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Pathway-based drug repositioning using causal inference

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Abstract

Background: Recent *in vivo* studies showed new hopes of drug repositioning through causality inference from drugs to disease. Inspired by their success, here we present an *in silico* method for building a causal network (CauseNet) between drugs and diseases, in an attempt to systematically identify new therapeutic uses of existing drugs.

Methods: Unlike the traditional 'one drug-one target-one disease' causal model, we simultaneously consider all possible causal chains connecting drugs to diseases via target- and gene-involved pathways based on rich information in several expert-curated knowledge-bases. With statistical learning, our method estimates transition likelihood of each causal chain in the network based on known drug-disease treatment associations (e.g. bexarotene treats skin cancer).

Results: To demonstrate its validity, our method showed high performance (AUC = 0.859) in cross validation. Moreover, our top scored prediction results are highly enriched in literature and clinical trials. As a showcase of its utility, we show several drugs for potential re-use in Crohn's Disease.

Conclusions: We successfully developed a computational method for discovering new uses of existing drugs based on casual inference in a layered drug-target-pathway-gene- disease network. The results showed that our proposed method enables hypothesis generation from public accessible biological data for drug repositioning.

Background

Despite the fast growth in drug research and development (R&D) such as chemical genomics technologies [1,2] and chemical libraries [3,4], the pharmaceutical R&D outputnew drugs brought to market-has significantly declined in recent decades. As reported in the most recent analysis, the number of new drugs approved per billion US dollars spent has halved approximately every 9 years since 1950 [5]. Discovering new uses for existing drugs, also known as drug repositioning, provides one possible solution to such a problem. The fact that existing drugs have already passed through development stages such as target validation and ADMET (absorption, distribution, metabolism, excretion and toxicity) characteristics analysis should



The traditional one drug-one target-one disease drug discovery model has been argued to more likely result in poor efficacy or unanticipated side effects by not taking into account the complexity of underlying mechanism [7,8]. Due to such limitations, network-based computational approaches were proposed recently, providing a new framework for identifying drug-repositioning opportunities. Keiser et al. predicted new targets for known drugs using drug chemical structures and their canonical biological targets, and the resulting novel drug-target network further connected drugs to new indications [9]. Li et al. measured drug pairwise similarity by combining similarity of drug chemical structures, similarity of target profiles, and interaction between target proteins [10]. Iorio et al. constructed a drug-drug similarity network using transcriptional responses (*i.e.*, gene expression profiles)



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following drug treatment [11]. Recent studies [12-14] compared the drug vs. disease gene expression profiles for identifying novel treatment relationships between drugs and diseases. Other kinds of network-based approaches for drug repositioning included literature mining [15] and shared pathway analysis [16].

Different from the aforementioned computational approaches, several recent studies demonstrated the feasibility of drug repositioning through manual analysis of causal associations in drug-involved pathways [17-20]. For example, Cramer et al. found that FDA approved anticancer drug bexarotene could be potentially used for Alzheimer's Disease (AD) treatment [19] based on molecular pathway examination and analysis. More specifically, they found bexarotene activates nuclear receptors PPAR (peroxisome proliferator-activated receptor) and LXR (liver × receptor) in coordination with RXR (retinoid \times receptor), thus up-regulating the expression of the ApoE (apoliporrotein E) gene. This process facilitates the clearance of A β (β -amyloid) from the brain, resulting in the alleviation of AD. In this example, the chain of causality between one drug and one disease was examined and inferred by domain experts who took advantage of the following knowledge in bexarotenerelated pathways: (1) drug-target (e.g., bexarotene is an RXR agonist); (2) target involved pathway (e.g., LXR:RXR activation pathway); (3) transcriptional responses in a given pathway (e.g., increased ApoE gene expression in the LXR:RXR activation pathway); (4) genetic mechanism of disease (e.g., ApoE is associated with AD).

Motivated by the success of manual pathway analysis for drug repositioning, we developed a new computational method for building a network of causal chains between drugs and diseases, allowing for computational drug repositioning. By taking advantage of the increasing amount of expert-curated biological knowledge in the public domain (e.g. pathway information in Pathway Commons [21]), we built a multi-layer causal network (CauseNet) consisting of chains from drug to target, target to pathway, pathway to downstream gene, and gene to disease. Furthermore, we used a statistical method to learn the transition likelihood of each causal chain in the network based on those known drug-disease treatment relationships. In the prediction stage, we identified novel drug re-uses using maximum likelihood estimation. Unlike the traditional causal chain models that relied on human examination of one drug target, pathway and gene at a time, our computational model allows us to investigate all possible causal links when connecting drugs to diseases at once. To our best knowledge, this is also the first attempt of using network-based causal inference in computational drug repositioning.

Methods

In Figure 1, we show a model of our proposed CauseNet which puts causal chains from drugs to diseases in a layered network. The nodes of CauseNet are organized in five layers: drug $D \{d_1, ..., d_n\}$, target $T \{t_1, ..., t_m\}$, pathway $P \{p_1, ..., p_n\}$, downstream genes $G \{g_1, ..., g_k\}$, and disease S $\{s_1, \dots, s_n\}$. Accordingly, from top to bottom the causal links between two layers represent (1) drug d acts on target t; (2) target t participants in pathway p; (3) pathway p affects the expression of downstream gene g; and (4) gene g is associated with disease s. To construct such a network, we integrated data from heterogeneous resources which contain expert-curated knowledge of relationships between drugs, molecules and diseases. Furthermore, we learn the transition weight for each causal link in the CauseNet to distinguish the likelihood of transitions between nodes based on the known treatment relationships between drugs and diseases (details in Section computing transition weights). For instance, if drug d_1 is known to treat disease s_{ν} , then the transition weights of the gold-colored links in Figure 1 should be promoted accordingly.

Constructing CauseNet

For constructing CauseNet, we extracted approved drugs and their targets from DrugBank [22], target-involvedpathways from Pathway Commons [21] and KEGG [23], downstream genes from Pathway Commons, and diseases and their associated genes from Comparative Toxicogenomics Database (CTD) [24]. Also from CTD, we assembled pairs of known drug-disease treatment relationships. Note that each pathway can mention information on a series of biological events such as biochemical reactions, physical interactions, transcriptional responses, and phosphorylation and enzyme catalysis. In this study, we focused on transcriptional responses (*i.e.*, up/down regulated expression of downstream genes) in a pathway.

Computing transition weights

We represent the constructed CauseNet as a directed graph G(V, E). The node set, $V(G)=\{D, T, P, G, S\}$, consists of five types of objects (*i.e.*, drug *D*, target *T*, pathway *P*, downstream gene *G* and disease *S*). The edge set is denoted as $E(G) \subseteq \{D \times T, T \times P, P \times G, G \times S\}$. A complete causal chain, $c = \langle d, t, p, g, s \rangle$, represents a 4-step path from drug d ($d \perp D$) to disease s ($s \perp S$)with a set of individual chains $E(c) = \{(d, t), (t, p), (p, g), (g, s)\} \subset E(G)$. All possible causal chains from drugs to diseases become the complete chain set *C*. We further use a subset of (treatment-enriched) chains C^* (*i.e.*, $C^* \subset C$) to represent the links between drug-disease pairs of known treatment relationships. For example, as shown in Figure 1, drug d_1 is linked to diseases s_2 and s_y through two separatechains



 $c_1 = \langle d_1, t_2, p_2, g_3, s_2 \rangle$ and $c_2 = \langle d_1, t_2, p_2, g_2, s_y \rangle$, where c_1 , $c_2 \not\subset C$ and $c_2 \not\subset C^*(d_1 \text{ is known to treat } s_y \text{ but not } s_2)$.

The graphs of the respective complete and enriched chain sets *C* and *C*^{*} are denoted as G(C) = G(V(C), E(C))and $G(C^*) = G(V(C^*), E(C^*))$, where $V(C^*) \subset V(C)$ and $E(C^*) \subset E(C)$. Given above, we can learn the transition weight $w(v_b, v_j)$ to represent the transition likelihood from node to towards treatment relationships $(\exists (v_b, v_b) \not \in E(C))$:

$$w(v_i, v_j) = \begin{cases} 1 + \frac{p(v_i \rightarrow v_j | G(C^*))}{p(v_i \rightarrow v_j | G(C))} & \text{if } (v_i, v_j) \in E(C^*) \\ 1 & \text{otherwise} \end{cases}$$
(1)

Where $p(v_i \rightarrow v_j | G(C^*))$ and $p(v_i \rightarrow v_j | G(C))$ are the transition probabilities from node v_i to node v_j in $G(C^*)$ and G(C), respectively. Let each chain graph $G(\bullet)$ be a Markov model. Thus the transition probability $p(v_i \rightarrow v_j | G(\bullet))$ is computed using maximum likelihood estimation:

$$p(v_i \to v_j \mid G(\bullet)) = \frac{N v_i, v_j}{N v_i, \bullet}$$
(2)

 $N\nu_i$, ν_j is the number of times that a transition $\nu_i \rightarrow \nu_j$ is observed in a chain set, and $N\nu_i$, is the total number of transitions originated from ν_i in the chain set.

Predicting novel treatment relationships between drugs and diseases

For each causal chain $c = \langle d, t, p, g, s \rangle$ in the global chain set (d C), we can estimate its likelihood L(c) based on the pre-computed transition weights in equation (1).

$$L(c) = \log(w(d, t) \cdot w(t, p) \cdot w(p, g) \cdot w(g, s))$$
(3)

Our prediction of a new indication of drug d_x for disease s_y is based on the final score $S(d_x, s_y)$ between drug d_x and disease s_y , which is the maximal likelihood of all possible chains from d_x to s_y :

$$S(d_x, s_y) = \max(L(c_{x,y})), \quad c_{x,y} \in \{ < d_x, t_{\bullet}, p_{\bullet}, g_{\bullet}, s_y > \} \quad (4)$$

 $c_{x,y}$ is a causal chain from drug d_x to disease s_y among all possible chains $C_{x,y} = \{ \langle d_x, t, p_y, g_y, s_y \rangle \}$. Note that alternatively, $S(d_xs_y)$ can also be measured simply by the number of successful chains from d_x to s_y : $|C_{x,y}|$. As shown below, we used such a method as a baseline for comparing our weighted method.

Results

Complete and treatment-enriched chain sets

Based on the CauseNet (see Section constructing CauseNet), we constructed a complete causal chain set C including 2,711,440 possible 4-step chains from 979 drugs, to 538 targets, to 207 pathways, to 1,122 downstream genes, to 1,650 diseases, corresponding to 389,945 possible drug-disease associations. A total of 6,268 such associations between 665 drugs and 583 diseases were labelled as known (i.e. found in CTD), resulting in a total of 135,936 chains to the treatment-enriched chain subset C^* .

Table 1 shows detailed statistics of the complete vs. enriched chain sets and their corresponding graph elements. For each edge in G(C), we calculated its transition weight based on equation 1 (see Section computing transition weights). Furthermore, we computed scores for each of the 389,945 possible drug-disease associations based on the maximal likelihood estimation of causal chains (equation 4) and ranked them accordingly. When treating the known 6,268 associations as the only positive instances, we calculated true positive rate (sensitivity) and false positive rate (1-specificity) of our results at different cut-off ranking scores. As plotted as a ROC curve in Figure 2(A), we obtained a high AUC score of 0.889, which suggests that the 6,289 known (positive) associations were indeed ranked high among all 389,945 pairs. Also in Figure 2(A), we show that our weighted inference method significantly outperformed the baseline method in AUC scores, which shows the value of computing weights for transition between nodes in our CauseNet.

Cross validation of therapeutic effect prediction

To further evaluate the validity of our method, we conducted a 10-fold cross validation by withholding 10% of the known treatment relationships in each fold and removing their connected chains accordingly. Figure 2 (B) shows the results of all ten ROC curves, with the average AUC score of 0.859 ± 0.006 with (CI = 0.95) (highlighted in blue). The best tradeoff between sensitivity (0.866) and specificity (0.760) is shown in red, which corresponds to 2.609 in our prediction score. After filtering known ones, 92,057 associations between 964 drugs and 1050 diseases have scores higher than 2.609. Additional File 1 lists the 92,057 predicted associations and all possible causal chains connecting the drug-disease associations via target-and gene-involved pathways.

We compared our method with the similarity-based methods [9,10] which assume that similar drugs are used for similar diseases' treatments. Drug pairwise similarity was measured by chemical 2D structure similarity (SIM_chem), drug target similarity (SIM_target), and linear combination of these two (SIM_combo) respectively. We applied the similarity-based methods to 602 small molecule drugs (with 2D chemical structure) in our CauseNet dataset. As can be seen in Figure 3, our method achieved a higher AUC score (0.866) than using chemical similarity (0.829), target similarity (0.841) or their combination (0.851).

Novel predictions in clinical trials and literature

We further evaluated our predictions by searching evidence in clinical trials and literature. About 1/3 were found in PubMed [25] (requiring three or more occurrences) and a relatively small percentage of our predictions (3,202) were found in ClinicalTrials.gov [26]. There are several main reasons for more evidence in the literature than in clinical trials: First, some predicted therapeutic uses are still in pre-clinical development and hence have not reached clinical trial stage. For example, we predicted anakinra to treat colorectal neoplasm with a high confidence score of 5.996.

Гable	1	Descriptive	statistics	of	global	and	treatment	enriched	chain set	ts
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		Complete chain set C	Enriched chain subset C*	
# of chains		2,711,440	135,936	
Node set	# of drugs D	979	655	
	# of targets T	538	397	
	# of pathways P	207	199	
	# of genes G	1,122	838	
	# of diseases S	1,650	583	
Edge set	D×T	2,953	2,074	
	T×P	2,922	2,004	
	P×G	2,772	2,179	
	G×S	6,496	3,954	
# of drug-disease	associations	389,945	6,268	

According to literature evidence [27], anakinra–a drug approved for treating rheumatoid arthritis–was recently found to be able to contribute to growth-inhibition of small tumors in mice with colon carcinoma. Second, clinical trials are not always registered in ClinicalTrails. gov. In our results, some highly scored predictions were found for novel uses of nadroparin–a drug outside of the U.S. market. Some trials have been launched for investigating these new uses in countries outside the U.S., with their studies reported in literature, but not in ClinicalTrials.gov.

To demonstrate the discriminative power of our prediction scores, we show in Figure 4 that in general the higher the prediction score and more likely the predicted association can be validated in ongoing clinical trial investigations and scientific publications. Hence, we believe such a score can greatly help others to use our prediction results for further investigations.





Investigations of drug repositioning opportunities for Crohn's Disease

Drug repositioning for poorly treated diseases is a promising strategy in drug discovery today because of the highly unmet need there [5]. In this study, we further explored drug repositioning opportunities for Crohn's disease (CD), a chronic inflammatory condition of the gastrointestinal tract, for which there is no known cure and most treatment options aim to relieve its symptoms such as rectal bleeding and diarrheal [28]. Every year, $10,000 \sim 47,000$ residents of North America are diagnosed with CD, and as many as 630,000 currently suffer from CD [29]. Epidemiology studies showed incidence of CD is highly influenced by geographic region and family history. Recently, genetic efforts have been made to explain these epidemiologic observations and to understand the underlying pathogenesis from the view of human genomics [30,31]. As a result, multiple CD susceptibility genes have been found such as IL23R, IL6, IL10, NLRP3, FN1, NCF4 and FPR2. These findings could lead to identifying novel therapeutic options for CD.

Figure 5 shows five selected CD drugs predicted by our method for CD and their exemplar causal chains found in our CauseNet. For example, anakinra, an approved rheumatoid arthritis drug, shows a high potential for CD treatment with a score of 5.26 in our method. Further analysis shows that anakinra works by binding receptor IL1R, which may influence multiple pathways like osteo-clast differentiation pathway and amoebiasis pathway, affecting CD genes NCF4 and FN1 respectively. Another highly scored drug is nedocromil (score = 4.00), a drug

approved for treating allergic conjunctivitis and asthma. Our method shows its potential therapeutic use in CD through acting on multiple targets HSP90AA1 and FPR1, affecting multiple pathways NOD-like receptor signaling pathway and staphylococcus aureus infection pathway, and further affecting multiple CD mechanism genes IL6, TNF, NLRP3, NOD2, FPR2 and IL10. This comprehensive evidence would greatly help experts generate hypotheses on the therapeutic values of these CD drug candidates which are worth further experimenting. We find that two drugs shown in Figure 5, adalimumab and prednisolone, have also been previously studied for CD [32,33].

Discussion

In this study, we propose a new computational drug repositioning approach by using causal chains in drug-disease networks (see Figure 1). Our method has the following important characteristics:

First, it provides a broad and semantic view of molecular causality between drugs and diseases. Unlike the traditional 'one drug-one target-one disease' model, we put all causality relationships between drugs and disease in a network view with five distinct layers. In the CauseNet construction, we integrated different types of data and semantic relationships between them from widely recognized and expert-curated resources. For example, when integrating pathway data, we focused on specific direction (downstream) and specific semantics (transcriptional response) relationships in an interested pathway by taking advantage of recent progress in pathway curation and standardization [21,34]. The resulting CauseNet laid down



a key foundation for further drug-disease relationship prediction.

Second, not only does our method find novel drugdisease treatment associations, but also scores and ranks each prediction accordingly. As shown in the crossvalidation experiment, our method is able to rank true associations generally at the top positions. Moreover, those highly scored drug-disease prediction results are found significantly enriched in clinical trials and biomedical literature. Hence, we believe that our weighted inference method is able to prioritize prediction results for further exploring drug repositioning opportunities.

Third, instead of being a black box, our method provides detailed and comprehensive molecular evidence supporting each prediction. As shown in the case study with Crohn's disease, the accompanying pathway evidence can

David	Selected Causal Chains Supporting This Prediction
Drug	(D Drug; T Target; P Pathway; G Downstream Gene; S Disease)
Anakinra	D Anakinra → T IL1R → P Osteoclast differentiation → G NCF4 → S CD
(score=5.29)	D Anakinra→ T IL1R→ P Amoebiasis→G FN1→ S CD
Rifabutin	 D Rifabutin → T HSP90AA1 → P NOD-like receptor signaling pathway → G IL6, TNF, NLRP3, NOD2 → S CD
(score=4.30)	D Rifabutin \rightarrow T HSP90AA1 \rightarrow P Glucocorticoid receptor regulatory network \rightarrow G IL6 \rightarrow S CD
Nedocromil	 D Nedocromil → T HSP90AA1 → P NOD-like receptor signaling pathway → G IL6, TNF, NLRP3, NOD2 → S CD
(score=4.00)	 D Nedocromil → T FPR1 → P Staphylococcus aureus infection → G FPR2, IL10 → S CD
Adalimumab	D Adalimumab→ T TNF, FCGR1A→ P Tuberculosis→ G TNF, IL6, IL10, IL23R → S CD
(score=3.64)	D Adalimumab → T FCGR1A, C1R, C1QA, C1S → P Staphylococcus aureus infection → G FPR2, IL10 → S CD
Prednisolone	D Prednisolone \rightarrow T NR3C1 \rightarrow P Glucocorticoid receptor regulatory network \rightarrow G IL6 \rightarrow S CD
(score=2.96)	D Prednisolone \Rightarrow T NR3C1 \Rightarrow P AP-1 transcription factor network \Rightarrow G IL6, IL10 \Rightarrow S CD
ure 5 Potential drugs for Crohi	s's Disease (CD) treatment.

support further human investigation. More importantly, such comprehensive pathway information could reveal unknown linkages between drugs and disease and help hypothesis generation on novel drug re-uses.

Lastly, our prediction results cover a wide range of diseases and drugs. For drugs, our repositioning results consist of both small molecule drugs (*e.g.*, rifabutin) and big molecules (*e.g.*, adalimumab), thus lifting the limitations of those methods that rely on 2D chemical structures or gene expression profiles of small molecules [9-14]. In addition, our method can identify drugs for a disease with no current treatments, making it different from similarity-based methods where predictions are always based on known uses of other drugs.

Like other knowledge-based methods, our approach relies on existing knowledge of drug-target, target-pathway, pathway-downstream gene, gene-disease, and drugdisease relationships. Despite increasing efforts in data curation and standardization, at present such information is still incomplete, thus limiting the prediction power of our method. For example, we extracted 1,239 targetinvolved pathways, but merely 209 of which contain transcriptional response relationships. Combining gene expression with pathway analysis to predict downstream genes is a hopeful strategy to help break the bottleneck [35]. We plan to investigate this issue in future work.

Conclusions

In this study, we successfully developed a computational drug repositioning method using pathway-based causal inference. Unlike the traditional 'one drug-one target-one disease' causal model, we systematically considered all possible causal chains connecting drugs to diseases via target- and gene-involved pathways. More specifically, we built a multi-layer causal network (CauseNet) consisting of chains from drugs to disease by integrating heterogeneous expert-curated biological resources in public domain. The transition likelihood of each causal edge in the CauseNet was estimated by learning known drug-disease treatment relationships. Furthermore, we predicated novel drug indications using maximum likelihood estimation of causal chains between drugs and diseases. In cross-validation experiments, our method achieved AUC score of 0.859 ± 0.006 with best tradeoff sensitivity = 0.866 and specificity = 0.760. When compared with a control group of drug uses, our drug repositioning results were found to be significantly enriched in both the biomedical literature and clinical trials. Additionally, in the Crohn's Disease case study, we demonstrated our method would provide more comprehensive evidence showing how drugs connect to diseases via pathways. We believe our method would greatly help experts generate hypotheses in drug discovery.

Additional material

Additional file 1: Predicted drug-disease associations. lists the 92,057 predicted associations and all possible causal chains connecting the drug-disease associations via target-and gene-involved pathways

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

JL and ZL conceived the whole study, participated in its design, analyzed the results and wrote the manuscript. JL collected the data, implemented the methods and performed the experiments. All authors read and approved the final manuscript.

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Declarations

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