Databases and ontologies

**CaNDis: a web server for investigation of causal relationships between diseases, drugs and drug targets**

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Abstract

**Motivation:** Causal biological interaction networks represent cellular regulatory pathways. Their fusion with other biological data enables insights into disease mechanisms and novel opportunities for drug discovery.

**Results:** We developed Causal Network of Diseases (CaNDis), a web server for the exploration of a human causal interaction network, which we expanded with data on diseases and FDA-approved drugs, on the basis of which we constructed a disease–disease network in which the links represent the similarity between diseases. We show how CaNDis can be used to identify candidate genes with known and novel roles in disease co-occurrence and drug–drug interactions.

**Availability and implementation:** CaNDis is freely available to academic users at http://candis.ijs.si and http://candis.insilab.org.

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**Supplementary information:** Supplementary data are available at Bioinformatics online.

1 Introduction

Protein–protein and protein–drug interaction networks are becoming increasingly important for drug development (Ashburn et al., 2004). Such biological interactions, whether predicted or experimental, are stored in various databases (Konec et al., 2017; Szklarczyk et al., 2019). However, the causal relationships between the interaction partners remain relatively unexplored. A causal relationship occurs when e.g. a transcription factor increases the expression of a particular protein. Knowledge of these relationships could open up new avenues for drug development and improve our understanding of the disease processes.

Tools to explore networks of causal relationships are available (Boué et al., 2015; Licata et al., 2020); however, the fusion of these networks with the knowledge about diseases and drugs into heterogeneous causal networks, should enable even higher understanding of diseases as well as more efficient drug discovery. In particular, such networks could be used for the study of disease co-occurrence, the simultaneous presence of two or more diseases in a patient (Hu et al., 2016).

We developed Causal Network of Diseases (CaNDis), a web server that enables visual exploration of a causal network of protein–protein interactions enriched with annotations on diseases and drugs. We also constructed a new disease–disease network. Here, each node is a subnetwork in the original causal network, in which all the nodes are linked to the same disease. A score is assigned to a link between a pair of diseases based on the connectivity of the two subnetworks, representing disease similarity. CaNDis is an intuitive web server based on a fast WebGL viewer (see sections ‘Software aspects and scaling’ and ‘Comparison with other tools’ in Supplementary Information). It facilitates drug discovery with respect to the identification of disease co-occurrence and drug–drug interactions at the level of drugs, genes, proteins and diseases.

2 The CaNDis web server

The CaNDis heterogeneous causal network is based on the CBN (Boué et al., 2015) and SIGNOR 2.0 (Licata et al., 2020) causal biological networks. It is extended with protein–RNA interactions from the PRD database (Fujimori et al., 2012) and drug–gene interactions from the DGe database (Cotto et al., 2018); gene nodes are superimposed with the gene–disease annotations from the DisGeNET database (Piñero et al., 2020). Causal pathways are merged into a single network containing the interacting proteins as nodes and the causal relationships, such as increase or decrease in expression levels, as links (see section ‘The CaNDis network’ in Supplementary Information).

Gene–disease annotations are added as attributes to the protein nodes and drugs and RNA are added as new nodes, which are linked...
to their respective target protein nodes. The obtained heterogeneous causal network contains annotations on more than 3000 complex diseases as well as on the FDA approved drugs to enable the investigation of diseases (i.e. co-occurrence of diseases) and drug–drug interactions (see section ‘Use in drug–drug interactions’ in Supplementary Information).

A new disease–disease network is also developed from the heterogeneous causal network (sections ‘Disease–disease network construction’ and ‘Normalized disease–disease similarity score’ in Supplementary Information), and is available on the web server.

3 Use for identification of genes with roles in disease co-occurrence

According to epidemiological studies, patients with central nervous system (CNS) disorders, that is, Alzheimer’s disease, Parkinson’s disease and schizophrenia, are less likely to develop certain cancers (colorectal, lung and prostate), and vice versa. This inverse co-occurrence is attributed to a set of proteins, including the proteins coded by the TP53 and BCL2 genes (Behrens et al., 2009; Ibañez et al., 2014; Tabarés-Seisdedos et al., 2009), both of which are down-regulated in cancers and up-regulated in CNS disorders. Using ‘colorectal carcinoma’ and ‘schizophrenia’ as the inputs to the CaNDis web server, we successfully identified the TP53 tumor suppressor gene and the BCL2 anti-apoptotic gene as those in common to both the diseases (BCL2 network is shown in Fig. 1). In addition, we also found proteins that have not yet been associated with either disease (Piñero et al., 2020), but which could play a role in this inverse disease co-occurrence due to their interactions with the protein coded by the BCL2 gene (the list of proteins is in the caption of Fig. 1). Literature search showed that some of the identified proteins are indeed associated with both diseases. For example, the oncoprotein coded by the DDIT3 gene (Engström et al., 2006) is associated with lung cancer (Li et al., 2015) and upregulated in schizophrenia (Umeda-Yano et al., 2013). Similarly, MAPK8 plays a role in both colorectal cancer and schizophrenia (Chandrasekaran et al., 2012; He et al., 2012; Sztakiewicz et al., 2014). This illustrates the use of the CaNDis web server’s innovative network visualization to quickly suggest candidate proteins with potential new roles in co-occurrence of diseases.

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