Targeting Plague Virulence Factors: A Combined Machine Learning Method and Multiple Conformational Virtual Screening for the Discovery of Yersinia Protein Kinase A Inhibitors

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Abstract: Yersinia spp. is currently an antibiotic resistance concern and a re-emerging disease. The essential virulence factor Yersinia protein kinase A (YpkA) contains a Ser/Thr kinase domain whose activity modulates pathogenicity. Here, we present an approach integrating a machine learning method, homology modeling, and multiple conformational high-throughput docking for the discovery of YpkA inhibitors. These first reported inhibitors of YpkA may facilitate studies of the pathogenic mechanism of YpkA and serve as a starting point for development of anti-plague drugs.

Many Gram-negative bacterial pathogens utilize a type III secretion system (TTSS) to inject effector proteins into the cytosol of host cells.1 These virulence factors play an important role in bacterial pathogenesis by modulating the host processes that regulate actin cytoskeletal assembly.2 With the emergence of antibiotic resistance and the threat of such bacteria being used as biological weapons, targeting virulence proteins for antibiotic design is attractive, as such compounds are unlikely to be cross-resistant or to induce resistance.3,4

Herein, we report our efforts on the discovery of inhibitors for the Yersinia protein kinase A (YpkA). YpkA is an essential virulence determinant in Yersinia spp., which includes the causative agent of plague.5 The protein contains a chaperone binding/membrane localization domain,6 a Ser/Thr kinase domain, a GDI-like domain that interacts with the Rho family of small GTPases, and a C-terminal subdomain responsible in part for actin binding and kinase activation.7,8 Because the kinase activity of YpkA has been shown to directly correlate to virulence by phosphorylating the small G protein Gtaq, inhibition of YpkA could yield new antiplague therapeutics.9,10

Protein kinase inhibitor design is a challenging problem because of the high similarity and plasticity of the catalytic site.11–14 In this study, we applied an approach combining a machine learning method and multiple conformational high-throughput docking for the discovery of YpkA inhibitors. The screening strategy employed is illustrated in Figure 1. First, we developed a machine learning support vector machine (SVM) model using a data set of known kinase inhibitors from a diverse kinase collection. The ligand-based SVM model was used as a kinase filter to prioritize the large size of chemical databases, and a target-focused library was obtained. Second, we constructed homology models of YpkA based on the MAPK templates and further performed MD simulations to sample different protein conformations characterized in the catalytic site to account for protein flexibility. Finally, with an ensemble of protein structures and the kinase inhibitor-enriched library, multiple conformational high-throughput docking was performed and a number of potent and selective inhibitors of YpkA have been successfully identified.

To develop a general kinase model for database filtering, a data set of known kinase inhibitors was seeded into a randomly selected chemical library serving as the training set. These kinase inhibitors were assembled from the literature and those in complex with a protein target deposited in the Protein Data Bank (PDB). In total, 364 kinase inhibitors were selected covering a diversity of known chemical scaffolds for both Ser/Thr kinases and Tyr kinases. The inactive data set comprising 4220 compounds was randomly selected from the MDDR database (Elsevier MDL, San Leandro, CA). Molecular descriptors were calculated with ADMET/Predictor consisting of 276 descriptors from the 3D structure (SimulationsPlus, Lancaster, CA). The use of ADMET molecular descriptors was anticipated to improve the druglikeness property of identified compounds, which is a crucial aspect in the late stage of drug development. The SVM model was derived from the molecular descriptors of the training set in distinguishing the active and inactive compounds. As shown in Figure 2, 319 out of 364 inhibitors were classified in the “positive” region, while only 15% of the
active compounds were misclassified as false negatives. To validate the model, a testing data set comprising 175 known kinase inhibitors and 669 inactive compounds was applied using the SVM model. 127 out of 175 active compounds were predicted correctly, yielding an enrichment of 70%. This result is promising and comparable to many other machine learning models published recently.15,16 Given the high efficiency of the SVM model, we then screened our in-house database collections consisting of more than 2 million compounds, and a kinase-focused library of ~200,000 compounds was obtained.

Because the structure of YpkA is unavailable, we constructed 3D models based on MAPK. YpkA shares about 20% homology to mammalian Ser/Thr kinases (Figure 3), but considering only the residues near the ATP binding site, the sequence identity to MAPK is 60%. Therefore, there is enough similarity to build a reliable model focused on the catalytic site. Two structural models were constructed on the basis of different templates of MAPK. Model A used the apo structures of p38 (PDB codes 1p38 and 1erk), while model B adopted ligand-bound complexes with induced fit at the ATP binding site (PDB codes 1a9u and 3erk). As shown in Figure 3, structural differences can be seen within these two models. Model A possesses a more open ATP binding pocket at the Glycine loop (G-loop), while the catalytic site in model B is closed with the G-loop flipping down. Because the conformational changes of the G-loop are sensitive to ligand perturbation, both are valid conformations for inhibitor design. We also examined the DFG motif, which is a key element in kinase inhibitor design.17 Although in YpkA the corresponding motif is DLG, His293 following the DLGL motif could potentially act in a similar manner to the Phe in mammalian kinases, namely, “His-in” and “His-out” as modeled in A and B.

To further examine the structural features of YpkA, we performed molecular dynamic simulations using both the apo and ATP bound models. The simulations were carried out in vacuo to permit more extended conformational changes. Analysis of the dynamics of the protein at different states revealed a number of active site residues that exhibited high flexibility (Figure 3). To sample a good representation of protein conformations for the subsequent ensemble docking, 500 conformers were extracted from 2.0 ns MD simulations and clustered according to a defined residue center at the active site. Five major clusters were obtained with model A and three clusters with model B. From the MD simulations and the docking studies we believe that the conformational changes of the active site residues represent to some extent the plasticity of the ATP binding site upon ligand binding.

With the model of YpkA and the SVM-enriched kinase inhibitor library, we then performed a multiple conformational high-throughput docking to search for YpkA inhibitors. The program FlexE was used, which accommodates multiple protein conformations in docking by forming new structural representatives.18 A total of eight conformers of YpkA sampled from MD simulations were used in FlexE docking. A kinase focused library consisting of ~200,000 compounds was subsequently docked into the ensemble of protein structures and ranked according to the FlexX score. To improve the hit selection, we also applied consensus scoring for the FlexX-docked complexes. The top 5% compounds were extracted and reranked using X-Score.19 The top 1000 compounds from both the original FlexX score and the X-Score were visually inspected in terms of overall fit, interactions in the binding site, as well as structural complexity and diversity. A total of 45 compounds were finally selected to experimentally test against YpkA. Seven of the 45 compounds showed complete inhibition at 225–450 µM, yielding a hit rate of 15%. The IC50 values of these compounds were determined by radiological assay with three compounds exhibiting inhibitory activities at 1.81, 5.87, and 9.72 µM and the remaining four having IC50 values below 50 µM. Examination of these active compounds revealed a diversity of chemical structure, as represented in Figure 4. Compound 1 possesses a scaffold of indolin-2, which is found in the derivatives of CDK2 inhibitors.20 Compound 2 belongs to the anthraquinone family, potent inhibitors of casein kinase-2.21 The structures of 3 and 4 are quite interesting, as they possess the functional group pyrimidine-2,4,6-trione. As can be seen in Figure 5, the binding
mode of pyrimidine derivative in YpkA resembles the adenosine moiety of the cofactor, involved in two H-bonding interactions with hinge residues Glu216 and Asp218.

We further evaluated the selectivity of the YpkA inhibitors by testing against two other kinases, MAPK and protein kinase C (PKC). It is not surprising that some compounds showed comparable inhibitory activities to MAPK, from which the homology models of YpkA were derived. For example, 1 showed the best inhibition of YpkA with an IC_{50} of 1.81 μM but also exhibited similar activity to MAPK with an IC_{50} of 2.45 μM. However, 2-4 are more selective to YpkA over MAPK and PKC with 5- to 10-fold better inhibition (Figure 4). The predicted interactions of 4 in the YpkA active site showed that the nitro group forms strong interactions with residue Arg221 at the end of hinge loop (Figure 5). Because an Asp or Glu residue is typically present at this position in mammalian serine/threonine kinases, interactions between the basic, positive charged Arg residue, and 4 may impart selectivity for YpkA. To the best of our knowledge, these are the first reported small-molecule inhibitors for YpkA, providing a means to investigate the mechanism of YpkA in bacterial pathogenesis, as well as a starting point for the design of potent and selective inhibitors as antiplague drugs. Although these results are promising, one must consider nonspecific inhibition due to the induction of protein aggregation. On the basis of our experimental results in Figure 4, the reported inhibitors are most likely not acting as aggregation agents and are specifically inhibiting YpkA. This is demonstrated because the compounds have little effect on PKC but are inhibitory to MAPK, from which our homology model was created.

In summary, we have described an integrated approach combining machine learning techniques and high-throughput docking for the discovery of Yersinia protein kinase A inhibitors. We have made use of the abundant resource of known kinase inhibitors and have developed a SVM model to prioritize these databases. With the construction of homology models and an ensemble of protein structures, we performed multiple conformational high-throughput docking on the target-focused library for the search of potent and selective inhibitors of YpkA. The combination of both ligand-based and structure-based knowledge of protein kinases has demonstrated high screening efficiency and reasonable speed, which has allowed us to characterize the first reported inhibitors of Yersinia protein kinase A. This integrated approach therefore provides a practical method to account for protein flexibility in a large-scale database for virtual screening of effective inhibitors of therapeutic targets.

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Supporting Information Available: The 3D structural models of YpkA, detailed methods on homology modeling, SVM clustering, MD simulations, virtual screening, and YpkA purification, inhibitor data, and experimental details for Figure 4. This material is available free of charge via the Internet at http://pubs.acs.org.

References

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