In silico-based identification of human α-enolase inhibitors to block cancer cell growth metabolically

Abstract: Unlimited growth of cancer cells requires an extensive nutrient supply. To meet this demand, cancer cells drastically upregulate glucose uptake and metabolism compared to normal cells. This difference has made the blocking of glycolysis a fascinating strategy to treat this malignant disease. α-enolase is not only one of the most upregulated glycolytic enzymes in cancer cells, but also associates with many cellular processes or conditions important to cancer cell survival, such as cell migration, invasion, and hypoxia. Targeting α-enolase could simultaneously disturb cancer cells in multiple ways and, therefore, is a good target for anticancer drug development. In the current study, more than 22 million chemical structures meeting the criteria of Lipinski’s rule of five from the ZINC database were docked to α-enolase by virtual screening. Twenty-four chemical structures with docking scores better than that of the enolase substrate, 2-phosphoglycerate, were further screened by the absorption, distribution, metabolism, excretion, and toxicity (ADMET) properties prediction. Four of them were classified as non-mutagenic, non-carcinogenic, and capable of oral administration where they showed steady interactions to α-enolase that were comparable, even superior, to the currently available inhibitors in molecular dynamics (MD) simulation. These compounds may be considered promising leads for further development of the α-enolase inhibitors and could help fight cancer metabolically.

Keywords: α-enolase inhibitor, virtual screening, molecular dynamics simulation, glycolysis, metabolism

Introduction

A hallmark of cancer cells is their ability for unlimited growth and proliferation. To maintain this exceptional competence in cell mass expansion, cancer cells outcompete normal cells in nutrient uptake by upregulating molecules in nutrient absorption and utilization to support their needs. The drastic difference in nutrient demand between cancer cells and normal cells has made blocking the nutrient supply a fascinating and important strategy in cancer treatment and has led to the development and application of antiangiogenesis agents to treat different types of cancer. Nonetheless, cancer cells escape the cytotoxicity of anti-vascular agents by disintegration of tumor mass into small pieces to avoid dependence on the nutrient supply from the vascular system. For this reason, instead of preventing nutrient entry from the outside of the cells, restriction of nutrient uptake and utilization from within the cells would be a better alternative and the ultimate strategy to deprive cancer cells of nutrients.

To accomplish rapid growth and proliferation, cancer cells preferentially grasp and use large quantities of glucose from the surrounding extracellular space. After glucose enters the cells, it is metabolized through glycolysis to provide energy, reducing power and various building blocks for downstream anabolic processes. To support the heaviest loading metabolic pathway in cancer cells, the expression of many glycolytic enzymes needs to be upregulated.

Keywords: glycolysis, α-enolase, virtual screening, molecular dynamics simulation, cytotoxicity, hypoxia

Related Articles

- Virtual screening of the human α-enolase inhibitor
- Metabolic pathways and tumor growth
- The role of glycolysis in cancer development
- Targeting α-enolase for cancer therapy

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enzymes is constantly upregulated. α-Enolase is one of the most upregulated glycolytic enzymes in cancer cells, and has been found to overexpress in types of cancer, including nasopharyngeal carcinoma, lung, breast, colon, prostate, and so on. It not only serves as the penultimate step in glycolysis, which catalyzes the conversion of 2-phosphoglycerate (2-PGA) to phosphoenopyruvate (PEP), but is also involved in many important cellular processes in tumorigenesis, such as gene transcription, and cell migration and invasion. Due to its pivotal roles in the metabolism and tumorigenesis of cancer cells, targeting α-enolase could strike cancer cells at multiple points simultaneously and is of great interest in anticancer drug development. Until today, several α-enolase inhibitors with various potencies designed on the basis of substrate or intermediate analogs have been reported, such as the D-tartrone semialdehyde phosphate (TSP) with a Ki value in the micromolar range, 3-aminoenolpyruvate phosphate (AEP) with a Ki value in the submicromolar range, and the most potent one, phosphonoacetohydroxamate (PhAH), with a Ki value in the picomolar range. Very recently, an antibiotic, SF-2312, produced by Micromonospora s. was identified through a similarity search to the structure of PhAH as a potent α-enolase inhibitor with a Ki value in the nanomolar range. However, these inhibitors show either poor stability or undesirable pharmacological properties in vivo, and none are favorable for further clinical development.

More effort is required to identify a useful α-enolase inhibitor for therapeutic use.

To identify a useful inhibitor for α-enolase, in silico-based virtual screening is adopted to identify the lead compounds for the α-enolase inhibitors from the ZINC database – a comprehensive public accessible chemical database. Several different methodologies, including contact fingerprint analysis, ADMET properties prediction, and MD simulation, were used to refine the dock screening results. Through this work, comprehensive molecular structure information on compounds fitting the catalytic pocket of human α-enolase was obtained. The entire workflow combined with the results of contact fingerprint, ADMET, and MD simulation give not only a direction to further optimization of the α-enolase inhibitor but also a template to develop other metabolic enzyme blockers to help fight cancer.

Materials and methods

α-Enolase structure preparation

The X-ray crystallographic structure of human α-enolase (2PSN) was retrieved from the RCSB Protein Data Bank. As the substrate does not co-crystallize with the protein, the chain A structure of 2PSN was modeled against chain A of three γ-enolase structures – 3UCC, 3UCD, and 3UFA – harboring 2-PGA and magnesium in the active site and in closed conformation to rebuild the α-enolase structure using comparative structure modeling by Modeller. The model with the lowest Discrete Optimized Protein Energy score (DOPE) was chosen for subsequent virtual screening.

Ligand data set selection

The chemical structure library was constructed by selecting compounds meeting the modified Lipinski’s rule of five (molecular weight range: 55–500 Dalton; hydrogen bond donor: <5, hydrogen bond acceptor: <10, octanol–water partition coefficient logP: −1–6; polar surface area: <140 Å; net charge: −2–2) from the ZINC database (http://zinc.docking.org/zinc). All Clean subset 2012-04-26, 2013-01-10, and 2013-12-18 versions, and combined with a non-redundant approved drug dataset (a kind gift by Dr Hongjiang Li, Chinese University of Hong Kong) constructed from the US (FDA), UK (NHS), EU (EMA), Japanese (NHI), and Canadian (HC) authorities to a total number of 3,167.

Molecular docking, virtual screening, and contact fingerprint clustering analysis

Molecular docking and screening were undertaken using the CUDA supported multithreading screening program, idock, in a server hosted by the Chinese University of Hong Kong (http://istar.cse.cuhk.edu.hk/idock). A grid box encompassing the amino acid side chains involved in catalytic process, including Glu167, Glu210, Lys343, and Lys394, and stabilized substrate and metal cofactors around the catalytic pocket, including Ala39, Ser40, His158, Gln166, Arg372, and Ser373, were used in the screening. The grid box parameters were set to values of 102.724 Å, 97.855 Å, and 23.498 Å for the grid box center and 10.000 Å × 13.248 Å × 10.764 Å for the box dimensions. For each chemical structure, nine docking conformations are generated, and the best docking score was used to generate its ranking. Chemical structures with a good docking score were further analyzed and classified according to their contact modes and strengths with amino acids of α-enolase using the AuPosSOM 2.1 web interface (https://www.biomedicale.univ-paris5.fr/aupossom). The clustering began with an initial raw training phase, in which α began at 0.2 and ended at 0, radius began at 6 Å and ended with 1 Å for 1,000 iterations. It was followed by a second phase of a more accurate clustering, in which α began at 0.02 and ended at 0, and radius began at 3 Å and ended with 1 Å for 10,000 iterations. The weak contact population was...
In silico analysis of ADMET properties

The absorption, distribution, metabolism, excretion, and toxicity (ADMET) properties of the potential inhibitors were predicted in silico using the admetSAR server (http://metlmmd.ecust.edu.cn:8000/predict).21

Molecular dynamics simulation

Binding stabilities of these potential α-enolase inhibitors are further evaluated by molecular dynamics simulation using NAMD and QwikMD.24,25 The structure of the ligand–enzyme complex with ligand in the best docking pose was simulated with CHARMM general force field for protein (https://www.charmm.org/charmm)26 and the SwissParam-generated ligand force field (http://www.swissparam.ch).27 The simulation was done at a salt concentration of 0.15 M and under an implicit solvent model. The simulation is started with an equilibration simulation from 60 K to 300 K for 1.242 ns, and continued with a production simulation for 50 ns at constant temperature of 300 K. The simulation results were saved at a frequency of 2 fs. The MD trajectory generated in the production simulation was used to calculate the root mean-square deviation (RMSD) of the protein backbone, and the nonbond interaction energy between the ligand and protein using the VMD1.9.3 toolkit.28 In RMSD analysis, frame 0 is set as the reference frame, and the structure of each frame is aligned to that of frame 0 to measure actual structural fluctuations during simulation.

Virtual screening using molecular docking

A total of 22,326,149 chemical structures meet the criteria of modified Lipinski’s rule of five from the All Clean subset of the ZINC database, and the approved drug data set were used in the virtual screening. Due to the upper limit of the input number for each run (1 million/run) and to search the molecular-weight hot zone of high score hits, a crude and rapid screening was undertaken within each 100-Da interval, using chemical structures randomly selected by the program until the number reached 1 million. Following the rapid

Results

Construction of an α-enolase structure model for virtual screening

The only human α-enolase structure (PDB ID: 2PSN) in the RCSB protein data bank is an apoenzyme, without being substrate-soaked. To check the difference between the catalytic pocket of 2PSN and an enolase structure co-crystallized with substrate, the chain A structure of the 2PSN was compared with the structure of the human neuron-specific enolase (γ-enolase), which contains 2-PGA and magnesium in the structure, using the UCSF Chimera implanted Matchmaker program.29 Although the two human enolases only share 84% identity and 92% similarity, the RMSD of the superimposed polypeptide backbones, and the five catalytic amino acids between the α-enolase and three γ-enolase structures ranged from 0.415 to 0.448 Å and from 0.242 to 0.538 Å, respectively. This result indicates the two human enolases not only share a high level of conservation in protein structure, but also have a rigid architecture of catalytic pocket even in the substrate-vacant state. To make the catalytic pocket of 2PSN better resemble the conformation in the catalytic process, the chain A structure of the 2PSN was fine-tuned against the three γ-enolase structures by the homology-based comparative modeling program, Modeller.30 The normalized DOPE scores (zDOPE) of the generated models ranged from −1.75 to −1.93. The model with the lowest zDOPE score, which has the lowest free energy and is in the most native-like state, was chosen for the subsequent virtual screening. Compared to the original 2PSN structure, the structure with the lowest zDOPE score has smaller RMSD value to the three γ-enolase structures, ranging from 0.204 to 0.243 Å for the polypeptide backbone and 0.183 to 0.494 Å for the five catalytic amino acids. The structure superposition of the original and the Modeller-rebuilt α-enolase structure as well as the γ-enolase structure (3UCC) are illustrated in Figure 1.

Virtual screening of the human α-enolase inhibitor

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In silico analysis of ADMET properties

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Molecular dynamics simulation

Binding stabilities of these potential α-enolase inhibitors are further evaluated by molecular dynamics simulation using NAMD and QwikMD.24,25 The structure of the ligand–enzyme complex with ligand in the best docking pose was simulated with CHARMM general force field for protein (https://www.charmm.org/charmm)26 and the SwissParam-generated ligand force field (http://www.swissparam.ch).27 The simulation was done at a salt concentration of 0.15 M and under an implicit solvent model. The simulation is started with an equilibration simulation from 60 K to 300 K for 1.242 ns, and continued with a production simulation for 50 ns at constant temperature of 300 K. The simulation results were saved at a frequency of 2 fs. The MD trajectory generated in the production simulation was used to calculate the root mean-square deviation (RMSD) of the protein backbone, and the nonbond interaction energy between the ligand and protein using the VMD1.9.3 toolkit.28 In RMSD analysis, frame 0 is set as the reference frame, and the structure of each frame is aligned to that of frame 0 to measure actual structural fluctuations during simulation.

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survey, it was found that the molecular weight of the hits with the idock score less than −5 kcal/mol were less than 300 Da. Based on this observation, a complete and comprehensive docking screening was carried out within the range between 55 and 300 Da. At the end of screening, 87 chemical structures were found having idock scores less than −6.5 kcal/mol, including 24 with scores less than −7 kcal/mol. Among the 87 chemical structures, four pairs are in different tautomeric or ionization states of the same compounds. ZINC12428227/ZINC06523895, ZINC12428226/ZINC05567035, and ZINC32599190/ZINC0528317 are three pairs that belong to this category among the top 24 hits with scores less than −7 kcal/mol. The structures of many compounds with idock scores less than −7 kcal/mol contain five-member ring structures, such as oxadiazole, triazole, and isoxazole (Figure 2). Glycerol 3-phosphate and malic acid are the only two compounds from the approved drug data set and their idock scores are −6.63 and −6.59 kcal/mol, respectively. The idock scores for the α-enolase substrate, 2-PGA and PEP, are −7.09 and −6.63 kcal/mol respectively, and that of the four known α-enolase inhibitors, AEP, TSP, PhAH, and SF-2312, are −8.01, −7.01, −7.06, and −8.63 kcal/mol, respectively. Although, the idock scores for the four known enolase inhibitors do not correlate well with the previously reported inhibitory potency, the Ki value, their high ranking orders and a low RMSD value (1.246 Å) between the 2-PGA conformations in the modeled crystal structure and the docking pose indicate the screening results are reliable and promising for further evaluation.

**Enzyme/inhibitor contact fingerprint analysis**

To further analyze these candidates, the 87 selected chemical structures are further classified according to their contact fingerprints to α-enolase using the AuPosSom 2.1, which calculates the strengths and interaction modes of the contact—functional moieties between the ligand and protein (Figure 3). Two enolase substrates, 2-PGA and PEP, and four known α-enolase inhibitors, AEP, TSP, PhAH, and SF-2312, were included in the analysis as the internal control of contact references to differentiate the contact modes. It can be found that all the candidates share similar binding modes

**Figure 2** The structures of the top 24 α-enolase inhibitor candidates with idock scores less than −7 kcal/mol. These chemical structures are grouped according to their common ring and side-chain structures. The structures in blue, green, orange, or purple areas contain 1,2,4-oxadiazole, 1,2,3-oxadiazole, isoxazole, or a six-member ring in the structures, respectively. Others contain imidazole, triazole, or tetrazole structures. The enolase substrates and known inhibitors are shown with dashed and solid-line rectangles, respectively. The names of chemical structures in different tautomeric or ionization states of the same compounds are labeled in green and have double-headed arrows between them. The names of compounds that passed the ADMET prediction are labeled in red. Structures of these compounds were drawn by ACDLabs ChemSketch (http://www.acdlabs.com/resources/freeware/chemsketch/).

**Abbreviation:** ADMET, absorption, distribution, metabolism, excretion, and toxicity.
Virtual screening of the human α-enolase inhibitor

with α-enolase, but with different extents of interactions to the amino acids involved in the catalytic process. Group 0, which includes 2-PGA and known α-enolase inhibitors, has the most intense and strongest interactions. More detail about contact modes between 2-PGA, AEP, PhAH, SF-2312, and chemical structures with the lowest docking scores of each contact group and α-enolase are illustrated in Figure S1. Since these chemical structures show strong interactions with amino acid residues involved in the α-enolase catalytic process, chemical structures with idock scores approximately equal to or less than that of 2-PGA are retained for downstream evaluation.

Prediction of ADMET properties

To eliminate compounds with unfavorable properties, which could potentially make experimental animals suffer in downstream preclinical tests or lead to early termination of clinical trial, the top 24 hits with idock score less than −6.5 kcal/mol were further evaluated for their ADMET properties, predicted by the admetSAR web server. (Table S1). In the absorption property analysis, all 24 hits are predicted to have good absorption by the intestine (HIA+, if HIA ≥30%), but only 10 of them are Caco-2+ (Papp ≥8×10⁻⁶ cm/s), which suggests that compounds with good absorption by the gastrointestinal tract after oral uptake are retained for downstream analysis. After entering the circulation, distribution should occur uniformly to every tissue and organ to ensure better treatment efficiency. In the distribution property analysis, the 10 chemical structures are predicted to have good ability to penetrate the blood–brain barrier (BBB+), which is important when the lesion is within the brain. Moreover, their good distribution is supported as they are predicted as non-substrates or non-inhibitors of P-glycoprotein (ABCB1) and renal organic cationic transporter (OCT2/SLC22A2), which facilitate efflux of xenobiotics from the cells. After entering the cells, keeping a drug at a sustainable level would ensure better target inhibition. Xenobiotic compounds entering cells are generally metabolized by cytochrome P450 enzymes (CYP450), which determine the concentration of the drug within cells. The analysis shows that the 10 chosen chemical structures are either non-substrates or non-inhibitors of CYP2D6, CYP2C9, CYP2C19, and CYP3A4, and seven of these are even predicted as CYP1A2 inhibitors. These results indicate that these candidates could be metabolized slowly and are able to stay within cells long enough to inhibit α-enolase activity. To ensure the safety and avoid potential adverse
effects after intake, the toxicity profiles of the 10 candidates are calculated from five different models. All of these chemical structures are predicted to be noncarcinogenic (Carc_I model),\textsuperscript{31} without cardiotoxicity (hERG model I and II),\textsuperscript{32,33} and with low acute oral toxicity (class III, LD50 value between 500 and 5,000 mg/kg). Unfortunately, three of them are classified as dangerous (TD50 ≤10 mg/kg/day) and one falls within the warning category (TD50 >10 mg/kg/day) based on the Carcinogenicity test model (Carc3_I model). Six of them are predicted to be mutagens based on the AMES test model.\textsuperscript{34} These unfavorable candidates are eliminated from the candidate list. At the end of the entire ADMET evaluation screening, four compounds, ZINC1304634, ZINC16124623, ZINC1702762, and ZINC72415103, passed the test and were elected for subsequent evaluation.

Molecular dynamics simulation
To evaluate the stabilities of α-enolase–ligand complexes under dynamic conditions, we conducted a molecular dynamics simulation using a protocol described in material and methods. The stability of the complex was determined by the RMSD of protein backbone using the 50 ns trajectory data and compared with the four known α-enolase inhibitors. All of these compounds finished the 50 ns simulation, except TSP, which was dislodged from the docked site of the α-enolase catalytic pocket at 7 ns (data not shown). This result probably echoes the inhibitory potency of the weak α-enolase inhibitor. Values of the protein backbone RMSD of AEP, PhAH, SF-2312, and the four candidates of the α-enolase inhibitors enzyme complex are shown in Figure 4A together with the average backbone RMSD values range from 2.24 Å to 2.96 Å, in which ZINC1304634- and AEP- enzyme complex delineate the lower and upper limits, respectively. This result indicates the four candidates of the α-enolase inhibitor bind to the α-enolase and induce structural fluctuations comparable to that of three known enolase inhibitors. To better compare the binding stabilities of these compounds to α-enolase, the nonbond interaction energies composed of electrostatics and Van der Waals forces between each protein backbone RMSD of the α-enolase protein backbone (A) and the nonbond interaction energies (B) between α-enolase and ligand in the complexes between 0 and 50 ns.

Figure 4 The root mean square deviation (RMSD) values of the α-enolase protein backbone (A) and the nonbond interaction energies (B) between α-enolase and ligand in the complexes between 0 and 50 ns.

Abbreviations: AEP, 3-aminoenolpyruvate phosphate; PhAH, phosphonoacetohydroxamate; RMSD, root mean-square deviation.
inhibitor and α-enolase were calculated. Interaction energies between these compounds and α-enolase are contributed mostly by electrostatics interactions (data not shown), and the average nonbond energies between α-enolase and AEP, PhAH, and SF-2312 are −126.84, −183.62, and −73.90 kcal/mol, respectively. The four candidates, ZINC1304634, ZINC16124623, ZINC1702762, and ZINC72415103, have comparable nonbond energies to that of the AEP, PhAH, and SF-2312, and have values of −118.22, −168.36, −112.10, and −264.83 kcal/mol, respectively (Figure 4B). This result indicates the four compounds are promising candidates for downstream in vitro testing.

### Cross-inhibition prediction

Reducing the occurrence of off-target interactions is important to manipulate the target enzyme activity precisely and reduce side effects. To evaluate the specificity of the four candidates of the α-enolase inhibitor, they are docked to the catalytic pockets of the other two enolase isoforms, β-enolase (PDB ID:2XSX, subunit A) and γ-enolase (PDB ID:3UCC, subunit A), using idock. The grid boxes for the β-enolase and γ-enolase structures were cloned from the α-enolase docking work by superimposing the protein structures and transforming the grid box center and dimensions of the α-enolase structure into those of β-enolase and γ-enolase, respectively. The docking results are shown in Table 1. It can be found that the idock values for the 2-PGA are smaller than PEP for each of the three enolase isoforms. This result may partially reflect the preferential favoring of the forward reaction for all enolase isoforms. Despite the fact that catalytic pockets of the enolase isoforms are almost identical, they seem to respond differently to the known enolase inhibitors due to β-enolase having the highest idock scores for two known α-enolase inhibitors. This result would imply that finding an isoform-specific enolase inhibitor is possible, because the score of PhAH for β-enolase is comparably higher than that of α- and γ-enolase. On the other hand, the possibility of a pan-enolase inhibitor can also be envisaged from the result of the recently discovered enolase inhibitory antibiotic, SF-2312, which has the lowest idock scores for the three enolase isoforms. The four candidates of α-enolase inhibitors show some degree of selectivity as the idock scores of these candidates for α- and γ-enolase are comparably lower than β-enolase, but the cross-inhibition to other enolase isoforms is expected.

### Conclusion

In silico drug screening from the commercially available molecular entities provides a feasible and promising route for researchers to solve health problems by ameliorating reliance on structure creation by synthesis. The current study harnesses computer-based virtual screening to identify inhibitors of human α-enolase, which are needed for cancer treatment. From more than 22 million chemical structures, and through a series of rational refinement steps, including contact fingerprint, ADMET properties and MD simulation, we identified four compounds, ZINC1304634, ZINC16124623, ZINC1702762, and ZINC72415103, as good candidates for further testing. It is hoped further optimizations of these compounds would eventually lead to an effective α-enolase inhibitor for clinical use. The entire workflow to identify the human α-enolase inhibitors in this work could also become a template for other researchers to discover inhibitors of their targets.

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### Disclosure

The authors report no conflicts of interest in this work.

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| Table 1 Cross-inhibition prediction |
|-------------------------------------|
|          | 2-PGA | PEP | AEP | PhAH | SF-2312 | 1304634 | 16124623 | 1702762 | 72415103 |
| α-enolase| −7.09  | −6.63| −8.01| −7.06| −8.63   | −7.34   | −7.21    | −7.03    | −7.00     |
| β-enolase| −6.14  | −5.81| −6.69| −5.96| −7.71   | −6.07   | −5.95    | −5.75    | −6.19     |
| γ-enolase| −6.38  | −5.96| −6.58| −7.90| −6.83   | −6.50   | −6.38    | −6.65    | −6.65     |

Notes: Cross-inhibition is predicted using the molecular docking program idock. The score (kcal/mole) calculates the binding affinity between ligand and enzyme.

Abbreviations: AEP, 3-aminoenolpyruvate phosphate; PEP, phosphoenopyruvate; PGA, phosphoglycerate; PhAH, phosphonoacetohydroxamate.
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Supplementary materials

**Figure S1** Contact mode analysis. The functional moieties involving in the contacts between (A) 2-PGA, (B) 3 enolase inhibitors, AEP, PhAH, and SF-2312, (C) compound structures with the lowest idock scores of each contact group, and α-enolase and their relative distances are analyzed and illustrated by LigPolt. Carbon, oxygen, nitrogen, and fluoride molecules are marked as white, red, blue, and green circles, respectively. Covalent bonds in ligands and amino acid residues of α-enolase are labeled in purple and orange solid lines, respectively. The light blue dot lines label the distance (in angstroms) of hydrogen bonds formed between the functional moieties of ligands and α-enolase. Hydrophobic interactions between ligands and α-enolase are depicted by the name of involving amino acid residues, which are labeled with dark green with dark red eyelashes pointing to the involved functional moiety of ligands. Tautomers of the same compound are labeled in green.

*Abbreviations:* AEP, 3-aminoenolpyruvate phosphate; PGA, phosphoglycerate; PhAH, phosphonoacetoxyhydroxamate.
Table S1 ADMET properties of potential α-enolase inhibitors with idock scores less than −7 kcal/mol

| Zinc ID     | idock score | HIA  | Caco-2 | BBB  | P-gp(S/I) | RenalOCT | CYP InP | hERG(I/II) | AMES | Carcinogen | AOT | Carcinogenicity |
|-------------|-------------|------|--------|------|-----------|----------|---------|------------|------|------------|-----|----------------|
| 12428226<sup>a</sup> | −8.32       | +    | +      | n/n  | −         | low(1A2+) | w/non   | +          | −    | III        | D   |                 |
| 5567035<sup>b</sup>  | −8.29       | +    | +      | n/n  | −         | low(1A2+) | w/non   | +          | −    | III        | D   |                 |
| 12428227<sup>b</sup> | −8.00       | +    | −      | n/n  | −         | low     | w/non   | +          | −    | III        | NR  |                 |
| 967421       | −8.00       | +    | +      | n/n  | −         | low(1A2+) | w/non   | +          | −    | III        | NR  |                 |
| 6523895<sup>b</sup> | −7.78       | +    | −      | n/n  | −         | low     | w/non   | ±          | −    | III        | NR  |                 |
| 3163100      | −7.52       | +    | +      | n/n  | −         | low(1A2+) | w/non   | +          | −    | III        | NR  |                 |
| 14980359     | −7.51       | +    | −      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 4343380      | −7.47       | +    | −      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 32599190<sup>a</sup> | −7.43       | −    | −      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 5282317<sup>a</sup> | −7.39       | +    | −      | n/n  | −         | low     | w/non   | +          | −    | III        | NR  |                 |
| 4357691      | −7.36       | +    | +      | n/n  | −         | low     | w/non   | +          | −    | III        | NR  |                 |
| 1304634      | −7.34       | +    | +      | n/n  | −         | low(1A2+) | w/non   | −          | −    | III        | NR  |                 |
| 1666582<sup>b</sup> | −7.26       | +    | −      | n/n  | −         | low(1A2+) | w/non   | +          | −    | III        | NR  |                 |
| 2258900      | −7.23       | +    | −      | n/n  | −         | low     | w/non   | +          | −    | III        | NR  |                 |
| 4342073      | −7.22       | +    | +      | n/n  | −         | low(1A2+) | w/non   | +          | −    | III        | D   |                 |
| 6854842      | −7.21       | +    | −      | n/n  | −         | low     | w/non   | +          | −    | III        | NR  |                 |
| 16124623     | −7.21       | +    | +      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 1063089<sup>a</sup> | −7.20       | +    | +      | n/n  | −         | low     | w/non   | +          | −    | III        | W   |                 |
| 1601705      | −7.13       | +    | −      | n/n  | −         | low     | w/non   | −          | −    | III        | W   |                 |
| 6666767      | −7.11       | +    | −      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 1295964      | −7.10       | +    | −      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 71605889     | −7.04       | +    | +      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 1702762<sup>a</sup> | −7.03       | +    | +      | n/n  | −         | low     | w/non   | −          | −    | III        | NR  |                 |
| 72415103     | −7.00       | +    | +      | n/n  | −         | low(1A2+) | w/non   | −          | −    | III        | NR  |                 |

Notes: Tautomers are labeled with asterisks (*), and different ionization states of the same compounds are labeled either with pounds ($) or ampersands (#). Idock score, kcal/mole, the binding affinity between ligand and enzyme; HIA, the probability of absorption by human intestine, HIA ≥ 30% is defined as HIA+; Caco-2, Caco-2 permeability value (Papp) ≥ 8 × 10<sup>−6</sup> cm/s is denoted as high permeability (+), otherwise as poor permeability (−); BBB, the probability of penetrating the blood–brain barrier; P<sub>gp(S/I)</sub>, P<sub>gp</sub> substrate or inhibitor. The probability of being a substrate or an inhibitor for P<sub>gp</sub> protein. The n means that the chance is low; CYP InP, cytochrome P450 inhibitory promiscuity. The probability of inhibiting cytochrome P450 isoforms, CYP450 1A2, 2C9, 2C19, 2D6, and 3A4; hERG(I/II), the probability of inhibiting human ether-a-go-go-related genes, which was predicted by two models (III). In model I, if a compound has a pIC50 > 6.0 μM, it is denoted as strong inhibitor (+), otherwise as weak inhibitor (−). In model II, if a compound has an IC50 > 50 μM, it is classified as an inhibitor, and otherwise as a non-inhibitor (−); AMES, The probability of being as mutagen; Carcinogen, the probability of being as carcinogen; AOT, compound in class III has LD50 value between 500 and 5,000 mg/kg. Carcinogenicity (three-class). Carcinogenic compounds with TD50 (tumorigenic dose rate 50) ≤ 10 mg/kg body wt/day were assigned as Danger (D), those with TD50 > 10 mg/kg body wt/day were assigned as Warning (W), and non-carcinogenic chemicals were assigned as Non-required (NR). The names of compounds that passed the ADME evaluation are labeled in bold.

Abbreviations: ADMET, absorption, distribution, metabolism, excretion, and toxicity; aOT, acute oral toxicity.