RESEARCH ARTICLE

PDTDB – AN INTEGRATIVE STRUCTURAL DATABASE AND PREDICTION SERVER FOR PLANT METABOLITES AND THERAPEUTIC DRUG TARGETS

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ABSTRACT

Understanding the biological phenomena of sequence, structure, function, metabolism, and molecular interactions of species is crucial to identify or analyze the biological problems in the ideal way. Integration of all data facilitates unified access to the key problem. PDTDB (Phytochemical and Drug Target DataBase) is an integrative structural database, which contains information about the plants (secondary metabolites, plant anatomy, side effects, and medicinal properties), ligands (3D molecular structure, SMILES string, side effects, and medicinal properties), therapeutic drug targets (3D molecular structure, sequence, mechanism of the drug target, disease nature, and disease symptoms), molecular dockings (target-ligand complex interaction, drug action, home remedies, and docking results), and structure activity relationship between diseases, therapeutic targets, plants, and phytochemicals. The database provides keyword or accession number search, similar phytochemical structures search using SMILES string or by sketching the structure in the chemical structure editor, phytochemical structures with similar molecular formula search, binding structure search, and browse and/or search database entries from the sortable table. It supports predicting physicochemical properties of sequence and structures, and interactive visualization of structures in various models. PDTDB is freely accessible at https://pdt.biogem.org.

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INTRODUCTION

Throughout the ages, natural products have formed as the major and valuable resource for health and medicine to the humans (Bagetta, 2012; Buss and Butler, 2010; Lahlou, 2007). Most of the drug discoveries of modern medicines were either extracted from natural products or inspired by them (Prasad and Tyagi, 2015; Ji et al., 2009; Li and Vederas, 2009). Natural product consists of compounds that are derived from natural sources such as plants, minerals, and animals (Natural Products Research, 2016; Lahlou, 2007; What Are the Pharmaceutical Sources of Drugs, 2016). Phytonutrients or phytochemicals are the organic bioactive chemical compounds present in the plants that have disease preventive properties (What Are Phytonutrients, 2016; Lahlou, 2013; Omotayo and Borokini, 2012; A-Z of Natural Foods and Nutritional Benefits, 2016). Natural products have the tendency to improve the body’s vital ability to balance and to heal itself (A Close Look at Naturopathy, 2016). Many research studies show that consuming phytonutrient rich foods and beverages prevent from disease (Phytochemicals’ Role in Good Health, 2016; Health Benefits of Fruits and Vegetables, 2016; A-Z of Natural Foods and Nutritional Benefits, 2016). But some of the phytochemicals loss or change its chemical structure during cooking foods, due to thermal decomposition and chemical reactions (Palermo et al., 2014). The success of natural products in drug discovery is, it consists of unique and more complex molecular structures while compared to synthetic molecules (Lahlou, 2013; Krishnamoorthy and Subramaniam, 2014). Due to its nature, selection of binding regions in the drug target for docking with phytochemicals is more (Lahlou, 2007; Lahlou, 2013).

Recently, many big pharmaceutical companies and R&D organizations have renewed their interest in discovering potential lead compounds from the natural products for life threatening diseases (Lahlou, 2007). In fact, it is estimated that about 49% of modern medicines is either natural products or directly derived (semi-synthetic) therefrom (Newman and Cragg, 2016; Newman and Cragg, 2007; Cragg and Newman, 2013). With the advances in modern drug discovery, three-dimensional molecular structure and pharmacokinetic properties of many phytochemicals were identified. Despite there are still many thousands out there which are yet to be
discovered (Lahlou, 2007). The environmental changes over years has destroyed many valuable medicinal plants and mineral resources.

Over the past decade, several commercial and open access 3D structural databases for natural products, drug targets, and molecular interactions were released with different specifications and large entries. Even though the databases have quantitative information, there no solid database containing both integrated data and properties prediction servers. For this purpose, we have been developing an integrative structural database which overcomes the problem.

**MATERIALS AND METHODS**

**Data Collection**

Information and/or image of the herbal plants/natural products, phytochemical compounds, drug targets, symptoms and causes of the diseases, and natural remedies to cure the diseases were collected and manually curated from the WebMd (https://www.webmd.com), eBooks, journal articles, blogs and other web pages through Google search. All information in the database are properly cross referenced by hyperlinks to the corresponding original source web page. The 3D phytochemical structures were retrieved from the PubChem (Kim et al., 2016) / DrugBank (Law et al., 2014) / ChEBI (Degtyarenko et al., 2008) / ChemSpider (Pence and Williams, 2010) / HMDB (Wishart et al., 2013), and 2D phytochemical images were obtained from Marvin JS version 16.7.4 (https://www.chemaxon.com) tool by manually drawing the structure or importing the structure file retrieved from the database. Some of the information about rare phytochemical compounds was retrieved from the book “Encyclopedia of Traditional Chinese Medicines: Molecular Structures, Pharmacological Activities, Natural Sources and Applications, (Vol. 1, 2, 3, 4, 5, and 6)” (Zhou et al., 2011). The unavailable 3D structure of phytochemical compounds was obtained manually by drawing the phytochemical structure in the JSDraw v4.0.5 (http://www.scililgence.com) tool and then the SMILES string is translated to the 3D molecule using CACTUS (https://cactus.nci.nih.gov/translate)/Balloon (Vainio and Johnson, 2007) / Open Babel (O’Boyle et al., 2011) / CORINA (Sadowski et al., 1994) tool.

Based on several literature studies, the therapeutic targets and binding phytochemical compounds were chosen. The 3D structure of drug targets was retrieved from the Protein Data Bank (PDB) (Berman et al., 2000). Details of binding affinities, interactions, drug-like properties, and drug action of drug target and phytochemicals were gathered from the BindingDB (Gilson et al., 2016), KEGG (Kanehisa and Goto, 2000), DrugBank (Law et al., 2014), PharmGKB (Hewett et al., 2002), STITCH (Kuhn et al., 2014), and SuperTarget (Hecker et al., 2012) database. Target-ligand interaction studies are carried out using CLC Drug Discovery Workbench version 3.0.1 (https://www.qiagenbioinformatics.com) and LigPlot’ (Laskowski and Swindells, 2011) tools.

**Database Construction**

The PDTDB was constructed on CentOS Enterprise v5.8 x86_64 Linux server with Apache HTTP Server v2.2.31, MySQL Community Server v5.5.50, PHP v5.4.45, and phpMyAdmin v4.0.10.14. The interactive web interface was designed and implemented using jQuery, AJAX, JavaScript, Dojo, HTML5, and CSS3. A JavaScript molecule viewer tool JSmol v14.2.9 (Hanson et al., 2013) is embedded in web pages to interactively display the phytochemical compounds in ball-and-stick model, drug targets in ribbon model, and target-ligand complex in docking interaction model. A Java chemical structure editor JSDraw version 4.0.5 (http://www.scililgence.com) is used for searching similar phytochemical structures.

![Figure 1. Workflow diagram and database schema of the PDTDB](image)

A simple workflow diagram representing the database architecture, database schema, entity relationship model of the database, entity integration, content in the database, and computation methods are shown in the Figure 1. The architecture of PDTDB is broadly categorized into graphical user interface (front-end) and server storage interface (back-end). In a graphical user interface, it consists of user interactive web pages, which are accessible for the users (can search, browse, and download data), group members (can add data, and predict data), and administrator (can add, modify and delete data; enable or upgrade modules; control user access; and server monitoring). The site administrator has direct access to the server storage interface and several modules in the PDTDB, through administrator portal. In a server storage interface, it consists of MySQL server which is connected to the phpMyAdmin. MySQL server stores the plant, ligand, target, disease, and docking information; and links to the files stored on the local disk.

**RESULTS AND DISCUSSION**

PDTDB is a tri-functional and integrated database, it serves as a plant database, drug target database, and docking database. Currently the PDTDB has published 61 records of medicinal plants/natural products (including major entities such as secondary metabolites, description and health benefits and image), 237 records of phytochemical ligands (including major entities such as SMILES, description and health benefits,
image, and 3D structure), 8 records of targets (including major entities such as description, sequence, 3D structure, and image), 8 records of diseases (including major entities such as description, symptoms, and image), and 12 records of target-ligand docking (including major entities such as score, no. of interactions, herbal recipe, drug action, 3D structure, and image). Each plant image represents leaf, fruit, stem, and/or seeds. Also there are nearly 307 therapeutic drugtarget and disease data, 1,657 plant data, 49,589 ligands, and 25 docking data under review. PDTDB is still under development, as we are currently focused on data curation and construction of several automatic prediction and validation servers to increase data accuracy and provide more information. We are constantly striving to complete the process within a short span. Also, server scripting for automatic integration and data extraction from the external databases or servers is under progress.

Accessing the Database

The web interface of PDTDB was designed for the benefit of both biologists and non-biologists, so that users can easily access the targeted information without much technical skill. The hyperlinks in the PDTDB web pages are annotated to provide additional information in the form of thumbnail images or tooltip text by simply moving the mouse pointer over the hyperlinked text. In the ‘Help’ menu of PDTDB, ‘Troubleshoot’, ‘Tutorial’, ‘PDTDB Entries’, and ‘Resources and Tools’ web pages are included to provide more information about accessing the database. The data entities in PDTDB comprises of ‘Target’, ‘Plant’, ‘Ligand’, and ‘Docking’. Information from the PDTDB can be accessed through the ‘Search’ page, ‘Browse’ page, ‘Binding’ page, and ‘Download’ page. A brief description about the web page interfaces and accessing methods are given in below sections.

1. Search page

In ‘Search’ page, there are various types of search methods available to retrieve the data from PDTDB, which are listed as follows:

- **Keyword Search** method allows users to perform text-based search the ‘keywords’ field of the database.
- **Unique ID Search** method allows users to perform accession number search on the ‘id’ field of the database. The format for PDTDB accession numbers are 5 letters (database name ‘pdtdb’) + 1 letter (types of entries ‘t’ – target, ‘p’ – plant, ‘l’ – ligand, and ‘d’ – docking) + 5 digits (serial number of the database entry in left padding with zero format). For instance, pdtdb00282.
- **Simply Structure Search** method allows users to perform SMILES string search on the ‘smiles’ field of a ligand table in the database. On pressing the ‘JSDraw’ button, a chemical structure editor is displayed. By drawing the 2D chemical structure in the editor, users can search on the ‘smiles’ field of a ligand table in the database or PubChem database.
- **Chemical Formula Search** method allows users to perform molecular formula search on the ‘formula’ field of a ligand table in the database. Chemical formula search is included in the ligand/docking result page near the section “Molecular Formula :”. By clicking the hyperlinked formula, it searches for matching records in the ligand table with similar chemical formula.

2. Browse page

The ‘Browse’ page contains four accordion panels with sortable tables inside. Each table in the accordion panel consists of the list of entries with the following fields: accession number, name, and alternative names/keywords. The rows in the table can be sorted in ascending/descending order according the fields, by clicking the column header in the table. Also, users can filter or shorten the list of entries in the table by typing a text in the search box. From the ligand table, to facilitate the formula search or sort, the superscript and subscript of the molecular formula were removed. For this reason, users can able to search or sort the molecular formula by simple text search. For example, C29H50O2 can be searched as linear text C29H50O2 or C29/H50/O2.

3. Binding page

Binding search allows users to predict the structure activity relationship (SAR) of the biomolecules. Currently PDTDB supports target-wise relationship and ligand-wise relationship prediction of biomolecules. The result page of binding search is displayed as a sortable table containing target-ligand interaction, target details, ligand detail, a list of plants containing the ligand, docking score, and target-ligand interaction bonds.

4. Download page

The ‘Download’ page consists of four download options, namely ‘Summary’, ‘Structures’, ‘Ligands’, and ‘Sequences’. In each section, AJAX enabled autocomplete text box is used to fetch the result instantly while typing the query. The allowed query search is Comma Separated Value (CSV) or Space Separated Value (SSV) formatted accession numbers. Users are allowed to download large numbers of sequence/structure/text files from the PDTDB as batches in compressed file format (.zip).

Figure 2. Network model of data integration in the PDTDB

The binding network model (Figure 2) represents the data integration of entities in PDTDB and structure activity relationship between diseases, therapeutic targets, plants, and phytochemicals.

Features of PDTDB

Apart from serving as a database, the PDTDB allows users to predict molecular interactions of target-ligand complex, predict physicochemical properties of sequence/structure, and search for similar 3D structures in PubChem and PDB database. The physicochemical properties prediction of sequences and
structures includes: (i) length of the protein sequence, (ii) molecular weight of protein and phytochemical compound, (iii) isoelectric point of the protein, (iv) amino acid composition, (v) total number of negatively and positively charged amino acids, (vi) atomic composition of the protein, (vii) molecular formula of the protein and phytochemical compound, (viii) total number of negatively and positively charged amino acids, (ix) instability index of the protein, (x) estimated half-life of the protein, (xi) grand average of hydropathicity (GRAVY) of the protein, and (xii) molecular interactions of protein-ligand complex.

The build-in three-dimensional molecular visualization tool enables interactive visualization of structures in various models, based on the type of the molecules (Figure 3).

**Conclusion**

PDTDB is an integrated, comprehensive, and open access resource for medicinal herbs/natural products, phytochemicals, drug targets, docking results, symptoms and causes of the diseases, and natural remedies to treat the diseases. All entities in the PDTDB are integrated by the related entries to provide more information. The 3D structure of all phytochemical and interacting target proteins can be accessed for each medicinal herbs/natural products. Also, other related interactions of target and phytochemical compounds can be achieved. In future, the PDTDB will be enhanced with more automatic prediction/validation server for physicochemical/pharmacokinetic properties, docking evaluation, visualization tool for target-ligand-plant-docking interaction network, and integrate with the external databases and prediction servers.

**Conflicts of interest**

The authors declare that they have no competing interests.

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