Identification of Phytochemicals Targeting c-Met Kinase Domain using Consensus Docking and Molecular Dynamics Simulation Studies

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Abstract c-Met receptor tyrosine kinase is a proto-oncogene whose aberrant activation is attributed to a lower rate of survival in most cancers. Natural product-derived inhibitors known as “fourth generation inhibitors” constitute more than 60% of anticancer drugs. Furthermore, consensus docking approach has recently been introduced to augment docking accuracy and reduce false positives during a virtual screening. In order to obtain novel small-molecule Met inhibitors, consensus docking approach was performed using Autodock Vina and Autodock 4.2 to virtual screen Naturally Occurring Plant-based Anti-cancer Compound–Activity–Target database against active and inactive conformation of c-Met kinase domain structure. Two hit molecules that were in line with drug-likeness criteria, desired docking score, and binding pose were subjected to molecular dynamics simulations to elucidate intermolecular contacts in protein–ligand complexes. Analysis of molecular dynamics simulations and molecular mechanics Poisson–Boltzmann surface area studies showed that ZINC08234189 is a plausible inhibitor for the active state of c-Met, whereas ZINC03871891 may be more effective toward active c-Met kinase domain compared to the inactive form due to higher binding energy. Our analysis showed that both the hit molecules formed hydrogen bonds with key residues of the hinge region (P1158, M1160) in the active form, which is a hallmark of kinase domain inhibitors. Considering the pivotal role of HGF/c-Met signaling in carcinogenesis, our results propose ZINC08234189 and ZINC03871891 as the therapeutic options to surmount Met-dependent cancers.

Keywords Consensus docking · c-Met inhibitor · Molecular dynamics simulation · Binding free energy · Natural products

Introduction

During recent years, a paradigm shift has occurred in cancer research. Identification of the molecular pathways involved in cancer initiation and progression will aid scientists to develop new therapies according to the molecular profile of patients [1]. Receptor tyrosine kinases (RTKs) are pharmacologically attractive targets for this purpose because they play a critical role in tumor formation. They regulate a variety of signal-transduction pathways within a cell [2], of which c-Met and its ligand, hepatocyte growth factor (HGF), mediate epithelial tissue remodeling, migration, morphogenesis, cell growth, differentiation, and angiogenesis [2, 3]. Deregulated activation of c-Met-signaling pathway through multiple mechanisms like activation, mutation, gene amplification, and heterodimerization has been reported to correlate with high tumor grade and lower rate of survival in most cancers [3, 4]. Moreover, in 2011, c-Met...
was found as a marker of pancreatic cancer stem cell, which made it a suitable therapeutic target for cancer treatment [5].

The Met receptor that belongs to the family of RTK is a heterodimer of an extracellular α-chain and a transmembrane β-chain. Upon binding of HGF, receptor dimerization and autophosphorylation of tyrosine residues of kinase domain in β sub-unit occurs, leading to recruitment of multiple effector proteins and promotes the activation of MAPK-signaling, PI3K/Akt-signaling, STAT-signaling, FAK-signaling pathways etc. Over the last decade, small-molecule inhibitors targeting ATP-binding site in kinase domain have been developed and tested in clinical trials [1, 6]. Two distinct binding patterns have been proposed for these ATP-competitive inhibitors: type I compounds (e.g., PF-023410661) bind to activated DFG-in conformation and interact with Met 1160 in the hinge region and Tyr 1230 in the activation loop. Although they are specific, they display limited activity toward Tyr 1230 mutation, while type II compounds (e.g., BMS-777607) are less selective and bind in the inactive DFG-out conformation formed by conformational changes of the Asp1222 side chain [2, 3, 7].

As evidenced by scientific literature, there is a great interest in the screening of natural products for drug development. It has been reported that natural products have great diversity and novel chemical structure which render them useful as new inhibitors. Accordingly, inhibitors derived from natural products have sometimes been called “fourth generation inhibitors” [8]. Actually, natural compounds and their derivatives constitute more than 60% of anticancer drugs [9]. In addition, phytochemicals have been used increasingly for many reasons including low cost, fewer adverse effects, and well-tolerated in the body. Thus, phytochemicals have been discovered as promising compounds for prevention, treatment, or reversal of drug resistance in cancer [10]. Moreover, previous investigations have revealed some flavonoids modulated HGF/c-Met axis in cancer cells; but the intermolecular contacts are unknown [11, 12].

Nowadays, molecular docking is a well-accepted computational method in structure-based drug design for identifying novel hit molecules, which employs a scoring function to evaluate ligand binding orientation and affinity [13–15]. In conventional molecular docking, if there is a difference in pose prediction identifying the correct pose will be impossible, while agreement in docked pose will increase accuracy and reduce false positive notably during a virtual screening. This is the objective of consensus docking, which enhances ligand pose prediction by more than one scoring algorithm [16]. Houston et al. reported that a significant improvement in pose prediction accuracy was achieved using Autodock and Vina in 82% of cases. Interestingly, this percentage increased up to 92% when three docking programs; Autodock, Vina, and Dock 6.5, were used simultaneously [16]. It is worthy of note that a Lamarckian Genetic Algorithm and an Amber-force-field-based scoring function have been implemented in Autodock and Vina, respectively [16, 17]. Berry et al. carried out a multi-step Autodock Vina-Glide-GOLD consensus docking procedure to identify novel inhibitors against human coronavirus validated by MD simulation [18]. However, a recent study by Tuccinardi et al focused on the reliability of the consensus docking results in binding pose prediction using ten docking procedures [19].

In this study, we used consensus docking procedure to screen Naturally Occurring Plant-based Anti-cancer Compound–Activity–Target (NPACT) database. For this purpose, information regarding NPACT entries were obtained from the ZINC database. Following consensus docking, MD simulation and molecular mechanics Poisson–Boltzmann surface area (MM-PBSA) binding energy calculation were performed to elucidate the binding mode of the hit compounds as novel c-Met kinase inhibitors.

Materials and Methods

Selection of Data Set

The three-dimensional, ready-to-dock format of 1574 ligands available in NPACT database was retrieved from the ZINC database (http://zinc.docking.org/) on January 18, 2015. ZINC has emerged as a widely-used database of purchasable compounds suitable for virtual screening [20]. Using physicochemical properties, 738 ligands were selected, which conformed to Lipinski’s rule of five [21] and Veber’s rules for drug-like molecules and oral bioavailability [22].

Structure Preparation

The crystal structures for active conformation of the c-Met kinase domain (PDB ID: 2RFS) [23] and inactive conformation (PDB ID: 1R0P) [24, 25] were retrieved from the Protein Data Bank (http://www.rcsb.org/pdb/home/home.do) and prepared for docking calculation by eliminating hetero atoms and water molecules. Thereafter, missing residues in protein structures were modified using MOD-ELLER v.9.11 [26] based on the original FASTA sequence. For the representative structures of c-Met, the Autodock Tools (ADT) 1.5.7 was used to merge non-polar hydrogens to carbon atoms and compute Gasteiger charges. Rotatable bonds and Gasteiger charges were also assigned by the prepare_ligand4.py utility of ADT for the selected ligands.
Consensus Docking

Autodock Vina 1.1.2 was used for protein–ligand docking against 2RFS and 1R0P structures with the default parameters [27]. The 500 top ranked ligands were further docked using Autodock 4.2.6 [28]. A grid box of 126 x 126 x 126 points with 0.153 Å spacing was defined by centering on the ligand in the active site. When running a docking using the Lamarckian GA, there were a number of parameters to set and, except the number of Lamarckian genetic algorithm runs was set to 100; other parameters were considered as their default values. In all docking studies, residues F1089, V1092, K1110, Y1159, M1160, M1211, D1222, M1229, and Y1230 in the c-Met ATP-binding site were allowed to be flexible. The lowest energy conformation in the Autodock Vina result and the largest cluster of Autodock 4.2 were selected as the resultant docked structure. The root mean square deviation (RMSD) calculation was used to assess the docking poses of Autodock Vina and Autodock 4.2 as described by Houston et al. [16]. Consequently, poses with RMSD more than 2.0 Å were rejected. To generate a molecular representation of each protein-ligand complex, PyMOL Molecular Graphics System v. 1.7.2 was used. ZINCPharmer (http://zincpharmer.csb.pitt.edu/) was also used to identify the pharmacophore features of the hit molecules [29].

Molecular Dynamics (MD) Simulation

Gromacs 4.5.3 [30] was used to perform MD simulation with the Amber03 force field parameters. To perform MD simulations, the best Autodock Vina result was considered as the initial conformation. Before the simulation, electrostatic potential (ESP) charges for drug candidates were computed employing Gaussian03 (B3LYP/6-31G* to optimize the geometry and #P HF/6-31G* SCF = Tight Pop = MK 10p (6/33 = 2) to calculate ESP); then, the restrained electrostatic potential (RESP) method was employed to assign atomic charges. To obtain ligand topology files based on generalized Amber force field, antechamber was used. A distance of 1 nm was set between a cubic box of TIP3P water molecules and the solute. Steepest descent algorithm was applied until the maximum force of the whole system was below 1000 kJ/mol. During 500 ps of NVT and NPT ensembles, all complexes were simulated to equilibrate at 300 K temperature and 1 bar pressure, respectively. Final simulation runs were carried out for 20 ns with a time step of 2 fs under V-rescale temperature and Parrinello–Rahman pressure coupling algorithms. The particle–mesh Ewald (PME) algorithm was utilized to calculate long-range electrostatics, while Linear Constraint Solver (LINCSC) method was used to constrain all covalent bond lengths. A 1.2 nm cutoff was set for van der Waals interactions. VMD 1.9.2 [31] and PyMOL Molecular Graphics System v. 1.7.2 were used to visualize trajectories and generate images, respectively.

MM-PBSA Binding Energy Calculation

Binding-free energy calculations based on MM-PBSA method were estimated using the g_mmpbsa tool [32]. Snapshots saved at every 10 ps from the last 5 ns of each MD trajectory were considered for calculating each energy term. The solute, solvent, and vacuum dielectric constants were set at 2, 80, and 1, respectively, for polar solvation calculation. The solvent-accessible surface area (SASA) model was also used to compute non-polar solvation energy.

Results and Discussion

Database Screening using Consensus Docking

A schematic procedure of consensus docking is shown in Fig. 1. A compilation of 1574 plant-derived compounds available in ZINC database, a currently widely-used database for virtual screening, was filtered according to Lipinski and Veber rules. These physico-chemical criteria can be employed to select desirable compounds having membrane permeability as well as to be easily absorbed in the human body. The processed ligand set was then docked through consensus docking method [16]. As mentioned earlier, this method is reasonable particularly when funds for purchasing compounds are limited. Calculation of RMSD between binding modes predicted by Autodock 4.2 and Vina for each complex produced a small set of 98 molecules. Of these 98 ligands, two hits were in line with good-docking score and estimated IC50, and the presence of essential interactions with key residues in the ATP-binding site were considered for further evaluation.

The 2D structures of two hit compounds alongside their molecular weight are shown in Table 1. ZINC08234189, a steroid lactone with a molecular weight of 470.61 g/mol, chemically consists of the 4-hydroxy-5,6-epoxy-2-en-1-one moiety and unsaturated lactone side chain which play crucial roles in biological activity [33].

With regard to ZINC03871891, it is an ortho-diphenolic diterpene which has 330.42 g/mol molecular weight. The structure of ZINC03871891 has an abietane carbon skeleton with a lactone moiety oriented across the central ring [34].

Docked Structures of the Hit Compounds

In order to reject false positives, it is crucial to evaluate the binding mode and affinity of the preliminary hit
compounds. Table 2 shows the docking results with associated interaction profile in details. Interestingly, ZINC08234189 exhibited higher affinity in comparison to ZINC03871891 with the inactive conformation of c-Met as evidenced by Autodock binding energy of $-12.91$ kcal/mol. Analysis of its interaction pattern showed that the phenolic hydroxyl group of ZINC08234189 engaged in two hydrogen bonds with the carboxyl group of P1158 and the amino group of M1160 in the hinge region while hydrophobic contacts were present with V1092 and M1211 in the central hydrophobic region. Other residues like I1084, A1108, L1157, and F1089 in the two hydrophobic subpockets, and M1229 and Y1230 in the activation loop are involved in the hydrophobic interactions (Fig. 2a).

In the case of ZINC03871891 bound to the active conformation, it did not form hydrogen bond with any residues in the hinge region, a typical characteristic of all kinase inhibitors targeting the ATP-binding site [25, 35]. Instead, hydrophobic interactions were found with M1160, suggesting a lower binding affinity with an IC$_{50}$ of 18.76 nM. Alternatively, residue V1092, L1140, L1157, and M1211 comprising the hydrophobic subpockets as well as D1222 (activation loop) are involved in hydrophobic interactions. The side chain of N1209 along with the carboxyl group of R1208 in the catalytic loop also formed a bidentate H-bond with this potent inhibitor as illustrated in Fig. 2b.

Furthermore, Fig. 3 shows the pharmacophoric features of the hit molecules. It revealed that O4 on compound ZINC08234189 had hydrogen donor (HD) and acceptor (HA) features, leading to hydrogen bonds with P1158 (O) and M1160 (N), which resulted in a strong interaction, while O6 on hydroxyl group had HD property. In addition, the scaffold of ZINC08234189 had five hydrophobic features. With regard to ZINC03871891, its backbone had four hydrophobic and one aromatic features, whereas O4 had HA property that tended to approach Y1230. Besides, O1 and O2 with HD and HA features formed hydrogen bonds with HD of R1208 (O) and N1209 (OD1) in the active site.

**MD Simulations**

Although we selected two complexes according to consensus docking: 1R0P bound with ZINC08234189 and 2RFS bound with ZINC03871891 to compare the stability and binding affinity of the aforementioned compounds computationally, the apo form of c-Met and all protein-ligand complexes were subjected to MD simulations and MM/PBSA binding energy calculation method.

Given that Autodock Vina is believed to be superior in ligand pose prediction than Autodock 4.2 [36, 37], docking results of Vina having lowest binding energy were chosen for MD simulation. It should be noted that docking simulation provides a single snapshot of interactions between ligand and protein both in rigid and semi-flexible docking. Thus, in order to gain a deeper insight into the effects of protein structural changes and flexibility on the complex interaction profile, a 20-ns MD simulation was performed. The C$\alpha$ RMSD in comparison with the initial structure indicated that 1R0P in the apo state experienced continuous fluctuation rising to 0.32 nm at the end of the simulation period. Upon binding of ZINC08234189 at the active site, the average values of conformational changes increased from 0.2 to 0.28 nm, while these changes were negligible for ZINC03871891 (Fig. 4a). In consideration of 2RFS without any ligand, the RMSD value increased to 0.26 nm.

Fig. 1 Flowchart of the consensus docking procedure followed to choose hit compounds with c-Met kinase inhibitory activity

Fig. 2a: Interaction pattern of ZINC08234189 in the active conformation of c-Met kinase.

Fig. 2b: Interaction pattern of ZINC03871891 in the active conformation of c-Met kinase.

Fig. 3: Pharmacophoric features of hit compounds.

Fig. 4a: RMSD comparison of 1R0P conformation with and without ligand.

Fig. 4b: RMSD comparison of 2RFS conformation with and without ligand.
for the first 4 ns. Thereafter, the receptor reached a plateau for the last 16 ns. Binding stability of 2RFS-ZINC03871891 as a function of time was also analyzed and it was shown that the system converged well with the range of 0.15–0.25 nm compared to ZINC08234189 (Fig. 4b). However, it is obvious that the RMSD value of 1R0P-ZINC08234189 was smaller than 2RFS-ZINC08234189. It was observed that around 3 and 5 ns RMSD value of ZINC08234189 underwent noticeable fluctuation due to the lactone ring rotation for 1R0P and 2RFS, respectively, and remained stable till the end (Fig. 4c, d). However, ZINC03871891 achieved relatively stable coordination throughout the entire simulation process. The same results were also observed after the initial and MD-averaged structures were superimposed (Fig. 5).

To investigate the residual mobility in ligand-bound and unbound forms, root mean square fluctuations (RMSF) of individual residues were also monitored in each case. As shown in Fig. 6, binding of hit molecules did not induce any remarkable changes in flexibility of ATP-binding site residues relative to the protein in the apo state. In particular, these fluctuations were minimal in any state of 2RFS. Thereafter, the protein structure compactness was analyzed using the g_gyrate utility of the Gromacs 4.5.3 software (Fig. 7, Table 3). It was shown that in its apo state, the protein was stable with no obvious structural expansion and

| Table 1 Chemical structures of the two final hit molecules |
| ZINC ID | Chemical formula | Molecular weight | Chemical structure |
|---------|-----------------|------------------|-------------------|
| 1       | ZINC08234189    | C_{28}H_{38}O_{6} | 470.61            |
| 2       | ZINC03871891    | C_{20}H_{26}O_{4} | 330.42            |

| Table 2 Docking scores and interaction profile of two final hit molecules with the inactive (1R0P) and active (2RFS) conformation of c-Met by Autodock |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| ZINC ID         | Docking energy (kcal/mol) | H-Bonds        | Hydrophobic interactions |
|                 | Autodock Vina   | Autodock 4.2   | Estimated IC_{50} (nM) |                   |
| 1R0P ZINC08234189 | −11.7          | −12.91         | 0.345             | P1158, M1160, I1084, R1086, F1089, V1092, A1108, L1140, L1157, Y1159, M1211, D1228, M1229, Y1230, D1231, K1232 |
| 2RFS ZINC03871891 | −10.4          | −10.54         | 18.76             | R1208, N1209, V1092, L1140, L1157, M1160, D1164, M1211, A1221, D1222, Y1230 |

for the first 4 ns. Thereafter, the receptor reached a plateau for the last 16 ns. Binding stability of 2RFS-ZINC03871891 as a function of time was also analyzed and it was shown that the system converged well with the range of 0.15–0.25 nm compared to ZINC08234189 (Fig. 4b). However, it is obvious that the RMSD value of 1R0P-ZINC08234189 was smaller than 2RFS-ZINC08234189. It was observed that around 3 and 5 ns RMSD value of ZINC08234189 underwent noticeable fluctuation due to the lactone ring rotation for 1R0P and 2RFS, respectively, and remained stable till the end (Fig. 4c, d). However, ZINC03871891 achieved relatively stable coordination throughout the entire simulation process. The same results were also observed after the initial and MD-averaged structures were superimposed (Fig. 5).

To investigate the residual mobility in ligand-bound and unbound forms, root mean square fluctuations (RMSF) of individual residues were also monitored in each case. As shown in Fig. 6, binding of hit molecules did not induce any remarkable changes in flexibility of ATP-binding site residues relative to the protein in the apo state. In particular, these fluctuations were minimal in any state of 2RFS. Thereafter, the protein structure compactness was analyzed using the g_gyrate utility of the Gromacs 4.5.3 software (Fig. 7, Table 3). It was shown that in its apo state, the protein was stable with no obvious structural expansion and
Fig. 2 Binding poses of the two hit molecules: a 1R0P in complex with ZINC08234189, and b 2RFS in complex with ZINC03871891. The N and C lobes of tyrosine kinase domain are shown in cyan and green, respectively. The predicted binding mode using Autodock 4.2 and Vina are displayed in blue and yellow stick representation, respectively. Key residues involved in interaction are represented in the stick format. The length of Hydrogen bonds colored as yellow dotted lines is indicated in Angstrom (color figure online).
contraction, whereas ZINC08234189 led to lower Rg value especially in complex with 1R0P within the range of 2.02–2.10 nm. In addition, an obvious expansion has occurred for ZINC03871891 bound to 2RFS after 6 ns.

Table 3 presents information regarding hydrogen bonds formed throughout the trajectory for all protein-ligand complexes. The percentage of H-bonds presence during the last 5 ns of MD simulation was measured to examine their
Fig. 5 Superimposed structures of c-Met inhibitors using Autodock Vina (green with a blue ligand) and MD-averaged structure of the last 5 ns (cyan with a yellow ligand) (color figure online).

Fig. 6 Root mean square fluctuations (RMSF) analysis of c-Met kinase domain residues in the presence and absence of hit molecules over 20 ns MD simulation period. a 1R0P, b 2RFS

Fig. 7 Radius of gyration of c-Met kinase domain in the presence and absence of hit molecules over 20 ns MD simulation period. a 1R0P, b 2RFS
stability, though only contacts populated with over 40% of the simulation time are listed.

As summarized in Table 3, ZINC08234189 formed hydrogen bonds with e-amino group of K1110 (79.12% occupancy) in 1R0P (inactive form), while ZINC08234189-2RFS complex formed two hydrogen bonds with main chain nitrogen and oxygen atoms of M1160 and P1158 in the hinge region, respectively, being the main determinant for the overall stability of the complex. For ZINC03871891-1R0P, two hydrogen bonds with the main chain oxygen atom of R1227 were detected with occupancy of about 95 and 65%, while only an amino group of M1160 participated in H-bond interaction between ZINC03871891 and 2RFS.

### Binding-Free Energy Analysis by MM-PBSA

Recent studies have reported MM/PBSA method as a simulation technique to estimate biomolecular interaction-free energy. Although it increases computational cost, it provides more accurate information than energy-scoring functions that have been used in docking softwares for rescoring top-ranked hits [17]. Five hundred snapshots extracted between 15–20 ns with equal intervals were selected as input for analysis as reported in Table 4. The results showed that both selected molecules had higher binding affinity of $-128.27 \pm 0.48$ and $-113.19 \pm 0.59$ kJ/mol for ZINC03871891 and ZINC08234189, respectively toward 2RFS than 1R0P, whereas the latter structure showed $-113.71 \pm 0.51$ and $-107.60 \pm 0.81$ kJ/mol for ZINC03871891 and ZINC08234189, respectively, though this difference was not considerable for ZINC08234189. In addition, ZINC03871891 has less negative binding energy than ZINC08234189 in both complexes.

Among the different energy terms that contributed to the protein-ligand binding energy, van der Waals ($E_{vdw}$), electrostatic, ($E_{ele}$), and SASA energy played a crucial role in binding energy and complex stability. Nevertheless, polar solvation energy ($G_{ps}$) has an opposite effect, causing binding energy to depend on its unfavorable positive value [38]. In this regard, van der Waals energy contributed more negative free energy than electrostatic energy in all protein–ligand complexes. Besides, in ZINC03871891-bound complexes, van der Waals interaction was predominant in total binding-free energy, whereas in ZINC08234189-bound complexes, the unfavorable contribution of polar solvation energy was significant.

### Conclusion

We used the consensus docking approach to virtually screen 1574 compounds retrieved from NPACT database against both active (2RFS) and inactive (1R0P) state of the c-Met kinase domain, yielding a selection of two hit molecules. Using a 20-ns MD simulation, the stability of each complex was evaluated. Our results showed that both protein and ligand backbone of ZINC08234189 achieved stability after 5 ns. Nevertheless, ZINC03871891 experienced stable conformation in each case during the entire simulation process. Given that hydrogen bond with residues of the hinge region (P1158, M1160) is a hallmark of kinase domain inhibitors, our analysis showed that both hit molecules formed hydrogen bonds with key residues in the active form. In summary, based on hydrogen bond analysis...
and MM-PBSA studies, we predicted that ZINC08234189 is a plausible inhibitor for the active state of c-Met. In line with our docking results, ZINC03871891 showed more effectiveness toward active c-Met kinase domain than the inactive form due to higher binding energy. The results disclosed here can be useful to overcome Met-addicted cancers.

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**Compliance with Ethical Standards**

**Conflict of Interest** The authors declare that they have no competing interests.

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