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Review Article

COMPUTER AIDED DRUG DESIGN: AN OVERVIEW

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ABSTRACT

Discovery and development of a new drug is generally known as a very complex process which takes a lot of time and resources. So now a day's computer aided drug design approaches are used very widely to increase the efficiency of the drug discovery and development course. Various approaches of CADD are evaluated as promising techniques according to their need, in between all these structure-based drug design and ligand-based drug design approaches are known as very efficient and powerful techniques in drug discovery and development. These both methods can be applied with molecular docking to virtual screening for lead identification and optimization. In the recent times computational tools are widely used in pharmaceutical industries and research areas to improve effectiveness and efficacy of drug discovery and development pipeline. In this article we give an overview of computational approaches, which is inventive process of finding novel leads and aid in the process of drug discovery and development research.

Keywords: computer aided drug discovery, structure-based drug design, ligand-based drug design, virtual screening and molecular docking.

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INTRODUCTION

Computational approaches in drug design, discovery and development process gaining very rapid exploration, implementation and admiration. Introducing a new drug in a market is a very complex, risky and costly process

in terms of time, money and manpower. Generally it is found that drug discovery and development process takes around 10-14 years and more than 1 billion dollars capital in total ¹.

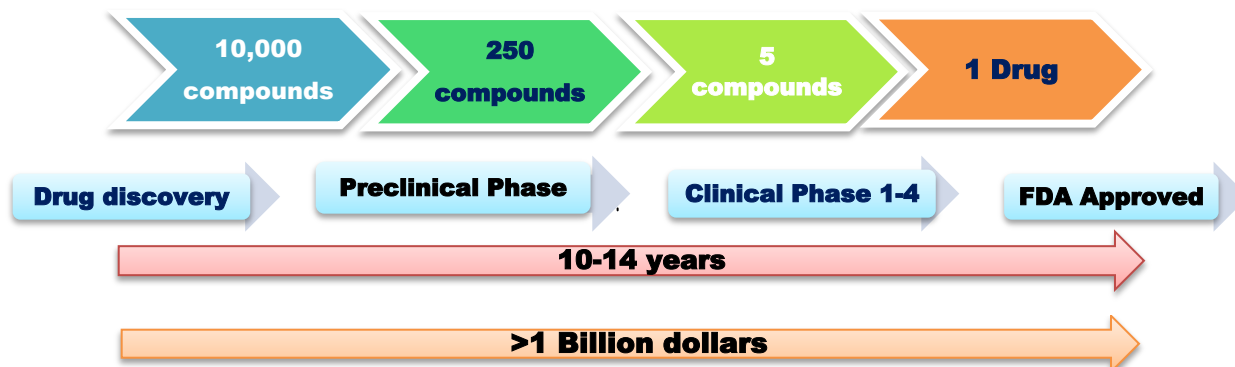


Figure 1: Traditional process of drug discovery and development.

So for reducing time, cost and risk borne factors computer aided drug design (CADD) method is widely used as a new drug design approach. It has been seen that by the use of CADD approaches we can reduced the

cost of drug discovery and development up to 50%². CADD consist use of any software program based process for establishing a standard to relate activity to structure³.

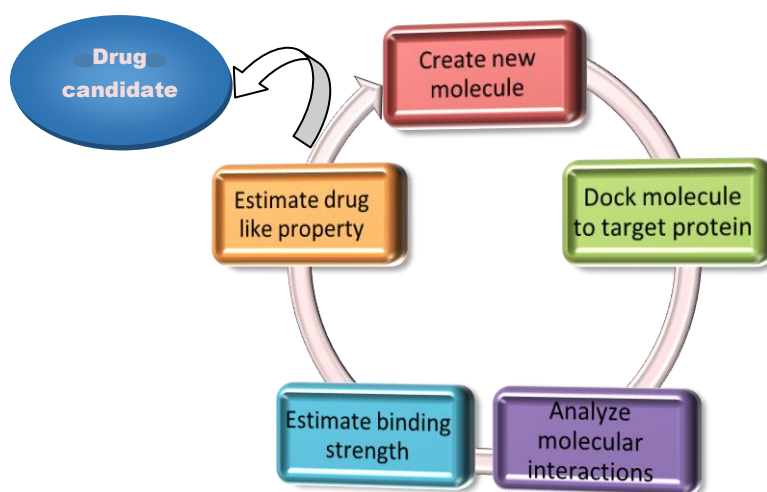


Figure 2: General Principle for Drug design through CADD.

Major types of approaches in CADD

There are mainly two types of approaches for drug design through CADD is the following:

1. Structure based drug design / direct approach
2. Ligand based drug design / indirect approach

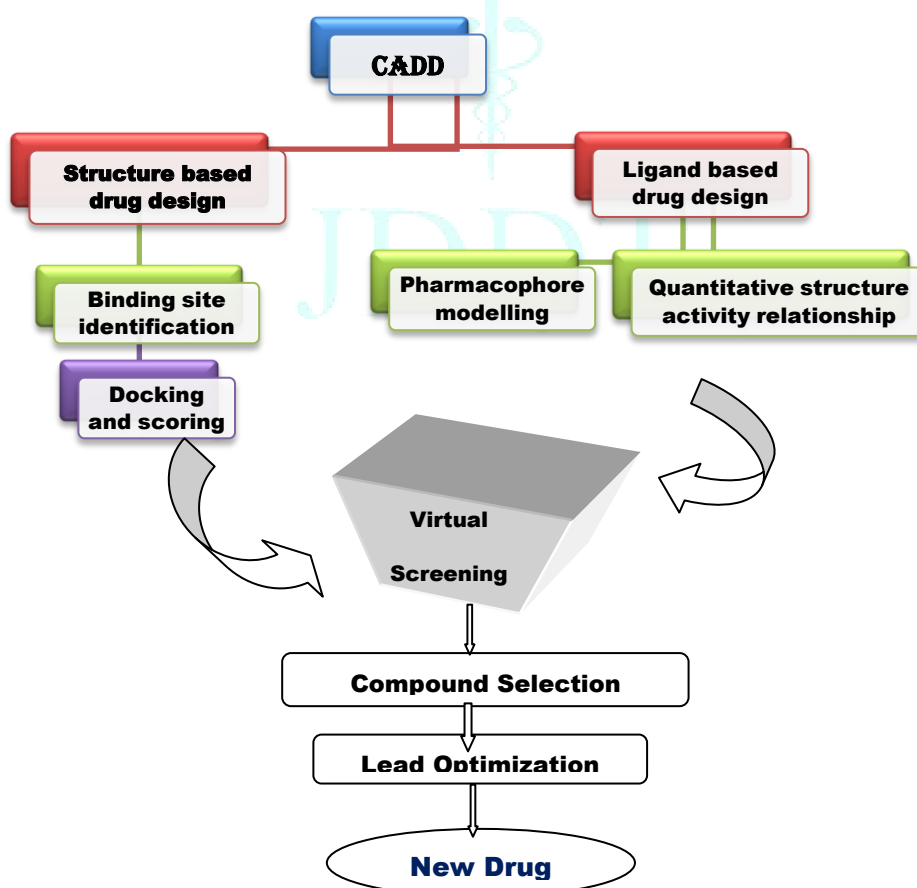


Figure 3: General Representation of workflow for CADD.

1. Structure-based drug design

In SBDD, structure of the target protein is known and interaction or bio-affinity for all tested compounds

calculate after the process of docking; to design a new drug molecule, which shows better interaction with target protein⁴.

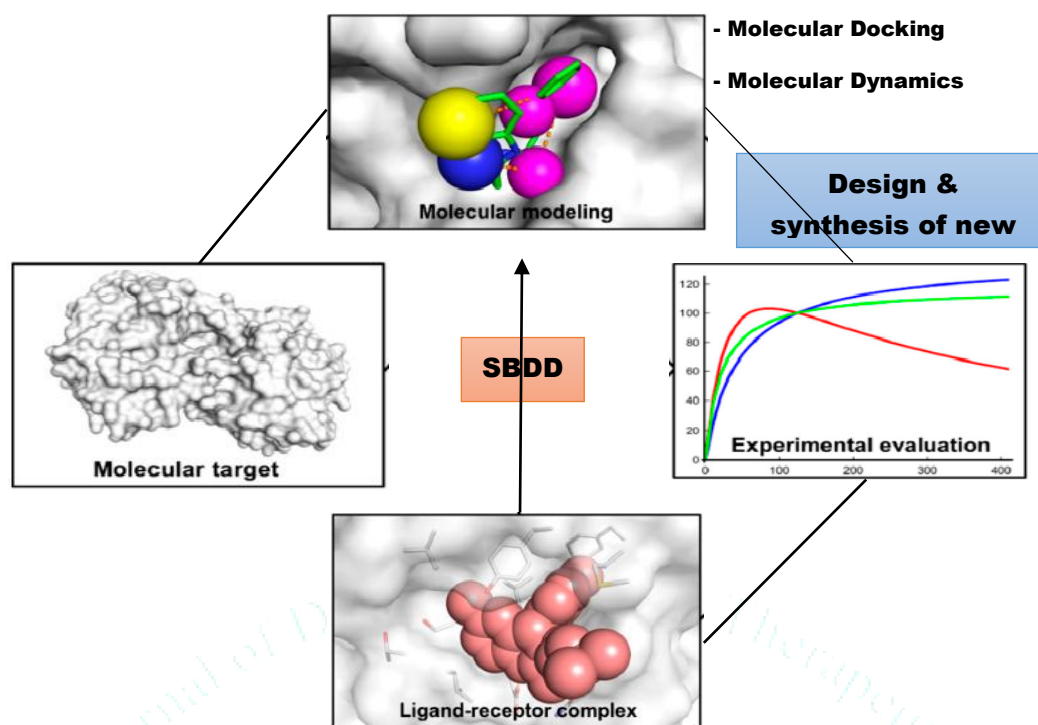


Figure 4: Layout of SBDD⁵.

Overview of the process involved in SBDD

SBDD runs through multiple cycles before the optimized lead reached into clinical trials. The first cycle comprises isolation, purification and structure determination of the target protein by one of three key methods: like X-ray crystallography, homology modeling or NMR. Using compounds comes through virtual screening of different databases are placed into a selected region (active site) of the protein. These compounds are scored and ranked on the bases of steric, hydrophobic, electrostatic interaction of these molecules

with the active site of target protein. Top ranked compounds are tested with biochemical assays.

Second cycle comprises structure determination of the protein in complex with the most optimistic lead of the first cycle, the one with minimum micro-molar inhibition in-vitro, and shows sites of the compound which can be optimized for further increment in the potency. After several additional cycles like synthesis of lead, further optimization of lead through complex structure of protein with lead compound, the optimized compounds generally show marked increment in the target specificity and binding affinity⁶.

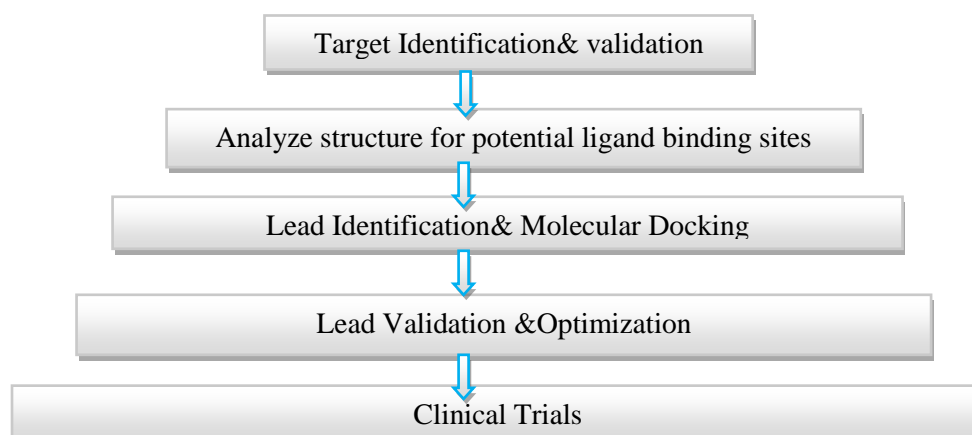


Figure 5: Steps involved in SBDD.

2. Ligand-Based drug design

In LBDD, 3D structure of the target protein is not known but the knowledge of ligands which binds to the desired

target site is known. These ligands can be used to develop a pharmacophore model or molecule which possesses all necessary structural features for bind to a target active site.

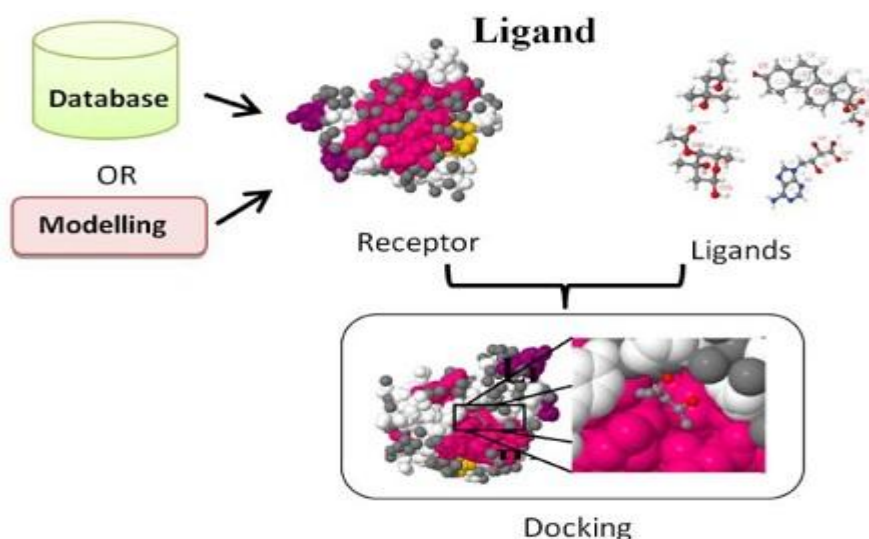


Figure 6: Outline of process involved in LBDD ⁷.

Generally ligand-based techniques are pharmacophore based approach and quantitative-structure activity relationships (QSARs). In LBDD it is assumed that compounds which having similarity in their structure also having the same biological action and interaction with the target protein ⁸.

Virtual screening

Virtual screening has been worked as a most convenient tool now a day to find out the most favorable bioactive compounds with the help of information about the protein target or known active ligands. In the recent time virtual screening is known as a mind blowing alternative of high-throughput screening mainly in terms of cost effectiveness and probability of finding most appropriate novel hit through filter the large of libraries of compounds ⁹.

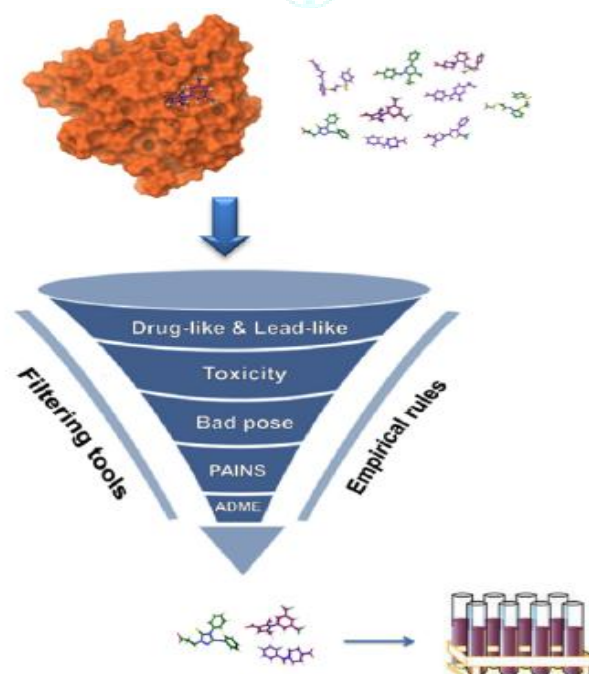


Figure 7: Overview of Virtual screening process ¹⁰.

There are generally two types of virtual screening approaches like structure-based virtual screening (SBVS) and ligand-based virtual screening (LBVS), SBVS method rely on the structure of target protein

active site and LBVS method is based on estimation of calculated similarity between the known active and compound come from databases.

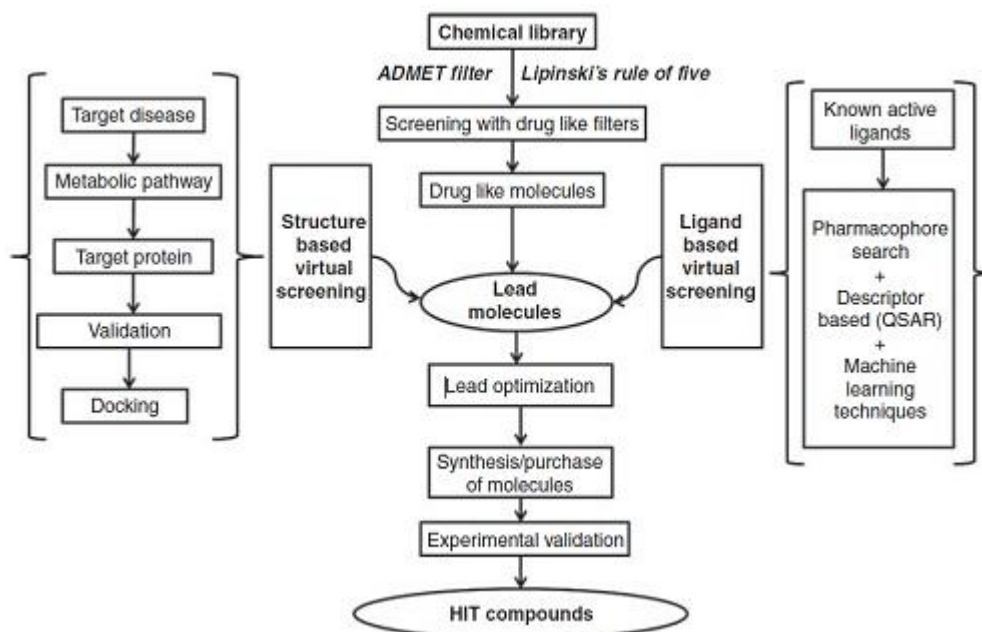


Figure 8: Schematic diagram of VS process for SBDD & LBDD ¹¹

Molecular docking

Molecular docking is *in-silico* method which predicts the placement of small molecules or ligands within the active site of their target protein (receptor). It is mainly

used to accurate estimation of most favorable binding modes and bio-affinities of ligands with their receptor, presently it has been broadly applied to virtual screening for the optimization of the lead compounds.



Figure 9: Process of Docking ¹²

Molecular docking methodology comprises mainly three goals which are interconnected to each other like: prediction of binding pose, bio affinity and virtual screening. In the molecular docking method the basis tools are search algorithm and scoring functions for creating and analyzing conformations of the ligand ¹³.

ADVANTAGES OF CADD

- Through it we can reduce the synthetic and biological testing efforts ¹⁴.
- It gives the most promising drug candidate by eliminate the compounds with undesirable properties (poor efficacy, poor ADMET etc.) through *in silico* filters ¹⁵.
- It is a Cost-effective, time saving, Rapid and automatic process.
- Through it we can know about the drug-receptor interaction pattern.

- It gives compounds with high hit rates through searching huge libraries of compounds *in silico* in comparison to traditional high throughput screening ¹⁶.
- These approaches minimize chances of failures in the final phase.

CONCLUSION AND FUTURE ASPECTS

Computer aided drug design is an efficient tool in the area of drug discovery and development, through it we can find the most promising drug candidate in a very cost-effective way. It always provides a hope for betterment in drug discovery area. In the past years through Computer aided drug design many impressive researches are achieved so it will play a very much important role in the near future. With the current achievement's, there is a promising future of computer aided drug design to aid drug discovery of many more curatives in future.

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