Discovery of Tyrosinase Inhibitors: Structure-Based Virtual Screening and Biological Evaluation

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Abstract

Tyrosinase (EC 1.14.18.1) plays an indispensable role in the rate-limiting steps of melanin biosynthesis, and its uncontrolled activity may result in various diseases, such as albinism, melanoma, freckles, etc. The inhibition of tyrosinase activity may provide a useful and efficient strategy to treat hyperpigmentation disorders. However, the widely used tyrosinase inhibitors, like α-arbutin, hydroquinone, and kojic acid, have many shortcomings, such as lower efficacy and much more side effects. Herein, we reported the use of homology modeling and multistep structure-based virtual screening for the discovery of novel tyrosinase inhibitors. In this study, 10 initial potential hits (compounds T1–T10) were evaluated for enzyme inhibition and kinetic study, with kojic acid being used as a control. Among them, the IC50 values of both T1 (11.56 ± 0.98 µmol/L) and T5 (18.36 ± 0.82 µmol/L) were superior to that of kojic acid (23.12 ± 1.26 µmol/L). Moreover, T1 and T5 were also identified as the effective noncompetitive tyrosinase inhibitors by the subsequent kinetic study. Above all, T1 and T5 may represent the promising drug candidates for hyperpigmentation therapy in pharmaceutical fields, as well as the effective whitening agents in cosmetic applications.

Keywords
► tyrosinase inhibitors
► hyperpigmentation
► homology modeling
► virtual screening
► kinetic study

# These authors contributed equally to this work.

received October 21, 2021 accepted December 7, 2021


ISSN 2628-5088.

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Georg Thieme Verlag KG, Rüdigerstraße 14, 70469 Stuttgart, Germany
Introduction

Tyrosinase (EC 1.14.18.1) is a type III metalloprotease and oxidoreductase containing two copper ions in the active center accompanied by six histidine residues. Tyrosinase is widespread in microorganisms, plants, animals, and humans, and plays the key role as the rate-limiting enzyme in melanin biosynthesis. Melanin contributes to cuticular hardening and exoskeletal pigmentation in animals, and enzymatic browning in fruits and vegetables. And the color of human skin is predominantly determined by the amount of melanin produced by skin melanocytes. Melanin can prevent the skin from the side effects of environmental factors such as ultraviolet radiation and oxidative stress. However, excessive accumulation of melanin results in pigmentation diseases including freckles, age spots, melanosis, malignant melanoma, etc. In this case, tyrosinase inhibitors, such as kojic acid, α-arbutin, hydroquinone, and azelaic acid, are considered as important skin-whitening agents or food preservatives in the fields of medicine, cosmetics, and food. However, they have certain side effects and cannot be widely used. Thus, developing novel efficient and selective inhibitors for tyrosinase remained significantly important.

Computer-aided drug design (CADD) represents an effective tool for drug discovery, which can largely reduce the cost of drug discovery and increase the feasibility of drug screening. It is acknowledged that virtual screening of molecular docking based on receptor structure is a common method in drug design. Virtual screening from a large database is based on the three-dimensional (3D) structure of receptor enzymes to discover novel ligands. The binding affinity of ligand–receptor complexes can be predicted according to the interaction between ligands and protein macromolecules, thus favoring the discovery of a better ligand structure. Obviously, computer technology can simulate a more real binding state of ligand and receptor, greatly reducing the number of the wet-lab ligand activity experiments and saving a lot of manpower, material, and financial resources. However, virtual screening based on receptor structure is often dominated by visual inspection of medicinal chemists in the later stage, and many target proteins are still short of a clear and complete crystal structure, which together greatly limit the wide application of the tool.

Herein, we reported a multiple-step virtual screening method of receptor structure, based on the homologous modeling of human tyrosinase. Potential active molecules were selected from the compound library constructed by our laboratory for tyrosinase biological activity test. In our study, two noncompetitive tyrosinase inhibitors (T1 and T5) with novel skeleton structures were obtained. Compounds T1 and T5 may be used as promising drug candidates for hyperpigmentation treatment, and whitening agents in cosmetic applications.

Results and Discussion

Homologous Modeling Results

Human tyrosinase belongs to a family of transmembrane glycoproteins and faces a lot of challenges toward stable crystallization at a high concentration. Until now, there is still no available cocrystal structure of human tyrosinase. A homology modeling strategy shrinks the gap between the amino sequence and the 3D structure of protein. In this study, a homology modeling was performed using the Swiss-model for the virtual screening. First, an amino acid sequence of human tyrosinase (P14679) was retrieved in the UniProt database. After importing the amino acid sequence, the results showed that the template system selected was human tyrosinase–related protein 1 (TYRP-1) (PDB ID: 5M8L), which had four chains A, B, C, and D and was the crystal structure of TYRP-1 and 2-acetamido-2-deoxy-β-D-glucopyranose. A chain was chosen by the system automatically. The total length of the chain sequence was 446, and the resolution was 2.35 Å. The homologous modeling of human tyrosinase was established by the Swiss-model. The sequence identity between human tyrosinase and TYRP-1 was 43.32% and the global model quality estimate value was 0.67. The amino acid residues of the homologous modeling were from His19 to Gln453, the total sequence length was 435, and both Met1-Glu18 and Asp454-Leu529 were missing. The active site of human tyrosinase was similar to that of mushroom tyrosinase (PDB ID: 2Y9X), which contained two metal ions coordinated with six histidine residues. On the basis of the mentioned protein preparation work above, we conducted the multiple-step virtual screening targeting tyrosinase.

Structure-Based Virtual Screening

Molecular docking plays a common and important role in the area of structural molecular biology and CADD. There are numerous successful examples obtained with the help of this method. Molecular docking can speed up the pace of drug development and lower down the costs. – Fig. 1 shows the process of a multiple-step virtual screening in discovering lead inhibitors of human tyrosinase. In the first turn, small-molecule compound databases including SPECS, ACDB, and MCD with more than 200,000 compounds were filtered by the Lipinski’s rule of five. It is well known that the molecule with the following features would be more likely to result in a real small-molecule drug: (1) a molecular mass less than 500 Da; (2) no more than 10 hydrogen bond acceptors; (3) no more than five hydrogen bond donors; and (4) log P (octanol–water partition coefficient) no greater than 5. In addition, all of these compounds are commercially available or synthesized in laboratory, and could be picked out for further biological evaluation. In this study, a total of 198,460 small-molecule compounds were obtained for the following docking-based virtual screening.

Next, the filtered databases were screened by molecular docking using the crystal structure of human tyrosinase in the HTVS (high-throughput virtual screening) module of Glide. The top 5,000 hits with a Glide docking score ranging from −7.97 to −3.52 kcal/mol were screened. Afterward, the 5,000 hits were docked precisely with human tyrosinase in the SP (standard precision) module of Glide. The top 150 candidates were selected for the docking pattern analysis, whose Glide docking scores were greater than −4.12
kcal/mol. We searched the professional advice of pharmaceutical chemists to classify the 150 candidates by visual inspection according to their structural features. In this process, the docking score, the interaction between key amino acids and ligands, the match degree of compound conformation and protein, the interaction patterns, and especially the novelty of molecular skeleton were highly considered. Consequently, the 10 hits with different skeletons were purchased from SPECS database and determined for the following tyrosinase inhibition assay (Fig. 2).

**Fig. 1** The workflow of discovering tyrosinase inhibitors based on a multistep structure-based virtual screening.

**Fig. 2** The chemical structures of 10 lead hits prepared for the biological activity assays.
Mushroom Tyrosinase Inhibition Assay

To carry out the tyrosinase inhibition assay, mushroom tyrosinase was used to detect the inhibition rate of 10 hits. Mushroom tyrosinase is commercially available, highly homologous to mammalian enzyme in the active center, and is one of the most common tyrosinase enzymes used today. Kojic acid is a well-known whitening agent in cosmetic field and a food additive in preventing enzymatic browning. Kojic acid inhibits tyrosinase activity, and was used as a positive control in the biological activity assay according to a reported study. The results of the preliminary screening of tyrosinase inhibitory activity (Fig. 3) showed that four hits (compounds T3, T4, T7, and T8) exerted no inhibitory potency at two concentrations. Four hits (compounds T2, T6, T9, and T10) showed moderate inhibitory activity at the concentration of 30 µmol/L. Obviously, compounds T1 and T5 exhibited stronger tyrosinase inhibitory potency than the positive control kojic acid. Tyrosinase inhibition rates of T1 and T5 were 90 and 70%, respectively, at the concentration of 30 µmol/L, while over 70 and 20%, respectively, at the concentration of 10 µmol/L. Thus, T1 and T5 were chosen for the further IC50 (half-maximal inhibitory concentration) measurement.

IC50 is a measure of the effectiveness of inhibitors in medicinal chemistry. IC50 is defined as the concentration of an inhibitor that inhibits up to 50% in such biosystem like enzyme. Then, a dose–response curve for T1 and T5 was constructed using GraphPad Prism 6 (version 6.01). The results showed that T1 and T5 dose-dependently inhibited the tyrosinase activity (Fig. 4A). And the IC50 values of T1 and T5 were 1.56 ± 0.98 and 18.36 ± 0.82 µmol/L, respectively, much lower than that of kojic acid (23.12 ± 1.26 µmol/L). Particularly, compound T1 exhibited the best inhibitory activity with more than twofold inhibitory potency in comparison to the control. In short, compounds T1 and T5 were effective tyrosinase inhibitors, and their inhibition mechanisms were further explored in the following study.

Fig. 3 The preliminary screening results of mushroom tyrosinase inhibitory activity of 10 lead hits performed at 10 and 30 µmol/L using L-tyrosine as the substrate. Data are presented as means (n = 3).

Fig. 4 Inhibition effect and inhibition mechanism assays of T1 and T5. (A) Inhibition rate curve against mushroom tyrosinase inhibitory. (B) Determining inhibition type for mushroom tyrosinase activity. Each experiment was repeated in three parallels.
Kinetic Study of Tyrosinase Inhibition
To identify the inhibition type of two lead hits, the relationship between substrate concentration and enzymatic reaction rate was illustrated by the GraphPad Prism 6 (version 6.01). An inhibitor was determined to be irreversible or reversible according to whether the two lines were parallel or intersected at origin. Fig. 4B shows that the three lines intersected at origin. T1 and T5 decreased the slope of the straight line, inhibited the enzyme activity, and led to the reduction in the catalytic efficiency of tyrosinase (instead of through reducing the effective enzyme amount). All above results suggested that T1 and T5 were reversible tyrosinase inhibitors.

Furthermore, the reaction rates of the lead hit with different concentrations of L-tyrosine were also collected, and a Lineweaver–Burk double-reciprocal plot was analyzed to evaluate the inhibition type of the two lead hits in depth. The kinetic study showed that both the curves of T1 and T5 intersected on the X-axis (Fig. 5), indicating that T1 and T5 were noncompetitive inhibitors of mushroom tyrosinase due to the decreased $V_{\text{max}}$ value and the unchanged $K_m$ (the Michaelis constant) value. Moreover, for each lead hit, the $K_i$ (the inhibitory constant) value was also calculated from the plot. The $K_i$ value of T1 and T5 was 11.35 and 17.7 µmol/L, respectively, indicating that in comparison to T5, T1 had a higher affinity with mushroom tyrosinase, which was consistent with the results from IC$_{50}$ assay (Table 1).

Molecular Docking Results
Then molecular docking analyses of T1 and T5 in silicon were performed using the protocol of Ligand Interaction Diagram of SchrödingerSuite and Protein–Ligand Interaction Profiler (PLIP). Our data suggested that T1 and T5 had a similar binding mode and site to human tyrosinase. Fig. 6A shows the docking results of T1, which shows that the ligand molecule was stabilized by intermolecular hydrogen bonds with Asp197, Glu203, and Gln378, and π–π stacking interactions with His367. In addition, the fluorine atom was oriented toward the metal ions in the active center and formed a halogen bond with Ser380, which was beneficial to stabilize the complex of the ligand and human tyrosinase. The ligand molecule could interact with the surrounding amino residues, including Phe347, His367, Ile368, and Val377, through the hydrophobic contact. Fig. 6B shows the docking results of T5, which displays that the carboxyl group could chelate with the metal ion in the active center, and the benzene ring might form π–π interaction with His202 and His367. The chlorine atom can interact with Gln378 through halogen interaction, while the benzylpiperidine moiety formed π–π interaction with Phe347. In addition, the ligand formed hydrogen bonds with...

Table 1 Mushroom tyrosinase inhibitory activity and kinetic assays of two lead hits

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC$_{50}$ (µmol/L)$^a$</th>
<th>$K_i$ (µmol/L)$^b$</th>
<th>Type of inhibition</th>
<th>Docking score (kcal/mol)$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1</td>
<td>11.56 ± 0.98</td>
<td>11.35</td>
<td>Noncompetitive</td>
<td>−5.576</td>
</tr>
<tr>
<td>T5</td>
<td>18.36 ± 0.82</td>
<td>17.7</td>
<td>Noncompetitive</td>
<td>−6.920</td>
</tr>
<tr>
<td>Kojic acid</td>
<td>23.12 ± 1.26</td>
<td>–</td>
<td>–</td>
<td>−4.996</td>
</tr>
</tbody>
</table>

$^a$Data are presented as means ± SEM from three parallels.

$^b$K$_i$ values were obtained at the concentration of 30 µmol/L of T1 and T5.

$^c$Docking scores were collected from SP module of Glide docking.
Ser375 and Lys306, as well as hydrophobic interactions with Asn364, His367, Ile368, and Val377, making the complex more stable. The two-dimensional interaction patterns between T1 and human tyrosinase (►Fig. 6C) and T5 and human tyrosinase (►Fig. 6D) are also illustrated. Our data showed that the docking scores of T1 (C0 5.576 kcal/mol) and T5 (C0 6.920 kcal/mol) were much higher than that of the control (C0 4.996 kcal/mol) with the optimal docking results being seen in T5 (►Table 1). Thus, we predicted that the chelation of carboxyl group and metal ion in the active site might be the main factor affecting the docking results. The interaction with metal ions or histidine residues in the active center might directly enhance the inhibitory effect on tyrosinase.

**Conclusion**

In summary, we have introduced a multistep virtual screening process using CADD methods including molecular docking and homology modeling. Compounds T1 and T5, along with other eight potential active compounds, were picked out through virtual screening from the database containing more than 200,000 small-molecule compounds. Excitedly, T1 and T5 were identified as effective noncompetitive reversible tyrosinase inhibitors. The IC50 values of the two inhibitors were 11.56 ± 0.98 and 18.36 ± 0.82 µmol/L, respectively, which were superior to the positive drug kojic acid (23.12 ± 1.26 µmol/L). Our experimental results provided a strong basis for the efficiency of our screening and docking strategies. Our study enriched the function study of human tyrosinase, suggested two promising drug candidates (T1 and T5) for the treatment of hyperpigmentation and malignant melanoma, and provided two potential whitening agents for the cosmetic field. Furthermore, the appropriate structure optimization of T1 and T5 and structure–activity relationship study need to be further studied in our laboratory; unfortunately, data are unavailable at present.

**Methods and Materials**

**Homology Modeling**

Homology modeling was performed according to previous studies.25 The 3D structure of human tyrosinase protein was simulated by the Swiss-model.26 The amino acid sequence of human tyrosinase was downloaded from the UniProt...
database (https://www.uniprot.org; ID: P14679).\textsuperscript{27} The sequence was copied to the Swiss-model Target Sequence and the project request was submitted. The server returned the homology modeling result after a period of time.

**Structure-Based Virtual Screening**

**Protein Preparation**

Homology modeling was used to construct the 3D structure of human tyrosinase. Human tyrosinase was optimized using the Protein Preparation Wizard module of Maestro (version 11.8) in SchrödingerSuite (Schrödinger, LLC, New York, NY, United States) with default parameters to remove water molecules, add hydrogen atoms, and fill in missing residues.\textsuperscript{28} The active site was defined in the active site containing metal ions through the Receptor Grid Generation module.

**Database Preparation**

The databases including SPECS, ACDB, and MCDB were filtered in accordance with Lipinski’s rule of five. The structures of all molecules were optimized by the LigPrep module with the OPLS_2005 force field being in an ionized state at pH 7.0 $\pm$ 2.0, and the default parameters being used to generate the 3D structure.\textsuperscript{29} The optimized database was saved as Maestro file format.

**Molecular Docking**

Molecular docking of human tyrosinase was performed by the Ligand Docking module. The protein structure was derived from homology modeling. First, the Glide module HTVS was used to roughly evaluate the binding affinity of the human tyrosinase and the ligands in the prepared database in the active site.\textsuperscript{30} Then, the top 5,000 hits were selected and docked precisely with human tyrosinase by SP and XP (Extra Precision) modules, leaving the top 150 leads.\textsuperscript{31} Analysis and visualization of the molecular docking results were completed by means of Ligand Interaction Diagram and PLIP tool.\textsuperscript{32}

**Mushroom Tyrosinase Inhibition Assay**

The enzyme inhibitory activity assay was performed according to previous studies.\textsuperscript{33,34} In brief, a total of 0.1 mL reaction system was added to a 0.5 mL centrifuge tube, which contained 50 µL sample (10 and 30 µmol/L, respectively) and mushroom tyrosinase (25 µL, 1 mmol/L). The mixed solution was maintained at 25°C for 20 minutes, and then L-tyrosine (25 µL, 2.5 U/µL) was used as a substrate and mixed well. The change of optical density (OD) value was measured within 5 minutes at 492 nm. DMSO-containing phosphate buffer (50 µL) was used as a vehicle control plate, while DMSO-containing phosphate buffer (75 µL) and L-tyrosine (25 µL, 2.5 U/µL) was used as a blank control plate. The inhibition rate was calculated with the following Eq. (1).

$$\text{Inhibition rate} (%) = \frac{\Delta \text{OD}_{\text{compound}} - \Delta \text{OD}_{\text{vehicle}}}{\Delta \text{OD}_{\text{blank}}} \times 100\% \quad (1)$$

where $\Delta \text{OD}_{\text{compound}}$ is the changed OD value after inhibitor treatment, $\Delta \text{OD}_{\text{vehicle}}$ is the OD value for the vehicle control plate, and $\Delta \text{OD}_{\text{blank}}$ is the OD value for the blank control.

**IC\textsubscript{50} Assay**

IC\textsubscript{50} assays for T1 and T5 were further assessed. Different concentrations (less than 1 mmol/L) of T1 and T5 were dissolved in DMSO, respectively. The changed OD value for each solution in the 0.1 mL reaction system was measured according to the method under the heading “Mushroom Tyrosinase Inhibition Assay.” The tyrosinase activity rate was calculated according to the inhibition rate of Eq. (1). The dose–response curve was drawn by GraphPad Prism 6 (version 6.01). The IC\textsubscript{50} value for each compound was obtained according to the dose–response curve.

**Kinetic Study**

The kinetic study was performed according to a reported study.\textsuperscript{35} We prepared 30 µmol/L of T1 and T5, and different concentrations of mushroom tyrosinase (0.78, 1.56, 3.125, 6.25, 12.5, 25, 50, and 100 U/well). The enzyme catalytic activity was evaluated using the method under the heading “Mushroom Tyrosinase Inhibition Assay.” The blank control was investigated in the same dose except for the addition of inhibitors. The curves were made by GraphPad Prism 6 (version 6.01) to illustrate the relationship between tyrosinase concentration and the tyrosinase enzyme activity.

The inhibition mechanisms of T1 and T5 were further studied. L-tyrosine was prepared at the concentration of 1,000, 667, 444, 296, 198, 132, 88, 59, and 39 nmol/L, respectively. The blank control was performed in the same dose except for the addition of inhibitors. The catalytic activity of tyrosinase enzyme was completed respectively according to the method under the heading “Mushroom Tyrosinase Inhibition Assay.” Lineweaver–Burk double-reciprocal plots were determined by the reciprocal 1/[S] (the concentration of L-tyrosine), the reciprocal 1/V (the enzyme catalytic activity for OD change), and the $K_m$ and maximum velocity ($V_{max}$) values were calculated later.\textsuperscript{36}

**Funding**

This work was supported by the National Natural Science Foundation of China (Grant No. 81922064, 22177083, 81803755, and 81874290).

**Conflict of Interest**

All authors declare no conflict of interest.

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Pharmaceutical Fronts Vol. 4 No. 1/2022 © 2022. The Author(s).

Discovery of Tyrosinase Inhibitors

Wang et al.


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