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## wSDTNBI: a novel network-based inference method for virtual screening†

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In recent years, the rapid development of network-based methods for the prediction of drug-target interactions (DTIs) provides an opportunity for the emergence of a new type of virtual screening (VS), namely, network-based VS. Herein, we reported a novel network-based inference method named wSDTNBI. Compared with previous network-based methods that use unweighted DTI networks, wSDTNBI uses weighted DTI networks whose edge weights are correlated with binding affinities. A two-pronged approach based on weighted DTI and drug-substructure association networks was employed to calculate prediction scores. To show the practical value of wSDTNBI, we performed network-based VS on retinoid-related orphan receptor  $\gamma t$  (ROR $\gamma t$ ), and purchased 72 compounds for experimental validation. Seven of the purchased compounds were confirmed to be novel ROR $\gamma t$  inverse agonists by *in vitro* experiments, including ursonic acid and oleanonic acid with IC $_{50}$  values of 10 nM and 0.28  $\mu$ M, respectively. Moreover, the direct contact between ursonic acid and ROR $\gamma t$  was confirmed using the X-ray crystal structure, and *in vivo* experiments demonstrated that ursonic acid and oleanonic acid have therapeutic effects on multiple sclerosis. These results indicate that wSDTNBI might be a powerful tool for network-based VS in drug discovery.

#### 1 Introduction

Lead discovery is an essential step in the pipeline of drug discovery.1 However, identification of lead compounds by in vitro and in vivo experiments is often expensive and timeconsuming. To improve efficiency and reduce costs, virtual screening (VS), a computer-aided technology aiming to identify promising compounds for targets of interest from chemical libraries, has been widely used in lead discovery.<sup>2,3</sup> Currently, there are two principal types of VS, namely, structure-based and ligand-based VS.3,4 Ligand-based VS can be further divided into several subtypes, such as pharmacophore-based,5 similaritybased, and machine learning-based. As their names imply, structure-based VS uses the information of three-dimensional (3D) structures of targets to identify new ligands, while ligandbased VS uses the information of ligands. To date, these two types of VS have achieved great success in drug discovery.3,9,10 Meanwhile, with the rapid development of network pharmacology, the drug discovery paradigm has been shifting from the linear mode of "one drug  $\rightarrow$  one target  $\rightarrow$  one disease" to the

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network mode of "multiple drugs  $\rightarrow$  multiple targets  $\rightarrow$  multiple diseases". <sup>11,12</sup> A series of network-based methods have been proposed for the prediction of drug-target interactions (DTIs), <sup>13</sup> which provides an opportunity for the emergence of a new type of VS, namely, network-based VS.

Representative network-based methods, including networkbased inference (NBI) methods, 14-16 random walk-based methods,17 and local community-based methods,18 are derived from link prediction algorithms in complex networks.19 Compared with structure-based methods, network-based methods do not rely on 3D structures of targets.13 Moreover, compared with supervised machine learning-based methods, network-based methods do not rely on experimentally determined inactive DTIs, namely, negative samples.<sup>13</sup> For example, our previously developed NBI only uses a known DTI network to predict potential targets for drugs in the DTI network.14 No other information such as 3D structures of targets and negative samples is required. The two improved versions of NBI, substructure-drug-target NBI (SDTNBI) and balanced SDTNBI (bSDTNBI),15,16 use both DTI and drug-substructure association (DSA) networks as inputs, and hence can predict potential targets for not only drugs within the DTI network, but also various compounds outside the DTI network. Based on the predicted DTIs, it is possible to explore the molecular mechanisms of the therapeutic and adverse effects of approved drugs and natural products, 15,20,21 and identify active compounds for targets of interest.15 Using bSDTNBI in combination with in vitro

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experiments, we discovered new compounds targeting estrogen receptor α,15 prostaglandin E2 receptor EP4 subtype,21 and NAD(P)H:quinone oxidoreductase 1 (NQO1).22 These applications indicate that network-based methods, especially our NBI methods, have the potential to be applied in VS.

Despite the successful applications of network-based methods, there is still a weakness, namely, prediction scores are not correlated with binding affinities. For each predicted DTI between a drug and a target, the prediction score only measures the probability that the drug can interact with the target, rather than the potency of the drug against the target. 14-16 A higher score does not mean a stronger interaction. This weakness makes it hard to quantitatively assess the activities of the screened compounds in network-based VS, just like scoring functions do in structure-based VS.23 The possible reason for this weakness is that network-based methods are usually based on unweighted DTI networks, whose edge weights are always equal to a constant (e.g., 1). Binding affinities have not been considered in network construction and network-based prediction vet. Hence, it is necessary to improve networkbased methods by introducing binding affinity data in order to make them more suitable for VS.

In this study, we developed a novel network-based method named weighted substructure-drug-target NBI (wSDTNBI). Compared with previous network-based methods that use unweighted DTI networks, wSDTNBI uses weighted DTI networks whose edge weights are correlated with binding affinities. As shown in Fig. 1, a two-pronged approach based on DTI and DSA networks was employed to calculate prediction scores for all possible DTIs. This improved method can output prediction scores correlated with binding affinities, and help to find more active compounds with higher activities for targets of interest. To show the practical value of wSDTNBI, we performed network-based VS on retinoid-related orphan receptor yt (RORyt, a specific isoform of RORy), and purchased 72 compounds for experimental validation. Seven of the purchased compounds were confirmed to be novel RORyt inverse agonists by in vitro experiments, including ursonic acid and oleanonic acid with IC50 values of 10 nM and 0.28 µM, respectively. Moreover, the direct contact between the best compound ursonic acid and RORyt was confirmed using the X-ray crystal structure, and in vivo experiments demonstrated that ursonic acid and oleanonic acid have therapeutic effects on multiple sclerosis (MS). The success rate of our network-based VS on RORyt (7/72  $\approx$  9.7%) is higher than the success rates of recent structure-based and deep learning-based VS on RORyt (only about 5%).24,25 These results indicate that wSDTNBI might be a powerful tool for network-based VS in drug discovery.

#### **Methods** 2

#### 2.1 Description of wSDTNBI

The schematic diagram of network-based VS with wSDTNBI is illustrated in Fig. 1. Just as its name implies, wSDTNBI uses a weighted DTI network and a DSA network as inputs to predict potential DTIs. In the weighted DTI network, nodes represent drugs and targets, and edges represent DTIs (Fig. 1a). A drug node and a target node are connected by an edge if the drug is known to interact with the target. The edge weights are not always equal to 1, but set to be positively correlated with binding affinities. In the DSA network, nodes represent drugs and substructures, and edges represent DSAs (Fig. 1b). A drug node and a substructure node are connected by an edge if the chemical structure of the drug contains the substructure. The DSA network generally contains DSAs for not only all drugs in the DTI network, but also novel compounds without any known targets, such as newly discovered natural products and newly synthesized compounds.

Assuming that there are a total of  $N_D$  drugs and novel compounds in the DTI and DSA networks, and a total of  $N_{\rm T}$ targets in the DTI network, the weighted DTI network can be represented as a  $N_D \times N_T$  adjacency matrix  $W_{DTI}$  (Fig. 1c). In this matrix,  $W_{\text{DTI}}(i,j)$  = the weight of the edge between drug  $D_i$  and target  $T_i$  if they are connected in the DTI network, otherwise = 0. Assuming that there are a total of  $N_S$  substructures in the DSA network, the DSA network can be represented as a  $N_{\rm D} \times N_{\rm S}$ adjacency matrix  $A_{DSA}$  (Fig. 1d). In this matrix,  $A_{DSA}(i,j) = 1$  if drug  $D_i$  and substructure  $S_i$  are connected in the DSA network, otherwise = 0. Based on the two networks represented by adjacency matrices  $W_{\rm DTI}$  and  $A_{\rm DSA}$ , prediction scores are calculated for all possible DTIs by the following two-pronged approach.

On one prong (Fig. 1, red arrows), our previously developed bSDTNBI is employed to calculate prediction scores. At first, the weighted DTI network is converted into an unweighted DTI network (Fig. 1e), whose adjacency matrix  $A_{DTI}$  can be obtained using the following equation:

$$A_{\text{DTI}}(i, j) = |\operatorname{sgn}(W_{\text{DTI}}(i, j))|$$

Then, bSDTNBI scores are calculated for all possible DTIs, based on a substructure-drug-target network constructed by integrating the unweighted DTI network and the DSA network. In the calculation of bSDTNBI scores, three tuneable parameters  $\alpha$ ,  $\beta$  and  $\gamma$  are used to address potential imbalances. <sup>15</sup> These bSDTNBI scores are subsequently normalized to lie between 0 and 1, and stored in a  $N_{\rm D} \times N_{\rm T}$  matrix  $S_{\rm norm}$  (Fig. 1f). A new parameter  $\delta$  is introduced to adjust the distribution of normalized scores. Specifically, assuming that  $S_{\text{bSDTNBI}}(i, j)$  is the bSDTNBI score of the DTI between drug  $D_i$  and target  $T_i$ , normalized bSDTNBI scores can be calculated using the following equations:

$$S_{\text{norm}}(i,j) = \frac{S_{\text{bSDTNBI}}(i,j)}{\max\{S_{\text{bSDTNBI}}(i,j),t(i)\}}$$

$$t(i) = \left(\delta + \sum_{l=1}^{N_{\mathrm{T}}} A_{\mathrm{DTI}}(i,l)\right) \text{th largest value in } S_{\mathrm{bSDTNBI}}(i,*)$$

On the other prong (Fig. 1, blue arrows), a similarity-based method is used to calculate prediction scores. At first, a  $N_{
m D}$  imes $N_{\rm D}$  drug similarity matrix is obtained by calculating similarity

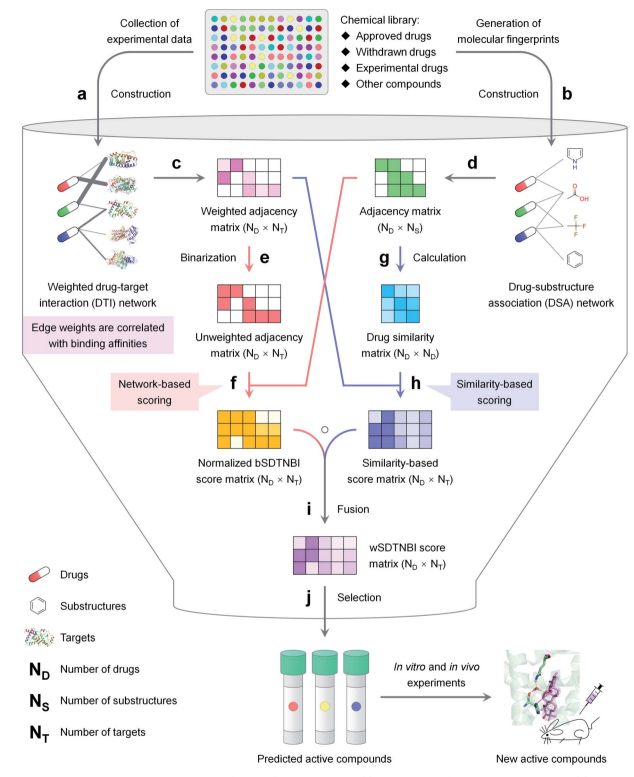


Fig. 1 The schematic diagram of network-based VS with wSDTNBI, including (a) construction of a weighted DTI network, (b) construction of a DSA network, (c) representing the weighted DTI network as an adjacency matrix, (d) representing the DSA network as an adjacency matrix, (e) converting the weighted DTI network into an unweighted DTI network, (f) calculation and normalization of bSDTNBI scores using the unweighted DTI network and the DSA network, (g) generation of a drug similarity matrix, (h) calculation of similarity-based scores using the weighted DTI network and the drug similarity matrix, (i) calculation of wSDTNBI scores by fusing normalized bSDTNBI scores and similarity-based scores, (j) selection of potential active compounds according to the calculated wSDTNBI scores. The details are described in the Methods and ESI† methods.

values for all possible drug pairs (Fig. 1g). Specifically, for each pair of drugs, two rows in  $A_{\rm DSA}$  corresponding to the two drugs are extracted as two vectors, and the similarity between the two drugs is measured using the Tanimoto coefficient:<sup>26</sup>

$$\frac{\vec{x} \cdot \vec{y}}{\left|\vec{x}\right|^2 + \left|\vec{y}\right|^2 - \vec{x} \cdot \vec{y}}$$

Then, similarity-based scores are calculated for all possible DTIs, and stored in a  $N_{\rm D} \times N_{\rm T}$  matrix  $S_{\rm sim}$  (Fig. 1h). Specifically, for each DTI between drug  $D_{\rm i}$  and target  $T_{\rm j}$ ,  $S_{\rm sim}(i,j)$  is set to the average of the edge weights of the DTIs between  $T_{\rm j}$  and its  $\varepsilon$  known ligands most similar to  $D_{\rm i}$  in the weighted DTI network. Compared with normalized bSDTNBI scores, these similarity-based scores may be more correlated with binding affinities.

The final prediction scores, which we called wSDTNBI scores, are obtained by fusing normalized bSDTNBI scores and similarity-based scores (Fig. 1i). Mathematically, the wSDTNBI score matrix  $S_{\text{wSDTNBI}}$  is the Hadamard product of the aforementioned two score matrices  $S_{\text{norm}}$  and  $S_{\text{sim}}$ :

$$S_{\text{wSDTNBI}} = S_{\text{norm}} \bigcirc S_{\text{sim}}$$

In general, only those DTIs having both high normalized bSDTNBI scores and high similarity-based scores will have high wSDTNBI scores. The wSDTNBI scores may help to identify active compounds with high binding affinities (Fig. 1j). The pseudo codes of wSDTNBI and previous network-based methods NBI, SDTNBI and bSDTNBI are provided in ESI† methods.

#### 2.2 Construction of network-based models

**2.2.1 Construction of DTI networks.** Drugs and related information were collected from DrugBank<sup>27</sup> (version 5.1.3), IUPHAR/BPS Guide to PHARMACOLOGY<sup>28</sup> (accessed in May 2019), KEGG DRUG<sup>29</sup> (accessed in May 2019) and Therapeutic Target Database<sup>30</sup> (accessed in May 2019). For each drug, the following three steps were performed using Open Babel<sup>31</sup> (version 2.4.1). Firstly, dative bonds were converted, and salt ions were removed. Secondly, the standardized chemical structures were represented in canonical SMILES format. Thirdly, three molecular descriptors were calculated, including molecular weight  $(M_{\rm W})$ ,  $\log P$  and the number of carbon atoms. Only those drugs that have  $M_{\rm W}$  <800 and more than one carbon atoms were retained.

Using the canonical SMILES of the retained drugs, DTI data with experimentally determined  $K_i$ ,  $K_d$ , half maximal inhibitory concentration (IC<sub>50</sub>), half maximal effective concentration (EC<sub>50</sub>), half maximal activity concentration (AC<sub>50</sub>) or potency values were extracted from ChEMBL<sup>32</sup> (version 25), BindingDB<sup>33</sup> (version 2019m4), IUPHAR/BPS Guide to PHARMACOLOGY<sup>28</sup> (accessed in May 2019) and PDSP  $K_i$  database<sup>34</sup> (accessed in May 2019). The extracted data were then filtered. Specifically, a DTI data item was retained if it met the following criteria: (i) the target is a protein from *Homo sapiens*; (ii) the target protein can be represented as an unique UniProt accession number; (iii) the

target protein was marked as "reviewed" in UniProt Knowledgebase<sup>35</sup> (accessed in June 2019); (iv) the interaction was not explicitly remarked as "inconclusive", "inactive" or "not active" in source databases.

Based on the processed DTI data, a local DTI network was constructed only using DTI data with  $K_{\rm i}$  values <10  $\mu$ M. The edge weights in the local network were set as  $-\log_{10}(K_{\rm i}/10~\mu{\rm M})$ . Meanwhile, a global DTI network was constructed using not only DTI data with  $K_{\rm i}$  values <10  $\mu$ M, but also DTI data with  $K_{\rm d}$ , IC<sub>50</sub>, EC<sub>50</sub>, AC<sub>50</sub> or potency values <10  $\mu$ M. The way of assigning edge weights in the global network was similar to that in the local network. When lacking  $K_{\rm i}$  values,  $K_{\rm d}$ , IC<sub>50</sub>, EC<sub>50</sub>, AC<sub>50</sub> or potency values were used as alternatives.

2.2.2 Construction of DSA networks. For each DTI network, nine DSA networks were constructed by generating nine types of molecular fingerprints for all drugs in the DTI network. Four of them, including Substructure (also known as FP4 in Open Babel,<sup>31</sup> 307 bits), MACCS (166 bits), PubChem (881 bits) and Klekota-Roth (KR, 4860 bits), were calculated by PaDEL-Descriptor<sup>36</sup> (version 2.21). The other five, including ECFP\_0, ECFP\_2, FCFP\_0, FCFP\_2 and FCFP\_4, were calculated by Discovery Studio (version 3.5).

**2.2.3 Model construction and optimization.** In addition to wSDTNBI, three previous network-based methods, namely, NBI, <sup>14</sup> SDTNBI, <sup>16</sup> and bSDTNBI, <sup>15</sup> were also used to construct prediction models. NBI models were constructed only using DTI networks, while the others were constructed using both DTI and DSA networks. The three parameters  $\alpha$ ,  $\beta$  and  $\gamma$  for each bSDTNBI model were optimized by combining grid search and 10-fold cross validation. Specifically, with a step length of 0.1, both  $\alpha$  and  $\beta$  were searched in the range of 0 to 1, and  $\gamma$  was searched in the range of -1 to 0. Based on the optimal parameters for bSDTNBI models, the remaining two parameters  $\delta$  and  $\varepsilon$  for each wSDTNBI model were further optimized by combining grid search and 10-fold cross validation.

#### 2.3 Evaluation of network-based models

2.3.1 Cross validation. The performance of all network-based models was evaluated by 10-fold cross validation, a method commonly used in previous studies of network-based DTI prediction. 14-16,21 In the 10-fold cross validation process for a model, DTIs in the DTI network for model construction were randomly divided into ten groups. One of them was extracted as the test set in turn, while the remaining networks were used as the training set. This resulted in ten pairs of training and test sets. For each pair of training and test sets, nodes which lost all its edges in the training set were removed from the two sets, and the following steps were performed to calculate evaluation indicators.

At first, all possible DTIs are predicted based on the training set. To avoid meaningless values, only those drugs that have DTIs in the test set were considered in the calculation of evaluation indicators. For each drug  $D_{\rm i}$ , its newly predicted DTIs were sorted by prediction scores in descending order, and divided into two classes. Specifically, the ones ranked in the top L were considered as positive, while the others were considered

as negative. After comparing the predicted positives and negatives with the test set, four numbers were calculated for drug  $D_i$ , including the number of true positives  $TP_i(L)$ , the number of false positives  $FP_i(L)$ , the number of true negatives  $TN_i(L)$  and the number of false negatives  $FN_i(L)$ . Denoting  $N'_D$  and  $N'_T$  as the numbers of drugs and targets considered in the calculation of evaluation indicators, recall R(L) and enhancement recall  $e_R(L)$ , two evaluation indicators depending on L, were calculated using the following equations:

$$R(L) = \frac{1}{N'_{\rm D}} \sum_{i=1}^{N'_{\rm D}} \frac{\text{TP}_i(L)}{\text{TP}_i(L) + \text{FN}_i(L)}$$

$$e_{\rm R}(L) = \frac{R(L)}{L/N'_{\rm T}}$$

Meanwhile, area under receiver operating characteristic curve (AUROC), an evaluation indicator independent of L, was calculated. The curve was obtained by plotting true positive rates (TPRs) against false positive rates (FPRs), after calculating a series of TPRs and FPRs at different L values using the following equations:

$$TPR(L) = \frac{\sum_{i=1}^{N'_{D}} TP_{i}(L)}{\sum_{i=1}^{N'_{D}} TP_{i}(L) + \sum_{i=1}^{N'_{D}} FN_{i}(L)}$$

$$FPR(L) = \frac{\sum\limits_{i=1}^{N_{D}^{'}} FP_{i}(L)}{\sum\limits_{i=1}^{N_{D}^{'}} FP_{i}(L) + \sum\limits_{i=1}^{N_{D}^{'}} TN_{i}(L)}$$

In addition, Pearson correlation coefficient (*r*), mean absolute error (MAE) and root mean square error (RMSE) were calculated by comparing the known edge weights of the DTIs in the test set with the prediction scores of these DTIs.

For each model, the aforementioned 10-fold cross validation process was repeated ten times, and each of the evaluation indicators was expressed as mean  $\pm$  standard deviation (SD).

2.3.2 External validation. Network-based models that showed good performance in 10-fold cross validation were used to calculate prediction scores between five nuclear receptors and their ligands. The five nuclear receptors are androgen receptor (UniProt accession number: P10275, gene symbol: AR), estrogen receptor  $\alpha$  (UniProt accession number: P03372, gene symbol: ESR1), peroxisome proliferator-activated receptor  $\alpha$  (UniProt accession number: Q07869, gene symbol: PPARA), peroxisome proliferator-activated receptor  $\gamma$  (UniProt accession number: P37231, gene symbol: PPARG) and retinoid X receptor  $\alpha$  (UniProt accession number: P19793, gene symbol: RXRA). The ligands were collected from ChEMBL³²² (version 25), and filtered

using the following criteria: (i) have not been used to construct the local and global DTI networks, (ii) have  $M_{\rm W}$  <800 and more than one carbon atom, and (iii) showed  $K_{\rm i}$  values <10  $\mu$ M on at least one of the five nuclear receptors.

In addition to network-based methods, molecular docking methods were also used to calculate prediction scores. The crystal structures of the five nuclear receptors were chosen according to DUD-E, 37 and downloaded from RCSB Protein Data Bank<sup>38</sup> (PDB codes: 2AM9 for AR, 1SJ0 for ESR1, 2P54 for PPARA, 2GTK for PPARG, and 1MV9 for RXRA). Each of the crystal structures was prepared by Protein Preparation Wizard in Schrödinger (version 2016; Schrödinger, LLC) with the following three steps. Firstly, the crystal structure was preprocessed using the default settings, and missing side chains were added by Prime (version 4.3; Schrödinger, LLC) if needed. Secondly, redundant chains and water molecules were deleted. Thirdly, the protonation states of residues were optimized at pH 7.0. Based on the prepared crystal structures, receptor grids were generated by Receptor Grid Generation in Schrödinger (version 2016; Schrödinger, LLC). In grid generation, the size of the enclosing box was set to maximum (36 Å) to ensure that most of the ligands in external validation sets can be docked into the prepared receptors. Then, the ligands of the five nuclear receptors were prepared by LigPrep in Schrödinger (version 2016; Schrödinger, LLC), and docked into the prepared receptors by Glide<sup>39</sup> (version 7.0; Schrödinger, LLC) in both standard precision (SP) and extra precision (XP) modes. The absolute values of docking scores were used as prediction scores.

Finally, the correlation between prediction scores and  $pK_i$  values was evaluated by the Pearson correlation test in R (version 3.3.3).

### 2.4 Construction of drug-target-immunological process network

The best wSDTNBI model Global-wSDTNBI-FCFP\_4 was used to predict 20 new targets for all drugs in the global DTI network. Known and predicted DTIs for the drugs marked as "approved" in DrugBank<sup>27</sup> and IUPHAR/BPS Guide to PHARMACOLOGY<sup>28</sup> were retained. Meanwhile, targets associated with immunological processes were collected from IUPHAR Guide to IMMUNOPHARMACOLOGY.<sup>28</sup> Then, a drug-target-immunological process network was constructed by integrating the DTIs for approved drugs and the target-immunological process associations. This network was visualized by Cytoscape<sup>40</sup> (version 3.4.0).

#### 2.5 Prediction and validation of RORyt ligands

**2.5.1 Network-based VS on RORγt.** The best wSDTNBI model Global-wSDTNBI-FCFP\_4 was used to perform network-based VS on RORγt (UniProt accession number: P51449, gene symbol: RORC). At first, a chemical library containing 730 natural products was obtained from TargetMol (Catalog No. L6000). After generating FCFP\_4 fingerprints for compounds outside the global DTI network, the newly obtained compound-substructure associations were added into the model to make it able to predict potential targets for these compounds. Then, the

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model was used to predict 20 new targets for not only the drugs within the global DTI network, but also compounds outside the global DTI network. If RORyt appeared in the newly predicted targets of a compound, the compound was regarded as a potential RORyt ligand. Finally, 72 of the predicted RORyt ligands were purchased from TargetMol for experimental validation.

2.5.2 Expression and purification of the RORγt-LBD. The human RORγt LBD (residues 262-518) was amplified by PCR from pCDNA2-FLAG-RORγt (a gift from Prof. Dan R. Littman, New York University). The RORyt LBD gene was subcloned into the pET-15b vector (Novagen, Madison, WI) with a N-terminal histidine-tag and transformed into Escherichia coli strain BL21 (DE3). The expression and purification of the recombinant RORγt-LBD protein was performed as described previously.41 The cells were cultured in lysogeny broth (LB) media supplemented with ampicillin (100 μg mL<sup>-1</sup>) at 37 °C up to an OD value of 0.8, and then induced with 0.1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) for protein expression. The incubation continued overnight with shaking at 16 °C. Cells were collected by centrifugation and resuspended in buffer A (20 mM Tris, pH = 7.0, 200 mM NaCl, 4% glycerol), followed by sonication. The lysate was centrifuged at 12 000 rpm for 30 min at 4 °C and the supernatant was loaded on a HisTrap nickel affinity column (GE Healthcare) pre-equilibrated with Buffer A. Protein was eluted with a gradient of 40-500 mM imidazole in buffer A. Fractions containing the protein of interest were pooled and dialyzed overnight against buffer B (20 mM Tris, pH = 7.0, 200 mM NaCl, 4% glycerol, 5 mM DTT). The RORγt-LBD protein was then concentrated using 10 kDa centrifugal filters from Merck Millipore (KGaA of Darmstadt, Germany) and analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). All the purification operations were performed at 4 °C.

2.5.3 Dual-luciferase reporter assay. To generate the RORγt-GAL4 fusion protein, human RORγt (from 97 to 516) was PCR-amplified from pCDNA2-FLAG-RORyt. The PCR product was ligated into the pFN11A (BIND) Flexi vector (Promega, Madison, WI) containing the yeast GAL4 DNA-binding domain upstream of the cloning site. This vector also expresses the Renilla luciferase under the control of SV40 promotor, allowing normalization for differences in transfection efficiency. The recombination plasmid was named RORyt-GAL4. The vector, pGL4.31 [luc2P/GAL4UAS/Hygro] (Promega, Madison, WI), contains five tandem GAL4 binding sites upstream of a minimal TATA box, which is upstream of a firefly luciferase gene that acts as a reporter for interactions between protein and the ligand.42 The pRL-SV40 Renilla luciferase control reporter vector was purchased from Promega (Madison, WI).

For luciferase activity assay, HEK293T cells were cotransfected with 2 µg ROR-GAL4, 2 µg pGL4.31 and 100 ng pRL-SV40 in a 6 cm dish using Lipofectamine 2000 (Invitrogen, USA). After 24 h of transfection, the cells were incubated with a variety of concentrations of compounds. After 24 h treatment, the cells were lysed with passive lysis buffer and luciferase activities were measured using a Dual-Glo Luciferase Assay

System (Promega, Madison, WI) using an EnVision Multilabel Plate Reader (PerkinElmer, Waltham, MA).

2.5.4 Homogeneous time resolved fluorescence (HTRF) assay. HTRF assay was used to evaluate the ability of compounds to interrupt the interaction of RORγt-LBD (262-518) and co-activator peptide SRC1-2. The recombinant RORγt-LBD protein was prepared as described aforementioned. Peptide SRC1-2 (sequence CPSSHSSLTERHKILHRLLQEGSPS) biotinylated at the N terminus was synthesized by GL Biochem Ltd (Shanghai, China). Europium-labelled anti-His antibody and XL665-labelled streptavidin were purchased from Cisbio (Codolet, France). For HTRF assay, 100 nM RORγt-LBD, 0.5 nM Eu-labelled anti-His antibody, 100 nM SRC1-2, 41.67 nM XL665labelled streptavidin, and 8-point serial dilutions of RORyt ligands from 0.01 nM to 100 µM were mixed and added to wells of low-volume 384-well white plates (PerkinElmer, Waltham, MA) with a final volume of 20 μL. All dilutions were prepared in 20 mM Tris, 200 mM NaCl, 5 mM dithiothreitol (DTT), 4% glycerol, pH = 7.0. After 2 h of incubation, the emissions were measured at 620 nm and 665 nm upon excitation at 330 nm. The inhibitory rates were calculated, and then concentrationresponse curves were generated and IC50 values were calculated by GraphPad Prism (GraphPad Software, CA, USA).

2.5.5 Intrinsic fluorescence-quenching based assay. Fluorescence quenching analysis was performed as described previously.43 Briefly, the purified RORγt-LBD protein was incubated with various concentrations of compounds at 4 °C for 1 h and fluorescence spectra were obtained using a Cary Eclipse fluorescence spectrophotometer (Agilent Technologies, CA, USA). The excitation wavelength was set at 280 nm and emission spectra ranging from 285 to 500 nm were recorded.

2.5.6 Isothermal titration calorimetry (ITC) assay. The binding affinity between RORyt-LBD and compounds was measured using an ITC200 instrument (Microcal, GE Healthcare, PA, USA) at 25 °C. The compounds and the protein were predissolved in the ITC buffer containing 20 mM Tris, 200 mM NaCl, 5 mM DTT, pH = 7.0. The final concentration of DMSO in the buffer is less than 1% of the total volume. Adding RORyt-LBD into compounds was performed by 19 identical injections of 2 µL with a duration of 4 s per injection spaced at intervals of 120 s between injections following an initial injection of 0.4 µL. The reaction heat of injecting protein into compounds was obtained, and the data were processed by the supplied MicroCal Origin (PA, USA). The thermodynamic parameters were calculated with the formula  $\Delta G = \Delta H - T\Delta S =$  $-RT \ln K$ , where  $\Delta G$  is the change in free energy,  $\Delta H$  is the change in enthalpy,  $\Delta S$  is the change in entropy, T is the experimental temperature, R is the gas constant, and K is the binding constant.

2.5.7 Drug affinity responsive target stability assay. Drug affinity responsive target stability (DARTS) assay was performed described previously.<sup>44</sup> RORγt-LBD (262-507)-SRC2.2 (KEKHKILHRLLQDSS) was subcloned into the pET-15b vector and the corresponding protein was named RORyt-LBD-SRC2.2. The expression and purification of the recombinant RORγt-LBD-SRC2.2 protein was performed as described for RORγt-LBD (residues 262-518). 60 μM RORγt-LBD-SRC2.2 was incubated with ursonic acid or oleanonic acid for 2 h at 4  $^{\circ}$ C. The samples were then warmed to 40  $^{\circ}$ C and proteolysed with 1  $\mu g$  pronase E (Yuanye Bio-Technology, Shanghai, China) for every 60  $\mu g$  protein for 10 min. Subsequently, the samples were resolved by SDS-PAGE, stained with Coomassie (SimplyBlue).

2.5.8 Cellular thermal shift assay. The cellular thermal shift assay (CETSA) was performed as described previously. 45 In brief, the human RORyt was amplified by PCR from pCDNA2-FLAG-RORγt, and then subcloned into the pCDNA3.0 vector. The recombinant pcDNA3.0-RORyt was transfected into HEK293T cells using Lipofectamine 2000 (Invitrogen, USA). After 12 h, the cells were cultured in fresh cell culture medium supplemented with 2 µM MG132 (Sigma-Aldrich, USA), and then incubated with ursonic acid (60 µM) or oleanonic acid (60 μM) for 2 h. After washing with phosphate-buffered saline (PBS), the cells were resuspended in PBS supplemented with 1% protease inhibitor (Sigma-Aldrich, USA), incubated at a series of temperatures for 5 min, and the cells were subsequently lysed by freeze-thaw three times using liquid nitrogen. Cell lysates were centrifuged at 12 000 rpm at 4 °C for 20 min. The supernatant samples were resolved by SDS-PAGE, and then transferred to a PVDF membrane. After incubation with the RORyt antibody and the horseradish peroxidase (HRP)-conjugated secondary antibody, the protein bands were analyzed using the ECL Western Blotting Substrate Kit (Thermo Fisher, USA).

2.5.9 Crystallization and structure determination. For cocrystallization, ursonic acid was solubilized in DMSO (50 mM) and added to a final concentration of 2 mM in the concentrated (10 mg ml<sup>-1</sup>) protein stock solution. The complex solution was incubated at 4 °C for 2 h, and then it was centrifuged at 12 000 rpm for 15 min prior to crystallization experiments. Diffraction-quality crystals of ligand-bound RORγt-LBD-SRC2.2 were grown by the sitting drop vapor diffusion technique. The reservoir solution contained 20% (w/v) PEG3350, 0.2 M CaCl<sub>2</sub>, and 5  $\mu$ M EDTA-2Na. Each 10  $\mu$ L sitting drop consisted of 5  $\mu$ L sample and 5 µL reservoir solution and was equilibrated against 0.5 ml reservoir solution at 20 °C. Crystals were flash-frozen in liquid nitrogen with cryoprotecant using corresponding reservoir condition supplemented with 30% glycerol. X-ray diffraction data were collected at 100 K on the synchrotron beamline BL19U1 at the Shanghai Synchrotron Radiation Facility (SSRF, Shanghai, China). Diffraction data were processed using XDS. The initial structure was determined by molecular replacement with CCP4 and PDB entry 4XT9 as an initial model. Multiple rounds of positional isotropic B-factor refinement were performed using phenix.refine followed by re-building in Coot. The coordinates and structure factors of the final model of ROR $\gamma$ t in complex with ursonic acid were deposited in PDB (PDB code: 6J3N).

## 2.6 Effects on experimental autoimmune encephalomyelitis (EAE)

**2.6.1 EAE disease induction and treatment.** All animal care and experimental procedures in this study were conducted in compliance with the protocol approved by the institutional Animal Care and Use Committee at East China University of

Science and Technology. Mice were housed in a flow cabinet with a 12 h light/dark cycle and fed with standard rodent chow and tap water ad libitum. For active EAE, typically 8-10 week old female C57BL/6 mice, purchased from the National Rodent Laboratory Animal Resources (Shanghai, China), were subcutaneously immunized in the flanks with 200 µg myelin oligodendrocyte glycoprotein peptide 35-55 (MOG<sub>35-55</sub>) (GL Biochem, Shanghai, China) emulsified in complete Freund's adjuvant (including M. tuberculosis H37Ra extract 4 mg ml $^{-1}$ , Beijing Biolead Biology Sci & Tech). 200 ng pertussis toxin was administered intraperitoneally (i.p.) to mice on day 0 and 2.46 The clinical signs of EAE mice were assessed daily using the following disease scoring system: 0, none; 1, limp tail or waddling gait with tail tonicity; 2, wobbly gait; 3, hindlimb paralysis; 4, hindlimb and forelimb paralysis; 5, death. 46 When indicated, ursonic acid and oleanonic acid were dissolved in 0.5% carboxymethylcellulose (CMC) at a final concentration of 5 mg ml<sup>-1</sup>. Mice were orally administered the suspension 200  $\mu$ L per mouse (50 mg kg<sup>-1</sup>) every day for 10 days from day 12. At day 21, the mice were killed and their spleens, spinal cords and brains were collected for further analysis.

2.6.2 Flow cytometry and cytokine detection. Preparation of individual cell suspensions from EAE mouse spleen for flow cytometric analysis was performed as described before. 47,48 Mice were killed and dealt with 75% alcohol pre-prepared, and then spleens were taken out and cut into fractionlets. Spleen tissue was crushed through the cell strainer in 2 mL HBSS and suspended cells were collected by centrifugation at 1500 rpm for 10 min. Ammonium-Chloride-Potassium (ACK) lysing buffer was used for the lysing of red blood cells in the mouse lymphocyte preps. Cells were washed with 3 mL HBSS three times, and then resuspended in RPMI-1640 with 10% FBS. For intracellular cytokine staining, the cells were stained with APC anti-mouse CD4 (eBioscience, CA, USA) at 4 °C for 1.5 h, and then fixed and permeabilized with 4% paraformaldehyde and 0.5% Triton-X100. Thereafter, the cells were stained with fluorophore-conjugated monoclonal antibody FITC-conjugated anti-mouse IL-17A (eBioscience, CA, USA) at 4 °C for 2.5 h. Flow cytometry was performed on a BD LSRII (BD Biosciences) instrument and analyzed using FlowJo (Tree Star, Inc., Ashland, OR).

2.6.3 *In vivo* imaging of brain inflammation. At day 9 post-administration, mice were shaved before the imaging to avoid the influence of autofluorescence and light scattering caused by hair. For blood–brain barrier permeability assay, Cy5.5 labeled bovine serum albumin (BSA-Cy5.5) with bright near infrared fluorescence was utilized as described before.<sup>49</sup> In brief, BSA-Cy5.5 (50 mg kg<sup>-1</sup>) was injected into the tail vein and optical imaging was performed approximately 6 h post-injection. The images were recorded using an IVIS Spectrum CT Imaging System (PerkinElmer, Inc., USA) with bright near infrared fluorescence probe. The IVIS settings were Epi-FI, Ex660/Em720, Binning 8, FStop 2, FOV D, Lamp Level High, and Height 1.50.

**2.6.4 ELISA.** The concentration of IL-17A in the spinal cord and brain supernatant was determined using an ELISA kit

according to the manufacturers' protocol (Neobioscience, Shenzhen, China).

2.6.5 Histochemical analysis. Mice were anaesthetized and perfused with PBS followed by 4% paraformaldehyde in PBS. For histochemical analysis, spinal cords were dissected from the spinal canal, processed into paraffin, and sectioned into 3  $\mu$ m slides. The axial sections were generated, randomly selected from each tissue block and stained with hematoxylin and eosin stain (H&E) and luxol fast blue (LFB).<sup>50</sup>

**2.6.6 Statistical analysis.** GraphPad Prism (GraphPad Software, CA, USA) was used to perform statistical analysis. All values were shown as mean  $\pm$  standard error of the mean (SEM) of three independent experiments. Student's *t*-test was used for pairwise comparisons of significance. \*P < 0.05 was considered statistically significant.

#### 3 Results

#### 3.1 Overview of network-based models

DTI networks are the basis of network-based models for DTI prediction. In this study, we constructed two comprehensive DTI networks by collecting experimental data from public databases. One is a local DTI network with 18 249 edges connecting 5560 drugs and 968 human target proteins, which was constructed only using DTI data with  $K_i$  values <10  $\mu$ M. The other is a global DTI network with 45 018 edges connecting 12 751 drugs and 1844 human target proteins, which was constructed using DTI data with  $K_i$ ,  $K_d$ ,  $IC_{50}$ ,  $EC_{50}$ ,  $AC_{50}$  or potency values <10 μM. These two DTI networks contain various types of drugs and target proteins. Drugs include not only approved drugs, but also withdrawn drugs, clinical drug candidates and experimental drug-like compounds. Target proteins include G protein-coupled receptors (GPCRs), nuclear receptors, enzymes, ion channels and transporters. In previous studies of networkbased DTI prediction, 14-16,21 edge weights in DTI networks were usually set to a constant (e.g., 1). Compared with them, edge weights in the two weighted DTI networks were greater than zero and positively correlated with binding affinities. In simple terms, the higher the edge weight is, the higher the binding affinity is. The detailed information of drugs (e.g., canonical SMILES, names, types, CAS registry numbers and Anatomical Therapeutic Chemical classification codes), targets (e.g., UniProt accession numbers, names and families) and DTIs (e.g., activity values and references) in the two DTI networks is provided in ESI data 1.†

We then constructed 18 DSA networks by generating nine types of fingerprints for all drugs in the two DTI networks. The nine types of fingerprints include four common fingerprints (FP4, KR, MACCS and PubChem) and five extended-connectivity fingerprints (ECFP\_0, ECFP\_2, FCFP\_0, FCFP\_2 and FCFP\_4). The statistics of the 18 DSA networks are given in ESI Table S1.† Based on the DTI and DSA networks, we constructed 18 wSDTNBI models, and optimized the parameters of these models by grid search. The optimized parameters are listed in ESI Table S2.† For the purpose of comparison, we also constructed models using previous network-based methods. Specifically, two NBI models were constructed based only on the

DTI networks, while 18 SDTNBI and 18 bSDTNBI models were constructed based on the DTI and DSA networks. Some of the models are freely available in our web server NetInfer<sup>51</sup> (http://lmmd.ecust.edu.cn/netinfer).

#### 3.2 Performance of network-based models

To compare wSDTNBI with previous network-based methods and find the best model, we evaluated the performance of all 56 models in this study by 10-fold cross validation. The evaluation indicators of local and global models, expressed as mean  $\pm$  SD, are given in ESI Tables S3 and S4,† respectively. At first, we focused on the performance of recovering missing DTIs, which was measured using three evaluation indicators: R,  $e_R$  and AUROC. Under the condition that 20 new DTIs were predicted for each drug (i.e., L = 20), 48 of the 56 models have R values >0.5 and  $e_{\rm R}$  values >24.3. For a model, the R value >0.5 means that the missing DTIs of each drug can be successfully recovered more than 50% in average, and the  $e_{\rm R}$  value >24.3 means that the observed R value is at least 24.3 times larger than the theoretical R value in random prediction. Meanwhile, all models have AUROC values >0.9, further indicating their good performance in recovering missing DTIs. Moreover, we found that the models constructed using FCFP\_4 fingerprints usually outperform the models constructed using other types of fingerprints, which is consistent with the finding in our previous study.22

We next focused on the correlation and difference between prediction scores and binding affinities, which were measured using the other three evaluation indicators: r, MAE and RMSE. As shown in Fig. 2, all NBI, SDTNBI and bSDTNBI models have r values very close to 0, which means that they cannot output prediction scores correlated with binding affinities. In contrast, the r values of local-wSDTNBI-FCFP\_4 and global-wSDTNBI-FCFP\_4, the best local and global models, are higher than 0.55 and comparable with those of molecular docking methods.<sup>23</sup> The high r values of local-wSDTNBI-FCFP\_4 and global-wSDTNBI-FCFP\_4 mean that they can output prediction scores positively correlated with binding affinities. Moreover, these two wSDTNBI models have MAE  $\leq$ 0.80 and RMSE  $\leq$ 1.07, indicating the small difference between prediction scores and binding affinities.

To further evaluate the correlation between prediction scores and binding affinities, we collected hundreds of nuclear receptor ligands outside the local and global DTI networks, including 717 AR ligands, 277 ESR1 ligands, 104 PPARA ligands, 345 PPARG ligands and 147 RXRA ligands. Using these nuclear receptor ligands, we further evaluated the performance of global-wSDTNBI-FCFP\_4 and the other two models constructed using FCFP\_4 fingerprints, global-bSDTNBI-FCFP\_4 and global-SDTNBI-FCFP\_4. Besides previous network-based methods, wSDTNBI was also compared with two commonly used molecular docking methods, Glide SP and XP.39 The correlation between prediction scores and  $pK_i$  values, measured using r and P values, are provided in ESI Fig. S1.† From the P values, we found that wSDTNBI performed well on all five nuclear receptors, while the other four methods only performed well on part of the nuclear receptors. From the r values, we found that

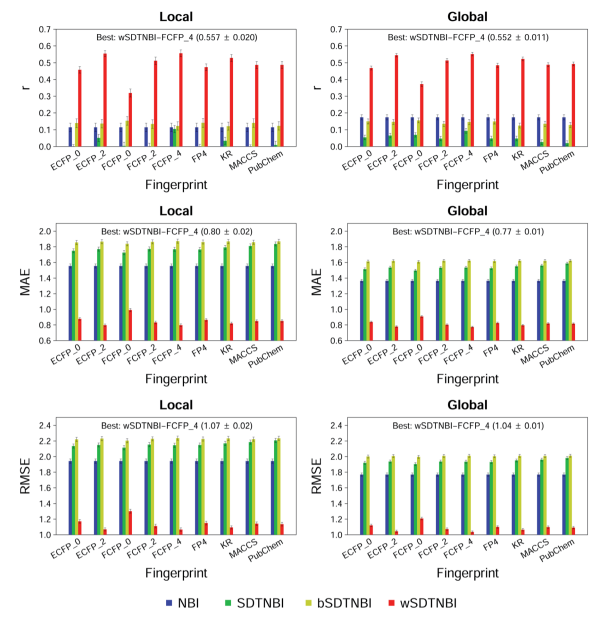


Fig. 2 Three evaluation indicators (*r*, MAE and RMSE) of network-based models in 10-fold cross validation. Results of the models constructed using NBI, SDTNBI, bSDTNBI and wSDTNBI are represented by blue, green, yellow and red bars.

wSDTNBI outperforms not only SDTNBI and bSDTNBI, but also Glide SP and XP.

In addition to evaluation indicators, we also focused on application ranges. Global models such as global-wSDTNBI-FCFP\_4 were constructed based on the global DTI network, and hence can predict active compounds for 1844 target proteins in the global network. Local models such as local-wSDTNBI-FCFP\_4 were constructed based on the local DTI network, and hence can only predict active compounds for 968 target proteins in the local network. It is obvious that the application ranges of global models are wider than those of local models.

Taken together, global-wSDTNBI-FCFP\_4 can output prediction scores correlated with binding affinities, and has

a wide application range. Hence, we recognized this wSDTNBI model as the best model. It is promising to apply global-wSDTNBI-FCFP\_4 in practical applications, such as network analysis and network-based VS.

#### 3.3 Analysis of predicted drug-target interactions

Using the best model global-wSDTNBI-FCFP\_4, we predicted 20 new targets for all 12 751 drugs in the global DTI network. Interestingly, several newly predicted DTIs can be validated using recently published articles. For example, abiraterone is known as an inhibitor of cytochrome P450 17A1 (UniProt accession number: P05093, gene symbol: CYP17A1) and an approved drug for the treatment of castration-resistant prostate cancer. <sup>52</sup> Herein, it was predicted to interact with AR. In a recent

study, abiraterone showed  $IC_{50}=7.6$  and  $9.4~\mu M$  in AR competitive binding and transcriptional assays, which means that abiraterone can directly bind to AR and inhibit AR transcriptional activity. Antimalarial drugs chloroquine and hydroxychloroquine were predicted to target sigma 1 receptor (UniProt accession number: Q99720, gene symbol: SIGMAR1). In a recent study, chloroquine and hydroxychloroquine were found to be sigma 1 receptor ligands with  $pK_i=7.1$  and 6.9, respectively. Besides these DTIs predicted for approved drugs, some DTIs predicted for natural products can also be validated

using the literature. For example, resveratrol and piceatannol were predicted to target 5-lipoxygenase (UniProt accession number: P09917, gene symbol: ALOX5). In a recent study, resveratrol and piceatannol were found to be 5-lipoxygenase inhibitors with  $IC_{50}=4.9$  and  $0.24~\mu\text{M}$ , respectively.<sup>54</sup> These results further indicate the reliability of global-wSDTNBI-FCFP\_4.

Our predicted DTIs may facilitate the discovery of new drugs for the treatment of complex diseases. Previous studies have demonstrated that innate immunity and inflammation play

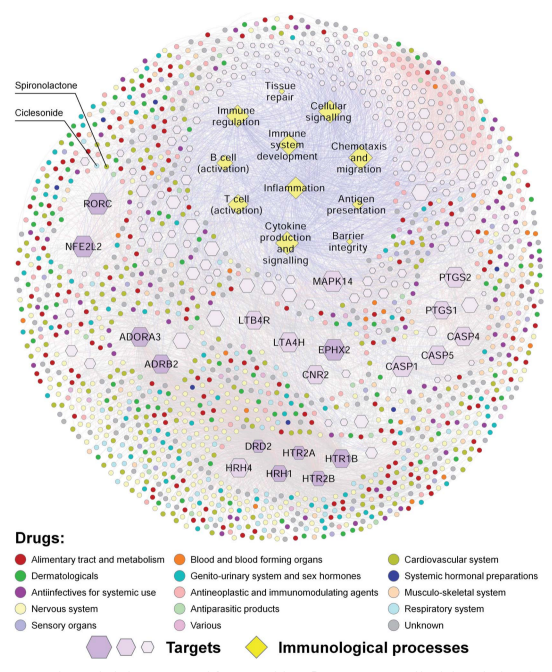


Fig. 3 The drug—target—immunological process network for approved drugs. Drugs are represented by circles and coloured according to their first-level Anatomical Therapeutic Chemical classification codes. Targets are represented by purple hexagons. Immunological processes are represented by yellow diamonds. Known and predicted DTIs are represented by light red and gray lines, respectively. Target-immunological process associations are represented by light blue lines.

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important roles in many age-related chronic diseases, such as neurodegenerative diseases, diabetes and cancer.<sup>28,55</sup> Meanwhile, autoimmune diseases, such as psoriasis, rheumatoid arthritis, inflammatory bowel disease and MS, are estimated to affect approximately 8-9% of the world population.<sup>56</sup> To explore the relationships among drugs, targets and immunological processes, we constructed a drug-target-immunological process network for approved drugs (Fig. 3). This network contains 11 immunological processes [including barrier integrity, inflammation, antigen presentation, T cell (activation), B cell (activation), immune regulation, tissue repair, immune system development, cytokine production and signaling, chemotaxis and migration, and cellular signaling], 247 human target proteins associated with these immunological processes, and 1297 approved drugs that have known or predicted interactions with these targets. Ten of the targets, such as RORyt (UniProt accession number: P51449, gene symbol: RORC), were connected with more than 100 approved drugs. Our drugtarget-immunological process network may help to explain the molecular mechanisms of the effects of approved drugs on immunological processes, and discover new drugs for the treatment of inflammatory and immune diseases.

To confirm this view, we used RORγt as an example. Previous studies have shown that targeting T helper 17 (TH17) cells is a promising strategy for the treatment of various autoimmune diseases.<sup>57</sup> As the "master" transcriptional factor of T<sub>H</sub>17 cells, RORγt orchestrates the differentiation program of T<sub>H</sub>17 cells, and has emerged as an attractive target for autoimmune diseases.58 In our drug-target-immunological process network, several approved drugs such as spironolactone were predicted to target RORy and associated with immunological processes such as the activation of T cells. A previous study has shown that mineralocorticoids such as aldosterone and deoxycorticosterone acetate (DOCA) promote autoimmune damage by enhancing T<sub>H</sub>17-mediated immunity.<sup>59</sup> A more recent study has shown that spironolactone can decrease DOCA-salt-induced organ damage in rats by blocking the activation of TH17 cells and the downregulation of regulatory T lymphocytes.<sup>60</sup> These studies suggest that some approved drugs such as spironolactone may target RORyt and have therapeutic effects on autoimmune diseases.

#### 3.4 Discovery of new RORyt ligands

Using the best model Global-wSDTNBI-FCFP\_4, we performed network-based VS on RORyt, the aforementioned nuclear receptor associated with autoimmune diseases. Considering that natural products are important sources of new drugs over the past few decades, 61,62 we predict 20 new targets for not only all 12 751 drugs within the global DTI network, but also 468 natural products outside the global DTI network. The prediction results of the 12751 + 468 = 13219 compounds are provided in ESI data 2,† which may provide useful information for researchers interested in VS and network pharmacology. According to the prediction results, we obtained 512 potential RORyt ligands and found that approximately 262 of them were purchasable from TargetMol. After considering the price and availability, we purchased 72 compounds from TargetMol for

experimental validation. Most of the purchased compounds are approved drugs and natural products.

A cell-based nuclear receptor RORγt-GAL4 reporter assay was used to confirm the activity of these compounds affecting the transcriptional activity of RORyt. Seven of the 72 purchased compounds  $(7/72 \approx 9.7\%)$  showed inhibition of the transcriptional regulation of RORyt with IC50 values ranging from 0.10 μM to 4.97 μM (Fig. 4a), including ursonic acid, oleanonic acid, ciclesonide, AKT inhibitor VIII, BX-471, spironolactone and veratramine. To validate the ability of these compounds to interact with RORyt directly, co-activator peptide SRC1-2 with a biotin label was used to evaluate the affinity of the seven compounds to RORyt by HTRF technology. All seven compounds showed dose dependent inhibition of RORyt binding to SRC1-2 with IC<sub>50</sub> values ranging from 10 nM to 19.81 μM (Fig. 4a), further indicating that they are potent RORγt inverse agonists.

#### 3.5 Further investigation of ursonic acid and oleanonic acid

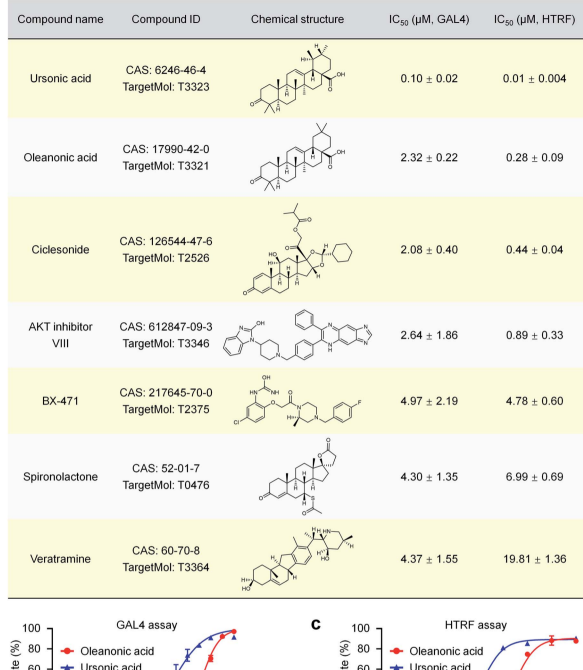
We turned to focus on ursonic acid and oleanonic acid, and the two natural products showed high activities in both dualluciferase reporter and HTRF assays (Fig. 4), which may provide useful information for further development.

Considering the presence of tryptophan residues in RORyt-LBD (Trp 314 and Trp 317), an intrinsic fluorescencequenching based assay was used to further verify the direct interactions between the two compounds and RORyt-LBD.63 The intrinsic fluorescence spectra were measured after RORyttreated with increasing concentrations compounds. Fig. 5a showed that ursonic acid and oleanonic acid dose-dependently induced fluorescence quenching of RORyt-LBD, indicating that they are capable of binding to RORγt-LBD.

The thermodynamic parameters of the interaction between RORyt-LBD and ursonic acid or oleanonic acid were examined by ITC. As shown in Fig. 5b, ursonic acid and oleanonic acid bound to ROR $\gamma$ t-LBD with  $K_d$  values of 3.04  $\mu$ M and 1.06  $\mu$ M, respectively. According to the changes in free energy and enthalpy ( $\Delta G$  and  $\Delta H$ ), the binding of ursonic acid and oleanonic acid to RORγt-LBD is based on hydrogen bonds, van der Waals bonds and hydrophobic effects as indicated by the negative binding enthalpy  $(\Delta H)$  and entropy factor  $(T\Delta S)$ . Although the  $K_d$  values of the two compounds are similar, ursonic acid has more strong hydrogen bonds and van der Waals bonds as indicated by the bigger enthalpy change. The ITC results indicated that ursonic acid and oleanonic acid could interact with RORyt-LBD directly.

Binding of small compounds may stabilize its target protein and reduce the protease sensitivity of its target protein.44 We also run DARTS44 assay to identify the binding target of ursonic acid and oleanonic acid. As shown in Fig. 5c, proteolysis of RORγt-LBD-SRC2.2 by the pronase E was clearly decreased by the presence of ursonic acid or oleanonic acid in a dosedependent manner, indicating that the binding of ursonic acid or oleanonic acid reduced the pronase E sensitivity of RORγt-LBD-SRC2.2.

a



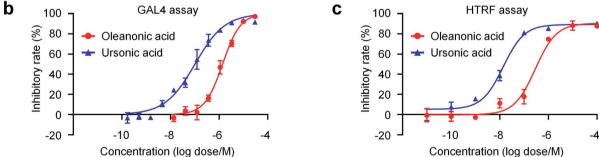


Fig. 4 Seven newly discovered ROR $\gamma$ t inverse agonists. (a) The names, structures and IC $_{50}$  values of these ROR $\gamma$ t inverse agonists. (b) The doseresponse curves of ursonic acid and oleanonic acid in GAL4 assay. (c) The doseresponse curves of ursonic acid and oleanonic acid in HTRF assay.

To further validate the interaction between ursonic acid or oleanonic acid and ROR $\gamma$ t in a cellular context, we performed CETSA<sup>64</sup> in the HEK293T cell line exogenously expressing

ROR $\gamma$ t. As shown in Fig. 5d, both ursonic acid and oleanonic acid significantly increased the  $T_{\rm m}$  values of ROR $\gamma$ t, suggesting that both of them stabilized ROR $\gamma$ t through the direct

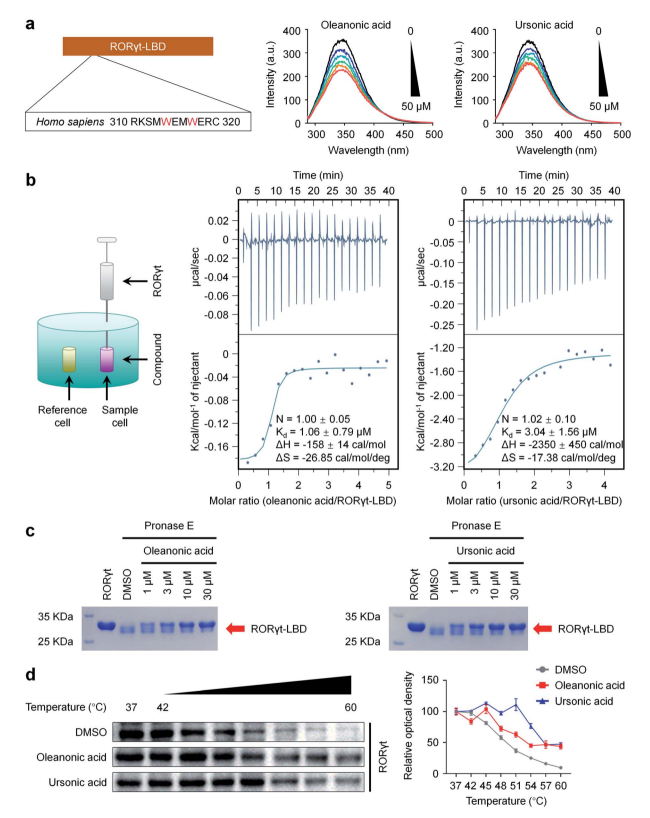


Fig. 5 The further experimental validation for ursonic and oleanonic acid binding to RORγt directly. (a) Binding of ursonic acid and oleanonic acid to RORγt-LBD was determined by intrinsic fluorescence quenching assay. RORγt-LBD (30 μM) was incubated with increasing concentrations of ursonic acid or oleanonic acid (0 μΜ, 10 μΜ, 20 μΜ, 30 μΜ, 40 μΜ, and 50 μΜ) for 30 min. The fluorescence emission (285–500 nm) was measured with an excitation wavelength of 280 nm. (b) Thermodynamic characterization of the interaction between RORyt-LBD and ursonic acid or oleanonic acid by ITC. Compounds were titrated with RORyt-LBD in ITC buffer as mentioned in the Methods. Binding curves were fitted as a single binding event. (c) Purified RORyt-LBD-SRC2.2 protein (60 μM) was incubated with indicative concentrations of compounds at 40 °C for 10 min and then subjected to SDS-PAGE and Coomassie (SimpleBlue)-staining. (d) CETSA showed that ursonic acid and oleanonic acid stabilized their target protein RORyt in HEK293T cells. Cells were treated with compounds for 2 h. Quantitative analysis according to the western blotting is shown in the right.

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interactions. All the above findings indicated that ursonic acid and oleanonic acid indeed bound with their target protein RORγt.

#### Crystal structure of RORyt in complex with ursonic acid

In addition, we obtained the crystal structure of RORγt-LBD-SRC2.2 in complex with ursonic acid at a resolution of 1.99 Å (PDB code: 6J3N) (Fig. 6a and ESI Table S5†), further confirming the direct interaction between ursonic acid and RORyt. The RORγt-LBD-SRC2.2 structure was solved in the space group  $P4_12_12$  with a polypeptide chain per symmetric unit. Common to most nuclear receptor LBDs, this RORyt-LBD-SRC2.2 structure presented a canonical three-layered helix sandwich folded by 12  $\alpha$ -helices and three  $\beta$ -strands (Fig. 6a). X-ray results showed that ursonic acid was buried in the canonical orthosteric ligand pocket (Fig. 6a) and formed van der Waals interactions with residues located in the orthosteric binding site, including Gln 286, Leu 292, Leu 324, Arg 367, Phe 377, Leu 391 and Ile 400 (Fig. 6b). Besides, Gln 286 and His 323 formed a network of hydrogen bonds with ursonic acid mediated by a water molecule (Fig. 6b). Ursonic acid failed to form hydrogen bonds with His 479 in H11 and Tyr 502 and Phe 506 in AF2, and hence behaved as an "agonist lock untouched" inverse agonist of RORyt.65 The crystal results provide useful information on structural modification of ursonic acid and help find new RORγt inverse agonists with higher activities.

#### 3.7 RORyt inhibition ameliorated MOG-induced chronic progressive EAE model

MS is a chronic inflammatory disease of the central nervous system that affects more than 2 million people worldwide,66 and one of the common causes of serious physical disability in young adults.67 As mentioned above, RORγt is a potential therapeutic target for autoimmune diseases such as MS. Our newly discovered RORyt inverse agonists may have therapeutic

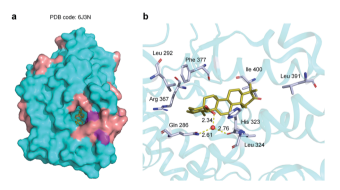


Fig. 6 X-ray structure of ursonic acid with RORγt-LBD-SRC2.2 (PDB code: 6J3N). (a) The overview of the crystal structure of ROR $\gamma$ t-LBD-SRC2.2 with ursonic acid. (b) The binding mode of ursonic acid in the ligand binding pocket. RORyt-LBD-SRC2.2 is shown as surface or cartoon with cyan  $\alpha$ -helices and purple  $\beta$ -strands. Ursonic acid is drawn as yellow sticks. Key residues and water molecules are shown as grey sticks and red spheres respectively. Hydrogen bonds are indicated by yellow dotted lines. Oxygen and nitrogen are colored by red and blue, respectively.

effects on MS. Hence, we investigated the in vivo effects of ursonic acid and oleanonic acid on the disease process of EAE, a most commonly used animal model of human MS.68

As shown in Fig. 7a, we induced EAE in female C57BL/6 mice with MOG<sub>35-55</sub> and PTX, and the mice were orally administrated usronic acid and oleanonic acid every day from day 12. We observed that EAE mice started to exhibit clinical symptoms at day 13 after immunization (n = 8) (Fig. 7b). Compared with the EAE group, a significant delay of the disease onset was found in oleanonic acid-treated group mice with a mean disease onset date of 15.3  $\pm$  0.2 days (n = 8). Surprisingly, better therapeutic effects were found in ursonic acid-treated group mice with an average onset date of 17.3  $\pm$  0.5 days (n = 8), which may be attributed to its better inhibitory activity. In the course of EAE, body weight variation also reflects disease severity. As expected, the ursonic acid and oleanonic acid administration significantly reversed EAE induced mice body weight loss (Fig. 7c).

Demyelinated lesions and inflammation in the peripheral white matter of spinal cord are the main pathological features of the EAE model.69 To further assess the disease condition of EAE, representative sections of spinal cord from each group mice were stained with LFB and H&E. Consistent with decreased clinical symptoms, immune cell infiltrations were reduced in white matter of spinal cords from ursonic acid and oleanonic acid treated-group mice (Fig. 7d). Moreover, bovine serum albumin coupled with Cy5.5 (BSA-Cy5.5) was used to assess the degree of blood-brain barrier (BBB) damage, which is one of the early pathophysiological hallmarks of EAE. 49,70 Compared with the EAE group, the visualization of brain BSA-Cy5.5 accumulation imaging exhibited that ursonic acid and oleanonic acid remitted the EAE-induced BBB disruption obviously (Fig. 7e), and ursonic acid showed better therapeutic effects than oleanonic acid, which is consistent with their inhibitory activities reflected in vitro.

Accumulating evidence has established that IL-17-producing CD4<sup>+</sup> T cells, also known as T<sub>H</sub>17 cells, play a critical role in the pathogenesis of EAE. 71 To examine whether RORγt inhibition by compounds affects the differentiation of T<sub>H</sub>17 cells in vivo, we thereby performed intracellular cytokine staining to determine the population of T<sub>H</sub>17 cells in the spleen of mice. As demonstrated in Fig. 7f, ursonic acid and oleanonic acid treatment resulted in a reduced frequency of pro-inflammatory T<sub>H</sub>17 cells as detected by CD4 and IL-17 dual-staining. Meanwhile, we also detected the IL-17 content in the spinal cord and brain supernatant by ELISA.72,73 Consistent with decreased TH17 population, clinical, ELISA analysis of mice tissue samples revealed that ursonic acid and oleanonic acid effectively reduced IL-17A protein expression in brains and spinal cords (Fig. 7g).

Taken together, ursonic acid and oleanonic acid ameliorated the MOG-induced chronic progressive EAE model by targeting pro-inflammatory T<sub>H</sub>17 cells, especially ursonic acid. Therefore, they may serve as lead compounds for the treatment of MS.

#### Discussion

In this study, we proposed wSDTNBI, a novel network-based method integrating weighted DTI and DSA networks for DTI

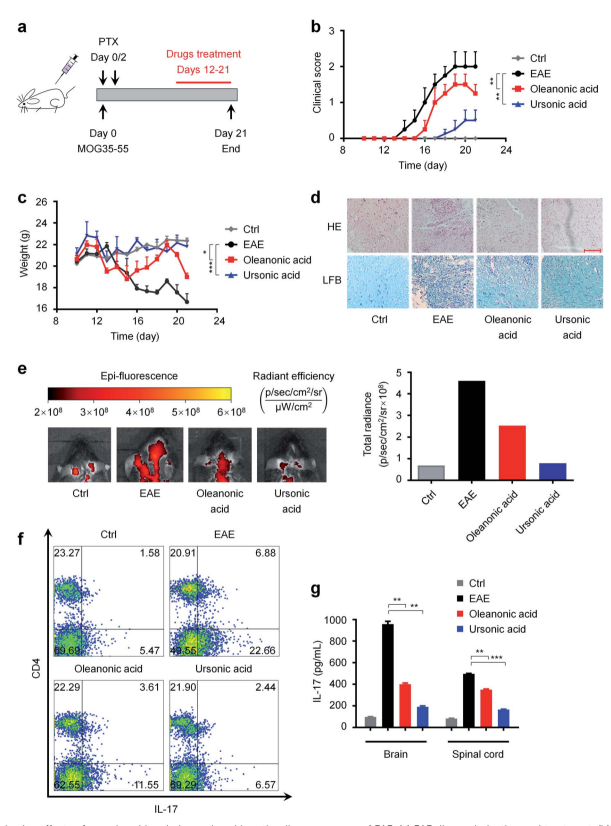


Fig. 7 In vivo effects of ursonic acid and oleanonic acid on the disease process of EAE. (a) EAE disease induction and treatment. (b) Disease severity assessed by the clinical scoring system ranging from 0 (no signs) to 5 (death). (c) Disease severity assessed by body weight variation. (d) Histochemical analysis of spinal cords of mice from different groups (scale bar =  $100 \mu m$ ). (e) EAE-induced BBB disruption reflected by BSA-Cy5.5 images in the brain. (f) Frequency of pro-inflammatory  $T_H17$  cells detected by CD4 and IL-17 dual-staining in the spleen of mice. (g) IL-17 expression in spinal cords and brains measured by ELISA.

prediction. wSDTNBI inherits the good features of previous network-based methods, including independent of 3D crystal structures and negative DTI samples. Furthermore, wSDTNBI can output prediction scores correlated with binding affinities, overcoming the limitation of previous network-based methods. This improvement makes it more suitable for VS, and lead to a new type of VS, namely, network-based VS. Compared with traditional methods, wSDTNBI have several obvious advantages.

Firstly, wSDTNBI can help to identify more active compounds with higher activities. In a previous study, Zhang et al. performed structure-based VS on RORγt.<sup>24</sup> Only one of 24 purchased compounds (1/24  $\approx$  4.2%) was validated by GAL4 reporter assay (IC<sub>50</sub> = 11.84  $\mu$ M). More recently, Zeng et al. developed a deep learning-based method for DTI prediction, and used the method to identify potential RORyt ligands.<sup>25</sup> Only one of 18 purchased compounds (1/18  $\approx$  5.6%) was validated by GAL4 reporter assay (IC<sub>50</sub> = 0.43  $\mu$ M). In this study, we performed network-based VS on RORyt using Global-wSDTNBI-FCFP\_4, and purchased 72 compounds for experimental validation. Seven of them (7/72  $\approx$  9.7%) showed IC<sub>50</sub> <10  $\mu$ M in the same GAL4 reporter assay, and the best compound ursonic acid even showed  $IC_{50} = 0.10 \mu M$  in GAL4 assay and  $IC_{50} = 10 nM$  in HTRF assay. Compared with the two previous VS studies on RORγt, 24,25 we obtained not only the highest success rate, but also the compound with the highest activity.

Secondly, wSDTNBI can help to identify some active compounds that other methods cannot. To show this, we inputted the chemical structures of the seven newly discovered RORyt ligands into six web servers for DTI prediction, and then investigated whether these web servers can correctly predict the interactions between these compounds and ROR $\gamma$ t. The six web servers include one pharmacophore-based, PharmMapper;<sup>74</sup> three similarity-based, SEA,75 ChemMapper,76 and SwissTargetPrediction;<sup>77</sup> and two machine learning-based, Target-Hunter,78 and TargetNet.79 After trying different parameter settings, still only three of the web servers, ChemMapper, SwissTargetPrediction and TargetHunter, reproduced a small number of the interactions, while the others totally failed (ESI Table S6†). Meanwhile, we also tried previous network-based methods. Unsurprisingly, only one, one and four compounds were predicted as potential RORyt ligands by NBI, SDTNBI and bSDTNBI, respectively (ESI Table S6†).

Thirdly, wSDTNBI can output comprehensive prediction results in a short time. In this study, we used the NVIDIA cuBLAS library, an GPU-accelerated implementation of the basic linear algebra subroutines (BLAS), to accelerate the matrix computation such as matrix multiplication. In three workstations with the hardware support of NVIDIA GeForce RTX 2080 Ti, RTX 2080 Super and GTX 1070, Global-wSDTNBI-FCFP\_4 only costs about 48, 53 and 125 seconds to calculate prediction scores for all possible interactions between 13 219 compounds and 1844 target proteins, which means that it can simultaneously perform VS on the 1844 targets in one or two minutes. As a comparison, we docked the seven newly discovered RORγt ligands into a crystal structure of RORγt by Glide SP and XP. However, the two docking experiments between seven

compounds and only one target cost approximately 7 and 50 minutes, respectively. The high speed allows large-scale VS to be performed using wSDTNBI. The comprehensive prediction results covering hundreds of targets may not only help to discard those compounds binding to the proteins closely associated with toxic or side effects [e.g., human ether-à-go-go-related gene (hERG) potassium channel], but also provide opportunities for multi-target drug discovery.

Despite the aforementioned advantages, there is still room for improvement of our method. wSDTNBI cannot be used for a protein without any known ligands, because this protein cannot be interlinked with the known DTI network. In addition, interaction types (*e.g.*, agonism or antagonism, activation or inhibition) are not considered yet. We are actively developing novel network-based methods to overcome these limitations. For example, we are trying to extend the prediction results to more proteins by introducing different types of protein sequence similarity and structure similarity data.

Besides the methodology we developed, the active compounds we discovered can also facilitate drug discovery and development. In this study, seven compounds were confirmed to be novel RORγt inverse agonists by in vitro experiments, and in vivo experiments demonstrated that two of them (ursonic acid and oleanonic acid) have therapeutic effects on MS, indicating that they may serve as lead compounds for the treatment of autoimmune diseases including MS. Considering that RORyt is a specific isoform of RORy, the newly discovered RORyt inverse agonists may be applied in other diseases associated with ROR $\gamma$ . For example, ROR $\gamma$  was found to drive the expression of AR and represent a potential therapeutic target for the treatment of prostate cancer in a previous study.80 As one of the newly discovered RORyt inverse agonists, spironolactone was found to reduce the incidence of prostate cancer in a recent study.81 The possible reason of this chemoprevention of prostate cancer is that spironolactone suppresses the expression of AR by inhibiting the transcriptional activity of RORγ. These studies indicate that the newly discovered RORyt inverse agonists may have therapeutic effects on prostate cancer. Moreover, we predicted absorption, distribution, metabolism, excretion and toxicity (ADMET) properties of the newly discovered RORyt inverse agonists, and provided the prediction results in ESI data 2.† Interestingly, most of the newly discovered RORyt inverse agonists, especially ursonic acid and oleanonic acid, were predicted to have not only good human intestinal absorption, but also low carcinogenicity, cardiotoxicity and hepatotoxicity. These predicted ADMET properties further suggest that our newly discovered RORγt inverse agonists are promising lead compounds for the treatment of autoimmune diseases and cancer.

#### 5 Conclusions

We developed a novel network-based method named wSDTNBI, which employed weighted DTI networks instead of unweighted DTI networks for the prediction of potential DTIs, and hence can help to find more active compounds with higher activities for targets of interest. In a network-based VS on RORγt, seven

of the 72 purchased compounds including ursonic acid and oleanonic acid were confirmed to be novel ROR $\gamma$ t inverse agonists by *in vitro* experiments. In addition, an X-ray crystal structure of ROR $\gamma$ t in complex with ursonic acid (PDB code: 6J3N) was obtained, and *in vivo* experiments showed the therapeutic effects of ursonic acid and oleanonic acid on MS. The success rate of our network-based VS on ROR $\gamma$ t is higher than the success rates of recent structure-based and deep learning-based VS on ROR $\gamma$ t. Our newly discovered ROR $\gamma$ t inverse agonists and the cocrystal structure may facilitate the discovery of more effective drugs for the treatment of autoimmune diseases including MS. Our methods and workflow can be easily extended to other targets for other diseases. We believe that wSDTNBI would become a powerful tool for network-based VS in drug discovery.

#### Data availability

The detailed information of drugs, targets and DTIs for model construction is provided in ESI data 1.† The prediction results of the best wSDTNBI model and the detailed information of the purchased compounds are provided in ESI data 2.† Some of the wSDTNBI models constructed in this study are freely available in our web server NetInfer (http://lmmd.ecust.edu.cn/netinfer). All other data are available upon reasonable request.

#### Author contributions

Yun Tang and Jin Huang supervised the research project. Zengrui Wu designed and implemented the wSDTNBI method. Zengrui Wu, Lulu Zheng, Zhuohang Yu, Weihua Li and Guixia Liu constructed network-based models, evaluated the performance of network-based models, performed network-based VS on RORγt and analyzed the prediction results. Hui Ma, Zehui Liu, Shuying Cao, Wenqing Fang, Lili Wu performed *in vitro* and *in vivo* experiments. Zengrui Wu, Hui Ma, Zehui Liu, Jin Huang and Yun Tang wrote the manuscript. All authors read and approved the final version of the manuscript.

#### Conflicts of interest

There are no conflicts to declare.

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