1 L-MolGAN: An improved implicit generative model for

2 generation of large molecular graphs

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8Abstract

9Deep generative models are used to generate arbitrary molecular structures with the desired 10chemical properties. MolGAN is a renowned molecular generation models that uses 11generative adversarial networks (GANs) and reinforcement learning to generate molecular 12graphs in one shot. MolGAN can effectively generate a small molecular graph with nine or 13fewer heavy atoms. However, the graphs tend to become disconnected as the molecular size 14increase. This poses a challenge to drug discovery and material design, where large molecules 15are potentially inclusive. This study develops an improved MolGAN for large molecule 16generation (L-MolGAN). In this model, the connectivity of molecular graphs is evaluated by 17a depth-first search during the model training process. When a disconnected molecular graph

18is generated, L-MolGAN rewards the graph a zero score. This procedure decreases the 19number of disconnected graphs, and consequently increases the number of connected 20molecular graphs. The effectiveness of L-MolGAN is experimentally evaluated. The size and 21connectivity of the molecular graphs generated with data from the ZINC-250k molecular 22dataset are confirmed using MolGAN as the baseline model. The model is then optimized for 23a quantitative estimate of drug-likeness (QED) to generate drug-like molecules. The 24experimental results indicate that the connectivity measure of generated molecular graphs 25improved by 1.96 compared with the baseline model at a larger maximum molecular size of 2620 atoms. The molecules generated by L-MolGAN are evaluated in terms of multiple 27chemical properties, QED, synthetic accessibility, and log octanol—water partition coefficient, 28which are important in drug design. This result confirms that L-MolGAN can generate 29various drug-like molecules despite being optimized for a single property, i.e., QED. This 30method will contribute to the efficient discovery of new molecules of larger sizes than those 31being generated with the existing method.

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33**Keywords:** deep learning, generative adversarial network, graph convolutional network, 34molecular graph

35**1. Introduction**

36Machine learning-based molecular design of drugs is used to efficiently determine the desired 37molecular structure in drug discovery. It also aids the automated search for unknown 38molecular structures of the desired properties and predict their physical properties without 39requiring the domain knowledge of organic chemistry. A renowned classical molecular design 40model is inverse quantitative structure-activity relationship (inverse-QSAR) [1]. Based on the 41QSAR model—an analytical model of the relationship between molecular structure and 42bioactivity, formulated using molecular descriptors quantifying the features of the molecular 43structure—inverse-QSAR performs a backward prediction of the molecular structure from the 44desired bioactivity. Therefore, to obtain a molecular structure with the desired bioactivity, it is 45necessary to select the appropriate molecular descriptors that are equivalent to the raw data of 46feature engineering in machine learning. However, it is difficult to identify the descriptors 47correlated with the desired bioactivity from the numerous available molecular descriptors, 48which is a core problem in inverse-QSAR analysis. 49Several molecular-structure search methods based on deep generative models, which generate 50new data with similar features as the original without the availability of predetermined 51feature vectors for the dataset, have been proposed and developed. Most adopt a graph-based 52approach in which the molecular structure is represented as a graph and are classified into 53two approaches in terms of the molecular generation process: sequential iterative process and

54one-shot generation [2].

55In the sequential iterative process, molecules are assembled stepwise by adding atoms and 56bonds to a predefined scaffold. The advantage of the generative model [3–6] when combined 57with the sequential iterative process is the assurance of chemical validity of the generated 58molecules. Thus, it is possible to obtain functional molecules by reliably generating larger 59molecules. However, the disadvantage of the sequential iterative process is the increased 60computational cost of verifying the valence, topological prediction of molecular structure, 61and graph isomorphism to calculate the reconstruction error when iteratively assembling 62molecules.

63In one-shot generation, a molecule is generated by determining the combination of atoms and 64bonds in a single step. The advantage of the generative model combined with one-shot 65generation [2,7–10] is the simplicity of its architecture and algorithm. Its computational cost 66is smaller than the sequential iterative process. Consequently, the generative model can be 67optimized in a short time. However, the one-shot generation method can only generate small 68molecular graphs because the number of possible connections between atoms in larger 69molecules increases quadratically, increasing the likelihood of the generation of chemically 70invalid molecules [8].

71One of the most successful generative models using the one-shot generation scheme is the 72molecular generative adversarial network (MolGAN) [7]. MolGAN generates small

73molecular graphs with the desired chemical properties by combining GANs [11] and 74reinforcement learning. It can generate chemically valid molecules if the number of heavy 75atoms used for molecular representation is nine or fewer. However, when this number 76exceeds nine, many disconnected molecular graphs are generated.

77To overcome this issue, we propose a large MolGAN (L-MolGAN), an improved version of 78the MolGAN model, for generating larger, more connected molecular graphs. Increasing the 79number of connected molecular graphs in MolGAN will lead to the rapid generation of large 80molecular graphs. We integrated into L-MolGAN a mechanism that enhances the generation 81of connected graphs in the generative process of MolGAN. The first stage of the model 82judges if the generated molecular graph is connected or disconnected. If the graph is 83disconnected, it will be penalized during model training. Consequently, the generation of 84disconnected molecular graphs is suppressed in the model optimization process.

85The contributions of this study are:

disconnections.

- 1. An improved MolGAN that produces large (up to 20 atoms), novel molecules without
- 88 2. A molecular graph expansion mechanism that penalizes, and consequently suppresses,
- the production of disconnected graphs.

90The remainder of this paper is organized as follows. Section 2 presents an overview of the 91proposed L-MolGAN and a method to represent the molecular graph and the framework of 92the original MolGAN. A method to improve the connectivity of molecular graphs generated

93by L-MolGAN is also described in this section. Then, in Sections 3 and 4, the effectiveness of 94the proposed model is validated by comparing its performance in generating new molecules 95with that of the original MolGAN using a publicly available dataset of drug-like molecules. 96Finally, the paper is concluded in Section 5.

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982. Method

992.1 Model Architecture

100MolGAN, which is the baseline model, consists of GANs (generator and discriminator) and a 101reward network. In this model, the molecular structure is represented by a graph. The L-102MolGAN adds a mechanism called molecular graph expansion to the baseline model that 103increases the number of generated connected molecular graph. The model architecture of L-104MolGAN is illustrated in Fig 1. The L-MolGAN differs from the original MolGAN only in 105terms of the molecular graph expansion mechanism, highlighted by the colored box in the 106figure.

107

108Fig 1 Model architecture of L-MolGAN for generating large molecular graph. It consists 109of a generator, a discriminator, a reward network, and a molecular graph expansion 110mechanism. Molecular graphs are generated by inputting into the generator vectors sampled

111from a prior distribution. The discriminator classifies the input molecular graph into 112generator-produced or dataset. The reward network predicts the chemical properties of the 113input molecular graph.

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115GANs were used to learn the molecular features of the training dataset, and the reward 116network was trained to predict the chemical properties of the given molecular graph. A multi-117layer perceptron (MLP) was adopted for all three components, the generator, the 118discriminator, and the reward network, similar to the baseline study by De Cao et al. [7]. In 119the following subsections, we shall explain the molecular representation and each network 120model, as well as the proposed modifications to the baseline model.

121**2.2 MolGAN**

1222.2.1 Molecular representation as a graph

123Studies related to the artificial generation of molecules using deep generative models [12–14] 124represented molecules as strings using the simplified molecular-input line-entry system 125(SMILES) [15]. The linear SMILES is in turn generated string using a recurrent neural 126network and long short-term memory. Thus, the molecule of interest was artificially 127produced.

128However, the inherent syntax of SMILES is complex, and the chemical structure and 129properties of a molecule can vary drastically with the order of the string and changes in a

130single character. In addition, the same molecule has multiple string representations, making it 131impossible to determine a unique SMILES [16, 17]. To avoid these problems, researchers 132have developed molecular graphs that represent molecules based on the graph theory. 133Molecular graphs are an intuitive, more robust representation of molecules compared with 134intermediate representations such as SMILES. In this study, the molecules were treated as 135labeled undirected graphs. A molecular graph was defined as G=(V,E), where E and V 136denote a set of edges and nodes, respectively. Each atom and each bond that make up a 137molecule correspond to a node $v_i \in V$ and an edge $(v_i, v_i) \in E$, respectively. The molecular 138graph consists of two types of matrix: the node feature matrix and the adjacency matrix. The 139node v_i ∈ V in the molecular graph G was defined by the one-hot vector x_i in T dimensions, 140where *T* represents the number of types of atoms. From this vector, the type of atom, which is 141an attribute of node v_i , can be determined. The node feature matrix is represented by 142aggregating all node feature vectors. The edge $(v_i, v_i) \in E$ in the molecular graph G indicates 143that nodes v_i and v_j are connected. In addition to the connections between nodes, the type of 144bond $y \in [1,...,Y]$ is considered in the molecular graph, where *Y* is the number of bond types.

145In this study, the node feature matrix $X = [x_1, ..., x_N]^T \in \mathbb{R}^{N \times T}$ and the adjacency matrix

146 $_{A \in \mathbb{R}^{N \times N \times Y}}$ were used to identify the types of atoms in all node sets of the molecular graph 147 $_{G}$ and the adjacency matrix.

1482.2.2 Generative adversarial networks

149GANs are deep generative models that aim to generate samples similar to a training set by 150approximating the model distribution to an empirical distribution. In computational molecular 151design, adversarial generation is an important strategy for producing molecular species 152similar to a given molecular dataset.

153GANs can be interpreted as an implicit generative model as it does not need assume a 154specific probability distribution for the model distribution when approximating the empirical 155distribution. This eliminates the need for an explicit likelihood function for approximating the 156probability distribution. On the one hand, the variational autoencoder (VAE) [18], a 157likelihood-based model, adopts a method to approximate the empirical and model 158distributions by assuming in advance the latter to be Gaussian and maximizing the evidence 159lower bound instead. On the other hand, GANs adopt a method to approximate the model **160**distribution to the empirical distribution by parameterizing the distribution with a deep neural 161network and estimating its density ratio. GANs mainly consist of two deep neural networks to 162approximate the distribution by density ratio estimation: generator G_{θ} , generates a new **163**sample $G(z;\theta)$ similar to the training sample x p_{data} by inputting a random number z p_z **164**obtained from a prior distribution p_z ; discriminator D_ω , which accurately identifies the input **165**data as a training sample x p_{data} or a sample $G(z;\theta)$ generated by the generator. Training **166**generator G_{θ} to generate samples similar to the empirical distribution means will yield worse

167identification results for the samples produced by the generator. In other words, the density 168ratio estimation problem is replaced by a classification problem, which can be effectively 169solved by deep neural networks are good. Therefore, these deep neural networks can be 170considered players in the minimax game of Equation 1, which shows the expected value of 171the cross-entropy error.

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$$173 \min_{\theta} \max_{\sigma} E_{x \ p_{dow}(x)} [\log D_{\varphi}(x)] + E_{z \ p_{z}(z)} [\log (1 - D_{\varphi}(G_{\theta}(z)))] \tag{1}$$

174

175In adversarial learning, the generator is trained to generate samples similar to the training set 176and misidentify them to the discriminator. In contrast, the discriminator is trained to correctly 177discriminate between the samples generated by the generator and those from the training set. 178With this process, the two models coevolve in adversary, with the generator minimizing the 179second term in Equation 1 and the discriminator maximizing the linear sum of the first and 180second terms. The alternate optimization the two neural networks through back-propagation, 181a sample is eventually generated such that the discriminator cannot distinguish between real 182and fake samples.

1832.2.3 Generator

184The generator can generate molecular graphs with the desired chemical properties. In this 185study, its architecture was a simple MLP with four layers. The number of units in each layer

186was 256, 512, 1024, and 2200, respectively. By inputting a random number z sampled from 187the standard normal distribution N(0,I) into the generator, we output the adjacency matrix \widetilde{A} 188and the node feature matrix \widetilde{X} representing the molecular graph. The output graph $\widetilde{G} = (\widetilde{A}, \widetilde{X})$ 189is a probabilistic complete graph, which is interpreted as a categorical distribution for the 190types of atoms and bonds. Here, \widetilde{A} contains the existence probabilities of the nodes and edges 191for each bond type, and \widetilde{X} the class probabilities of the nodes. To enable its transformation 192into a chemically valid molecular graph, the discrete graph G = (A, X) was obtained using the 193argmax function on the output probabilistic complete graph $\widetilde{G} = (\widetilde{A}, \widetilde{X})$. The adjacency matrix

194was defined as $A \in \mathbb{R}^{N \times N \times Y}$, and the node feature matrix as $X = [x_1, ..., x_N]^T \in \mathbb{R}^{N \times T}$.

195The maximum number of nodes in the molecular graph was set to N = 20, and the number of 196bond types to Y = 5. The five types of bonds are single bond, double bond, triple bond, 197aromatic bond, and no bond. The number of types of atoms was set to T = 10: carbon, 198nitrogen, oxygen, fluorine, phosphorus, sulfur, chlorine, bromine, and iodine, and one-199padding symbol. Thus, the maximum number and types of atoms and bond types were 200restricted. These constraints shall be used to determine the generator architecture. The 201dimensions of the output adjacency and output node feature matrices were represented by $202N \times N \times Y$ (i.e., $20 \times 20 \times 5$) and $N \times T$ (i.e., 20×10), respectively.

203To output the two types of matrices simultaneously, an output layer is required to output the

2042200-dimensional vector, which is the sum of the number of elements of the adjacency and 205 node feature matrices. The number of units in the output layer depends on these constraints. 206The random number inputs to the generator had 256 dimensions. Based on the results of 207existing research, the number of units in each hidden layer was set as a multiple of the 208number of dimensions of the input random numbers. The 2200-dimensional vector output 209from the generator was split into two vectors—2000- and 200-dimensional vectors—to create 210the adjacency and node feature matrices. These divided vectors were then transformed into 211the dimensionality of each defined matrix. Consequently, the output molecular graph is a 212complete probabilistic graph. 213The final output molecular graph is a chemically valid molecular graph. Therefore, the 214argmax function was used to break the weak bonds in the complete graph. The output of this 215operation on the adjacency matrix \widetilde{A} is the adjacency matrix A binarized at [0,1]. The node

214argmax function was used to break the weak bonds in the complete graph. The output of this 215operation on the adjacency matrix \widetilde{A} is the adjacency matrix A binarized at [0,1]. The node 216feature matrix \widetilde{X} was also binarized using the same process. Finally, a new molecular graph 217with the correct valence was generated through the optimized molecular generation process. 218However, this adversarial generation process only generates molecular species similar to the 219training set. Moreover, it is necessary to introduce methods to improve the properties of the 220generated molecules such as reinforcement learning, which uses a deterministic policy in the 221process of molecule generation. We incorporated the deep deterministic policy gradient 222method [19] into a generative model to optimize the non-differentiable chemical indices

223based on the literature. The stochastic policy is expressed as $\pi(a \vee s; \theta)$. This denotes the 224policy π_{θ} that probabilistically selects action a for state s. In this case, θ is a parameter used 225when the policy is being modeled. The deterministic policy μ_{θ} is the policy $a = \mu_{\theta}(s)$, where 226action a is uniquely determined for a certain state s. This policy is optimized by updating θ to 227maximize the behavioral value function for this behavior. In this study, the policy was G, and 228state s was represented as a random number z. Thus, for a random number z, the molecular 229graph is uniquely generated according to the deterministic policy. In the deep deterministic 230policy gradient method, the deterministic policy and action value functions were 231approximated using a deep neural network. Therefore, a property prediction neural network, 232which can be trained using gradients, was introduced into the action value function for 233calculating rewards. The rewards can then be used to generate molecules with 234indistinguishable chemical properties. These properties can be maximized by varying the 235policy parameters in the direction of the approximated action value gradient. 236By formulating these series of processes, we trained the generator such that the objective 237function $L(\theta)$ in Equation 2 was minimized. A molecular graph with the desired chemical

238properties similar to the training data was generated by minimizing the linear combination of

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$$L(\theta) = \lambda \cdot L_{RL} + (1 - \lambda) \cdot L_{GAN}, \qquad (2)$$

239the GAN loss, L_{GAN} , and the reinforcement learning loss, L_{RL} :

243where λ is a hyperparameter that balances between adversarial learning and property 244optimization. This tunable parameter takes values in the range of $\lambda \in [0,1]$.

2452.2.4 Discriminator and Reward network

246The architecture of these two neural networks, discriminator and reward network, were 247implemented by a simple MLP with three layers. The three hidden layers of both the 248discriminator and the reward network had 512, 256, and 2 units, respectively. The input 249molecules were discriminated by the discriminator as the training set or molecules sampled 250by the generator. The chemical properties of the input molecules were predicted using the 251reward network. In the generation process, the discriminator outputs the discrimination rate 252of authenticity based on the feature vector of the entire molecular graph, and the reward 253network outputs the predicted score of the chemical property. The generator can be optimized 254by feeding back the outputs.

255However, a simple MLP cannot directly handle the graph structure data. Therefore, it would 256be necessary to develop a graph convolution operation specific to the graph structure data 257before inputting molecular graphs into the two models. The type of bond between atoms must 258be considered when convoluting the molecular graph. Therefore, based on the literature, we 259used a relational graph convolution operation that considers the attributes of the edges on a 260graph [20, 21]. This operation uses the adjacency matrix to convolute the node information

261 for each edge attribute as follows:

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263
$$h_i^{(l+1)} = \sigma\left(\sum_{r \in R} \sum_{j \in N_i^r} \frac{1}{|N_i|} W_r^{(l)} h_j^{(l)} + W_0^{(l)} h_i^{(l)}\right)$$
(3)

264

265where h_i^l is the feature representation of node v_i in the lth layer, R is the set of relations, and $266N_i^r$ is the set of nodes connected by the relation r in node v_i . Thus, a linear transformation was 267performed by extracting the neighboring node information for each relation. The self-loop 268was convolved similarly.

269Finally, a nonlinear transformation was performed over the input signal by the activation 270function σ , and the feature representation $h_i^{(l+1)}$ of the l+1st layer was output. The convolution 271of a node uses its own information as well as information from its neighboring nodes. The 272output of the hidden layer was recursively used same as in a neural network by accumulating 273the convolutions. Finally, each convolved node information was aggregated into a single 274feature representation. Each time the convolution operation was repeated, the neighboring 275node information was convolved; thus, a global feature representation revealing the entire 276graph was obtained from the local features.

277The generator and discriminator were used to facilitate the adversarial learning of the 278molecular generation model. The discriminator was trained to maximize Equation 1. The 279parameters of the generator was updated via backpropagation through the discriminator to the

280generator. The generator and the reward network were used to optimize the chemical 281properties of molecules. The generator used the reward network output as a reward, and the 282parameters of the two models were updated using the deep deterministic policy gradient 283method. In addition, the reward network was trained by back-propagating the error between 284the output of the reward network and the estimated property using RDKit [22], a 285chemoinformatics tool. Reinforcement learning for chemical properties optimization was 286performed once for every three iterations of adversarial learning. The parameters of the 287reward network were fixed in adversarial learning, whereas the parameters of the 288discriminator are fixed in the chemical properties optimization.

2892.2.5 Molecular graph generation using the trained MolGAN

291new molecules by inputting random numbers sampled from the standard normal distribution 292into the generator. Changes in these numbers resulted in different molecular graphs. This 293allowed the generator to generate not only known molecules but also unknown ones included 294in the training dataset.

2952.3 L-MolGAN and Molecular Graph Expansion Mechanism

296According to the literature [7, 8], the number of nodes in the generated molecular graph is 297small, which is the problem we aim to solve. Earlier studies evaluated the MolGAN under the 298condition that only nine heavy atoms can be used to produce a molecular graph without

299disconnection. However, this limit is not practical in drug discovery, especially for larger 300molecules because the more the atoms, the more the disconnected graphs. To solve this 301problem, we propose modifications to the MolGAN.

302We suppress the generation of disconnected graphs by penalizing them during the model 303training process. The detailed algorithm is as follows: 1) for each generated graph, its 304connectivity is checked by depth-first search (DFS) and 2) if the graph is disconnected, its 305chemical property score is set to zero as a penalty; otherwise, its score is predicted by the 306reward network. This is similar to the general training process. DFS is a recursive and 307exhaustive algorithm used to search all nodes of a graph or a tree. With DFS, the entire graph 308is traversed by starting at a certain node in the molecular graph and following the edges. If all 309the nodes in the graph can be reached, the graph is considered connected. Repeated 310penalizations to a disconnected graph will suppress its generation and increase the number of 311connected graphs generated. We refer to these modifications as the "molecular graph 312expansion mechanism," and rename the resulting improved MolGAN as L-MolGAN.

313**3. Experiment**

314We shall investigate the effectiveness of the L-MolGAN by comparing it with the baseline 315MolGAN. In all experiments, we set the QED as a singular objective to derive new drug 316candidates and trained two modes (i.e., the model training was performed to optimize QED 317score with an RL objective). Its effectiveness was evaluated in terms of 1) how well it works

318for large molecular graph generation and 2) how many novel drug-like molecules it 319generates. The general settings of the model training and its evaluation metrics shall be 320described in Subsections 3.1 to 3.3. Then, three different numerical experiments shall be 321described in the Subsections 3.4 to 3.6.

3223.1 Dataset

323In this study, ZINC-250k [23], a renowned molecular datasets, was used in the experiments to 324generate molecular graphs. ZINC-250k is made up of 250 000 commercial drug-like 325molecules randomly selected from the ZINC database. The maximum number of constituent 326heavy atoms of a molecule in ZINC-250k is 38. Particularly, a subset of ZINC-250k was 327sampled by randomly choosing 15 000 molecules from ZINC-250k, with the maximum 328number of constituent heavy atoms limited to 20, which is approximately twice the molecular 329size of that used in the baseline study by De Cao et al. [7].

330**3.2 Evaluation metrics**

331We employed the generally used indices of validity, novelty, and uniqueness to evaluate the 332molecular generation model. Validity is the percentage of chemically valid molecules among 333the generated molecules. Note that validity is not a measure of the connectivity of molecules 334but only the valence of atoms. Novelty is the percentage of valid molecules among the 335generated molecules not included in the training data. In this study, these molecules were

336defined as novel molecules. Uniqueness is the percentage of generated molecules that are 337valid as well as unique. This measure indicates the degree of diversity among the molecules 338generated. Furthermore, species, the number of unique and connected molecular graphs, was 339introduced to clearly represent the number of unique molecules that were derived. The ideal 340molecular generation model should generate novel, valid, and connected molecules. 341Additionally, connectivity, which is the percentage of connected graphs, is one of the most 342important metrics introduced in this study. It indicates the percentage of valid and connected 343molecular graphs among the ones generated. 344Furthermore, three chemical indicators were used to evaluate the chemical properties of the 345generated molecules, QED [24], solubility, and synthetic accessibility (SA) score. In QED, 346drug-like properties were calculated using a weighted geometric mean based on the 347distribution of multiple drug-properties data. Solubility indicates the degree of hydrophilicity 348of a molecule, which was quantified by the logP coefficient. This coefficient is defined as the 349logarithm of the concentration ratio of different solvents [25]. The SA score indicates the ease 350of synthesis of a molecule [26]. In this experiment, all chemical-property scores were

353**3.3 Model training**

354MolGAN and L-MolGAN were trained using the Adam optimizer [27] with a learning rate of

351manipulated to take values in the range of [0,1]. Note that the property scores of molecules

352with disconnected graphs in the L-MolGAN were set to zero as a penalty.

3550.0001 to optimize the QED for all the experiments. Mini-batch training was conducted to 356stabilize the learning. The batch size was set to 100. With an early stopping strategy, the 357model training was terminated when the average change in loss during 10 epochs was less 358than 1.0% or when the maximum number of epochs (300) had been reached. Mode collapse 359[28], a situation where similar data are generated regardless of the arbitrariness of numbers 360input to the generator, is one of the crucial issues in GANs. To circumvent this issue, we used 361mini-batch training and the early stopping strategy mentioned above.

362In another study [7], researchers terminated model training when the uniqueness score fell

362In another study [7], researchers terminated model training when the uniqueness score fell 363below 2.0%. However, this cause the generated molecules to become more homogenous 364because several epochs would be solely dedicated to satisfying the termination criterion. 365Therefore, we focused on the average loss change in the training process to determine the 366termination criterion, rather than thresholding for each property score.

Experiment I: Parameter study of learning balance

369Properties optimization and adversarial learning, affects the characteristics of the generated 370molecules. The optimal choice of λ for the molecular generation model was determined 371through this experimental task. The value of λ was varied from 0.0 to 1.0 in increments of 3720.2, and the model was trained in five trials for each value.

373As explained earlier, we optimized the QED to generate drug-like molecules. The trained

374model that maximized the sum of validity, novelty, uniqueness, and QED scores was selected 375as the reference to evaluate the performance of the molecular generation model.

3763.5 Experiment II: Performance comparison of proposed

377 method with existing method

378The proposed method and the baseline model (MolGAN) were compared in terms of 379performance using the evaluation metrics described in Subsection 3.2. The representative 380model for each method was chosen through a parametric study of λ .

3813.6 Experiment III: Generation and evaluation of novel

382 **molecules**

383The proposed method was evaluated in terms of the number of novel drug-like molecules that 384can be derived. Here, new molecules were generated by inputting into the pretrained 385generator random numbers sampled 5000 times from the standard normal distribution. The 386chemical properties of the generated molecules were evaluated using RDKit. Novel 387molecules with the desired chemical properties were identified from the generated molecules. 388The 20 molecules with the highest QED scores were chosen, and their chemical properties 389were examined in terms of novelty and ease of synthesis.

390In addition, in drug discovery and materials design, the generated molecules should not only 391satisfy a single property (such as the QED) but also possess other properties such as 392synthesizability. Therefore, SA and logP were chosen in addition to the QED to evaluate the 393molecules generated in this experiment. However, there is a tradeoff between QED and logP 394[29]. There is no single best molecule but several 'good' molecules that exist within the 395envelope of all the generated molecules. Here, we refer to them as 'dominant molecules' and 396chose them in terms of the three chemical properties, QED, SA, and logP, for each 397combination of two of the three properties. Furthermore, we classified them into hydrophiles 398and lipophiles based on the logP score and verified if the dominant molecules possessed both 399the properties. Both hydrophilicity and lipophilicity are important properties considered in 400drug design.

4014. Results and Discussion

402**4.1** Experiment I: Parametric study of learning balance λ

403The effect of λ on the molecules generated by the L-MolGAN are summarized in Table 1. 404The table indicates that the mean value of validity increases with an increase in λ , while its 405standard deviation decreases. The same tendency was observed for the connectivity and QED 406scores. In contrast, the uniqueness decreased as λ increased. We believe this could have been 407caused by the occurrence of mode collapse in the model training, as reported by a

408conventional study [7]. Consequently, as λ increases, the generative model becomes more 409susceptible to mode collapse. This would also affect the diversity of the generated molecules, 410as evidenced by the decrease in the standard deviation of every molecular property index with 411an increase in λ .

Table 1 Comparison of properties of molecules generated at different λ **by the proposed** 414**method.** Each value indicates the mean and the standard deviation of each metric for five 415trials.

	Validity	Uniquen	Novelty	Connecti				
λ					QED	SA	logP	Species
	[%]	ess [%]	[%]	vity [%]				
0.0	28.62 ±	19.66 ±	100.00 ±	59.82 ±	$0.62 \pm$	0.29 ±	$0.54 \pm$	72.60 ±
	6.43	6.53	0.00	15.5	0.03	0.05	0.05	41.67
0.2	80.72 ±	8.46 ±	100.00 ±	85.57 ±	$0.77 \pm$	$0.21 \pm$	$0.59 \pm$	138.00 ±
	2.71	2.40	0.00	9.10	0.04	0.11	0.06	41.74
0.4	94.70 ±	4.10 ±	100.00 ±	95.26 ±	$0.82 \pm$	$0.12 \pm$	$0.60 \pm$	72.60 ±
	3.34	4.92	0.00	4.92	0.02	0.05	0.04	41.67
0.6	95.00 ±	3.21 ±	100.00 ±	94.85 ±	$0.85 \pm$	0.16 ±	$0.62 \pm$	76.80 ±
	3.31	1.09	0.00	5.58	0.03	0.07	0.03	20.98
0.8	98.79 ±	0.30 ±	100.00 ±	99.81 ±	$0.82 \pm$	$0.10 \pm$	$0.55 \pm$	10.20 ±
	1.88	0.11	0.00	0.24	0.05	0.06	0.08	5.12
1.0	96.79 ±	0.04 ±	100.00 ±	100.00 ±	$0.86 \pm$	$0.16 \pm$	$0.55 \pm$	1.60 ±
	0.21	0.02	0.00	0.0	0.05	0.09	0.11	0.89

417The average QED score from the training data was 0.76 ± 0.12 . The QED score of the newly 418generated molecules for $\lambda=0.0$ was smaller than the average. The scores with other λ values 419were greater than the average. Validity, connectivity, and QED for $\lambda=0.0$ were remarkably 420smaller than those of the other settings. The generator should be trained not only to improve 421the chemical property score, but also to suppress the generation of invalid molecules during 422the optimization. However, at $\lambda=0.0$, the model training was completely dedicated to the 423adversarial learning of the generative model, rather than the chemical properties optimization. 424It is plausible that the overall performance at $\lambda=0.0$ was the weakest because of the 425generation of several invalid graphs. 426From these results, we chose $\lambda=0.6$ as the optimal value, which maximizes the total values 427of all the considered evaluation metrics. Furthermore, a single representative model, which

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431Fig 2 Distribution of QED scores of molecules generated by the proposed method at the 432different λ . The black solid line labeled as 'ZINC subset' indicates the distribution of 433molecules included in the training dataset. Kernel density estimation has been used to depict 434the QED distribution. Only the connected molecular graphs are used for the density 435estimation.

428had the largest total value among the five trials for $\lambda = 0.6$, was chosen. This setting shall be

429used for all subsequent experiments with MolGAN as well as L-MolGAN.

437Fig 2 shows the distribution of QED scores of molecules generated by the proposed method, 438estimated using the best-performing generative model for each λ value. Additionally, 'ZINC 439subset' indicates the distribution of molecules included in the training dataset. 440We shall focus on the peak of each distribution to determine the effect of λ . Because each 441distribution has multiple peaks, we shall focus only on the highest one. As λ increases, the 442peak shifts to a higher QED score. Specifically, when λ was larger than 0.4, the peak shifted 443to a higher QED score than that of the training data. From the fact that the model was trained 444to maximize the QED score, we confirmed that the model was well-optimized. Note that a 445narrower distribution was obtained owing to the mode collapse with a larger λ .

4464.2 Experiment II: Performance comparison of the proposed

447 **method with existing methods**

448Table 2 lists the results of the performance comparison between MolGAN and L-MolGAN in 449generating molecules. The results confirm the validity, connectivity, and QED scores of the 450L-MolGAN were better than those of the MolGAN.

452Table 2 Comparison of molecules generated by MolGAN and L-MolGAN

Model Val	lidity Uniqueness	Novelty Connectivity	OED	SA logP	Species
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	[%]	[%]	[%]	[%]				
MolGAN	94.53	5.97	100.00	48.12	0.85	0.47	0.60	44.00
L-MolGAN	98.91	4.88	100.00	94.32	0.88	0.23	0.66	88.00

454Particularly, connectivity of graphs saw an improvement of 1.96 times in L-MolGAN over 455the existing model. However, the uniqueness and SA of L-MolGAN were worse than those of 456MolGAN. This indicates that the molecules generated by MolGAN were more diverse than 457L-MolGAN and are relatively easy to synthesize. However, the lower connectivity score of 458MolGAN indicates the presence of several disconnected molecules in the generated 459molecules. In this regard, the L-MolGAN is more effective in generating valid as well as 460connected molecules than the MolGAN.

461In addition, the lower SA score in the L-MolGAN suggests that the model generates 462molecular graphs with more complex molecular structures because they have mostly 463connected nodes. Both models generated entirely novel molecules that did not exist in the 464training data; however, MolGAN had a connectivity score of 48.12%. In contrast, the L-465MolGAN achieved a higher connectivity score of 94.32%. These results suggest that the 466proposed model generates larger, more novel drug-like molecules, and has more practical 467implications for drug discovery compared with the existing method.

469Fig 3 Distributions of the QED score in the molecules generated by the representative 470models of MolGAN and L-MolGAN. Kernel density estimation has been used to depict the

471QED distribution. Only the connected molecular graphs were used for the density estimation.

472

473Fig 3 illustrates the distribution of QED scores estimated from the molecules generated by the 474L-MolGAN and MolGAN. These distributions only represent the connected molecular 475graphs.

476The average QED values of the training data, MolGAN, and L-MolGAN were 0.76, 0.81, and 4770.88, respectively. In Fig 3, the training data and MolGAN have a single peak, whereas L-478MolGAN has two. In addition, the peak positions of the training data and MolGAN were 479similar. However, the distribution of MolGAN is narrower than that of the training data and 480has higher QED scores than the training data. These results indicate that MolGAN has been 481successful at chemical properties optimization.

482A closer look at the distribution in L-MolGAN in Fig 3 reveals two large peaks; one is close 483to the peak position of the MolGAN and the training data, while the other is located where 484the QED score is higher. We hypothesize that the peak in MolGAN and one of the peaks in L-485MolGAN were close to those of the training data because the optimization of chemical 486properties was strongly affected by the properties of mode of the QED distribution in the 487training data.

488Another peak in the L-MolGAN was located where the QED score was greater than 0.9.
489Notably, in this range of QED, the distributions of MolGAN and training data contained few
490molecules. This indicates that the L-MolGAN can exploit molecular graphs with better QED

491scores than the MolGAN. We assumed this was so because the L-MolGAN generated many 492connected molecular graphs. Improvement in connectivity would contribute to the generation 493of substituents and molecular skeletons with higher QED scores. Moreover, because the QED 494score is based on the physical properties of a molecular graph, it can be even calculated for 495disconnected graphs. For this reason, contrary to our presupposition, the chemical properties 496would be optimized for disconnected graphs as well. These results suggest that the L-497MolGAN can overcome this issue in MolGAN.

4984.3 Experiment III: Generation and evaluation of novel

molecules

500Fig 4 illustrates a two-dimensional description of the best 20 molecules with the highest QED 501scores generated by the optimized generator of L-MolGAN.

Fig 4 Two-dimensional representation of 20 molecules with the best QED scores. The 504numbers at the bottom of each molecule represent the corresponding QED, logP, and SA 505scores.

507Most molecules contained one or two sulfur atoms in their structures. In addition, many had 508QED scores of 0.9 or higher, indicating that their chemical properties were superior to those

509of the ZINC dataset. However, because their SA scores were significantly small, the 510molecules generated may be unrealistic. Therefore, we focused on the relationship between 511the structure of the generated molecules and their synthesizability.

5121,3-Thiazole was included as the common substructure of the top 20 molecules. Thiazole is a 513nitrogen-containing five-membered heterocyclic compound, which is a common skeleton in 514molecules used in pharmaceuticals and agrochemicals. The bicyclic ring skeleton was also 515found to be a common substructure within thiazole-containing molecules with an SA of 0.0. 516This skeleton is composed of five carbon or sulfur atoms bridging the carbon atoms at the 2 517and 5 positions of the thiazole ring. Nine out of the top twenty molecules had these 518characteristics. The bicyclic ring skeleton is difficult to synthesize because of the high steric 519strain of the molecule.

520

521Fig 5 Three-dimensional representation of the thiazole and bicyclic skeletons.

522

524skeletons included in the generated molecules. Because the thiazole ring skeleton has a planar 525structure, the atoms and substituents (i.e., the two methyl groups) in the ring lie on the same 526plane because the thiazole ring is aromatic. However, the planarity of the thiazole ring and its 527surroundings in the bicyclic ring skeleton is broken. The planarity of the thiazole ring was not 528maintained at the 2 and 5 carbon positions. This steric strain is affected by the number of

529atoms to be bridged. Therefore, we concluded that they were unrealistic due to the steric 530strain caused by the bicyclic framework. However, the L-MolGAN could generate these 531molecules, which have not yet been discovered. Therefore, it was worth an attempt to 532synthesize them as drug candidates.

533Fig 6 highlights the dominant molecules chosen in the QED–SA space. The dotted line 534indicates the envelope of the generated molecules. We confirmed that a tradeoff between the 535QED and SA scores, and the existence of eight dominant molecules. The dominant molecules 536[A] to [F] with a high QED score had a common cyclic substructure, whereas molecules [G] 537and [H] with low QED scores were chain-like. This suggested the potential of heterocyclic 538compounds as oral drugs. It is plausible that the synthesis became difficult when the ring 539structure contained two or more heteroatoms. Consequently, molecules [A] to [C] exhibited 540an SA of less than 0.2, and that of molecules [D] to [F] approximately 0.5.

541

542Fig 6 Dominant molecules identified through QED and SA scores ([A] to [H]). The points 543indicate all the molecules generated by the L-MolGAN. The dotted line indicates the 544envelope of the generated molecules.

545

546Subsequently, the dominant molecules were selected in SA–logP space, as shown in Fig 7.
547The molecule [H] was also chosen in the QED–SA space. We only reported molecules with
548higher lipophilicity based on the logP score. The molecule [H] was also chosen in the QED–

549SA space. The dominant molecules with higher lipophilicity in the QED–logP space are also 550shown in Fig 8. The molecule [B] was also chosen in the QED–SA space. Several dominant 551molecules were sulfur-containing compounds.

Fig 7 Dominant molecules with higher lipophilicity identified through SA and logP554**scores ([H], [I] and [J]).** The points indicate all the molecules generated by the generator of 555L-MolGAN. The dotted line indicates the envelope of the generated molecules.

Fig 8. Dominant molecules with higher lipophilicity identified through QED and logP558**scores ([B], [K] to [Q]).** The points indicate all the molecules generated by the L-MolGAN.
559The dotted line indicates the envelope of the generated molecules.

561The dominant molecules with higher hydrophilicity in the QED–logP space are shown in Fig 5629. Molecules [A] and [C] are also chosen in the QED–SA space. Fig 10 indicates molecules 563in the SA–logP space. Molecules [E], [G], and [H] were already chosen in the QED—SA 564space. In addition, the molecule [R] was also chosen in the QED—logP space.

566Fig 8 Dominant molecules with higher hydrophilicity identified through QED and logP 567scores ([A], [C], [R] to [U]). The points indicate all the molecules generated by the L-568MolGAN. The dotted line indicates the envelope of the generated molecules.

569

570Fig 9 Dominant molecules with higher hydrophilicity identified through SA and logP

571scores ([E], [G], [H], [R] and [V]). The points indicate all the molecules generated by the L-

572MolGAN. The dotted line indicates the envelope of the generated molecules.

573

574These results revealed the generation of a variety of dominant molecules by the L-MolGAN.

575Although the model was trained to optimize only the QED for drug discovery, a variety of

576molecules were identified among several combinations of the three chemical properties.

577Additionally, there was no best single molecular graph that simultaneously optimized the

578three chemical properties or their combinations. This motivated us to search for a variety of

579molecular graph among conflicting optimization goals of plural chemical properties. Future

580studies should apply a multi-objective optimization framework to the proposed method to

581search for dominant molecules with higher chemical property scores and more diverse

582chemical structures and properties.

583

5845. Conclusions

585The performance of the MolGAN deteriorates when generating a molecular graph with a

586molecular size larger than nine atoms, owing to the increase of disconnected graphs. This

587introduces challenges in the design of drugs with large molecules. We addressed this

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32

588challenge by adding to the MolGAN a molecular graph expansion mechanism that penalizes 589disconnected graphs and referred to it as L-MolGAN. The L-MolGAN improved the number 590of connected graphs generation on the ZINC-250k molecular dataset by a factor of 1.96, 591compared with the MolGAN. We also confirmed the generation of a variety of drug-like 592molecules by the L-MolGAN, even though it was optimized for a single property, i.e., QED. 593The L-MolGAN shall contribute to the efficient discovery of new molecules larger than those 594generated by the MolGAN.

595

596Acknowledgements

597We would like to thank Editage s[http://www.editage.com] for editing and reviewing this 598manuscript for English language. No funding to declare.

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