668±81 (3)	544±503
Decoder	19	149±94	0.686±0.055	104±69	547±215 (10)	731±113 (9)	897±83 (7)	594±172
Decoder	20	137±135	0.693±0.046	78±56	555±200 (9)	740±181 (9)	855±145 (5)	514±399

Table 11: Decoder batch size 16.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Decoder	0	2±3	0.55±0.1	2±2	810±93 (7)	983±0(1)	Failed (0)	78±25
Decoder	2	66±50	0.796±0.037	59±41	602±158 (10)	799±106 (9)	921±3 (2)	8±7
Decoder	3	84±66	0.77±0.037	64±44	536±170 (10)	769±122 (9)	919±44 (4)	28±24
Decoder	4	71±44	$0.74\pm0.102$	62±41	632±118 (10)	780±82 (9)	977±36 (3)	22±12
Decoder	5	154±93	0.748±0.052	122±70	439±151 (10)	679±128 (10)	907±92 (8)	90±90
Decoder	6	116±94	0.748±0.039	86±64	517±165 (10)	728±158 (10)	904±126 (5)	73±42
Decoder	7	108±85	0.747±0.051	71±50	510±222 (10)	740±127 (9)	868±48 (4)	126±63
Decoder	8	108±94	0.708±0.109	72±57	538±164 (10)	742±116 (9)	887±87 (4)	150±72
Decoder	9	78±83	0.687±0.116	51±55	614±244 (10)	790±150 (8)	890±62 (3)	242±139
Decoder	10	120±128	0.691±0.042	74±73	663±170 (9)	768±169 (8)	805±65 (4)	344±218
Decoder	11	146±134	0.727±0.038	110±100	609±169 (9)	725±166 (9)	829±132 (5)	389±199
Decoder	12	119±127	0.704±0.047	76±68	624±185 (9)	779±176 (9)	828±110 (4)	363±256
Decoder	13	183±177	0.696±0.031	97±80	484±227 (9)	671±216 (9)	753±144 (5)	498±412
Decoder	14	146±111	0.673±0.055	88±60	572±240 (10)	737±162 (9)	850±87 (6)	702±387
Decoder	15	146±100	$0.64\pm0.123$	108±79	623±141 (10)	772±150 (10)	867±70 (6)	774±414
Decoder	16	209±173	0.688±0.043	155±130	530±124 (9)	654±161 (9)	813±170 (7)	1369±777
Decoder	17	190±168	0.662±0.109	154±149	571±207 (10)	674±179 (9)	746±162 (5)	1096±883
Decoder	18	226±138	0.668±0.052	174±115	550±156 (10)	646±131 (9)	802±118 (8)	1540±986
Decoder	19	232±154	$0.648 \pm 0.07$	168±96	564±152 (10)	681±161 (10)	781±147 (7)	1693±1165
Decoder	20	258±200	0.636±0.077	166±103	448±223 (9)	589±179 (8)	763±177 (8)	1741±1020

Table 12: Decoder batch size 8.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Decoder	0	57±64	0.621±0.222	37±36	554±137 (9)	766±178 (7)	912±52 (3)	368±164
Decoder	2	120±76	0.745±0.055	97±59	497±207 (10)	667±110 (8)	913±62 (7)	39±22
Decoder	3	93±60	0.73±0.06	74±45	530±166 (10)	759±87 (9)	918±22 (4)	128±82
Decoder	4	111±49	0.741±0.036	79±34	467±170 (10)	737±101 (10)	950±32 (7)	173±81
Decoder	5	79±82	0.724±0.044	59±54	609±123 (8)	805±101 (8)	901±72 (3)	283±179
Decoder	6	138±112	0.72±0.062	96±78	608±162 (10)	737±138 (9)	843±81 (5)	400±222
Decoder	7	197±165	0.688±0.064	149±131	502±287 (10)	684±237 (10)	758±112 (6)	820±1051
Decoder	8	219±179	0.68±0.063	132±120	475±201 (8)	581±127 (7)	763±136 (7)	840±900
Decoder	9	194±144	0.651±0.049	153±118	496±157 (8)	627±149 (8)	791±109 (7)	1059±864
Decoder	10	183±200	0.684±0.055	130±130	571±201 (9)	654±217 (8)	789±205 (6)	944±597
Decoder	11	141±123	0.581±0.166	96±84	617±198 (9)	662±142 (7)	801±97 (5)	1715±1380
Decoder	12	133±196	0.574±0.149	92±135	665±291 (9)	699±268 (7)	664±209 (3)	1604±1130
Decoder	13	331±151	0.664±0.095	271±143	418±230 (10)	503±88 (9)	711±107 (9)	2030±1408
Decoder	14	164±152	$0.602\pm0.06$	125±109	620±257 (9)	714±194 (8)	825±133 (6)	2628±1665
Decoder	15	281±242	0.661±0.054	230±185	496±243 (9)	589±251 (9)	663±201 (7)	2482±1515
Decoder	16	213±191	0.58±0.143	180±176	512±245 (9)	596±223 (8)	730±186 (6)	3113±2436
Decoder	17	252±186	0.622±0.072	203±167	614±231 (10)	615±169 (8)	735±139 (7)	3278±1894
Decoder	18	81±113	0.595±0.064	69±97	630±232 (7)	759±209 (7)	862±102 (3)	2811±2415
Decoder	19	136±171	0.611±0.062	119±154	645±195 (7)	708±180 (6)	771±142 (4)	2886±2066
Decoder	20	98±139	0.54±0.075	91±136	736±195 (7)	785±160 (6)	813±140 (3)	3190±2113

Table 13: Mamba batch size 64.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Mamba	0	0±0	_	0±0	946±41 (2)	Failed (0)	Failed (0)	0±1
Mamba	2	2±1	0.580±0.086	2±1	817±244 (10)	Failed (0)	Failed (0)	0±0
Mamba	3	9±6	0.734±0.068	9±6	659±234 (9)	942±34 (4)	Failed (0)	1±1
Mamba	4	6±3	0.672±0.114	6±3	652±297 (10)	1040±7 (2)	Failed (0)	2±2
Mamba	5	9±5	0.697±0.113	9±5	640±210 (10)	995±30 (5)	Failed (0)	3±3
Mamba	6	17±11	$0.770\pm0.041$	17±11	656±119 (10)	960±90 (9)	Failed (0)	6±4
Mamba	7	19±6	0.769±0.027	18±6	623±152 (10)	957±65 (9)	Failed (0)	7±3
Mamba	8	29±15	0.786±0.035	27±15	545±176 (10)	917±82 (10)	Failed (0)	12±8
Mamba	9	21±10	0.755±0.075	20±10	585±192 (10)	938±57 (9)	Failed (0)	26±23
Mamba	10	34±22	0.785±0.028	28±15	486±176 (10)	884±91 (10)	Failed (0)	30±21
Mamba	11	18±8	0.757±0.044	17±7	550±203 (10)	937±31 (8)	Failed (0)	37±21
Mamba	12	22±17	0.727±0.051	20±15	629±234 (10)	876±53 (6)	Failed (0)	72±68
Mamba	13	33±33	0.739±0.090	29±28	561±222 (10)	915±120 (10)	1020±0(1)	62±28
Mamba	14	47±39	0.701±0.138	30±15	540±242 (10)	839±94 (8)	980±0(1)	127±56
Mamba	15	60±88	0.725±0.117	31±17	585±225 (10)	866±143 (10)	726±0(1)	136±112
Mamba	16	46±40	0.661±0.170	29±22	614±193 (10)	865±104 (9)	978±33 (2)	199±89
Mamba	17	43±24	0.727±0.054	30±13	538±185 (10)	866±101 (10)	Failed (0)	174±77
Mamba	18	51±42	0.732±0.056	40±32	621±219 (10)	838±111 (9)	995±34 (2)	262±99
Mamba	19	49±40	0.723±0.048	36±25	633±218 (10)	829±123 (8)	975±0(1)	241±73
Mamba	20	77±68	0.695±0.088	46±32	549±241 (9)	771±146 (8)	940±76 (3)	385±180

Table 14: Mamba batch size 32.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Mamba	0	0±0	_	0±0	773±189 (4)	Failed (0)	Failed (0)	4±2
Mamba	2	12±7	0.744±0.060	12±7	644±199 (10)	933±29 (5)	Failed (0)	3±2
Mamba	3	16±9	0.759±0.050	15±9	640±158 (10)	912±45 (6)	Failed (0)	8±7
Mamba	4	30±15	0.797±0.029	29±15	579±140 (10)	879±86 (10)	Failed (0)	11±5
Mamba	5	38±23	0.718±0.151	35±21	695±159 (10)	833±83 (8)	Failed (0)	24±9
Mamba	6	44±37	0.770±0.044	41±34	564±145 (10)	861±110 (9)	1000±3 (2)	42±17
Mamba	7	52±43	0.750±0.047	46±37	539±174 (10)	848±123 (10)	996±11 (2)	68±28
Mamba	8	76±51	0.775±0.025	67±45	515±108 (10)	794±85 (10)	923±30 (2)	90±49
Mamba	9	64±47	0.755±0.083	53±38	546±143 (10)	808±116 (10)	959±45 (2)	140±106
Mamba	10	96±76	0.768±0.028	75±54	553±186 (10)	782±161 (10)	949±84 (5)	165±63
Mamba	11	87±60	$0.732\pm0.045$	62±40	592±218 (10)	741±105 (8)	936±31 (3)	303±152
Mamba	12	118±60	0.680±0.130	67±21	500±159 (10)	730±132 (10)	932±61 (6)	280±151
Mamba	13	92±60	$0.742\pm0.082$	74±43	578±226 (10)	771±98 (9)	940±39 (4)	353±104
Mamba	14	166±75	0.748±0.041	121±54	458±97 (10)	659±64 (10)	901±78 (8)	483±202
Mamba	15	139±94	0.755±0.033	106±72	456±141 (10)	740±127 (10)	847±54 (5)	488±167
Mamba	16	136±75	0.740±0.039	97±54	571±131 (10)	742±119 (10)	899±50 (6)	769±354
Mamba	17	186±88	0.696±0.058	138±83	510±103 (10)	683±88 (10)	871±76 (8)	937±677
Mamba	18	214±87	0.723±0.059	169±81	540±113 (10)	672±88 (10)	862±84 (9)	1027±554
Mamba	19	242±109	0.686±0.041	184±104	493±133 (10)	661±116 (10)	819±109 (9)	1376±596
Mamba	20	187±78	0.706±0.038	152±67	557±101 (10)	714±80 (10)	892±79 (9)	1183±413

Table 15: Mamba batch size 16.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Mamba	0	3±4	0.417±0.161	2±2	545±232 (7)	982±0 (1)	Failed (0)	91±32
Mamba	2	39±29	0.761±0.047	34±23	609±165 (10)	829±117 (9)	Failed (0)	46±31
Mamba	3	61±51	0.771±0.051	50±39	498±193 (10)	797±118 (9)	953±15 (3)	71±28
Mamba	4	52±33	0.779±0.031	42±23	581±102 (10)	817±112 (10)	970±0(1)	139±59
Mamba	5	69±38	0.764±0.052	54±28	542±93 (10)	807±76 (10)	988±17 (3)	178±90
Mamba	6	138±46	0.759±0.039	110±42	456±89 (10)	693±75 (10)	919±36 (7)	286±137
Mamba	7	174±95	0.737±0.059	127±83	427±177 (10)	643±102 (10)	858±77 (7)	395±147
Mamba	8	209±95	0.751±0.030	137±60	461±151 (10)	617±135 (10)	817±71 (8)	482±214
Mamba	9	202±98	0.735±0.032	137±80	389±112 (10)	631±102 (10)	841±92 (8)	518±237
Mamba	10	306±57	0.714±0.035	206±34	387±148 (10)	555±66 (10)	761±58 (10)	1110±636
Mamba	11	306±92	0.716±0.039	237±85	403±136 (10)	554±93 (10)	761±100 (10)	1341±596
Mamba	12	266±100	0.723±0.041	199±83	392±126 (10)	590±100 (10)	806±111 (10)	1312±666
Mamba	13	327±108	$0.722\pm0.043$	258±101	428±111 (10)	549±111 (10)	741±116 (10)	1508±780
Mamba	14	318±109	0.695±0.061	246±117	416±164 (10)	535±148 (10)	736±123 (10)	1776±912
Mamba	15	284±74	0.691±0.052	219±42	442±67 (10)	584±87 (10)	785±82 (10)	2629±939
Mamba	16	293±112	0.672±0.053	209±77	483±145 (10)	570±136 (10)	767±130 (10)	2284±1011
Mamba	17	344±115	0.656±0.047	278±92	462±113 (10)	563±98 (10)	725±121 (10)	3512±1227
Mamba	18	281±155	0.640±0.082	216±125	464±174 (9)	595±155 (9)	730±93 (8)	2885±1344
Mamba	19	307±115	0.624±0.084	238±102	491±146 (10)	579±133 (10)	750±119 (10)	3318±1347
Mamba	20	352±69	0.673±0.046	294±61	403±102 (10)	525±81 (10)	714±79 (10)	3331±1454

Table 16: Mamba batch size 8.

Model	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats
Mamba	0	3±2	0.43±0.133	2±1	498±322 (8)	Failed (0)	Failed (0)	940±234
Mamba	2	69±32	0.755±0.059	56±28	453±176 (10)	780±78 (10)	992±8 (2)	214±72
Mamba	3	156±113	0.745±0.035	109±70	452±221 (10)	659±143 (9)	792±83 (5)	282±120
Mamba	4	200±117	0.748±0.046	125±64	402±208 (10)	602±150 (10)	859±145 (9)	425±160
Mamba	5	240±102	0.719±0.062	195±102	429±191 (10)	596±136 (10)	805±108 (9)	1195±687
Mamba	6	298±167	0.706±0.052	212±122	405±190 (10)	557±197 (10)	736±170 (9)	1420±632
Mamba	7	328±116	0.662±0.107	246±112	332±142 (10)	489±131 (10)	727±124 (10)	1657±947
Mamba	8	356±142	0.671±0.029	304±119	380±158 (10)	514±144 (10)	699±167 (10)	2340±806
Mamba	9	359±135	0.682±0.054	298±115	439±140 (10)	536±161 (10)	663±102 (9)	2974±1394
Mamba	10	368±164	0.692±0.032	305±154	391±234 (10)	485±99 (9)	658±125 (9)	2829±1290
Mamba	11	321±148	0.636±0.048	280±137	415±154 (10)	561±153 (10)	720±145 (9)	3515±1592
Mamba	12	335±148	0.637±0.055	285±148	425±162 (10)	564±178 (10)	687±135 (9)	4060±1694
Mamba	13	260±158	0.579±0.121	213±139	505±168 (10)	602±141 (9)	744±130 (8)	3691±1790
Mamba	14	290±120	0.608±0.047	235±89	463±213 (10)	583±150 (10)	765±127 (10)	4505±1968
Mamba	15	343±157	0.621±0.069	317±149	367±140 (10)	534±159 (10)	706±166 (10)	4196±1064
Mamba	16	320±214	0.61±0.095	293±199	450±210 (10)	560±241 (9)	602±141 (7)	5035±1995
Mamba	17	233±131	0.611±0.059	219±131	552±165 (10)	665±147 (10)	806±130 (9)	3728±1946
Mamba	18	270±205	0.617±0.061	256±200	516±155 (10)	628±191 (10)	705±201 (7)	5378±2020
Mamba	19	168±164	0.632±0.070	139±121	468±221 (8)	604±233 (8)	805±193 (6)	4740±2181
Mamba	20	256±196	0.539±0.190	245±192	462±225 (9)	531±233 (8)	642±156 (7)	4476±2383

sample efficiency (we note that the explainability aspect is still applicable). In the original work, a batch size of 64 was used and a Patience of 5. Under these parameters, the earliest that Beam Enumeration can execute is 320/1000 oracle calls, which is almost 1/3 the budget already. Moreover, Beam Enumeration decreases diversity and decreasing batch size and increasing augmentation rounds also decreases diversity. *Too much* decrease in diversity may be detrimental even with oracle caching. In this subsection, we systematically study the effect of Beam Enumeration when used in conjunction with decreasing batch size and augmentation rounds in a series of hypotheses.

# Based on observations from batch size and augmentation rounds grid-searches, the following design decisions were made in this subsection:

- Augmentation rounds capped at 5 as diversity generally decreases more substantially past this point. Beam Enumeration itself will decrease diversity, so this is a preemptive measure against detrimental diversity-induced mode collapse.
- Investigate batch sizes of 64 and 32. Since Beam Enumeration executes on improved reward over successive epochs, lower batch sizes would likely increase performance variance too much.
- 3. Focus only on RNN model as experiments will be the fastest (less repeated SMILES). If benefits are observed, move to Decoder and Mamba models. For clarity, repeated SMILES are not detrimental, as we have shown in the previous subsections but they add some wall time (this is insignificant when compared to expensive oracles).
- 4. Beam Enumeration can pool improbable substructures. There is a Patience Limit denoting the number epochs permitted where the entire generated batch is filtered. This limit was 100,000 in this work. This does not add that much wall time and surpassing the limit is not indicative of the experiment failing. However, we enforce this upper bound in case it occurs (seldom) to manage wall times since we are performing grid searches.
- 5. Use Minimum Structure Size = 15, unless otherwise stated. Enforcing larger substructure extraction was found to improve sample efficiency in the original work <sup>22</sup>

# B.4.1 Hypothesis 1

Beam Enumeration's Patience parameter is dependent on the mean reward of the sampled batch. With lower batch sizes, variance increases, such that executing Beam Enumeration may be *too variable*.

Proposed solution. Increase Beam Enumeration's default Patience (5) to mitigate lower batch size variance. We note that increasing Patience means that more of the oracle budget needs to be consumed before Beam Enumeration executes for the first time. First explore Batch sizes = [64, 32].

**Observations.** Across batch sizes = [64, 32] and all Patience = [5, 6, 7, 8, 9, 10], sample efficiency does not improve in a statistically significant manner (Tables 17 and 18). Using Beam Enumeration also leads to notably higher variance and decreased diversity.

Table 17: Beam Enumeration batch size 64 with Structure and Minimum Size 15. Filter Limit is the number of times that no SMILES contained the pool substructure in 100,000 generation epochs. Patience N/A indicates just Augmented Memory and no Beam Enumeration.

Patience	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats	Filter Limit
N/A	0	0±0	_	0±0	584±251 (5)	Failed	Failed	1±1	N/A
N/A	2	15±9	0.775±0.073	15±9	644±173 (10)	941±58 (8)	Failed	$0\pm0$	N/A
N/A	3	33±42	0.788±0.043	32±40	613±96 (10)	927±128 (9)	993±0(1)	$0\pm0$	N/A
N/A	4	32±16	0.813±0.024	31±16	527±198 (10)	880±90 (10)	Failed	$0\pm0$	N/A
N/A	5	40±14	0.812±0.023	39±13	459±177 (10)	862±68 (10)	Failed	0±0	N/A
5	0	2±2	_	2±2	687±232 (7)	Failed	Failed	17±21	0
5	2	29±68	0.688±0.044	22±48	555±185 (8)	887±182 (4)	866±0(1)	15±27	1
5	3	110±75	0.754±0.024	81±52	488±79 (10)	711±99 (10)	902±79 (4)	20±21	0
5	4	86±82	0.702±0.045	58±53	504±205 (10)	739±193 (9)	912±76 (3)	14±15	0
5	5	94±41	0.745±0.027	68±30	436±167 (10)	739±88 (10)	970±30 (4)	15±17	0
6	0	2±3	_	2±2	581±205 (7)	958±0 (1)	Failed	25±29	0
6	2	20±20	0.619±0.168	16±15	659±226 (10)	809±27 (4)	Failed	9±10	0
6	3	82±84	0.73±0.039	52±44	520±84 (10)	777±134 (10)	863±131	19±26	0
6	4	83±91	0.723±0.074	62±62	508±233 (9)	737±130 (8)	874±93	19±21	0
6	5	84±52	0.693±0.049	54±30	449±169 (10)	771±131 (10)	973±44	38±56	0
7	0	2±2	_	2±2	599±238 (6)	Failed	Failed	15±17	0
7	2	40±43	0.661±0.161	32±34	579±137 (10)	836±112 (8)	1000±28 (2)	9±10	0
7	3	121±120	0.719±0.038	80±69	546±66 (10)	735±131 (10)	803±75 (3)	27±30	0
7	4	69±64	0.701±0.098	45±39	560±249 (10)	726±84 (7)	941±55 (2)	12±18	0
7	5	61±34	0.735±0.055	43±21	467±188 (10)	796±77 (10)	1026±4 (2)	11±15	0
8	0	1±2	_	1±1	556±225 (5)	1010±0(1)	Failed	24±32	0
8	2	80±90	0.697±0.074	51±60	604±153 (10)	775±119 (8)	882±94 (3)	8±11	0
8	3	79±86	0.714±0.028	58±67	579±88 (10)	769±131 (9)	920±139 (3)	7±6	0
8	4	68±85	0.671±0.044	45±55	537±202 (10)	786±115 (6)	902±49 (3)	20±23	0
8	5	88±61	0.711±0.098	64±45	459±184 (10)	757±118 (9)	960±33 (4)	15±27	0
9	0	1±1	_	1±1	564±226 (5)	Failed	Failed	11±11	0
9	2	49±53	0.7±0.119	36±34	620±171 (10)	826±115 (8)	953±12 (2)	2±4	0
9	3	87±81	$0.739 \pm 0.034$	53±38	599±92 (10)	787±100 (10)	935±122 (3)	9±11	0
9	4	65±49	0.688±0.08	48±41	518±187 (10)	798±88 (10)	910±0(1)	11±17	0
9	5	99±84	0.694±0.098	60±51	459±180 (10)	774±80 (10)	907±93 (3)	19±27	0
10	0	1±1	_	1±1	564±226 (5)	Failed	Failed	11±11	0
10	2	49±53	0.7±0.119	36±34	620±171 (10)	826±115 (8)	953±12 (2)	2±4	0
10	3	87±81	0.739±0.034	53±38	599±92 (10)	787±100 (10)	935±122 (3)	9±11	0
10	4	65±49	0.688±0.08	48±41	518±187 (10)	798±88 (10)	910±0(1)	11±17	0
10	5	99±84	0.694±0.098	60±51	459±180 (10)	774±80 (10)	907±93 (3)	19±27	0

# B.4.2 Hypothesis 2

The use of "Structure" substructure is too biased when operating in an already biased environment: increasing augmentation rounds and under a low oracle budget.

**Proposed solution.** Investigate "Scaffold" substructure which is less biased.

Observations. Across batch sizes = [64, 32] and all Patience = [5, 6, 7, 8, 9, 10], sample efficiency does not improve in a statistically significant manner (Tables 19 and 20). Variance decreases relative to "Structure" which is in agreement with the hypothesis that "Structure" is more biased.

# B.4.3 Hypothesis 3

In the original Beam Enumeration<sup>22</sup> work, enforcing a Structure Minimum Size for extracted substructures improves sample efficiency across all hyperparameter combinations (and is statistically significant). The results so far suggest that this observation does not hold when optimizing under a particularly low oracle budget (1000 calls). Thus far, experiments were aimed at mitigating the Beam Enumeration bias either by tuning the Patience parameter or by changing the Substructure Type. Another method to mitigate bias is by not enforcing a Structure Minimum Size. In this scenario, Scaffold substructure should be used as Structure substructure tends to extract small functional groups (as observed in the original work).

**Proposed solution.** Investigate "Scaffold" substructure without enforcing Structure Minimum Size.

**Observations.** Across batch sizes = [64, 32] and all Patience = [5, 6, 7, 8, 9, 10], sample efficiency *sometimes* improves (Tables 21 and 22). Variance is also manageable but the performance improve-

Table 18: Beam Enumeration batch size 32 with Structure and Minimum Size 15. Filter Limit is the number of times that no SMILES contained the pool substructure in 100,000 generation epochs. Patience N/A indicates just Augmented Memory and no Beam Enumeration.

Patience	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats	Filter Limit
N/A	0	0±0	_	0±0	798±101 (5)	Failed	Failed	1±1	N/A
N/A	2	43±25	0.825±0.029	42±24	608±151 (10)	844±90 (9)	Failed	0±0	N/A
N/A	3	52±34	0.81±0.059	51±32	522±141 (10)	789±100 (9)	1018±0(2)	0±1	N/A
N/A	4	87±33	0.82±0.018	83±31	466±120 (10)	740±77 (10)	987±30 (4)	1±3	N/A
N/A	5	98±57	0.817±0.027	89±50	408±184 (10)	714±136 (10)	915±20 (4)	1±2	N/A
5	0	2±4	0.611±0.074	2±3	776±155 (4)	983±0 (1)	Failed	43±30	0
5	2	18±27	0.666±0.077	15±19	705±173 (8)	857±104 (4)	Failed	9±9	0
5	3	26±20	0.652±0.076	19±11	618±88 (10)	850±108 (7)	Failed	16±18	0
5	4	65±64	0.695±0.092	54±53	604±214 (10)	742±124 (6)	936±55 (3)	65±90	0
5	5	99±110	0.713±0.046	66±61	452±216 (10)	741±173 (9)	870±146 (4)	64±56	0
6	0	2±5	0.655±0.051	2±4	614±213 (4)	836±0(1)	Failed	39±27	0
6	2	36±49	0.691±0.096	32±47	625±188 (9)	834±139 (7)	943±31 (2)	9±9	0
6	3	60±58	0.662±0.124	47±53	574±148 (10)	811±146 (10)	895±81 (2)	93±220	0
6	4	67±52	0.654±0.185	54±43	592±214 (10)	740±133 (8)	934±50 (3)	114±154	0
6	5	66±70	0.68±0.059	50±44	530±209 (10)	822±141 (9)	933±69 (3)	65±70	0
7	0	1±2	_	1±2	686±161 (6)	Failed	Failed	83±78	0
7	2	49±60	0.699±0.101	41±56	601±156 (10)	821±152 (8)	923±93 (2)	18±20	0
7	3	47±46	0.67±0.107	37±36	623±198 (9)	810±161 (8)	994±16 (3)	20±21	0
7	4	41±45	$0.686\pm0.058$	33±42	588±81 (9)	838±94 (9)	905±0(1)	53±43	0
7	5	76±76	0.698±0.111	66±74	531±210 (10)	776±128 (8)	866±69 (2)	126±325	0
8	0	16±37	_	14±33	749±210 (8)	668±194 (2)	949±0(1)	109±163	0
8	2	33±48	0.691±0.049	24±33	692±144 (9)	856±142 (6)	974±35 (2)	15±18	0
8	3	50±30	0.675±0.068	40±22	636±109 (10)	803±84 (8)	Failed	39±49	0
8	4	104±104	0.73±0.056	84±96	406±128 (10)	696±149 (9)	879±141 (4)	30±36	0
8	5	42±30	0.7±0.051	32±18	506±186 (10)	848±95 (10)	974±0 (1)	30±45	0
9	0	7±12	_	6±10	713±201 (7)	848±1 (2)	Failed	68±50	0
9	2	36±34	$0.686\pm0.052$	28±28	559±138 (10)	812±96 (7)	1015±0(1)	29±28	0
9	3	81±89	$0.668\pm0.102$	52±52	598±186 (10)	732±159 (7)	826±49 (3)	23±19	0
9	4	158±103	$0.723\pm0.041$	104±63	432±104 (10)	639±115 (10)	868±106 (7)	60±78	0
9	5	91±66	0.707±0.036	57±35	453±194 (10)	763±131 (10)	928±65 (4)	40±29	0
10	0	2±3	_	2±3	768±107 (5)	1003±0 (1)	Failed	93±97	0
10	2	55±54	0.722±0.027	44±40	559±156 (10)	807±149 (10)	836±0 (1)	26±39	0
10	3	86±46	0.705±0.063	67±36	478±143 (10)	678±114 (9)	962±33 (4)	41±50	0
10	4	99±77	0.705±0.048	63±43	474±162 (10)	693±91 (9)	944±113 (4)	58±86	0
10	5	110±100	0.715±0.039	80±78	430±164 (10)	750±142 (10)	881±107 (4)	57±55	0

ments, when observed, is much less than with lower batch size and higher augmentation rounds (for instance Mamba batch size 16 and augmentation rounds 10).

**Conclusions.** Based on the grid-search results, Beam Enumeration can *sometimes* improve sample efficiency when using "Scaffold" structure and without enforcing Structure Minimum Size. However, the improvements are minor, such that it would be better to use small batch sizes with high augmentation rounds. Thus, we do not further experiment with Beam Enumeration in this work.

# B.5 Hallucinated Memory: Is it beneficial to allocate a portion of the oracle budget to hallucination?

In this section, we investigate coupling GraphGA <sup>63</sup> to Saturn. GraphGA in itself a sample-efficient generative algorithm <sup>20</sup> and was recently used in the GEAM model proposed by Lee et al. <sup>13</sup> which achieves impressive MPO performance. Previously work <sup>80</sup> found that coupling a GA in RL can encourage diverse sampling. In the previous sections, we have identified Mamba with batch size 16 and 10 augmentation rounds as the best hyperparameters so far. The improved sample efficiency comes at a trade-off in diversity. The objective in the experiments to follow is to investigate whether allocating a portion of the oracle budget to GraphGA generation (which we call "hallucinating") is beneficial in recovering diversity while maintaining sample efficiency.

Before presenting the grid-search results, we describe the GraphGA integration further. GraphGA is only activated when the replay buffer is full (100 SMILES). Once full, at every epoch thereafter, the replay buffer itself is treated as the parent population to generate new SMILES. These new SMILES are then concatenated with the sampled batch (16 SMILES) and used to update the agent. Importantly, these hallucinated SMILES are also deposited into the replay buffer (if they possess higher reward).

Table 19: Beam Enumeration batch size 64 with Scaffold and Minimum Size 15. Filter Limit is the number of times that no SMILES contained the pool substructure in 100,000 generation epochs. Patience N/A indicates just Augmented Memory and no Beam Enumeration.

Patience	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats	Filter Limit
N/A	0	0±0	_	0±0	584±251 (5)	Failed	Failed	1±1	N/A
N/A	2	15±9	0.775±0.073	15±9	644±173 (10)	941±58 (8)	Failed	$0\pm0$	N/A
N/A	3	33±42	0.788±0.043	32±40	613±96 (10)	927±128 (9)	993±0(1)	$0\pm0$	N/A
N/A	4	32±16	0.813±0.024	31±16	527±198 (10)	880±90 (10)	Failed	0±0	N/A
N/A	5	40±14	0.812±0.023	39±13	459±177 (10)	862±68 (10)	Failed	0±0	N/A
5	0	5±17	0.726±0.0	5±15	653±275 (3)	819±0 (1)	Failed	48±31	0
5	2	14±22	0.616±0.182	13±21	635±226 (7)	850±131 (3)	Failed	36±29	0
5 5 5	3	21±26	0.675±0.116	18±22	647±198 (8)	852±88 (5)	Failed	19±26	0
5	4	20±30	0.6±0.122	18±26	592±262 (9)	869±108 (4)	1038±0(1)	28±19	0
5	5	33±27	0.692±0.082	29±25	506±208 (10)	875±101 (8)	Failed	33±37	0
6	0	$0\pm1$	0.399±0.0	0±0	433±98 (4)	Failed	Failed	98±99	0
6	2	9±16	0.656±0.072	7±13	713±237 (8)	864±82 (2)	Failed	$30\pm25$	0
6	3	16±19	0.645±0.072	14±18	662±152 (8)	905±103 (5)	Failed	27±30	0
6	4	15±23	0.644±0.069	14±22	466±185 (8)	884±137 (4)	Failed	23±16	0
6	5	24±28	0.599±0.139	21±22	583±293 (10)	849±83 (5)	1014±0 (1)	35±38	0
7	0	0±1	_	0±1	459±139 (4)	Failed	Failed	82±47	0
7	2	10±10	$0.64\pm0.072$	9±10	666±180 (9)	911±76 (3)	Failed	37±59	0
7	3	27±31	0.659±0.119	23±23	648±153 (9)	880±122 (7)	1041±0(1)	11±8	0
7	4	20±19	0.634±0.125	19±18	575±249 (10)	853±72 (5)	Failed	46±59	0
7	5	14±13	0.676±0.096	12±10	519±267 (10)	932±75 (6)	Failed	24±32	0
8	0	0±0	_	0±0	383±53 (3)	Failed	Failed	36±23	0
8	2	10±13	0.665±0.131	10±12	654±201 (8)	910±85 (4)	Failed	15±19	0
8	3	30±48	0.693±0.031	29±46	624±164 (9)	863±129 (6)	901±0(1)	24±21	0
8	4	29±43	0.667±0.095	23±30	571±268 (9)	745±98 (4)	981±0(1)	20±26	0
8	5	40±47	0.665±0.093	35±45	450±168 (10)	879±95 (9)	920±0 (1)	43±74	0
9	0	0±0	_	0±0	500±207 (4)	Failed	Failed	31±29	0
9	2	20±36	0.683±0.055	19±36	683±226 (9)	825±84 (3)	1005±0(1)	8±9	0
9	3	41±34	0.675±0.08	34±28	654±155 (10)	849±134 (8)	Failed	25±22	0
9	4	16±14	0.647±0.093	13±11	573±240 (10)	917±39 (5)	Failed	10±11	0
9	5	39±24	0.707±0.083	34±22	456±172 (10)	829±67 (9)	Failed	8±9	0
10	0	3±8	_	3±7	519±171 (5)	851±0 (1)	Failed	16±26	0
10	2	16±19	0.674±0.07	13±15	599±144 (9)	905±95 (5)	Failed	17±20	0
10	3	32±38	0.703±0.074	26±27	621±107 (10)	861±129 (8)	961±0(1)	5±7	0
10	4	18±15	0.682±0.087	16±15	529±202 (10)	876±81 (7)	Failed	5±8	0
10	5	37±31	0.711±0.057	30±20	456±172 (10)	829±68 (8)	996±0(1)	23±42	0

Finally, 100 SMILES are hallucinated and either 5 or 10 are selected. The selection criteria are **Random** or **Tanimoto Distance**. Random selects at random while Tanimoto Distance selects via maximum fingerprint *dissimilarity* to the replay buffer. Our rationale is that dissimilar new SMILES will help encourage diversity since Augmented Memory heavily biases towards the replay buffer SMILES.

#### 1089 The grid-search investigated the following hyperparameter settings:

1. Fix Mamba with batch size 16

- 2. Augmentation Rounds = [5,20]
- 3. GA with Random and Tanimoto Distance selection criterion
- 4. Select 5 or 10 hallucinations at every epoch

The reason we increased the augmentation rounds back to 20 in our grid-search is because if indeed the GA recovers diversity, then the "augmentation tolerability" of Saturn would probably be increased. Higher augmentation rounds lead to more repeated SMILES precisely due to overfitting. If new high reward SMILES *refresh* the replay buffer, Saturn may be more tolerable to higher augmentation rounds to potentially further improve sample efficiency. The results of the grid-search are presented in Tables 23 and 24.

**Observations.** The results show that coupling a GA to the replay buffer does not improve sample efficiency. However, we make several interesting observations. Firstly, the number of repeated SMILES *notably* drops and IntDiv1<sup>71</sup> recovers. This is in agreement with our hypothesis and previous work <sup>80</sup> that coupling a GA to RL can recover diversity. Secondly, hallucinating SMILES does indeed lead to some replacement of the replay buffer, and hence, these SMILES are necessarily

Table 20: Beam Enumeration batch size 32 with Scaffold and Minimum Size 15. Filter Limit is the number of times that no SMILES contained the pool substructure in 100,000 generation epochs. Patience N/A indicates just Augmented Memory and no Beam Enumeration.

Patience	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats	Filter Limit
N/A	0	0±0	_	0±0	798±101 (5)	Failed	Failed	1±1	N/A
N/A	2	43±25	0.825±0.029	42±24	608±151 (10)	844±90 (9)	Failed	0±0	N/A
N/A	3	52±34	0.81±0.059	51±32	522±141 (10)	789±100 (9)	1018±0(2)	0±1	N/A
N/A	4	87±33	0.82±0.018	83±31	466±120 (10)	740±77 (10)	987±30 (4)	1±3	N/A
N/A	5	98±57	0.817±0.027	89±50	408±184 (10)	714±136 (10)	915±20 (4)	1±2	N/A
5	0	0±0	_	0±0	852±141 (2)	Failed	Failed	119±78	0
5	2	25±38	0.65±0.109	23±35	698±191 (8)	779±127 (4)	959±0(1)	57±67	0
5	3	33±59	0.629±0.073	26±44	636±148 (8)	867±133 (6)	871±0(1)	88±123	1
5	4	57±68	0.666±0.032	44±51	648±163 (9)	834±128 (7)	952±70 (3)	118±104	0
5	5	50±69	0.649±0.038	33±39	498±268 (9)	855±170 (8)	890±3 (2)	89±46	0
6	0	2±6	_	2±6	788±161 (3)	840±0 (1)	Failed	174±112	0
6	2	25±59	0.618±0.148	16±36	672±240 (7)	694±238 (3)	706±0 (1)	53±55	1
6	3	35±47	0.667±0.119	27±35	702±189 (8)	789±93 (5)	974±0 (2)	52±43	0
6	4	46±66	0.653±0.068	39±56	656±127 (9)	831±144 (6)	945±67 (2)	135±206	0
6	5	57±76	0.584±0.157	45±59	571±274 (8)	668±83 (4)	907±7 (3)	101±113	0
7	0	14±27	0.551±0.116	10±17	663±109 (5)	814±130 (3)	Failed	106±58	0
7	2	19±41	0.657±0.121	12±24	660±127 (6)	894±136 (5)	929±0(1)	34±23	0
7	3	38±51	0.636±0.115	28±30	650±161 (10)	812±131 (6)	863±0(1)	45±33	0
7	4	36±36	0.652±0.109	26±21	700±151 (10)	811±76 (7)	981±0(1)	67±49	0
7	5	46±45	0.608±0.108	39±40	485±204 (9)	810±50 (6)	991±5 (2)	237±244	0
8	0	0±0	_	0±0	794±302 (4)	Failed	Failed	149±100	0
8	2	34±45	0.625±0.105	30±39	696±175 (9)	777±105 (5)	901±0(1)	57±46	0
8	3	53±77	0.543±0.174	42±61	652±213 (9)	715±141 (5)	836±6 (2)	57±87	1
8	4	30±53	0.631±0.092	24±39	684±235 (9)	781±165 (3)	957±51 (2)	54±43	0
8	5	90±101	0.632±0.124	70±74	556±248 (9)	706±127 (6)	879±78 (4)	179±158	0
9	0	0±0	_	0±0	733±157 (3)	Failed	Failed	175±142	0
9	2	20±37	0.61±0.124	15±25	643±237 (8)	849±152 (4)	967±0(1)	61±69	0
9	3	28±25	0.639±0.09	23±20	661±121 (10)	819±78 (6)	Failed	53±60	0
9	4	67±63	0.66±0.105	55±56	605±203 (9)	783±126 (8)	906±58 (2)	92±65	0
9	5	55±73	0.618±0.13	36±41	513±225 (9)	779±149 (6)	877±74 (2)	150±206	0
10	0	2±5		1±3	835±154 (4)	890±0 (1)	Failed	93±68	0
10	2	5±4	_	4±3	680±196 (8)	960±0 (1)	Failed	58±52	0
10	3	32±48	0.636±0.143	31±47	572±171 (10)	880±130 (7)	900±0(1)	30±36	0
10	4	44±32	0.693±0.059	34±26	503±195 (10)	811±126 (9)	965±0(1)	107±125	0
10	5	51±55	0.581±0.206	36±37	584±317 (9)	712±88 (5)	949±34 (2)	156±239	1

are high reward. Thirdly, rarely are the hallucinated SMILES the best in the buffer. Finally, we note that hallucinated SMILES are generated off-policy and agent updates may be more meaningful with importance sampling <sup>111</sup>, which we did not explore this this work.

# **B.6** Saturn: Final Hyperparameters

1105

1108

1113

The most sample-efficient hyperparameter settings, on average, are: **Mamba with batch size 16**and 10 augmentation rounds. The results in the immediate previous section shows that the GA can
recover diversity, which can be a useful setting that can easily be activated on and off depending on
the oracle setting.

# C Mechanism of Augmented Memory and Mamba

In this subsection, we show additional results supporting our statement on Augmented Memory's <sup>21</sup> mechanism: Augmented Memory squeezes the likelihood of generating the Buffer *molecules* such that it becomes probable to generate *some* SMILES representation of them. In the main text, the experiment to show likelihood squeezing was as follows: starting from the pre-trained Mamba model, generate molecules until the Buffer is full and then save the agent state before and after Augmented Memory. Every augmented Buffer SMILES was also saved. This experiment isolates the effect of Augmented Memory on a *clean* pre-trained model.

The first set of additional results we show is the same experiment but we first allow the agent 500 oracle calls of optimization on the test experiment. Our intention is to show that later in the run, Augmented Memory still makes generating the Buffer *molecules* more likely (Fig. C3). There are

Table 21: Beam Enumeration batch size 64 with Scaffold and no Minimum Size enforced. Filter Limit is the number of times that no SMILES contained the pool substructure in 100,000 generation epochs. Patience N/A indicates just Augmented Memory and no Beam Enumeration.

Patience	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats	Filter Limit
N/A	0	0±0	_	0±0	584±251 (5)	Failed	Failed	1±1	0
N/A	2	15±9	0.775±0.073	15±9	644±173 (10)	941±58 (8)	Failed	0±0	0
N/A	3	33±42	0.788±0.043	32±40	613±96 (10)	927±128 (9)	993±0(1)	0±0	0
N/A	4	32±16	0.813±0.024	31±16	527±198 (10)	880±90 (10)	Failed	0±0	0
N/A	5	40±14	0.812±0.023	39±13	459±177 (10)	862±68 (10)	Failed	0±0	0
		0±0		0±0		Failed	Failed		
5	0		0.744+0.069		307±0 (1)			0±0	0
5	2	15±12	0.744±0.068	14±11	678±227 (10)	930±70 (5)	Failed	0±0	0
5	3	38±14	0.791±0.026	37±14	552±70 (10)	824±44 (9)	Failed	0±0	0
5	4	43±45	0.791±0.021	42±43	516±230 (10)	839±132 (9)	918±0 (1)	0±0	0
5	5	55±33	0.77±0.073	50±30	467±197 (10)	811±81 (9)	961±0 (1)	0±1	0
6	0	0±0		0±0	594±268 (5)	Failed	Failed	0±0	0
6	2	28±23	0.752±0.053	26±21	671±190 (10)	880±72 (6)	Failed	0±0	0
6	3	44±28	$0.782 \pm 0.032$	42±24	584±120 (10)	832±64 (9)	1006±0(1)	0±0	0
6	4	41±37	0.778±0.028	39±36	571±241 (10)	874±118 (9)	959±0(1)	0±0	0
6	5	54±21	0.794±0.025	49±17	453±169 (10)	827±72 (10)	Failed	0±0	0
7	0	0±0	_	0±0	567±234 (5)	Failed	Failed	0±1	0
7	2	27±13	0.778±0.072	27±13	603±148 (10)	880±80 (9)	Failed	0±0	0
7	3	47±33	0.797±0.027	44±30	586±73 (10)	859±113 (10)	1035±1 (2)	0±0	0
7	4	48±23	0.799±0.017	45±20	498±176 (10)	828±87 (10)	Failed	0±0	0
7	5	51±23	0.793±0.023	48±21	463±190 (10)	854±72 (10)	Failed	0±0	0
8	0	0±0	_	0±0	383±53 (3)	Failed	Failed	0±0	0
8	2	20±12	0.755±0.072	20±12	637±153 (10)	929±62 (8)	Failed	0±0	0
8	3	39±32	0.793±0.021	38±31	593±85 (10)	882±111 (10)	962±0 (1)	0±0	0
8	4	47±30	0.793±0.024	45±29	544±208 (10)	873±75 (10)	1013±0 (1)	0±0	0
8	5	69±28	0.803±0.019	64±22	446±162 (10)	789±73 (10)	991±0 (1)	0±0	0
						. ,			
9	0	0±0		0±0	656±281 (6)	Failed	Failed	0±0	0
9	2	16±10	0.761±0.041	16±10	640±166 (10)	946±48 (6)	Failed	0±0	0
9	3	52±60	0.798±0.021	49±55	619±106 (10)	847±107 (10)	847±0 (1)	0±0	0
9	4	50±25	0.802±0.01	48±22	505±177 (10)	846±79 (10)	1004±0 (1)	0±0	0
9	5	54±26	0.792±0.024	50±24	450±165 (10)	809±55 (9)	Failed	0±0	0
10	0	$0\pm0$	_	0±0	636±260 (6)	Failed	Failed	0±0	0
10	2	21±17	0.739±0.091	21±17	643±178 (10)	920±78 (8)	Failed	$0\pm0$	0
10	3	46±48	0.791±0.024	43±43	613±99 (10)	853±115 (9)	899±0(1)	0±0	0
10	4	44±35	0.783±0.041	42±33	541±222 (10)	858±89 (9)	990±0(1)	$0\pm0$	0
10	5	48±18	0.792±0.024	45±15	456±173 (10)	853±50 (10)	Failed	0±0	0

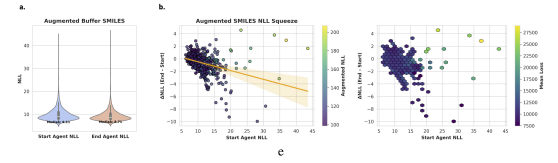


Figure C3: Mamba (batch size 16, augmentation rounds 10) after running for 500 oracle calls of the illustrative example and isolating the effect of Augmented Memory. **a.** Augmented Memory makes the likelihood of generating SMILES in the Buffer more likely. **b.** Augmented forms of the Buffer SMILES become more likely, but still regularized by the prior.

Table 22: Beam Enumeration batch size 32 with Scaffold and no Minimum Size enforced. Filter Limit is the number of times that no SMILES contained the pool substructure in 100,000 generation epochs. Patience N/A indicates just Augmented Memory and no Beam Enumeration.

Patience	Aug. Rounds	Yield	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Repeats	Filter Limit
N/A	0	0±0	_	0±0	798±101 (5)	Failed	Failed	1±1	0
N/A	2	43±25	0.825±0.029	42±24	608±151 (10)	844±90 (9)	Failed	$0\pm0$	0
N/A	3	52±34	0.81±0.059	51±32	522±141 (10)	789±100 (9)	1018±0 (2)	0±1	0
N/A	4	87±33	0.82±0.018	83±31	466±120 (10)	740±77 (10)	987±30 (4)	1±3	0
N/A	5	98±57	0.817±0.027	89±50	408±184 (10)	714±136 (10)	915±20 (4)	1±2	0
5	0	0±1	_	0±1	783±134 (3)	Failed	Failed	0±1	0
5	2	38±28	0.796±0.03	35±25	504±111 (9)	828±115 (9)	Failed	1±1	0
5	3	63±44	0.762±0.073	57±38	593±170 (10)	763±82 (8)	988±29 (3)	1±2	0
5	4	87±57	0.779±0.038	72±43	540±145 (10)	764±139 (10)	958±48 (5)	2±4	0
5	5	106±61	0.784±0.031	84±41	467±187 (10)	718±109 (10)	960±41 (6)	1±2	0
6	0	1±3	_	1±3	837±135 (3)	998±0 (1)	Failed	2±2	0
6	2	40±33	0.761±0.078	36±29	609±149 (9)	811±64 (7)	1014±0(1)	1±2	0
6	3	49±23	0.796±0.03	46±21	585±104 (10)	839±101 (10)	Failed	1±2	0
6	4	57±41	0.783±0.031	53±37	557±187 (10)	771±82 (8)	987±10 (3)	1±2	0
6	5	106±85	0.776±0.05	85±55	508±241 (10)	718±151 (9)	927±94 (5)	3±6	0
7	0	0±0	_	0±0	741±222 (5)	Failed	Failed	1±1	0
7	2	43±27	0.79±0.037	41±26	631±182 (10)	799±77 (8)	Failed	$0\pm0$	0
7	3	84±67	0.79±0.021	73±56	578±188 (10)	781±117 (9)	937±42 (4)	0±1	0
7	4	74±43	0.785±0.041	69±37	574±149 (10)	789±111 (10)	948±39 (2)	1±3	0
7	5	121±52	0.786±0.033	105±39	422±155 (10)	673±90 (10)	898±52 (5)	4±9	0
8	0	3±5	_	3±5	683±213 (5)	882±0 (1)	Failed	2±3	0
8	2	44±39	0.713±0.166	40±30	629±177 (10)	778±97 (7)	995±0(1)	1±4	0
8	3	69±43	0.794±0.039	65±40	530±183 (10)	778±104 (9)	975±8 (3)	0±2	0
8	4	75±39	0.795±0.033	66±30	547±142 (10)	770±118 (10)	981±29 (3)	1±1	0
8	5	103±55	0.761±0.091	90±49	488±221 (10)	693±142 (9)	961±39 (7)	4±5	0
9	0	2±4	_	2±4	805±127 (4)	915±0 (1)	Failed	1±1	0
9	2	41±23	$0.79\pm0.022$	40±22	572±132 (10)	839±95 (10)	Failed	0±0	0
9	3	59±34	0.81±0.021	54±31	520±110 (9)	778±68 (9)	993±0(1)	0±1	0
9	4	101±60	0.799±0.025	89±45	515±142 (10)	725±104 (10)	944±91 (4)	1±1	0
9	5	128±61	0.792±0.022	102±41	425±179 (10)	684±93 (10)	919±51 (6)	2±2	0
10	0	0±1	_	0±1	822±160 (4)	Failed	Failed	1±1	0
10	2	53±45	0.795±0.025	49±44	515±129 (9)	793±106 (9)	973±30 (2)	2±5	0
10	3	86±63	0.759±0.119	73±46	553±179 (10)	720±62 (8)	956±69 (4)	0±1	0
10	4	89±35	0.794±0.034	77±26	464±132 (10)	743±51 (10)	984±27 (4)	3±5	0
10	5	123±58	0.795±0.031	105±44	434±177 (10)	704±102 (10)	949±59 (8)	2±2	0
	-						2 12 = 27 (0)		-

cases when a large loss magnitude does not make the sequence more likely to be generated. This could occur for instance when the likelihood under the prior is extremely low (large NLL) where the intended behavior is actually to regress the agent back towards the prior. In these cases, the large loss could make the update less stable for the parameter updates.

Next, the main text results showed that Mamba (batch size 16, augmentation rounds 10) exhibits "hop-and-locally-explore" behavior but what about RNN (batch size 16, augmentation rounds 10)? We show that the RNN model also begins to exhibit this behavior but to a lesser extent (Fig. C4), in agreement with the enhanced likelihood convergence observed for Mamba (Appendix B.1).

We now focus on Mamba (batch size 16, augmentation rounds 10) and present additional results to qualitatively and quantitatively demonstrate "hop-and-locally-explore" behavior. Firstly, we supplement the main text Fig. 2e. The figure shows the intra- and inter-chunk similarities across chunks of generated molecules. Specifically, the test experiment was run with an oracle budget of 3,000 and this generated set is chunked. To provide a more granular inspection into the generative behavior, we chunk this set into 30 chunks (each 100 SMILES) instead of 10 chunks (each 300 SMILES) in the main text. Mamba (batch size 16, augmentation rounds) exhibits notably higher intra-chunk similarity and even inter-chunk similarity at this more granular chunking level (Fig. C5a). We further supplement these quantitative results with a qualitative inspection. Looking at **unique** molecules generated at adjacent epochs, common substructures are shared (Fig. C5b highlights), displaying a "neighborhood-like" exploration.

Table 23: Mamba batch size 16 with GraphGA <sup>63</sup> applied on the replay buffer. The hallucinated SMILES were selected at *Random*. **Hall. Yield** is the yield from GraphGA. **Buf. Replace** is the number of times a hallucinated SMILES replaced another SMILES in the buffer. This means that it was better than the top-100 SMILES generated in the run so far. **Buf. Best** is the number of times the hallucinated SMILES was better than the top-1 in the buffer.

GA Random	Aug. Rounds	Hall. Yield	Total Yield	Buffer Replace	Buffer Best	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Sampled Repeats	Hall. Repeats
5	5	9±7	54±43	91±13	2±1	0.756±0.043	45±33	538±212 (10)	812±114 (9)	989±27 (3)	58±39	5±3
5	6	21±10	88±56	92±11	3±1	0.773±0.046	68±41	457±122 (10)	729±103 (10)	936±83 (3)	57±29	6±3
5	7	11±9	57±42	90±17	3±2	0.73±0.063	49±37	619±125 (10)	795±116 (9)	988±13 (3)	122±50	6±3
5	8	14±11	63±42	95±15	3±2	0.758±0.044	49±25	574±166 (10)	793±96 (10)	916±0(1)	177±80	6±3
5	9	20±15	106±75	92±14	2±1	0.767±0.03	86±55	531±128 (10)	733±121 (10)	833±57 (3)	207±101	9±5
5	10	21±11	113±61	93±19	2±1	$0.742\pm0.04$	83±38	496±158 (10)	690±118 (10)	910±59 (5)	257±143	7±3
5	11	15±11	102±69	89±13	3±2	0.739±0.031	69±43	552±141 (10)	730±116 (10)	887±62 (4)	308±116	7±3
5	12	29±17	139±83	101±13	3±1	0.781±0.025	101±55	488±104 (10)	666±92 (10)	856±76 (5)	339±153	9±4
5	13	25±14	144±97	97±15	3±1	0.727±0.048	94±50	463±209 (10)	658±155 (10)	843±99 (6)	511±226	10±4
5	14	36±22	176±82	102±18	3±2	0.742±0.038	133±56	475±121 (10)	640±110 (10)	863±92 (8)	691±333	13±7
5	15	42±17	208±65	104±18	4±2	0.746±0.06	167±58	401±115 (10)	595±89 (10)	844±91 (10)	693±319	13±8
5	16	34±9	187±77	100±20	5±2	0.744±0.055	150±59	421±119 (10)	624±106 (10)	829±83 (8)	789±465	10±5
5	17	33±25	181±95	99±14	3±1	0.75±0.042	127±64	469±142 (10)	664±132 (10)	838±86 (8)	830±417	10±6
5	18	35±18	164±57	102±24	4±2	0.727±0.038	133±54	459±105 (10)	637±76 (10)	872±66 (8)	881±389	16±16
5	19	30±16	190±76	103±16	3±1	0.744±0.046	145±51	467±123 (10)	630±113 (10)	822±59 (8)	1072±465	12±9
5	20	44±18	247±83	96±10	3±1	0.748±0.034	185±60	380±144 (10)	566±115 (10)	761±59 (9)	1310±512	14±6
10	5	12±10	44±44	141±13	3±1	0.77±0.066	35±29	478±206 (10)	802±133 (9)	888±0(1)	24±14	8±5
10	6	16±13	44±34	139±7	4±2	0.784±0.023	37±29	534±139 (10)	812±87 (9)	936±0(1)	38±19	8±4
10	7	14±9	43±27	139±23	4±2	0.739±0.109	37±23	594±117 (10)	800±54 (9)	Failed	61±34	9±4
10	8	20±16	55±41	148±13	4±2	0.771±0.026	46±30	520±114 (10)	805±129 (10)	924±0(1)	71±30	9±4
10	9	22±18	70±51	143±19	4±2	0.753±0.04	57±42	520±174 (10)	788±149 (10)	952±44 (3)	113±58	11±7
10	10	17±16	65±63	148±19	4±2	0.714±0.104	48±37	539±183 (10)	758±141 (9)	773±0(1)	138±69	11±6
10	11	18±11	57±47	140±21	5±1	0.761±0.031	42±29	605±139 (10)	789±104 (9)	931±38 (2)	192±90	10±7
10	12	37±37	88±79	165±26	4±1	0.734±0.092	70±59	591±142 (10)	716±119 (9)	882±110 (3)	222±106	17±14
10	13	$29\pm25$	84±84	150±22	3±1	0.727±0.078	61±51	502±195 (10)	737±169 (9)	842±52 (3)	260±134	13±7
10	14	29±16	97±64	149±14	5±2	0.756±0.046	72±44	456±217 (10)	733±164 (10)	908±9 (5)	271±116	9±6
10	15	37±24	102±64	161±13	4±1	$0.759\pm0.03$	85±48	480±184 (10)	688±162 (10)	913±77 (5)	336±182	19±10
10	16	40±22	110±60	157±18	5±3	0.754±0.028	91±50	432±200 (10)	691±149 (10)	913±55 (6)	361±185	15±10
10	17	34±22	103±62	156±28	5±2	0.75±0.048	80±47	529±154 (10)	704±117 (9)	916±45 (6)	467±214	15±8
10	18	25±15	91±52	148±22	5±1	$0.745\pm0.03$	64±31	562±102 (10)	750±88 (10)	927±42 (4)	572±322	17±10
10	19	25±14	88±46	145±17	6±2	0.75±0.036	71±39	563±127 (10)	751±114 (10)	948±33 (5)	603±236	16±9
10	20	38±24	136±80	148±19	6±1	0.748±0.059	95±48	444±150 (10)	626±117 (9)	867±90 (6)	781±360	13±5

Table 24: Mamba batch size 16 with GraphGA <sup>63</sup> applied on the replay buffer. The hallucinated SMILES were selected by highest *Tanimoto Distance*. **Hall. Yield** is the yield from GraphGA. **Buf. Replace** is the number of times a hallucinated SMILES replaced another SMILES in the buffer. This means that it was better than the top-100 SMILES generated in the run so far. **Buf. Best** is the number of times the hallucinated SMILES was better than the top-1 in the buffer.

GA Random	Aug. Rounds	Hall. Yield	Total Yield	Buffer Replace	Buffer Best	IntDiv1	Scaffolds	OB 1	OB 10	OB 100	Sampled Repeats	Hall. Repeats
5	5	12±11	68±60	84±16	2±1	0.77±0.05	57±46	532±244 (10)	752±125 (8)	913±51 (3)	50±35	17±7
5	6	8±8	61±73	83±13	1±1	0.763±0.041	51±57	602±171 (10)	834±151 (10)	890±110 (2)	62±36	17±11
5	7	15±8	68±46	90±10	4±2	0.776±0.035	60±38	610±62 (10)	797±86 (10)	855±0(1)	122±59	17±8
5	8	11±8	89±61	77±13	2±1	0.765±0.031	72±45	473±120 (10)	753±116 (10)	888±42 (3)	156±84	14±8
5	9	22±17	123±86	88±8	2±1	0.757±0.049	97±66	471±187 (10)	712±164 (10)	872±96 5)	309±150	16±7
5	10	18±15	97±79	87±14	2±1	0.758±0.045	78±57	544±183 (10)	748±158 (10)	901±107 (4)	317±133	16±9
5	11	18±14	92±60	84±15	2±2	0.785±0.031	78±49	560±130 (10)	749±97 (10)	846±42 (2)	314±126	20±9
5	12	26±17	146±101	90±10	2±1	0.772±0.043	109±70	491±165 (10)	684±184 (10)	838±124 (6)	418±220	22±15
5	13	21±15	114±77	90±19	2±1	$0.74\pm0.053$	97±62	494±200 (10)	706±134 (9)	912±71 (6)	494±218	19±13
5	14	28±24	158±95	91±21	2±1	0.756±0.042	131±82	505±152 (10)	681±152 (10)	846±85 (7)	682±355	27±20
5	15	39±20	189±98	97±8	3±1	0.752±0.074	151±76	415±159 (10)	600±176 (10)	818±103 (8)	698±382	28±14
5	16	45±30	189±110	100±29	2±2	0.788±0.042	152±91	456±171 (10)	630±168 (10)	784±98 (7)	771±329	33±16
5	17	29±22	166±89	95±13	3±1	0.76±0.053	124±58	506±145 (10)	652±130 (10)	874±102 (8)	733±343	26±15
5	18	17±12	114±75	88±16	3±2	0.686±0.104	87±50	549±154 (10)	668±86 (8)	913±65 (6)	911±412	30±20
5	19	16±14	117±86	73±22	2±2	0.708±0.101	94±70	559±169 (10)	706±153 (9)	862±117 (5)	1287±520	24±23
5	20	32±16	183±72	85±17	3±2	0.752±0.072	151±60	417±161 (10)	628±111 (10)	878±102 (10)	1241±508	22±13
10	5	13±13	39±39	127±17	3±2	0.768±0.065	35±34	551±214 (9)	765±155 (7)	942±0(1)	34±15	19±8
10	6	11±10	43±34	128±17	2±1	0.76±0.064	41±32	556±156 (10)	777±99 (7)	Failed	34±20	16±8
10	7	13±8	41±28	138±12	3±2	0.767±0.066	38±27	550±140 (10)	835±106 (9)	997±0(1)	62±43	19±9
10	8	12±9	41±26	138±13	2±2	0.751±0.093	36±22	575±156 (10)	786±123 (9)	Failed	75±41	21±9
10	9	18±12	56±35	129±20	3±2	0.764±0.072	48±30	527±156 (10)	732±79 (8)	991±0(1)	117±78	19±9
10	10	10±12	42±46	133±14	3±2	0.775±0.055	32±31	660±225 (10)	797±127 (7)	870±0 (1)	158±80	15±7
10	11	10±8	39±39	124±18	3±1	0.713±0.109	32±30	626±173 (10)	828±124 (7)	964±0 (1)	181±93	30±23
10	12	16±19	63±64	139±18	3±1	0.733±0.123	53±56	534±207 (10)	731±113 (8)	897±107 (2)	236±106	29±23
10	13	20±19	67±63	140±21	3±2	0.732±0.117	50±41	542±228 (9)	746±139 (8)	902±38 (3)	300±150	30±19
10	14	15±13	61±50	128±21	2±1	0.714±0.114	49±41	589±175 (10)	770±102 (8)	924±22 (2)	365±210	26±15
10	15	28±25	80±71	144±22	5±1	0.762±0.033	68±58	599±160 (10)	741±129 (8)	925±100 (4)	366±228	32±19
10	16	30±28	89±77	152±28	5±2	0.765±0.07	74±63	563±186 (10)	719±167 (9)	832±34 (3)	376±188	35±24
10	17	30±25	101±80	147±16	3±1	0.787±0.028	77±58	532±182 (9)	719±173 (9)	880±45 (5)	503±237	42±25
10	18	16±13	54±39	137±33	3±2	0.721±0.071	43±31	543±152 (10)	811±112 (9)	926±0 (1)	609±309	48±59
10	19	21±12	83±54	129±15	3±2	0.761±0.034	64±41	495±135 (9)	738±121 (9)	920±40 (4)	620±259	30±17
10	20	16±17	54±44	133±24	2±1	0.761±0.044	46±34	524±206 (9)	796±86 (8)	925±0 (1)	747±416	32±17



1143

1144 1145

1147

1148

1149

1150

1151

1152

1153 1154

1155

1156

1157

1158

1159

1160

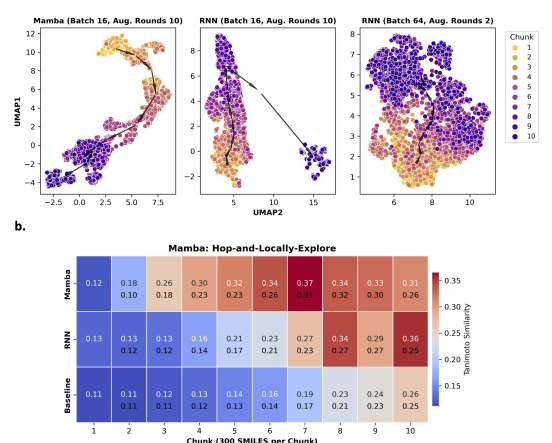


Figure C4: Mamba and RNN (both batch size 16, augmentation rounds 10) and baseline Augmented Memory (batch size 64, augmentation rounds 2). **a.** 3,000 oracle budget test experiment chunked into 300 SMILES. UMAP embedding of the agent chemical space traversal (arrows are the centroid of each chunk). **b.** Mamba exhibits a "hop-and-locally-explore" behavior where the intra-chunk Tanimoto similarity (top values) are higher than RNN. The bottom value is the inter-chunk similarity.

# C.1 Is "Hop-and-Locally-Explore" Always Good?

The results in the main text and this section so far provide evidence that Mamba with batch size 16 and 10 augmentation rounds exhibits local exploration behavior. We hypothesize that sample efficiency improves because "similar molecules, on average, exhibit similar properties". But is this always true? In the test experiment, it is straightforward to see that this indeed holds true. Cross-referencing Fig. C5b, small changes to the molecular graphs should still display high polar surface area which is the objective. However, oracles we care about are physics-based simulations. In the main text results and later in the Appendix for Part 2 and Part 3 additional results, we show that this behavior is beneficial for sample efficiency. The physics-based oracles used in this work are AutoDock Vina 88 and QuickVina 2<sup>90</sup> which run molecular docking. The question we pose is: are these oracles too permissive? Such that the optimization landscape is smooth <sup>82</sup>. As we push towards higher-fidelity oracles such as QM/MM and free energy simulations <sup>15,18</sup>, it is expected that they will be more stringent and demand more specificity. This means that the current hypothesis of "similar molecules, on average, exhibit similar properties" may be loosened. Whether this turns out to be detrimental or not in high-fidelity oracle settings remains to be empirically tested which we leave for future work. By characterizing the behavior of Saturn and understanding what exactly Augmented Memory is doing, it is possible to adapt the current model accordingly. For example, decreasing augmentation rounds relaxes the "hop-and-locally-explore" behavior, which could be advantageous for high-fidelity oracles.

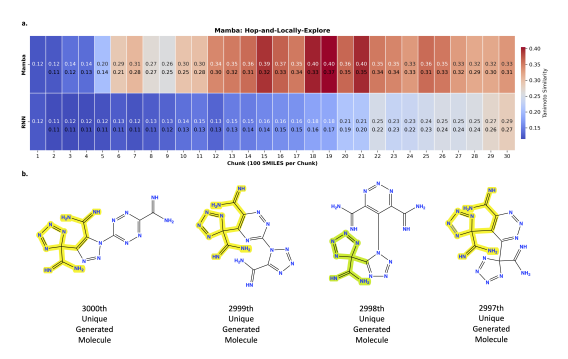


Figure C5: Mamba (batch size 16, augmentation rounds 10) and baseline Augmented Memory (batch size 64, augmentation rounds 2) which is labelled as **RNN**. **a.** 3,000 oracle budget test experiment **chunked into 100 SMILES**. Mamba exhibits a "hop-and-locally-explore" behavior where the intra-chunk Tanimoto similarity (top values) are higher than RNN. The bottom value is the inter-chunk similarity. **b.** Qualitative examples of unique molecules generated at adjacent epochs. Many substructures are shared and the model generates in the local neighborhood. Yellow highlights are exact substructures shared while green indicates a portion.

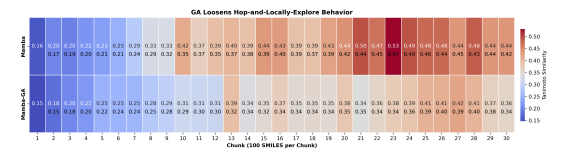


Figure C6: Mamba (batch size 16, augmentation rounds 10) with and without GA <sup>63</sup> activated. The experiment is the Part 3 MPO objective (docking against parp1).

# 1162 C.2 Genetic Algorithm Loosens "Hop-and-Locally-Explore Behavior"

In our investigations of applying a GA on the replay buffer, we show that while sample efficiency does not improve, diversity recovers. To quantitatively show why, we plot the chunk similarity for an experiment from Part 3 on the parp1 target with and without the GA activated (Fig. C6). The Mamba model in both cases uses batch size 16 and 10 augmentation rounds. With the GA activated, the intra-chunk similarities decrease, thus loosening the locally exploration behavior and is the reason why diversity recovers.

# 1169 D Part 2: Transferability of Sample Efficiency to Physics-based Oracles

This section contains information on the Autodock Vina 88 docking protocol and additional results.

All results are averaged across 10 seeds (0-9 inclusive).

# 1172 D.1 Docking Protocol

All protein receptor structures were pre-processed from the raw PDB.

#### 1174 The following were removed:

(x, y, z) = (9.93, 5.85, -9.58).

- 1. Duplicate protein chains and duplicate ligands.
- 1176 2. Co-factors.
- 1177 3. Ions.

1184

- 4. All waters.
- Next, Schrödinger's Protein Preparation Wizard <sup>113,114</sup> with default parameters was used to pre-process the structure. PROPKA hydrogen-bond network optimization was performed at pH 7.4 and energy
- minimization with OPLS3e force-field <sup>115</sup>. Below are details on the docking grids generated from the pre-processed PDBs.
- DRD2 Dopamine Type 2 Receptor. The PDB ID is 6CM4<sup>84</sup> and the docking grid was centered at
- MK2 MK2 Kinase. The PDB ID is  $3KC3^{86}$  and the docking grid for the extracted monomer was centered at (x, y, z) = (-61.62, 30.31, -21.9).
- AChE Acetylcholinesterase. The PDB ID is  $1EVE^{85}$  and the docking grid was centered at (x, y, z) = (2.78, 64.38, 67.97).
- Docking. The search box for all grids was 15Å x 15Å x 15Å and docking was executed through
- 1190 DockStream<sup>8</sup>. All generated molecules were first embedded using the RDKit Universal Force Field
- (UFF) 116 with the maximum convergence set to 600 iterations. Docking was parallelized over 16 CPU
- cores (since the generative model's batch size was 16). The cores were Intel(R) Xeon(R) Platinum
- 1193 8360Y processors.

### 1194 D.2 Additional Results

- In the main text, results were shown at the 0.8 reward threshold. In this section, we also show
- results for Saturn-RNN (batch size 16, augmentation rounds 10) and for the 0.7 reward threshold
- 1197 (Tables 25 and 26). At the 0.7 reward threshold, Saturn-RNN's performance is almost identical
- to Saturn. However, at the 0.8 reward threshold, Saturn (using Mamba) is more performant. We
- highlight that although at times, the difference may be small, it can be highly practically relevant when
- using expensive oracles, e.g., 50 docking calls may be inconsequential but 50 molecular dynamics
- 1201 simulations can be costly. Both Saturn-RNN and Saturn outperform baseline Augmented Memory.
- 1202 Finally, adding a GA on top of Saturn recovers diversity but sample efficiency decreases.

# 1203 **D.3 Compute Time**

Due to insufficient GPU resources, we ran all experiments in this section on CPU. Averaged across all targets and across all 10 replicates, the wall times were as follows: 172 minutes (approximately

Table 25: Docking MPO with 1,000 oracle budget. Baseline is vanilla Augmented Memory  $^{21}$ . All metrics are computed at the 0.7 reward threshold. IntDiv1 is the internal diversity, scaffolds is the number of unique Bemis-Murcko scaffolds, OB is Oracle Burden (oracle calls required to generate N unique molecules). The number in parentheses in the OB statistics represent how many runs out of 10 were successful. The mean and standard deviation across 10 seeds (0-9 inclusive) is reported. Saturn-RNN is RNN with batch size 16 and augmentation rounds 10.

Model	Yield (†)	IntDiv1 (↑)	Scaffolds (↑)	OB 1 (↓)	OB 10 (↓)	OB 100 (↓)
DRD2						
Baseline Saturn-RNN Saturn Saturn-GA	$630 \pm 45$ $818 \pm 22$ $850 \pm 23$ $804 \pm 26$	$\begin{array}{c} 0.858 \pm 0.006 \\ 0.821 \pm 0.011 \\ 0.784 \pm 0.015 \\ 0.817 \pm 0.022 \end{array}$	$585 \pm 43$ $671 \pm 56$ $677 \pm 51$ $685 \pm 56$	$57 \pm 2(10)$ $14 \pm 1(10)$ $14 \pm 1(10)$ $14 \pm 1(10)$	$57 \pm 2(10)$ $31 \pm 6(10)$ $35 \pm 7(10)$ $35 \pm 7(10)$	$279 \pm 32(10)$ $219 \pm 16(10)$ $199 \pm 20(10)$ $199 \pm 19(10)$
MK2 Kinase						
Baseline Saturn-RNN Saturn Saturn-GA	$431 \pm 32$ $704 \pm 25$ $702 \pm 43$ $636 \pm 29$	$\begin{array}{c} 0.863 \pm 0.005 \\ 0.833 \pm 0.013 \\ 0.811 \pm 0.022 \\ 0.827 \pm 0.019 \end{array}$	$406 \pm 26$ $525 \pm 32$ $519 \pm 69$ $506 \pm 68$	$57 \pm 2(10)$ $14 \pm 1(10)$ $17 \pm 6(10)$ $17 \pm 6(10)$	$74 \pm 26(10)$ $43 \pm 9(10)$ $52 \pm 12(10)$ $52 \pm 12(10)$	$396 \pm 37(10)$ $282 \pm 19(10)$ $282 \pm 31(10)$ $291 \pm 31(10)$
AChE						
Baseline Saturn-RNN Saturn Saturn-GA	$801 \pm 27$ $909 \pm 21$ $906 \pm 15$ $874 \pm 21$	$\begin{array}{c} 0.867 \pm 0.006 \\ 0.842 \pm 0.006 \\ 0.816 \pm 0.014 \\ 0.841 \pm 0.008 \end{array}$	$759 \pm 30$ $772 \pm 73$ $742 \pm 76$ $732 \pm 48$	$57 \pm 2(10)$ $14 \pm 1(10)$ $14 \pm 1(10)$ $14 \pm 1(10)$	$57 \pm 2(10)$ $25 \pm 6(10)$ $27 \pm 4(10)$ $27 \pm 4(10)$	$201 \pm 29(10)  163 \pm 19(10)  158 \pm 13(10)  158 \pm 14(10)$

Table 26: Docking MPO with 1,000 oracle budget. Baseline is vanilla Augmented Memory  $^{21}$ . All metrics are computed at the 0.8 reward threshold. IntDiv1 is the internal diversity, scaffolds is the number of unique Bemis-Murcko scaffolds, OB is Oracle Burden (oracle calls required to generate N unique molecules). The number in parentheses in the OB statistics represent how many runs out of 10 were successful. The mean and standard deviation across 10 seeds (0-9 inclusive) is reported. Saturn-RNN is RNN with batch size 16 and augmentation rounds 10.

Model	Yield (↑)	IntDiv1 (↑)	Scaffolds (†)	OB 1 (↓)	OB 10 (↓)	OB 100 (↓)
DRD2						
Baseline Saturn-RNN Saturn Saturn-GA	$22 \pm 7$ $185 \pm 40$ $369 \pm 62$ $209 \pm 55$	$\begin{array}{c} 0.774 \pm 0.019 \\ 0.745 \pm 0.022 \\ 0.671 \pm 0.050 \\ 0.745 \pm 0.041 \end{array}$	$22 \pm 7$ $148 \pm 47$ $310 \pm 70$ $189 \pm 57$	$143 \pm 75(10)$ $128 \pm 94(10)$ $93 \pm 53(10)$ $96 \pm 56(10)$	$733 \pm 120(10)$ $440 \pm 72(10)$ $391 \pm 56(10)$ $403 \pm 75(10)$	Failed $854 \pm 63(10)$ $663 \pm 55(10)$ $806 \pm 84(10)$
MK2 Kinase						
Baseline Saturn-RNN Saturn Saturn-GA	$0.2 \pm 0.4$ $2.5 \pm 3.4$ $14.9 \pm 14.1$ $6.1 \pm 6.5$	$\begin{array}{c} - \\ 0.414 \pm 0.213 \\ 0.454 \pm 0.212 \\ 0.415 \pm 0.202 \end{array}$	$0.2 \pm 0.4$ $2.5 \pm 3.4$ $14.1 \pm 13.2$ $5.5 \pm 5.5$	$836 \pm 186(2)$ $642 \pm 91(6)$ $677 \pm 186(9)$ $678 \pm 140(9)$	Failed $999 \pm 0(1)$ $861 \pm 108(6)$ $911 \pm 11(2)$	Failed Failed Failed Failed
AChE						
Baseline Saturn-RNN Saturn Saturn-GA	$173 \pm 19$ $419 \pm 38$ $480 \pm 79$ $343 \pm 57$	$\begin{array}{c} 0.843 \pm 0.009 \\ 0.804 \pm 0.019 \\ 0.757 \pm 0.020 \\ 0.809 \pm 0.013 \end{array}$	$170 \pm 18$ $338 \pm 55$ $400 \pm 96$ $287 \pm 50$	$57 \pm 2(10)$ $21 \pm 11(10)$ $32 \pm 24(10)$ $32 \pm 25(10)$	$189 \pm 52(10)$ $165 \pm 60(10)$ $185 \pm 82(10)$ $187 \pm 80(10)$	$776 \pm 58(10)$ $531 \pm 36(10)$ $508 \pm 80(10)$ $565 \pm 80(10)$

3 hours) for Augmented Memory<sup>21</sup>, 246 minutes (approximately 4 hours) for Saturn-RNN, 1,426 minutes (approximately 24 hours) for Saturn, and 1,111 minutes (approximately 18.5 hours) for Saturn-GA. There is such a large discrepancy in run time due to repeated SMILES (which do not impose additional oracle calls) that still require backpropagation. Moreover, the runs with Mamba take so much longer because the GPU implementation is highly optimized (we use the official code from https://github.com/state-spaces/mamba). When run on GPU, the difference in wall time between Saturn-RNN and Saturn (Mamba) are not significant.

# **E** Part 3: Benchmarking Saturn

In this section, we detail how Saturn was pre-trained for benchmarking, the procedure we followed to reproduce GEAM <sup>13</sup>, and additional results. We ensured exact reproducibility by using GEAM's official code: https://anonymous.4open.science/r/GEAM-45EF. For running Saturn with GEAM's objective function, all the oracle code was taken, without modification, from the same repository.

#### 1219 E.1 Saturn ZINC 250k Pre-training

GEAM pre-trained on ZINC 250k <sup>89</sup> and provide the dataset in their repository. We used this dataset as is for Saturn pre-training (Mamba model).

#### 1222 The pre-training parameters were:

- 1223 1. Training steps = 50 (each training step entails a full pass through the dataset)
- 1224 2. Seed = 0
- 1225 3. Batch size = 512
- 1226 4. Learning rate = 0.0001
- 5. Train with SMILES randomization <sup>25</sup> (all SMILES in each batch was randomized)

#### 1228 Mamba model:

- 1. Vocabulary size = 66 (including the 2 added tokens for <START> and <END>)
- 1230 2. 5,272,832 parameters
- 3. Used checkpoint from epoch 50 (NLL = 28.10, Validity (10k) = 95.2%)

All Saturn experiments were run on a single workstation equipped with an NVIDIA RTX A6000 GPU and AMD Ryzen 9 5900X 12-Core CPU. The total run time for Saturn across all targets was 41.5 hours (total of 50 runs: 5 targets, 10 seeds each).

# 1235 E.2 Reproducing GEAM's Results

We followed the instructions directly in GEAM's README: https://anonymous.4open.
science/r/GEAM-45EF/README.md. We trained the FGIB with seed 0. Everything else was
run with their default parameters. In the original work, 3 replicates were run but the seeds were not
specified. In our comparisons, we run GEAM across 10 seeds (0-9 inclusive) using an NVIDIA V100
GPU with a Xeon-Gold processor (2.1 GHz and 20 cores) CPU. The reason why a different GPU was
used in GEAM experiments compared to Saturn is due to CUDA compatibility in GEAM's code.

### 1242 E.3 GEAM's MPO Objective

1243 GEAM optimized for the following objective:

$$R(x) = \widehat{DS}(x) \times QED(x) \times \widehat{SA}(x) \in [0, 1]$$
(22)

 $\widehat{DS}$  is the normalized QuickVina 2<sup>90</sup> docking score (Eq. 23), QED<sup>87</sup> is the quantitative estimate of drug-likeness, and  $\widehat{SA}$  is the normalized synthetic accessibility score <sup>91</sup> (Eq. 24).

$$\widehat{DS} = -\frac{DS}{20} \tag{23}$$

$$\widehat{SA} = \frac{10 - SA}{9} \tag{24}$$

#### 1246 E.4 Saturn-Jaccard

In GEAM <sup>13</sup>, the "Novel" in **Novel Hit Ratio** enforces molecules to possess < 0.4 Tanimoto similarity to ZINC 250k <sup>89</sup>. GEAM achieves this by use of their genetic algorithm which directly uses GraphGA <sup>63</sup>. The crossover and mutation operations promote diversity. Otherwise, generative models are pre-trained to model the training data distribution. This means that generated molecules would not necessarily be *very* dissimilar to the training data, especially if the training data actually possesses "good" molecules already. By virtue of pre-training on a selected dataset, we implicitly assume that the pre-training dataset is "good" for our task, otherwise, we probably should not pre-train on this data. This is the rationale on why ChEMBL <sup>79</sup> and ZINC 250k <sup>89</sup> are popular pre-training datasets:

they contain bio-active molecules. To satisfy GEAM's "Novel" criterion, we take the base Saturn 1255 model and first teach it to generate molecules that are dissimilar to the ZINC 250k dataset which 1256 was used for pre-training. The objective function is then defined as minimizing the max Tanimoto 1257 similarity to any molecule in ZINC 250k. This experiment was run with an oracle budget of 1500 and 1258 took about 10 minutes. The resulting **Saturn-Jaccard** model generates molecules with low Tanimoto 1259 similarity to ZINC 250k. Starting from this model, we run GEAM's case study and the results from 1260 1261 this are reported in the main text and here in the Appendix. We finally note that this criterion is somewhat arbitrary and we do it so we can exactly match GEAM's experiments. 1262

# E.5 Quantitative Supplementary Results

1263

1264

1265

1266

1267

1268

1269

1270

1275 1276

1277

1278

1280

1281

1282

1283

1284

1285

In this section, we present supplementary benchmarking results and show additional results for Saturn-GA.

Table 27: Hit Ratio (%). Results are from Lee et al. <sup>12</sup> except GEAM, datasets, and Saturn which we ran across 10 seeds (0-9 inclusive). The mean and standard deviation are reported. Best results (statistically significant at the 95% confidence level) are bolded.

Method			Target Protein		
	parp1	fa7	5ht1b	braf	jak2
Datasets					
ZINC 250k 89	$3.993 \pm 0.355$	$1.097 \pm 0.192$	$24.26 \pm 0.622$	$1.020 \pm 0.193$	$6.183 \pm 0.344$
ChEMBL 33 <sup>79</sup>	$6.077 \pm 0.453$	$1.830 \pm 0.240$	$24.163 \pm 0.715$	$2.073 \pm 0.181$	$9.013 \pm 0.562$
Generative Models					
REINVENT 23	$4.693 \pm 1.776$	$1.967 \pm 0.661$	$26.047 \pm 2.497$	$2.207 \pm 0.800$	$5.667 \pm 1.067$
JT-VAE 45	$3.200 \pm 0.348$	$0.933 \pm 0.152$	$18.044 \pm 0.747$	$0.644 \pm 0.157$	$5.856 \pm 0.204$
GraphAF 93	$0.822 \pm 0.113$	$0.011 \pm 0.016$	$6.978 \pm 0.952$	$1.422 \pm 0.556$	$1.233 \pm 0.284$
MORLD 94	$0.047 \pm 0.050$	$0.007 \pm 0.013$	$0.893 \pm 0.758$	$0.047 \pm 0.040$	$0.227 \pm 0.118$
HierVAE 95	$1.180 \pm 0.182$	$0.033 \pm 0.030$	$0.740 \pm 0.371$	$0.367 \pm 0.187$	$0.487 \pm 0.183$
GraphDF 99	$0.044 \pm 0.031$	$0.000 \pm 0.000$	$0.000 \pm 0.000$	$0.011 \pm 0.016$	$0.011 \pm 0.016$
FREED 11	$4.860 \pm 1.415$	$1.487 \pm 0.242$	$14.227 \pm 5.116$	$2.707 \pm 0.721$	$6.067 \pm 0.790$
FREED-QS 11	$5.960 \pm 0.902$	$1.687 \pm 0.177$	$23.140 \pm 2.422$	$3.880 \pm 0.623$	$7.653 \pm 1.373$
LIMO 100	$0.456 \pm 0.057$	$0.044 \pm 0.016$	$1.200 \pm 0.178$	$0.278 \pm 0.134$	$0.711 \pm 0.329$
GDSS 101	$2.367 \pm 0.316$	$0.467 \pm 0.112$	$6.267 \pm 0.287$	$0.300 \pm 0.198$	$1.367 \pm 0.258$
MOOD 12	$7.260 \pm 0.764$	$0.787 \pm 0.128$	$21.427 \pm 0.502$	$5.913 \pm 0.311$	$10.367 \pm 0.616$
Augmented Memory 21	$16.966 \pm 3.224$	$2.637 \pm 0.860$	$52.016 \pm 2.302$	$8.307 \pm 1.714$	$21.548 \pm 4.938$
GEAM 13	$45.158 \pm 2.408$	$20.552 \pm 2.357$	$47.664 \pm 1.198$	$30.444 \pm 1.610$	$46.129 \pm 2.073$
Ours					
Saturn	$\bf 57.981 \pm 18.537$	$14.527 \pm 9.961$	$68.185 \pm 3.400$	${\bf 38.999 \pm 10.114}$	$60.827 \pm 11.502$
Saturn-GA	$55.597 \pm 5.617$	$16.711 \pm 6.761$	$63.112 \pm 4.316$	$34.284 \pm 10.345$	$58.625 \pm 6.982$
Saturn-Jaccard	$77.674 \pm 7.127$	$23.119 \pm 6.852$	$78.433 \pm 1.029$	$30.258 \pm 12.315$	$83.012 \pm 6.678$

Hit Ratio (%). Table 27 shows the Hit Ratio (%) results. Random sampling of 3,000 molecules from common datasets (ZINC 250k<sup>89</sup> and ChEMBL 33<sup>79</sup>) are included as baselines. The results show that only GEAM<sup>13</sup> and Saturn outperform these baselines with both methods performing similarly overall. With the exception of a few targets where performance differs (significant at the 95% confidence level), Saturn notably exhibits higher variance which is expected given the small batch size (16). One way to mitigate high variance is to use a larger batch size, as this makes the approximation for the expected reward less noisy. Next, we show that the Saturn-Jaccard agent displays notably high Hit Ratios but do not present this in the main results as the purpose of the Jaccard agent is to generate hits that have less than 0.4 Tanimoto similarity to the ZINC 250k 89 training dataset. It is difficult to predict a priori a favorable chemical space to move the agent. However, this result is interesting as it suggests that this simple additional pre-training which took minutes via curriculum learning (CL), makes the agent more suited for the docking tasks. Finally, we show that using the GA (Saturn-GA) is a straightforward solution to recover diversity. From Part 1 and Part 2 experiments, activating the GA comes at the expense of some sample efficiency but interestingly, this is not the case here (Table 28). Moreover, Saturn-GA also decreases variance in this case study (Table 27). Based on these results, it would actually be beneficial to activate the GA in this case, but it is difficult to know a priori the best configuration, thus we report the out-of-the-box hyperparameters (without GA) in the main text based on tuning on the test experiment in Part 1.

**Novel Hit Ratio** (%). Table 29 shows the Novel Hit Ratio (%) results with all additional metrics, mirroring the main text table. Similar to the main text results, Mamba-Jaccard agent generates significantly more molecules passing the strict filter and also much faster (fewer oracle calls).

Table 28: Strict Hit Ratio (%) (QED > 0.7 and SA < 3) additional results. GEAM and Saturn results are across 10 seeds (0-9 inclusive). OB is Oracle Burden (oracle calls required to generate N unique molecules). The number in parentheses in the OB statistics represent how many runs out of 10 were successful. The mean and standard deviation are reported. Best results (statistically significant at the 95% confidence level) are bolded.

Method			Target Protein		
	parp1	fa7	5ht1b	braf	jak2
GEAM <sup>13</sup> - Presented in Main Text					
Strict Hit Ratio (↑)	$6.510 \pm 1.087$	$2.106 \pm 0.958$	$8.719 \pm 0.903$	$3.685 \pm 0.524$	$7.944 \pm 1.157$
IntDiv1 (↑)	$0.766 \pm 0.017$	$0.709 \pm 0.043$	$0.799 \pm 0.017$	$0.751 \pm 0.023$	$0.763 \pm 0.021$
#Circles (↑)	$14 \pm 3$	$7\pm2$	$25 \pm 3$	$11 \pm 2$	$18 \pm 2$
OB (1) (\(\psi\))	$250 \pm 157(10)$	$433 \pm 209(10)$	$114 \pm 112(10)$	$355 \pm 96(10)$	$230 \pm 117(10)$
OB (10) (\( \psi \))	$743 \pm 52(10)$	$1446 \pm 404(10)$	$531 \pm 38(10)$	$892 \pm 144(10)$	$537 \pm 70(10)$
OB (100) (\( \psi\))	$2106 \pm 202(10)$	$2927 \pm 0(1)$	$1527 \pm 110(10)$	$2674 \pm 163(6)$	$1606 \pm 218(10)$
Saturn (ours) - Presented in Main Text					
Strict Hit Ratio	$55.102 \pm 18.027$	$13.887 \pm 9.723$	$64.730 \pm 3.717$	$37.250 \pm 9.615$	$55.903 \pm 13.613$
IntDiv1 (↑)	$0.596 \pm 0.049$	$0.592 \pm 0.066$	$0.685 \pm 0.021$	$0.597 \pm 0.042$	$0.638 \pm 0.034$
#Circles (↑)	$5 \pm 0$	$3 \pm 1$	$17 \pm 3$	$4 \pm 0$	$7\pm1$
OB (1) (\( \psi \))	$139 \pm 96(10)$	$352 \pm 206(10)$	$21 \pm 7(10)$	$291 \pm 143(10)$	$88 \pm 56(10)$
OB (10) (\( \psi \))	$518 \pm 92(10)$	$924 \pm 247(10)$	$105 \pm 23(10)$	$581 \pm 123(10)$	$348 \pm 96(10)$
OB (100) (\psi)	$956 \pm 259(10)$	$1776 \pm 551(10)$	$441 \pm 44(10)$	$1057 \pm 187(10)$	$785 \pm 191(10)$
Saturn-GA (ours) - Newly presented here					
Strict Hit Ratio	$47.146 \pm 4.952$	$13.187 \pm 6.340$	$53.055 \pm 3.764$	$28.377 \pm 9.703$	$49.528 \pm 5.463$
IntDiv1 (†)	$0.659 \pm 0.023$	$0.636 \pm 0.039$	$0.724 \pm 0.022$	$0.625 \pm 0.047$	$0.676 \pm 0.041$
#Circles (↑)	$8 \pm 2$	$4\pm1$	$22\pm4$	$6 \pm 1$	$12 \pm 2$
OB (1) (\(\psi\))	$121 \pm 71(10)$	$350 \pm 203(10)$	$20 \pm 6(10)$	$242 \pm 194(10)$	$91 \pm 43(10)$
OB (10) (↓)	$467 \pm 114(10)$	$912 \pm 168(10)$	$110 \pm 36(10)$	$582 \pm 177(10)$	$375 \pm 120(10)$
OB (100) (\psi)	$937 \pm 136(10)$	$1852 \pm 349(10)$	$499 \pm 85(10)$	$1266 \pm 486(10)$	$861 \pm 123(10)$

Table 29: Strict Novel Hit Ratio (%) (QED > 0.7 and SA < 3). GEAM and Saturn results are across 10 seeds (0-9 inclusive). OB is Oracle Burden (oracle calls required to generate N unique molecules). The number in parentheses in the OB statistics represent how many runs out of 10 were successful. The mean and standard deviation are reported. Best results (statistically significant at the 95% confidence level) are bolded.

Method			Target Protein		_
	parp1	fa7	5ht1b	braf	jak2
GEAM 13					
Strict Hit Ratio (↑)	$4.018 \pm 0.849$	$1.676 \pm 0.836$	$5.338 \pm 0.789$	$2.621 \pm 0.464$	$5.930 \pm 1.151$
IntDiv1 (†)	$0.768 \pm 0.019$	$0.710 \pm 0.047$	$0.793 \pm 0.019$	$0.753 \pm 0.026$	$0.763 \pm 0.026$
#Circles (↑)	$13 \pm 2$	$5 \pm 2$	$21 \pm 3$	$11 \pm 2$	$16 \pm 3$
OB (1) (\(\psi\))	$319 \pm 175(10)$	$502 \pm 209(10)$	$253 \pm 159(10)$	$419 \pm 102(10)$	$242 \pm 124(10)$
OB (10) (↓)	$857 \pm 86(10)$	$1625 \pm 380(10)$	$689 \pm 77(10)$	$1047 \pm 136(10)$	$616 \pm 83(10)$
OB (100) (↓)	$2633 \pm 202(9)$	Failed	$2221 \pm 224(10)$	$2942 \pm 0(1)$	$2005 \pm 268(10)$
Saturn-Jaccard (ours)					
Strict Novel Hit Rate	$47.405 \pm 8.593$	$17.130 \pm 5.538$	$50.445 \pm 6.334$	$18.228 \pm 9.438$	$45.185 \pm 13.321$
IntDiv1 (↑)	$0.595 \pm 0.029$	$0.600 \pm 0.030$	$0.559 \pm 0.032$	$0.520 \pm 0.040$	$0.567 \pm 0.041$
#Circles (†)	$2 \pm 0$	$2 \pm 0$	$2 \pm 0$	$1 \pm 0$	$1 \pm 0$
OB (1) (\(\psi\))	$26 \pm 17(10)$	$98 \pm 53(10)$	$15 \pm 0(10)$	$164 \pm 137(10)$	$18 \pm 7(10)$
OB (10) (\( \psi \))	$177 \pm 38(10)$	$320 \pm 69(10)$	$31 \pm 5(10)$	$388 \pm 156(10)$	$70 \pm 13(10)$
OB (100) (\(\psi\))	$562 \pm 94(10)$	$1051 \pm 251(10)$	$223 \pm 50(10)$	$1041 \pm 58\dot{5}(9)$	$402 \pm 196(10)$
Saturn-Jaccard-GA (ours)					
Strict Novel Hit Rate	$29.801 \pm 11.603$	$11.895 \pm 5.197$	$40.261 \pm 8.168$	$17.845 \pm 7.943$	$37.498 \pm 11.200$
IntDiv1 (†)	$0.621 \pm 0.041$	$0.596 \pm 0.030$	$0.613 \pm 0.042$	$0.640 \pm 0.040$	$0.606 \pm 0.034$
#Circles (†)	$3\pm1$	$2\pm1$	$3\pm1$	$3\pm1$	$3\pm1$
OB (1) (↓)	$36 \pm 38(10)$	$216 \pm 232(10)$	$15 \pm 0(10)$	$181 \pm 122(10)$	$17 \pm 5(10)$
OB (10) (\( \psi \)	$205 \pm 65(10)$	$556 \pm 275(10)$	$27 \pm 5(10)$	$472 \pm 135(10)$	$96 \pm 13(10)$
OB (100) (\psi)	$703 \pm 113(10)$	$1490 \pm 460(9)$	$272 \pm 39(10)$	$1367 \pm 561(10)$	$480 \pm 84(10)$

However, the diversity notably drops (much more than the Mamba agent without Jaccard distance training presented in the main text). However, diversity is particularly low. We first not that when moving to high-fidelity oracles where satisfying the objective function equates to higher true positive hit rates, low diversity need not be detrimental. We additionally run an experiment with the GA activated and we see diversity recovers, but is still notably lower than GEAM. Moreover, the sample efficiency drops notably here compared to without GA, but is still much more performant than GEAM in finding hits faster. Finally, to recover more diversity, one could make the Diversity Filter <sup>77</sup> more stringent. In this work, a bucket size of 10 was used (allow 10 of the same scaffold to be generated before truncating the reward to 0). Decreasing the bucket size to 5 or even lower, may recover more diversity.

#### E.6 Saturn: Architecture Scaling.

1297

In the main text Part 1, we investigated *why* Mamba (5.2M) outperforms LSTM $^{26}$  RNN (5.8M) and decoder transformer  $^{27,28}$  (6.3M). Augmented Memory  $^{21}$  squeezes the likelihood of generating augmented forms of *any* replay buffer *molecules*. Increased capacity to match this distribution directly leads to the "hop-and-locally-explore" behavior which improves sample efficiency. We note that our observations are for optimization landscapes that are not *too rough*  $^{81,82}$ . It is difficult to know *a priori* the roughness of optimization and also whether the benefits of "hop-and-locally-explore" behavior is beneficial in higher-fidelity oracle settings. We leave this for future work.

Based on these observations, we investigate scaling benefits for the LSTM RNN and decoder 1305 transformer models. Increasing model size can lead to lower loss convergence, which in this case, 1306 means modelling the conditional token distribution of the SMILES<sup>30</sup>. One may argue that this is 1307 simply a hyperparameter tuning which we missed. However, the purpose of this work is in the 1308 goal-directed learning setting where we want to tune the model's distribution towards desirable 1309 molecules. If desirable molecules are already in the training data, minimal optimization is required. 1310 Moreover, it is difficult to know a priori whether matching the training distribution very closely is 1311 strictly advantageous for an arbitrary MPO objective, unless we have an enormous amount of data, 1312 by the law of large numbers. Therefore, all pre-trained models (priors) in this work were trained until loss flattens out and Validity (fraction of valid SMILES generated) is high.

In this section, we scale up the LSTM RNN and decoder transformer models to around 25M to make the *distribution learning capability* approach Mamba (5.2M). We use the training loss for this, where similar loss convergence is taken as the proxy. We first present the exact model parameter counts, hyperparameters, and training details.

#### LSTM RNN 24.7M:

1. Seed = 0

1319

1320

1332

- 1321 2. Parameters = 24,741,442
- 3. Vocabulary Size = 66
- 4. Embedding Dimension = 256
- 5. Hidden Dimension = 512
- 6. Number of Layers = 12
- 1326 7. Dropout = 0.0
- 8. Layer Normalization = False
- 9. Train Epochs = 300
- 1329 10. Batch Size = 512
- 1330 11. Learning Rate = 0.0001
- 1331 12. Final NLL Loss at Epoch 300 = 29.318

#### Decoder 25.3M:

- 1333 1. Seed = 0
- 1334 2. Parameters = 25,306,178
- 3. Vocabulary Size = 66
- 4. Embedding Dimension = 256
- 5. Hidden Dimension = 1024
- 6. Number of Layers = 32
- 7. Number of Heads = 16
- 1340 8. Dropout = 0.0
- 9. Train Epochs = 100
- 1342 10. Batch Size = 512
- 1343 11. Learning Rate = 0.0001

#### 12. Final NLL Loss at Epoch 100 = 26.963

In addition, we scale up Mamba to 16M and 21M and also present the exact model parameter counts, hyperparameters, and training details. For these two models, we intentionally train until the loss is at similar values (NLL = 26) which suggests both models have learned the training distribution to a similar extent. Optimization then starts from a similar distribution.

#### 1349 Mamba 15.8M:

1344

1345

1346

1348

1355

1362

- 1350 1. Seed = 0
- 1351 2. Parameters = 15,785,728
- 3. Vocabulary Size = 66
- 4. Embedding Dimension = 256
- 5. Number of Layers = 36
  - 6. Use RMSNorm = True
- 7. Residual in fp32 = True
- 8. Fused AddNorm = True
- 9. Train Epochs = 100
- 1359 10. Batch Size = 512
- 1360 11. Learning Rate = 0.0001
- 1361 12. Final NLL Loss at Epoch 92 = 26.003

#### **Mamba 21.0M**:

- 1363 1. Seed = 0
- 1364 2. Parameters = 21,041,920
- 3. Vocabulary Size = 66
- 4. Embedding Dimension = 256
- 5. Number of Layers = 48
- 1368 6. Use RMSNorm = True
- 7. Residual in fp32 = True
- 1370 8. Fused AddNorm = True
- 9. Train Epochs = 100
- 1372 10. Batch Size = 512
- 11. Learning Rate = 0.0001
- 12. Final NLL Loss at Epoch 75 = 25.993

Hit Ratios (%). Table 30 shows the Hit Ratios of compared models. Saturn outperforms baseline Augmented Memory and GEAM. In terms of architecture scaling, we show decoder transformer and RNN approach Mamba performance but are still less performant. Scaling up Mamba does not necessarily lead to better results, as there is notably even higher variance.

Sample Efficiency Metrics Table 31 presents the Strict Hit Ratios for compared models. While 1379 GEAM outperforms baseline Augmented Memory for the Hit Ratio, the results here show that the 1380 optimization capability of baseline Augmented Memory exceeds that of GEAM. Saturn outperforms 1381 both Augmented Memory and GEAM to generate more hits and also finds them faster (lower 1382 OB). Next, we investigate architecture scaling again, but this time, under the strict filter. decoder 1383 transformer (25.3M) approaches Mamba (5.2M) performance and outperforms it in many tasks (Fig. 1384 31), trading off even more diversity. Variance is also higher. However, we believe this is an interesting 1385 observation as Augmented Memory's mechanism is squeezing the likelihood of augmented sequences. 1386 By simply scaling up the architecture and enabling the model to converge to this distribution, sample 1387 efficiency improves. This directly draws parallel to NLP LLMs where scaling improves downstream 1388 performance on many tasks, when trained on next token prediction 117. Finally, while scaling up the 1389 architecture to the parameter counts we have investigated adds negligible generation time, Mamba 1390 (5.2M) is parameter-efficient in its synergistic behavior with Augmented Memory. 1391

Table 30: Architecture scaling experiments: Hit Ratio (%) metrics. GEAM <sup>13</sup> and Saturn results are across 10 seeds (0-9 inclusive). The mean and standard deviation are reported.

Method	Target Protein							
	parp1	fa7	5ht1b	braf	jak2			
Datasets								
ZINC 250k 89	$3.993 \pm 0.355$	$1.097 \pm 0.192$	$24.26 \pm 0.622$	$1.020 \pm 0.193$	$6.183 \pm 0.344$			
ChEMBL 33 <sup>79</sup>	$6.077 \pm 0.453$	$1.830 \pm 0.240$	$24.163 \pm 0.715$	$2.073 \pm 0.181$	$9.013 \pm 0.562$			
Generative Models								
Augmented Memory 21	$16.983 \pm 3.221$	$2.641 \pm 0.868$	$52.046 \pm 2.327$	$8.354 \pm 1.727$	$21.604 \pm 4.958$			
GEAM <sup>13</sup>	$49.597 \pm 3.078$	$21.988 \pm 2.968$	$51.765 \pm 1.463$	$33.086 \pm 1.673$	$51.228 \pm 3.132$			
Ours								
Saturn-Mamba 5.2M	$57.981 \pm 18.537$	$14.527 \pm 9.961$	$68.185 \pm 3.400$	$38.999 \pm 10.114$	$60.827 \pm 11.502$			
Saturn-Mamba 15.8M	$56.088 \pm 9.899$	$18.804 \pm 13.980$	$68.322 \pm 3.885$	$38.699 \pm 19.841$	$61.320 \pm 18.673$			
Saturn-Mamba 21.0M	$56.299 \pm 16.583$	$23.764 \pm 19.280$	$65.015 \pm 6.060$	$32.018 \pm 12.584$	$59.175 \pm 20.689$			
Saturn-Decoder 25.3M	$61.732 \pm 16.032$	$21.058 \pm 13.940$	$68.340 \pm 5.094$	$37.399 \pm 12.632$	$65.470 \pm 12.628$			
Saturn-RNN 24.7M	$52.914 \pm 9.955$	$13.254 \pm 7.276$	$63.799 \pm 3.249$	$33.805 \pm 8.694$	$54.165 \pm 7.445$			

# **E.7** Qualitative Supplementary Results

 In this section, we show random generated molecules from Saturn that pass the Strict Filter (Fig. E7). All molecules possess QuickVina  $2^{90}$  docking scores better than the median of known actives  $^{12}$  while possessing QED  $^{87} > 0.7$  and SA score  $^{91} < 3$ . We further highlight two points: firstly, there may be some particularly large rings that are undesirable from a chemistry perspective, even though QED and SA score permits them. Saturn is an optimization engine and if specific chemistry is desired, including it into the MPO objective will steer the agent away from this chemical space. In this work, a concrete example of this is in the main text Part 3 experiments where the Saturn pre-trained model was additionally pre-trained via curriculum learning  $^{81}$  to generate molecules dissimilar to the ZINC  $250k^{89}$  training data to satisfy the *Novel* metric defined Lee et al  $^{12,13}$ . This example shows the flexibility of Saturn. Secondly, as stereochemistry was not purged from the vocabulary, Saturn can generate stereoisomers.

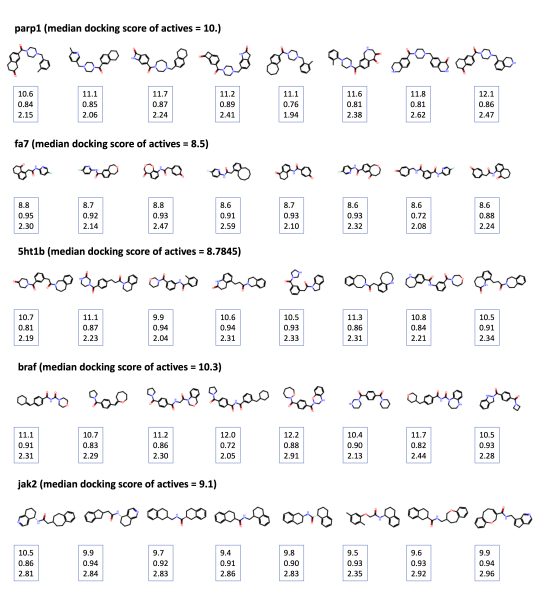


Figure E7: Example Saturn generated molecules passing the Strict Filter for all 5 targets: parp1, fa7, 5ht1b, braf, and jak2. The scores are annotated from top to bottom, QuickVina  $2^{90}$  docking score, QED<sup>87</sup>, and SA score  $^{91}$ .

Table 31: Architecture scaling experiments: Strict Hit Ratio (%) (QED > 0.7 and SA < 3). GEAM and Saturn results are across 10 seeds (0-9 inclusive). OB is Oracle Burden (oracle calls required to generate N unique molecules). The number in parentheses in the OB statistics represent how many runs out of 10 were successful. The mean and standard deviation are reported.

Method			Target Protein		
	parp1	fa7	5ht1b	braf	jak2
GEAM 13					
Strict Hit Ratio (↑)	$6.510 \pm 1.087$	$2.106 \pm 0.958$	$8.719 \pm 0.903$	$3.685 \pm 0.524$	$7.944 \pm 1.157$
IntDiv1 (†)	$0.766 \pm 0.017$	$0.709 \pm 0.043$	$0.799 \pm 0.017$	$0.751 \pm 0.023$	$0.763 \pm 0.021$
#Circles (↑)	$14 \pm 3$	$7\pm2$	$25 \pm 3$	$11 \pm 2$	$18 \pm 2$
OB (1) (\(\psi\))	$250 \pm 157(10)$	$433 \pm 209(10)$	$114 \pm 112(10)$	$355 \pm 96(10)$	$230 \pm 117(10)$
OB (10) (\( \psi \)	$743 \pm 52(10)$	$1446 \pm 404(10)$	$531 \pm 38(10)$	$892 \pm 144(10)$	$537 \pm 70(10)$
OB (100) (\(\psi\))	$2106 \pm 202(10)$	$2927 \pm 0(1)$	$1527 \pm 110(10)$	$2674 \pm 163(6)$	$1606 \pm 218(10)$
Augmented Memory 21	2100 ± 202(10)	2021 ± 0(1)	1021 ± 110(10)	2011 ± 100(0)	1000 ± 210(10)
Strict Hit Ratio	$13.486 \pm 3.033$	$1.757 \pm 0.805$	$43.824 \pm 2.124$	$6.920 \pm 1.734$	$17.884 \pm 4.636$
IntDiv1 (†)	$0.748 \pm 0.019$	$0.718 \pm 0.047$	$0.779 \pm 0.007$	$0.685 \pm 0.022$	$0.772 \pm 0.013$
#Circles (↑)	$20 \pm 5$	$9 \pm 2$	$54 \pm 6$	$8 \pm 1$	$27 \pm 3$
OB (1) (\(\psi\)	$173 \pm 149(10)$	$503 \pm 313$	$61 \pm 1(10)$	$329 \pm 152$	$80 \pm 28(10)$
OB (1) (\(\psi\)) OB (10) (\(\psi\))	$686 \pm 214(10)$	$1776 \pm 257(10)$	$117 \pm 51(10)$	$1173 \pm 375(10)$	$420 \pm 54(10)$
OB (100) (\$\dagger\$) OB (100) (\$\dagger\$)	$1836 \pm 174(10)$	$2867 \pm 0(1)$	$657 \pm 80(10)$	$2396 \pm 139(9)$	$1499 \pm 109(10)$
ОВ (100) (\$)	1630 ± 174(10)	2807 ± 0(1)	037 ± 60(10)	2390 ± 139(9)	1499 ± 109(10)
Ours					
Saturn-Mamba 5.2M					
Strict Hit Ratio	$55.102 \pm 18.027$	$13.887 \pm 9.723$	$64.730 \pm 3.717$	$37.250 \pm 9.615$	$55.903 \pm 13.613$
IntDiv1 (†)	$0.596 \pm 0.049$	$0.592 \pm 0.066$	$0.685 \pm 0.021$	$0.597 \pm 0.042$	$0.638 \pm 0.034$
#Circles (†)	$5\pm0$	$3 \pm 1$	$17 \pm 3$	$4 \pm 0$	$7\pm1$
OB (1) (↓)	$139 \pm 96(10)$	$352 \pm 206(10)$	$21 \pm 7(10)$	$291 \pm 143(10)$	$88 \pm 56(10)$
OB (10) (↓)	$518 \pm 92(10)$	$924 \pm 247(10)$	$105 \pm 23(10)$	$581 \pm 123(10)$	$348 \pm 96(10)$
OB (100) (\(\psi\))	$956 \pm 259(10)$	$1776 \pm 551(10)$	$441 \pm 44(10)$	$1057 \pm 187(10)$	$785 \pm 191(10)$
Saturn-Mamba 15.8M					
Strict Hit Ratio	$52.093 \pm 12.503$	$18.064 \pm 13.932$	$63.740 \pm 5.623$	$37.350 \pm 19.173$	$59.372 \pm 18.465$
IntDiv1 (†)	$0.587 \pm 0.033$	$0.587 \pm 0.068$	$0.662 \pm 0.042$	$0.568 \pm 0.064$	$0.633 \pm 0.035$
#Circles (†)	$6\pm 2$	$3 \pm 1$	18 ± 3	$4 \pm 1$	9 ± 2
OB (1) (\(\psi\)	$157 \pm 112(10)$	$223 \pm 167(10)$	$25 \pm 10(10)$	$204 \pm 115(10)$	$54 \pm 43(10)$
OB (10) (\( \psi \)	$406 \pm 111(10)$	$691 \pm 151(10)$	$108 \pm 31(10)$	$634 \pm 180(10)$	$266 \pm 50(10)$
OB (100) (\( \psi \))	$905 \pm 204(10)$	$1491 \pm 389(8)$	$421 \pm 61(10)$	$1220 \pm 410(10)$	$786 \pm 254(10)$
Saturn-Mamba 21.0M					
Strict Hit Ratio	$54.297 \pm 16.480$	$23.021 \pm 19.064$	$61.307 \pm 5.991$	$30.972 \pm 12.605$	$57.013 \pm 20.601$
IntDiv1 (†)	$0.590 \pm 0.041$	$0.535 \pm 0.056$	$0.655 \pm 0.042$	$0.560 \pm 0.060$	$0.605 \pm 0.046$
#Circles (†)	$6 \pm 1$	$4 \pm 1$	$17 \pm 3$	$4 \pm 1$	8 ± 1
OB (1) (1)	$167 \pm 73(10)$	$316 \pm 236(10)$	$28 \pm 13(10)$	$235 \pm 138(10)$	$68 \pm 78(10)$
OB (10) (\( \psi \)	$425 \pm 91(10)$	$710 \pm 314(10)$	$115 \pm 44(10)$	$556 \pm 147(10)$	$335 \pm 118(10)$
OB (100) (\$\dagger\$)	$831 \pm 147(10)$	$1446 \pm 629(9)$	$432 \pm 69(10)$	$1134 \pm 282(10)$	$798 \pm 340(10)$
· / · · · /	,	,	. ,	,	` /
Saturn-Decoder 25.3M					
Strict Hit Ratio	$59.560 \pm 15.480$	$20.195 \pm 13.394$	$65.202 \pm 5.847$	$35.857 \pm 12.228$	$62.874 \pm 11.810$
IntDiv1 (†)	$0.615 \pm 0.034$	$0.575 \pm 0.078$	$0.658 \pm 0.031$	$0.614 \pm 0.045$	$0.590 \pm 0.062$
#Circles (↑)	$6 \pm 1$	$3 \pm 1$	$13 \pm 3$	$4 \pm 1$	$6 \pm 1$
OB (1) (\( \psi \))	$98 \pm 81(10)$	$242 \pm 160(10)$	$18 \pm 5(10)$	$248 \pm 81(10)$	$52 \pm 37(10)$
OB (10) (\(\psi\))	$375 \pm 131(10)$	$797 \pm 227(10)$	$92 \pm 29(10)$	$515 \pm 98(10)$	$320 \pm 63(10)$
OB (100) (↓)	$769 \pm 165(10)$	$1698 \pm 507(10)$	$378 \pm 43(10)$	$1101 \pm 216(10)$	$722 \pm 140(10)$
Saturn-RNN 24.7M					
Strict Hit Ratio	$50.586 \pm 9.574$	$12.731 \pm 7.211$	$60.331 \pm 3.294$	$32.380 \pm 8.503$	$51.819 \pm 7.247$
IntDiv1 (†)	$0.654 \pm 0.023$	$0.642 \pm 0.042$	$0.719 \pm 0.018$	$0.636 \pm 0.030$	$0.693 \pm 0.027$
#Circles (↑)	8 ± 2	$4\pm1$	$25 \pm 5$	$7\pm1$	$12 \pm 2$
OB (1) (\(\psi\)	$126 \pm 99(10)$	$384 \pm 289(10)$	$27 \pm 19(10)$	$186 \pm 170(10)$	$50 \pm 52(10)$
OB (10) (\( \psi \)	$465 \pm 71(10)$	$1243 \pm 273(10)$	$111 \pm 41(10)$	$714 \pm 214(10)$	$305 \pm 100(10)$
OB (100) (\(\psi\))	$1045 \pm 148(10)$	$2150 \pm 311(10)$	$487 \pm 61(10)$	$1404 \pm 269(10)$	$935 \pm 130(10)$

# **NeurIPS Paper Checklist**

#### 1. Claims

1405

1406

1407

1408

1409

1410

1411

1412

1413

1414

Question: Do the main claims made in the abstract and introduction accurately reflect the paper's contributions and scope?

Answer: [Yes]

Justification: The paper starts by elucidating the mechanism of SMILES augmentation and experience replay and how Mamba synergistically leverages this. The remanining paper benchmarks the model against previous works.

#### Guidelines:

• The answer NA means that the abstract and introduction do not include the claims made in the paper.

- The abstract and/or introduction should clearly state the claims made, including the contributions made in the paper and important assumptions and limitations. A No or NA answer to this question will not be perceived well by the reviewers.
- The claims made should match theoretical and experimental results, and reflect how much the results can be expected to generalize to other settings.
- It is fine to include aspirational goals as motivation as long as it is clear that these goals
  are not attained by the paper.

#### 2. Limitations

Question: Does the paper discuss the limitations of the work performed by the authors?

Answer: [Yes]

Justification: Limitations are discussed in the Conclusion section.

#### Guidelines:

- The answer NA means that the paper has no limitation while the answer No means that the paper has limitations, but those are not discussed in the paper.
- The authors are encouraged to create a separate "Limitations" section in their paper.
- The paper should point out any strong assumptions and how robust the results are to violations of these assumptions (e.g., independence assumptions, noiseless settings, model well-specification, asymptotic approximations only holding locally). The authors should reflect on how these assumptions might be violated in practice and what the implications would be.
- The authors should reflect on the scope of the claims made, e.g., if the approach was
  only tested on a few datasets or with a few runs. In general, empirical results often
  depend on implicit assumptions, which should be articulated.
- The authors should reflect on the factors that influence the performance of the approach. For example, a facial recognition algorithm may perform poorly when image resolution is low or images are taken in low lighting. Or a speech-to-text system might not be used reliably to provide closed captions for online lectures because it fails to handle technical jargon.
- The authors should discuss the computational efficiency of the proposed algorithms and how they scale with dataset size.
- If applicable, the authors should discuss possible limitations of their approach to address problems of privacy and fairness.
- While the authors might fear that complete honesty about limitations might be used by
  reviewers as grounds for rejection, a worse outcome might be that reviewers discover
  limitations that aren't acknowledged in the paper. The authors should use their best
  judgment and recognize that individual actions in favor of transparency play an important role in developing norms that preserve the integrity of the community. Reviewers
  will be specifically instructed to not penalize honesty concerning limitations.

# 3. Theory Assumptions and Proofs

Question: For each theoretical result, does the paper provide the full set of assumptions and a complete (and correct) proof?

Answer: [NA]

Justification: No formal proofs are introduced in this work.

- The answer NA means that the paper does not include theoretical results.
- All the theorems, formulas, and proofs in the paper should be numbered and crossreferenced.
- All assumptions should be clearly stated or referenced in the statement of any theorems.
- The proofs can either appear in the main paper or the supplemental material, but if they appear in the supplemental material, the authors are encouraged to provide a short proof sketch to provide intuition.
- Inversely, any informal proof provided in the core of the paper should be complemented
  by formal proofs provided in appendix or supplemental material.

• Theorems and Lemmas that the proof relies upon should be properly referenced.

#### 4. Experimental Result Reproducibility

Question: Does the paper fully disclose all the information needed to reproduce the main experimental results of the paper to the extent that it affects the main claims and/or conclusions of the paper (regardless of whether the code and data are provided or not)?

Answer: [Yes]

Justification: In the provided anonymous code-base, instructions and prepared files are provided to reproduce the experiments. Checkpoint models are also provided. We further detail the exact GPUs and CPUs we used in the Appendix.

#### Guidelines:

- The answer NA means that the paper does not include experiments.
- If the paper includes experiments, a No answer to this question will not be perceived well by the reviewers: Making the paper reproducible is important, regardless of whether the code and data are provided or not.
- If the contribution is a dataset and/or model, the authors should describe the steps taken to make their results reproducible or verifiable.
- Depending on the contribution, reproducibility can be accomplished in various ways. For example, if the contribution is a novel architecture, describing the architecture fully might suffice, or if the contribution is a specific model and empirical evaluation, it may be necessary to either make it possible for others to replicate the model with the same dataset, or provide access to the model. In general, releasing code and data is often one good way to accomplish this, but reproducibility can also be provided via detailed instructions for how to replicate the results, access to a hosted model (e.g., in the case of a large language model), releasing of a model checkpoint, or other means that are appropriate to the research performed.
- While NeurIPS does not require releasing code, the conference does require all submissions to provide some reasonable avenue for reproducibility, which may depend on the nature of the contribution. For example
- (a) If the contribution is primarily a new algorithm, the paper should make it clear how to reproduce that algorithm.
- (b) If the contribution is primarily a new model architecture, the paper should describe the architecture clearly and fully.
- (c) If the contribution is a new model (e.g., a large language model), then there should either be a way to access this model for reproducing the results or a way to reproduce the model (e.g., with an open-source dataset or instructions for how to construct the dataset).
- (d) We recognize that reproducibility may be tricky in some cases, in which case authors are welcome to describe the particular way they provide for reproducibility. In the case of closed-source models, it may be that access to the model is limited in some way (e.g., to registered users), but it should be possible for other researchers to have some path to reproducing or verifying the results.

#### 5. Open access to data and code

Question: Does the paper provide open access to the data and code, with sufficient instructions to faithfully reproduce the main experimental results, as described in supplemental material?

Answer: [Yes]

Justification: In the provided anonymous code-base, instructions and prepared files are provided to reproduce the experiments. Checkpoint models are also provided.

- The answer NA means that paper does not include experiments requiring code.
- Please see the NeurIPS code and data submission guidelines (https://nips.cc/public/guides/CodeSubmissionPolicy) for more details.

- While we encourage the release of code and data, we understand that this might not be possible, so "No" is an acceptable answer. Papers cannot be rejected simply for not including code, unless this is central to the contribution (e.g., for a new open-source benchmark).
- The instructions should contain the exact command and environment needed to run to reproduce the results. See the NeurIPS code and data submission guidelines (https://nips.cc/public/guides/CodeSubmissionPolicy) for more details.
- The authors should provide instructions on data access and preparation, including how
  to access the raw data, preprocessed data, intermediate data, and generated data, etc.
- The authors should provide scripts to reproduce all experimental results for the new
  proposed method and baselines. If only a subset of experiments are reproducible, they
  should state which ones are omitted from the script and why.
- At submission time, to preserve anonymity, the authors should release anonymized versions (if applicable).
- Providing as much information as possible in supplemental material (appended to the paper) is recommended, but including URLs to data and code is permitted.

#### 6. Experimental Setting/Details

Question: Does the paper specify all the training and test details (e.g., data splits, hyperparameters, how they were chosen, type of optimizer, etc.) necessary to understand the results?

Answer: [Yes]

Justification: Exact details on the model hyperparameters and step-by-step training data pre-processing are provided in the Appendix.

#### Guidelines:

- The answer NA means that the paper does not include experiments.
- The experimental setting should be presented in the core of the paper to a level of detail
  that is necessary to appreciate the results and make sense of them.
- The full details can be provided either with the code, in appendix, or as supplemental material.

### 7. Experiment Statistical Significance

Question: Does the paper report error bars suitably and correctly defined or other appropriate information about the statistical significance of the experiments?

Answer: [Yes]

Justification: Every experiment was run across 10 seeds (0-9 inclusive) and all metrics report the mean and standard deviation. "Best" performance is reported via t-test 95% significance.

- The answer NA means that the paper does not include experiments.
- The authors should answer "Yes" if the results are accompanied by error bars, confidence intervals, or statistical significance tests, at least for the experiments that support the main claims of the paper.
- The factors of variability that the error bars are capturing should be clearly stated (for example, train/test split, initialization, random drawing of some parameter, or overall run with given experimental conditions).
- The method for calculating the error bars should be explained (closed form formula, call to a library function, bootstrap, etc.)
- The assumptions made should be given (e.g., Normally distributed errors).
- It should be clear whether the error bar is the standard deviation or the standard error
  of the mean.
- It is OK to report 1-sigma error bars, but one should state it. The authors should preferably report a 2-sigma error bar than state that they have a 96% CI, if the hypothesis of Normality of errors is not verified.

- For asymmetric distributions, the authors should be careful not to show in tables or figures symmetric error bars that would yield results that are out of range (e.g. negative error rates).
- If error bars are reported in tables or plots, The authors should explain in the text how
  they were calculated and reference the corresponding figures or tables in the text.

#### 8. Experiments Compute Resources

Question: For each experiment, does the paper provide sufficient information on the computer resources (type of compute workers, memory, time of execution) needed to reproduce the experiments?

Answer: [Yes]

Justification: Details on the exact GPUs and CPUs used and also the wall times are presented in the Appendix.

#### Guidelines:

- The answer NA means that the paper does not include experiments.
- The paper should indicate the type of compute workers CPU or GPU, internal cluster, or cloud provider, including relevant memory and storage.
- The paper should provide the amount of compute required for each of the individual experimental runs as well as estimate the total compute.
- The paper should disclose whether the full research project required more compute than the experiments reported in the paper (e.g., preliminary or failed experiments that didn't make it into the paper).

#### 9. Code Of Ethics

Question: Does the research conducted in the paper conform, in every respect, with the NeurIPS Code of Ethics https://neurips.cc/public/EthicsGuidelines?

Answer: [Yes]

Justification: We conform with the NeurIPS Code of Ethics as detailed in the URL.

#### Guidelines:

- The answer NA means that the authors have not reviewed the NeurIPS Code of Ethics.
- If the authors answer No, they should explain the special circumstances that require a deviation from the Code of Ethics.
- The authors should make sure to preserve anonymity (e.g., if there is a special consideration due to laws or regulations in their jurisdiction).

#### 10. Broader Impacts

Question: Does the paper discuss both potential positive societal impacts and negative societal impacts of the work performed?

Answer: [Yes]

Justification: We add a section after the Conclusion to discuss Broader Impacts.

- The answer NA means that there is no societal impact of the work performed.
- If the authors answer NA or No, they should explain why their work has no societal
  impact or why the paper does not address societal impact.
- Examples of negative societal impacts include potential malicious or unintended uses (e.g., disinformation, generating fake profiles, surveillance), fairness considerations (e.g., deployment of technologies that could make decisions that unfairly impact specific groups), privacy considerations, and security considerations.
- The conference expects that many papers will be foundational research and not tied to particular applications, let alone deployments. However, if there is a direct path to any negative applications, the authors should point it out. For example, it is legitimate to point out that an improvement in the quality of generative models could be used to generate deepfakes for disinformation. On the other hand, it is not needed to point out that a generic algorithm for optimizing neural networks could enable people to train models that generate Deepfakes faster.

- The authors should consider possible harms that could arise when the technology is being used as intended and functioning correctly, harms that could arise when the technology is being used as intended but gives incorrect results, and harms following from (intentional or unintentional) misuse of the technology.
- If there are negative societal impacts, the authors could also discuss possible mitigation strategies (e.g., gated release of models, providing defenses in addition to attacks, mechanisms for monitoring misuse, mechanisms to monitor how a system learns from feedback over time, improving the efficiency and accessibility of ML).

#### 11. Safeguards

Question: Does the paper describe safeguards that have been put in place for responsible release of data or models that have a high risk for misuse (e.g., pretrained language models, image generators, or scraped datasets)?

Answer: [NA]

Justification: The released models and case studies are not directly applicable for malicious misuse. Focus is on therapeutics design.

#### Guidelines:

- The answer NA means that the paper poses no such risks.
- Released models that have a high risk for misuse or dual-use should be released with necessary safeguards to allow for controlled use of the model, for example by requiring that users adhere to usage guidelines or restrictions to access the model or implementing safety filters.
- Datasets that have been scraped from the Internet could pose safety risks. The authors should describe how they avoided releasing unsafe images.
- We recognize that providing effective safeguards is challenging, and many papers do
  not require this, but we encourage authors to take this into account and make a best
  faith effort.

# 12. Licenses for existing assets

Question: Are the creators or original owners of assets (e.g., code, data, models), used in the paper, properly credited and are the license and terms of use explicitly mentioned and properly respected?

Answer: [Yes]

Justification: A license is provided in the anonymous repository. In the code-base, some code was adapted from existing code-bases which are licensed under Apache 2.0. For these code files, we explicitly state where the code was adapted from.

### Guidelines:

- The answer NA means that the paper does not use existing assets.
- The authors should cite the original paper that produced the code package or dataset.
- The authors should state which version of the asset is used and, if possible, include a URL.
- The name of the license (e.g., CC-BY 4.0) should be included for each asset.
- For scraped data from a particular source (e.g., website), the copyright and terms of service of that source should be provided.
- If assets are released, the license, copyright information, and terms of use in the
  package should be provided. For popular datasets, paperswithcode.com/datasets
  has curated licenses for some datasets. Their licensing guide can help determine the
  license of a dataset.
- For existing datasets that are re-packaged, both the original license and the license of the derived asset (if it has changed) should be provided.
- If this information is not available online, the authors are encouraged to reach out to the asset's creators.

#### 13. New Assets

Question: Are new assets introduced in the paper well documented and is the documentation provided alongside the assets?

	Α	F <b>X</b> 7
1676	Answer:	res

1677

1678

1679

1680

1681

1682

1683

1684 1685

1686

1687

1688

1689

1690

1691

1692

1693

1694

1695

1696

1697

1698

1699

1700

1701

1702

1703

1704

1705

1706

1707

1708

1709

1710

1711

1712

1713

1714

1715

1716

1717

1718

1719

1720

Justification: The released anonymous code is licensed under Apache 2.0.

#### Guidelines:

- The answer NA means that the paper does not release new assets.
- Researchers should communicate the details of the dataset/code/model as part of their submissions via structured templates. This includes details about training, license, limitations, etc.
- The paper should discuss whether and how consent was obtained from people whose asset is used.
- At submission time, remember to anonymize your assets (if applicable). You can either create an anonymized URL or include an anonymized zip file.

# 14. Crowdsourcing and Research with Human Subjects

Question: For crowdsourcing experiments and research with human subjects, does the paper include the full text of instructions given to participants and screenshots, if applicable, as well as details about compensation (if any)?

Answer: [NA]

Justification: Neither crowdsourcing nor human subjects were involved.

#### Guidelines:

- The answer NA means that the paper does not involve crowdsourcing nor research with human subjects.
- Including this information in the supplemental material is fine, but if the main contribution of the paper involves human subjects, then as much detail as possible should be included in the main paper.
- According to the NeurIPS Code of Ethics, workers involved in data collection, curation, or other labor should be paid at least the minimum wage in the country of the data collector.

# 15. Institutional Review Board (IRB) Approvals or Equivalent for Research with Human Subjects

Question: Does the paper describe potential risks incurred by study participants, whether such risks were disclosed to the subjects, and whether Institutional Review Board (IRB) approvals (or an equivalent approval/review based on the requirements of your country or institution) were obtained?

Answer: [NA]

Justification: Neither crowdsourcing nor research with human subjects were involved.

- The answer NA means that the paper does not involve crowdsourcing nor research with human subjects.
- Depending on the country in which research is conducted, IRB approval (or equivalent) may be required for any human subjects research. If you obtained IRB approval, you should clearly state this in the paper.
- We recognize that the procedures for this may vary significantly between institutions and locations, and we expect authors to adhere to the NeurIPS Code of Ethics and the guidelines for their institution.
- For initial submissions, do not include any information that would break anonymity (if applicable), such as the institution conducting the review.