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Keywords
ascomycete; C-terminal plug; laccase; proton transfer; redox potential

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(Received 8 March 2011, revised 20 April 2011, accepted 27 April 2011)

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Structural data are available in the Protein Data Bank database under the accession numbers 3PPS and 2VDZ

Structured digital abstract
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Laccases have recently been extensively studied by X-ray crystallography. The first complete laccase structures were published in 2002, and crystal structures from at least 10 different organisms have now been reported. Most of the structures are from sporophoral basidiomycota fungi: Coprinus cinereus [10], Trametes versicolor [11,12], Rigidoporus lignosus [13], Cerrena maxima [14], Coriolus zonatus [15], Lentinus tigrinus [16], Trameter trogii [17], Trametes hirsuta [18], and Coriolopsis gallica. Structures of bacterial laccases or multicopper oxidases are also available, including the spore coat protein A from Bacillus subtilis [20], a copper efflux operon from E. coli [21], and the more recently published novel two-domain laccases [22–24]. The other phylum of the fungi, the Ascomycota or sac fungi, is much less studied, and only the crystal structure of Melanocarpus albomyces laccase (MaL) has been solved [25].

The fold of the three-domain laccases is composed of three β-barrel domains that are assembled around a catalytic dinuclear copper–copper pair, and are paramagnetic, and can be detected by EPR spectroscopy. The T1 copper is responsible for the strong absorption at 600 nm. The T1 and T2 coppers form the phenolic substrate-binding site of the enzyme, whereas the T2 and the T3-pair coppers form ortho trogii [17], Trametes hirsuta [18], and Coriolopsis gallica. Structures of bacterial laccases or multicopper oxidases are also available, including the spore coat protein A from Bacillus subtilis [20], a copper efflux operon from E. coli [21], and the more recently published novel two-domain laccases [22–24]. The other phylum of the fungi, the Ascomycota or sac fungi, is much less studied, and only the crystal structure of Melanocarpus albomyces laccase (MaL) has been solved [25].

The fold of the three-domain laccases is composed of three β-barrel domains that are assembled around two catalytic copper-binding sites (Fig. 1). The active sites are formed by four copper cations, which are divided into three different types – type 1 (T1), type 2 (T2), and type 3 (T3) – by their characteristic spectroscopic features. The T1 copper of the laccases and redox potentials of the substrates at pH 4.5 and pH 6.0 are provided, together with the redox potential differences (ΔEp) between the T1 coppers of the laccases and the substrates. ND, not determined.

### Table 1. Kinetic parameters for rMaL, TaLcc1, and ThL, measured with 2,6-DMP, syringic acid and methyl syringate in 25 mM succinate buffer at pH 4.5 and in 40 mM Mes buffer at pH 6.0 (25 °C). Structural formulas of the substrates are presented. Redox potentials (Ep) of T1 coppers of the laccases and redox potentials of the substrates at pH 4.5 and pH 6.0 are provided, together with the redox potential differences (ΔEp) between the T1 coppers of the laccases and the substrates. ND, not determined.

<table>
<thead>
<tr>
<th></th>
<th>2,6-DMP</th>
<th>Syringic acid</th>
<th>Methyl syringate</th>
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<tr>
<td></td>
<td>pH 4.5</td>
<td>pH 6</td>
<td>pH 4.5</td>
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<tr>
<td></td>
<td>E° = 0.53 V</td>
<td>E° = 0.40 V</td>
<td>E° = 0.57 V</td>
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<tr>
<td>rMaL (Ep = 0.48 V)</td>
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<tr>
<td>ΔEp (V)</td>
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<td>-0.09</td>
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<tr>
<td>Km (μM)</td>
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<td>9.5 ± 0.9</td>
<td>122 ± 11</td>
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<tr>
<td>Vmax (ΔA min⁻¹·nmol⁻¹)</td>
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<td>119 ± 1</td>
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<tr>
<td>TaLcc1 (Ep = 0.51 V)</td>
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<tr>
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<td>ThL (Ep = 0.78 V)</td>
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<td>ΔEp (V)</td>
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<td>Vmax (ΔA min⁻¹·nmol⁻¹)</td>
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<td>91 ± 2</td>
<td>8.3 ± 0.4</td>
</tr>
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</table>
scopic features. The T1 copper is responsible for the characteristic blue colour of these enzymes, and has strong absorption at 600 nm. The T1 and T2 coppers are paramagnetic, and can be detected by EPR spectroscopy. The T3 coppers form an antiferromagnetically coupled dinuclear copper–copper pair, and are therefore EPR silent, although these coppers cause absorbance at 330 nm. The loops surrounding the T1 copper form the phenolic substrate-binding site of the enzyme, whereas the T2 and the T3-pair coppers form the trinuclear site that is responsible for binding and reduction of the molecular oxygen. The reduction of oxygen to two water molecules requires the transfer of four electrons [26,27]. The rate-limiting step for the catalysis is apparently the transfer of the first electron from the substrate to the T1 copper in laccase. The suitability of a chemical compound as a laccase substrate depends on two factors. First, the substrate must dock at the T1 copper site, which is mainly determined by the nature and position of substituents on the phenolic ring of the substrate. Second, the redox potential ($E^\circ$) of the substrate must be low enough, as the rate of the reaction has been shown to depend on the difference between the redox potentials of the enzyme and the substrate ($\Delta E^\circ$) [28–31].

This study presents the crystal structure of a novel laccase (TaLcc1) from the ascomycete fungus *Thielavia arenaria* [32]. The molecular mass of the enzyme is ~80 kDa (based on SDS/PAGE), and it shows multiple bands in IEF. The pH optimum is 5.5, but the enzyme retains substantial activity at pH 7. The three-dimensional structure of TaLcc1 shows both similarities to and differences from the analogous structures of the ascomycete laccase (asco-laccase) MaL, thus giving a comprehensive view of the structure and function of asco-laccases.

**Results and Discussion**

**Overall structure**

The crystal structure of TaLcc1 was solved to 2.5-Å resolution from pseudomerohedrally twinned crystals by molecular replacement, using the recombinant MaL expressed in *Trichoderma reesei* [rMaL; Protein Data Bank (PDB) code 2Q9O] [33] as a model. The real space group was $P2_1$, and it was mimicking orthorhombic ($\beta = 90.3^\circ$). This led us to the solution with four molecules in an asymmetric unit, with a Matthews coefficient probability of 2.61 Å$^3$/Da$^{-1}$ and a solvent content of 52.9%. Interestingly, molecules A and B (and C and D) formed a similar weak dimer as previously reported for rMaL [33]. Thus, the asymmetric unit contained two weak TaLcc1 dimers (Fig. 1).

The crystal structure of TaLcc1 contained 564 amino acids. The overall structure was similar to that of other fungal laccases, especially the only known asco-laccase structure from *M. albomyces* [25]. Protein monomers of the two asco-laccases could be superimposed with an rmsd of 0.65 Å for 558 Ca atoms. The fold is composed of three cupredoxin-like domains, called A (1–160), B (161–340), and C (340–564) (Fig. 1A,B), or sometimes referred to in the literature as domains I, II, and III. In TaLcc1, three disulfide bridges located in domain A (Cys5–Cys13), in domain B (Cys298–Cys332) and between domains A and C (Cys115–Cys545) stabilize the fold.

Most laccases are glycoproteins, with typically 3–10 glycosylation sites per monomer, although the functional role of the carbohydrates is not clear. Glycosylation has been suggested to be involved, for example, in the stabilization of the catalytic centre, giving protection against hydrolysis, and improving the...
thermostability of the enzyme [34]. On the basis of the sequence of TaLcc1, there are eight putative N-glycosylation sites (Asn89, Asn202, Asn217, Asn247, Asn290, Asn337, Asn376, and Asn396), and carbohydrate residues were found on six of these sites (Asn89, Asn202, Asn217, Asn247, Asn290, and Asn376) in our crystal structure (Fig. 1A). The carbohydrate composition slightly varied between the molecules in the asymmetric unit; however, the glycans attached to Asn89 (G1 in Fig. 1A) and to Asn202 (G2 in Fig. 1A) were consistent in all four molecules. These two glycans seem to have a clear stabilizing effect on the multidomain protein structure. The glycan on Asn89 was located alongside the C-terminal tail between all three domains (Fig. 2A), and had two hydrogen bonds with Ser180 and Asn555. The glycan on Asn202 was in the groove between the β-barrels of domains A and B (Fig. 2B), and had six hydrogen bonds, three to the main chain carbonyls at Asn6, Leu168, and Val170, and three to the side chains of Arg11, Arg71, and Tyr216. The catalytic centres were arranged in similar way as previously reported for MaL [25]. In the mononuclear centre, the T1 copper was coordinated to two ND1 atoms of His residues. The residues in the axial positions of the mononuclear centre were Leu and Ile. The trinuclear centre had two type 3 coppers (T3 and T3’), each being coordinated to three nitrogen atoms of His residues. The T2 copper was coordinated to two nitrogens of His residues. On the basis of the electron density, we refined one oxygen atom (probably a hydroxide) between the type 3 coppers in molecules B, C and D in the asymmetric unit of the crystal structure. However, the electron density among the coppers in molecule A was stronger than in the other molecules, and had a slightly elliptic shape towards the T2 copper. On the basis of these observations, we decided to refine a dioxygen molecule at this site. Another oxygen atom (most likely a hydroxide or a water molecule) was coordinated to the T2 copper on the opposite side (in the T2 solvent channel). No chloride was observed, even though the purified enzyme was in Tris/HCl buffer. In the crystal structure of MaL/rMaL, a chloride is bound to the T2 copper, whereas in other published laccase crystal structures, an oxygen atom, most likely in a hydroxide ion, is reported to be here.

**T2 solvent channel**

The water channel leading to the trinuclear centre from the side of the T2 copper, between domains A and C, can be found in all fungal laccases except in rMaL, where His98 blocks the access. The T2 cavity is surrounded by acidic Asp residues (Fig. 3), which have been suggested to provide the protons required for dioxygen reduction in Fet3p multicopper oxidase [35]. In our TaLcc1 structure, His98 was replaced by Arg99, orientated such that it formed the surface of the solvent channel. Therefore, the access through the channel was unhindered in TaLcc1 (Fig. 3A). It is possible that His98 in rMaL may also rotate to another conformation to open the T2 channel (Fig. 3B). On the basis of protein structure libraries, the ‘open conformation’ would be the second most favoured conformation. On the other hand, we did not observe any trace of the movements on the His98 residue in our MaL/rMaL.
crystal structures. In our near-atomic crystal structure of rMaL, His98 exists in its oxidized form, possibly because of the oxidative stress [33]. The oxidation of the His residues probably affects its ability to change the side chain conformation, which may have implications for the catalytic function of this laccase.

Role of the C-terminus

In addition to the T2 solvent channel, a so-called T3 solvent channel is generally reported in basidiomycete laccases. The T3 solvent channel gives the solvent access to the trinuclear centre. However, the channel is blocked by the C-terminal end of the amino acid chain in MaL/rMaL [25,33]. Similarly, in the structure of TaLcc1, the last four amino acids (DSGL) penetrate inside the channel. This is known as a C-terminal plug or a C-terminal tail. On the basis of the crystal structure, the mature TaLcc1 enzyme lacks 40 residues at the N-terminus and 13 residues at the C-terminus as compared with the coded sequence. It has been previously reported that the gene sequence of rMaL codes for 623 residues, but the secreted mature enzyme lacks 50 residues at the N-terminus and 14 residues at the C-terminus [36]. The C-terminal extension containing the last 14 (13 in TaLcc1) residues is post-translationally cleaved, and thus the active forms of both enzymes have DSGL as the last four amino acids penetrating into the channel.

The C-terminal processing has been reported for asco-laccases of different origins [37–39]; furthermore, the C-terminus of the mature asco-laccases is highly conserved, suggesting that the DSGL/V/I plug is most likely a characteristic feature of asco-laccases. Basidiomycete laccases do not generally have this type of C-terminus. However, R. lignosus laccase (RL) [13] has a C-terminal DSGLA sequence. Among the known basidiomycete laccases, RIL is phylogenetically the closest to asco-laccases. Although the last amino acids of RIL are not visible in the crystal structure, it is unlikely that the C-terminus of RIL would be long enough to form such a plug, as the last visible amino acid (Asn494) is located on the surface of the molecule and is rather far away from the trinuclear site. Therefore, the C-terminal sequence of RIL might be more of an evolutionary relic than a functional feature of the enzyme.

The actual role of the C-terminus in asco-laccases has been unclear. However, we have recently shown that a mutation in the C-terminus of rMaL affects both the activity and the stability of the enzyme [40]. The Leu559 → Ala mutation greatly reduced the turnover number for ABTS, whereas the turnover number for the phenolic substrates was not significantly altered. In addition, deletion of the four last amino acids (delDSGL) of rMaL resulted in a practically inactive form of the enzyme [40]. Therefore, it is obvious that the C-terminal amino acids are critical for the function of asco-laccases. Furthermore, the C-terminal extension (the amino acids after the cleavage site) has been shown to affect the secretion process and folding of asco-laccases [41].

Very recently, studies on a CotA laccase from B. subtilis (BsL) have provided evidence for Glu498 near the T3 coppers participating in the catalytic
basidiomycete laccases are both group of asco-laccases and the conserved Asp of the C-terminus. The C-terminal carboxylate is strong enough to assist in the proton transfer is the carboxyl group of the so-called SDS gate, which is conserved in asco-laccases but not detected in basidiomycete laccases or the fungal species located between the T3 coppers. Glu498 of BsL is 4.7 Å from this oxygen (Fig. 4). It is plausible that aso-laccases use the C-terminal carboxylate in TaLcc1, whereas Pro195 and Val428 make the cavity in TaLcc1 wider in the other direction. In addition, the loop with Val428 has an additional Ile427 in TaLcc1. This function of the enzyme, possibly by promoting proton transfer [4-45]. The fungal laccase structures have no transfer in this position, but basidiomycete fungal laccases have a conserved Asp in close proximity, i.e. Asp456 in T. versicolor laccase. In asco-laccases, the group and basidiomycete laccases use the conserved Asp residue and the conserved Asp from the C-terminus. The C-terminal carboxylate assists the proton flow. Laccases from different organisms might thus have adopted different strategies to assist in the proton transfer for the dioxygen reduction.

Fig. 4. The trinuclear centres of (A) TaLcc1, (B) rMaL, (C) TvL, and (D) BsL. The distance of the putative catalytic carboxyl group from the oxygen species between the T3 coppers is shown.

Oxidation of phenolic substrates

The substrate-binding pocket of TaLcc1 is similar to that in MaL, but there are clear differences in both the group and basidiomycete laccases. In TaLcc1 (PDB code 3FU8), the corresponding residue is an Asn, and it has been suggested that this contributes to assist in the trinuclear centre [33]. In TaLcc1, Ser143, Ser511 and Asp561 form the SDS gate, which possibly assists the proton flow. Laccases from different organisms might thus have adopted different strategies to assist in the proton transfer for the dioxygen reduction.
loop resembles ‘the extended jut’ reported in LacB of *Trametes* sp., which was also suggested to be involved in substrate recognition [44].

In TaLcc1, 10 hydrophobic residues (Ala193, Leu297, Leu363, Phe371, Ile427, Val428, Leu430, Trp508, and His509) and one hydrophilic residue, Asp236, form the binding pocket. The most evident difference between MaL and TaLcc1 is in this putative catalytic amino acid: TaLcc1 has an Asp236, instead of the Glu235 observed in MaL. Most basidiomycete laccases, such as *T. versicolor*, *L. tigrinus* and *C. cinereus* laccases, have Asp residues here. In the crystal structure of the basidiomycete *T. hirsuta* laccase (PDB code 3FPX), the corresponding residue is an Asn, and it has been suggested that this contributes to the high catalytic constants of *T. hirsuta* [18]. However, the purified laccase from *T. hirsuta* (ThL) (UniProt Knowledgebase accession number Q02497) used in our experiments has an Asp here [45].

In order to understand the oxidation of phenolic compounds in the binding pockets of laccases, the kinetic behaviour of TaLcc1, rMaL and ThL on three phenolic compounds [2,6-dimethoxyphenol (2,6-DMP), syringic acid, and methyl syringate] was studied (Table 1). The dimethoxy phenolic substrates have different para-substituents and different redox potentials. On the basis of our crystal structure of rMaL with 2,6-DMP [46], and the *T. versicolor* laccase (TvL) complex structure with 2,5-xylidine [11], the para-substituents would point out from the binding pocket and therefore not affect the substrate binding. The rate of laccase-catalysed reactions is thought to increase as the redox potential difference ($\Delta E^\circ$) between the T1 copper and the substrate increases. In TaLcc1, the redox potential of the T1 copper is slightly higher (0.51 V) than that in rMaL (0.48 V), but not as high as in ThL (0.78 V); thus, it would be expected that the kinetic data for TaLcc1 would fit in between the data of rMaL and ThL. However, our kinetic data clearly show that this is not the case, suggesting that the redox potential difference is not the only factor contributing to the rate of substrate oxidation (Table 1).

The kinetics of substrate oxidation by laccases has also been shown to be pH-dependent [47]. At higher pH values, phenolic substrates have lower $E^\circ$ values, whereas $E^\circ$ for the T1 copper of laccases seems to be unaffected by varying pH [28]. As consequence, when $\Delta E^\circ$ increases at the higher pH, the reaction rate is increased, but the inhibitory effect of hydroxide also increases. A typical feature of basidiomycete laccases is
their acidic pH optima, whereas asco-laccases generally work in a more neutral range with phenolic compounds [48,49]. Therefore, kinetic studies were carried out at pH 4.5, which is more optimal for ThL, and at pH 6.0, which is, in general, more optimal for TaLcc1 and rMaL (Fig. S1). Because both the pH dependence and the difference in redox potential affect the kinetics of the laccases, we concluded that rMaL and TaLcc1 were able to oxidize syringic acid at pH 6.0, mainly owing to the favourable pH, whereas ThL could oxidize the same substrate, mainly because of the large difference in redox potential. However, the effect of the difference in redox potential outweighs the effect of pH for substrates with high $E^\circ$, such as methyl syringate. Oxidation of methyl syringate was only possible with the high redox potential ThL, whereas the kinetic parameters for this substrate could not be determined with TaLcc1 or rMaL at either pH value (Table 1).

Interestingly, TaLcc1 showed a lower $K_m$ for 2,6-DMP at pH 6.0 than at pH 4.5 (Table 1). Despite the small difference in redox potentials of the two asco-laccases and their very similar pH optimum profiles, the affinity of TaLcc1 for 2,6-DMP was lower than the affinity of rMaL for the same substrate at pH 4.5. The similar pH profiles and $\Delta E^\circ$ values for the two asco-laccases do not explain the three-fold increase in reaction rate of rMaL with syringic acid at pH 6 as compared with pH 4.5. These differences in kinetic behaviour between TaLcc1 and rMaL must therefore be attributable to the variations in several residues forming the binding pocket, most likely Asp236, Ala193 and Val428 observed in TaLcc1 instead of Glu235, Pro192 and Phe427 observed in rMaL. Our mutagenesis studies with MaL have demonstrated that Glu235 → Asp mutation of the catalytic residue clearly increases the $K_m$ value for phenolic substrates while not affecting the $k_{cat}$ value. Furthermore, both the $K_m$ and $k_{cat}$ values were clearly affected by the Glu235 → Thr mutation, suggesting the importance of the carboxylic group for the catalytic activity [46]. In addition, Phe427 in rMaL (Val428 in TaLcc1) might be involved in placing substrate molecules into the correct orientation for oxidation. In T. hirsuta, the loops forming the substrate-binding pocket are completely different, possibly accounting for the clear differences in reaction kinetics between basidiomycete laccases and asco-laccases.

**Dimerization**

In the TaLcc1 crystal structure, molecules A and B (and C and D) of the asymmetric unit form a weak dimer (Fig. 6). On the basis of calculations performed

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**Fig. 6.** (A) Cartoon representation of dimers of TaLcc1 (blue) and rMaL (green). 2,6-DMP ligands (purple) are presented as they are in the rMaL complex structure (PDB code 3FU8). A surface representation of TaLcc1 (light blue) shows a small central channel that provides access for the substrates. (B, C) The contact amino acids at the dimeric interface in TaLcc1 (B) and rMaL (C). The residues from molecule A are shown in purple, and those from molecule B in yellow. The hydrogen bonding residues (according to Protein Interfaces, Surfaces and Assemblies) have been labelled, as have the hydrophobic residues with the closest contacts.
with the Protein Interfaces, Surfaces and Assemblies service [50–52], the buried surface area for the weak dimers, AB and CD, is 3.2% (658 Å²) and 3.3% (667 Å²) of the total surface area, respectively. In this weak dimer, the loop areas surrounding the phenolic substrate-binding pockets are packed together (Fig. 6B). Similar dimerization has been reported in the crystal structure of MaL [33]. In MaL, one of the key residues for the dimeric interaction is Phe427, located at the edge of the substrate-binding pocket. This residue might be involved in the orientation or the docking of the substrate molecules. In MaL, the Phe residues from two molecules are packed face-to-face. In TaLcc1, the corresponding loop is longer, and the interacting residues are Ile427 and Val428 (Fig. 6B). As a consequence, the T1–T1 copper distance is slightly longer in TaLcc1 (28 Å) than in MaL (27 Å), and the surface contact area is also slightly smaller (667 Å²) than that on MaL (796 Å² for 2Q9O).

It is possible that the weak dimers of MaL and TaLcc1 are the so-called ‘transient dimers’, which exist in solution as a mixture of monomers and dimers in a concentration-dependent manner [53]. It is noteworthy that the dimeric composition in MaL and TaLcc1 is very similar, suggesting that the ability to form dimers may have functional meaning. Interestingly, the two substrate-binding sites are packed against each other, and there is a shared cavity in the interface of the dimer. This cavity is enclosed in the MaL structure, whereas in the TaLcc1 structure there is a clear solvent channel in the interface between protein molecules that provides free access of the substrate molecules to the binding sites (Fig. 6A). However, the cavity itself is more compact in TaLcc1 than in MaL, owing to the loop containing Ile427 and Val428. The other narrow solvent channel reported earlier for rMaL is also visible in TaLcc1 [46]. The observed dimer in the crystalline state favours smaller phenolic substrates without large substituents in the para-position. The docking simulations based on the crystal structure for TaLcc1 reveal that substrates with large groups at the para-position clash with the loop containing Ile427 and Leu428. The corresponding loop is shorter in MaL, and the cavity is also more spacious.

Industrial utilization

Laccases have wide reaction capabilities and possess great biotechnological potential, because of their broad substrate specificity. Laccases can be utilized in many industrial applications, including biopulping, textile dye bleaching, bioremediation, biological fuel cells, and sensors. The stability and activity over broad pH and temperature ranges are desired properties for industrial enzymes. With respect to industrial applications, the ascomycete fungal laccase TaLcc1 is an efficient enzyme, particularly in denim bleaching, even at high temperatures and at neutral pH [32].

In general, asco-laccases possess a wider optimal pH range than basidiomycete laccases; however, the catalytic ability of asco-laccases in less acidic conditions has not yet been fully clarified on the basis of the available laccase structures. It could be that the adaptation of slightly different methods for proton transfer in asco-laccases and basidiomycete laccases (and in bacterial laccases) is responsible for the differences in the pH optimum range of laccases. In addition, the trinuclear site in asco-laccases is more protected, owing to the C-terminal plug; this might reduce the effect of hydroxide inhibition. Both TaLcc1 and MaL are also rather thermostable as compared with many other laccases. The stabilization of both the N-termini and C-termini of TaLcc1 and MaL might be a reason for the higher thermal stability. The extended C-terminus of asco-laccases is buried inside the solvent channel, and the extended N-terminus is stabilized by an additional disulfide bridge. In addition, both termini interact with carbohydrates bound to the protein structure.

Asco-laccases typically have middle redox potentials (TaLcc1, 0.51 V), whereas many basidiomycete laccases have very high redox potentials (ThL, 0.78 V), resulting in enhanced oxidation power. In the future, rational design methods could be used for tuning the redox potential of asco-laccases, or to increase the stability and the optimal pH range of high redox potential basidiomycete laccases. However, more studies are needed to understand how the enzyme structure contributes to the industrially desired properties.

Experimental procedures

Purification

TaLcc1 was produced at Roal Oy (Rajamäki, Finland), with Tr. reesei as host [32]. The culture supernatant was concentrated, buffer-exchanged, applied to a weak anion exchange column (DEAE Sepharose FF) in 5 mM Tris/HCl buffer (pH 8.5), and eluted with a linear 0–100 mM sodium sulfate gradient. Active fractions were pooled on the basis of ABTS activity, and were concentrated and desalted (Vivaspin, MWCO 10 000 Da). Typically, the laccase was further purified with a high-resolution anion exchange column (Resource Q) pre-equilibrated in 5 mM Tris/HCl.
buffer (pH 8.5). The bound proteins were eluted with a linear sodium sulfate gradient (0–100 mM). Active fractions were eluted at sodium sulfate concentrations between 5 and 40 mM, and concentrated. Subsequently, the buffer was changed to Tris/HCl (20 mM, pH 7.2). The protein yield from the purification was 6%, and the purification factor was 1.5.

MaL was overproduced in *Tr. reesei*, and purified basically as described previously [54]. ThL, assigned with sodium sulfate, and 0.1 M cistron was always under 0.1 mm. The crystals were approximately 0.3–1 mm long and 0.15–0.5 mm wide, and the thickness of the crystals was 2292

**Crystal structure of a Thielavia arenaria laccase**

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**Crystallization**

TaLcc1 was crystallized at room temperature with the hanging drop vapour diffusion method. Two microlitres of protein solution at a concentration of 9 mg mL⁻¹ and 2 µL of crystallization solution were equilibrated against 500 µL of reservoir solution. Initial screens were made with Crystal Screen I by Hampton Research. Optimization of the molecular weight of poly(ethylene glycol) and its concentration, together with pH, led us to the final crystallization condition of 7.5% poly(ethylene glycol) 3350, 0.2 M ammonium sulfate, and 0.1 M sodium acetate (pH 4.4). The streak seeding method with an equilibration time of ~10 h was used to obtain better-quality crystals. The crystals grew as thin plates, which made them difficult to handle. The crystals were approximately 0.3–1 mm long and 0.15–0.5 mm wide, and the thickness of the crystals was always under 0.1 mm.

**Data collection and structure refinement**

Before data collection, TaLcc1 crystals were quickly soaked in a cryoprotectant solution containing the reservoir solution with 25% glycerol. The crystal was then picked up with nylon loops and flash-frozen in liquid nitrogen. Data were collected at 100 K with synchrotron radiation at the European Synchrotron Radiation Facility (Grenoble) on beamline ID14-1, using an ADSC Q210 charge-coupled device detector. The data were indexed and integrated in MOSFLM [56], and scaled to 2.5-Å resolution in SCALA [57] from CCP4 [58]. The structure factors were created with TRUNCATE [59] from CCP4. A summary of the processing statistics is given in Table 2. The rather high *R*_merge might be attributable to twinning or, additionally, to anisotropic diffraction patterns of the crystal.

The data were analysed with XTRIAGE from the PHENIX package [60]. The suggested lattice was monoclinic, but the highest possible lattice seemed to be orthorhombic. The multivariate *L*-tests strongly indicated that the data were twinned. The *Z*-score was 10.9, whereas the value should be under 3.5 for good-quality to reasonable-quality data. The twin operator for this case was (h, −k, −l), and the estimated twin fraction was about 0.3 (Table 2). In addition, we also noticed a rather large off-origin peak, indicating some pseudotranslational symmetry that is most likely involved in the twinning with noncrystallographic symmetry.

The structure was solved by molecular replacement with the rMaL structure (75% sequence identity) as a model. Molecular replacement was performed with Phaser [61] from the CCP4 package and the rMaL (PDB code 2QPO) coordinates as a model. Only in space group *P*2₁ were rotation and translation solutions found that showed reasonable crystal packing. The *R*_values for the first round of refinement were *R*_work = 26.7% and *R*_free = 34.9%. When a twin operator was included, *R*_value and *R*_free were decreased to 19.2% and 25.2%, respectively. Refinement of the model and twin fraction were carried out with PHENIX, and model building in COOT [62]. The final *R*_values after the refinement were *R*_work = 18.1% and *R*_free = 22.4%, and the twin fraction was refined to 0.36. Despite the twinning,

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a surprisingly good-quality and continuous electron density map was observed. Noncrystallographic symmetry restraints were used during the refinement, and we also tried to release them, but this resulted in high B-factors for atoms in molecules C and D. Validation was performed with SFCHECK [63] from the CCP4 package, with 92.9% of all residues being in the most favourable region of the Ramachandran plot (Table 2).

**Kinetic data**

Kinetic constants ($K_m$ and $V_{\text{max}}$) for rMaL, TaLcc1 and ThL were determined on 2,6-DMP, syringic acid, and methyl syringate, at both pH 4.5 and pH 6.0, in 25 mM succinate buffer and 40 mM Mes buffer, respectively (Table 1). Kinetic measurements were performed in microtitre plates with a Varioskan kinetic plate reader (Thermo Electron Corporation, Waltham, MA, USA). The reactions were started by addition of substrate, and the rate of substrate oxidation was measured by monitoring the change in absorbance over 5 min. All of the measurements were performed in duplicate, with eight substrate concentrations (0.008–1.333 mM for 2,6-DMP and syringic acid; 0.008–2.800 mM for methyl syringate). The kinetic parameters were obtained by curve-fitting analysis with GRAPHPAD PRISM by curve-fitting analysis with graphite surface.

**Acknowledgements**

We are grateful to M. Paloheimo from Roal Oy for providing us with samples of TaLcc1. The ESRF is thanked for provision of synchrotron facilities. We also thank the staff members at beamline ID14-1 for their skilled assistance, and B. Hillebrant-Chellaoui at VTT for skilful help with the TaLcc1 purification. Roal Oy and the Academy of Finland (Project 115085) supported the work at Joensuu, and part of the VTT work was carried out with financial support from the Marie Curie EU-project ‘Enzymatic tailoring of protein interactions and functionalities in food matrix’, PRO-ENZ (MEST-CT-2005-020924), and from the National Technology Agency of Finland, Tekes project 40522/02.

**References**


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**Supporting information**

The following supplementary material is available:

Fig. S1. The pH activity profiles of rMaL, TaL and ThL for 1.7 mM 2,6-DMP, syringic acid and methyl syringate.

This supplementary material can be found in the online version of this article.

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