Topography-enhanced molecular graph representation for anti-breast cancer drug selection

Yue Gao1,2†, Songling Chen1,2†, Junyi Tong3† and Xiangling Fu1,2*

Abstract

Background: Breast cancer is currently one of the cancers with a higher mortality rate in the world. The biological research on anti-breast cancer drugs focuses on the activity of estrogen receptors alpha (ERα), the pharmacokinetic properties and the safety of the compounds, which, however, is an expensive and time-consuming process. Developments of deep learning bring potential to efficiently facilitate the candidate drug selection against breast cancer.

Methods: In this paper, we propose an Anti-Breast Cancer Drug selection method utilizing Gated Graph Neural Networks (ABCD-GGNN) to topologically enhance the molecular representation of candidate drugs. By constructing atom-level graphs through atomic descriptors for each distinct compound, ABCD-GGNN can topologically learn both the implicit structure and substructure characteristics of a candidate drug and then integrate the representation with explicit discrete molecular descriptors to generate a molecule-level representation. As a result, the representation of ABCD-GGNN can inductively predict the ERα, the pharmacokinetic properties and the safety of each candidate drug. Finally, we design a ranking operator whose inputs are the predicted properties so as to statistically select the appropriate drugs against breast cancer.

Results: Extensive experiments conducted on our collected anti-breast cancer candidate drug dataset demonstrate that our proposed method outperform all the other representative methods in the tasks of predicting ERα, and the pharmacokinetic properties and safety of the compounds. Extended result analysis demonstrates the efficiency and biological rationality of the operator we design to calculate the candidate drug ranking from the predicted properties.

Conclusion: In this paper, we propose the ABCD-GGNN representation method to efficiently integrate the topological structure and substructure features of the molecules with the discrete molecular descriptors. With a ranking operator applied, the predicted properties efficiently facilitate the candidate drug selection against breast cancer.

Keywords: Graph neural network, Breast cancer, Molecular representation, Drug prediction, Bioinformatics, Deep learning, Feature engineering, Decision support system
Background

Breast cancer is currently one of the most common cancers in the world with a higher fatality rate. According to the related statistics, more than 2 million new cases of breast cancer were diagnosed, where 0.6 million cases died. It accounted for about 15% of all cancer deaths among women worldwide [1]. Meanwhile, drug development is a process with long period and high candidate attrition rate. It was reported that the attrition rate of drug candidates has reached 90% [2]. Therefore, the research on anti-breast cancer drug with the assistance of in-silico tools is an urgent task pending for solutions.

The research on breast cancer is closely related to estrogen receptors [3, 4]. Studies have found that estrogen receptor alpha (ERα) is expressed in no more than 10% of normal breast epithelial cells, but about 50%-80% of breast tumor cells; and the experimental results of mice deficient in the ERα gene show that ERα does play a very important role in the development of the breast. At present, anti-hormone therapy is often used in breast cancer patients with ERα expression, which regulates the level of estrogen in the body by regulating the activity of estrogen receptors. Therefore, ERα is considered an important target for the treatment of breast cancer, and compounds that can antagonize the activity of ERα may be candidate drugs for the treatment of breast cancer [5–7].

In order for a compound to be a candidate drug, in addition to having good biological activity, it also needs to have good pharmacokinetic properties and safety in the human body, collectively known as ADMET (Absorption, Distribution, Metabolism, Excretion, Toxicity) properties [8]. Among them, ADME mainly refers to the pharmacokinetic properties of the compound, which describes the law of the concentration of the compound in the organism over time, and T mainly refers to the toxic and side effects that the compound may produce in the human body. No matter how active a compound is, if its ADMET properties are poor, for example, it is difficult to be absorbed by the human body, or the metabolism rate in the body is too fast, or it has some toxicity, then it is still difficult to become a drug, so ADMET properties need to be optimized.

At present, in the field of drug research, regarding time and cost consuming [9, 10], Quantitative Structure-Activity/Property Relationship (QSAR/QSPR) model is one of the most representative in-silico prediction tools to evaluate biological activity and ADMET properties of candidate drug compounds. By leveraging the disease-related targets, e.g. ERα, and modeling them as dependent variables, QSAR/QSPR models can predict new compound molecules with better biological activity, physicochemical property, and toxicological responses, and realize preliminary virtual screening for drugs.

With the development of the field of bioinformatics, diverse machine learning based methods have been proposed and applied into QSAR/QSPR modeling for drug property prediction [11–14]. The process can be generally divided into three stages. The first stage is traditional machine learning method represented by linear regression [15], random forest [16], and support vector machine (SVM) [17]. Such representation methods are dependent on hand-craft discrete features from the descriptors and the fingerprints of molecules to model the ADMET properties [18, 19], which is, however, time-consuming and inefficient. The second stage is sequential-based deep learning method represented by CNN [20–23] and LSTM [24]. Such methods can map the structure of compounds into a sequential dimension and aggregate the molecule-level features. Given their remarkable performance improvement compared with traditional machine learning
methods, in recent years, sequential-based deep learning methods have been the most popular in-silico methods for QSAR/QSPR modeling. However, existing sequential-based deep learning methods are still based on hand-craft discrete features from the descriptors and the fingerprints of molecules, which means that these methods cannot further reflect the topological characteristics implicit in the molecular structure.

Recently, the popularity of graph neural networks in the bioinformatics community brings potential to further enhance the molecular representation, which is the third stage of QSAR/QSPR modeling [25–30]. Graph neural networks are naturally suitable for modeling topological structure of non-Euclidean data like molecule and can realize global feature extraction from the global structure [31–35]. Currently, a series of graph-based deep learning methods have been proposed for molecular representation and applied for QSAR/QSPR modeling. For example, Duvenaud et al. [36] proposed convolutional networks on graphs to represent molecular fingerprints, which mapped the features of fingerprints into molecular structure via graph convolution operations. In terms of graph based ADMET prediction, Feinberg et al. [37] utilized a modified graph convolutional networks to model ADMET properties at Merck. Montanari et al. [38] demonstrated that graph convolutional neural networks are much more competitive to predict physicochemical ADMET endpoints; Feinberg et al. [39] proposed PotentialNet which applied graph convolution neural networks to conduct multi-task ADMET property prediction.

Although many variants of graph neural networks have been developed for molecular representation and ADMET prediction, limitations of existing methods still exist. First, existing graph-based methods only map the descriptors into a global molecule-level structure, which means that they may fail to mine the intrinsic knowledge implicit in the key chemical substructures of the molecules. The significance of biological substructure within a compound is neglected. Second, compared with feature engineering-based machine learning models, GNN models are generally less sensitive to the source of atomic descriptors [40, 41], which means that graph-based methods are less explainable and are not good at representing known explicit knowledge. Therefore, existing graph-based methods fail to integrate the implicit topological knowledge with the explicit discrete descriptor knowledge. Third, most of the existing graph-based methods for ADMET modeling are modified from graph convolution neural network. Such methods follow the transductive learning strategy, which is more computationally expensive and time-consuming compared with inductive learning strategy. Meanwhile, in terms of QSAR/QSPR modeling task, most of the existing methods focus on ADME or ADMET property prediction, while neglect the prediction of biological activity. In addition, to our knowledge, there is still no graph-based QSAR/QSPR model focusing on anti-breast cancer drug selection.

Inspired by the recent progress claimed above, in this paper, we propose the ABCD-GGNN representation method to topologically realize QSAR/QSPR model for ERα and ADMET prediction. ABCD-GGNN can topologically learn both the structure and substructure representations of molecules and deeply integrate them with discrete molecular descriptor representation, which strongly enhances the molecular representation performance and can realize inductive prediction on activity, property, and toxicity. In addition, we design a whole framework of anti-breast cancer drug selection based on
ABCD-GGNN with a decision-support setting. With an extra ranking operator applied based on the predicted properties from ABCD-GGNN, selection of candidate drugs against breast cancer can be efficiently facilitated, which may hugely benefit the research on anti-breast cancer drugs. The contributions of this paper are threefold:

- We propose an Anti-Breast Cancer Drug selection method utilizing Gated Graph Neural Networks (ABCD-GGNN), which topologically learns both the implicit structure and substructure characteristics of a candidate drug, and integrates with explicit discrete molecular descriptors to better generate a molecular-level representation. As a result, activity, property, and toxicity of the candidate drugs can all be inductively predicted.
- We design a whole framework of anti-breast cancer drug selection based on ABCD-GGNN to automatically assist researchers with a decision-support setting. To our best knowledge, this is the first work aiming to deal with anti-breast cancer drug development via graph-based deep learning method.
- Extensive experiments conducted on our collected anti-breast cancer candidate drug dataset demonstrate the outstanding performance of our proposed ABCD-GGNN representation method and the rationality of our designed framework for candidate drug selection.

**Methods**

In this section, we first introduce the candidate drug dataset we collect. Then, we illustrate the implementation of our anti-breast cancer drug selection method based on ABCD-GGNN step by step. As the pipeline shown in Fig. 1, our designed drug selection process can be decomposed into four stages: 1) topological molecular graph representation based on GGNN which integrates both structure and substructure characteristics of the molecule, 2) discrete property representation based on machine learning algorithm, 3) integration of the molecular representation of ABCD-GGNN and prediction for ERα and ADMET, and 4) candidate drug selection based on our designed ranking operator.

**Dataset**

To evaluate the efficiency of our proposed method, we collect a dataset containing 1974 organic compounds that may be the candidate drugs of anti-breast cancer. The dataset provides the simplified molecular input line entry system (SMILES) and 729 molecular descriptors of each organic compound. The 729 molecular descriptors include diverse descriptions on the characteristics of molecule in two-dimension and three-dimension. The dataset labels the ERα value expressed as pIC50 for each organic compound. Meanwhile, to objectively evaluate the pharmacokinetic properties and the safety of each organic compound, the dataset quantifies them with 5 property labels: absorption, distribution, metabolism, excretion, and toxicity (ADMET). In our collected dataset, the 5 properties are referred to 5 metrics: Caco-2, CYP3A4, human Ether-a-go-go Related Gene (hERG), and Human Oral Bioavailability (HOB), respectively. Due to the page limit of the paper, we present a detailed illustration of a candidate drug sample in Additional file 1: Table 1 of the Appendix section.
Topological molecular graph representation

In the stage of topological molecular graph representation, graph neural networks are adopted to atomically model the structure of a drug so as to learn the topological molecular features three-dimensionally for the final representation of ABCD-GGNN. With the atom node information globally interacted in the graph structure, both topological structure and substructure features can be well represented and integrated. We first illustrate the implementation of the topological structure representation. Then we illustrate how topological substructure representations are generated and integrate with the topological structure feature to enhance the topological molecular representation. The whole framework of the topological molecular graph representation based on ABCD-GGNN is shown in Fig. 2.

Atom-level topological structure graph construction

Graph construction is the kernel stage for the topological graph representation. Given that a graph is denoted as $G = (V, E)$ where $V$ ($|V| = n$) is the set of graph nodes and $E$ is the set of graph edges. In terms of atom-level graph construction for candidate drugs, $V$ denotes the atom set in a molecule and $E$ denotes the chemical bond set in a molecule.
In terms of the feature initialization for each atom node, here we summarize 8 atomic descriptors from the corresponding SMILES and 729 molecular descriptors, which are atom type, number of bonds, formal charge, chirality, hydrogen bound number, hybridization, aromaticity, and atom mass. The detailed descriptions on the 8 atomic descriptors are listed in Table 1. Every atomic descriptor is transferred into a one-hot vector and are concatenated to form a 39-dimension vector as the initialization of an atom feature.

In terms of edge construction for each molecular graph, we construct an adjacent matrix $A \in \mathbb{R}^{|V| \times |V|}$ to describe the connection relationship between atom nodes. The element in $A$, e.g., $a_{ij}$, is the connection type between $i$-th node and node $j$-th node. The connection type varies among 0, 1, 2, 3, and 1.5, which denotes the bond type: single bond, double bond, triple bond, and aromatic hydrocarbon, respectively.

### Table 1 Descriptions of components of the feature initialization for the atomic nodes

| Atomic descriptor          | Description                                                                 | Vector size       |
|----------------------------|-----------------------------------------------------------------------------|-------------------|
| Atom type                  | 12 types of atoms in the 200 molecules of the dataset                       | 12-digit 0/1 vector |
| Number of bonds            | The number of chemical bonds that the atom participates in                  | 6-digit 0/1 vector |
| Formal charge              | The integer-form electric nucleus of the atom                               | 5-digit 0/1 vector |
| Chirality                  | CW, CCW, unspecified, or other                                              | 4-digit 0/1 vector |
| Hydrogen bound number      | Atomic bound hydrogen atom charge                                          | 5-digit 0/1 vector |
| Hybridization              | sp, sp2, sp3, sp3d, or sp3d2                                               | 5-digit 0/1 vector |
| Aromaticity                | Whether the atom is part of an aromatic hydrocarbon                         | 1-digit 0/1 vector |
| Atom mass                  | The mass of the atom                                                        | A normalized number between 0 and 1 |

In terms of edge construction for each molecular graph, we construct an adjacent matrix $A \in \mathbb{R}^{|V| \times |V|}$ to describe the connection relationship between atom nodes. The element in $A$, e.g., $a_{ij}$, is the connection type between $i$-th node and node $j$-th node. The connection type varies among 0, 1, 2, 3, and 1.5, which denotes the bond type: single bond, double bond, triple bond, and aromatic hydrocarbon, respectively.

#### Graph-based global information interaction

Getting the molecular graph constructed, we then employ GGNN [42] to realize the global information interaction between the atom nodes. GGNN learns node representations through neural networks with gated recurrent units (GRU), so that information from neighborhood can be fused and enrich the own representation. Information fusion between nodes strengthens continuously with the interaction time $t$ increased.
and can finally achieve global information interaction of the whole topological structure. In this way, we can finally get a topological structure representation for a candidate drug. Detailed interaction functions are listed as follow:

\[ o^t = (W_o A + b_o)h^{t-1} \]
\[ z^t = \sigma \left( W_z o^t + U_z h^{t-1} + b_z \right) \]
\[ r^t = \sigma \left( W_r o^t + U_r h^{t-1} + b_r \right) \]
\[ \tilde{h}^t = \tanh \left( W_h o^t + U_h \left( r^t \odot h^{t-1} \right) + b_h \right) \]
\[ h^t = \tilde{h}^t \odot z^t + h^{t-1} \odot (1 - z^t) \]

where \( \sigma \) is the sigmoid function, and parameters \( W, U, \) and \( b \) are trainable weights and biases. \( o^t \) denotes the information that a node could receive from its adjacent neighbors in time step \( t \). \( z^t \) and \( r^t \) are functions that control update gate and reset gate, respectively, which determine to what degree the neighborhood information contributes to the current node embedding.

**Topological molecular representation readout**

With the topological structure representation of the distinct molecule updated, we then aggregate the atom-level representations into a molecule-level representation in the readout stage. The readout functions are designed as follow:

\[ h_{w,d} = \sigma \left( f_1 \left( h^t_w \right) \right) \odot \tanh \left( f_2 \left( h^t_w \right) \right) \]

\[ h_G = \frac{1}{|W| + 1} \sum_{w \in W} h_w + \text{Maxpool}(h_1, \ldots, h_w, h_d) \]

where \( f_1 \) and \( f_2 \) are two multilayer perceptrons (MLP) which perform as a soft attention weight and a non-linear feature transformation, respectively.

The readout functions are designed as above with the intention to reflect the truth that all atom node representations contribute to the information aggregation by getting through averaging function and a max-pooling function, while only part of atom nodes with higher weights distributed by attention mechanism contribute more [34]. Consequently, here we get the topological structure representation of the molecule \( h_G \) for further prediction.

**Substructure graph construction and integration**

Subgraphs are believed to imply significant attribute characteristics that may further extract and enhance the original graph representation [43], especially to the graph
representation of molecules whose substructures represent scaffolds of molecule which should imply much attribute knowledge.

Therefore, we additionally extract the substructures from SMILES of the molecules via the SMILES pair encoding algorithm. Given $G_{\text{sub}} = S_1, \cdots, S_n$ denotes the subgraph set of n substructures extracted from the graph $G$. Then, we construct the atom-level subgraphs and get through the global interaction via GGN and representation readout operations as the original graph does. Consequently, we get a substructure-level representation set $H_{\text{sub}} = h_{S_1}, \cdots, h_{S_n}$.

Considering that the contributions different substructures make to the molecular representation are uneven, here we adopt an attention mechanism to dynamically adjust the weights of the original graph and each subgraph. In this way, both molecular graph representation and diverse substructure graph representations get deeply integrated. In other words, the topological graph representations of the candidate drugs are strongly enhanced. Detailed formulas of the attention mechanism and the feature integration is shown below:

$$w_j = \frac{\exp(e_j)}{\sum_{k \leq |H_{\text{sub}}|+1} \exp(e_k)}, \quad e_j = c^T \tanh \left( W h_j + b \right) \quad (8)$$

$$h = w_0 \times h_G + w_1 \times h_{S_1} + \cdots + w_n \times h_{S_n} \quad (9)$$

where $w_0, \cdots, w_n$ is distributed attention weights. $c$, $W$, and $b$ are trainable parameters to be learned. Consequently, here we finally get the topological molecular graph representation $h$ that deeply integrate the structure and substructure characteristics of the molecule.

**Discrete molecular descriptor representation**

Molecular descriptors are the discrete expression of a molecule which may imply the potential chemical properties as a candidate drug. Given that the anti-breast cancer candidate drug dataset provides 729 molecular descriptors of all the candidate drugs, which is a quite large number. Here we first employ XGBoost algorithm to select the descriptors that count more. Then, we further reduce the dimensionality of the integrated molecular descriptor representation to realize the molecular descriptor representation readout.

**Discrete molecular descriptor selection**

Considering the redundancy and sparsity of the raw molecular descriptors, we believe it is necessary to select the more property-related descriptors with the help of machine learning method. Therefore, here we apply XGBoost, a decision-tree-based ensemble Machine Learning algorithm that uses a gradient boosting framework, to select the top 50 property-related descriptors for further feature integration and readout.

In terms of the implementation of XGBoost, we first set the objective function, i.e., the loss function as below:
where $L(\phi)$ is the differentiable convex loss function, which represents the gap between the predicted value $\hat{y}_i$ and the target value $y_i$ to avoid under-fitting; the function $\sum_k \Omega(f_k)$ can reduce the complexity of the model. The additional regularization term helps avoid overfitting. When the regularization parameter is set to 0, the goal is back to the traditional gradient tree boosting algorithm. Since the model is trained by addition, the prediction at time step $t$ equals the prediction at time step $t-1$ plus the function at time step $t$. The formula is shown below:

$$y^{(t)}_i = \sum_k f_k(x_i) = y^{(t-1)}_i + f_t(x_i)$$  \hspace{1cm} (12)$$

Second, we utilize Taylor expansion formula for approximation.

$$L(\phi) \approx \sum_i \left[ l \left( y_i, \hat{y}^{(t-1)}_i \right) + g_i f_t(x_i) + \frac{1}{2} h_i f_t^2(x_i) \right] + \Omega(f_t)$$  \hspace{1cm} (13)$$

where $g_i = \frac{\partial \hat{y}^{(t-1)}_i}{\partial y^{(t-1)}_i}$ and $h_i = \frac{\partial^2 \hat{y}^{(t-1)}_i}{\partial y^{(t-1)}_i^2}$ are the first partial derivative and the second partial derivative, respectively.

To make each sample on a leaf node, the node score is defined as $f_t(x) = W_q(x)$. The optimal weight is defined as $W_{q} = -\frac{G_j}{\sum_j G_j}$ according to the quadratic function to find the most value formula, where $G_j = \sum_i g_i, H_j = \sum_i h_i$. Thus, the optimal function value is defined as obj = $-\frac{1}{2} \sum_j G_j^2 + \gamma T$ and can rank the most properties-related molecular descriptors consequently. Discrete molecular descriptor representation readout With the 50 molecular descriptors selected, we then concatenate them in to a 50-digit vector as a molecule-level representation. Since the contribution of each descriptor, as is ranked by XGBoost algorithm above, should be uneven, we adopt the attention mechanism to dynamically adjust the weight of each digit. Then, to further integrate the discrete molecular descriptor representation so as to better integrate with the topological molecular representation, we reduce the dimensionality of the molecular descriptor representation in to a 39-digit vector with a fully connected layer, which make the two representation readouts in the same size. The formulas are shown below:

$$w_j = \frac{\exp(e_j)}{\sum_{k \leq |m|} \exp(e_k)}, \quad e_j = c^T \tanh(Wm_j + b)$$  \hspace{1cm} (14)$$

$$m = [w_0 \times m_0, w_1 \times m_1, \cdots, w_n \times m_{|m|}]$$  \hspace{1cm} (15)$$

$$h_m = Wm + b$$  \hspace{1cm} (16)$$

where $h_m$ is the representation readout of the discrete molecular descriptors.
Metric Prediction

Based on the topological graph representation and the molecular descriptor representation, a final representation of anti-breast candidate drug can be integrated to predict both the ERα value and the ADMET properties.

Topological and discrete property representation integration

To adaptively adjust the contribution the topological graph representation and molecular descriptor representation make to the prediction result, we design the hyperparameter $\lambda \in (0, 1)$ to weight and integrate the two types of features as the formula shown below:

$$h_{ABCD-GGNN} = \lambda h + (1 - \lambda) h_m$$

(17)

where $h_{ABCD-GGNN}$ is the final integrated representation of the anti-breast candidate drugs.

In this stage, we can claim that the molecular representation based on ABCD-GGNN is completed.

Prediction and training process

We treat the prediction of ERα value and ADMET properties as a regression task and a two-class classification task, respectively. In terms of ERα value prediction, the representation $h_{ABCD-GGNN}$ gets fed into a fully connected layer. Parameters are trained through the mean square error.

$$\hat{y}_{ER\alpha} = W h_{ABCD-GGNN} + b$$

(18)

$$Loss = \frac{1}{m} \sum_{i=1}^{m} (y_{ER\alpha} - \hat{y}_{ER\alpha})^2$$

(19)

where $W, b$ denote trainable parameters, $m$ denotes the batch size, and $y_{ER\alpha}$ denotes the ground truth value of ERα.

In terms of ADMET properties prediction, the representation $h_{ABCD-GGNN}$ gets fed into a softmax layer to make prediction. Parameters are trained through the cross-entropy function.

$$\hat{y}_{ADMET} = \text{softmax}(W h_{ABCD-GGNN} + b)$$

(20)

$$Loss = - \sum_{i} y_{ADMET} \log (\hat{y}_{ADMET})$$

(21)

where $W, b$ denote trainable parameters and $y_{ADMET}$ denotes the i-th element of the one-hot label.


**Candidate drug selection**

To comprehensively consider both the two types of attributes when evaluating the potential of the candidate drugs, here we design a ranking operator consisting of feature binning and scorecard. By scoring each candidate drug, a ranking list can be generated, which can efficiently facilitate the research on anti-breast cancer drug selection.

**Feature binning**

Since the ADMET properties are binary while the ERα value is a continuous value, here we select chi-square binning. The adjacent intervals are the smallest chi-square value are merged together until the definite stopping criterion is met. We set the chi-square threshold (obtained from the significance level and degree of freedom), and calculate the chi-square for each pair of adjacent values as the formula shown below:

\[
\chi^2 = \sum_{i=1}^{2} \sum_{j=1}^{2} \frac{(A_{ij} - E_{ij})^2}{E_{ij}}
\]

where \( A_{ij} \) is the feature number of the j-th class attribute in the i-th interval, and \( E_{ij} \) is the expectation of \( A_{ij} \).

**Setup of scorecard**

To set up the scorecard for the candidate drug ranking, we first calculate the corresponding score of the attribute as below:

\[
\text{score} = \sum_{i=1}^{n} \left( \text{woe}_i \times \beta_i + \frac{a}{n} \right) \times \text{factor} + \frac{\text{offect}}{n}
\]

where \( \text{woe}_i = \ln \frac{p_1}{p_0} \) is the woe value calculated based on the results of binning and denotes the difference between the response value and the non-response value, \( \beta_i \) is the regression coefficient, \( a \) is the regression intercept term, factor is the scale factor, and \( \text{offect} \) is the offset.

Finally, the calculation formula of the scorecard is defined as below to get the scores of the candidate drugs.

\[
\text{score} = \sum_{i=1}^{n} \left( \text{woe}_i \times \beta_i + \frac{a}{n} \right) \times \text{factor} + \frac{\text{offect}}{n}
\]

**Results**

In this section, 1) we first evaluate the performance of our proposed ABCD-GGNN on our collected anti-breast cancer candidate drug dataset and compare them with other representative models. 2) Then, we make extensive characteristics analysis and ablation study to demonstrate the effectiveness and contribution each stage makes for the ABCD-GGNN representation. 3) Finally, we demonstrate the biological
rationality of applying the ABCD-GGNN prediction results into the ranking operator for candidate drug selection.

**Performance of ABCD-GGNN**

**Baselines and evaluation metrics**

Keeping track of the representation methods applied in the study on drug prediction, we compare the representation performance of ABCD-GGNN with those of representative baseline models, which can be categorized into two types: 1) traditional machine learning methods, for example, Linear Regression and Random Forest for ER$\alpha$ prediction and SVM for ADMET prediction; 2) deep learning methods, for example, Bi-LSTM and Graph-CNN for ADMET prediction. Detailed descriptions of these baselines are shown as follow:

- **Linear Regression** a representative supervised learning method. Based on one or more independent variables, linear regression can model a best-fitting relationship for regression problem.
- **Random Forest** an ensemble learning method that constructs decision trees during training. It can realize prediction on the mean prediction of trees for regression tasks by utilizing random subspace method and bagging during tree construction.
- **SVM** a traditional supervised learning method. By maximizing the margin between data samples, SVM can perform well on both regression and classification problem.
- **Bi-LSTM** a representative sequential deep learning method which consists of two LSTMs: one forward and the other backwards direction. Bi-LSTM effectively capture the contextual information in time dimension.
- **Graph-CNN** one of the most representative graph neural network method. By combining CNN with spectral theory, Graph-CNN is more advantageous in dealing with the discriminative feature extraction of signals in the discrete spatial domain and can better describe the intrinsic relationship between different nodes of the graph.

To better reflect the performance of the compared models, in the ER$\alpha$ prediction task, we adopt the of mean square error loss (MSE) and R-Square (R2) as the evaluation metric, while in the ADMET prediction task, we adopt the of mean square error loss (MSE) and R-Square (R2) as the evaluation metric, Precision, Recall, F-score (F1), Area Under the ROC Curve (AUC), and Area Under the Precision-Recall curve (AUPR) as the evaluation metric.

**Experimental settings**

In terms of the detailed dataset setting, we keep the ratio of positive samples and negative samples close to 1:1 for each property. In addition, we utilize ten-fold cross-validation to evaluate the performance of all the compared methods. Positive and negative samples are kept balanced in each fold. We divide the dataset in a ratio of 8:1:1 as training set, validation set, and test set, respectively. The hyperparameters were tuned according to the performance on the validation set. Empirically, we set the learning rate as 0.01 with Adam optimizer and the dropout rate as 0.5. The interaction step of GGNN is set as 2. The hyper parameter $\lambda$ is set as 0.6.
The performance of the compared models on the prediction of ERα and ADMET are presented on Tables 2 and 3, respectively. It can be observed that our proposed ABCD-GGNN outperforms all the representative models on the two prediction methods. Specifically, in the ERα prediction task, ABCD-GGNN achieves the lowest loss value and highest R2 value, which means that the prediction results of our proposed model can better fit the expected ERα value with lower error. In the ADMET prediction task, ABCD-GGNN achieves the highest performance on Precision, Recall, F1, AUC, and AUPR, and prevails other models in a large margin. Therefore, it can be concluded that our proposed ABCD-GGNN representation method achieve a splendid performance on the property prediction for anti-breast cancer candidate drug.

### Table 2 Performance comparison on the prediction of ERα

| Model          | MSE | R2  |
|----------------|-----|-----|
| Linear Regression | 2.156 | 0.276 |
| Random Forest  | 0.5147 | 0.6133 |
| SVM            | 0.6878 | 0.6273 |
| ABCD-GGNN      | 0.4811 | 0.7741 |

We run all models 10 times and report the mean test MSE and R2.

### Table 3 Performance comparison on the prediction of ADMET

| Model         | Dataset | Precision | Recall | F1     | AUC   | AUPR |
|---------------|---------|-----------|--------|--------|-------|------|
| SVM           | MN      | 0.7843    | 0.6709 | 0.6943 | 0.7957 | 0.8209 |
|               | HOB     | 0.7733    | 0.7498 | 0.7607 | 0.8104 | 0.6239 |
|               | hERG    | 0.8080    | 0.7589 | 0.7791 | 0.8239 | 0.8494 |
|               | CYP3A4  | 0.8397    | 0.7998 | 0.8133 | 0.8518 | 0.8591 |
|               | Caco-2  | 0.8453    | 0.7807 | 0.8068 | 0.8552 | 0.7525 |
| BiLSTM        | MN      | 0.8226    | 0.7310 | 0.7537 | 0.8195 | 0.7731 |
|               | HOB     | 0.7462    | 0.7008 | 0.7165 | 0.7711 | 0.7337 |
|               | hERG    | 0.8350    | 0.7914 | 0.7968 | 0.8452 | 0.8196 |
|               | CYP3A4  | 0.8838    | 0.8627 | 0.8741 | 0.9129 | 0.8952 |
|               | Caco-2  | 0.8134    | 0.7954 | 0.8021 | 0.8533 | 0.8258 |
| Graph-CNN     | MN      | 0.8629    | 0.8293 | 0.8461 | 0.8710 | 0.8623 |
|               | HOB     | 0.8110    | 0.7635 | 0.7824 | 0.8369 | 0.8061 |
|               | hERG    | 0.8495    | 0.8690 | 0.8556 | 0.9081 | 0.8585 |
|               | CYP3A4  | 0.8913    | 0.8827 | 0.8840 | 0.9304 | 0.8731 |
|               | Caco-2  | 0.8479    | 0.8227 | 0.8306 | 0.8740 | 0.8881 |
| ABCD-GGNN     | MN      | 0.9255    | 0.9613 | 0.9430 | 0.9714 | 0.9862 |
|               | HOB     | 0.8637    | 0.8804 | 0.8712 | 0.9130 | 0.9273 |
|               | hERG    | 0.8914    | 0.8839 | 0.8842 | 0.9303 | 0.9456 |
|               | CYP3A4  | 0.9474    | 0.9163 | 0.9355 | 0.9487 | 0.9322 |
|               | Caco-2  | 0.8828    | 0.8832 | 0.8829 | 0.9296 | 0.9134 |

We run all models 10 times and report the mean test precision, recall, F1, AUC, and AUPR.
Characteristics analysis and ablation study

Runtime analysis of the compared methods

We conduct the experiments to calculate the mean runtime of ABCD-GGNN and other compared baselines on both ERα value prediction and ADMET property prediction tasks. All experiments are conducted on NVIDIA GeForce RTX 2070. All deep learning methods are set with early stopping. Detailed statistics are shown in Table 4. It can be seen that all deep learning methods take more time compared with traditional machine learning methods. In addition, our proposed ABCD-GGNN takes the most runtime, but the runtime of ABCD-GGNN is still on the same order of magnitude as the other deep learning methods. Since all the prediction tasks are conducted through inductive representation learning, overall, the runtimes of all these methods are acceptable.

Ablation study of the two representation modules in ABCD-GGNN

To demonstrate the effectiveness of both representation readout: discrete descriptor representation and topological graph representation, we take ablation study on the ADMET prediction task. The results are shown in Table 5. It can be seen that the performance of ABCD-GGNN is better than any single representation readout, which demonstrates that
both representation readouts contribute to the final representation and are complementary to each other. Meanwhile, the two representation modules are effectively integrated according to the hyper parameter $\lambda$.

**Ablation study of the hyper parameter $\lambda$.**

In addition, since our designed hyper parameter $\lambda \in (0, 1)$ controls the trade-off between the two views of representation, we also conduct the ablation study to seek the optimal value of $\lambda$ for anti-breast cancer candidate drug selection. Figure 3 exhibits the performance of ABCD-GGNN with a varying $\lambda$ on ADMET prediction tasks. $\lambda = 1$ means we only utilize the topological molecular graph representation, and $\lambda = 1$ means we only utilize the discrete property representation. On all the five property prediction tasks, the precision is consistently higher with larger $\lambda$ value. This can be explained by the high performance of topological molecular graph representation. The model reaches its best when $\lambda = 0.6$, performing slightly better than only utilizing topological molecular graph representation.

**Ablation study of the pooling operation in the readout stage**

We designed fusion strategy in the readout stage of ABCD-GGNN, which utilizes both average pooling and max pooling operations to better represent each compound. To demonstrate the effectiveness of the fusion of the two pooling operations, we take the ablation study in terms of the pooling operation selection as is shown in Table 6. It can be seen that our designed fusion strategy does contribute to better representation performance for ADMET prediction tasks. Meanwhile, the average pooling and max pooling operations are complementary to each other.
Ablation study of the interaction step in molecular graph representation

Interaction step is the key parameter which controls the global information interaction of molecular graph representation. Therefore, we conduct the ablation study to seek the optimal number of interaction step for anti-breast cancer candidate drug selection. Figure 4 presents the performance of molecular graph representation with a varying number of the graph layer on ADMET prediction tasks. The result reveals that with the increment of the layer, a node could receive more information from high-order neighbors and learn its representation more accurately. Nevertheless, the situation reverses with a continuous increment, where a node receives from every node in the graph and becomes over-smooth. On all the five property prediction tasks, the representation method overall reaches its best when interaction step is set as 2.

The effect of XGBoost feature selection

In the stage of discrete molecular descriptor representation, a XGBoost is adopted to select the top 50 molecular descriptors, which is intended to reduce the redundancy of the original 729 molecular descriptors. To demonstrate the effectiveness of the XGBoost, we conduct analysis on the 50 molecular descriptors from the XGBoost. The scores of and the heatmap of the selected 50 molecular descriptors are shown in Fig. 5a,
b, respectively. It can be seen that the correlation between the selected descriptors are commonly low, which fits our expectation that the 50 molecular descriptors should be in low redundancy.

**The effect of the ranking operator**
We also conduct result analysis to demonstrate the biological rationality of the ranking operator for the final candidate drug selection. We first comprehensively consider the predicted value of the model’s biological activity value and the classification value of the ADMET property, and perform a cluster analysis on it, as shown in Fig. 6. For example, to analyze the results of cluster analysis, SMILES35 and SMILES33 are classified into one category, and SMILES3 and SMILES2 may also be the same category. Figure 7 shows
the quantitative evaluation of the anti-breast cancer ability of the compounds based on the scoring mechanism, where the horizontal axis arranges the compounds in the order of the cluster analysis results in Fig. 6, and the vertical axis represents the scoring of the compounds in this article. It can be seen that the compounds with similar scores are close in the horizontal direction, that is, they are also classified in the same category (with similar properties) in the cluster analysis. For example, two compounds of SMILES35 and SMILES33 belong to the same class and have similar scores. In other words, the ranking operator can make a reasonable quantitative assessment of the compound’s anti-breast cancer ability based on the classification prediction results of the compound.

Discussion

We evaluated the effectiveness of ABCD-GGNN in predicting ERα, and the pharmacokinetic properties and safety of the compounds, by benchmarking on compound dataset containing SMILES and 729 molecular descriptors. In contrast to previous studies, ABCD-GGNN focuses on learning the the structure and substructure characteristics of a candidate drug topologically, and integrating with discrete molecular descriptors to form a more optimal molecular-level representation of feature of a drug.

The experimental results of our method ABCD-GGNN confirm two perspectives to improve the performance of methods for predicting the properties of molecular compounds. From a computational perspective, advanced artificial intelligence methods such as graph neural networks can be utilized to construct a better representation of molecular compound properties based on the structure and substructure of molecules. From a biological perspective, effective integration of structural and substructural features of molecules and other characteristics that reflect the properties of molecules (i.e., molecular descriptors) can better model the characteristic expression of molecular compounds and help researchers understand the biological mechanisms involved. Conclusions above are based on the facts that 1) molecular descriptors can determine the biological activity of compounds as independent variables; 2) graph neural networks enable global feature extraction to further enhance the molecular representation; and 3) as illustrated in Table 5, the ablation experimental results demonstrated that the integration of topological features and discrete descriptor features can further enhance the performance of molecular representation.

If a large number of molecular descriptor classes are available, we suggest using a regression model to evaluate the correlation of descriptors with compound properties and the coupling between descriptors, so as to reduce the redundancy and sparsity of the original molecular descriptors. We analyzed the original 729 molecular descriptors using the XGBoost model, and the results are shown in Table 1, where 50 molecular descriptors with low redundancy status were selected, and they had the highest correlation with the compound properties.

For the selection of anti-breast cancer drugs, we suggest a ranking operator consisting of feature binning and scorecard to select the appropriate anti-breast cancer drugs statistically. Figure 7 shows the quantitative evaluation of the anti-breast cancer ability of the compounds based on the scoring mechanism. Compounds with similar scores can remain similar in the clustering analysis, implying that the ranking operator can
comprehensively consider $\text{ER}_\alpha$, and the pharmacokinetic properties and safety of the compounds, which consists with the biological significance.

In summary, in this paper, we give full consideration to the high correlation between $\text{ER}_\alpha$ expression and breast cancer, and the significance of ADMET properties of a compound. By employing the ABCD-GGNN representation method, our designed framework can integrate multi-view features of compounds and efficiently select candidate drugs for researchers for further drug discovery. Given the universality and adaptability of molecular representation methods, it is expectable that such framework, with corresponding modification, can also be utilized for the research on other drug selection and contribute to intelligent administration in the pharmacology community.

**Conclusion**

In this paper, we propose the ABCD-GGNN representation method aiming at topologically representing the features of anti-breast cancer candidate drugs and predicting the $\text{ER}_\alpha$ value and ADMET properties of the organic compounds. With the ranking operator employed, research on the drug selection can be facilitated based on these significant metrics. Our proposed ABCD-GGNN representation method topologically learns both the implicit structure and substructure characteristics of a candidate drug and then deeply integrate them with explicit discrete molecular descriptors to strongly enhance the molecule-level representation. Extensive experiments conducted on our collected anti-breast cancer candidate drug dataset demonstrate that our proposed model outperforms all the other representative methods. Extended analysis also proves the biological rationality of our designed anti-breast cancer candidate drug selection strategy.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12859-022-04913-6.

**Acknowledgements**

This work was supported in part by Beijing Natural Science Foundation under Grants M22012, in part by the National Natural Science Foundation of China under Grant 72274022 and Grant 82071171, and in part by BUPT Excellent Ph.D. Students Foundation under Grant CX2022133.

**Author Contributions**

YG wrote the main manuscript text and led the method design and experiment implementation. SC and JT participated in the data preprocessing, experiment implementation, and results analysis. XF participated in the method design and revision of this paper. All authors read and approved the final manuscript.

**Availability of data and materials**

The datasets used and/or analysed during the current study available from the corresponding author on reasonable request.

**Declaration**

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.
Competing interests
The authors declare that they have no conflict of interest.

Received: 28 May 2022   Accepted: 24 August 2022
Published online: 19 September 2022

References
1. Bray F, Ferlay J, Soerjomataram I, Siegel RL, Torre LA, Jemal A. Global cancer statistics 2018: Globocan estimates of incidence and mortality worldwide for 36 cancers: in 185 countries. CA Cancer J Clin. 2018;68:394–424.
2. Peng Y, Lin Y, Jing X-Y, Zhang H, Huang Y, Luo GS. Enhanced graph isomorphism network for molecular admet properties prediction. IEEE Access. 2020;8:168344–60.
3. Hajijoo M, Damavandi B, HooshSadat M, Sangi F, Mackey JR, Ciss CE, Greiner R, Damaraju S. Breast cancer prediction using genome wide single nucleotide polymorphism data. BMC Bioinform. 2013;14:3–3.
4. Li X, Triung BMT, Xu T, Liu L, Li J, Le TD. Uncovering the roles of micromRs/micrnas in characterising breast cancer subtypes and prognosis. BMC Bioinform. 2021;22:1–22.
5. Hondermarck H, Vercoutter-Edouart AS, Révillion J, el-Yazidi-Belkoura I, Nurcombe V, Peyrat JP. Proteomics of breast cancer for marker discovery and signal pathway profiling. PROTEOMICS. 2001;1:1216–32.
6. Waks AG, Winer EP. Breast cancer treatment: a review. JAMA. 2019;321:288–300.
7. Rodriguez JC, Merino GA, Llera AS, Fernández EA. Massive integrative gene set analysis enables functional characterization of breast cancer subtypes. J Biomed Inf. 2019;93:103157. https://doi.org/10.1016/j.jbi.2019.103157.
8. John S, Thangapandian S, Arooj M, Hong J-C, Kim K, Lee KW. Development, evaluation and application of 3d qsar pharmacophore model in the discovery of potential human renin inhibitors. BMC Bioinform. 2011;12:4–4.
9. Parvathareni V, Kulkami NS, Muth A, Gupta V. Drug repurposing: a promising tool to accelerate the drug discovery process. Drug Discov Today. 2019;24:2076–85.
10. Arrowsmith JE. Trial watch: phase iii and submission failures: 2007–2010. Nat Rev Drug Discov. 2011;10:87–87.
11. Lee K, Lee M, Kim D. Utilizing random forest qsar models with optimized parameters for target identification and its application to target-fishing server. BMC Bioinform. 2017;18:75–86.
12. Gaur AS, Nagamani S, Tanneeru K, Druzhilovskiy D, Rudik A, Poroikov V, Narahari Sastry G. Molecular property diagnostic suite for diabetes mellitus (mpdsuite): an integrated web portal for drug discovery and drug repurposing. J Biomed Inf. 2018;85:114–25. https://doi.org/10.1016/j.jbi.2018.08.003.
13. Cui C, Ding X, Wang D, Chen L, Xiao F, Xu T, Zheng M, Luo X, Jiang H, Chen K. Drug repurposing against breast cancer by integrating drug-exposure expression profiles and drug-drug links based on graph neural network. Bioinformatics. 2021;37:2930–7.
14. Li X, Fourches D. Smiles pair encoding: a data-driven substructure tokenization algorithm for deep learning. J Chem Inf Model. 2021;61(4):1360–9.
15. Riau BPR, Afendi FM, Anisa R. Selection of compound group to identify the authenticity one of jamu product using the group lasso for logistic regression. J Phys Conf Ser. 2019;1341:092020.
16. Allouzi I, Ahmed MB. A 3d-cnn and svm for multi-drug resistance detection. In: CLEF 2018.
17. Matsumoto A, Aoki S, Ohwada H. Comparison of random forest and svm for raw data in drug discovery: prediction of radiation protection and toxicity case study. Int J Mach Learn Comput. 2016;6:145–8.
18. Fodeh SJ, Tiwari A. Exploiting medline for gene molecular function prediction via nfm based multi-label classification. J Biomed Inf. 2018;86:160–6. https://doi.org/10.1016/j.jbi.2018.08.009.
19. Wang Z, Wang Z, Huang Y, Lu L, Fu Y. A multi-view multi-omics model for cancer drug response prediction. Appl Intell. 2022.
20. Hirohara M, Saito Y, Koda Y, Sato K, Sakakibara Y. Convolutional neural network based on smiles representation of compounds for detecting chemical motif. BMC Bioinform. 2018;19:83–94.
21. Haneczok J, Delijewski M. Machine learning enabled identification of potential sars-cov-2 3clpro inhibitors based on fixed molecular fingerprints and graph-cnn neural representations. J Biomed Inf. 2021;119:103821–103821.
22. Moniz JRA, Pal CJ. Convolutional residual memory networks. arXiv: 1606.05262, 2016.
23. Krizhevsky A, Sutskever I, Hinton GE. Imagenet classification with deep convolutional neural networks. Commun ACM. 2012;60:84–90.
24. Kong W, Dong ZY, Jia Y, Hill DJ, Xu Y, Zhang H. Short-term residential load forecasting based on lstm recurrent neural network. IEEE Trans Smart Grid. 2019;10:841–51.
25. Berahmand K, Nasiri ES, Rostami M, Forouzandeh S. A modified deepwalk method for link prediction in attributed social network. Computing. 2021;103:2227–49.
26. Du Z-H, Wu Y-H, Huang Y-A, Chen J, Pan G-Q, Hu L, You Z, Li J. Graph Isl: an attention-based graph embedding model for predicting tf-target gene interactions. Brief Bioinform. 2022;23(3):bbac148.
27. Su X-R, Hu L, You Z, Hu P, Wang L, Zhao B. A deep learning method for repurposing antiviral drugs against new viruses via multi-view nonnegative matrix factorization and its application to sars-cov-2. Brief Bioinform. 2022;23(1):bbab526.
28. Haneczok J, Delijewski M. Machine learning enabled identification of potential sars-cov-2 3clpro inhibitors based on fixed molecular fingerprints and graph-cnn neural representations. J Biomed Inf. 2021;119:103821–103821. https://doi.org/10.1016/j.jbi.2021.103821.
29. Kipf TN, Welling M. Semi-supervised classification with graph convolutional networks. In: 5th International Conference on Learning Representations, ICLR 2017, Toulon, France, April 24–26, 2017, Conference Track Proceedings. OpenReview.net, 2017. https://openreview.net/forum?id=SJU4ayYgl.
30. Liu T, Cui J, Zhuang H, Wang H. Modeling polypharmacy effects with heterogeneous signed graph convolutional networks. Appl Intell. 2021;51:8316–33.
31. Rostami M, Forouzandeh S, Berahmand K, Soltani M, Shahsavari M, Oussalah M. Gene selection for microarray data classification via multi-objective graph theoretic-based method. Artif Intell Med. 2022;123:102228.
32. Rostami M, Forouzandeh S, Berahmand K, Soltani M. Integration of multi-objective pso based feature selection and node centrality for medical datasets. Genomics. 2020;112(6):4570–84.
33. Wang Y, Min Y, Chen X, Wu J. Multi-view graph contrastive representation learning for drug-drug interaction prediction. In: Proceedings of the Web Conference 2021.
34. Gao Y, Fu X, Liu X, Zhou K, Wu J. Smp-graph: Structure-enhanced unsupervised semantic graph representation for precise medical procedure coding on emrs. In: 2021 IEEE International Conference on Bioinformatics and Biomedicine (BIBM), 2021;1303–1308.
35. Gao Y, Fu X, Ouyang T, Wang Y. Eeg-gcn: spatio-temporal and self-adaptive graph convolutional networks for single and multi-view eeg-based emotion recognition. IEEE Signal Processing Letters. 2022;29:1574–8.
36. Duvenaud DK, Maclaurin D, Aguilera-Iparraguirre J, Gómez-Bombarelli R, Hirzel TD, Aspuru-Guzik A, Adams RP. Convolutional networks on graphs for learning molecular fingerprints. 2015; arXiv: 1509.09292.
37. Feinberg EN, Joshi EM, Pande VS, Cheng AC. Improvement in admet prediction with multitask deep featureization. J Med Chem. 2020;63:8835–48.
38. Montanari F, Kuhnke L, ter Laak A, Clevert D-A. Modeling physico-chemical admet endpoints with multitask graph convolutional networks. Molecules. 2019;25:44.
39. Feinberg EN, Sheridan R, Joshi EM, Pande VS, Cheng AC. Step change improvement in admet prediction with potential deep featureization. 2019; arXiv: 1903.11789.
40. Sun M, Zhao S, Gilvary C, Elemento O, Zhou J, Wang F. Graph convolutional networks for computational drug development and discovery. Brief Bioinform. 2020;21(3):919–35.
41. Chen C, Ye W, Zuo Y, Zheng C, Ong SP. Graph networks as a universal machine learning framework for molecules and crystals. Chem Mater. 2019;31(9):3564–72.
42. Li Y, Tarlow D, Brockschmidt M, Zemel RS. Gated graph sequence neural networks. CoRR 2016. arXiv: 1511.05493.
43. Feng H, Zhang L, Li S, Liu, Yang T, Yang P, Zhao J, Arkin IT, Liu H. Predicting the reproductive toxicity of chemicals using ensemble learning methods and molecular fingerprints. Toxicol Lett. 2021;340:4–14.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.