Efficacy and Toxicity Assessment of Different Clinically Used Small Molecular Tyrosine Kinase Inhibitors by Computational Molecular Docking Method

Keywords: Angiogenesis; Toxicity; Antiangiogenic drug; Small molecular antiangiogenic drug; VEGF; Tyrosine kinase; Her-2

Abstract

Along with the antibody based antiangiogenic [AAG] drugs, several small molecular antiangiogenic (sAAG) drugs are also in clinical practice for the treatment of different cancers. Though population based clinical trials have provided their efficacy; however, comparative efficacies of these drugs at molecular level are not available. The drug-target network is highly important to understand the efficacy and toxicity of drugs in terms of systems biology. In this study, multiply molecular docking between several sAAG drugs and different target receptors are carried out. Our data suggest that though gefitinib is quite potent in targeting different receptors involved in angiogenesis, but it may impose some serious side effects within physiological system. Our data also suggest that one sAAG drug have some preferential binding affinity to one receptor while have lower binding capacity for other receptor. Hence different sAAG drugs may produce different types of toxicity within the physiological system. For example, lapatinib has highest binding affinity towards NO synthetase, while erlotinib has lowest binding affinity to this enzyme. Similarly, sunitinib has highest binding affinity towards beta-2 receptor, while lapatinib has lowest affinity to this receptor. Hence this work provides the rationality of switching between different sAAG drugs during the course of treatment in the control of toxicity by one sAAG drug without discontinuation of therapy and management of toxicity by other drugs.

Introduction

Though angiogenesis is a physiological process; however almost all types of solid tumors survive by utilizing this process. Hence, development of different therapeutic strategies that can control of tumor angiogenesis becomes an important aspect in clinical oncology [1,2]. For several types of cancer, different antiangiogenic (AAG) drugs are generally used as an adjuvant therapy that is, in combination with the conventional MTD (maximal tolerable dosing strategy) chemotherapy. Such applications have shown positive results with reduced (chemotherapy related) toxicities [3]. Therapy with different AAG drugs are not only supplement the ongoing chemotherapy (with docetaxel, platinum based therapies, paclitaxel, vinorelbine and gemcitabine) but also becomes important when the conventional chemotherapies have failed or are not tolerated [4]. Generally different AAG drugs are targeted towards receptors that are involved with tyrosine kinase (TK) mediated cell proliferation [5-7].

Though different antibody based AAG drugs have developed and are in clinical use; however, development of different small molecular drugs that act as AAG becomes the first hand choice for cancer therapy in clinics [8,9]. There are several advantages of using small molecular AAG (sAAG) drug over antibody based AAG drugs. These drugs are more thermo-stable and can be stored without constant refrigeration, and do not agglutinate or precipitate. They may have multiple targets, inhibit multiple kinases and target many growth factors to inhibit angiogenesis. Reports suggest that most of the small molecular tyrosine kinase inhibitors have an immunomodulatory role [10]. Perhaps the greatest advantage of sAAG drugs in future could be used for targeted therapies by inserting in liposomes [11] or can be applied through MCT (metronomic dosing strategy, where chemotherapeutic drugs are applied with dose-dense mode i.e., very low doses but with a frequent intervals) strategy; hence can be applied orally [12-18]. There are several sAAG drugs now available for clinical use and the list is growing in number; however, most of them have a molecular weight ranges from 130 to 600.

Among different sAAG drug, Imatinib mesylate (STI 571; Gleevec) was the first discovered sAAG drug that has clinical efficacy. It inhibits the Bcr-Abl tyrosine kinase (TK), the constitutive abnormal TK created by the Philadelphia chromosome t(9;22) (q34;q11) abnormality in chronic myeloid leukemia (CML). Imatinib also inhibits the receptor tyrosine kinases for platelet derived growth factor (PDGF) and stem cell factor (SCF), called c-kit mediated cellular events (Table 1) [6,19-23]. A number of studies prove the antiangiogenic potential of imatinib. Imatinib reduces the angiogenic factors to inhibit angiogenesis. Reports suggest that most of the small molecular tyrosine kinase inhibitors have an immunomodulatory role [10]. Perhaps the greatest advantage of sAAG drugs in future could be used for targeted therapies by inserting in liposomes [11] or can be applied through MCT (metronomic dosing strategy, where chemotherapeutic drugs are applied with dose-dense mode i.e., very low doses but with a frequent intervals) strategy; hence can be applied orally [12-18]. There are several sAAG drugs now available for clinical use and the list is growing in number; however, most of them have a molecular weight ranges from 130 to 600.
lesser extent, CYP1A2, CYP2D6, CYP2C9, and CYP2C19. The main metabolism occurs in the liver mitochondria and is mediated by several isozymes of the cytochrome P450 system, including CYP3A4 and, to a lesser extent, CYP1A2, CYP2D6, CYP2C9, and CYP2C19. The main metabolite, N-demethylated piperezine derivative, is also active. The major route of elimination is in the bile and feces; only a small portion of the drug is excreted in the urine.

Gefitinib (446.90) EGFR associated TK

Daily oral dose of 250 mg (45 hrs) It is primarily metabolized by CYP3A4.

Types of cancer that are recommended for treatment

Chronic phase of CML

First-line treatment of metastatic NSCLC, maintenance treatment of NSCLC whose diseases has not advanced after four cycles of platinum-based chemotherapy, first-line treatment of metastatic pancreatic cancer in combination with gemcitabine, myeloproliferative disorder, poor tolerance to gefitinib

Erlotinib (393.43) EGFR (ErbB1) associated TK and JAK2V617F, blocks cell cycle progression in G1 phase

Daily oral dose of 150 mg It is primarily metabolized by CYP3A4.

Lapatinib (581.05) EGFR (both ErbB1 and ErbB2) as well as Akt-overexpressing cell

Once dose of 1250 mg in every two weeks in combination with capecitabine (~24 hrs) Lapatinib undergoes metabolism, predominantly by liver mitochondrial CYP450 enzyme system specially CYP3A4 and CYP3A5, with minor contributions from CYP2C19 and CYP2C8 to a variety of oxidized metabolites, to form a variety of metabolites.

Metastatic breast cancer with HER2+ used in combination with capcitabine and effective even in cases failed with taxane and trastuzumab.

Sorafenib (464.82) VEGF cell surface kinases (KIT, FLT-3, VEGFR-2, VEGFR-3, and PDGFR-B), and Raf/MEK/Erk pathway including intracellular kinases (CRAF, BRAF and mutant BRAF)

Daily oral dose of 400 mg, no dose adjustment is required on the basis of patient age, gender or body wt. (25-48 hrs) Sorafenib is metabolized primarily by Cytochrome CYP3A4 as well as glucuronidation (mediated by UGT1A9), 77% of the dose excreted in feces and 19% of the dose excreted in urine as glucuronidated metabolites. At steady state the metabolites of sorafenib are detected in plasma. Pyridine N-oxide, the main circulating metabolite of sorafenib in plasma, shows in vitro potency similar to that of sorafenib.

Recurrent, metastatic, progressive, differentiated thyroid carcinoma (DTC) refractory to radioactive iodine treatment; advanced renal cell carcinoma, hepatocellular carcinoma.

Sunitinib (398.47) Multi-targeted TKs

Oral dose of 50 mg for 4 weeks (drug: 40-60 hrs; metabolite: 80-110 hrs) Sunitinib is metabolized by the liver mitochondrial cytochrome P 450 system namely by CYP3A4 to its primary active metabolite which is also metabolized by CYP3A4 enzyme. Its primary active metabolite is also active.

Renal cell carcinoma, advanced pancreatic neuroendocrine tumors, advanced or metastatic gastrointestinal stromal tumor with disease progression or intolerance to imatinib mesylate.


Table 1: FDA approved different TK inhibitors that act as AAG drugs.

<table>
<thead>
<tr>
<th>Drug (MW)</th>
<th>Drug Target</th>
<th>Recommended Dose (half-life)</th>
<th>Metabolism</th>
<th>Types of cancer that are recommended for treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imatinib</td>
<td>TK (BCR-ABL, PDGF, c-Kit)</td>
<td>Oral dose of 400-800 mg (16 hrs)</td>
<td>Metabolism occurs in the liver mitochondria and is mediated by several isozymes of the cytochrome P450 system, including CYP3A4 and, to a lesser extent, CYP1A2, CYP2D6, CYP2C9, and CYP2C19. The main metabolite, N-demethylated piperezine derivative, is also active. The major route of elimination is in the bile and feces; only a small portion of the drug is excreted in the urine.</td>
<td>Chronic phase of CML</td>
</tr>
<tr>
<td>Gefitinib</td>
<td>EGFR associated TK</td>
<td>Daily oral dose of 250 mg (45 hrs)</td>
<td>It is primarily metabolized by CYP3A4.</td>
<td>NSCLC</td>
</tr>
<tr>
<td>Erlotinib</td>
<td>EGFR (ErbB1) associated TK and JAK2V617F, blocks cell cycle progression in G1 phase</td>
<td>Daily oral dose of 150 mg</td>
<td>It is primarily metabolized by CYP3A4.</td>
<td>First-line treatment of metastatic NSCLC, maintenance treatment of NSCLC whose diseases has not advanced after four cycles of platinum-based chemotherapy, first-line treatment of metastatic pancreatic cancer in combination with gemcitabine, myeloproliferative disorder, poor tolerance to gefitinib</td>
</tr>
<tr>
<td>Lapatinib</td>
<td>EGFR (both ErbB1 and ErbB2) as well as Akt-overexpressing cell</td>
<td>Once dose of 1250 mg in every two weeks in combination with capecitabine (~24 hrs)</td>
<td>Lapatinib undergoes metabolism, predominantly by liver mitochondrial CYP450 enzyme system specially CYP3A4 and CYP3A5, with minor contributions from CYP2C19 and CYP2C8 to a variety of oxidized metabolites, to form a variety of metabolites.</td>
<td>Metastatic breast cancer with HER2+ used in combination with capcitabine and effective even in cases failed with taxane and trastuzumab.</td>
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<td>VEGF cell surface kinases (KIT, FLT-3, VEGFR-2, VEGFR-3, and PDGFR-B), and Raf/MEK/Erk pathway including intracellular kinases (CRAF, BRAF and mutant BRAF)</td>
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<td>Recurrent, metastatic, progressive, differentiated thyroid carcinoma (DTC) refractory to radioactive iodine treatment; advanced renal cell carcinoma, hepatocellular carcinoma.</td>
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<td>Multi-targeted TKs</td>
<td>Oral dose of 50 mg for 4 weeks (drug: 40-60 hrs; metabolite: 80-110 hrs)</td>
<td>Sunitinib is metabolized by the liver mitochondrial cytochrome P 450 system namely by CYP3A4 to its primary active metabolite which is also metabolized by CYP3A4 enzyme. Its primary active metabolite is also active.</td>
<td>Renal cell carcinoma, advanced pancreatic neuroendocrine tumors, advanced or metastatic gastrointestinal stromal tumor with disease progression or intolerance to imatinib mesylate.</td>
</tr>
</tbody>
</table>

IV: Intravenous administration; std: standard; CT: chemotherapy; rst: resistant; TK: Tyrosine kinase; wt: Weight
improved survival of NSCLC patients after failure of at least one chemotherapy regimen with previously treated (with other chemotherapy) and in patients with untreated pancreatic cancer (Table 1) [6,21,32]. Erlotinib has recently been shown to be a potent inhibitor of JAK2V617F, a mutant of TK activity. JAK2V617F is found in most patients with polycythemia vera (PV) and a substantial proportion of patients with idiopathic myelofibrosis or essential thrombocythemia. The study suggests that erlotinib may be used for treatment of JAK2V617F-positive PV and other myeloproliferative disorders [33]. It is primarily metabolized by CYP3A4. It's common side effects are rash, diarrhea, dyspnea, dry skin, back pain, chest pain, conjunctivitis, mucosal inflammation, pruritus, paronychia, arthralgia, musculoskeletal pain [34].

**Lapatinib** (Tykerb) is a member of the 4-anilinoquinazoline class of kinase inhibitors. It binds to human EGFR type-1 (HER1/EGFR/ERBB1) and type-2 (HER2/ERBB2) and prevent receptor autophosphorylation upon ligand binding by TKs [35]. In combination with capecitabine, it is effective for the treatment of patients with advanced or metastatic breast cancer with overexpress HER2+ and who have received prior therapy including an anthracycline, a taxane, and trastuzumab (Table 1) [36]. At a clinically relevant concentration, lapatinib is a substrate and inhibitor of the transporter enzymes ABCG2 or BCRP (a wild type Breast Cancer Resistance Protein), P-gp (or ABCB1, ATP Binding Cassette subfamily B member 1) and also an inhibitor of OATP1B1 (Organic Anion Transporter Pump family 1B1) or Solute carrier organic anion transporter family member 1B1 (SLCO1B1) [37]. Lapatinib undergoes metabolism, predominantly by liver mitochondrial CYP450 enzyme system specially CYP3A4 and CYP3A5, with minor contributions from CYP2C19 and CYP2C8 to a variety of oxidized metabolites, to form a variety of metabolites. Side effects of lapatinib include hepatotoxicity (increase in liver enzymes and cholestatic liver damage by disrupting of hepatocyte bile salt efflux and a decreased of left ventricular ejection fraction (LVEF). 10% patients experience diarrhea, rash, nausea and fatigue when lapatinib is applied in combination with letrozole [38-41]. Lapatinib has also been associated with interstitial lung disease and pneumonitis in monotherapy or in combination with other chemotherapies [38].

**Sorafenib** (Nexavar, BAY 43-9006) is unique in targeting the Raf/Mek/Erk pathway. By inhibiting these kinases, genetic transcription involving cell proliferation and angiogenesis is inhibited. However it also inhibit a variety of kinase receptors, including VEGFR, EGFR, and PDGFR kinases; so it is recommended for different types of cancer (Table 1) [6,42,43] Sorafenib is metabolized primarily by CYP3A4 as well as glucuronidation (mediated by UGT1A9): 77% of the dose excreted in feces and 19% of the dose excreted in urine as glucuronidated metabolites. At steady state the metabolites of sorafenib are detected in plasma. Pyridine N-oxide, the main circulating metabolite of sorafenib in plasma, shows in vitro potency similar to that of sorafenib. Side effects include cardiac ischemia, infarction, hypertension, hemorrhage, hand-foot skin reaction, rash, Stevens-Johnson syndrome, and toxic epidermal necrolysis, gastrointestinal perforation, QT interval prolongation, drug induced hepatitis, hypoalbuminemia, elevated amylase, thrombocytopenia, hypocalcaemia [44].

**Sunitinib** (Sutent) is a multi-targeted receptor tyrosine kinase
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Figure 3: Docked conformation of sorafenib (in violet color) (A) and sunitinib (in blue color) (B) to Her-2 receptor (in cyan color).

Figure 4: Docked conformation of lapatinib (in magenda color) to androgen (A) and estrogen-α (B) receptor, receptor proteins are in cyan color.

(TK) (>80 kinases) inhibitor that was approved by the FDA for the treatment of different cancers. Sunitinib was evaluated for its inhibitory activity against a variety of receptor protein kinases which include PDGFRα and PDGFRβ; VEGFR1, VEGFR2 and VEGFR3; stem cell factor receptor (KIT), Fms-like TK-3 (FLT3), colony stimulating factor receptor Type 1 (CSF-1R), and the glial cell-line derived neurotrophic factor receptor (RET) [45-47]. Sunitinib is metabolized by the liver mitochondrial CYP450 system namely CYP3A4 enzyme. Its primary active metabolite is also active. As sunitinib targets many different receptors, leads to many of its side effects. In >20% cases, the most common side effects are fatigue, asthenia, diarrhea, fever, diarrhea, nausea, mucositis/stomatitis, vomiting, dyspepsia, abdominal pain, constipation, hypertension, peripheral edema, rash, hand-foot syndrome, skin discoloration, dry skin, hair color changes, altered taste, headache, back pain, arthralgia, extremity pain, cough, dyspnea, anorexia, and bleeding. The potentially serious adverse reactions include hepato-toxicity, left ventricular dysfunction, QT interval prolongation, hemorrhage, hypertension, thyroid and adrenal dysfunction.

Here attempt has been made to find out the molecular rationale of side effects and/or off-target toxicities within the physiological system by these small molecular AAG drugs.

Materials and Methods

All data used in the present work are collected from the data resources that are available in the public domain and majority of the used data and its resources are already mentioned in an early work [48]; however, the newly added data (along with its resources) are mentioned in Table 2. The protein/macromolecular 3D structural data and the 3D drug structures are downloaded from the Protein Data Bank (PDB) (http://www.rcsb.org/pdb/home/home.do) [49] and the Drug Bank (http://redpoll.pharmacy.ualberta.ca/drugbank/) database [50].

For studying the binding of drug to protein/receptor, present work is performed with docking software Autodock 3.0 (The Scripps Research Institute, www.scripps.edu) with the help AutoDockTools (ADT), that allows the user to interact with AutoDock from a Graphic User Inter-face (GUI) [51]. Obtained PDB files of macromolecules and ligands are further prepared by using the following steps as mentioned in the early works [52-53].

1. Preparing the macromolecular PDB files - as the PDB files often contained added waters, these water molecules were files were selected as HOH from a string and deleted after the warning. Polar

<table>
<thead>
<tr>
<th>Receptor/Drug</th>
<th>Code/Accession No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gefitinib</td>
<td>DB00317</td>
</tr>
<tr>
<td>Erlotinib</td>
<td>DB00530</td>
</tr>
<tr>
<td>Lapatinib</td>
<td>DB01259</td>
</tr>
<tr>
<td>Sorafenib</td>
<td>DB00398</td>
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<td>Sunitinib</td>
<td>DB01268</td>
</tr>
<tr>
<td>Flumenazil</td>
<td>DB01205</td>
</tr>
<tr>
<td>Cimetidine</td>
<td>DB00501</td>
</tr>
</tbody>
</table>
hydrogens with no bond order and Kollman charges were added [by default; ADT adds Kollman charges for a peptide (determined by checking whether all of its component residue names appear in the standard set of 20 commonly occurring amino acids) and Gasteiger charges if not so]. Finally, solvation parameters were added and the files saved as molecule. pdbqs (where “q” and “s” represent charge and solvation, respectively).

2. Preparation of the ligand file - Generally all hydrogens are added and non-polar hydrogens are merged. Gasteiger charges added unless the ligand is also a peptide in which case the above procedure would be followed. ADT automatically takes care of solvation and checks for aromatic carbon atoms and hence lone pairs and non polar hydrogens merged. ADT then determined the best root (the best root is the atom in the ligand with the smallest or largest sub-tree; in case of a tie, if either atom is in a cycle, it is picked as the root, and if neither or both is in a cycle, the first found is picked). Next we defined rotatable bonds in the ligand, making all amide bonds non-rotatable, and set the number of active torsions to fewest atoms. The ligand file was then saved with a ligand.out.pdbq extension (“q” representing charge).

3. Preparation of the grid and the grid parameter file - For the calculation of docking interaction energy, 3D grid box was created
in which the protein molecule was enclosed. The grid volume was set to be large enough to allow the ligand to rotate freely; even with it’s full extended conformation. The parameters required to create such a grid were stored in the grid parameter file, molecule.gpf.

4. Now autogrid3 was run to create a map for every atom type in the ligand and create the corresponding macromolecular file with the extension molecule.glg either from Run of the GUI or the command line autogrid3 -p molecule.gpf -l molecule.glg and when finished it writes Successful Completion.

5. Preparation of Docking Parameter File - The macromolecular pdbqs and ligand.out.pdbq files were read. The search methods of AutoDock include the Monte Carlo simulated annealing method, the genetic algorithm, local search, and the hybrid genetic algorithm with local search. The latter is also referred to as the Lamarckian genetic algorithm because offspring’s are allowed to inherit the local search adaptations of their parents, and this was the chosen algorithm for the analysis. The docking job can similarly be run from Run of the GUI or from command line autodock3 -p molecule.dpf -l molecule.dlg and when finished, Successful Completion was written the PDB file and Analyze open docking log is done and choosing different color for receptor and ligand the conformations could be played by energy. The dlg files can be opened in a terminal and each run’s (the number of runs can be fixed by the user) final docked energy, Gibbs free energy, Inhibition Constant is written. Also given in the file are the RMSD values. A conformation can be chosen and it’s coordinates written to run the next set of docking and this should be done till Gibb’s free energy is no longer significantly reduced.

6. Diagrams were drawn with the protein and the ligand attached to it in its lowest energy conformer. Protein and ligand can be made of different width and color and the background white. The image can be saved in various formats of which we chose the tif format.

Results

Rigorous docking experiments had been performed to assess the comparative efficacy between clinically used TK inhibitors (drugs) and their cross-reactivity to different receptors and/or enzymes within the physiological system. The TK drugs, the receptors, and the corresponding docking results are listed in Table 3.

Among the studied small molecular AAG drugs we have found that gefitinib most strongly binds to the VEGF receptor (-11.6 kcal/mol); whereas erlotinib has less binding affinity to the VEGF (Table 3 and Figure 1). However, sunitinib & imatinib have high binding affinity to the VEGF; whereas lapatinib and sorafenib have less binding affinity.

![Figure 8: Docked conformation of sunitinib (in blue color) to estrogen-α (A) and beta-1 (B) receptor, receptor proteins are in cyan color.](image)

<table>
<thead>
<tr>
<th>Receptor/Enzyme</th>
<th>Drug/Ligand</th>
<th>Known antagonist</th>
<th>Known agonist</th>
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<td>Androgen</td>
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<td>Dopamine-2</td>
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<td>Angiotensin II type 1 (AT1)</td>
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<td>Her-2/Neu</td>
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</tr>
<tr>
<td></td>
<td>-5.97</td>
<td>-7.74</td>
<td>-7.36</td>
</tr>
</tbody>
</table>

Table 3: Docked energies of drug – receptor/enzyme (Kcal/mol).

*NNA = N ω -nitro-L-arginine
This signifies that the use of sunitinib and imatinib would be helpful in TK mediated cancer treatment management with more effective with gefitinib. The effect of gefitinib is more pronounced due to high half-life (48 hrs) (Table 1). So cancers like NSCLC, chronic phase of CML, renal cell carcinoma, advance pancreatic neuroendocrine tumor would be better managed with this drugs, but less effective in DTC, breast cancer with Her2+, advanced renal cell carcinoma, hepatocellular carcinoma. As it is less effective in metastatic breast cancer with Her2+ so, combination with capetibatinib is suggested for lapatanib.

Among the studied drugs our results indicate that binding affinities of the drugs to the TK receptor are in following order: imatinib < gefitinib < sunitinib < soratibin < erlotinib < lapatinib. So imatinib has the highest binding affinity to TK receptor; hence it may be the ideal choice for treatment of CML (Figure 2). Gefitinib has almost the same binding affinity, hence may be recommended for others myeloproliferative disorder and NSCLC. Likewise sunitinib and sorafenib are recommended for different epithelial cancers with recurrent metastatic, or progressive cancers of thyroid, kidney, liver and GI system.

Though lapatinib is used for Her2+ breast cancer; however, our data suggested that gefitinib or sorafenib or sunitinib would be most effective (Figure 3). Probably this may be the reason that lapatinib is used in combination with capetabatinib for the treatment of Her2+ breast cancer.

Among the studied small molecular AAG drugs (Table 3) we have found that lapatinib most strongly binds to the androgen receptor (-13.2 kcal/mol); however it has least binding capacity to the Dopamin-2 receptor (Figure 4). The highest binding energy of lapatinib to androgen receptor is more than the antagonist cyproterone to androgen receptor these signifies that use of lapatinib may interefere sexual function and libido, but its effect on nausea and vomiting would be less due to less binding affinity to dopamine-2 receptor. Its binding affinity to NO synthase receptor is higher than the antagonist NNA, so it may cause cardiac dysfunction. Interestingly the binding affinity of lapatinib toward estrogen-a is high; hence it would be effective in estrogen positive breast/ovarian cancer (Figure 4). But due to less binding affinity of Beta-1 and Beta-2 receptor there is less chance of its effect of heart rate by this drug. It has a higher binding capacity to the GABA-A, GABA-B, histamine-2 angiotensin-II type-1 and VEGF, so the use of this drug may cause acidity sensation of pain and hypertension.

Imatinib has a high binding affinity to the androgen, histamine-2, angiotensin-II type-1 and most probably similar binding affinity to GABA-A, GABA-B, NO synthase, estrogen-α, VEGF; whereas it has less binding affinity to beta-1, beta-2, dopamine-2 (Figure 5). These data signify that use of imatinib may interfere sexual function, hypertension, sensation of pain, acidity, cardiac dysfunction but heart rate, nausea, vomiting would be less due to less binding affinity to the concerned receptors.

The binding affinities of gefitinib to different receptors are in the following order: androgen < GABA-B < GABA-A < estrogen-α < histamine-2 < NO synthase < angiotensin-II type-I < beta-1 < beta-2 < dopamine-2 (Table 3). So gefitinib binds most strongly with androgen receptor while moderately binds with estrogen-α receptor (Figure 6). So it may cause anaphylaxis and lung problems, chest pain, cardiac disease, eye pain but due to less binding affinity to dopamin-2, nausea, vomiting would be less. Erlotinib has also less binding affinity to dopamine-2, so it may less effect on nausea and vomiting but it would be effect on cardiac dysfunction due to high binding affinity to the NO Synthase (Figure 7).

The binding affinity of sorafenib is high to GABA-A and GABA-B receptors and less binding affinity to beta-2 and dopamine-2 receptor, so it can cause sensation of pain, acidity and cardiac problem but nausea and vomiting hardly occurs.

Sunitinib has high binding affinity to the estrogen-α and less binding affinity to beta-1 (Figure 8). This signifies it would be effective in estrogen receptor positive (ER+) breast/ovarian cancer but lesser effect on heart rate.

Discussion

To circumvent the toxic side effect of conventional chemotherapy as well as to make better control over tumor growth different antibody based AAG drugs were developed. Primarily Bevacizumab and trastuzumab were developed; both of them are now in clinical use. Though it was apprehended that they don’t possess any toxic side effect but clinical data suggest that these drugs have toxic side effects of the physiological system. We have performed Molecular docking experiment between these drugs and array of receptors and enzyme molecules of the physiological system, our data supports the clinical findings of toxicity by these drugs [48].

For clinical feasibility like storage and application to patients through different routes, different small molecular drugs have also been developed. Though several empirical based experimental as well as clinical data are available regarding their efficacy, however no comparative data in quantitative terms are available. Here we have performed computational docking method to make an assessment regarding their efficacy in quantitative terms for small molecular AAG drugs. We have found that binding efficacy of Imatinib (to TK receptor) is most potent. Gefitinib binds to TK receptor almost the same efficacy with imatinib; however, binding capacity of gefitinib to VEGF, Her-2 is the most potent. Among the studied small molecular AAG drugs; however, lapatinib has least binding affinity to TK receptor and Her-2 and erlotinib has least binding affinity to VEGF receptor. The overall finding of this work suggests that gefitinib is the most potent broad spectrum SAAG drug.

Similar to the antibody based AAG drugs several reports are available which indicate that these small molecular AAG drugs also have some toxic side effects. This may be due to cross-reactivity of these drugs to the physiological system. Previously it has been shown that different small molecular chemotherapeutic drugs when applied on a long term basis may produce off target binding ability and hence produce toxic side effect [52] and computational docking method could be the way out to prove this in a quantitative terms so that off-target points can be graded.

Off target toxicity of a drug can be assessed if that particular drug binds to a receptor with the same binding efficacy compare to its known antagonist/agonist. Though gefitinib can be regarded
as the potent small molecular AAG drug, however its binding capacity is either more or almost same to the known antagonist/ agonist of the studied receptor/enzyme molecules. Imatinib may has a strong cross reactivity to almost all the studied receptor/enzyme namely androgen, beta-2, estrogen-α, GABA-B receptor, histamine, angiotensin-II type-I receptor and NO Synthase. Though sunitinib can be consider as an effective sAAG drug after gefitinib but it has also very high binding capacity to beta-2, dopamin-2, estrogen-α, GABA-B receptors. Lapatinib has intermediate binding affinity to VEGF receptor but it has a strong binding affinity to GABA-A receptor. Our data suggest that erlotinib has less affinity to receptors involved in angiogenesis and also less toxicity potential compared to other small molecular AAG drugs. Interestingly though sorafenib has lesser binding affinity compared to gefitinib to TK, VEGF and Her-2, and it has also very lesser amount of binding affinity to the studied physiological receptor. Hence, this study would help in rationalizing to switch over from one drug to another during the intermittent period of toxicity, thereby discontinuation of therapy or adoption of other drug for management may be prevented. While make an attempt to study the off target toxicity an interesting finding may be concluded from our study is that these small molecular AAG drugs can be used for estrogen positive breast/ovarian cancer.

Androgen receptor is most effected by lapatinib and then imatinib < gefitinib < sunitinib < sorafenib < erlotinib in a sequential manner. So this result may hind toward the cause of fatigue by the use of lapatinib. Beta-1 receptor is mostly affected by gefitinib while beta-2 receptor mostly affected by sunitinib; however, almost same effects were noted for other drugs. This finding indicates cardiac dysfunction, ventricular dysfunction, pulmonary hypertension and peripheral edema. Dopamine-2 receptor is mostly affected by sunitinib also and all other drugs with almost same efficiency. This indicate that why nausea and vomiting occurs during application of small molecular AAG drugs. Lapatinib, sunitinib, gefitinib affects estrogen-α in the almost same efficiency. This may indicate the cause of fatigue, cholesterol liver damage, hepatocyte damage, dyspnea, adrenal dysfunction, anaphylaxis reaction, breathing problem that occurs after application by these drugs. When gefitinib is used, GABA receptor (both GABA-A and GABA-B) are affected mostly, this hind toward the cause of interstitial lung disease with chest pain by the use of this drug, similar inference can be drawn with Lapatinib. Histamin-2 receptor is mostly affected by imatinib and gefitinib followed by sunitinib, lapatinib, sorafenib, erlotinib and binding efficiency of all of these drugs are much more to their antagonist/ agonist, these indicate that why anorexia, nausea, loss of appetite is commonly observed by the application of any of these drugs. Binding to angiotensin-II type-I receptor of the drugs are in the order: imatinib < gefitinib < sunitinib < lapatinib < sorafenib and majority of them; and binding energy all of these drugs are more than their agonists. Therefore it may be concluded that why majority of the drugs causes either hypertension or myocardial infarction. By our study it is noted that most of the sAAG drugs bind efficiency to the NO synthase, compare to the binding efficiency of their corresponding antagonist/ agonist. Though cardiac dysfunction, myocardial infarction are notable side effect in sorafenib treatment; however, in general fatigue, hypertension and peripheral edema are seen as side effect during treatment with lapatinib, imatinib and sunitinib may be due to the effect of these drugs on NO synthase.

The drug-target network is highly important to understand the efficacy and toxicity in terms of systems biology. This study may help in an understanding to take precautionary steps to prevent the toxic side effects; if possible, different dynamical models are also developed to pin-point the onset of toxicity for long term use of chemotherapeutic drugs. Hence the grading approach of organs/ tissue specific toxicity may provide strong impetus to the dynamical model in the assessment of very onset of development by any particular drug. This way the observed efficacy and toxicity from this study would be fully understandable. Hence to explore the off-target toxicities by these small molecular AAG drugs has of relevance.

References

18. Majumder D, Mukherjee A (2013) Multi-scale modeling approaches in systems biology towards the assessment of cancer treatment dynamics: adoption of
35. (2013) Highlights of prescribing information. Astellas Pharma US.
36. BAX: ligand summary. Protein Data Bank.
40. BAX: ligand summary. Protein Data Bank.
42. B49: ligand summary. Protein Data Bank.
44. BAX: ligand summary. Protein Data Bank.

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