1	An interface-based molecular generative framework for
2	protein-protein interaction inhibitors
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29 Abstract

Protein-protein interactions (PPIs) play a crucial role in many biochemical processes and biological processes. Recently, many structure-based molecular generative models have been proposed. However, PPI sites and compounds targeting PPIs have distinguished physicochemical properties compared to traditional binding pockets and drugs, it is still a challenging task to generate compounds targeting PPIs by considering PPI complexes or interface hotspot residues. In this work, we propose a specifically molecular generative framework based on PPI interfaces, named GENiPPI. We evaluated the framework and found it can capture the implicit relationship between the PPI interface and the active molecules, and can generate novel compounds that target the PPI interface. Furthermore, the framework is able to generate diverse novel compounds with limited PPI interface inhibitors. The results show that PPI interfacebased molecular generative model enriches structure-based molecular generative models and facilitates the design of inhibitors based on PPI structures.

57 **Main**

A vast network of genes is inter-linked through protein-protein interactions and is 58 critical component of almost every biological process under physiological conditions, 59 and can be ubiquitous in many living organisms and biological pathways¹⁻⁴. Modulation 60 of PPIs expands the drug target space and has enormous potential in drug discovery. In 61 homo sapiens, it is estimated that the entire interactome comprises between 130,000 to 62 930,000 binary PPIs⁵⁻⁷. Despite significant efforts in developing modulators of PPIs, 63 drug design and development for PPI targets, especially targeting the PPI interfaces, 64 remains challenging^{6,8-12}. Structure-based rational design serves as an important tool 65 for the discovery of lead compounds in drug discovery¹³⁻¹⁷. Traditional drug targets and 66 PPIs targets have different bio-chemical features (**Table 1**)^{11,18-22}, so conventional drugs 67 68 and PPIs inhibitors have different physicochemical properties and drug-like properties (Table 1)^{11,23-30}. Given their differences, developing molecular generative models of 69 different paradigms are essential for the drug design of different target types^{11,19,31}. 70

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72 Generative artificial intelligence(AI) is enable to model the distribution of training samples and generate novel samples^{32,33}. In drug discovery, generative AI can accelerate 73 drug discovery by generating novel molecules with desired properties. Numerous 74 excellent review articles have summarized the development in this field^{16,17,34-41}. 75 Molecular generative models in drug design can be roughly divided into three 76 categories: ligand-based molecular generative (LBMG) models, 77 structurebased(pockets or binding sites) molecular generative models (SBMG), and fragment-78 79 based molecular generative models (FBMG), among which SBMG models have received much attention. ^{17,39,42}. Currently, some significant methods in structure-based 80 molecular generative models can be found in ⁴³⁻⁵¹, molecular generative models for PPI 81 structures or PPI interfaces have been rarely reported in the literature. In recent years, 82 classical machine learning⁵²⁻⁵⁴, active learning⁵⁵, and deep learning-assisted methods ⁵⁶ 83 84 is better screening and design of PPIs inhibitors have been explored, and ligand-based molecular generative models of PPI inhibitors have been reported⁵⁷. There are few 85

86 structure-based molecular generative models for PPIs targets have not been sufficiently

87 explored.

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In this study, we developed a conditional molecular generative framework based on 89 90 protein-protein interaction interfaces (named GENiPPI) for the design of PPI interface inhibitors. The framework was developed by a conditional Wasserstein generative 91 92 adversarial network (cWGAN) with convolutional neural networks (CNNs), integrated 93 graph attention networks (GATs) and long short-term memory (LSTM). It was designed to efficiently capture the relationship between PPI interface with active/inactive 94 compounds to train conditional molecular generative models (Fig. 1). As demonstrated 95 by the conditional evaluation, GENiPPI is an effective architecture for capturing the 96 97 implicit relationships between the PPI interface and active compounds. In summary, GENiPPI represents a potent deep learning framework for structure-based design of 98 PPI inhibitors. 99

100

101 **Results**

102 Generation of molecules targeting the PPI interface

Here, we introduce GENiPPI a modular deep learning framework for the design of structure-based PPIs inhibitors (**Fig. 1**). GENiPPI is composed of four main modules: GATs module ⁵⁸⁻⁶⁰ for representation learning of the protein complex interface, CNNs module for molecular representation learning, cWGAN module ⁶¹ for conditional molecular generation, and molecular captioning network module for SMILES strings decoding (as shown in Supplementary Figs.1, Figs.2, Figs.3 and Figs.4, respectively).

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Our framework undergoes four steps to accomplish the generation of molecules targeting the PPI interface. In the first step, we use GATs module designed for the protein complex interface is to effectively capture the nuanced atomic-level interaction characteristics inherent to the protein complex interface region. Next, we use CNN

module to provide a representation of the compound that contains voxel and electronic 115 density information in three-dimension space ⁶². And, the cGAN module is designed 116 to generate compounds that target PPI interfaces using features from the protein 117 complex interface region to regulate the inputs ⁶³. The cGAN module consists of a 118 generator, a discriminator, and a conditional network. The generator takes a Gaussian 119 random noise vector, and the protein complex interface features to generate a vector in 120 the molecular embedding space, the discriminator evaluates whether the generated 121 122 molecule embedding corresponds to a real or generated molecule, and the conditional network evaluates whether the molecule embedding matches the protein complex 123 interface features. Finally, we use the molecular captioning network, which is made by 124 a 3D CNNs and a recurrent LSTM ⁶⁴to decode molecular representations. The 125 126 molecular representation generated by the generator is fed as input to the 3D convolutional network with the LSTM subsequently decoding the SMILES strings. 127

128

129 **Conditional evaluation**

130 First, we verified the validity of the conditions that act as conditional molecular generative models for the protein complex interfaces. For this purpose, we selected 131 three PPI targets: MDM2(mouse double minute 2)/p53, Bcl-2(B-cell lymphoma 2)/Bax 132(Bcl-2 associated X), and BAZ2B(Bromodomain adjacent to zinc finger domain protein 133 134 2B)/H4(histone) for conditional evaluation. We generated 10,000 validated molecules each by the GENiPPI framework and calculated the drug-like metrics of the generated 135compounds: QED²⁷, QEPPI^{28,29} and Fsp3(fraction of sp3 carbon atoms)⁶⁵. We 136 compared the QED, QEPPI, and Fsp3 distributions of the active compounds and the 137 138 generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4 (Fig. 2). As shown, the distributions of drug-like properties were similar between the generated compounds 139 and the active compounds for the three PPI interface targets (Fig.2a, Fig.2b, and 140 Fig.2c), while different distributions of drug-like properties were observed between the 141 142 generated compounds based on different targets (Fig.2d, Fig.2e, and Fig.2f). The 143 results demonstrate the effectiveness of the PPI interface in conditioning the molecular 144 generative model. The drug-like properties of the framework generated compounds 145 migrate relative to those of the compounds in the training dataset, indicating that the 146 framework captures the distributions of the training dataset and generates novel 147 compounds.

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149 Model performance

In order to gain insight into the performance of the GENiPPI framework and to compare 150 151 it with other molecular generative models. We benchmarked our method by the MOSES platform⁶⁶, a leading benchmark platform of molecular generation. We trained all 152models on the full training dataset and randomly sampled 30,000 molecules. We 153utilized models and hyperparameters provided by the MOSES platform, such as an 154 Adversarial Autoencoder(AAE)⁶⁷, character-level recurrent neural networks 155(CharRNN)⁶⁸, Variational Autoencoder(VAE)⁶⁹, LatentGAN⁷⁰ and ORGAN⁷¹. To 156 validate the higher quality of the molecules generated by the conditioned model, we 157 compared them with molecules sampled from the GENiPPI framework and the 158 159 GENiPPI-noninterface framework without the conditioned module. We found that molecules generated by the conditioned GENiPPI framework were superior to other 160 models in novelty and diversity. 161

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As shown in Table 2, the GENiPPI framework has advantages in terms of uniqueness, 163 novelty, and diversity over the GENiPPI-noninterface. The GENiPPI framework 164 performs better overall in molecular generation. Compared with LatentGAN and 165 ORGAN, GENiPPI offers more benefits in terms of validity and diversity. While all 166 167 molecular generative models have their unique advantages in various performance comparison. However, the molecular generative models tailored to specific tasks, 168 especially those based on PPI structure, have more advantages and inspirations from 169 the GENiPPI framework. To understand the similarities and differences between the 170 molecular distributions generated by the GENiPPI framework and other models. We 171 172 compared the distribution of molecular properties of the Testset, iPPI-DB inhibitor, and the generated molecular datasets of AAE, CharRNN, VAE, LatentGAN, GENiPPI(noninterface) and GENiPPI(Supplementary Figs.4). The generated compounds have similar distributions of physicochemical properties to the compounds from the training set. While most of the iPPI-DB inhibitors have QED values lower than 0.5, most of them have QEPPI values higher than 0.5.

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179 Chemical space exploration

180 To better obtain an estimate of the chemical space distribution of the model generated molecules with the active compounds in the training datasets, we evaluated the 181 chemical drug-like space of the generated compounds by calculating t-distributed 182 random neighbourhood embedding (t-SNE) maps of MACCS fingerprint ⁷². The t-SNE 183 184 is a dimensionality reduction method used for data points visualization in two or threedimensional space by mapping high-dimensional data to a lower dimension ^{73,74}. By 185 this method, similar compounds are clustered to visualize the high-dimensional 186 chemical space of the compounds. The distribution of the generated compounds and 187 188 active compounds in chemical drug-like space by t-SNE visualization (Fig.3a, Fig.3b, and Fig.3c). The generated drug-like compounds not only share the chemical space 189 with the active compounds, but are also homogeneously mixed in the two-dimensional 190 space. The generated compounds show a similar chemical drug-like space to that of the 191 192 active compounds under 2D topological fingerprint. Adding the three dimensions of 193 compounds contributes to the design of promising drug-like compounds^{30,75}. We performed PMI shape analysis on the generated compounds and compared them with 194 drug-like compounds from DrugBank and iPPI-DB(Fig.3d). Many of the approved 195 compounds are either rod or disk shaped, and the generated drug-like compounds 196 library has a similar three-dimensional space. The PBF distribution of the library of 197 generated drug-like compounds is about 0~2 Å(Fig.3e). The results show that many of 198 the generated drug-like compounds are derived from relatively planar molecular 199 scaffolds. Moreover, we evaluated the ability of the model to generate target-specific 200 201 compounds by chemical space maps. To assess the overlapping of drug-like chemical

space, we utilized Tree MAP (TMAP)⁷⁶ to create the 2D projection(**Fig.3f**). Each point corresponds to a compound and is colored by its target label. The dark and light colors denote the generated compounds and the active compounds in the training set. These results suggest that our GENiPPI model can generate compounds that are similar to the active compounds in the training set and have novel structures. The results show that the framework enriches and expands the chemical space of PPI-targeted drug-likeness compounds.

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210 Few-shot molecular generation

Because of the huge consumptive costs involved in data collection, only a small amount 211 of labeled biomedical data are usually available. The process of drug design and 212 optimization often faces the problem of low data⁷⁷. The lack of effectively labeled data 213 tends to diminish the practical performance of most deep learning frameworks for drug 214 design. To perform generalized molecular generative design with limited labeled data, 215 it has been a trending topic in the few-shot generative community^{78,79}. The GENiPPI 216 217 model was applied to generate a virtual compound library for the heat shock protein 90 - cell division cycle 37(Hsp90-Cdc37) interaction interface. By training the model on 218 the PPI structure of Hsp90-Cdc37 (PDB ID: 1US7) and seven disruptors, we sampled 219 500 valid compounds. The similarity between active disruptors and generated 220 221 compounds of Hsp90/Cdc37 in the chemical space was visualized by t-SNE projection maps(Fig.4a). After few-shot learning, the generated compounds were mostly 222 distributed around the active disruptor, which demonstrated the effectiveness of few-223 shot learning in navigating through the targeted chemical space. We performed 224 pharmacophore-based matching by considering DCZ3112(a novel triazine derivative 225 that disrupts Hsp90-Cdc37 interactions) as a reference molecule⁸⁰. The top 5 generated 226 molecules have similar pharmacophore and shape features with DCZ3112(Fig.4b), 227 demonstrating the potential of the model to be applied to low-data PPI targets. Fig. 4c, 228 shows the hot spot amino acid residues at the PPI interface of the Hsp90-Cdc37 protein 229 230 complex(PDB ID: 1US7). We performed molecular docking for prediction of the

231 binding poses(Fig.4e) of DCZ3112 with the Hsp90-Cdc37 complex by the UCSF DOCK6.9 program⁸¹. The structure of the Hsp90-Cdc37 complex with DCZ3112 232 highlights the hydrogen bond interactions with amino acid residues:Arg32A, Glu33A, 233 234 Ser36A, Ser115A, Gly118A, Gln119A, and Arg167B(Fig.4e), which may be the major 235 energy contributors to protein-ligand interactions. The generated compounds were performed molecular docking together with DCZ3112, and selected compounds with 236 reasonable binding modes and higher binding affinity by visual inspection for 237 238 interaction pattern analysis. The generated compounds of GENiPPI not only obtained the better docking score than the active compounds, but also reproduced the interactions 239 with the key residues of the PPI interface. The generated compounds also formed 240 halogen bonds, salt bridges and π -cation interactions to improve the binding affinity of 241 the generated compounds to the target interface(Fig.4f). In conclusion, by analyzing 242 the interaction patterns between the generated compounds and the PPI interface, 243 GENiPPI learned the implicit interaction rules between the active compounds and the 244 PPI interface. 245

246

247 **Discussion**

We developed the GENiPPI framework, which combines protein-protein interaction 248 (PPI) interfaces features and conditional molecular generative model to generate novel 249 250 modulators for PPI interfaces. We validated the ability of GENiPPI framework to learn the implicit relationship between PPI interfaces and active molecules through 251 conditional evaluation experiments. GENiPPI used GATs to extract key features of PPI 252253 interfaces, and searched molecules by conditional wGAN with specific constraints. We 254 compared GENiPPI with various evaluation settings and benchmarks to demonstrate 255 its practical potential.

256

257 Despite the promising results, our framework has some limitations that can be 258 addressed in future work. We have not tested the model on a large number of receptor-259 ligand pairs of PPIs, which may affect its the generalization ability. The reason is that

the current PPI has relatively little data of drug-PPI target complexes than the traditional 260 261 dataset of drug-target complexes. Furthermore, the current framework does not incorporate the 3D structural information of ligand-receptor interactions of PPIs. and 262 263 There are still many ways to improve representation learning, balance training speed of molecular generative models and the diversity of generated molecules. Several 264 potential directions could further improve GENiPPI: (1) collecting and cleaning higher 265 quality data pairs for model development and testing; (2) fusing of molecular chemical 266 267 language models and pre-trained models of protein-protein structural features to finetune the datasets of receptor-ligand of PPIs to enhance the model generalization, 268 novelty and diversity of the generated compounds; (3) incorporating structural 269 270 information of PPIs into fragment-based molecular generative models is also a 271promising direction; (4) change the architecture of the model or combining it with deep reinforcement learning to generate novel compounds with better binding affinity. 272 Therefore, we will collect more data to further develop an enhanced version of 273 GENiPPI by combining novel representation learning methods and deep generative 274 275 approaches. In summary, the GENiPPI framework brings encouraging advances in PPI structure-based molecular generative tasks and presents a tool for rational drug design 276 277 in finding modulators of macromolecule-macromolecule interactions.

278

279 Methods

280 Datasets

We first investigated PPI targets that were annotated with sufficient compound 281 bioactivity data for training and evaluation of our model ⁸². For this study, we selected 282 10 validated PPI drug targets that cover the binding interface(Supplementary Table). 283 These targets are E3 ubiquitin-protein ligase Mdm2, apoptosis regulator Bcl-2, BAZ2B, 284 285 apoptosis regulator Bcl-xL, BRD4 bromodomain 1 BRD4-1, CREB-binding protein (CREBBP), ephrin type-A receptor 4 (EphA4), induced myeloid leukemia cell 286 287 differentiation protein Mcl-1, and menin. In addition, we randomly selected a subset of 250,000 compounds as additional inactive compounds from the ChEMBL ⁸³ dataset 288

that was used as part of the training datasets. A detailed data preprocessing can be found

- in Supplementary Note A.
- 291

292 Model strategy and training

293 Graph attention networks of protein-protein interaction interface

In this section, the representation learning of protein-protein complexes interfaces is 294 inspired by the work in protein docking model evaluation⁶⁰, which designed a double-295 graph representation to capture the interface features and interactions of protein-protein 296 complexes (Supplementary Figs.1). The extracted interface region is constructed as 297 two graphs (G^1 and G^2) for representing the interfacial information and the residues 298 involved in the two proteins participating in the interaction. A graph G can be defined 299 as G = (V, E, and A), where V is the set of nodes, and E is the set of edges between 300 them, and A is the adjacency matrix for mapping the association between the nodes of 301 the graph, which numerically denotes the connectivity of the graph. If the graph G has 302 N nodes, the dimension of the adjacency matrix A of the graph is N^*N , where $A_{ij} > 0$ 303

if the *i*-th node is connected to the *j*-th node, and $A_{ij} = 0$ otherwise. The graph G^1 describes the coding of the atomic types of all residues in the interface region, and its adjacency matrix A^1 describes the classification of interatomic bonding types for all residues at the interface region, which only considers the covalent bonds between atoms of interface residues within each subunit as edges. Therefore, it is defined as follows:

 $A_{ij}^{1} = \begin{cases} 1 & if \text{ atom } i \text{ and atom } j \text{ are connected by a covalent bond or } if i=j \\ 0 & \text{otherwise} \end{cases}$

The graph G^2 links both covalent bonds (thus including G^1) and non-covalent residue interactions as edges. The adjacency matrix A^2 for G^2 describes both covalent bonds and non-covalent interactions between atoms within the range of 10.0 Å to each other. The non-covalent atom pairs are defined as those which are closer than 10.0 Å to each other. It is defined as follows:

315
$$A_{ij}^{2} = \begin{cases} A_{ij}^{1}, & \text{if } i, j \in \text{receptor } or \ i, j \in \text{ligand} \\ e^{\frac{-(d_{ij}-\mu^{2})}{\sigma}}, & \text{if } d_{ij} \leq 10\text{ Å and } i \in \text{receptor and } j \in \text{ligand}; \\ or \ if \ d_{j} \leq 10 \text{ Å and } j \in \text{receptor } and \ i \in \text{ligand} \\ 0, & otherwise \end{cases}$$

where d_{ij} represents the distance between the *i*-th and the *j*-th atoms of all atoms of all residues in the interaction region. μ and σ are learnable parameter with initial values of 0.0 and 1.0, respectively. The formula $e^{-(d_i j - \mu)^2/\sigma}$ decays with increasing distance between atoms.

320

The graph representation is more flexible and natural to encode interactive information and adjacent(local) relationships. For the node features of the graph, we considered the physicochemical properties of the atoms. We used the same features from the previous work^{60,84,85}. Then, the feature vector of the nodes is 23 in length and was embedded into 140 features by a one-layer fully connected (FC) network.

326

The constructed graphs are used as the input for GATs. The graph consists of adjacency matrices A^1 , A^2 , node matrices N_{mn}^1 , N_{pq}^2 , and the node features, $x^{in} =$ $\{x_1^{in}, x_2^{in}, \dots, x_N^{in}\}$ and $x \in \mathbb{R}^F$, where F is the dimensionality of the node features. For the input graph of x^{in} , the pure graph attention coefficients are defined as follows, which represent the relative importance between the *i*-th and *j*-th nodes:

$$e_{ij} = x_i^T E x_j' + x_j^T E x_i',$$

where x'_i and x'_j are the transformed feature representations defined by $x'_i = W x_i^{in}$ and $x'_j = W x_j^{in}$. $W, E \in \mathbb{R}^{F \times F}$ are learnable matrices in the GATs. e_{ij} and e_{ji} become identical to satisfy the symmetrical property of the graph by adding $x_i^T E x_j^T$ and $x_i^T E x_i'$. The coefficient will only be computed for *i* and *j* where $A_{ij} > 0$.

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338 The attention coefficients will also be calculated for the elements in the adjacency

matrix. For the elements (i, j), they are defined in the following form:

340
$$a_{ij} = \frac{\exp(e_{ij})}{\sum_{j \in N_i} \exp(e_{ij})} A_{ij},$$

where a_{ij} represents the normalized attention coefficient between the *i*-th and *j*-th node pairs, while e_{ij} is the computed symmetric graph attention coefficient. N_i denotes the set of neighbors for the *i*-th node, which includes the interacting node *j* with $A_{ij} > 0$. The purpose here is to define attention by considering both the physical structure A_{ij} and the normalized attention coefficient e_{ij} of the interactions simultaneously.

346

Based on the attention mechanism, the new node features of each node are updated in consideration of its neighboring nodes, which is a linear combination of the neighboring node features and the final attention coefficient a_{ii} :

$$x_i'' = \sum_{j \in N_i} a_{ij} x_j',$$

Making the use of the GATs mechanism described previously, we applied four layers 351 of GATs to process the node embedding information of the neighboring nodes and 352 output the updated node embedding. For two adjacency matrices A^1 and A^2 , we use a 353 354 shared GAT. the initial input to the network is the atomic feature. Working with two matrices A^1 and A^2 , we have $x_1 = GAT(x^{in}, A^1)$ and $x_2 = GAT(x^{in}, A^2)$. In order to 355 focus only on the intermolecular interactions at the interface of the input protein-protein 356 357 complex, we obtain the final node embedding by subtracting the embeddings of the two graphs. By subtracting the updated embedding x_1 from x_2 , we can capture aggregated 358 information on intermolecular interactions from only the other nodes in the protein-359 protein complex interface. The output node feature is therefore defined as: 360

 $x^{out} = x^2 - x^1,$

362 After that, the updated x^{out} became x^{in} to iteratively increase the information through 363 the three following GATs layers. After the four GATs layers updated the node 364 embeddings, the node embedding of the entire graph was summed up as the overall

365 intermolecular interaction representation of the protein-protein complex:

366
$$x_{graph} = \sum_{k \in G} x_k$$

³⁶⁷ Finally, the FC layers were applied to the x_{graph} to obtain a [4,4,4] vector as features

- 368 of the protein-protein interface.
- 369

370 Molecular representation

For each SMILES string, a 3D conformer is generated using RDKit ⁸⁶ and optimized using the default settings of the MMFF94 force field. The molecular structure information is then extracted into a 35Å grid centered at the geometric center of the molecule using the HTMD package⁸⁷. The atoms of the molecule are discretized into a 1 Å cubic grid, and eight channels are considered to compute voxelized information. Finally, the electronic density of the molecules 9th channel is calculated using the original molecule method in Multiwfn(**Supplementary Figs.2**) ⁸⁸.

378

379 Conditional Wasserstein generative adversarial networks

The generator takes a conditional vector and a noise vector sampled from a Gaussian 380 381 distribution as inputs. The PPI interface features([1,4,4,4], vector shape) are concatenated with a noise vector of size [9, 4, 4, 4] and input to a 4-layer transposed 382 383 convolutional neural network (CNNs) with 256, 512, 1024, and 1024 filters, respectively. The first three layers downsample the array size using concatenation 384 convolution (s=2). For all convolutions, we use a kernel size of 4, and the Leaky ReLU 385 386 is used as an activation function after convolution. BatchNorm3d is applied between convolution and activation operations to normalize the values of each channel of each 387 388 sample.

389

390 The discriminator consists of a 4-layer sequential convolutional neural network (CNNs)

with 256, 512, 1024, and 1024 filters, respectively. The first three layers downsample

392 the array size using concatenation convolution (s=2). For all convolutions, we use a

kernel size of 4, and the Leaky ReLU ($\alpha=0.2$) is used as an activation function after 393 394 convolution. InstanceNorm3d is applied between convolution and activation operations 395 to normalize the values of each channel of each sample.

396

The physical and spatial features of the compounds are derived from the molecular 397 representation learning module, and the PPI interface features are obtained from the 398 GATs module of the protein complex interface. They are used to estimate the matching 399 400 probability between molecules and PPI interface features(Supplementary Figs.3).

401

Molecular captioning network 402

In this section, we will describe how to decode the generated molecular representation 403 into a SMILES strings. Our work is inspired by shape-based molecular generation^{89,90}, 404 which designs a combination network of convolutional neural networks (CNNs) and 405 Long Short-Term Memory (LSTM) ⁶⁴ to generate SMILES strings. In brief, the 406 molecular captioning network consists of a 3D CNNs and a recurrent LSTM. The 407 408 molecular representation generated by the generator is fed as input to the 3D CNNs, and the output of the 3D CNN is fed into the LSTM to decode the SMILES strings 409 (Supplementary Figs.4). 410

411

Model training 412

The conditional generative adversarial network is trained with Wasserstein loss. The 413 loss functions for the generator $(G_{(0(z,c))})$ and discriminator $(D_0(x))$ are: 414

415

$$\begin{split} L_{x_0} &= E_{iy_{xx}}[-D_y(x)] + E_{z_{xx},iy_{yx}}[D_{yy_{yx}}(G_{zy}(z,c))] + \lambda E_{iy_1}[(\| \nabla_{zx}D_y(\hat{x}) \|_z - 1)^2], \\ I_{x_{xx}} &= E_{z_{xx},ixx_{yx}}\left[-D_z \left(G_{zy}(z,c) \right) - \alpha lo \, g \big(f_u(G_u(z,c),c) \big) \right] \end{split}$$

(**A** (

417

where x and c are molecular representations and PPI interface features, respectively, 418 sampled from the true data distribution p_{real} , z is a random noise vector sampled from 419 a Gaussian distribution (p_z) , and f_0 is a function that evaluates the probability that a 420

421 PPI interface feature corresponds to a molecular representation. λ and α terms are 422 regularization parameters, both empirically set to 10. λ term weighs the effect of the 423 gradient penalty on discriminator loss. α term weighs the effect of the effect of f_0 on 424 the loss of the generator.

425

The model was trained for 50,000 calendar hours with a batch size of 8 (65 steps per 426 calendar hour). The discriminators were updated after each step, while the generators 427 428 were updated every 30 steps. The network was trained using the RMSprop optimizer with a learning rate of $1 \times 10-4$ for the generator and discriminator. during training, we 429 monitored the similarity between real and generated molecular representations using 430 Fréchet distances. The weights of the conditional networks were pre-trained on a binary 431 cross-entropy loss and frozen during GAN training. Training was performed on a single 432 NVIDIA A40 GPU, and all neural networks were built and trained using Pytorch 1.7.1 433 ⁹¹ and Tensorflow 2.5 ⁹². 434

435

436 Molecular generation

437 After the model has been trained, the embedding information of the protein-protein 438 complex interface is used to guide the model to generate novel molecules from the 439 latent space. The maximum sampling strategy was used in the LSTM, meaning that the 440 SMILES strings are generated by selecting the next token based on the highest 441 prediction probability⁸⁹.

442

443 **Evaluation settings**

444 **Conditional evaluation metrics**

In this study, the key is to evaluate the effectiveness of the proposed framework of protein-protein interaction interface-based conditional molecular generation. Therefore, we sampled the same number of valid molecules for the three PPI targets. Then we calculated the QED values and Fsp3 by RDKit⁸⁶ and calculated the QEPPI values by the QEPPI package (https://github.com/ohuelab/QEPPI) for the generated compounds and others, and plotted the density distribution for comparing the differences of drug-

451 likeness.

452

453 **MOSES evaluation metrics**

To evaluate the performance of our proposed conditional molecule generation framework, we used the evaluation metrics of validity, uniqueness, novelty and diversity provided by the MOSES platform⁶⁶, which are defined as follows:

457 Validity: Molecules defined as valid in the generated molecules.

458 Validity =
$$\frac{N_{\text{valid}}}{N_{\text{generalated}}}$$

459 Uniqueness: The proportion of unique molecules found among the generated valid460 molecules.

461 Uniqueness =
$$\frac{N_{\text{unique}}}{N_{\text{valid}}}$$

462 Novelty: The generated molecules are not to be covered in the training set.

463 Novelty =
$$\frac{N_{\text{novel}}}{N_{\text{unique}}}$$

FCD(Fréchet ChemNet Distance): To detect whether the generated molecules are diverse and whether they have chemical and biological properties that are similar with the real molecules⁹³.

467

468 Molecular shape

To evaluate the shape space of molecules, we used two widely adopted molecular 469 descriptors to represent the three dimensions of molecular structure: principal moment 470 of inertia (PMI)⁹⁴ and the best-fit plane (PBF)⁹⁵. The PMI descriptor classifies the 471 geometric shape of molecules into the degree of rod-shaped (linear shape, such as 472 acetylene), disk-shaped (planar shape, such as benzene), and sphere (spherical shape, 473 such as adamantane). The normalized PMI ratios (NPRs) are plotted in two-474dimensional triangle and then used to compare the shape space covered by different sets 475 of molecules, evaluating and visualizing the diversity of the molecular shape associated 476 with a given set of molecules³⁰. PBF is a three-dimensional descriptor that represents 477

the deviation of a molecule from a plane. The PBF descriptor is the mean distance of

each heavy atom from the best-fit plane passing via all heavy atoms 95 .

480

481 Tree MAP

To explore and explain the chemical space by unsupervised visualization of highdimensional data⁷⁶, we calculated MinHash fingerprint⁹⁶ vectors for active compounds and generated compounds. Then tmap⁷⁶ and faerun⁹⁷ were utilized to construct twodimensional projections of Tree MAP (TMAP).

486

487 **Protocol for few-shot generation**

Targeting the Hsp90-Cdc37 PPI interface is recognized as an important option for
cancer therapy. The crystal structure of the Hsp90-Cdc37 protein complex (PDB ID:
1US7) is available for molecular docking⁹⁸. In addition, known Hsp90-Cdc37 PPI
disruptors were collected for training of few-shot generative. They are DCZ3112,
Celastrol , FW-04-804, Sulforaphane, Withaferin A, Platycodin D, Kongensin A ⁹⁹.

OpenPharmacophore(https://github.com/uibcdf/OpenPharmacophore) was utilized to create pharmacophore models and virtual screening. The protein structures were processed by using UCSF Chimera¹⁰⁰, the program DOCK6.9 was used for semiflexible docking⁸¹, and PyMOL¹⁰¹ was used to create the figures. A detailed docking protocol can be found in Supplementary Note B.

498

499 Data Availability

500 The datasets are available at Github (<u>https://github.com/AspirinCode/GENiPPI</u>). The 501 data implementation will be provided upon acceptance of the manuscript for publication. 502

503 Code Availability

All the codes are freely available at Github (https://github.com/AspirinCode/GENiPPI).
The code implementation will be provided upon acceptance of the manuscript for
publication.

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509

510 Ethics declarations

511 The authors declare no competing interests.

512

513 **Competing Interests Statement**

514 The authors have declared no competing interests.

515

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Figure. legends 742

Fig. 1. Generation of molecules targeting PPI. 3D structural information of the 743 protein-protein complex interface is represented as a graph. Feature representation of 744 the interface region is captured by using a graph attention neural networks. The 745 746 representation of the voxel and electron density of the compound is encoded by a 3D convolutional neural networks (CNNs). A conditional Wasserstein generative 747 adversarial networks is trained to generate molecular embeddings with interface 748 749 features as conditions. Generator: takes interface features and random noise vectors to 750 generate molecular embeddings for the input features. Discriminator: calculates the probability that a molecule is from a real or a fake molecule. Condition: controls or 751 752 regulates the generation of molecules constrained by a specific protein-protein interface. Finally, a long short-term memory (LSTM) networks parse SMILES strings from 753 754 molecular representation.

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Fig. 2: Results of conditional evaluation. (a) The OED, OEPPI and Fsp3 distribution of 756 active compounds and compounds generated by the GENiPPI framework for 757 MDM2/p53; (b) The QED, QEPPI and Fsp3 distribution of active compounds and 758 compounds generated by the GENiPPI framework for Bcl-2/Bax; (c) The QED, QEPPI 759 760 and Fsp3 distribution of active compounds and compounds generated by the GENiPPI framework for BAZ2B/H4; (d) The QED distribution of generated compounds for 761 MDM2/p53, Bcl-2/Bax and BAZ2B/H4; (e) The QEPPI distribution of generated 762763 compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; (f) The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; 764

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Fig. 3: Chemical space exploration. (a) The t-SNE visualization of active compounds 766 767 and generated compounds for MDM2/p53; (b) The t-SNE visualization of active compounds and generated compounds for Bcl-2/Bax; (c) The t-SNE visualization of 768 active compounds and generated compounds for BAZ2B/H4; (d) The PMI ternary 769 density plots of generated compounds, small molecule drugs of DrugBank, and iPPI-770 771 DB inhibitors. Top left: propyne, bottom: benzene, and the top right: adamantane; (e) The molecular three-dimensionality distribution of the generated molecules was 772 visualized with NPR descriptors and PBF descriptors. (f) TMAP visualization of active 773 774 compounds and generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4.

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Fig. 4: Few shot molecular generation analysis. (a) The t-SNE visualization of the 776 distribution of active compounds and generated compounds for Hsp90/Cdc37; (b) 777 Comparison of the pharmacophore of the generated molecules with the reference 778 779 molecule(DCZ3112); (c) PPI interface region(in green) of the Hsp90(in palecyan)/Cdc37(in lightpink) complex; (d) The complex structure of DCZ3112(in 780 781 green) and Hsp90(in palecyan)-CDC37(in lightpink) modeled by molecular docking (PDB ID: 1US7); (e) The binding poses of generated compounds(in green) and 782 Hsp90(in palecyan)-CDC37(in lightpink) modeled by molecular docking (PDB ID: 783 1US7). Hydrogen bonds are displayed as blue dotted lines. π -cation Interactions are 784 785 displayed as orange dotted lines.

Figure 1



Fig. 1. Generation of molecules targeting the PPI interface. 3D structural information of the protein-protein complex interface is represented as a graph. Feature representation of the interface region is captured by using a graph attention neural networks. The representation of the voxel and electron density of the compound is encoded by a 3D convolutional neural networks (CNNs). A conditional Wasserstein generative adversarial networks is trained to generate molecular embeddings with interface features as conditions. Generator: takes interface features and random noise vectors to generate molecular embeddings for the input features. Discriminator: calculates the probability that a molecule is from a real or a fake molecule. Condition: neural or regulates the generation of molecules constrained by a specific protein-protein interface.



Fig. 2. Results of conditional evaluation. (a) The QED, QEPPI and Fsp3 distribution of active compounds and compounds generated by the GENiPPI framework for MDM2/p53; **(b)** The QED, QEPPI and Fsp3 distribution of active compounds and compounds generated by the GENiPPI framework for Bcl-2/Bax; **(c)** The QED, QEPPI and Fsp3 distribution of active compounds and compounds generated by the GENiPPI framework for Bcl-2/Bax; **(d)** The QED distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(d)** The QEPPI distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4; **(f)** The Fsp3 distribution of generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4.



Fig. 3. Chemical space exploration. (a) The t-SNE visualization of active compounds and generated compounds for MDM2/p53; (b) The t-SNE visualization of active compounds and generated compounds for Bcl-2/Bax; (c) The t-SNE visualization of active compounds and generated compounds for BAZ2B/H4; (d) The PMI ternary density plots of generated compounds, small molecule drugs of DrugBank, and iPPI-DB inhibitors. Top left: propyne, bottom: benzene, and the top right: adamantane; (e) The molecular three-dimensionality distribution of the generated molecules was visualized with NPR descriptors and PBF descriptors. (f) TMAP visualization of active compounds and generated compounds for MDM2/p53, Bcl-2/Bax and BAZ2B/H4.

Figure 4



Fig. 4. Few shot molecular generation analysis. (a)The t-SNE visualization of the distribution of active compounds and generated compounds for Hsp90/Cdc37; (b) Comparison of the pharmacophore of the generated molecules with the reference molecule(DCZ3112); (c)PPI interface region(in green) of the Hsp90(in palecyan)/Cdc37(in lightpink) complex; (d)The complex structure of DCZ3112(in green) and Hsp90(in palecyan)-CDC37(in lightpink) modeled by molecular docking (PDB ID: 1US7); (e)The binding poses of generated compounds(in green) and Hsp90(in palecyan)-CDC37(in lightpink) modeled by molecular docking (PDB ID: 1US7); Hydrogen bonds are displayed as blue dotted lines. π-cation Interactions are displayed as orange dotted lines.

Table. 1.

PPI interfaces	Binding sites		
Target properties			
Large surface area (1000-6000 Å ²)	Small surface (300-1000 Å ²) Hydrophobic		
Preference for Trp (W), Tyr (Y), and Arg (R) as PPI hotspot residues; subpockets	Large volume (~260 Å ³)		
Shallow, flat, flexible	Pocket, cliff		
Hydrophobic, featureless, undruggability	Diverse properties		
Chemical space			
MW≥ 400	MW≤ 500		
$LogP \ge 4$	LogP≤ 500		
$HBA \ge 4$	$HBA \leq 10$		
number of rings: ≥ 4	$HBD \leq 5$		
Ro4 Morelli's rules	Lipinski's Rule of 5 (Ro5)		
Quantitative estimate of drug-likeness scores			
QEPPI	QED		

Table 1. Comparisons between PPI interfaces and binding sites

Table. 2.

Table 2. Valid, unique, novelty and FCD of sampling SMILES after training. We sampled 30,000 SMILES each time.

Model	valid	Unique@1k	Unique@10k	novelty	FCD	
		• ©	• •	·	Test	TestSF
AAE	0.881	1.000	0.995	0.995	8.573	9.117
CharRNN	0.985	0.999	0.988	0.994	8.7564	8.952
VAE	0.834	1.000	0.996	0.994	7.703	8.141
LatentGAN	0.724	1.000	0.999	0.998	7.595	8.160
ORGAN	0.609	0.996	0.994	0.999	39.800	41.158
GENiPPI(noninterface)	0.999	0.997	0.975	0.997	7.653	8.132
GENiPPI	0.999	0.998	0.977	0.998	7.450	7.884