Isolation and partial nucleotide sequence of the laccase gene from *Neurospora crassa*: Amino acid sequence homology of the protein to human ceruloplasmin

(multicopper oxidase/copper binding sites/genomic cloning/protein evolution)

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**ABSTRACT** The laccase (benzenediol:oxygen oxidoreductase, EC 1.10.3.2) gene from *Neurospora crassa* was cloned and part of its nucleotide sequence corresponding to the carboxy-terminal region of the protein has been determined. The gene was cloned by cDNA synthesis with a laccase-specific synthetic deoxyundecanucleotide as primer and poly(A) RNA isolated from cycloheximide-treated *N. crassa* cultures as template. Based on the nucleotide sequence of the cDNA obtained, a unique 21-mer was synthesized and used to screen a genomic DNA library from *N. crassa*. Five different positive clones were isolated and shown to share an overlapping DNA region with the same pattern of restriction sites. Sequence analysis of the common 1.36-kilobase *Sal I* fragment revealed an open reading frame of 726 nucleotides. The amino acid sequence deduced is in complete agreement with the primary structures of several tryptic peptides isolated previously from *N. crassa* laccase. The analyzed carboxy-terminal region of laccase exhibits a striking sequence homology to the carboxy-terminal region of the third homology unit of the multicopper oxidase ceruloplasmin and to a smaller extent, to the low molecular weight blue copper proteins plastocyanin and azurin. Based on amino acid sequence comparison between these proteins, putative copper ligands of *N. crassa* laccase are proposed. Moreover, these data further support the hypothesis that the small blue copper proteins and the multicopper oxides have evolved from the same ancestral gene.

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Laccase (benzenediol:oxygen oxidoreductase, EC 1.10.3.2) belongs to the multicopper oxidases containing three distinct Cu centers known as type 1 or blue, type 2 or normal, and type 3 or coupled binuclear (1). The enzyme catalyzes the four-electron reduction of molecular oxygen to water with the concomitant oxidation of organic substrates. Laccases have been isolated from different sources such as microorganisms (2–8) and plants (9–11). Physical characterization of the proteins indicates a great variety in molecular weight, carbohydrate content, and quaternary structure. With the exception of the laccase from *Agaricus bisporus* and the tetrameric *Podospora anserina* laccase 1, all native enzymes bind four copper ions per molecule, representing one type 1, one type 2, and one type 3 Cu center (12). In contrast, the other multicopper oxidases, ceruloplasmin and ascorbate oxidase, contain more than four copper ions per molecule (13–17). Hence, laccase provides the most simple system to study the active site structure and reactivity of multicopper oxidases.

*Neurospora crassa* laccase is excreted into the culture medium after induction with low protein amounts of synthesis inhibitors (18). The enzyme has a molecular mass of 64 kDa, with a carbohydrate content of 11–12% (4, 5). In an attempt to determine the primary structure of *N. crassa* laccase and to study its molecular mechanisms of biosynthesis and excretion, we set out to clone the structural gene. Here we describe the strategy used to clone this gene and present part of its nucleotide sequence. The amino acid sequence deduced is compared to that of other blue copper proteins, such as ceruloplasmin, plastocyanin, and azurin.

**MATERIALS AND METHODS**

**Organism and Conditions of Culture.** *N. crassa* wild-type strain FGSC 321 (Fungal Genetics Stock Center, Kansas City, KA) was grown and induced with 2.8 μM cycloheximide as described by Froehner and Eriksson (18).

**Protein Chemical Methods.** Extracellular *N. crassa* laccase was isolated essentially as described (5) from the culture medium collected 96 hr after induction. Apolaccase was prepared by dialysis of the native enzyme against 5% (vol/vol) formic acid at 4°C for at least 24 hr. The lyophilized metal-free protein was modified by S-carboxymethylation (19). Tryptic digestion of the carboxymethylated apolaccase was performed in 0.1 M ammonium bicarbonate (pH 8.5) with 2% (wt/wt) trypsin (treated with 1:1-tosylamido-2-phenyl-ethyl chloromethyl ketone; Worthington) at 37°C. After 1 hr, a second portion of trypsin was added and the mixture was further incubated for 3 hr at 37°C. Peptides were purified by HPLC on a LiChrosorb RP-18 column (0.46 × 25 cm). The buffer system used was buffer A/0.1% (wt/vol) trifluoroacetic acid, pH 1.9, and buffer B [same as A except with 60% (vol/vol) acetonitrile]. Chromatography was carried out using a linear gradient of buffer B (0.5% per min) at 20°C with a flow rate of 1 ml/min. Absorbance was recorded at 220 nm. The peptides obtained were further characterized by amino acid analysis (Durrum Chemical, Palo Alto, CA, D-500 amino acid analyzer) and by automated Edman degradations (Beckman Sequenator model 890-B, updated).

**Isolation of Poly(A) RNA.** Total cellular RNA was isolated from lyophilized mycelia harvested 48 hr after induction by the method of Lucas et al. (20). Poly(A) RNA was selected by oligo(dT)-cellulose chromatography according to Aviv and Leder (21).

**Synthesis of Oligodeoxynucleotides and Primer Extension.** Oligodeoxynucleotides were synthesized according to the phosphotriester method (22, 23). Dimers were used as starting products, and synthesis was carried out on a manual solid-phase DNA synthesizer (Bachemgentecl, Torrance, CA). Oligomers were finally deprotected and subsequently purified by gel filtration on Sephadex G-25 (Pharmacia), followed by gel electrophoresis (20% acrylamide/8 M urea) and chromatography on NACS 52 (Bethesda Research Laboratories).

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Abbreviation: kb, kilobase(s).

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cDNA was synthesized essentially as described by Nathans and Hogness (24) using a sequence-specific oligonucleotide as primer and poly(A) RNA as template. High molecular weight primer extension products were separated by chromatography on Sepharose S-300 (Pharmacia).

Screening of a Genomic DNA Library. A *N. crassa* genomic DNA library in the plasmid vector pRK9 (25) was provided by M. Schechtmant (Syracuse University). This library was used to transform *Escherichia coli* LE 392. Ampicillin-resistant transformants were grown on nitrocellulose filters (Schleicher & Schuell) as described by Hanahan and Meselson (26). Filter replicas were screened by colony hybridization (27), using a 5' labeled synthetic oligodeoxynucleotide as hybridization probe.

Recombination Plasmid DNA Isolation and Restriction Endonuclease Digestions. Plasmid DNA was isolated by the method of Birnboim and Doly (28). Restriction enzymes were purchased from Boehringer Mannheim or Anglian Biotechnology (Colchester, UK). Digestions were carried out using assay conditions recommended by the suppliers. The resulting DNA fragments were electrophoresed on agarose and transferred to nitrocellulose (Schleicher & Schuell) (29).

DNA Sequencing. DNA sequence analysis was carried out by either the chemical cleavage (30) or the chain-termination technique (31). Single-stranded templates for the chain-termination method were obtained by molecular cloning of isolated subfragments in the single-stranded bacteriophage vectors M13mp18 or M13mp19 (32). DNA sequences were analyzed with a computer program (33).

RESULTS AND DISCUSSION

Synthesis of Laccase-Specific cDNA by Primer Extension and Screening of a Genomic Neurospora DNA Library. Synthetic oligonucleotide primers provide a powerful tool for the efficient detection and transcription of specific mRNA species and thus have been often used to isolate cDNA (e.g., see refs. 34 and 35) or genomic DNA (e.g., see refs. 36 and 37) clones. To design a unique oligonucleotide suitable for priming laccase-specific cDNA synthesis, tryptic peptides were prepared from the carboxymethylated apoprotein. Purified fractions isolated by HPLC were subjected to sequence analysis (data not shown). From the tryptic decapeptide T3 with the amino acid sequence Ala-Phe-Asn-Asp-Cys-Asp-Ala-Trp-Arg-C (Fig. 1A), a unique 11-mer (Fig. 1C) coding for residues 5–8 within the peptide was deduced. This primer

1 2 3 4 5 6 7 8 9 10
N- Ala Phe Asn Asp Asp Cys Asp Ala Trp Arg - C
1' 2' 3' 4' 5' 6' 7' 8' 9' 10'

was selected over the other seven possibilities (Fig. 1B) based on the published codon usage in other *N. crassa* nuclear genes (25, 38, 39). Furthermore, possible mismatches were only of the G-U type, building a “wobble” base pair (40), which has been shown to approximate an A-U base pair in stabilizing efficiency (41). Poly(A) RNA isolated from cycloheximide-treated *N. crassa* cultures served as a template for 11-mer-primed cDNA synthesis. The RNA used has earlier been shown to contain laccase mRNA by immunological analysis of *in vitro* translation products. High molecular weight primer extension products were subjected to sequence analysis by the chemical cleavage technique (30) (Fig. 1D). The amino acid sequence deduced from the specifically primed cDNA (Fig. 1E) was in complete agreement with the partially known primary structure of *N. crassa* laccase from the priming site toward the amino terminus (Fig. 1A). This allowed the synthesis of a unique 21-mer (Fig. 1F), which was used as a hybridization probe to screen a partial *Sac3A* genomic DNA library from *N. crassa* (25). From ~6 x 10⁴ colonies, seven positive clones were isolated. Examination of their plasmid DNA by restriction mapping revealed two pairs of them (nos. 53 and 81, 102 and 113) to be identical. Although the five positive clones nos. 10, 53, 62, 101, and 102 differ in size and the pattern of restriction sites, they were shown to share an overlapping DNA region. *Sal I* restriction endonuclease digests of the five different positive clones are presented in Fig. 2. All clones contain a common 1.36-kilobase (kb) *Sal I* fragment, which strongly hybridizes with the 32P-labeled 21-mer. Hence, this fragment was structurally further characterized.

DNA Sequence Analysis. The 1.36-kb *Sal I* fragment common to all five different positive clones (Fig. 2) was subjected to DNA sequence analysis with the strategy outlined in Fig. 3. Both the chemical cleavage (30) and the chain-termination (31) techniques were used. The 1.36-kb *Sal I* fragment was found to contain an open reading frame of 726 nucleotides starting with the triplet GTA of one of the two *Sal I* restriction sites and ending with the termination signal TAG (Fig. 4), thus encoding the carboxyl-terminal part of a protein. The correct 21-mer (Fig. 1F) used to screen the genomic *N. crassa* DNA library could be localized readily. It is interesting to note that the designed 11-mer used for primer extension (Fig. 1C) obviously worked with one G-U mismatch. The amino acid sequence deduced from the coding region agrees completely with the primary structures of six previously isolated tryptic peptides (T₁–T₆) of *N. crassa* laccase (Fig. 4). Hence, these results clearly indicate that the deduced amino acid sequence represents part (40–50%) of the
primary structure of \textit{N. crassa} laccase. This conclusion is further supported by the data of a preliminary sequence analysis of laccase B from the white rot fungus \textit{P. versicolor} (42). Among three fragments characterized, the active site peptide Leu-His-Cys-Ile-Leu-Asp-Glu indicates clear homology to the \textit{N. crassa} laccase sequence Met-His-Cys-Ile-Leu-Trp. The alignment of the two peptides reveals four amino acid residues to be identical and two to be isofunctional (Leu and Met, Phe and Trp).

**Sequence Comparison of \textit{N. crassa} Laccase with Other Blue Copper Proteins.** In Fig. 5, part of the amino acid sequence of \textit{N. crassa} laccase is compared with homologous parts of human ceruloplasmin (43) and with the primary structures of plastoxygenase from \textit{P. nigra} var. \textit{italica} (44) and azurin from \textit{Pseudomonas aeruginosa} (45). Ceruloplasmin is so far the only multicopper oxidase with a completely known primary structure (43). The single polypeptide chain consists of 1046 amino acid residues and exhibits a 3-fold internal homology in amino acid sequence (46, 51). Six (or seven) copper ions are bound per molecule—namely, two type 1, one type 2, and three (or four) type 3 (13–15). Plastoxygenase and azurin are low molecular weight blue copper proteins binding only one type 1 copper ion per molecule (52, 53). Amino acid sequences of azurin and plastoxygenase from a wide variety of species have been determined (52, 53). Both plastoxygenase and azurin from \textit{P. aeruginosa} have been crystallized and their three-dimensional structures as well as their copper ligands are known (49, 50). Plastoxygenase and azurin have been shown to be divergently related (54), and several reports have suggested that the small blue copper proteins and the multicopper oxidases share a common ancestry (47, 55, 56). The amino acid sequence comparison shown in Fig. 5 further supports this hypothesis. The primary structure of \textit{N. crassa} laccase is highly homologous to the carboxyl-terminal part of ceruloplasmin (CpC) and to a smaller extent to the other two homology units (CpP, CpM) of ceruloplasmin and to plastoxygenase and azurin. In all six polypeptides compared, five amino acid residues are invariant (one Cys, two Gly, two His) and four are isostructural having a conserved aromatic (three Phe, Trp, or Tyr) or hydrophobic (one Leu or Met) character. Among the invariant ones, three represent ligands of the type 1 copper in plastoxygenase (His-37, Cys-84, and His-87) and azurin (His-46, Cys-112, and His-117). The fourth blue copper ligand in plastoxygenase and azurin is Met-92 and Met-121, respectively. The corresponding residues Met-690 and Met-1031 are found only in CpM and CpP and represent the putative fourth binding sites for the two type 1 copper ions of human ceruloplasmin (51). Surprisingly, however, \textit{N. crassa} laccase lacks a methionine residue at the homologous position. Hence, this finding implicates a type 1 copper coordination different from plastoxygenase and azurin, and possibly from ceruloplasmin. In this context, it is interesting to note that stellacyanin, a small blue copper protein isolated from \textit{Rhus vernicifera} (57) is also devoid of methionine. The nature of its fourth copper ligand is presently still a matter of debate (58).

As mentioned above, the highest overall sequence homology is found when \textit{N. crassa} laccase is compared with the carboxyl-terminal region of human ceruloplasmin. Thus, 114 amino acid residues of the third homology unit (CpC) of ceruloplasmin can be aligned with part of the primary structure of laccase to give 27 identical and 12 isostructural amino acid residues (Fig. 5). Most interesting, two short
stretches show extensive sequence homology. A computer analysis using the unitary matrix indicated maximal homology scores of 6 and 9 for the segments presented in Fig. 6 A and B, respectively (N. Takahashi and F. W. Putnam, personal communication). The remarkable similarity of these two regions suggests that they play an important role in the formation of the active site of multicopper oxidases. In particular, they could provide ligands to all three different types of copper in both enzymes. This suggestion is consistent with the observation of Raju (59) that 50% of the non-blue copper of the ceruloplasmin molecule resides in the carboxy-terminal cryptic peptide of 11 kDa. As pointed out above, at least three invariant type 1 copper binding residues are present within these two stretches. The fourth type 1 copper ligand in N. crassa laccase is currently not known (see above). It is conceivable, however, that the nearby Met-169 of N. crassa laccase might be involved in copper binding. The idea that also type 2 or type 3 copper ligands could be provided by the two homologous regions shown in Fig. 6 is based on earlier reports. The sequence containing the histidine cluster of ceruloplasmin (Fig. 6 A) has been shown to be homologous to that providing ligands to the non-blue copper site in Cu/Zn-superoxide dismutase (51). Furthermore, using the coordinates of the known three-dimensional structure of poplar plastocyanin (49), Rydén recently presented a three-dimensional model of the copper active site of human ceruloplasmin (47). The model involves the carboxy-terminal region (amino acid residues 932–1040; ref. 43) and proposes His-975, Cys-1021, His-1026, and Met-1031 as type 1; His-1022 and His-1028 as type 2; and His-978, His-980, His-982, and His-1020 as type 3 copper ligands. The amino acid sequence comparison of ceruloplasmin with laccase supports this model with respect to the type 1 and the type 3 copper ligands. However, no histidine residue is found in

![Fig. 5. Amino acid sequence comparison between part of N. crassa laccase (Lac) and homologous parts of other blue copper proteins. Cp, human ceruloplasmin (43); CpN, first homology unit residues 234–344; CpM, second homology unit residues 595–705; CpC, third homology unit residues 933–1046; Pc, poplar plastocyanin (44); Az, P. aeruginosa azurin (45). The three homology units CpN, CpM, and CpC of ceruloplasmin were aligned as described by Takahashi et al. (46). CpC and Pc were aligned according to Rydén (47) and Pc and Az were aligned according to Murata et al. (48) with some modifications maximizing the sequence homology of each primary structure to that of N. crassa laccase. The positions of the four copper ligands in Pc (49) and Az (50) are marked by triangles. The number on the left of each sequence identifies the position within the protein of the first amino acid residue shown. For Lac, the numbers refer to valine as 1 according to Fig. 4. Amino acid residues are abbreviated using the one-letter code. Dashes represent gaps introduced for alignments. Boxes relate amino acid residues of Lac with identical residues in CpN, CpM, CpC, Pc, and/or Az. Shaded areas, isofunctional residues (D and E; K and R; T and S; F, W, and Y; I, L, M, and V).]

![Fig. 6. Amino acid sequence comparison of two highly homologous regions (A and B) of N. crassa laccase (Lac) and the carboxyl-terminal homology unit (CpC) of human ceruloplasmin (46). Numbers on the left of each sequence identify the positions within the proteins of the first residues shown. For Lac, numbers refer to valine as 1, according to Fig. 4. Identical amino acid residues are boxed and isofunctional amino acid residues are shaded (see Fig. 5). Potential ligands to the three different types of copper are indicated by *1, *2, and *3, respectively. The assignment of the copper ligands for N. crassa laccase is based on that proposed earlier for CpC (47).]
laccase that is homologous to the suggested type 2 copper binding His-1028 of ceruloplasmin. Hence, our data implicate that either the structures of the type 2 site in ceruloplasmin and laccase are different or the homologous type 2 ligands are closer to the amino terminus than suggested. It is clear that the complete amino acid sequence of *N. crassa* laccase and those of other multicopper oxidases are required to fully define the amino acid residues involved in the binding of the active site copper.

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