A central goal of systems pharmacology is to understand how biological systems respond to bioactive molecules. Most drugs achieve their therapeutic effect by interacting selectively with target proteins, which can be classified into major classes based on molecular function: enzymes, G-protein coupled receptors (GPCRs), ion channels (voltage-gated and ligand-gated), nuclear hormone receptors, catalytic receptors, and transporters. The binding of a ligand contributes to the conformation change of the target protein, which then affects the biochemical and physiological function linked to the target. The target also interacts with other molecules to affect cellular activity at the pathway level. Most drug targets can directly participate in cellular activity except GPCRs. The signal of ligand binding is passed from GPCRs to two types of transducer molecules: G-protein and \( \beta \)-arrestin, which then affect cellular activity by G-protein dependent and independent signaling, respectively.

Current efforts to systematically study pathways in response to drugs can be classified into two categories. Studies in the first category analyze gene expression datasets derived from \textit{in vitro} drug-induced experiments to identify differentially expressed genes, and then use enrichment analysis methods, such as gene set enrichment analysis, to find significantly enriched pathways. Such methods are facilitated by publicly available datasets like Connectivity Map, L1000, which contain collections of gene-expression profiles from cultured human cells treated with thousands of bioactive small molecules. However, a major drawback of this method is that due to the dominating noise inherent to microarray technology, many unrelated pathways will be drawn into

**WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC?**

- Understanding downstream functional consequences of pharmacologically targeted proteins is essential to drug design. Current approaches investigate molecular effects under tissue-naive assumptions. Many target proteins, however, have tissue-specific expression. A systematic study connecting drugs to target pathways in \textit{in vivo} human tissues is needed. We introduced a data-driven method that integrates drug-target relationships with gene expression, protein-protein interaction, and pathway annotation data. We applied our method to four independent genomewide expression datasets and built 467,396 connections between 1,034 drugs and 954 pathways in 259 human tissues or cell lines. We validated our results using data from L1000 and Pharmacogenomics Knowledgebase (PharmGKB), and observed high precision and recall. We predicted and tested anticoagulant effects of 22 compounds experimentally that were previously unknown, and used clinical data to validate these effects retrospectively. Our systematic study provides a better understanding of the cellular response to drugs and can be applied to many research topics in systems pharmacology.

**WHAT QUESTION DID THIS STUDY ADDRESS?**

- Many target proteins have tissue-specific expression limiting the usefulness of current methods. We present a systematic study connecting drugs to specific target pathways in human tissues.

**WHAT DOES THIS STUDY ADD TO OUR KNOWLEDGE?**

- We introduced a data-driven method that integrates drug-target relationships with gene expression, protein-protein interaction, and pathway annotation data. We applied our method to four independent genomewide expression datasets and built 467,396 connections between 1,034 drugs and 954 pathways in 259 human tissues or cell lines. The connections enabled us to predict and experimentally test anticoagulant effects of 22 compounds that were previously unknown.

**HOW MIGHT THIS CHANGE DRUG DISCOVERY, DEVELOPMENT, AND/OR THERAPEUTICS?**

- Our systematic study provides a better understanding of the cellular response to drugs and can be applied to many research topics in systems pharmacology, such as side effects prediction, drug repurpose, etc.
In most cases, hundreds of pathways are identified as significant, which will not provide much useful information for pharmacologists studying the drug. In addition, these drug-induced experiments are usually conducted on in vitro cancer cell lines, in which the expression of many genes is quite different from in vivo human tissues. Studies in the second category collect experimentally validated evidence from published articles and manually curate pathways to describe the pharmacological process of drugs, such as Pharmacogenomics Knowledgebase (PharmGKB), Kyoto Encyclopedia of Genes and Genomes (KEGG), etc. However, the pathways are usually limited to a small number of well-described pharmacokinetic or pharmacodynamic pathways and exclude many basic cellular processes involved in signaling and metabolism.

Cell-type plays an important part in the selectivity of drugs. Many drug target proteins are found to have tissue-specific expression. Even the same target protein can trigger distinct downstream effect in various cell types. For example, despite ubiquitous expression across the heart, calcium channels play a more important role in the action potential of cardiac pacemaker cells, which causes these cells to be more sensitive to calcium channel blocking agents than other cardiac cells. Understanding the mode of action of drugs from a tissue and cell type-specific perspective is crucial to drug discovery and development.

In this article, we developed a data-driven method to connect Drugs to target Pathways by the Tissue Expression (DATE). Our main hypothesis is that, for a drug to affect the function of a pathway in a tissue, the target protein must be associated with the pathway, and both the target and the pathway must be highly expressed in the target tissue. We validated our method against independent standards, tested the robustness of our results, and then applied DATE to drug safety prediction and drug-repositioning. We investigated a wide range of physiological drug effects, including anticoagulation activity, where we predicted 132 drugs with potential anticoagulation activity and selected 22 newly predicted compounds to validate experimentally. Our results provide evidence that targeting ADRB2, a GPCR that regulates the activation of major platelet integrin and levels of cytosolic calcium, may contribute to anticoagulation in whole blood. In addition, we integrated experimental results with clinical side effects of compounds to show that predicted compounds are more likely to have significant anticoagulation activity. Our results can be accessed at http://tatonettilib.org/resources.html.

**MATERIALS AND METHODS**

**Connecting drugs to target tissues**

Four tissue-specific gene expression datasets were analyzed in this study. We refer to them as U133A (microarray), NC160 (microarray), HPM_PRT (MS), and GTEX (RNA-seq). The data were preprocessed as detailed in **Supplementary Methods**. There were 5,016 connections between drugs and target proteins that were obtained from Drugbank version 3.0. A drug was connected to a target tissue if the target protein is both highly and specifically expressed in the tissue, as detailed in **Supplementary Methods**.

**Identifying the target pathways of drugs**

We used Reactome pathways with sizes between 5 and 500 as the data source for pathways. The class information of target proteins was integrated from three resources: GtoPdb, ENZYME, and Uniprot (**Supplementary Methods**). Target proteins that are GPCRs were connected to downstream pathways using our previously published method predicting GPCR downstream signaling pathways using the tissue expression (GOTE). Non-GPCR target proteins were connected to the annotated pathways that are highly expressed in the target tissue. We further filtered the parent pathway from results if it was connected to a drug in the same tissue together with its descendant pathway.

**Calculating the target, chemical, and indication similarity of drugs**

The similarity of target protein (or pathway) between two drugs was defined as the Jaccard similarity between two sets of target proteins (or pathway-tissue connections). The chemical similarity of two drugs was defined as the Tanimoto coefficient between the simplified molecular-input line-entry systems (SMILES) of two drugs. The indication similarity of two drugs was defined as the Jaccard similarity between two sets of indications from an ensemble resource linking medications to their indications named MEDI (**Supplementary Methods**).

**Predicting adverse events of drugs with target pathways of drugs**

A reference standard between 4 adverse events and a total of 149 drugs were used (**Supplementary Methods**). We scored the filtered pathways of each drug by the number of target proteins in the pathway and used scores as features to predict the adverse event of drugs. For each adverse event, we trained a binary random forest classifier with 500 trees. The out-of-bag probability of each drug was used to evaluate the performance of the classifier. To account for the stochastic nature of random forests, we ran the analysis for 100 times and used the average as the final results.

**Validation of drug-tissue-pathway connections using the L1000 dataset**

The expression dataset of 653,394 compound treatment experiments were downloaded from lincscloud.org. A reference standard containing positive controls (pathways with significant change in expression) and negative controls (pathways without significant change in expression) was created for each connection between a drug and a cell line (**Supplementary Methods**).

**Predicting drugs with potential effect on coagulation activity**

Twenty Reactome pathways that describe cellular activities in the coagulation process were manually selected from the 44 hemostasis pathways. Two additional PharmGKB pathways were added (**Supplementary Table S1**). We identified 132 drugs that are connected to the 22 pathways in three types of tissues: platelets, whole blood, and liver.
A series of "Maximum assay to measure coagulation and side effects from the OFFSIDES database."

Among those, 188 screened drugs were included. Of all 5,016 drug targets from the database, 337 were annotated with at least one side effect. We ranked them by the median of all the genes. To account for the variation in the absolute expression of different genes, the expression of each target was normalized to a score between 0 and 1. A proportion of tissues in which each drug target is highly expressed when compared to the median of all the genes. To account for the variation in the absolute expression of different genes, the expression of each gene is normalized by the baseline. Each box on the Y-axis represents one target class. The X-axis shows the tissue-specificity of proteins belonging to the target class. "All_drug_targets" represents the combination of all the target classes. "All_genes" represents all the genes in the human genome.

We extracted 695 drugs with the side effect of bleeding (many coagulation factors as well as red blood cells are synthesized by the liver) are defined as predicted drugs with effect on coagulation activity. As a reference set, 22 known anticoagulants were found through the Anatomical Therapeutic Chemical (ATC) code with the first four digits of "B01A."

Coagulation experiment and data process
We conducted an in vitro assay to measure coagulation activity of compounds, as described previously. A series of thrombin activity was measured 0–300 minutes after the treatment of each compound. The outliers of positive or negative controls on each plate were filtered out using the Median Absolute Deviation method (MAD). "Maximum ratio" of a compound was calculated by dividing the maximum activity to the average maximum activity of negative controls located on the same plate as the compound, then normalized to a score between 0 and 1.

Defining a threshold of anticoagulation activity
combining measured score and clinical side effect
We extracted 695 drugs with the side effect of bleeding (UMLS CUI: C0019080), 417 from side effect resource (SIDER) and 396 from the OFFSIDES database. Among those, 188 screened drugs were included. Of all 388 screened drugs, 337 were annotated with at least one side effect from two databases. We ranked them by maximum ratio from lowest to highest. Cutoff values from 5th to 95th percentile were used to divide all drugs into two groups. Odds ratios (ORs) of drugs with a bleeding side effect were calculated for each cutoff value (Supplementary Figure S1). As the percentile varies from 5th to 95th, the OR first rapidly increases to a maximum of 1.97 at the 17th percentile, then gradually declines close to 1. Thus, the 17th percentile cutoff, with corresponding maximum ratio of 0.775 was defined as the threshold of significant anticoagulation activity.

RESULTS
Drug target proteins have tissue-specific expression
We classified all 5,016 target proteins into eight major classes (Figure 1a, Supplementary Table S2). The two largest groups are enzymes (29.7%) and GPCRs (25.5%). The 13.4% target proteins which cannot be classified into eight groups. Odds ratios (ORs) of drugs with a bleeding side effect were calculated for each cutoff value (Supplementary Figure S1). As the percentile varies from 5th to 95th, the OR first rapidly increases to a maximum of 1.97 at the 17th percentile, then gradually declines close to 1. Thus, the 17th percentile cutoff, with corresponding maximum ratio of 0.775 was defined as the threshold of significant anticoagulation activity.

Figure 1a, Supplementary Table S2

Figure 1b–e

Supplementary Figure S2
specifically expressed in 31% of all tissues in U133A, 5.6% in NCI60, 20% in HPM_PRT and 32% in GTEx, significantly less than a median around 50% for all the genes ($P < 2.2e^{-16}$). Distinct classes of target proteins show different levels of tissue-specificity. The G-protein coupled receptor, ion channel, and nuclear hormone receptor are highly tissue-specific, whereas enzyme, catalytic receptor, and transporter are less tissue-specific.
Using tissue-specific expression of target proteins to predict target pathways of drugs

We used DATE (Figure 2) to predict target pathways of drugs using four datasets. We found 467,396 significant connections between 1,034 drugs and 954 pathways in 259 human tissues (Table 1). Each drug was connected to an average of 112 pathways and 27 tissues when combining the results of four datasets. The standard deviation is greater than the mean in all four datasets, suggesting that the number of connected pathways and tissues varies greatly by drug. We identified 115,905 unique connections between drugs and pathways. There were 41,098 (35.5%) connections that are recurrent in at least two datasets and 3,871 (3.34%) connections that are recurrent in all four datasets. We further filtered the results using hierarchy of Reactome pathways to minimize redundancy. The median size of predicted pathways was significantly reduced after filtering (P < 0.05; Supplementary Figure S3).

Many pathways were connected to more than one drug (Supplementary Table S3). The number of drugs that each pathway was connected to is positively correlated with pathway size (P < 2.2e-16; Supplementary Figure S4). A drug and a pathway can be connected in multiple tissues in which the target proteins are highly expressed. The number of target proteins is positively correlated with the number of target proteins in the pathway (P < 2.2e-16; Supplementary Figure S5).

Distinct datasets show consistency in the predicted target pathways of drugs

We compared the similarity of results in the three distinct datasets derived from normal human tissues: U133A, HPM_PRT, and GTEx. There were 7,510 connections between drugs and pathways that are recurrent in all three datasets (Figure 3a), and 1,032 connections between drugs, pathways, and tissues that are recurrent in all three datasets (Figure 3b).

For each drug, we calculated the pairwise Jaccard similarity of connected pathways among different datasets. The average pairwise Jaccard similarity is 0.46 ± 0.03 (U133A vs. HPM_PRT), 0.48 ± 0.02 (U133A vs. GTEx), and 0.55 ± 0.03 (HPM_PRT vs. GTEx), all significantly greater than null distribution 0.0099 ± 0.0015 (P < 2.2e-16; Figure 3c). The same analysis was performed to compare the pairwise Jaccard similarity of connected pathways in the same tissue among datasets. The average pairwise Jaccard similarity is 0.054 ± 0.007 (U133A vs. HPM_PRT), 0.045 ± 0.007 (U133A vs. GTEx), and 0.036 ± 0.006 (HPM_PRT vs. GTEx), all significantly greater than null distribution 0.00014 ± 0.00005 (P < 2.2e-16; Figure 3d).

Hormonal drugs show high tissue-specificity

We classified drugs according to their ATC class and defined the tissue-specificity of each class as the inverse number of tissues targeted by the class. Figure 3e and Supplementary Figure S6 show the tissue-specificity of each ATC class, as well as their target tissues. Hormonal drugs have the highest tissue-specificity in all four datasets. Ten of 84 (11.9%) tissues in U133A are targeted by hormonal drugs. Most of the 10 tissues belong to the endocrine system, such as the adrenal glands, hypothalamus, and pituitary glands. Other ATC classes have lower tissue-specificity, especially the alimentary tract, dermatological, sensory, and cardiovascular, which target >90% of the tissues in all four datasets.

Distinct classes of drugs show preference in different category of pathways

Reactome pathways can be classified into 25 general categories, representing various biological processes. For every ATC class, we showed the proportion of drug-pathway connections that belong to each pathway category (Figure 3f; Supplementary Figure S7), and identified the enriched categories (Supplementary Methods; Supplementary Table S4). All ATC classes are enriched with at least one pathway category except sensory and various. Blood, hormonal, immunological, and nervous are all enriched with five pathway categories, with hemostasis, gene expression, cell cycle, and neuronal system being the most significant category, respectively.
We also classified drugs by the class of target proteins (Figure 3g; Supplementary Figure S7; Supplementary Table S4). All target classes are enriched with at least two pathway categories except nuclear hormone receptor, which is uniquely enriched with gene expression. GPCR and enzyme are each enriched with seven and six pathway categories with no overlaps between them. Both ligand-gated ion channel and voltage-gated ion channel are enriched with the neuronal system. In addition, voltage-gated ion channel is also enriched with developmental biology. Other ion channel (second messenger-regulated) is most significantly enriched with membrane trafficking. Transporter is most significantly enriched with transmembrane transport of small molecules.

Drugs with similar chemical structure or indications target similar pathways

We tested whether drugs with similar chemical structure or medical indication will target similar pathways in the same tissue. We found that the similarity of target pathways increases exponentially with chemical similarity of drugs and remains consistently higher than the similarity of target proteins (Figure 4a–d). By contrast, the similarity of target pathways increases linearly with indication similarity (Figure 4e–h).

Using target pathways improves the performance of drug safety predictions

We hypothesized that the target pathways would better explain the adverse events of drugs than simply the target alone. To validate the hypothesis, we used target pathways or proteins as features and trained binary random forest classifiers to predict four adverse events of drugs. The performance of each classifier was shown by the receiver operating characteristic (ROC) curve in Figure 4i–l. Except for myocardial infarction, target pathways are better predictors than target proteins. For example, using target pathways in NCI60 to predict kidney failure can achieve an area under the curve (AUC) of 0.975, which improves the performance of target proteins by 15%. Similarly, using target pathways in GTEx to predict liver failure can achieve an AUC of 0.829, which improves the performance by 33.7%.
The number of features used in the classifier varies by the adverse events and by the datasets used for prediction (Supplementary Table S5). After filtering the features by importance score, we found that the number of features with importance >0 varies from 57 to 162. No correlation between the number of features and the AUC of classifier was found. “Synthesis of 15-eicosatetraenoic acid derivatives,” “synthesis of prostaglandins and thromboxanes,” and “arachidonic acid metabolism” are important pathway predictors for gastrointestinal bleeding, kidney failure and myocardial infarction. In addition, “metabolism of angiotensinogen to angiotensin” is important in predicting kidney failure. “Trafficking of GluR2-containing AMPA receptors” and “peptide hormone metabolism” are important in predicting liver failure.

**Predicted pathways go through expression change after drug treatment in L1000 experiments**

We hypothesized that the expression of target pathways will change after the treatment of drug in the predicted tissue. We validated this using a reference standard among the drug, pathway, and cell line built from the L1000 dataset. Of all 76,191 connections, 281 connections (4%) between 92 drugs and 16 cell lines can be evaluated using the standard. Precision and recall were calculated using the reference standard (Figure 5a,b). The precision of our results is 0.38 ± 0.04, significantly outperforming null distribution 0.28 ± 0.03 (P = 6.51e-05). The recall of our results is 0.0087 ± 0.0021, which significantly outperforms the null distribution 0.0027 ± 0.0004 (P = 6e-08).
DATE recaptures the drug-pathway connections in PharmGKB

PharmGKB\textsuperscript{14} provides a reference standard of 187 connections between 143 drugs and 92 pathways that describe the pharmacodynamics and pharmacokinetics of the drug. We used DATE to connect drugs to the 92 pathways and compared our results to PharmGKB. Altogether we built 29,078 connections between 776 drugs and 88 pathways in 223 human tissues (Supplementary Table S6). Of 143 drugs from PharmGKB, 129 drugs (90\%) can be evaluated. Precision and recall were calculated using the reference standard (Figure 5c-d). The precision of combined (combining all four datasets) and recurrent (appearing in at least two datasets) results are 0.25 ± 0.06 and 0.35 ± 0.08, respectively, which significantly outperforms the null distribution of 0.01 ± 0.01 (P < 1.0 e-11). The recall of combined and recurrent results are 0.44 ± 0.08 and 0.38 ± 0.08, respectively, which significantly outperforms the null distribution of 0.04 ± 0.03 (P < 1.0 e-15).

DATE accurately predicts coagulation effects in experimental assays

We predicted 132 drugs with coagulation effects (Supplementary Table S7) based on their connections to a manually curated set of 22 coagulation-related pathways (Supplementary Table S1). Predicted drugs are enriched with known anticoagulants (P = 2.09 e-07; Supplementary Table S8).

We conducted one unbiased experimental screen of 388 drugs (Supplementary Table S9) to measure the coagulation activity using a high-throughput \textit{in vitro} assay developed previously.\textsuperscript{36} A “maximum ratio” score was calculated to represent coagulation activity (Supplementary Table S10). Screened drugs were not enriched with known anticoagulants (P = 0.16; Supplementary Table S8). We grouped all 388 screened drugs into the following categories: (1) Reactome: 26 predicted drugs connected to Reactome pathways; (2) PharmGKB: 41 predicted drugs connected to PharmGKB pathways; (3) tissue-naïve: 50 drugs that were connected to 22 coagulation-related pathways only in tissues other than coagulation tissues (these drugs can be predicted by tissue-naïve methods, but not DATE); and (4) others: 325 drugs that were not predicted to have anticoagulation effects. The average maximum ratio of predicted drugs using PharmGKB (0.76 ± 0.08) are lower than that of both tissue-naïve drugs (0.85 ± 0.02) and unpredicted drugs (0.85 ± 0.01), with \( P \) value of 0.014 and 0.013 (Figure 6a).

Meanwhile, we combined the measured coagulation activity of drugs with clinical side effects concerning drug-induced bleeding. A maximum ratio of 0.775 was defined as the threshold of significant anticoagulation activity, which maximizes the OR of drugs with bleeding side effects (OR = 1.97; \( P = 0.018 \); Supplementary Figure S1). We found 34\% of Reactome and 19\% of PharmGKB predicted drugs below this threshold (Figure 6a), both of which are greater than unpredicted drugs (15\%). Among the five Reactome or PharmGKB predicted drugs with the lowest maximum ratio, four of them have the side effect of bleeding (Figure 6c).

Targeting ADRB2 may cause anticoagulation effect in whole blood

Sixty-nine predicted drugs were not included in the initial screen. These unpredicted drugs were enriched with seven known anticoagulants (\( P = 6.44 e-06 \); Supplementary Table S8). We selected 22 compounds from the other 62 drugs that have not been validated on the coagulation activity (Supplementary Methods; Supplementary Table S11) and performed a prospective validation of these predictions. The average maximum ratio of 22 compounds is 0.81 ± 0.14 (Figure 6b), which is significantly lower than that of negative controls 1.0 ± 0.09 (\( P = 0.018 \)). Ten compounds (45\%) have significant anticoagulation activity according to the defined threshold. The five drugs with the lowest maximum ratio are labetalol, norepinephrine, procto-terol, alpenolol and sulcrafate (Figure 6c), which are primarily used to treat cardiovascular diseases, such as hypertension and chest pain. Labetalol, norepinephrine, and sulcrafate were reported to have the side effect of...
bleeding. Labetalol, norepinephrine, procaterol, and alprenolol mutually target ADRB2, a beta-2-adrenergic receptor. Procaterol is a selective agent of ADRB2, whereas the other four drugs are nonselective agents with other adrenergic receptor targets (e.g., ADRA1A, ADRA1B, and ADRB1).

Using DATE, we connected ADRB2 to three hemostasis pathways in whole blood: “integrin alphaIIb beta3 signaling,” “reduction of cytosolic Ca\(^{2+}\) levels,” and “factors involved in megakaryocyte development and platelet production.”

DISCUSSION

Target proteins of a drug function as a bridge that connects the drug to cellular activity. We found that many target proteins have tissue-specific expression, indicating the importance of taking tissue-specificity into account while identifying drug target pathways. This consideration is absent in current systematic methods. We presented a data-driven method, DATE, to systematically connect drugs to target pathways in human tissues. We applied DATE to four expression datasets and validated our results by comparing to two standards: (1) a weak standard created with the L1000 expression dataset; and (2) a stringent tissue-naive standard from PharmGKB. The Y-axis shows the proportion of compounds with significant anticoagulation activity in a and b. (a) Boxplot with jitter showing the coagulation activity of 6 groups: (1) positive control: argatroban; (2) Reactome: 26 predicted drugs connected to Reactome pathways; (3) PharmGKB: 41 predicted drugs connected to PharmGKB pathways; (4) tissue-naive: 50 drugs that can be predicted by tissue-naive methods, but not DATE; (5) others: the other unpredicted 325 drugs; and (6) Negative control: DMSO. The Y-axis shows the proportion of compounds with significant anticoagulation activity in a and b. (b) Boxplot with jitter showing the coagulation activity of three groups: (1) positive control: argatroban; (2) 22 newly predicted compounds that have not been screened on the coagulation activity in a and b; and (3) negative control: DMSO. (c) A table of predicted drugs with significant anticoagulation activity in a and b. Only the top five drugs in each group (PharmGKB, Reactome, and 22 compounds) were shown here. Full results can be found in Supplementary Table S10. P, P value; RR, reporting ratio; NA, compound was not studied in SIDER or OFFSIDES. Because only OFFSIDES provides a P value for each pair of drug and side effect, whereas SIDER only provides the mapping between them, some compounds will have blank profiles in the last two columns if they were only reported in SIDER.

The drug-pathway-tissue connections we built can be applied to address many important research topics in molecular systems pharmacology. For example, we used the connected pathways as features to predict four common adverse events, and saw an improvement of performance when compared to using the target proteins as features in three of the four adverse events tested. In
myocardial infarction, the molecular mechanism can be well explained by target proteins with high importance score, which lead to the better performance of target proteins. For instance, PTGS2 that shows the highest importance score in predicting myocardial infarction, is a cardioprotective protein that alleviates ischemia/reperfusion injury and mediates late phase preconditioning.\textsuperscript{30–42} We also predicted 132 drugs to have an effect on the coagulation process. An unbiased experimental screen showed that the predicted drugs are more likely to have anticoagulation activity compared with those drugs only predicted by tissue-naive methods or other unpredicted drugs. This is because our tissue-specific analysis can connect drugs to their target tissues or potential sites of action and provide more comprehensive knowledge about the mechanism of action. Three predicted drugs with lowest coagulation activity are known anticoagulants and several others can act as a vitamin-K antagonist, which reduces blood-clotting. We performed prospective validation on 22 predicted compounds that have not been used as anticoagulants, and found that their coagulation activity is significantly lower than the negative controls. Among the 22 compounds, several have been reported to affect coagulation in previous research, such as labetalol\textsuperscript{43,44} and alpenrolen.\textsuperscript{45,46} Four newly predicted compounds with lowest coagulation activity share a mutual target protein, ADRB2. This suggests that targeting ADRB2 may cause an anticoagulation effect in whole blood, opening potential new avenues for drug discovery research. In addition, we integrated experimental screen results with clinical data and defined the 17th percentile of the maximum ratio as the threshold of significant anticoagulation activity. Drugs with a maximum ratio below the threshold are nearly two times more likely to cause bleeding. We found that predicted drugs exhibit a higher probability of significant anticoagulation activity when compared to unpredicted drugs, which further validates our predictions. Shown in Figure 6c, many predicted drugs have already been reported with the bleeding side effect in OFFSIDES or SIDER, such as misoprostol and norepinephrine, etc. Meanwhile, several other drugs, such as sulfasalazine, mandelone, procaterol, or alpenrolen have not been reported with the bleeding side effect yet. Our analysis provided evidence that such drugs may have significant anticoagulation activity, thus, the patient should be monitored for potential bleeding risk.

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