Identification of potent histone deacetylase 8 inhibitors using pharmacophore-based virtual screening, three-dimensional quantitative structure–activity relationship, and docking study

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Abstract: In recent years, histone deacetylases (HDACs) have been considered one of the promising targets for cancer chemotherapy. In the present study, a six-featured pharmacophore model with two hydrogen bond acceptors (AA), two hydrogen bond donors (DD), and two aromatic rings (RR) was developed. A predictive three-dimensional quantitative structure–activity relationship model was generated using the pharmacophore models obtained. The model has an excellent correlation coefficient and good predictive ability, as shown by the significant statistical parameters for both the training set ($R^2 = 0.9565$, standard deviation $= 0.1171$, $F = 99$, and number of ligands in training set = 21) and test set ($Q^2 = 0.8468$, Pearson’s $R = 0.9363$, number of ligands in test set = 9) molecules. The pharmacophore model was employed for the virtual screening of molecules with HDAC8 activity. The screening resulted in 366 hits with predicted activity as HDAC8 inhibitors. The hits obtained from the virtual screening were subjected to a molecular docking study to identify the potent inhibitors that binds to the active site with high affinity. The molecular docking study of known inhibitors and their analysis showed that the crucial interacting amino acid residues of HDAC8 are TYR-306, HIS-142, PHE-152, TYR-100, HIE-180, PHE-207, and Zn-388. On the basis of fitness score, predicted activities, XP Glide score, ADME results, and interacting amino acid residues, ten structurally diverse hits were reported in this paper as HDAC8 inhibitors.

Keywords: pharmacophore, atom-based 3D QSAR, virtual screening, docking, ADME, HDAC8 inhibitors

Introduction

Histone deacetylases (HDACs) play an important role in the regulation of histone and nonhistone proteins; hence, they are considered to be crucial in many biological processes, such as the regulation of gene expression, the regulation of transcription,¹ cell cycle progression, and cell survival.²,³ Deacetylation of histones is connected with transcriptional repression, together with a reduction in the expression of tumor suppressor genes.⁴ Due to their role in various biological functions, HDAC inhibition has become a promising epigenetic target for the treatment of cancer.⁵ HDAC inhibitors (HDACis) are structurally diverse and are classified into various groups as hydroximates,⁶–⁸ cyclic tetrapeptides,⁹ benzamides,¹⁰,¹¹ electrophilic ketones,¹² and carboxylic acids.¹³ Two HDACis (vorinostat and romidepsin) have been approved by the United States Food and Drug Administration for the treatment of cutaneous T-cell lymphoma.¹⁴ However, treatment with HDACis has demonstrated narrow clinical benefit for patients with...
solid tumors, prompting the exploration of novel treatment combinations with other cancer therapeutics.15

The human HDACs family consists of 18 different members16 and on the basis of phylogenetic analysis, the eukaryotic HDACs have been classified into classes I, II, III, and IV.17 Among them, HDAC8 belongs to the class I group. HDAC8 is differentially expressed and associated with various cancers. It was reported that HDAC8 is relevant in neuroblastoma;18 its inhibition induces apoptosis in T-cell cancers such as leukemia.19,20 The significant concentrations of HDAC8 found in colorectal, cervical, and gastric cancers cells21–22 indicate that HDAC8 inhibitors may be chemotherapeutic agents for colon, cervical, and gastric cancers. Thus, the development of selective HDAC8 inhibitors is of high interest in the etiology and treatment of various cancers. Most of the HDACis available to date are designed in such a way as to interact with Zn metal.23–25 The HDACs inhibitors consist of three distinct structural motifs: Zn-binding group, a hydrophobic linker, and a cap group.26 Two types of cap groups are found in HDACis: a small planar group or a cyclic tetrapeptide group.27 The inhibitors of HDAC with a small cap group are inactive if the functional group is a carboxylic acid.28 The inhibitors, which contain a cap group of cyclic tetrapeptide and a carboxylic acid functional group,29,30 have proven to be crucial for HDAC inhibition. Most of the inhibitors reported so far belong to the hydroxamic acid group, which interacts with zinc present in the active site.2 Hydroxamic acid shows strong chelation with Zn ion; for this reason, hydroxamic acid could present with metabolic and pharmacokinetic challenges.31 Hence, inhibitors devoid of the hydroxamic acid group have become increasingly desirable.

In this study, our aim was to find nonhydroxamate inhibitors by searching a database. The resulted hits were then subjected to a molecular docking study to determine the binding affinity with the receptor. The best hits obtained from the docking studies were subjected to the adsorption, distribution, metabolism, and excretion (ADME) study to compare the drug-likeness properties with standard drugs. The hits with a high fitness score, a high Glide score, high predicted activity, and drug-like properties will be selected and further taken up for the development of potential and selective HDAC8 inhibitors.

**Materials and methods**

**Dataset**

The HDAC8 inhibitors used in this study, with their corresponding half maximal inhibitory concentration (IC50) values were obtained from the literature of same assay series.32 The IC50 values of all HDAC8 inhibitors were converted into their negative logarithm of IC50 (pIC50), which spanned 1.65 orders of magnitude (5.67–7.32). All of the compounds with pIC50 values were randomly grouped into a training set of 23 molecules and a test set of nine molecules (Figure 1). The distribution of activity data versus the number of compounds is shown in Figure 2 and confirms that the test set is a true representative of the training set. The X-ray crystal structures of HDAC8 were retrieved from the RCSB protein data bank for the molecular docking studies.33 All of the computational work was carried out using an HP Z820 Workstation running over CentOS 6.3.

**Ligand preparation, protein preparation, and grid generation**

All of the structures of HDAC8 inhibitors were initially drawn using a two-dimensional (2D) sketcher (Maestro 9.6) and then converted to three-dimensional (3D) structures. All of these molecules were prepared using the LigPrep module of the Schrödinger suite (LigPrep, version 2.6, Schrödinger, LLC). The force field used was OPLS_2005. All of the possible ionic states at the target pH of 7.2±2 were generated using Epik. Possible stereo isomers were also generated for each ligand with low energy ring conformations, one per ligand. The X-ray crystal structures of HDAC8 (Protein Data Bank identifier [PDB ID]: 1T64; 1T67; 1VKG; 3SFF; and 2V5X) complexed with coligands were prepared using the “Protein Preparation Wizard” (Schrödinger Suite 2013 Protein Preparation Wizard; Epik version 2.5 and Prime version 2.2, Schrödinger, LLC; 2013). Prepared protein structures were subsequently used for grid generation using the “Receptor Grid Generation” panel. The receptor grid box was defined with a 15Å size from the center of the selected co-crystallized ligand. The root-mean-square deviation (RMSD) values between the docked conformation and co-crystallized ligands of 1T64, 1T67, 1VKG, 3SFF, and 2V5X were 0.3571, 1.7528, 0.4160, 1.0505, and 0.9651, respectively. 1T64, with the lowest RMSD value, was selected for the docking studies.

**Pharmacophore identification**

PHASE (PHASE, version 3.5, Schrödinger, LLC; 2013)14 provides six built-in pharmacophoric features: hydrogen bond acceptor (A); hydrogen bond donor (D); hydrophobic group (H); negatively ionizable (N); positively ionizable (P); and aromatic ring (R). Common pharmacophoric sites were created by keeping the default number of features. In the present study, an initial analysis showed that three chemical features (hydrogen bond acceptor [A], hydrogen bond donor
Figure 1 (Continued)
[D], and aromatic ring [R]) could effectively map all inhibitors in the dataset. In the pharmacophore hypotheses scoring techniques, each pharmacophore and its associated ligand were treated temporarily as a reference in order to assign a score; the hypotheses were ranked according to the following scores: the alignment score, the vector score, and the volume score. To determine the common pharmacophore hypothesis, all compounds with pIC_{50} values $\geq 6.983$ were considered to be active and those with pIC_{50} values $= 5.764$ were considered to be inactive. The quality of alignment was measured by survival score.

Building atom-based 3D QSAR models

The top three pharmacophore hypotheses (AADHRR.106, AADDRR.1695, and AADDHR.476) were selected to build atom-based 3D quantitative structure-activity relationship (QSAR) models (Table 1). The selected hypothesis contained hydrogen bond donors (D), hydrogen bond acceptors (A), and aromatic ring features (R), as shown in Figure 3. The fitness scores of the most active compound (1) with hypothesis AADHRR.106, AADDRR.1695, and AADDHR.476 were 3.0, 3.0, and 2.98, respectively, while the fitness scores of the least active compound (32) with same hypothesis were
2.69, 2.72, and 2.91, respectively. The training set molecules were randomly selected to constitute an atom-based 3D QSAR model; grid spacing was set to 1Å. Four partial least squares factors were used to avoid over-fitting of the results. The best QSAR model was selected for further validation. In the present study, the number of pharmacophoric features was six. The probability of matching the pharmacophoric features with the database molecules reduced with increasing numbers of features. Hence, low fitness scored molecules (fitness score ≥0.75) were also considered for the docking studies (Figure S1).

**Validation of the QSAR model**

The 3D QSAR model is validated using both the internal and external validation methods. The prediction ability of the model is evaluated by the well-known “leave-one-out” cross-validation method. For comparison between the predicted and experimental activities of the training set, $R^2$ (squared correlation coefficient) was used; for a good model, it should be ≥0.70. A high $R^2$ in the training set shows good external validation, but good internal validation does not indicate the high predictive ability of an external test set. Therefore, for a reliable predictive model, external validation is also essential. In this model, to determine the predictive power of the external validation root-mean-square error (RMSE), $Q^2$ (test set correlation), and Pearson’s $R$ (between the predicted and observed activity for the test set) were used. For a good predictive model, RMSE values should be low (<0.30), $Q^2 > 0.60$, and Pearson’s $R$ should be >0.8. The other criteria for a good model are that $R^2 - Q^2$ should not be more than 0.30, the standard deviation (SD) should be small, the highest $F$ value, and lowest P-value containing factor should be considered. Based on these statistical criteria (Table 2), hypothesis AADDRR.1695 was selected for the 3D QSAR model generation.

**Molecular docking studies**

Molecular docking studies for all known inhibitors, as well as hits obtained from pharmacophore-based virtual screening, were performed with a previously prepared receptor grid using the default options of Glide version 6.0, Schrödinger, LLC. The best docking pose was selected based on the XP Glide score.
**Table 2** Statistical parameters for the best three pharmacophore hypotheses

<table>
<thead>
<tr>
<th>Sl number</th>
<th>PDB ID</th>
<th>Superposition of docked coligand (orange) on originally bound conformation of ligand (green) in X-ray ligand–enzyme complex</th>
<th>RMSD</th>
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**Note:** Bound ligands of different crystal structures of HDAC8 are 1IT64: trichostatin A; 1IT67: M344; 1VKG: CRA-19156; 3SFF: IDI; 2V5X-V5X.

**Abbreviations:** RMSD, root-mean-square deviation; SI, serial number; PDB, Protein Data Bank; HDAC8, histone deacetylase 8.

Validation of the docking study

X-ray crystal structures of HDAC8 inhibitors were used to evaluate the ability of an XP Glide docking program. The RMSD was calculated by superimposition between the docking and cocrystalline pose of the coligand. The RMSD values of the coligands of HDAC8 (PDB ID: 1T64; 1T67; 1VKG; 3SFF; and 2V5X) were 0.3571, 1.7528, 0.4160, 1.0505, and 0.9651, respectively (Table 3). Glide was able to predict the right binding disposition of all ligands with an RMSD <3 Å, which indicated that Glide was able to reproduce the native conformation successfully.

Virtual screening

In drug discovery, the virtual screening of a database is an important part. The PHASE module of Schrödinger 9.6 was used for virtual screening of a 3D database. The best pharmacophore model, AADDR.1695, was used for the virtual screening of a 3D chemical database to identify potential lead molecules that could be further developed as HDAC8 inhibitors with high activity. The Phase Database (Phase Commercially Available Compounds) was prepared by sound data format files of commercially available compounds obtained from the following vendors: Asinex (Winston-Salem, NC, USA); Bionet (Key Organics Inc., Bedford, MA, USA); ChemDiv (eMolecules, Inc., La Jolla, CA, USA) (chemdiv.emolecules.com); Enamine (Monmouth Junction, NJ, USA); LifeChem (Life Chemicals Inc., Niagara-on-the-Lake, ON, Canada); Maybridge (Thermo Fisher Scientific, Waltham, MA, USA); Specs (Zoetermeer, the Netherlands); and TimTec (Newark, DE, USA); containing approximately 4.3×10^6 molecules. Initially, the database was filtered to remove duplicate molecules and molecules with highly undesirable properties. The resulting database was used to perform pharmacophore-based virtual screening, as depicted in the flowchart (Figure 4).
ADME prediction of identified hits
The ADME study was performed to determine the drug-likeness properties of the identified HDAC8 inhibitors using the Schrödinger module, QikProp 3.5 (Schrödinger, LLC, New York, NY, USA). QikProp provides ranges of values to compare particular molecular properties with those of 95% of known drugs. The predicted absorption, distribution, metabolism, elimination, and toxicity (ADMET) properties of hits with acceptable ranges are listed in Table 4.

Results and discussion
Twenty-four pharmacophore hypotheses were generated with AADHRR.106, AADDRR.1695, AADDHR.476, and so on, as shown in Table 1. The top three hypotheses (AADDR.1695, AADHRR.106, and AADDHR.476) with high survival scores and survival-inactive scores were chosen for QSAR model building. All the statistical parameters of the three models are summarized in Table 2. Of the three best hypotheses, AADDR.1695 yielded a statistically significant atom-based 3D QSAR model and was thus chosen for further study. Scatter plots of the observed and predicted activities for both the training and test set molecules are shown in Figure 5.

3D QSAR analysis
The 3D QSAR visualization generated by PHASE for highest active (1) and lowest active (32) compounds suggests that the blue cube regions are favorable for the introduction of the hydrogen bond donor group (D), and the orange cubes depict regions that are favorable for hydrogen bond acceptors (A) (Figure 6). Introducing these groups in the specified regions could increase the HDAC8 inhibitory activity.

Figure 7 shows the regions that are favorable and unfavorable for hydrophobic substitutions, which are mapped on both molecules 1 and 32 among the training set. The regions that favor bulky substitutions are indicated as green cubes, while the unfavorable regions are shown as purple cubes. The mapping of these regions on molecule 32 shows that hydrophobic group unoccupied in favorable regions explains its least activity.

Figure 8 shows the favorable and unfavorable regions for substitution with electron withdrawing groups on compound 1 and compound 32. The blue regions indicate that the presence of the electron withdrawing group in these regions will favor HDAC8 inhibitory activity, and the orange cube region surrounding the ligands indicates that these regions are unfavorable for substitution with electron withdrawing groups.

In sum, the favorable regions for the introduction of the hydrogen bond donor (A), hydrophobic, and electron withdrawing groups are shown as blue, green, and reddish-brown dotted regions, respectively (Figure 9).

Docking analysis
Docking studies revealed that both the inhibitors and virtual screening (VS) hits interact with active site residues through hydrogen bonds, Zn metal coordination, as well as through $\pi - \pi$ and hydrophobic interactions. The 2D ligand interactions with different amino acid residues of HDAC8, the XP Glide score, and the Zn ligand interaction distances of both inhibitors and the ten identified hits are listed in Figures S2 and S3, respectively.

The analysis of docking interactions of both inhibitors and VS hits (Table 5) suggests that the major hydrophobic interactions occurred between the inhibitors and amino acid residues (PHE-152 [number of interactions {NOI} =14], PHE-207 [NOI=11], HIE-180 [NOI=10], TYR-100 [NOI=9], PHE-208 [NOI=6], and HIS-143 [NOI=3]). The hydrogen bond donor interactions were observed with amino acid residues HIS-142 (NOI=18), GLY-151 (NOI=7), GLY-140 (NOI=6), TYR-100 (NOI=3), HIS-143 (NOI=3), TYR-306 (NOI=23), PHE-208 (NOI=3), and HIE-180 (NOI=2) amino acid residues.
Hydroxamic acid functional groups of all the inhibitors bind Zn metal in the active site.

The docking analysis of hits showed that the majority of the hydrophobic contacts occurred with TYR-306 (NOI=11), hydrogen bond donor interactions occurred with amino acid residues like GLY-151 (NOI=9), and hydrogen bond acceptor interactions occurred with TYR-306 (NOI=3). The amide functional groups of most of the hits interacted with the Zn$^{2+}$ ion in the active site. The XP Glide score of identified hits (range: $-10.596$ to $-12.969$) was better than the XP Glide score of known inhibitors (range: $-8.221$ to $-11.129$) (Figures S2 and S3). All of the selected hits interacted with the Zn-like coligands of 1T64, which indicates their potential affinity to the receptor (Figure S4).

**ADME analysis**

The ADME parameters of the identified VS hits were within the acceptable range intended for human use (Table 4).
Table 4 ADME properties of identified hits with recommended range

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<th>QPlogPoct</th>
<th>QPlogPo/w</th>
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</table>

Notes: Recommended range for 95% known drugs. Number of stars: 0–5; dHB: 0.0–6.0; aHB: 2.0–20.0; QPlogPw: 4.0–45.0; QPlogPoct: 8.0–35; QPlogPo/w: −2.0 to 6.5; QPlog5: −6.5 to 0.5; QPlogHERG: below 5; QPlogCaco: <25 poor (poor predicted apparent Caco-2 cell permeability in nm/sec) and >500 great (great predicted apparent Caco-2 cell permeability in nm/sec); QPlogBB: <30 to 1.2; QPMPMDCK: <50 poor (poor predicted apparent MDCK cell permeability in nm/sec) and >500 great (high predicted apparent MDCK cell permeability in nm/sec); QPlogKp: −8.0 to −1.0; QPlogCaco: <1.5 to 1.5; HOA: 1 low, 2 medium, 3 high; PHOA: >80% is high and <25% is low; rule of five: maximum, 4.0.

Abbreviations: ADME, adsorption, distribution, metabolism, and excretion; IC50, half maximal inhibitory concentration; dHB, estimated number of hydrogen bonds that would be donated by the solute to water molecules in an aqueous solution; aHB, estimated number of hydrogen bonds that would be accepted by the solute from water molecules in an aqueous solution; QPlogPw, predicted water/gas partition coefficient; QPlogPo/w, predicted octanol/water partition coefficient; QPlog5, predicted aqueous solubility, logarithm S in mol dm−3; QPlogHERG, predicted IC50 value for blockage of HERG K+ channels; QPMPMDCK, predicted apparent MDCK cell permeability in nm/sec; MDCK, Madin-Darby canine kidney; QPlogKp, predicted skin permeability; QPlogCaco, prediction of binding to human serum albumin; HOA, human oral absorption; PHOA, percent human oral absorption.

Table 5 Summary of the number of HPB, HBD, HBA, and interactions with Zn

<table>
<thead>
<tr>
<th>32 known inhibitors</th>
<th>10 selected inhibitors</th>
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<td>HIS-142</td>
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<td>ASP-267</td>
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<tr>
<td>Zn-388</td>
<td>28</td>
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</table>

Abbreviations: HPB, hydrophobic; HBD, hydrogen bond donor; HBA, hydrogen bond acceptor.

Therefore, the VS hits obtained could be potential leads for the development of HDAC8 inhibitors. Experimental studies could be taken up further to assess their development.

Conclusion

Six-feature predictive pharmacophore models were generated using a training set of 21 compounds for the virtual
screening of a chemical database. The screening of the database yielded 366 hits with fitness scores >0.75. A statistically significant atom-based 3D QSAR model was built using the six-feature pharmacophoric hypothesis, AAD-DRR.1695. The QSAR model visually identified important structural features, such as the hydrogen bond donor (D), the hydrophobic interaction, and the electron withdrawing effect responsible for improved activity. Most of the HDAC8 inhibitors explored to date were from hydroxamic acid, which showed strong chelation with a Zn ion. For this reason, hydroxamic acid could present metabolic and pharmacokinetic challenges. The inhibitors reported in this paper are from the nonhydroxamic acid group. Finally, based on pharmacophore matching, predicted activity, the XP Glide score, and ADME filtration, ten structurally diverse hits were reported as potential HDAC8 inhibitors. These hits may be further developed as novel potential and selective inhibitors. Further, compound synthesis and in vitro inhibitory activity will be reported in due course.

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**Disclosure**

The authors report no conflicts of interest in this work.

**References**

Supplementary materials

Figure S1 Mapping of ten identified hits with pharmacophore matching and fitness score with compound ID.

Abbreviation: Fitness, fitness score.
piC\textsubscript{50} = 7.3279; XPGS = -10.012; HBD: HIS-142 (2.14 Å); HBA: TYR-306 (1.83 Å), HIE-180 (2.06 Å); HPB: PHE-208, HIE-180, TYR-100; Zn…O=C: 2.05 Å

piC\textsubscript{50} = 7.1674; XPGS = -9.783; HBD: HIS-142 (1.82 Å), GLY-140 (2.04 Å); HPB: HIS-143; Zn…O=C: 2.16 Å

piC\textsubscript{50} = 6.9871; XPGS = -9.672; HBD: HIS-142 (2.28 Å); GLY-151 (1.87 Å); HBA: TYR-306 (1.87 Å), HIE-180 (2.37 Å); HPB: HIE-180, PHE-208, TYR-100; Zn…O=C: 2.02 Å

piC\textsubscript{50} = 6.9829; XPGS = -9.615; HBD: GLY-140 (2.40 Å); HPB: PHE-152, PHE-207; Zn…O=C: 2.02 Å

piC\textsubscript{50} = 6.9430; XPGS = -10.666; HBD: GLY-151 (2.15 Å); HPB: PHE-152, TYR-306; Zn…O=C: 2.14 Å

piC\textsubscript{50} = 6.8569; XPGS = -9.688; HBD: HIS-140 (2.44 Å); HBA: TYR-306 (2.01 Å); HPB: PHE-152, PHE-207; Zn…O=C: 2.10 Å

piC\textsubscript{50} = 6.8507; XPGS = -9.674; HBD: HIS-142 (2.06 Å); HBA: TYR-306 (1.77 Å); HPB: PHE-152, HIE-180, TYR-100; Zn…O=C: 2.36 Å, Zn…NH: 2.50 Å; Zn…OH: 2.24 Å

piC\textsubscript{50} = 6.8326; XPGS = -10.328; HBD: GLY-151 (2.07 Å); HBA: TYR-306 (2.48 Å); HPB: TYR-100; Zn…O=C: 2.24 Å

Figure S2 (Continued)
$pIC_{50} = 6.7851; XPGS = -8.528; HBD: HIS-142 (2.46 Å); HBA: TYR-306 (1.81 Å); HPB: PHE-152, HIE-180; Zn......O=C: 2.47 Å, Zn......NH: 2.47 Å, Zn......OH: 2.28 Å

$piC_{50} = 6.7375; XPGS = -9.945; HBA: TYR-306 (2.02 Å); HPB: PHE-152, PHE-207, TYR-306; Zn......O=C: 2.18 Å

$piC_{50} = 6.6968; XPGS = -10.313; HBD: HIS-142 (2.07 Å); HBA: TYR-306 (1.78 Å); HPB: PHE-208; Zn......OH: 2.24 Å

$piC_{50} = 6.5800; XPGS = -9.459; HBD: GLY-151 (2.47 Å); HBA: TYR-306 (2.20 Å), PHE-208 (2.06 Å); HPB: HIE-180; Zn......O=C: 2.15 Å

Figure S2 (Continued)
pIC₅₀ = 6.4341; XPGS = -10.352; HBD: HIS-142 (1.95 Å); HBA: TYR-306 (1.77 Å); HPB: PHE-152, PHE-207; Zn……O=C: 2.11 Å

pIC₅₀ = 6.3178; XPGS = -9.539; HBD: HIS-142 (2.04 Å), HIS-143 (2.09 Å); HPB: TYR-100, HIE-180, Zn……O=C: 2.17 Å; Zn……NH: 2.39 Å

pIC₅₀ = 6.2992; XPGS = -10.062; HBD: HIS-142 (2.12 Å), HIS-143 (1.88 Å); HBA: TYR-306 (1.84 Å); HPB: PHE-152, HIE-180, Zn……O=C: 2.39 Å, Zn……NH: 2.49 Å; Zn……OH: 2.24 Å

pIC₅₀ = 6.1979; XPGS = -10.381; HBD: GLY-140 (2.47 Å), TYR-100 (2.16 Å); HPB: PHE-152, PHE-207; Zn……O=C: 2.02 Å

pIC₅₀ = 6.1706; XPGS = -9.485; HBD: TYR-100 (2.24 Å); HBA: TYR-306 (2.08 Å); HPB: PHE-207; Zn……O=C: 2.05 Å

pIC₅₀ = 6.1598; XPGS = -10.093; HBD: GLY-151 (2.19 Å); HBA: TYR-306 (2.08 Å); HPB: PHE-152, HIE-180, Zn……O=C: 2.06 Å

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Figure S2 (Continued)
Figure S2 Experimental activity ($\text{plC}_{50}$), XPGS, type of interaction (HBD and HBA), and Zn (interaction distance) of known inhibitors.

**Abbreviations:** $\text{plC}_{50}$, negative logarithm of half maximal inhibitory concentration; XPGS, XP Glide score; HBD, hydrogen bond donor; HBA, hydrogen bond acceptor.
Figure S3 Predicted activity (pIC₅₀), XPGS, type of interaction (HBD and HBA), and Zn (interaction distance) of ten potent hits.

Abbreviations: pIC₅₀, negative logarithm of half maximal inhibitory concentration; XPGS, XP Glide score; HBD, hydrogen bond donor; HBA, hydrogen bond acceptor; Fitness, fitness score.
Figure S4 (Continued)
Figure S4 Superposition of reported hits on the coligand of the X-ray ligand–enzyme complex of 1T64 and RMSD values.

Note: Coligand is shown in green.

Abbreviation: RMSD, root-mean-square deviation.

CACPD2011a-0001270398; RMSD = 6.8264

CACPD2011a-0001734794; RMSD = 6.4913