Crystallographic evidence that dinuclear copper center of tyrosinase is flexible during catalysis

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Running Title: Structure of tyrosinase complexed with a caddie protein

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At high resolution, we determined the crystal structures of copper-bound and metal-free tyrosinase in a complex with ORF378 designated as a “caddie” protein because it assists with transportation of two Cu(II) ions into the tyrosinase catalytic center. These structures suggest that the caddie protein covers the hydrophobic molecular surface of tyrosinase and interferes with the binding of a substrate tyrosine to the catalytic site of tyrosinase. The caddie protein, which consists of one six-stranded β-sheet and one α-helix, has no similarity with all proteins deposited into Protein Data Bank.

Although tyrosinase and catechol oxidase are classified into the type-3 copper protein family, the latter enzyme lacks monooxygenase activity. The difference of the catalytic activity is based on the structural observation that a large vacant space is present just above the active center of tyrosinase, and that one of the six His ligands for the two copper ions is highly flexible. These structural characteristics of tyrosinase suggest that, in the reaction that catalyzes the ortho-hydroxylation of monophenol, one of the two Cu(II) ions is coordinated by the peroxide-originated oxygen bound to the substrate. Our crystallographic study shows evidence that the tyrosinase active-center formed by dinuclear copper is flexible during the catalysis.

Tyrosinase (EC 1.14.18.1), which belongs to a protein family having the catalytic center formed by dinuclear copper, catalyzes the ortho-hydroxylation of monophenol and the subsequent oxidation of the diphenolic product to the resulting quinone (1). A series of reactions occurs under the concomitant reduction of molecular oxygen to water. The quinone product is a reactive precursor for the synthesis of melanin pigments. Tyrosinase, which is contained in vegetables, fruits, and mushrooms, is a key enzyme in the browning that occurs upon bruising or long-term storage. In mammals, the enzyme is responsible for skin pigmentation abnormalities, such as flecks, and defects (2). Recently, the enzyme was reported to be linked to Parkinson’s and other neurodegenerative diseases (3, 4). Thus, tyrosinase is quite significant in the fields of medicine, agriculture, and industry. In the cosmetic industry, the development and screening of potent inhibitors of tyrosinase are especially attractive. However, the determination of the three-dimensional structure of tyrosinase has never been accomplished until now.

Tyrosinase is classified into the type-3 copper protein family, as are catechol oxidase and the respiratory pigment hemocyanin. During the catalytic reaction, the type-3 copper center of tyrosinase exists in three redox forms (1). The deoxy-form [Cu(I)-Cu(I)] is a reduced species, which binds oxygen to give the oxy-form [Cu(II)-O₂-Cu(II)]. In the oxy-form, molecular oxygen is bound as peroxide in a μ-η⁴:η¹ side-on bridging mode, which destabilizes the O-O bond and activates it. The met-form [Cu(II)-Cu(II)] is assumed as a resting enzymatic form, where Cu(II) ions are normally bridged to a small ligand, such as a...
water molecule or hydroxide ion.

Catechol oxidase oxidizes ortho-diphenols to the corresponding quinones but lacks monooxygenase activity. Hemocyanin acts as an oxygen carrier in arthropods and mollusks. Since the crystal structure of tyrosinase has been undetermined until now, the catalytic mechanism at the atomic level has not yet been elucidated. To build structural models of tyrosinase, it would be practical to reference the structural information from the type-3 copper protein members (5-11). As a common feature in the active site of these structures, each of two closely spaced copper ions is coordinated by three His residues through $\text{N}^\text{H}$ nitrogen atoms. Although the active centers of the type-3 proteins are similar in both their overall structure and their ability to bind to molecular oxygen, their enzymatic functions differ. This is believed to result from a variation in the substrate-binding pocket or the accessibility of the substrate to the active site.

Many bacteria in the genus *Streptomyces* produce a melanin-like pigment (12-17). We have cloned a melanin-synthesizing gene from *Streptomyces castaneoglobisporus* HUT 6202, which forms an operon composed of two cistrons (12, 13): an open reading frame (orf) consisting of 378 nucleotides, designated orf378, was located just upstream of the tyrosinase gene designated *tyrC*. An ORF378-like protein, for example, ORF438 in *S. antibioticus*, is commonly present in some *Streptomyces* strains, and it has been suggested that it acts as a Cu(II)-carrier protein (18, 19). In fact, when the *S. castaneoglobisporus* tyrosinase is overexpressed using an *E. coli* host-vector system (20), the co-expression of ORF378 is indispensable to obtain tyrosinase that has catalytic activity: the transport of Cu(II) ions to the catalytic center of tyrosinase, which is mediated by ORF378, may be necessary to convert tyrosinase to its active form. In the present study, we call ORF378 a “caddie” protein because it carries cupric ions for tyrosinase.

To study tyrosinase crystallographically, much effort was given to achieving the crystallization of the active tyrosinase; however, as a result of the high hydrophobicity of tyrosinase, the crystallization has never been successful. In fact, the crystallization of membrane proteins, which are difficult to dissolve at high concentration, has been unsuccessful without the help of an antibody specific to the protein (21-24). In the present study, we succeeded in crystallizing tyrosinase with the help of the caddie protein ORF378. We determined the crystal structure of tyrosinase in a complex with ORF378 by the multiple isomorphous replacement method including the anomalous scattering effect (MIRAS). Furthermore, we obtained the met-form of Cu(II)-bound tyrosinase complexed with ORF378 by soaking the native crystal in a CuSO$_4$ solution and that of its deoxy- and oxy-forms by soaking the met-form crystal in a solution containing NH$_2$OH and H$_2$O$_2$, respectively. In the solution state, ORF378 is liberated from the complex by the addition of Cu(II); however, in the crystalline state, it is not. This might be due to the molecular packing effect in the crystal. In the present study, we show the structure of tyrosinase, which was determined as the first crystal structure among all tyrosinases from prokaryote and eukaryote.

**Materials and Methods**

**Crystallography of copper-free tyrosinase complexed with ORF378** - The expression and purification of the complex have been described previously (20). Before crystallization, the protein solution was dialyzed against a 20 mM Tris-HCl buffer (pH 7.9) containing 0.2 M NaCl and concentrated to 10 mg/ml using the Macrocep 3K (PALL). Crystals of tyrosinase in complex with ORF378 were obtained by the sitting-drop vapor diffusion method using two different crystallization conditions. A precipitant solution containing 20% PEG 4000, 0.1 M ammonium chloride, and 0.1 M Na-Hepes (pH 7.0) generated monoclinic $P2_1$ crystals with unit cell dimensions of $a = 58.9$ Å, $b = 93.2$ Å, $c = 65.6$ Å, and $\beta = 93.8^\circ$, with two complexes per asymmetric unit. On the other hand, a precipitant solution containing 20% PEG 3350, 0.2 M sodium nitrate, and 0.1 M Na-Hepes (pH 6.5) generated orthorhombic $P2_12_12$ crystals with unit cell
dimensions of \(a = 64.7\ \text{Å}, b = 96.7\ \text{Å},\) and \(c = 54.6\ \text{Å},\) with one complex per asymmetric unit. Since orthorhombic crystals diffracted to higher resolution than monoclinic ones and could be easily prepared by the micro-seeding method, later crystallographic analysis was mainly conducted using orthorhombic crystals.

All X-ray data were collected while the crystals were cooling under a nitrogen gas stream (100 K). Before flash cooling, crystals were soaked in a cryoprotectant solution containing a 5-15\% higher concentration of PEG 3350 or PEG 4000 than the precipitant solution. Moderate-resolution data of the native crystals (monoclinic and orthorhombic) and heavy-atom derivatives (orthorhombic) were collected using a high-speed Rigaku R-AXIS VII imaging plate detector with CuK\(\alpha\) radiation generated by Rigaku FR-E SuperBright. The diffraction intensities on the imaging plate were integrated and scaled by the program CrystalClear. Very high-resolution data of the orthorhombic crystal up to 1.20 \(\text{Å}\) were collected using synchrotron radiation adjusted to a wavelength of 0.6199 \(\text{Å}\) from the station BL41XU at SPring-8, Japan. Since saturation of the diffraction intensity occurred in the lower-resolution bin, a second data set of the same crystal was collected using an attenuated beam, and the two data sets were then merged. The diffraction intensities were integrated on the CCD camera (Mar research) using the program Mosflm, and the scaling was done using the program Scala equipped in the CCP4 program suite (25).

The phase problem was solved using the SHARP program (26). Heavy-atom derivatives were prepared by soaking the crystal in a reservoir solution containing each heavy-atom reagent. Two derivatives (Hg and Au) were used for the phasing. The initial phase set was obtained using MIRAS, and, subsequently, the density was modified using the solvent-flattening and histogram-matching method. The resulting Fourier map enabled us to build an initial model using the program Xfit in the XtalView software package (27). The model was refined by a combination of the simulated annealing (28) and conventional restrained refinement methods (29) using the CNS program (30). A subset of 5\% of the reflections was used to monitor the free R-factor \(R_{\text{free}}\) (31). Each refinement cycle included refinement of the positional parameters and individual isotropic \(B\)-factors, correction using the flat bulk solvent model, revision of the model using the omit map, and addition of solvent molecules. When the \(R\)-factor fell below 20\%, the native data were changed to the 1.20 \(\text{Å}\) resolution data. Further rebuilding, addition of solvent molecules, modeling of an alternative conformation, and refinement using the program SHELXL-97 (32) yielded the current model. The current model contains two protein subunits, 393 water molecules, and 4 nitrate ions. In the tyrosinase part, the N-terminal residue and the C-terminal region including the artificially added polyhistidine tag were invisible in the electron density map. Therefore, the current tyrosinase model consists of Thr\(^2\)-Leu\(^{274}\) residues. On the other hand, the Ala\(^{40}\)-Ala\(^{50}\) and Gly\(^{71}\)-Pro\(^{122}\) residues in ORF378, which explain only 52\% of the total residues (73\% of the residues in the mature portion), can be confirmed by the electron density map. The high flexibility of the leader peptide domain (Met\(^1\)-Ala\(^{40}\)) is not surprising, as it is deleted after the secretion in the original cell but does not function in the \textit{E. coli} system. Similarly, a large portion of the invisible C-terminal region contains the artificially added polyhistidine sequence. But, the flexibility of the Arg\(^{60}\)-Gly\(^{70}\) residues in ORF378 may be associable with the transportation of copper ions, as discussed later. No residues, except Phe\(^{103}\) and Asn\(^{188}\) of tyrosinase, are in the forbidden region of the Ramachandran plot (33). Three Pro residues (Pro\(^{169}\), Pro\(^{253}\), and Pro\(^{353}\)) in tyrosinase form \textit{cis} peptide bonds to each previous residue.

The structure of the tyrosinase complexed with the ORF378 in the monoclinic crystal was determined by the molecular replacement method using the program Amore in the CCP4 program suite (25). The start model used was a complex determined in the orthorhombic cell. The initial coordinates of the two complexes were obtained by rotation and translation function analyses, and refinement was then performed using the program CNS (30). Each complex
structure in the final model is almost the same as the previous structure refined in the orthorhombic cell. Most importantly, the undefined regions in the orthorhombic crystalline system remain uncertain in the monoclinic one. This indicates that the flexibility of the regions represents a molecular feature but is not derived from the molecular packing effect in the crystalline lattice. Details of the data collection, phasing, and refinement statistics are shown in Table I.

Crystallography of the met-form of Cu(II)-bound tyrosinase complexed with ORF378 - To investigate the copper-binding site in the model and the structural change caused by Cu(II) incorporation, we soaked the crystal in the reservoir solution containing 1 mM CuSO₄ for the given times. Soaking in Cu(II) for more than 18 hr successfully resulted in the complete incorporation of the ion to the active site. Two Cu(II) ions (Cu⁰ and Cu⁶⁺) were clearly defined in the active site of tyrosinase (Figs. 1B and 1C): Cu⁰ binds to the Ne nitrogen atoms of His¹⁹⁰, His¹⁹⁴, and His²¹⁶, and Cu⁶⁺ binds to those of His¹⁸⁰, His¹⁸⁴, and His²¹⁶. Interestingly, the distance between the two Cu(II) ions in the catalytic center gradually shortened as the soaking time increased. By Cu(II)-soaking for approximately 60 hr, the distance converged to a constant value of 3.3 Å. Until now, only two crystal structures are available for the met-form of type-3 copper proteins: one is the subunit-II of Limulus polyphemus hemocyanin (PDB code: 1LL1), and the other is Ipomoea batatas catechol oxidase (PDB code: 1BT3) (11). In the first case, the electron density was assigned to the presence of two bridging water molecules, while the presence of only one water molecule was assumed in the latter case. In our experiments, Cu(II)-soaking for less than 60 hr gave a 1BT3-like met-form, whereas soaking for a longer time gave a 1LL1-like met-form. Hereafter, we name the former and the latter as met-forms I and II, respectively. Moreover, two to three additional Cu(II) ions were identified, since the B-factors were extremely low when they were assumed to be water. One of the additional ions, called the third Cu(II) ion in the text, binds to the Ne atom of His⁸² of ORF378 and is most frequently identified in the Cu(II)-bound structures obtained by soaking in Cu(II) solution for a short time (less than 80 hr). With the increase of the soaking time, the third Cu(II)-binding site was partially occupied, and the fourth Cu(II)-binding site alternatively emerged. With further soaking, the Cu(II) ions were found only in the fourth Cu(II)-binding site. The fourth Cu(II) ion binds to the Ne atom from His⁸² and His⁹⁷ of ORF378 and the sulfur atom from Met⁸⁴ of ORF378 similarly to the type-1 copper-binding protein. The fifth Cu(II) ion binds to the Ne atom of His¹⁸⁰ of tyrosinase, and the sixth locates in the fully solvent region. The fifth and sixth Cu(II) ions are not constantly found among the Cu(II)-soaked structures. Details of the data collection and refinement statistics are shown in Table I.

Crystallography of the copper-bound tyrosinase complexed with ORF378 prepared by soaking in a solution containing the redox reagents - To determine the structure of deoxy-form of tyrosinase, we anaerobically soaked the crystal in a reservoir solution containing 10 mM NH₂OH for 10 min, after soaking in 1 mM CuSO₄ for 69 hr. Soaking in NH₂OH for about 5 min changed the crystal color from pale blue to pale yellow. Further soaking for more than 20 min generated cracks in the crystal, and the quality of the diffraction data became worse. On the other hand, the incorporation of a copper ion was not observed by soaking the crystal in a reservoir solution containing both 1 mM CuSO₄ and 10 mM NH₂OH for 40 hr. This means that, although the Cu(II) ion is introduced into the active site of tyrosinase, the Cu(I) ion is not. The distance between Cu⁰ and Cu⁶⁺ at the active site in the deoxy-form was clearly longer than that in the met-form II. Since the electron density for a bridging solvent atom is rather strong, it is not likely to be a water molecule (Fig. 1D). However, we assumed that the atom was a water molecule because a more proper assignment was impossible.

On the other hand, we aerobically soaked the crystal in a reservoir solution containing 10 mM H₂O₂ for 20 min after soaking in 1 mM CuSO₄ for 76 hr. After 5 min of soaking, the crystal color changed...
from pale blue to pale green. After additional soaking for more than 30 min, several cracks were observed in the crystal, and the quality of the diffraction data was worse. A bridging solvent atom might be a peroxide ion because the assignment gave clearer electron density than the assignment of two water molecules seen in the met-form II (Fig. 1E). Therefore, the current structure is suggested to be the oxy-form of tyrosinase in complex with ORF378. Details of the data collection and refinement statistics are shown in Table I.

RESULTS AND DISCUSSION

Overview of the structure - We determined the first crystal structure of tyrosinase, which is complexed with the caddie protein ORF378, at 1.2-1.8 Å resolution (Table I). Tyrosinase, complexed with ORF378, is ellipsoid in shape and has dimensions of 40×55×60 Å (Fig. 2).

Tyrosinase takes α-helical structures with the core of the enzyme, which is formed by a four-helix-bundle (α2, α3, α6, and α7 helices). The catalytic dinuclear copper center is lodged in the helical bundle (Fig. 2). Each of the two copper ions in an active site is coordinated by three His residues (Figs. 1B-1E), which are derived from the four helices of the α-bundle except His54. One copper ion (designated CuA) is coordinated by His38, His54, and His63. His38 and His63 are located in the middle of α2 and α3, respectively. The second copper ion (CuB) is coordinated by His190, His194, and His216. The residues His190 and His194 are at the beginning and in the middle of α6, respectively, and His216 is in the middle of α7. This dicopper center is located at the bottom of the large concavity as a putative substrate-binding pocket (Fig. 3A), which is formed by the hydrophobic residues. In addition to the helical structure, tyrosinase has a few β-structures, as judged from the backbone torsion angles. In these, only the N- and C-terminal β-strands form a sheet structure.

Although the amino acid sequence of tyrosinase respectively has only 25.3% and 26.0% identities with those of the Ipomoea batatas catechol oxidase (11) and the odg domain of the Octopus dofleini hemocyanin (10), which have a type-3 copper center (Fig. 4), its overall structure is quite similar to theirs (Fig. 5A). The Rms value of the positional differences of the main-chain atoms in tyrosinase and the Ipomoea catechol oxidase is 0.94 Å for 138 matched residues, and that in tyrosinase and the Octopus hemocyanin is 1.34 Å for 172 residues. Among these three proteins, a high degree of conservation is observed in the core domain composed of the α-bundle. The tyrosinase and hemocyanins from Panulirus interruptus (5, 6) and Limulus polyphemus (7-9) show no significant homology and no resemblance in their structures, but the catalytic core domains of these proteins are superimposable.

ORF378, a “caddie protein” that covers the molecular surface of tyrosinase, seems to prevent substrate binding to the active site of tyrosinase (Fig. 3B). This could imply that the folding of the catalytic center of tyrosinase occurs at an earlier step than that for the other parts of the enzyme in the process of maturation. ORF378 has one six-stranded β-sheet, largely twisted at strand β4, and one α-helix, which is surrounded by a twisted sheet. ORF378 has no sequence similarity with proteins structurally determined. According to the results obtained using the DALI program (34), however, ORF378 has a significant structural similarity with the SH2 domain (Fig. 5B), which is contained in the proteins with respect to the signal transduction, such as a growth factor-bound protein (35), tyrosine kinase (36, 37), tyrosine phosphatase (38), a T-cell signal transduction molecule (39), and phospholipase C (40). However, the typical SH2 domain has five-stranded β-sheets and two α-helices flanking the sheet. Since the N-terminal α-helix in the SH2 domain is replaced by a β1-strand in ORF378, the latter protein consists of a six-stranded β-sheet. Judging from the similarity between ORF378 and the SH2 domain, these proteins might have evolved from the same ancestor. ORF378 and the SH2 domain have a key process in common with tyrosine: ORF378 assists in the folding of tyrosinase, which is necessary to metabolize tyrosine toward melanin synthesis, and some SH2 domain-harboring proteins catalyze the phosphorylation or dephosphorylation of the
tyrosine residue.

Interaction between tyrosinase and ORF378 -
The decreased molecular surface area on the
complex formation between tyrosinase and
ORF378 is calculated to be 980 Å², which is
apparently lower than that caused by the
normal dimer formation, suggesting that the
interaction between both the proteins is
considerably weak. This conclusion is
reached on the basis of the observation that
several residues in the interaction surface of
the copper-free tyrosinase/ORF378 complex,
such as Ile42, His54, and Arg140 of tyrosinase
and Val94 of ORF378, take disordered
conformations. In addition, amino acid
residues, defined in the electron density, are
only 52% of the total residues in ORF378.

The interaction between tyrosinase and
ORF378, which is mediated by Tyr98 from
ORF378 (Fig. 3B). The side-chain of Tyr98
is accommodated in the substrate-binding
pocket of tyrosinase. The phenol ring is
stacked with the imidazole ring of His54 of
tyrosinase, which is one of the CuA ligands
(Fig. 6A). In addition, its phenolic hydroxyl forms a hydrogen bond with solvent
atoms (water molecule or peroxide ion),
which form a bridge with CuA and CuB in
the active center of tyrosinase (Figs. 7B-7E).
Furthermore, when compared to the catechol
oxidase in a complex with a potent inhibitor,
phenylthiourea, the Tyr98 ring is aligned
perfectly with the aromatic ring of the
inhibitor (Figs. 6A and 6B). These results
suggest that Tyr98 of ORF378 functions as a
competitive inhibitor to the substrate tyrosine.

Structural change of the tyrosinase active site -
We analyzed the structural change on five
forms of tyrosinase complexed with ORF378:
one is a copper-free form, and the others are
two met-forms (I and II) and the oxy- and
dehydroxy-forms of the copper-bound complex
(Figs. 1 and 7). Cu(II) ions are observed to
release ORF378 from the complex in solution
but not in crystal. This may be due to the
molecular-packing effect in the crystal, which
prevents the dissociation of ORF378.

However, a slight rotational movement of
ORF378 is observed by the addition of Cu(II)
ions in the crystalline state. This movement
is likely to exhibit the dissociation process of
ORF378 from the complex. Structural
changes at the active center of tyrosinase are
also observed (Fig. 7). In the copper-free
tyrosinase, the side-chain of His54, as one of
the CuA ligands, is flexible, as proved by the
presence of the disordered structure (Fig. 1A).

The flexibility of His54 may be caused by the
absence of a thioether bond, as found in
catechol oxidase (Fig. 6B) and hemocyanin
(Fig. 6C). On the other hand, the side-chain
is more rigid in the copper-bound tyrosinase
(Figs. 1B-1E). However, even in the
copper-bound structures, the His54 atoms
have higher B-factors than those of the other
five copper ligand residues. Due to the high
mobility of His54, CuA is less stable than CuB.

By soaking the tyrosinase complexed
with ORF378 in a CuSO₄ solution for over
18 hr, the met-form was obtained (Figs. 1B
and 1C). We observed that the distance
between the two Cu(II) ions present in the
met-form was gradually shortened to 3.3 Å
by increasing the soaking time. However,
when the soaking time was less than 60 hr,
the time at which the met-form I is formed,
the corresponding distance is approximately
3.9 Å. As a result, the Cu(II) ions were
observed to form a bridge with one water
molecule. On the other hand, under the soaking condition, which gives the met-form II where the distance was converged to 3.3 Å, the copper ions formed bridges with two water molecules. For tuning up the catalytic reaction of tyrosinase, it may be necessary that the distance between CuA and CuB in the catalytic center is flexible. The conversion of the met-form I to the met-form II induced a more stable conformation at the catalytic center. In this study, we examined the flexibility of His54 and CuA and CuB, with respect to the mean B-factors for all protein atoms. As a result, we found that the flexibility of CuB is almost constant, whereas that of CuA and His54 is largely varied. In the met-form II, CuA and His54 have structural stability, like CuB.

In the met-form I (Fig. 7B), six His ligands and a bridging solvent complete the four-coordinated trigonal coordination for both Cu(II) atoms with His54 and His190 in the apical position of the coordination polyhedron for CuA and CuB, respectively. On the other hand, in the met-form II (Fig. 7C), CuA is in the distorted tetragonal-pyramidal coordination with His38, His54, and two waters in the equatorial position and His54 in an axial position. Since a sixth coordination site locates in the interior of the enzyme, it is not bound to a solvent atom. CuB is also in distorted tetragonal-pyramidal coordination with His194, His190, and two waters in the equatorial position and His190 in an axial position, but the vacant sixth coordination site is located in the substrate-binding pocket.

We formed the deoxy-form of the Cu(I)-bound tyrosinase by the addition of NH2OH under anaerobiosis after Cu(II)-soaking for 69 hr (Fig. 1D). Upon soaking for 69 hr, the time at which the met-form II is formed, the distance between two Cu(II) ions is at a minimum. However, the distance in the deoxy-form increased to 4.1 Å. The space expanded between the copper ions may be useful to accommodate dioxygen in the catalytic center. A bridging solvent is equidistant (2.3 Å) from CuA and CuB, and the coordination geometry for the copper ions in the deoxy-form is similar to that in the met-form I (Fig. 7D). This is different from what was observed in the active center of the deoxy-form of catechol oxidase (11): that is, the distance between the bridging solvent and CuA is shorter than that between the solvent and CuB. Furthermore, the catalytic center in the deoxy-form is structurally stable, like that of the met-form II. We point out that although the distance between CuA and CuB of the catalytic center in the met-form I is approximately the same as that in the deoxy-form, the stability of the former’s catalytic center is obviously lower than that of latter’s one.

In contrast, the oxy-form of the Cu(II)-bound tyrosinase was formed by the addition of hydrogen peroxide (H2O2) after Cu(II)-soaking for 76 hr (Fig. 1E). Although it is difficult to distinguish the oxy-form from the met-form II by the electron density, the distance between the two bridging-solvent atoms of the former form is somewhat shorter than that of the latter one. Therefore, we assumed that the density for the bridging-solvent is a peroxide ion. In fact, the oxy-form of tyrosinase formed by the addition of H2O2 was confirmed by its typical UV-vis spectrum at 350 nm in solution. In the oxy-form structure, the peroxide ion binds in a bridging side-on \( \mu-\eta^2: \eta^1 \)-binding mode (Fig. 7E). The distance between the Cu(II) ions in the oxy-form is 3.4 Å, and the coordination geometry is almost identical to that in the met-form II. However, dicopper center of the oxy-form is more unstable than that of the met-form II. The difference electron density around the His54 side-chain indicates that the residue is disordered, as seen in the copper-free form. This observation may correlate with the high reactivity of the oxy-form of tyrosinase.

Structural evidence that ORF378 functions as a Cu(II) transporter for tyrosinase - We have suggested that ORF378 serves as a transporter of Cu(II) ions to the catalytic site of apotyrosinase (copper-free tyrosinase). This study presents structural evidence that ORF378 plays a role in the transport of two Cu(II) ions to apotyrosinase, as demonstrated by the X-ray diffraction analysis of about 30 crystals of copper-bound tyrosinase complexed with ORF378 prepared under different soaking conditions. The structural analysis of these crystals demonstrated that the complex has Cu(II)-binding sites in
addition to a catalytic center in which two Cu(II) ions are accommodated.

The third Cu(II) ion, which is most frequently identified in the crystals soaked in a Cu(II) solution for less than 80 hr, binds to the N€ atom from His\textsuperscript{82} of ORF378. This site is on the molecular surface of ORF378 (Fig. 8B), and the N\=O atom of the His\textsuperscript{82} side-chain forms a hydrogen-bond with the N\=O atom of His\textsuperscript{97} of ORF378 by the mediation of the nitrate ion, which is derived from a precipitant solution. However, by soaking crystals in a CuSO\textsubscript{4} solution for a longer time, the third binding site for the Cu(II) ion was partially occupied, and the fourth Cu(II) ion, which binds to the N\=H atoms from His\textsuperscript{82} and His\textsuperscript{97} of ORF378, alternatively emerged (Fig. 8C). The fourth Cu(II) ion was also found to bind to the sulfur atom from Met\textsuperscript{84} of ORF378 in a manner similar to that of the type-1 copper-binding protein. To make the fourth Cu(II)-binding site, the imidazole rings of His\textsuperscript{82} and His\textsuperscript{97} must be rotated around the bond between their C\=N and C\=O atoms. By soaking crystals in a CuSO\textsubscript{4} solution for 6 months, the fourth binding site for the Cu(II) ion was completely occupied, and the third binding site was emptied. Thus, we could observe the movement of the additional Cu(II) ion in a time-dependent manner.

The imidazole ring of His\textsuperscript{97} of ORF378 is located near the side-chain of His\textsuperscript{54} of tyrosinase. Especially, in the copper-free tyrosinase complexed with ORF378, the alternative His\textsuperscript{54} side-chain of tyrosinase is closer to the His\textsuperscript{97} side-chain of ORF378 (Fig. 8A). The clustered His residues, which extend from the molecular surface of ORF378 to the active center of tyrosinase, may bind to Cu(II) atoms in the transferring process of the metal ions to the catalytic center.

It has been reported that, when His\textsuperscript{102} and His\textsuperscript{117} of \textit{S. antibioticus} ORF438, which correspond to His\textsuperscript{82} and His\textsuperscript{97} of \textit{S. castaneoglobisporus} ORF378, were replaced by Leu and Asp, respectively, the copper-transferring activity of ORF438 was reduced (19). A copper ion, which is illustrated as the third position in Fig. 3B, may first binds for transportation of copper ions to tyrosinase to His\textsuperscript{82} located on the surface of ORF378. In general, an atmosphere for metal-binding should take negative charge, whereas electrostatic potential around the third position for Cu(II)-binding is neutral. This suggests that the third position is not actually located on the surface of ORF378. ORF378 contains four His residues (His\textsuperscript{64}, His\textsuperscript{65}, His\textsuperscript{66}, and His\textsuperscript{68}), but the residues are invisible in the electron density map. Since the disordered residues are likely to be positioned near the third copper-binding site, they might function as initial copper-binding site.

**Comparison with proteins having a type-3 copper center** - Tyrosinase belongs to the protein family having a type-3 copper center, as do catechol oxidase (11) and hemocyanin (5-10). The reasons for the different functions exhibited by these proteins pose an interesting query. For example, hemocyanin plays a significant role in the transportation of oxygen into arthropods and mollusks. The protein has a domain that shields the access to the dicopper center of the protein (Fig. 6C). Since the domain interferes with the binding of the substrate, hemocyanin can play only one role, namely, that of an oxygen-transporter. The dicopper center of both catechol oxidase and tyrosinase is near the molecular surface, a useful location to ensure that a substrate can access each catalytic site (Figs. 6A and 6B).

Both tyrosinase and catechol oxidase catalyze the oxidation of \textit{ortho}-diphenols to the corresponding quinones, but the latter enzyme lacks the monooxygenase activity to convert monophenols to \textit{ortho}-diphenols. The structure and function of tyrosinase have been compared with those of catechol oxidase. One of the most attractive differences may be the vacant space of the substrate-binding pocket. The Phe\textsuperscript{261} side-chain in the active site of catechol oxidase, which is located just above the Cu\textsuperscript{A} site, seems to partially prevent substrate binding (Fig. 6B). In contrast, the substrate-binding pocket of tyrosinase has a larger vacant space above the dicopper center, if ORF378, complexed with tyrosinase, is liberated from tyrosinase (Fig. 6A).

His\textsuperscript{54} of tyrosinase, which is one of the Cu\textsuperscript{A} ligands, differs obviously from the corresponding His\textsuperscript{109} of catechol oxidase from sweet potato. In copper-free
tyrosinase, complexed with ORF378, the His\(^{54}\) residue takes two conformations: one is exactly the same as the conformation observed in the copper-bound form, but the other, a side-chain of His\(^{54}\), protrudes toward the molecular surface (Fig. 7A). In catechol oxidase (11), His\(^{109}\) is fixed by the unusual covalent bond formed between the C\(_{\beta}\) atom of His\(^{109}\) and the sulfur atom of Cys\(^{92}\) (Fig. 6B). The catalytic specificity of tyrosinase or catechol oxidase may depend on the flexibility of His\(^{54}\) or the inflexibility of His\(^{109}\), respectively.

The thioether-bond is also found in the structure of hemocyanin. In the Octopus hemocyanin (10), the thioether-bond is formed with a C\(_{\beta}\) atom of His\(^{256}\), which is the counterpart of His\(^{109}\) of catechol oxidase (Fig. 6C). However, the sulfur atom is provided from a structurally different Cys residue. As judged from the sequence similarity to the Octopus hemocyanin, tyrosinases from Neurospora crassa (45) and Agaricus bisporus (46) might have the thioether-bond in the active site (Fig. 4). The hypothesis will be confirmed by the determination of their crystal structures.

Implication for the catalytic mechanism - Tyrosinase can catalyze the ortho-hydroxylation of monophenol and the conversion of the ortho-diphenols to the corresponding quinones. Based on the crystal structure of tyrosinase obtained in the present study, we can describe the tyrosinase-specific catalytic mechanism in detail (Fig. 9). At first, a peroxide ion, which forms a bridge with two Cu(II) ions in the oxy-form of tyrosinase, acts as a catalytic base. As a result, a proton is abstracted from the phenolic hydroxyl. Subsequently, the deprotonated oxygen atom of monophenol binds to Cu\(^{1}\) at the sixth coordination site. At this time, Cu\(^{1}\) is hexa-coordinated by a tetragonal bipyramidal cage, and an ortho-carbon of the substrate approaches the peroxide ion. One of two peroxide oxygens is then added to the ortho-carbon of monophenol. This monooxygenase reaction would be accelerated by the formation of a stable intermediate, in which newly generated oxygen atoms of diphenol bind to Cu\(^{1}\). To form this state, His\(^{54}\), which is an axial ligand to Cu\(^{1}\), must be released from the current position. This assumption is derived from the flexible feature of the residue His\(^{54}\) in the copper-free and Cu(II)-bound oxy-forms. Simultaneously, His\(^{54}\) can act as a catalytic base for the deprotonation from the substrate. The resulting intermediate has the advantage of easy translation of electrons, resulting in the formation of the deoxy-form of tyrosinase and quinone.

Our proposed scheme does not fit the case of catechol oxidase (11) because the bidentate intermediate cannot be formed due to the fixed conformation of His\(^{109}\), which corresponds to His\(^{54}\) of tyrosinase, and the presence of the Phe\(^{251}\) lying just above the Cu\(^{A}\) site, which is vacant in the tyrosinase structure (Figs. 6A and 6B). There is a consensus (1, 44) that the oxy-form of tyrosinase can catalyze both the monooxygenase and oxidase reactions, while the met-form lacks the monooxygenase activity. This can be explained as follows: the bidentate intermediate formation, which is permitted only by the oxy-form, is essential for the monooxygenase reaction, but the mono-dentate intermediate, which is formed by the met-form, is sufficient for an oxidase reaction, as proposed for catechol oxidase. Therefore, some compounds, which bind to two Cu(II) ions in the bidentate form, might be a potent inhibitor of tyrosinase.

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FOOTNOTES

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The atomic coordinates have been deposited in the Protein Data Bank with accession codes 1WX5, 1WXC, 1WX3, 2AHK, 2AHL, and 1WX2 for the monoclinic and orthorhombic copper-unbound tyrosinase/ORF378 complex, met-form I, met-form II, deoxy-form, and oxy-form of copper-bound tyrosinase complexed with ORF378, respectively.

FIGURE LEGENDS

**Fig. 1. Electron density around the dicopper center of tyrosinase.** (A) Electron density map around the dicopper center of copper-free tyrosinase complexed with ORF378. To emphasize the density, $F_o-F_c$ omit map was computed after removal of six His residues participating in copper binding and one water molecule. (B and C) Electron density map around the dicopper center of the met-forms I (B) and II (C) of tyrosinase complexed with ORF378. The omit map was computed after removal of six His residues, two copper ions, and the bridging molecule. (D and E) Electron density map around the dicopper center of the deoxy- (D) and oxy-forms (E) of tyrosinase complexed with ORF378, prepared by the treatment with NH$_2$OH and H$_2$O$_2$, respectively. The omit map was computed after removal of six His residues, two copper ions, and the bridging molecule (water and peroxide ion).

**Fig. 2. Overall structure of tyrosinase complexed with ORF378.** Stereo representation of the ribbon view of the copper-bound form of the *S. castaneoglobisporus* tyrosinase complexed with ORF378. Tyrosinase and ORF378 are shown in red and blue, respectively. Four of the six identified copper ions are indicated by green spheres. Two of them are located at the dicopper center of tyrosinase, and the third copper ion is bound at the molecular surface of ORF378. The fourth copper ion exists in the interior of ORF378.

**Fig. 3. Molecular surfaces of tyrosinase (A) and ORF378 (B).** (A): area surrounded by a circle is revealed under magnification. The area contains the substrate-binding pocket of tyrosinase. Red and blue colors show the negative and positive electrostatic potential, respectively. (B): orange, cyan and green colors represent tyrosinase, ORF378 and copper ions, respectively. Tyr$^{98}$ of ORF378 is present in the substrate-binding pocket of tyrosinase.

**Fig. 4. Multiple sequence alignment of the type-3 copper proteins.** TY_SCA indicates the amino acid sequence of the *S. castaneoglobisporus* tyrosinase. Other sequences are taken from the GenBank: TY_HUM: human tyrosinase, TY_MUS: *Agaricus bisporus* (mushroom) tyrosinase, TY_NCR: *Neurospora crassa* tyrosinase, CO_POT: potato catechol oxidase, and HC_OCT: octopus hemocyanin. Structurally superimposable residues in CO_POT and
HC_OCT are shown by capital letters. The identical and conserved residues are displayed in black and gray shades, respectively. The asterisks indicate the His residues participating in copper binding.

**Fig. 5.** Structural similarity. (A) Stereo representation of the superposition of tyrosinase and structurally homologous proteins, potato catechol oxidase (PDB code 1BT1) and octopus hemocyanin (PDB code 1JS8). Red, blue, and green indicate the backbone trace of tyrosinase, catechol oxidase, and hemocyanin, respectively. The yellow sphere indicates the two copper ions in the catalytic center. To emphasize similarity, the C-terminal domain of hemocyanin is omitted from the figure. (B) Stereo-view of the superposition of ORF378 and the SH2 domain in the growth-factor-bound protein 2 (PDB code 1GRI). Red and blue indicate the backbone trace of ORF378 and the SH2 domain in the growth-factor-bound protein 2, respectively.

**Fig. 6.** The active centers of tyrosinase and of structurally homologous proteins. (A) Active center of the met-form I of tyrosinase complexed with ORF378. Carbon atoms from the residues of tyrosinase and ORF378 are shown in orange and cyan, respectively. (B) Active center of the inhibitor-bound potato catechol oxidase. Carbon atoms from catechol oxidase and inhibitor (phenylthiourea, PTU) are shown in orange and purple, respectively. (C) Active center of the oxy-form of the octopus hemocyanin. Carbon atoms from hemocyanin are shown in orange.

**Fig. 7.** Conformational changes of the active center of tyrosinase caused by copper and dioxygen binding. (A), (B), and (C) indicate the active center of the copper-free form, met-form I, and met-form II of tyrosinase complexed with ORF378, respectively. (D) and (E) indicate the active center of the deoxy- and oxy-forms of tyrosinase complexed with ORF378, respectively. The carbon atoms from the residues of tyrosinase and ORF378 are shown in orange and cyan, respectively.

**Fig. 8.** Third and fourth copper-binding sites and clustered His residues. (A) shows the partial atomic models of the copper-free tyrosinase complexed with ORF378. Ile$^{42}$ and His$^{54}$ of tyrosinase and Val$^{94}$ of ORF378 have disordered structures. The molecule bridging between His$^{54}$ and His$^{97}$ of ORF378 is a nitrate ion. (B) and (C) show the partial atomic models of the met-form of tyrosinase complexed with ORF378 prepared by soaking in a CuSO$_4$ solution for 37 hr and 6 months, respectively. In (B), the third Cu(II) binds to the N$_e$ atom of His$^{82}$ of ORF378. In (C), although the third Cu(II)-binding site was hardly occupied, the fourth Cu(II) ion, which binds to the N$_e$ atoms from His$^{82}$ and His$^{97}$ of ORF378 and the sulfur atom from Met$^{84}$ of ORF378, was observed. Carbon atoms from the residues of tyrosinase and ORF378 are shown in orange and cyan, respectively.

**Fig. 9.** Structure-based catalytic mechanism of tyrosinase. The oxy-form of tyrosinase catalyzes the conversion of monophenol to the corresponding quinone through the ortho-diphenol formation. In this scheme, His$^{54}$ is released from the Cu$^+$ site, resulting in the formation of the bidentate intermediate. The met- and oxy-forms of tyrosinase can catalyze the conversion of ortho-diphenol to the corresponding quinone. This reaction should progress similarly to that of catechol oxidase.
| Data collection | Native-1 | Hg | Au | Native-2 | Native-3 | Met-I | Met-II | Deoxy | Oxy |
|----------------|---------|----|----|---------|---------|------|-------|-------|-----|
| Space group    | P2_12   | P2_12 | P2_12 | P2_12 | P2_12 | P2_12 | P2_12 | P2_12 | P2_12 |
| Soaking conditions | Reagent   | EMTS<sup>a</sup> | KAu(CN)<sub>2</sub> | CuSO<sub>4</sub> | CuSO<sub>4</sub> | CuSO<sub>4</sub>/NH<sub>2</sub>OH | CuSO<sub>4</sub>/H<sub>2</sub>O<sub>2</sub> |
| Concentration (mM) | 10       | 1   | 1   | 1       | 1/10    | 1/10  | 1/10  | 1/10  | 1/10 |
| Time           | 24 hr    | 21 hr | 37 hr | 6 month | 69 hr/10 min | 76 hr/20 min | 69 hr/10 min | 76 hr/20 min |
| X-ray          | R-AXIS VII | R-AXIS VII | R-AXIS VII | BL41XU | BL41XU | R-AXIS VII | R-AXIS VII | R-AXIS VII |
| Wavelength (Å) | 1.54184  | 1.54184 | 1.54184 | 0.9000 | 0.9000 | 1.54184 | 1.54184 | 1.54184 |
| Resolution (Å) | 1.70     | 1.80  | 2.20 | 2.62    | 1.20    | 1.71  | 1.60  | 1.00  |
| Unique reflection | 55,522  | 32,466 | 18,150 | 46,512 | 104,674 | 78,955 | 37,596 | 47,758 | 32,279 |
| Completeness (%) | 99.1 (97.2) | 99.5 (99.2) | 99.6 (99.8) | 99.6 (99.8) | 97.8 (99.2) | 97.3 (99.2) | 97.7 (99.4) | 99.1 (99.8) |
| R<sub>merge</sub> (%)<sup>a, b</sup> | 3.6 (1.0) | 5.0 (1.0) | 9.6 (2.0) | 12.5 (2.0) | 7.4 (2.0) | 3.7 (2.0) | 6.7 (2.0) | 5.0 (2.0) |
| R<sub>i</sub> (%)<sup>c</sup> | 10.8 (2.0) | 13.5 (5.0) | 7.3 (2.8) | 7.3 (3.4) | 5.9 (2.0) | 6.4 (2.4) | 15.3 (5.7) | 11.6 (2.5) |
| Phasing         | Heavy atom sites | 2 | 3 |
| Isomorphous     | 8.651     | 0.749 |
| Anomalous       | 8.250     | 0.225 |
| Refinement      | Resolution (Å) | 30.2–2.0 | 30.4–2.0 | 30.1–2.0 | 30.1–2.0 | 30.1–2.0 | 30.1–2.0 |
| Used reflections | 45,936    | 100,937 | 75,945 | 39,981 | 44,216 | 32,193 |
| R<sub>work</sub> (%) | 20.1     | 18.0  | 19.1 | 19.4    | 21.5   | 21.0  |
| R<sub>free</sub> (%) | 29.1     | 20.8  | 19.1 | 19.4    | 21.5   | 21.0  |
| Rms deviations from ideality<sup>c</sup> | 0.006     | 0.013 | 0.006 | 0.007   | 0.008  | 0.009 |
| Atomic B (Å<sup>2</sup>) | 1.2      | 2.3   | 1.3  | 1.2     | 1.5    | 1.3  |

<sup>a</sup>Values in parentheses are for the highest resolution bin.  
<sup>b</sup>R<sub>merge</sub> = \( \frac{\sum|I_i - \langle I \rangle|}{\sum I} \), where \( I \) is the observed intensity and \( \langle I \rangle \) is the mean value of \( I \).  
<sup>c</sup>Ideal values are defined by Engh & Huber (47).  
<sup>d</sup>EMTS means ethylmercuricthiosalicylic acid.
Figure 1
Figure 2
Figure 3
Figure 4
Figure 6
Figure 8
Crystallographic evidence that dinuclear copper center of tyrosinase is flexible during catalysis
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