Farnesyl Diphosphate Synthase Inhibitors from In Silico Screening

Steffen Lindert1,2,* , Wei Zhu3, Yi-Liang Liu3, Ran Pang4, Eric Oldfield3,5 and J. Andrew McCammon1,2,6,7

1Department of Pharmacology, University of California San Diego, La Jolla, CA, 92093, USA
2NSF Center for Theoretical Biological Physics, La Jolla, CA, 92093, USA
3Center for Biophysics & Computational Biology, University of Illinois at Urbana-Champaign, Urbana, IL, 61801, USA
4School of Molecular and Cellular Biology, University of Illinois at Urbana-Champaign, Urbana, IL, 61801, USA
5Department of Chemistry, University of Illinois at Urbana-Champaign, Urbana, IL, 61801, USA
6Howard Hughes Medical Institute, University of California San Diego, La Jolla, CA, 92093, USA
7Department of Chemistry & Biochemistry, National Biomedical Computation Resource, University of California San Diego, La Jolla, CA, 92093, USA
*Corresponding author: Steffen Lindert, slindert@ucsd.edu

The relaxed complex scheme is an in silico drug screening method that accounts for receptor flexibility using molecular dynamics simulations. Here, we used this approach combined with similarity searches and experimental inhibition assays to identify several low micromolar, non-bisphosphonate inhibitors, bisamidines, of farnesyl diphosphate synthase (FPPS), an enzyme targeted by some anticancer and antimicrobial agents and for the treatment of bone resorption diseases. This novel class of farnesyl diphosphate synthase inhibitors have more drug-like properties than existing bisphosphonate inhibitors, making them interesting pharmaceutical leads.

Key words: drug discovery, molecular modeling, virtual screening

Received 5 December 2012, revised 16 January 2013 and accepted for publication 9 February 2013

The enzyme farnesyl diphosphate synthase (FPPS) has been identified as an interesting target for antitumor and anti-infective drug leads. Farnesyl diphosphate synthase is a key enzyme in the mevalonate isoprenoid biosynthesis pathway and is responsible for the condensation of dimethylallyl diphosphate with isopentenyl diphosphate (IPP) to form geranyl diphosphate (GPP) and, subsequently, farnesyl diphosphate (FPP) (1,2). Farnesyl diphosphate is important as a substrate in subsequent steps to synthesize key molecules like cholesterol and ergosterol. Pharmacologically, FPPS has gained importance in the treatment of malignant bone disease, as inhibiting FPPS blocks excessive bone resorption in osteoclasts by causing apoptosis (3). However, inhibition of FPPS has implications beyond bone disease as preclinical research has shown direct antitumor activity in a variety of human cancers (4,5). Additionally, the isoprenoid biosynthesis pathway is essential for bacterial cell wall biosynthesis as the synthesis of peptidoglycan depends on formation of lipid I and lipid II from undecaprenyl phosphate, an isoprenoid derived from FPP. Farnesyl diphosphate synthase’s role in ergosterol biosynthesis also makes it an interesting target in the search for drug leads against Chagas disease and the leishmaniases, neglected tropical diseases that affect approximately 10 million individuals.a

Historically, bisphosphonates were the first FPPS inhibitors identified (6,7) and were developed as bleaching herbicides that block carotenoid biosynthesis. It was then shown that bisphosphonate bone resorption drugs such as alendronate (Fosamax) targeted FPPS, and it appeared (8–10) that bisphosphonates might be the only efficient inhibitors of the enzyme. From a pharmaceutical perspective, bisphosphonates have several undesirable features for anti-infective or anticancer drug leads in that they are highly polar as well as being prone to rapid removal from the circulatory system by binding to the bone mineral (11). There has thus been interest in the development of more apolar bisphosphonates and even non-bisphosphonate FPPS inhibitors (12–14). In particular, a fragment-based approach identified several non-bisphosphonate FPPS inhibitors that targeted a new, allosteric binding site (3). The pocket is mainly defined by helices C, G, H and a part of helix J, with some residues from the B-C loop, the H-I loop, helix A and the C-terminal loop contributing to a lesser extent. The pocket itself is amphipathic in nature. It has a hydrophobic base and rear side, centered on residues F206, F239, L344, I348 and Y10. The opposite side is polar with several positively charged (K57, R60, K347) as well as polar (N59, T63) residues. These non-bisphosphonate FPPS inhibitors may represent novel antitumor and antibiotic drug leads as they are not expected to bind to bone mineral (11). To build on this work, we carried out a virtual screening study targeting the FPPS allosteric binding site. For this, we used the relaxed complex scheme (RCS), an in silico drug screening method that accounts for receptor...
flexibility using molecular dynamics simulations (15–17). A previously reported MD and docking study on FPPS did not target the allosteric site (12). Virtual screens were performed with AutoDock Vina (18) and Glide (19,20) on crystal structure data as well as numerous structures from a FPPS molecular dynamics simulation. A neural network rescoring was performed to optimize the ranking of known inhibitors, and 10 consensus predictions were screened experimentally yielding one hit, which was further improved by a similarity search, yielding three low (1.8–2.5) micromolar leads. To our knowledge, this is the first successful virtual screen into the FPPS allosteric site.

**Methods and Materials**

**Crystal structures and structural ensemble from molecular dynamics simulations**

We carried out a virtual screen of the FPPS allosteric site using the crystal structures described by Jahnke et al. (3). In addition, we carried out a second virtual screen using representative snapshots from an MD simulation of FPPS. The setup for the MD simulation is described in detail in (12). Frames every 20 ps were extracted from the MD trajectories; the frames were aligned using all Cα atoms in the protein and subsequently clustered by RMSD using GROMOS++ conformational clustering (21). The chosen RMSD cutoff resulted in 23 clusters that reflected most of the trajectory. The central members of each of these clusters were chosen to represent the protein conformations within the cluster and, thereby, the conformations sampled by the trajectory. The central member of a cluster (also referred to as ‘cluster center’) is the structure that has the lowest pairwise RMSDs to all other members of the cluster.

**Docking and rescoring of known non-bisphosphonate allosteric site inhibitors**

To assess the abilities of the docking software, the 12 ligands described in (3) were docked. For those compounds where no crystal structure information was available, the ChemDraw file was converted to PDB format using Open Babel (22). For the AutoDock Vina screens, pdb2pqr (23,24) was used to add hydrogen atoms to the crystal structure receptor. The AutoDock scripts (25) prepare_ligand4.py and prepare_receptor4.py were used to prepare ligand and receptor PDQBT files. A docking grid of size 18.0 Å × 18.0 Å × 18.0 Å, centered on the position of the ligand in the allosteric site, was used for docking. For Glide docking, the ligands were prepared using LigPrep, and the receptors were prepared using the tools provided in the Maestro Protein Preparation Wizard and the Glide Receptor Grid Generation.

For rescoring of AutoDock Vina docked poses, we used the python implementation of NNScore 1.0 in combination with a consensus of the top three scoring networks (12.net, 16.net and 20.net).

**Receiver operating characteristics analysis**

A receiver operating characteristics—area under the curve (ROC-AUC) analysis (26) was performed on all known allosteric site crystal structures as well as the 23 MD cluster centers. For this, the eight FPPS allosteric site inhibitors with IC50 values <100 μM from (3) were combined with the Schrödinger decoy library [1000 compounds with average molecular mass approximately 400 Da (19,20)]. All compounds in the decoy set were assumed to be inactive. Both AutoDock Vina and Glide were then used to dock the 1008 compounds into the allosteric sites of all 32 receptor structures. The compounds were ranked by their AutoDock Vina scores and Glide XP docking scores, and AUC values were calculated from the ROC analysis.

**Virtual screen of NCI diversity set II**

The virtual screen was performed using the National Cancer Institute (NCI) diversity set II, a subset of the full NCI compound database. Ligands were prepared using LigPrep, adding missing hydrogen atoms, generating all possible ionization states, as well as tautomers. The final set used for virtual screening contained 1541 compounds. Docking simulations were performed with both AutoDock Vina (18) and Glide (19,20,27). An additional rescoring was performed on the AutoDock Vina results using NNScore. Finally, the individual Glide rankings and NNScore results were combined to form a consensus list of compounds that scored well with both methods.

![Figure 1: Set of 12 compounds known to bind to the farnesyl diphosphate synthase (FPPS) allosteric site. These ligands were used as positive controls and benchmark compounds to optimize the virtual screens.](image-url)
**Experimental inhibition assay**

Human FPPS was expressed and purified and inhibition assays carried out as described previously (14). Briefly, FPPS inhibition assays were carried out using 96-well plates with a 200-μL reaction mixture in each well. The condensation of GPP (100 μM final) and IPP (100 μM final) was monitored at room temperature using a continuous spectrophotometric assay for phosphate-releasing enzymes (28). The reaction buffer contained 50 mM Tris–HCl (pH 7.4), 1 mM MgCl₂ and 0.01% Triton X100. The compounds investigated were preincubated with enzyme for 30 min at room temperature. The IC₅₀ values were obtained by fitting dose–response curve using PRISM 4.0 (GraphPad Software Inc., La Jolla, CA, USA, www.graphpad.com).

**Results and Discussion**

**Docking known non-bisphosphonate inhibitors into the FPPS allosteric site**

For control and benchmarking purposes, the 12 ligands (1–12, Figure 1) described in (3) were docked using AutoDock Vina (18) and Schrödinger’s Glide (19,20,27). These compounds have IC₅₀ values between 80 nM and 500 μM, and all are thought to target a previously unreported allosteric binding site. Compound 11, the most potent inhibitor with a published structure (PDB-ID 3N6K), has an IC₅₀ of 200 nM. No structure was published for compound 12, the best (80 nM) inhibitor.

The bound pose of compound 11, as well as the relative binding affinities of the 12 compounds, was then used as positive control to fine-tune the virtual screen parameters. First, compound 11 was docked into the allosteric site of 3N6K using AutoDock Vina. The top scoring model (predicted binding affinity −7.8 kcal/mol) recaptured the published binding pose to within 0.8 Å RMSD (Figure 2). Similarly, Glide correctly found the experimentally determined binding pose of compound 11 (0.6 Å RMSD, Figure 2). This establishes that both AutoDock Vina and Glide can correctly predict bound poses for the FPPS allosteric site which, as is apparent from Figure 2A, B, is distinct from the bisphosphonate (zoledronate) or IPP binding sites.

A much harder task is to computationally predict the relative binding affinities of multiple known binders. To address this question, all 12 compounds were docked into 3N6K and the 23 ensemble structures from MD, using AutoDock Vina. Encouragingly, compound 12 (the most potent compound) scored best, with a predicted binding affinity of −9.3 kcal/mol. AutoDock Vina was not, however, able to properly recapture the relative affinities of the 12 compounds. More specifically, 3 and 4 had high predicted affinity (rank 4 and 2, respectively), while 9 and 10 were only predicted to be weak binders.

We thus next used a neural network approach [NNScore 1.0, (29)] to rerank the compounds. NNScore has been applied to the docked poses of compound 11 into the farnesyl diphosphate synthase (FPPS) allosteric site (green; PDB ID code 3N6K), superimposed on zoledronate and isopentenyl diphosphate (IPP)-bound structure (cyan; PDB ID code 2F8Z). (A) The RMSD between the crystallographic (green) and docked pose for 11 is 0.8 and 0.6 Å, using AutoDock Vina (purple) and Glide (yellow), respectively. Also shown for reference are zoledronate (in the allylic binding site) and IPP (in the homoaallylic binding site; PDB ID code 2F8Z). (B) Expanded view of the ligand binding sites in (A).
developed to characterize the binding affinities of docked protein–ligand complexes by distinguishing between well-docked, high-affinity ligands and well-docked, low-affinity decoy compounds, through neural-network-based rescoring. Testing this approach by rescoring the AutoDock Vina docked poses of the 12 compounds we found that the relative ranking among the compounds (as well as the relative rankings against a large drug database) improved considerably. Now, compounds 9, 10 and 12 were the top three scoring docked compounds, so we then rescored all the AutoDock Vina virtual screen results with NNScore.

All 12 compounds were also docked into the 3N6K crystal structure and the 23 ensemble structures from MD, using Glide. Here, 11 was the top scoring compound (with a predicted binding affinity of $-7.5 \text{ kcal/mol}$). Also, Glide identified 9, 10 and 11 as the top three scoring compounds. Given the excellent internal ranking, no rescoring was performed on the Glide docking results.

**Receiver operating characteristics analysis of FPPS structures for enriching known active compounds**

In addition to the analysis of the ability of the 3N6K crystal structure and the 23 MD cluster centers to rank 5–12 by activity, a ROC-AUC analysis (26) was performed on all crystal structures (3) as well as the 23 MD cluster centers. The highest AUC (0.50) for the AutoDock Vina analysis was obtained with the 3N6K crystal structure, and a number of cluster centers (clusters 23, 13, 7, 17 and 6) also performed quite well in identifying active compounds. Interestingly, the Glide analysis showed a bias toward the crystal structures: the average AUC was 0.65, while the average AUC for the cluster centers was 0.35. The best AUC results were with 3N49 (AUC = 0.76) and 3N6K (AUC = 0.69). Figure 3 shows the ROC curves for 3N6K using both AutoDock Vina and Glide. Based on these results and the ranking analysis described above, we elected to use 3N6K (the structure of the most potent inhibitor) as well as the 23 MD cluster centers for the virtual screen.

**Virtual screen of NCI diversity set II**

The 3N6K FPPS crystal structure and the 23 structurally representative snapshots from the FPPS MD simulation were then used as receptors for the RCS docking protocol (15–17). Potentially including additional crystal structures might have improved the screening results. This will be the focus of future studies. The data set used in the virtual screen was the NCI diversity set II. The rationale behind using two docking programs (AutoDock Vina and Glide) was that a consensus result would have a better chance of producing good leads. The Glide docking results were ranked according to the predicted docking score, and the AutoDock Vina results were rescored with NNScore since, as discussed previously, this strategy gave the best ranking for the set of known allosteric site inhibitors. A consensus score was used to build up a list of compounds that scored well with both methods. The top 10 compounds from this list had either an NNScore value of >0.71 (the NNScore value for control compound 11) or a Glide score of <$-7.46 \text{ kcal/mol}$ (the Glide score for control compound 11) and were selected for experimental investigation. Upon confirmation of the experimental activity of 13, a computational similarity search of the entire NCI database was then performed based on 13, using a Tanimoto index of 90% or higher as the search criterion. The receptors that contributed to the good consensus score of compound 13 were cluster centers 10 (Vina/NNScore) and 18 (Glide). So while the inclusion of all cluster centers into the RCS docking analysis may seem like a drawback in light of the AUC analysis, it was vital for identification of compound 13. This underscores the power of the RCS method to identify compounds that a screen into the crystal structure.
alone would not have identified. The docked pose of compound 13 shows several stabilizing interactions such as \( \pi \)-stacking interactions with F251 and hydrogen bonds with N49 and R50. Obtaining a crystal structure is the focus of active ongoing research, so that we decided to not include the docked pose here until confirmed by X-ray crystallography.

Experimental results

The top 10 compounds identified by the virtual screen were tested experimentally. Compound 13 (Figure 4) was the only experimental hit and had an \( IC_{50} \) value of 109 \( \mu \)M. Following the similarity search, additional compounds from the NCI database were screened. Three compounds having \( IC_{50} \) values in the approximately 2–3 \( \mu \)M range were identified: 14, \( IC_{50} \) 1.8 \( \mu \)M; 15, \( IC_{50} \) 1.9 \( \mu \)M; and 16, \( IC_{50} \) 2.5 \( \mu \)M. Additionally, there were a number of other compounds found to have \( IC_{50} \) values in the low micromolar range: 17, \( IC_{50} \) 7.0 \( \mu \)M; 18, \( IC_{50} \) 10.7 \( \mu \)M; 19, \( IC_{50} \) 13.7 \( \mu \)M; 20, \( IC_{50} \) 20.3 \( \mu \)M; 21, \( IC_{50} \) 21.0 \( \mu \)M; 22, \( IC_{50} \) 22.3 \( \mu \)M; and 23, \( IC_{50} \) 35.0 \( \mu \)M. Table 1 summarizes the experimentally determined \( IC_{50} \) results. The search for improved compounds will be an important extension of this work. The most active compound investigated so far is a bisamidine containing a central hydrophobic biphenyl core. Polar substitutions into this central hydrophobic core region abolish activity. The lead compounds have \( IC_{50} \) values that are larger than those found with the bisphosphonate zoledronate (\( IC_{50} = 0.2 \) \( \mu \)M in this assay). However, as they lack the bisphosphonate feature, they are likely to have longer residence times in plasma, because they will not bind to bone mineral, as well as better cell permeability.

Table 1: Enzyme inhibition results

| Compound | Human FPPS IC\(_{50}\) (\( \mu \)M) |
|----------|-----------------------------------|
| 13       | 109                               |
| 14       | 1.8                               |
| 15       | 1.9                               |
| 16       | 2.5                               |
| 17       | 7.0                               |
| 18       | 10.7                              |
| 19       | 13.7                              |
| 20       | 20.3                              |
| 21       | 21.0                              |
| 22       | 22.3                              |
| 23       | 35.0                              |

FPPS, farnesyl diphosphate synthase.

Figure 4: Structures of the bisamidine inhibitors. Shown for convenience (in parentheses) are the National Cancer Institute (NCI) code numbers.
Conclusions

A number of leads for non-bisphosphonate FPPS inhibitors have been identified in a RCS virtual screen of the allosteric binding site. The most potent leads, 14–16, were all bisamidines with IC_{50} values in the approximately 2–3 μM range that also satisfy Lipinski’s rule of five (30) (Suite 2012: QikProp, version 3.5; Schrödinger, LLC, New York, NY, USA, 2012). In other work, we have also found that 14 is also an inhibitor of undecaprenyl diphosphate synthase (from Staphylococcus aureus) with an approximately 100 nM IC_{50} (31), opening up the possibility of developing dual FPPS/UPPS inhibitors. How these compounds bind to FPPS remains, however, to be determined, as they are clearly larger than the largest and most potent allosteric site inhibitor, 12. Other bisamidines are known to be potent antibacterials (32,33), and it has been proposed that some bind to DNA (33), while others inhibit bacterial cell wall biosynthesis (34). The results presented here suggest that in some cases, FPPS inhibition may be another target, with multisite targeting being of particular interest in the context of decreasing the likelihood of the development of drug resistance.

The results of this study also suggest possible avenues of optimization. We propose the following steps that can improve the next generation of virtual screens on the FPPS allosteric site: (i) A RCS approach seemed helpful (given that the best scoring models for compound 13 were reported for docking into cluster centers 10 and 18). In the future, simulations on the actual allosteric site–bound conformation (3N6K) might, however, give an even clearer picture of the dynamics of this binding site. (ii) Future studies should use the previously determined binding affinities as well the affinities reported in this manuscript for a more comprehensive AUC analysis. (iii) Inclusion of more than one crystal structure may improve results as well. (iv) A pocket volume analysis of the allosteric binding site may be able to identify additional structures that can be included in the AUC analysis and, possibly, future screens. (v) Results of future 3N6K simulations should be used to screen for additional druggable hot spots on FPPS: this could be followed up by more virtual screens targeting the identified sites.

Acknowledgments

We thank Jacob Durrant and William Sinko for interesting discussions concerning the dynamics and function of both FPPS and UPPS, as well as other members of the McCammon group for useful discussions. This work was supported by the United States Public Health Service (NIH grant CA158191, GM31749), the National Science Foundation, the Howard Hughes Medical Institute, the National Biomedical Computation Resource, and the NSF Supercomputer Centers. Computational resources were supported, in part, by the National Science Foundation grant PHY-0822283 and the Center for Theoretical Biological Physics. S. L. was supported by the American Heart Association (12POST11570005) and the Center for Theoretical Biological Physics.

References

1. Mao J., Mukherjee S., Zhang Y., Cao R., Sanders J.M., Song Y., Meints G.A., Gao Y.G., Mukkamala D., Hodgk M.P., Oldfield E. (2006) Solid-state NMR, crystallographic, and computational investigation of bisphosphonates and farnesyl diphosphate synthase-bisphosphate complexes. J Am Chem Soc;128:14485–14497.
2. Stresing V., Daubine F., Benzaid I., Monkkonen H., Clezardin P. (2007) Bisphosphonates in cancer therapy. Cancer Lett;257:16–35.
3. Jahnke W., Rondeau J.M., Cotesta S., Marzinzik A., Pelle X., Geiser M., Strauss A. et al. (2010) Allosteric non-bisphosphonate FPPS inhibitors identified by fragment-based discovery. Nat Chem Biol;6:660–666.
4. Buhaescu I., Izzedine H. (2007) Mevalonate pathway: a review of clinical and therapeutic implications. Clin Biochem;40:575–584.
5. Gnant M., Milnerstich B., Schipinger W., Luschin-Ebengreuth G., Postberger S., Menzel C., Jakesz R. et al. (2009) Endocrine therapy plus zoledronic acid in premenopausal breast cancer. N Engl J Med;360:679–691.
6. US Patent No. 5,756,423 May 26, 1998. US Patent No. 5,756,423 May 26, 1998.
7. Cromartie T.H., Fisher K.J., Grossman J.N. (1999) The discovery of a novel site of action for herbicidal bisphosphonates. Pestic Biochem Physiol;63:114–126.
8. Oldfield E. (2010) Targeting isoprenoid biosynthesis for drug discovery: bench to bedside. Acc Chem Res;43:1216–1226.
9. van Beek E., Pieterman E., Cohen L., Lowik C., Papapoulos S. (1999) Farnesyl pyrophosphate synthase is the molecular target of nitrogen-containing bisphosphonates. Biochem Biophys Res Commun;264:108–111.
10. Rondeau J.M., Bitsch F., Bourgi E., Geiser M., Hemmig R., Kroemer M., Lehmann S., Ramage P., Rieffel S., Strauss A., Green J.R., Jahnke W. (2006) Structural basis for the exceptional in vivo efficacy of bisphosphonate drugs. ChemMedChem;1:267–273.
11. Oldfield E., Lin F.Y. (2012) Terpene biosynthesis: modularity rules. Angew Chem Int Ed Engl;51:1124–1137.
12. Durrant J.D., Cao R., Gorfe A.A., Zhu W., Li J., Sankovsky A., Oldfield E., McCammon J.A. (2011) Non-bisphosphonate inhibitors of isoprenoid biosynthesis identified via computer-aided drug design. Chem Biol Drug Des;78:323–332.
13. Sanders J.M., Song Y., Chan J.M., Zhang Y., Jennings S., Kosztowski T., Odeh S. et al. (2005) Pyridinium-1-yl bisphosphonates are potent inhibitors of farnesyl diphosphate synthase and bone resorption. J Med Chem;48:2957–2963.
14. Zhang Y., Cao R., Yin F., Hudock M.P., Guo R.T., Krysiak K., Mukherjee S. et al. (2009) Lipophilic bis-phosphonates as dual farnesyl/geranylgeranyl diphasphate synthase inhibitors: an X-ray and NMR investigation. J Am Chem Soc;131:5153–5162.

15. Amaro R., Baron R., McCammon J. (2008) An improved relaxed complex scheme for receptor flexibility in computer-aided drug design. J Comput Aided Mol Des;22:693–705.

16. Lin J.H., Perryman A.L., Schames J.R., McCammon J.A. (2002) Computational drug design accommodating receptor flexibility: the relaxed complex scheme. J Am Chem Soc;124:5632–5633.

17. Lin J.H., Perryman A.L., Schames J.R., McCammon J.A. (2003) The relaxed complex method: accommodating receptor flexibility for drug design with an improved scoring scheme. Biopolymers;68:47–62.

18. Trott O., Olson A.J. (2009) AutoDock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization, and multithreading. J Comput Chem;31:455–461.

19. Friesner R.A., Banks J.L., Murphy R.B., Halgren T.A., Klicic J.J., Mainz D.T., Repasky M.P., Knoll E.H., Shelley M., Perry J.K., Shaw D.E., Francis P., Shenkin P.S. (2004) Glide: a new approach for rapid, accurate docking and scoring. 1. Method and assessment of docking accuracy. J Med Chem;47:1739–1749.

20. Halgren T.A., Murphy R.B., Friesner R.A., Beard H.S., Frye L.L., Pollard W.T., Banks J.L. (2004) Glide: a new approach for rapid, accurate docking and scoring. 2. Enrichment factors in database screening. J Med Chem;47:1750–1769.

21. Christen M., Hunenberger P.H., Bakowies D., Baron R., Burgi R., Geerke D.P., Heinz T.N., Kastenholz M.A., Krautler V., Oostenbrink C., Peter C., Trzesniak D., Van Gunsteren W.F. (2005) The GROMOS software for biomolecular simulations: GROMOS96. J Comput Chem;26:1719–1751.

22. O’Boyle N.M., Banck M., James C.A., Morley C., Vandermeersch T., Hutchison G.R. (2011) Open Babel: an open chemical toolbox. J Cheminform;3:33.

23. Dolinsky T.J., Czodrowski P., Li H., Nielsen J.E., Jensen J.H., Klebe G., Baker N.A. (2007) PDB2PQR: expanding and upgrading automated preparation of biomolecular structures for molecular simulations. Nucleic Acids Res;35:W522–W525.

24. Dolinsky T.J., Nielsen J.E., McCammon J.A., Baker N.A. (2004) PDB2PQR: an automated pipeline for the setup of Poisson–Boltzmann electrostatics calculations. Nucleic Acids Res;32:W665–W667.

25. Morris G.M., Huey R., Lindstrom W., Sanner M.F., Belew R.K., Goodsell D.S., Olson A.J. (2009) AutoDock4 and AutoDockTools4: automated docking with selective receptor flexibility. J Comput Chem;30:2785–2791.

26. Lee H.S., Choi J., Kufareva I., Abagyan R., Filkov A., Yang Y., Yoon S. (2008) Optimization of high throughput virtual screening by combining shape-matching and docking methods. J Chem Inf Model;48:489–497.

27. Friesner R.A., Murphy R.B., Repasky M.P., Frye L.L., Greenwood J.R., Halgren T.A., Sanschagrin P.C., Mainz D.T. (2006) Extra precision glide: docking and scoring incorporating a model of hydrophobic enclosure for protein-ligand complexes. J Med Chem;49:6177–6196.

28. Webb M.R. (1992) A continuous spectrophotometric assay for inorganic phosphate and for measuring phosphate release kinetics in biological systems. Proc Natl Acad Sci USA;89:4884–4887.

29. Durrant J.D., McCammon J.A. (2010) NNScore: a neural-network-based scoring function for the characterization of protein-ligand complexes. J Chem Inf Model;50:1865–1871.

30. Lipinski C.A., Lombardo F., Dominy B.W., Feeney P.J. (2001) Experimental and computational approaches to estimate solubility and permeability in drug discovery and development settings. Adv Drug Deliv Rev;46:3–26.

31. Zhu W., Zhang Y., Sinko W., Hensler M.E., Olson J., Molohon K.J., Lindert S. et al. (2013) Antibacterial drug leads targeting isoprenoid biosynthesis. Proc Natl Acad Sci USA;110:123–128.

32. Butler M.M., Williams J.D., Peet N.P., Moir D.T., Panchal R.G., Bavari S., Shinabarger D.L., Bowlin T.L. (2010) Comparative in vitro activity profiles of novel bis-indole antibacterials against gram-positive and gram-negative clinical isolates. Antimicrob Agents Chemother;54:3974–3977.

33. Panchal R.G., Ulrich R.L., Lane D., Butler M.M., Houseweart C., Opperman T., Williams J.D., Peet N.P., Moir D.T., Nguyen T., Gussio R., Bowlin T., Bavari S. (2009) Novel broad-spectrum bis-(imidazolinylindole) derivatives with potent antibacterial activities against antibiotic-resistant strains. Antimicrob Agents Chemother;53:4283–4291.

34. Opperman T.J. (2010) Poster Session, 50th International Conference on Antimicrobial Agents and Chemotherapy, September 12–15, Boston.

Note

WHO|Parasitic Diseases|Leishmaniasis http://www.who.int/vaccine_research/diseases/soa_parasitic/en/index3.html.