MOLECULAR DOCKING STUDY ON 1H-(3,4-d) PYRAZOLOPYRIMIDINES AS CYCLIN DEPENDANT KINASE (CDK2) INHIBITORS

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Abstract
Objective: CDK2 inhibitors are implicated in several carcinomas viz. Carcinoma of lung, bladder, sarcomas and retinoblastoma. Pyrazolopyrimidines, being purine bioisosters inhibit more than one type of kinase. In this study, we are studying some novel derivatives of 1H-pyrazolo [3,4-d] pyrimidines not reported earlier. The objective of the present study is an attempt towards design and development of 1H[3,4-] pyrazolo-pyrimidines as CDK2 inhibitors through rational drug design.

Methods: The present study has been done on CDK2 structure, PDB ID, 3WBL, co-crystallized with ligand PDY from RCSB protein data bank. A series of seventeen 1H-Pyrazolo [3,4-d] pyrimidines feasible for synthesis was docked on the said CDK2 receptor using Auto Dock 4 version, 1.5.6. Outputs were exported to discovery studio 3.5 client for visual inspection of the binding modes and interactions of the compounds with amino acid residues in the active sites.

Results: The results of docking studies revealed that the present series of 1H-Pyrazolo[3,4-d] pyrimidines is showing significant binding through hydrogen bonding, hydrophobic, pi and Van der waals interactions similar to the ligand PDY. Some conserved H-bond interactions comparable to bioisosters and pyrazolo-pyrimidines not reported earlier. The objective of the present study is an attempt towards design and development of 1H[3,4-] pyrazolo-pyrimidines as CDK2 inhibitors through rational drug design.

Conclusion: The series designed and docked can be further developed by synthesis and in vitro and in vivo activity. The receptor inhibitory activity can also be checked by specific receptor assays.

Keywords: Cyclin-dependent kinase-2 inhibitors (CDK2 inhibitors), 3WBL, Pyrazolopyrimidines, AutoDock 4

INTRODUCTION

Protein kinases are enzymes that modify other proteins by chemically adding γ phosphate groups and are known to alter the function of target proteins. Protein kinases remove a phosphate group from ATP and covalently attach it to an amino acid that has a free hydroxyl group [1, 2]. This usually results in a functional change of the target protein (substrate), by changing enzyme activity, cellular location or association with other proteins. Since the discovery of the first kinase in early 50’s, 518 kinases have been discovered till date [3]. This corresponds to around 2% of the human genome and tells the extraordinary importance of this family in regulating the biological events. Up to 30% of proteins can be modified by kinase activity, which are known to regulate most of the cellular pathways, especially those involved in signal transduction, cycle progression, growth, differentiation and apoptosis. Protein kinases are turned ‘ON or OFF’ by Phosphorylation, by binding of activator proteins or inhibitory proteins, respectively, or small molecules, or by controlling their location in the cell relative to their substrates. Among them cyclin-dependent kinases (CDKs), Aurora, Ptk-1, Bub1, BubR1, Mps1, Nek kinases and checkpoint kinases, Chk1 and Chk2 are being looked upon as potential anticancer targets [4, 5, 5a].

PKs are classified on the basis of specificity to the amino acid they phosphoraylate and sequences in the catalytic domain as Tyrosine kinases (PTK), Serine-threonine Kinase and Atypical kinases. Kinases have two structurally and functionally distinct lobes N-lobe and C-lobe, that show unique contributions in both catalysis and regulation of the enzyme activity ( fig. 1). The N and C lobes are connected to each other with a flexible C helix loop forming a fold over the nucleotide and position the γ-phosphate of ATP for catalysis. It is unique, dynamic regulatory element in the protein kinase molecule. Both PTK and Serine threonine kinase have a common catalytic core, which consists of a glycine-rich N-terminal ATP binding pocket and a central conserved core consisting of aspartic acid residues which play a vital role in catalytic activity [3].

Fig. 1: Structure of the conserved protein kinase core [3] A. Protein kinase with a characteristic bilobal fold. The N-lobe with five β strands (steal colored) and αC-helix, mostly helical (colored red). The ATP molecule is bound to a deep cleft between the lobes. Catalytic loops (yellow coloured). B. N-lobe structure with three conserved glycines is shown as red spheres. C. C lobe, with catalytic and regulatory machinery bound to the rigid helical core.
CDKs are a type of second messenger-dependent serine threonine kinases. They phosphorylate serine or threonine followed by proline. They are active only in the presence of cyclins which form the regulatory subunit. Timely destruction of a particular CDK after their function is over is done through 26S proteasome by ubiquination. Numerous Ubiquitin molecules covalently attach to the protein that is to be destructed [6]. When the level of cyclins rises they form a heterodimeric complex with CDKs forming a catalytic complex. These complexes also get activated by phosphorylation of specific sites on CDK thereby forming CK (CDK-activating Kinases). These activated proteins drive the cell cycle from one phase to another e.g., from G1 to S or G2 to M phase. The CDK subfamily comprises of thirteen members (CDK1 to CDK13) which have a direct impact on transcription, translation and neuronal functions [7].

Overexpression of CDK2 has been seen in colorectal and lung cancer whereas CDK4 is shown to be overexpressed in sarcomas [8]. Overexpressed CDK4 leads to inactivation of retinoblastoma protein pRb leading to carcinogenesis. There are about 1-4 CDK2 inhibitors under clinical trials at various stages (fig. 2) with very promising IC50 values [7].

IH-Pyrazolo [3,4-d] pyrimidines derivatives have gained considerable attention due to their reported anticancer and antileukemic activities [9-13,14]. The biological effect is attributed to their cyclic-dependent kinase inhibitor [15-17] and tyrosine kinase inhibitory activity [18-19]. The scaffold has also shown potent xanthine oxidase inhibitors and adenosine receptor antagonistic activity [20]. The condensed heterocyclic derivatives have also shown radioprotective activities [10]. Extensive studies have shown pyrazolo[3,4-d]pyrimidines to have anticancer potential due to multiple target inhibition of epidermal growth factor (EGFR) inhibitors, mammalian target of rapamycin (mTOR) inhibitors [21], Src or dual Src/Abl inhibitors [22], cyclin-dependent kinase (CDK) inhibitors, glycogen synthase kinase-3b (GSK-3b) inhibitors [23-25], xanthine oxidase inhibitors through modulating oxygen stress in cancer cells [26]. Some pyrazolo [1.5α]-1,3,5 triazines, as purine biosofteres, have also shown potent CDK2 inhibitory activity [27]. The inhibitors have been reported to be placed in the ATP pocket of CDK2 receptor. They interact with hydrophobic interaction and make H-bonds with Kinase, especially with a backbone of Glu 81 and Leu83. These compounds have shown promising activity IC50 of 7 μmols [28]. Substituted Indoles [29] and benzopyrans (chromenes) [30] have shown good anticancer activity. Series of 1,3,4 Oxadiazoles substituted with benzimidazole and other aromatic groups have shown promising antitumour activity [31]. Substituted pyrazoles, as well as pyrimidines, have reported promising anticancer potential [32, 33]. A condensed moiety of the two heterocycles, pyrazolopyrimidine, which is also bioisosteric to purine has been selected for exploring their potential as anticancer agents as specific CDK2 inhibitors. This pyrazolopyrimidine series was taken up for molecular docking study using AutoDock 4 program and further for planned for synthesis and activity assay. This docking study gave us new insights into the CDK2 and helped us to understand the interactions of our compounds with various amino acids in the active site of the receptor.

MATERIALS AND METHODS

All computational studies were carried out using AutoDock 4.0 with MGL tools 1.5.6 on a 2.0 GHz Intel, core i5 processor with 2GB RAM and 2 TB hard disk with Windows 7 operating system. Ligand 2D structures were drawn using ChemDraw Ultra 7.0 (Chem Office 2010). Chem3D Ultra 7.0 was used to convert 2D structure into 3D. Minimized energy to a minimum RMS gradient of 0.100 was set in each iteration. All structures were saved as pdb file format for input into AutoDockTools (ADT) version 1.5.6. All the ligand structures were then saved in PDBQT files format, for input into AutoDock version 1.5.6.

For the molecular docking study, protein PDB 1D was 3WBL of CDK2 was obtained from the RCSB protein data bank. For protein preparation, the co-crystallized ligand PDI, (N7-(4 ethoxyphenyl)-6-methyl-N5-(piperidin-3yl)pyrazolo[3-5a] pyrimidine-5,7-diamine) on CDK2 was removed, all hydrogen atoms were added, lower occupancy residue structures were deleted, and any incomplete side chains were replaced using the ADT version 1.5.6. Further ADT was used to remove crystal water, Gasteiger charges were added to each atom and non-polar hydrogen atoms were merged with the protein structure. The distance between donor and acceptor atoms that formed hydrogen bond was defined as 1.9 Å with a tolerance of 0.5 Å and the acceptor-hydrogen-donor angle was not less than 120°. The structures were then saved in PDBQT file format, for input into AutoDock version 1.5.6.

The compounds were docked by creating a grid box with a dimension of 40 × 40 × 40 Å and centred on 28.946, 64.147, 8.5596 around the binding site of ligand PDI on CDK2 receptor using AutoDock tools. The centre of the box was set at PDI and grid energy calculations were carried out. For AutoDock docking calculation, default parameters were used and 10 docked conformations were generated for each compound.

In order to verify the reproducibility of the docking calculations, the bound ligand (PDI) was extracted from the complexes and redocked with above docking protocol. Docking protocol was validated by docking the PDI, which reproduced the PDI bioactive conformer with RMSD less than 1 (fig. 3). The outputs were exported to discovery studio for visual inspection of the binding modes and interactions of the compounds with amino acid residues in the active site.

![Fig. 2: Some CDK2 inhibitors under clinical trials](image-url)
RESULTS AND DISCUSSION

Serine threonine kinase (CDK2), a clinically validated target for the treatment of various lymphomas and sarcomas has received considerable interest from the scientist in design and development of newer anticancer drugs. Molecular docking studies were performed on 17 structurally diverse, proposed, substituted 1H-Pyrazolo[3,4-d] pyrimidines on CDK-2 (PDB ID- 3WBL) using AutoDock 4 model. Docking results show notable interactions of ligands and receptor that include H bond interactions with backbone carbonyl group of Asp A: 86 and Leu A: 83, hydrophobic and pi alkyl interactions with amino acids Leu A: 134, Val A: 18 and Ala A: 31, Van der Waals interactions with Gly A: 11, amide and pi staked amide interactions with ILE A: 10. Wherever halogens are present (Fluorine) they are seen to interact with GlyA: 11. Some unfavourable interactions are also seen with Val 18 (with fluorne) of IIb, Leu 83 (with pyrimidine ring). Leu 133(with phenyl ring) of IIIe, and Asp A: 145 (with Pyrazolopyrimidines) scaffold of IIe. Asp A: 145 shows both favourable and unfavourable interactions with OCH3 of Ic and Id. The binding free energy of Ligand PDY with the receptor is -7.93 Kcal, which is comparable to binding free energy between -6.85 Kcal/mol (Ii) to -8.64 Kcal/mol(Ii) of the proposed series. Compounds Ig, IIb, IIIa, IIIb and IIIId, show lower binding free energy as well as better Ki than ligand PDY and can be looked upon for further development.

Similar docking studies are carried out on CDK2 receptors on benzopyrazole derivatives [34] show binding interactions with Lys 33 and Leu 83, docking studies on 1H pyrazole derivatives [35] have shown H-bond interactions with Ile 10, Lys 89 and Asp145. Studies carried out on oxindoles, indole and purine derivatives show H-bond interactions with the carbonyl group of Gln 131, Leu 83, Lys 89 and Asp86 with very good predicted and actual IC50 [36]. A potent CDK2 inhibitor developed with cycloalkyl amino thiazolyl scaffold (BMS-387032) has shown 2 H bonds with LeuB3 in the active site. This compound has proven to be an effective and selective inhibitor and has entered phase I human clinical trials [37]. The conserved interactions that our compounds show suggest that these proposed set of compounds can be developed as potent CDK2 inhibitors.

Various docked conformations and active Site interactions with CDK-2 receptor have been shown in fig. 3. Further details of important amino acid interactions with CDK2 receptor with Predicted Ki values are shown in table 1 and key interactions of compounds at the binding site with bond lengths in Å are shown in table 2.
Table 1: Important amino acid interactions with CDK2 receptor with predicted Ki values

| S. No. | Name | Structure | Ki (μmol) | Binding free energy Kcal/mol | Important interactions-interacting residues |
|--------|------|-----------|-----------|-------------------------------|---------------------------------------------|
| 1.     | PDY  | ![PDY](image) | 1.54      | -7.93                         | H-bonds-ASP A: 86, LEU A: 83, Alkyl-LEU A: 34, ALA A: 31, VAL A: 18, C-H-HIS A: 84 |
| 2.     | Ia   | ![Ia](image) | 3.42      | -7.46                         | H-bonds-HIS A: 83, LEU A: 298, LEU A: 83, pi-Alkyl-LEU A: 134, ALA A: 31, VAL A: 18, pi-anion-ASP A: 86, C-H-GLN A: 131, Van der waals-GLY A: 11 |
| 3.     | Ib   | ![Ib](image) | 6.53      | -7.07                         | H-bonds-LYS A: 33, GLN A: 131, ASN A: 132, ASP A: 145, Alkyl and pi-Alkyl-ALA A: 144, VAL A: 18, ALA A: 31, PHE A: 82, LEU A: 134, C-H-ASP A: 86 |
| 4.     | Ic   | ![Ic](image) | 1.4       | -7.99                         | H-bonds-LEU A: 83, Alkyl and pi-Alkyl-ILE A: 10, LEU A: 134, VAL A: 18, ALA A: 31, LYS A: 33, ALA A: 144, C-H-ASP A: 145, pi-Sigma-PHE A: 89, Unfavourable bumps-ASP A: 145 |
| 5.     | Id   | ![Id](image) | 3.14      | -7.51                         | H-bonds-LEU A: 83, Alkyl and pi-Alkyl-LEU A: 134, ALA A: 31, LYS A: 33, ALA A: 144, VAL A: 18, C-H-ASP A: 145, C-H-ASP A: 145, pi-Sigma-PHE A: 89, Unfavourable bumps-ASP A: 145 |

Fig. 3: Docked conformations and active site interactions with CDK-2 receptor
| S. No. | Name | Structure | Ki (μmol) | Binding free energy Kcal/mol | Important interactions-interacting residues |
|-------|------|-----------|-----------|-----------------------------|---------------------------------------------|
| 8.    | Ig   |           | 2.3       | -7.69                       | H-bonds-Asp A: 86, Lys A: 33, Leu A: 83; pi alkyl-Val A: 18, Leu A: 134, Ala A: 144; pi Sigma IA: 10; Unfavourable bumps - Leu A: 10 |
| 9.    | Ih   |           | 3.13      | -7.51                       | H-bonds - Lys A: 33, Leu A: 83, Asp A: 86; pi alkyl-Val A: 18, Ala A: 144; pi Sigma Ile A: 10, Gly A11 |
| 10.   | Ij   |           | 9.55      | -6.85                       | H-bonds - Asp A: 86 and Leu A: 83; Amide pi stacked - Ile A: 10; pi alkyl-Val A: 18, Leu A: 134, Ala A: 144; C-H Ala A: 31; Van der waals - Gly A: 11 |
| 11.   | Ila  |           | 6.9       | -7.04                       | H-bonds - Leu A: 83 and Lys A: 89; Alkyl and pi-alkyl-Ile A: 10, Val A: 18, Ala A: 31 and Ala A: 144; pi-alkyl stacked - Phe A: 82; Unfavourable bumps - Leu A: 83 |
| 12.   | Ilb  |           | 1.71      | -7.87                       | H-bonds - Lys A: 33, Glu A: 131, and Asp A: 145; Alkyl and pi-alkyl-Ile A: 10, Val A: 18, Phe A: 82, Ala A: 31, Ala A: 144; pi Sigma-Phe A: 80 |
| 13.   | Iic  |           | 3.36      | -7.47                       | H-bond - Asn A: 132; Alkyl and pi-alkyl-Ile A: 10, Ala A: 31, Phe A: 82, Val A: 18 and Leu A: 134; pi Cationic Lys A: 33; Unfavourable bumps - Asp A: 145 |
| 14.   | Illa |           | 2.89      | -7.56                       | H-bond - Ile A: 10, Leu A: 83; Alkyl and pi-alkyl-Val A: 18, Ala A: 31, Lys A: 33, Phe A: 80, Leu A: 134, Ala A: 144; pi-alkyl stacked - Phe A: 82, Halogen-GLN A: 131, Asp A: 145 |
| 15.   | Illb |           | 0.460     | -8.64                       | H-bond - Lys A: 33, Leu A: 83 and Asp A: 86; Amide pi stacked - Ile A: 10; pi alkyl and alkyl-Val A: 18, Halogen-GLY A: 11; Unfavourable bumps - VAL A: 18 |

| S. No. | Name | Structure | Ki (μmol) | Binding free energy Kcal/mol | Important interactions-interacting residues |
|-------|------|-----------|-----------|-----------------------------|---------------------------------------------|
| 16.   | Ilc  |           | 2.36      | -7.68                       | H-bond - GLN A: 85, Asp A: 86, Leu 298; Amide pi stacked - Ile A: 10; pi alkyl and alkyl-Val A: 18, Ala A: 31 and Leu A: 134; C-H bond - His A: 84; Van der waals - GLY A: 11 |
| 17.   | Ild  |           | 1.33      | -8.01                       | H-bond - Leu A: 83; pi alkyl and alkyl-Ile A: 10, Val A: 18, Ala A: 31, Lys A: 33, Phe A: 80, Leu A: 134 and Ala A: 144; Halogen-GLN A: 131 |
| 18.   | Illc |           | 10.12     | -6.89                       | H-bond - Leu 298; Amide pi stacked - GLN A: 85; pi alkyl and alkyl-Val A: 18, Ala A: 31, Phe A: 82, Ile A: 10, Ala A: 144 and Leu A: 134; Van der waals - Asp A: 86; Unfavourable bump - Leu A: 83 and Leu 133; pi cationic Lys A: 33; Halogen - His A: 84 |

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CONCLUSION
This work has shown interesting results in terms of interactions with various amino acids in the active site of the CDK2 receptor, binding free energy and predicted Ki values. This series can be further taken up for the development of selective and effective CDK2 inhibitors as anticancer agents.

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CONFLICT OF INTERESTS
Declared none

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