SerpentinaDB: a database of plant-derived molecules of *Rauvolfia serpentina*

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**Abstract**

**Background:** Plant-derived molecules (PDMs) are known to be a rich source of diverse scaffolds that could serve as a basis for rational drug design. Structured compilation of phytochemicals from traditional medicinal plants can facilitate prospection for novel PDMs and their analogs as therapeutic agents. *Rauvolfia serpentina* is an important medicinal plant, endemic to Himalayan mountain ranges of Indian subcontinent, reported to be of immense therapeutic value against various diseases.

**Description:** We present SerpentinaDB, a structured compilation of 147 *R. serpentina* PDMs, inclusive of their plant part source, chemical classification, IUPAC, SMILES, physicochemical properties, and 3D chemical structures with associated references. It also provides refined search option for identification of analogs of natural molecules against ZINC database at user-defined cut-off.

**Conclusion:** SerpentinaDB is an exhaustive resource of *R. serpentina* molecules facilitating prospection for therapeutic molecules from a medicinally important source of natural products. It also provides refined search option to explore the neighborhood of chemical space against ZINC database to identify analogs of natural molecules obtained as leads. In a previous study, we have demonstrated the utility of this resource by identifying novel aldose reductase inhibitors towards intervention of complications of diabetes.

**Keywords:** Plant-derived molecules, *Rauvolfia serpentina*, Database, Drug discovery, Virtual screening, ADMET

**Background**

Plants have evolved to produce a diverse repertoire of secondary metabolites which have been used as a source of remedial agents [1, 2]. Medicinal plant extracts have been known for their efficacy against various diseases, and are classically used to discover drug-like molecules. Phytomedicines continue to play a central role in the health management systems in developing countries which include 65% of Indian population. Recent World Health Organization (WHO) review estimates that almost 80% of world's population depends on traditional medicines [3]. These indicators have impelled WHO to incorporate phytomedicines in health care systems. PDMs have also been recognized to provide specific substructures or scaffolds that make them comparable to trade drugs and their potential utilization in combinatorial chemistry [4].

Therefore, there is ample scope for rationalizing the process of drug discovery by prospecting for plant-derived molecules (PDMs) with virtual screening approach. PDMs could be effectively used to systematically extract unique molecular scaffolds, which could further be chemically elaborated to generate novel leads and to screen molecules from drug-like libraries [5, 6]. Computational approaches, such as molecular docking, ligand-based virtual screening, and molecular dynamics (MD), have been widely used in modern drug discovery to explore drug-receptor interactions, and have been able to restrain the number of PDMs that confront experimental validation that ultimately reducing the cost of drug development [1, 7–9]. Thus, hypothesis driven implementation of such pharmaco-informatics pipeline hastens the rate of drug discovery of natural molecules and their simpler mimetics with better pharmacological properties.

*Rauvolfia serpentina* is an important medicinal plant endemic to the Himalayan mountain range of Indian
subcontinent and South-East Asian countries [10]. Plant extracts of R. serpentina have been reported to be of therapeutic value against various diseases (Table 1) including hypertension, intestinal disorders, eye diseases, cuts, wounds, splenic diseases, uterine contraction, headache, and skin diseases [11]. Its extracts have also been reported with a broad range of therapeutic effects such as antioxidant, antiaging, anti hypertensive, anticancerous, antimalarial, antiinflammatory, anti fibrillar, anthelmintic, antiarrhythmic, anticholinergic, antidysentry, antidiarrhoeal, antihypotensive, anticontractile, antipyretic, antidiuretic, sympathomimetic, and antipsychotic [10–12]. Knowing the potential efficacy of R. serpentina PDMs and their derivatives, its phytochemical space could be effectively explored for systematical extraction of unique molecular scaffolds and their derivatives [6]. This strategy has been followed to identify ‘2 PDM leads’ and their 16 structural analogs as potent aldose reductase inhibitors (Additional file 1) [6].

With the aim of providing a comprehensive resource for rational prospection of R. serpentina PDMs towards drug discovery, we compiled an extensive, structured database of its molecules. After a thorough literature survey, details of PDMs were manually compiled and curated. We present a database, SerpentinaDB (Fig. 1), which is structured to include plant part source, chemical name, chemical class, IUPAC (International Union of Pure and Applied Chemistry) names, SMILES (Simplified Molecular-Input Line-Entry System) notations, and 3D (3-Dimensional) structures for 147 PDMs with all associated references (Additional file 2). These 3D structures are present in the form of mol2 file format that is amenable for conversion into other file formats that are accepted by various drug discovery softwares. It also provides several physicochemical descriptors of these PDMs which are indicators of their drug-like properties. Hence access to repertoire of PDMs like SerpentinaDB can be of considerable advantage to academia as well as industry.

Table 1 Therapeutic properties of Rauvolfia serpentina extracts reported for various diseases. Disease details include simple name of the disease and its broad classification. Citations for disease associations are provided in Additional file 2.

| S. No. | Disease name         | Disease type   | S. No. | Disease name         | Disease type   |
|-------|----------------------|----------------|-------|----------------------|----------------|
| 1     | Cancer               | Immunological  | 15    | Pneumonia            | Pathological   |
| 2     | Leukemia             | Immunological  | 16    | Asthma               | Immunological  |
| 3     | AIDS                 | Immunological  | 17    | Rheumatism           | Immunological  |
| 4     | Diabetes mellitus    | Digestive      | 18    | Anasarca             | Epidermal      |
| 5     | Hypopolipidemia      | Digestive      | 19    | Helminthias          | Pathological: Parasitic |
| 6     | Alzheimer’s disease  | Neurological    | 20    | Cholera              | Pathological: Bacterial |
| 7     | Schizophrenia        | Neurological    | 21    | Cardiac arrhythmia   | Circulatory    |
| 8     | Skin cancer          | Immunological   | 22    | Diarrhea             | Pathological   |
| 9     | Prostate cancer      | Immunological   | 23    | Tachycardia          | Circulatory    |
| 10    | Hypertension         | Circulatory     | 24    | Supraventricular tachysystole | Circulatory |
| 11    | Fever                | Infectious      | 25    | Thyrotoxicosis       | Immunological  |
| 12    | Insect bite          | Infectious      | 26    | Allergy              | Immunological  |
| 13    | Dysentery            | Digestive       | 27    | Meningitis           | Pathological   |
| 14    | Malaria              | Pathological: Parasitic | 28    | Encephalitic psychosis | Pathological |
software (https://www.chemaxon.com), and further saved into mol2 file format. These files were subjected to energy minimization with Merck Molecular Force Field (MMFF94) using OpenBabel v2.3.1 software [25]. Mol2 format is desirable due to ease of conversion into other molecular formats accepted by drug discovery softwares. To remove redundant entries, data from all resources were merged, and an extensive library of PDMs was compiled. A total of 147 molecules, reported to be extracted from various plant parts, were present in final dataset (Fig. 2a) and broadly classified into different chemical classes (Fig. 2b). Representative PDM of each of seven chemical classes is provided in the supplementary file (Additional file 3). A separate entry was created for molecules that were obtained from more than one plant part leading to 227 such individual entries. Of all these PDMs, mol2 files for 5 of them could not be obtained due to unavailability of both structure as well as IUPAC. Physicochemical properties of the PDMs, such energy, atomic contribution to the partition coefficient (AlogP), distribution coefficient (logD), molecular formula, molecular mass, molecular solubility, and molecular weight (MW), acid dissolution constant (pKa), number of aromatic bonds, number of aromatic rings, and radius of gyration, hydrogen bond acceptor (HBA) count, hydrogen bond donor (HBD) count, number of H acceptor, number of H donor, number of H acceptor (Lipinski), number of H donor (Lipinski), number of H bonds, solvent accessible surface area (SASA), and surface area as well as ADMET (ADMET Solubility, ADMET Solubility Level, ADMET BBB, and ADMET BBB Level) properties, were obtained using molecular property finder tool under small molecules category from the Discovery Studio v4.0 (Accelrys, San Diego, USA). Figure 3 illustrates the statistics of various physicochemical properties, such as MW (Fig. 3a), HBA as well as HBD (Fig. 3b), and molecular volume (MV) (Fig. 3c).

Data architecture and Web interface
SerpentinaDB is hosted in a Server at the Indian Institute of Technology Jodhpur on a Dell Power Edge R910 server running a Linux operating system (Red Hat version 5.5). A total of seven data tables were created to house compiled data. SerpentinaDB implements MySQL, an object-relational database management system (RDBMS) for its backend performance. Web browser interface was created using HTML, CSS, Ajax, JavaScript, and jQuery, which connects MySQL terminal using several PHP scripts. A

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**Fig. 1** Features of SerpentinaDB interface. (a) Demonstration of accessible search options: (1) Plant part, (2) Chemical class as well as plant part, and (3) Physicochemical properties based search of PDMs. (b) Result of input query with list of associated PDMs: this page further provides information of IUPAC, SMILES, physicochemical properties, and 3D visualization with associated references. User can download mol2 file and 2D structure of PDM for given query. (c) Similarity search of selected PDM against ZINC is also available at user-defined percentage.
JMol visualizer (http://www.jmol.org/) and ZINC database (http://zinc.docking.org/) has been embedded in Graphical User Interface (GUI) to provide a 3D visualization and percentage similarity search against ZINC, respectively, for all PDMs. The GUI is designed to be user friendly for data query and extraction, and has been tested in all major browsers (Chrome, Firefox, Safari, and Internet Explorer) and OS platforms.

**Data access**

SerpentinaDB can be explored for PDMs in a number of ways through querying the database with a simple text search tool that provides various options for searching. There are three search sections available to the user with several constraints in each. Search can be performed with (i) plant part, (ii) chemical class, and (iii) physicochemical properties (Fig. 1a). Physicochemical properties search option has advanced search query options for user to select PDMs in a particular range based on MW, number of HBA, number of HBD, and number of aromatic rings. The result for given query is presented in the same page (Fig. 1b) along with information such as PDM ID, plant part, chemical name, chemical class, IUPAC names, SMILES notations, and 3D structure of PDM with associated references. Clicking the drop down arrow provides details of physicochemical and ADMET properties. Two separate links to download mol2 file and 2-Dimensional structure of PDM for given query has been provided. Also, a JMol visualizer (http://www.jmol.org/) has been embedded in GUI to provide a 3D visualization of PDM which can be further downloaded to mol2 file.

Also, each PDM from result page can be searched, to explore the neighborhood of chemical space, against ZINC database to identify analogs of natural molecules at different percentage similarity cut-off (default 90 %). In order to perform this search ZINC database, a curated collection of commercially available chemical compounds [26], is hyperlinked to result page for each PDM separately to return their structural analogs. During this similarity
Fig. 3 Distribution of molecular descriptors of *R. serpentina* plant-derived molecules. (a) Molecular weight. (b) Hydrogen bond acceptor and Hydrogen bond donor. (c) Molecular volume.
search natural molecules are used as scaffolds to search for similar mimetics which may have equivalent biological properties. Thus, SerpentinaDB serves as a portal to facilitate the use of natural chemical diversity for drug discovery through prospection of direct novel leads and their analogs.

Utility and discussion

SerpentinaDB provides comprehensive information of *R. serpentina* PDMs as a structured and integrated library. This database was developed to facilitate prospection of therapeutic molecules from this medicinally important plant. Existing repositories of natural compounds, such as NPACT [27], SuperNatural [28], Herb Ingredients’ Targets [29], and CamMedNP [30], focus on different utilitarian aspects of PDM libraries. While some of these databases emphasize on a specific disease or target-compound interactions, others cover plants of specific geography. SerpentinaDB contains natural molecules of *R. serpentina* which is an important Himalayan medicinal plant reported for various pharmacological properties. While the reported efficacy of *R. serpentina* extract against hypertension has been explored very well to identify specific therapeutic PDMs, its potential against a host of other disorders (Table 1) is hitherto not pinned down to specific molecules. This database can facilitate prospection of novel leads for these disorders from the repertoire of natural molecules.

Natural molecules have been recognized to provide specific scaffolds that make them comparable to trade drugs and their potential utilization in combinatorial chemistry [4]. The MW distribution of PDMs present in SerpentinaDB has been found to follow Gaussian
distribution and peaked in the range of 300-450 Da (Fig. 3a) which is similar to drug-like molecules of previously reported libraries of natural products [31]. Significant number of PDMs have HBA in the range of 3–5 with a sharp decline thereafter, as desired from drug-like molecules (Fig. 3b). Similarly, HBDs of PDMs have a peak at 2 with a sharp drop for higher values, as desired (Fig. 3b). SerpentinaDB PDMs have maximum density in the ‘Lipinski region of interest’ reflecting their drug-like properties and hence their utility in prospection of novel leads. The relevance of SerpentinaDB in drug discovery has been demonstrated with the virtual screening protocol, molecular dynamics, and ZINC similarity search for potential inhibitors of aldose reductase, a target for complications of diabetes [6]. This hypothesis driven prospection study yielded two indole alkaloids as well as their structural analogs as potential AR inhibitors (Fig. 4) [6]. This protocol serves as a demonstration of utility of SerpentinaDB for rational search of therapeutic molecules and highlights its relevance [32, 33]. Future extensions of SerpentinaDB may include 3D structure similarity search and disease associations for each PDM.

Conclusions
SerpentinaDB is an exhaustive resource of R. serpentina molecules facilitating prospection for therapeutic molecules from a medicinally important source of natural products. Pharmacoinformatics pipeline involving virtual screening to perform docking of molecules against disease specific target to identify inhibitors. Hence, compilation of such datasets is essential step towards in-silico drug discovery that hastens the process of prospection of novel leads from natural repertoire with drug-like properties in terms of their biological behavior and toxicity.

Availability and requirements
SerpentinaDB is available at http://home.iitj.ac.in/~bagler/webservers/SerpentinaDB/. Browsers need to be installed with latest JAVA plugins. For more support please consult the FAQs section of SerpentinaDB.

Additional files
Additional file 1: Aldose reductase inhibitors (‘2 PDM leads’ and their ‘16 structure analogs’) obtained using pharmacoinformatics pipeline in the study performed by Pathania et al., 2013 [6].
Additional file 2: Details of therapeutic properties of phytochemicals from R. serpentina against various disorders along with associated references.
Additional file 3: Representative PDMs from each of seven chemical classes of R. serpentina.

Abbreviations
PDMs: Plant-derived molecules; WHO: World Health Organization; MD: Molecular dynamics; 3D: 3Dimensional; 1D: 1Dimensional; 2D: 2Dimensional; IUPAC: International Union of Pure and Applied Chemistry; SMILES: Simplified molecular-input line-entry system; DNP: Dictionary of Natural Products; MWFF94: Merck Molecular Force Field; ALOGP: Partition coefficient; LogD: Distribution coefficient; MW: Molecular weight; PKa: Acid dissociation constant; HBA: Hydrogen bond acceptor; HBD: Hydrogen bond donor; SASA: Solvent accessible surface area; MV: Molecular volume; ADMET: Absorption, distribution, metabolism, excretion, and toxicity; RDBMS: Relational database management system; GUI: Graphical User Interface; OS: Operating System.

Competing interests
The authors declare no conflicts of interests.

Authors’ contributions
GB and SP conceived the idea. SP performed data collection and generated 3D structures. All authors contributed to data analysis, discussion of results as well as writing of the paper. SMR created the database architecture and interface. This work is part of PhD project of SP.

Authors’ information
SP and VR are the PhD students of bioinformatics focusing on hypothesis driven in-silico drug discovery specifically from natural molecules. SMR is master’s student and worked as an intern at IIT Jodhpur, and is keenly interested in developing database architecture and interface. GB is an interdisciplinary researcher with basic training in physics, computer science and computational biology. As the group leader of Complex Systems Laboratory at IIT Jodhpur, he has been involved in rational drug discovery from medicinal plants which he initiated while working as a scientist at CSIR-Institute of Himalayan Bioresource Technology.

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