SuperPain—a resource on pain-relieving compounds targeting ion channels

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ABSTRACT

Pain is more than an unpleasant sensory experience associated with actual or potential tissue damage: it is the most common reason for physician consultation and often dramatically affects quality of life. The management of pain is often difficult and new targets are required for more effective and specific treatment. SuperPain (http://bioinformatics.charite.de/superpain/) is freely available database for pain-stimulating and pain-relieving compounds, which bind or potentially bind to ion channels that are involved in the transmission of pain signals to the central nervous system, such as TRPV1, TRPM8, TRPA1, TREK1, TRESK, hERG, ASIC, P2X and voltage-gated sodium channels. The database consists of ~8700 ligands, which are characterized by experimentally measured binding affinities. Additionally, 100 000 putative ligands are included. Moreover, the database provides 3D structures of receptors and predicted ligand-binding poses. These binding poses and a structural classification scheme provide hints for the design of new analgesic compounds. A user-friendly graphical interface allows similarity searching, visualization of ligands docked into the receptor, etc.

INTRODUCTION

Ion channels are proteins forming a pore that allows the flow of ions across membranes. Ion channels are voltage or ligand gated. Some of these proteins help nerve cells to transmit pain signals to the central nervous system and are therefore promising targets for the development of pain therapeutics.

Transient receptor potential channels (TRPs) are a family of 28 human cellular ion channels, varying in homology to each other but all with six transmembrane regions in common, and are nonselectively permeable to cations. There are seven subfamilies that can be divided into two groups. Group 1 includes TRPC, TRPV, TRPA, TRPM, TRPN, and group 2 comprises TRPP and TRPML (1). Compounds perceived as hot stimulate the vanilloid receptor (TRPV). At the same time, this receptor plays a crucial role in pain mediation and is therefore an interesting drug target. Known pungent chemicals with high receptor affinity such as capsaicin or resiniferatoxin were lead structures in drug development toward desensitization (2). Recent research focuses on novel analgesic mechanisms like positive allosteric modulation (3) and broad-spectrum TRP antagonists (4).

The mechanism of cold-induced analgesia was unclear until the discovery of TRPM8. This receptor is also known as the cold or menthol receptor (5). It is activated by cold temperatures and cooling agents, such as menthol or icilin, allowing the entry of Na⁺ and Ca²⁺ to the cell (6). Two modulating mechanisms are generally discussed. Whereas antagonists physically block the receptor for cold and menthol, agonists activate TRPM8 and generate a cooling sensation. Selective ligands could be used as a new generation of analgesic drugs in neuropathic pain (7,8).

TRPA1 plays a key role in chemical sensing in the inflammatory pain pathway. Many small molecules, including ingredients like wasabi, horseradish, garlic and mustard oil, can activate the channel. Recently, it has been shown that desensitizing TRPA1 could help in the treatment of neuropathic pain (9,10).

The human Ether-à-go-go Related Gene (hERG) channel or KCNH2 is a voltage-gated potassium channel. It has been the focus of pharmaceutical research for years because the inhibition of hERG potassium channels by drugs can lead to cardiac arrhythmia (11). In 2010, Stary and colleagues published a homology model (12) and a group from Canada investigated structural mechanisms of state-dependent

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drug binding (13). The different binding sites for ions and compounds are defined and the International Conference of Harmonization set up guidelines for drug development. Therefore, new screening methods for the prediction of drug liability to the hERG channel have been developed (14).

Voltage-gated sodium channels are activated through action potential firing. They represent the target for local anesthetic agents. It is challenging to find selective inhibitors of sodium channels in the pain pathway. Genotyping of families suffering from congenital indifference to pain identified mutations in gene coding for Nav1.7 channels (15). A lot of research is performed to find specific blockers to the treatment of pain and epilepsy (16).

There are some complementary resources on ion channels. Multiple analysis for voltage-gated potassium channels from different species is compiled in the voltage-gated potassium channel database VKCDB (17). Protein–protein interactions (PPIs) are the focus of the TRIP database (18): Shin and colleagues manually curated 653 PPIs for mammalian TRP channels. MoleOnline 2.0 (19) is a web server that provides interactive channel analysis to identify active sites.

The International Union of Basic and Clinical Pharmacology (IUPHAR) database provides information on human and rodent receptors (20). There is a need for a specific resource for pain-relieving compounds targeting ion channels and their 3D homology models so researchers can identify targets and putative ligands.

MATERIALS AND METHODS

Here, we shortly describe our methods used. If you are interested in further details, please read the ‘About’ page on our Web site.

Compounds

To identify pain-related receptors and ligands, scientific literature was screened via text mining and subsequent manual evaluation. Therefore, we first downloaded Medline/PubMed data from the NCBI FTP site in xml format. Using the search engine library Apache Lucene (http://lucene.apache.org) and a tool kit for processing text with computational linguistics (http://alias-i.com/lingpipe), the data was indexed. The search engine, written in Java, dynamically queries the indexed data and results in a structured query language (SQL) file containing the textmining hits. An example for a query for the literature search is as follows: [lidocaine (TI) AND ic50 (TI)] OR [lidocaine (abstract) AND ic50 (abstract)]. Found hits were put in a preliminary database and are displayed with keywords highlighted in different colors. In the manual evaluation process, confirmed experimentally determined affinities were moved to the final database. In some cases, the full text of the article was checked. Further information regarding the molecules was retrieved from the PubChem database (21). PubChem is a freely available database, which is provided by the National Center for Biotechnology Information (NCBI). It contains detailed information on ~30 million compounds. All those compounds and additional information were put in a MySQL database. Additional affinities were found via a search in BindingDB (22). BindingDB is a database on experimentally determined protein–ligand interactions. It provides ~782,000 affinities for 6500 protein targets.

Compound clustering

The experimentally determined ligands had to be clustered regarding their similarity to each other by means of a K-means algorithm. As in the K-means algorithm, the number of clusters has to be defined the algorithm was slightly modified. To ensure that the most similar compounds are members of one cluster, a neighbor-joining algorithm was used. The R package with heatmap.2 was used to display the compound similarities in a heatmap.

In silico screening

Putative ligands were indentified through a similarity search. Therefore, the structural fingerprints of each ligand were calculated and used for a similarity search within the PubChem Compound database. The similarity between experimentally determined ligands and potential ligands was calculated with the Tanimoto coefficient. Depending on the chemical topological properties it gives values between zero (no similarity) and one (identical). Compounds with a coefficient >0.90 were classified as putative ligands and included in the database.

Receptor structure

Currently, no radiographic crystallographic structures are available for most of the receptors, but there is some information on the active sites of some of the molecules. To perform dockings a homology model for TRPV1 had to be created. The template structure for the homology modeling of TRPV1 and the models of P2X and ASIC were retrieved from the Protein Data Bank (PDB). The PDB is a freely available database on 3D structures of proteins and nucleic acids (23,24). Models are downloadable and provide coordinates of each atom within the molecule. A homology model of the hERG channel was retrieved from a research group in Germany (12).

Docking

To obtain an understanding of binding mechanisms, the homology model of TRPV1 and the models of P2X, ASIC and hERG were imported into Accelrys Discovery Studio (25). About 6000 ligands were docked into the binding sites with the integrated Docking Tool LibDock. LibDock is a high-throughput docking algorithm. Based on polar and apolar interaction sites up to 1000 ligand conformations were positioned in the binding site. The five best-ranked poses regarding energetic conditions are displayed on the Web site. ‘JSmol’ was used to implement a molecular viewer, which is JavaScript based.
Web site

The Web site is based on PHP (http://www.php.net/); web access is enabled through Apache HTTP Server (http://httpd.apache.org/). We recommend a recent version of Mozilla Firefox or Google Chrome; alternative browsers like Microsoft Internet Explorer and Apple Safari were tested with some configurations. JavaScript must be enabled, as it is required for all features of the site.

RESULTS

Compound database

A total of 100,000 putative channel modulators were included of which 12,000 are purchasable and 8700 ligands have measured binding affinities.

Compound clustering

A structural clustering of the 8700 compounds with an internal similarity (Tanimoto) above 0.7 was performed and resulted in clusters with member sizes up to 52 compounds. For a better visualization, the member size was limited to 30. The neighbor-joining algorithm resulted in 684 clusters. The similarities are displayed in interactive heatmaps.

Receptor structure

The homology model of TRPV1 was based on the known structure of the Kv1.2 potassium channel, which exhibits significant similarity and a high resolution (2.4 Å). The structure was downloaded in PDB format (PDB-ID: 2R9R). Homology modeling was carried out in accordance with Fernandez-Ballester et al. (26) using the SWISS-MODEL server alignment mode. PyMOL was used to create a tetramer and manual refinement was carried out with Accelrys Discovery Studio.

Docking

The docking of 6000 TRPV1, P2X,ASIC and hERG ligands in >1000 different binding poses revealed five binding poses for each ligand. Figure 1 shows capsaicin in the binding site of TRPV1 in an active interactive view. These binding poses were chosen regarding energetic conditions. The results are embedded into the database. Each ligand can be displayed in different manners, as well as the receptor itself (ball and stick, spacefill, etc.). Holding the left mouse button while moving the mouse leads to a rotation. A right click brings up a drop-down menu for zooming, measurements, color adjustments, etc. Safari or Tablet users can use their common gestures to navigate through the docking results. Each binding pose can be exported as a graphic (PNG) or file (MOL).

Usage of the database

There are different ways to use the database and browse through the data. Figure 2 summarizes the main functionalities of the Web site.

‘Receptor’ holds information on pain-related ion channels. The homology model of TRPV1 and docking results can be found there.

There are two search boxes in the ‘Compounds’ section. Specific compounds can be found through a property search by typing in the name, PubChem ID, IUPAC or SMILES. It is also possible to search for a specific target such as TRPV1 or hERG or to select certain features like purchasability, IC50, EC50, molweight, rotatable bonds, etc. On the results page, information on compounds including vendors is listed in a table. This table can be used as a starting point for a similarity search or to

Figure 1. Capsaicin in TRPV1. Capsaicin was put into the homology model of TRPV1 using a high-throughput docking algorithm. Each docked ligand is shown in the interactive view.
compare structures directly by viewing the compound clusters. In the ‘Cluster’ section all clusters are listed with one compound of the cluster. A search box enables quick access to a distinct cluster. Clicking on a compound also enables browsing through different clusters. The heatmaps are diagonally divided to compare similarity (Tanimoto) and affinity (IC50). Mouseover shows the structure of the compared compounds. Clicking leads to a comparison page with information about (co-) affinities, as well as chemical information and vendors.

DISCUSSION

SuperPain aims at providing a comprehensive resource on ligands for pain-related ion channels. Acid sensing ion channels (ASICs) belong to the degenerin-epithelial sodium channel superfamily and are voltage independent. They are activated by extracellular acidosis and are involved in different processes, such as taste, mechanosensation and nociception (27). Recently, it has been shown that mambalgin, a three-finger peptide from the venom of the black mamba, suppresses pain in mice without toxicity and fewer side effects than morphine (28). These findings show that ASICs are promising targets in the treatment of pain, especially to avoid addiction problems (29).

In traditional Chinese medicine, tetramethylprazine, sodium ferrulate and puerarin are used in the therapy of pain. It has been found that these compounds target some P2X receptors. These ligand-gated ion channels open in response to binding of adenosine 5'-triphosphate (ATP). Advances in radiographic crystallography allow in silico ligand docking and P2X receptors are becoming therapeutically important drug targets (30).

Further potassium channels of interest are the TWIK-related spinal cord potassium channel (TRESK) and TREK-1. TRESK is a two-pore domain potassium channel that is mainly expressed in dorsal root and trigeminal ganglia (31). This channel is involved in acute and chronic pain and plays an important role in migraine pathogenesis, which makes it a promising target (32). TREK-1 is an interesting target because of its co-expression with TRPV1 and its involvement in polymodal pain perception (33). The present intensive research on 3D structures will improve the development of channel modulators. For example, there are different binding sites for agonists and antagonists (34). The binding sites of the ion channels are rather large, which allows binding of a variety of compounds if certain physiochemical property conditions are fulfilled. This is reflected by the great diversity of the pain-related compounds. They show a low mean similarity of 0.4 compared with other targets like PARP (0.6). It is also important to find out more about the interactions or co-inhibition of different receptors for pain. In the database, they can be found by choosing ‘MultiTarget’ from the ‘Compound Properties’ search box. For example, menthol is known to be an inhibitor of TRPM8 with a half maximal effective concentration (EC50) of 29 000 nM. At the same time, it has also inhibitory potency on TRPA1 with an EC50 of 28 400 nM. Cannabigerol is a compound that occurs naturally in hemp strains. In contrast to other cannabinoids, it is nonpsychoactive because it is not only a β2-adrenergic receptor agonist, but also a mild CB1 receptor antagonist.
Although the compound is not well-studied, it has been found to lower the intraocular pressure (36) and there is some research on treating inflammatory bowel diseases with cannabigerol (37). ‘SuperPain’ stores affinities for TRPA1, TRPM8 and TRPV1 for the compound. These experimentally determined co-affinities suggest that these multi-target compounds might be promising for the development of pain therapeutics (38).

Availability

‘SuperPain’ is publicly available via http://bioinformatics.charite.de/superpain and should be used under the terms of the Creative Commons Attribution-Noncommercial-Share Alike 3.0 License.

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