Research Article

Isolation and Physicochemical Characterization of Laccase from *Ganoderma lucidum*-CDBT1 Isolated from Its Native Habitat in Nepal

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Abstract

At present, few organisms are known to and capable of naturally producing laccases and white rot fungi are one such group. In the present study, three fungal species, namely, *Ganoderma lucidum*-CDBT1, *Ganoderma japonicum*, and *Lentinula edodes*, isolated from their native habitat in Nepal were screened for laccase production, and *G. lucidum*-CDBT1 was found to express highest levels of enzyme (day 10 culture media showed 0.92 IU/mg total protein or 92 IU/mL laccase activity with ABTS as substrate). Lignin extract from rice straw was used in Olga medium for laccase production and isolation from *G. lucidum*-CDBT1. Presence of lignin (5 g/L) and copper sulfate (30 μM) in the media increased the extracellular laccase content by 111% and 114%, respectively. The laccase enzyme produced by *G. lucidum*-CDBT1 was fractionated by ammonium sulfate and purified by DEAE Sepharose anion exchange chromatography. The purified enzyme was found to have a molecular mass of 43 kDa and exhibit optimal activity at pH 5.0 and 30°C. The isolated laccase was thermally stable for up to 70°C for 1 h and exhibited broad pH stability. The kinetic constants, $K_m$, $V_{max}$, and $K_{cat}$, determined using 2,2'-azinobis-(3-ethylbenzothiazoline-6-sulfonic acid) as substrate were found to be 110 μM, 36 μmol/min/mg, and 246 min$^{-1}$, respectively. The isolated thermostable laccase will be used in future experiments for delignification process.

1. Introduction

Laccases (benzenediol: oxygen oxidoreductase, EC 1.10.3.2) are copper containing enzymes that catalyze one-electron oxidation of a wide variety of substrates such as diphenols, polyphenols, diamines, aromatic amines, and other electron rich compounds, using molecular O$_2$ [1]. They belong to the family of proteins that include ascorbate oxidase, ceruloplasmin, and bilirubin oxidase. These enzymes are believed to utilize a free radical-catalyzed reaction mechanism in which the substrate forms an unstable free radical that further undergoes nonenzymatic reactions including hydration and disproportionation reactions [2].

Laccases are distributed among some plants [3, 4], fungi [3], and bacteria [5, 6]. Literature review shows that laccases are widely distributed among the prokaryotes. For example, laccases from *Azospirillum lipoferum* [7], *Marinomonas mediterranea* [8], *Streptomyces griseus* [9], *E. coli* [10], *Bacillus subtilis* [11], and many more bacteria have been purified and characterized. The crystal structures for several bacterial laccases have also been published [12–14]. Laccases are recognized for their wide spread applications including ethanol production, food industry, dye bleaching, paper and pulp processing, and production of value added chemicals from lignin [12]. Other applications of laccases include their use in detection of catecholamine neurotransmitters (dopamine,
norepinephrine) and glucose dehydrogenase-coupled oxidation of morphine. Some fungal laccases have also been shown to detoxify fungal metabolites, for example, aflatoxin B1, and accordingly are useful in the field of food microbiology from the view point of food processing and preservation [15].

Among the fungi, the wood-rotting basidiomycetes produce different kinds of extracellular oxidoreductases including laccases, peroxidases, and oxidases that generate $\text{H}_2\text{O}_2$ [16]. The laccases from wood-rotting basidiomycetes are of significant interest as they are able to utilize a wide spectrum of carbon sources, including intermediates of lignin degradation, phenols, and heterocyclic compounds. Like bacterial laccases, the fungal enzymes are also involved in efficient degradation of lignin and various types of dyes such as azo, heterocyclic, reactive, and polymeric dyes [17–19]. Lignin is the most recalcitrant components in lignocellulosic biomass that encases the cellulose and hemicellulose, thus necessitating pretreatment of lignocellulosic biomass to liberate fermentable sugars to produce liquid biofuels. Given the lignin depolymerizing activity of laccases, they can serve as efficient biocatalysts for the pretreatment of lignocellulosic biomass to isolate fermentable sugars and other products [17–19].

Chemical pretreatment processes are available to release the fermentable sugars; however, they are expensive, difficult to operate, and environmentally unfriendly. Further, furan derivatives, weak acids, and phenolic compounds formed after lignin breakdown during chemical delignification have inhibitory effects on fermentation process [20, 21]. On the other hand, biological methods provide safer and economic alternatives to obtain fermentable sugars from lignocellulosic biomass. One such alternative is depolymerization of lignin with the aid of laccases. There are numerous fungal species that grow on wooden logs and most of them secrete laccases to overcome the lignin barrier and thus obtain the energy from cellulose and hemicelluloses [3, 17–19]. The mountainous regions of Nepal are a rich source of fungi growing on logs [22]. In this regard, the Central Department of Biotechnology at Tribhuvan University, Kathmandu, Nepal, has collected, developed seed cultures and archived a number of fungal species. We believe that these fungal species have adapted to their natural environment of Nepal and some of them may be efficient laccase producers. Accordingly, as part of this study, we have screened three fungal species to identify potent laccase producers, optimized enzyme production conditions, and purified and characterized a laccase enzyme from *G. lucidum*-CDBT1.

2. Materials and Methods

2.1. Materials. ABTS [2,2′-azino-bis(3-ethylbenothiazoline-6-sulfonic acid)] and guaiacol were purchased from Hi-media Pvt. Ltd., New Delhi, India. DEAE Sepharose was purchased from Sigma Chemical Co., St. Louis, MO, USA. Protein molecular weight markers were purchased from Genei Pvt. Ltd., Bangalore, India. All other reagents and chemicals used were of analytical grade available locally.

2.2. Fungal Species Cultures. Pure cultures of *Ganoderma lucidum*-CDBT1 (Figure 1), *Ganoderma japonicum*-CDBT2, and *Lentinula edodes*-CDBT3 (locally known as Shiitake) preserved at 4°C were obtained from Central Department of Biotechnology, Tribhuvan University, Nepal. They were maintained by subculturing them in potato dextrose agar (PDA) Petri-plates by incubating the cultures at 25°C for 72 h in the case of *Ganoderma* species and 120 h for *Lentinula* species. Regarding the fungal species used herein, although not fully characterized, we do believe that they are novel strains adapted to growing in high altitudes of Nepal.

2.3. Screening of Fungal Species for Laccase Production. PDA-agar plates supplemented with guaiacol (0.02%) (wt/wt), 1-naphthol (5 mM), and tannic acid (0.5%) (wt/wt) and inoculated with various fungal cultures were used for screening of laccase production by the selected fungal species. All cultures were incubated at 25°C. Laccase secretion was monitored by visual color change in the plates, due to oxidation of screening agents, for several days [28, 29]. Biotic and abiotic cultures were used as positive and native controls, respectively. A reddish brown color was formed when secreted laccases react with guaiacol, a deep purple color was formed when they react with 1-naphthol, and a brown color was formed when laccases react with tannic acid. Guaiacol was added to the

Figure 1: *Ganoderma lucidum*-CDBT1 fruiting bodies (courtesy of Central Department of Biotechnology): (a) ventral surface and (b) dorsal surface.
Laccase enzyme activity was 2.5. Laccase Enzyme Assay. The precipitated lignin was washed, vacuum filtered, and the pH to 3.0 and incubated overnight to precipitate lignin. In the second step, the liquid was further titrated with acid to drop the pH to 7.0 and vacuum filtered to remove silica. In the first step, the pH was decreased to 7.0, and vacuum filtered to separate from the solids. Lignin was isolated from the black liquor by two-step treatments. In the first step, the pH was adjusted to 1.15 mL with 0.1M of sodium acetate buffer, pH 5.0. Reaction was started by the addition of enzyme and oxidation of ABTS was monitored spectrophotometrically at 420 nm for 90 seconds using quartz cuvettes. The amount of enzyme required to oxidize 1 μmol ABTS per minute is considered equivalent to 1 unit of enzyme activity (ε420 = 36000 M⁻¹ cm⁻¹ and path length l = 1 cm) [31].

2.5. Laccase Enzyme Assay. Laccase enzyme activity was measured at room temperature using ABTS as substrate (1 mM solution prepared in 0.1M of sodium acetate buffer, pH 5.0). A typical reaction mixture consisted of 350 μL of enzyme preparation (isolated or purified and appropriately diluted) and 350 μL of 1M ABTS and the final volume was adjusted to 1.15 mL with 0.1M of sodium acetate buffer, pH 5.0. Reaction was started by the addition of enzyme and pH 7.0. Finally, the desalted enzyme factions were subjected to a DEAE Sepharose anion exchange column chromatography at pH 7.0 as described previously [25]. The protein fractions collected were analyzed for laccase activity, and the fractions with high laccase activity were pooled and subjected to SDS- and native-PAGE to assess its purity and determine its molecular mass.

2.7. Purification of Laccase. Laccase secreted into the culture media was purified in three different steps. First, the cultures were vacuum filtered using Whatman number 1 filter paper. Thus obtained supernatant was centrifuged at 10,000 xg for 10 min to remove any debris and the clear supernatant was further used for purification process [33]. Second, the supernatant was subjected to ammonium sulfate fractionation (40% to 80%). The precipitated protein was isolated by centrifugation (10,000 xg/10 min). The precipitates were reconstituted in 0.1M sodium acetate buffer, pH 5.0, and screened for the presence of laccase activity as described above. The fractions with high laccase activity were subjected to dialysis overnight against 0.1M sodium phosphate buffer pH 7.0. Finally, the desalted enzyme factions were subjected to a DEAE Sepharose anion exchange column chromatography at pH 7.0 as described previously [25]. The protein fractions collected were analyzed for laccase activity, and the fractions with high laccase activity were pooled and subjected to SDS- and native-PAGE to assess its purity and determine its molecular mass.

2.8. Electrophoresis. Molecular mass of laccase was determined by SDS-PAGE using standard molecular mass markers (Genei India Pvt. Ltd.) according to Laemmli’s method [34]. Polyacrylamide gels (6% stacking gel and 10% separating gel) loaded with denatured and reduced protein preparations were subjected to electrophoresis at 25 mA for 3 hours at room temperature. Proteins were visualized on PAGE gels using EZ-Visi Blue protein staining solution [5]. Native-PAGE was also as described above except the gels were run at 4°C; protein samples were not denatured or reduced, and SDS was omitted from the gels. The native-PAGE gels were stained with 5 mM ABTS solution and 1% guaiacol solutions for 15 min to visualize laccase bands on the gels.

2.9. Protein Quantitation. Protein content in various preparations used in this study were determined by Bradford method [35] using BSA as standard.

2.10. Physicochemical Characterization of Laccase. The pH and temperature optima for the purified laccase were determined as follows. Purified laccase enzyme activity was determined by varying the pH 2.0–8.0 or the temperature (20–80°C) of the reaction under standard assay conditions described above. To investigate the pH stability, the purified enzyme was preincubated in buffers of pH 2.0–8.0 at room
3.2. Optimization of Laccase Production Media: Effect of Lignin, Copper Sulfate, Temperature, and pH. Presence of lignin and copper sulfate is known to promote production/secretion of laccase. In this regard the culture media were supplemented with 5 g/L lignin (isolated from rice straw). This resulted in 111% increase in the secretion of laccase as compared to a control culture supplemented with glucose (Figure 3). The effect of copper sulfate on laccase production was determined by varying its concentration (10–50 μM) in the culture medium. Maximum laccase production (93 IU/mL on 10th day; a 114% increase as compared to the blank) was obtained at a concentration of 30 μM copper sulfate (Figure 3). G. lucidum-CDBT1 cultures were further subjected to temperature and pH optimization studies to determine the conditions required for the optimal secretion of laccase. The optimal temperature and pH needed to produce high levels of laccase were found to be 30°C (89 IU/mL on day 10) and pH 5.0 (92 IU/mL on day 10), respectively (Figure 3). Secretion of detectable levels of the enzyme activity begins as early as day 2 and reaches a maximum at around day 10. Enzyme activity decreased sharply as the pH increased from 5.0 towards the neutral range or the temperature increased towards 50°C (Figure 3).

3.3. Purification of Laccase. The culture filtrate obtained from culture media subjected to ammonium sulfate precipitation led to enrichment of laccase in the 70% ammonium sulfate fraction. The precipitate dissolved in sodium acetate buffer, pH 5.0, and dialyzed overnight was further purified by DEAE Sepharose anion exchange chromatography, Table 1. The column was developed using a linear gradient of 0 to 0.1M sodium chloride at pH 7.0. The peak fractions were pooled and analyzed by native- and denaturing-PAGE (Figure 4). The ion-exchange chromatography purified laccase protein pool showed the presence of a single protein band on SDS-PAGE gels, with a molecular weight of 43 kDa (Figure 4(a)). Native-PAGE gels stained with ABTS and guaiacol as substrates also indicated the presence of one isozyme of laccase (Figures 4(b) and 4(c)) in the purified fraction. The extracts, as judged by native-PAGE, showed the presence of 3 laccase isozymes and the purified enzyme corresponds to isozyme 3 (the major isoform produced).

3.4. Partial Characterization of Purified Laccase: pH and Temperature Optima and pH and Thermal Stability. The influence of pH and temperature on laccase activity was determined by varying pH of the reaction mixture from pH 2.0 to 8.0, while the influence of temperature on laccase activity was determined by varying the temperature of the reaction between 20 and 80°C and at pH 5.0 (pH optima). The pH

| Enzyme                                           | Total activity (U) | Protein (µg/mL) | Specific activity (U/µg) | Fold-purification |
|--------------------------------------------------|--------------------|-----------------|--------------------------|-------------------|
| Culture media-filtrate                           | 97.22              | 105.7           | 0.92                     | 1.0               |
| Ammonium sulfate fraction (after overnight dialysis) | 3115.74            | 35.9            | 86.85                    | 94.4              |
| DEAE-anion exchange chromatography                | 2662.04            | 13.3            | 201.00                   | 218.0             |

*Preparation of the culture media-filtrate containing laccase and its ammonium sulfate fractionation and DEAE-anion exchange chromatography to isolate laccase were as described in Materials and Methods. Laccase enzyme was monitored using 1 mM ABTS as substrate at room temperature (−25°C) and protein concentrations in various fractions were determined by Bradford assay described in Materials and Methods.

2.11. Data Analysis. Microsoft Excel and Prism GraphPad V 5.00 computer programs were used for data analysis. All values reported are average of triplicate experiments.

3. Results

3.1. Screening Fungal Species for Laccase Activities. Three white rot fungal species, namely, G. lucidum-CDBT1, G. japonicum-CDBT2, and L. edodes-CDBT3 (Shiitake), were screened for the secretion of laccase using PDA plates supplemented with 1-naphthol, tannic acid, and guaiacol [28, 29]. The oxidative polymerization of guaiacol forming reddish brown zones in the medium, oxidation of tannic acid to brown color [29], and oxidation of 1-naphthol to a deep purple complex [28] are a visual confirmation for the presence/secretion of laccase enzyme. In this test, all species tested, that is, G. lucidum-CDBT1, G. japonicum-CDBT2, and L. edodes-CDBT3 (Shiitake), gave visual colors described above indicating secretion of extracellular laccase into the culture media subjected to ammonium sulfate precipitation and DEAE-anion exchange chromatography to isolate laccase as described in Materials and Methods. Laccase enzyme was monitored using 1 mM ABTS as substrate at room temperature (−25°C) and protein concentrations in various fractions were determined by Bradford assay described in Materials and Methods.

3.2. Optimization of Laccase Production Media: Effect of Lignin, Copper Sulfate, Temperature, and pH. Presence of lignin and copper sulfate is known to promote production/secretion of laccase. In this regard the culture media were supplemented with 5 g/L lignin (isolated from rice straw). This resulted in 111% increase in the secretion of laccase as compared to a control culture supplemented with glucose (Figure 3). The effect of copper sulfate on laccase production was determined by varying its concentration (10–50 μM) in the culture medium. Maximum laccase production (93 IU/mL on 10th day; a 114% increase as compared to the blank) was obtained at a concentration of 30 μM copper sulfate (Figure 3). G. lucidum-CDBT1 cultures were further subjected to temperature and pH optimization studies to determine the conditions required for the optimal secretion of laccase. The optimal temperature and pH needed to produce high levels of laccase were found to be 30°C (89 IU/mL on day 10) and pH 5.0 (92 IU/mL on day 10), respectively (Figure 3). Secretion of detectable levels of the enzyme activity begins as early as day 2 and reaches a maximum at around day 10. Enzyme activity decreased sharply as the pH increased from 5.0 towards the neutral range or the temperature increased towards 50°C (Figure 3).

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and temperature optima for laccases were found to be pH 5.0 and 30°C, respectively (Figures 5(a) and 5(b)). Laccase was active in a wide range of pH values. At pH values above pH 5.0, the enzyme activity decreased gradually by 50% at pH 8 (Figure 5(a)). Laccase activity was relatively stable in the range of pH 3.0–7.0. The enzyme activity declined when the temperature was increased from 30 to 80°C (Figure 5(b)). Purified laccase was incubated in buffers of varying pH between pH 2.0 and 8.0 for up to 180 min to determine its pH stability. Similarly, the purified enzyme dissolved in 0.1 M sodium acetate, pH 5.0, was incubated at varying temperatures (40–80°C) for up to 180 min to determine its thermal stability. Laccase was found to be most stable at pH 5.0 and 30°C (Figures 5(c) and 5(d)). Laccase activity decreased significantly (93%) after 180 min at pH 8.0 and it was fairly stable between pH 3.0 and 7.0 for up to 180 min. While the enzyme was relatively stable below 50°C, activity decreased significantly when the temperature was 70°C or higher. In fact, laccase activity was completely lost within an hour at 80°C (Figures 5(c) and 5(d)).

3.5. Determination of Kinetic Parameters. The kinetic parameters ($K_m$, $V_{max}$, and $K_{cat}$) for the purified laccase enzyme were determined using varied concentrations of ABTS as substrate and they were found to be 110 μM, 36 μmol/min/mg, and 246 min$^{-1}$, respectively.

4. Discussion

Some of the most important factors that influence secretion of laccase into culture media by *Ganoderma* species include (i) pH and temperature for their growth, (ii) addition of suitable amounts of sugars, for example, glucose (carbon source),
(iii) addition of lignin (inducer), and (iv) an appropriate concentration of copper sulfate (inducer). The optimal pH conditions under which fungal laccases are secreted into culture media vary between pH 3.0 and 7.0. This also depends on the type of substrate used in the activity assays. For example, when ABTS is used as substrate, the pH optima are more in acidic range, pH 3.0–5.0 [1]. In our study, laccase secretion was found to be maximal at pH 5.0. Accordingly, all of the cultures used for the production of laccase from *G. lucidum*-CDBT1 in this study were performed at pH 5.0. Further, the enzyme was secreted into the culture media over a wide range of pH, suggesting versatility of the enzyme for applications in many biotechnological processes. The decrease in activity at higher pH is most probably due to the binding of a hydroxide anion to type 2 and 3 copper centers in laccase, which in turn inhibits the binding of oxygen to the copper centers and thus decreased laccase activity [35, 36].

*G. lucidum*-CDBT1 cultures used in this study secreted maximal laccase into the culture medium at 30°C. This has been supported by the fact that *G. lucidum* is a mesophilic fungus and others have shown that *G. lucidum* secretes maximal laccase at 25°C [25].

Addition of glucose to culture media has been shown to influence laccase synthesis [37]. The concentration of glucose...
used in these studies varied from 0.1 to 1% [38]. The *G. lucidum*-CDBT1 cultures used herein were optimized to grow in relatively low concentrations (0.4%) of glucose containing media, and under these conditions *G. lucidum*-CDBT1 culture produced relatively high levels of laccase, while high concentrations of sugars may satisfy the nutrient demands of *G. lucidum* for growth but do not necessarily promote optimal secretion of laccase. In certain species, for example, *Trametes pubescens*, laccase production was, in fact, inhibited by high concentrations of glucose [39]. Additionally, it has also been shown that high concentrations of glucose trigger synthesis of extracellular polysaccharides that will interfere with the extraction of laccase from culture media [40]. Given the above factors, 0.4% glucose was used while optimizing our cultures.

Addition of lignin to culture media has also been shown to enhance laccase secretion [41]. Our cultures were optimized to grow and secrete maximal amount of laccase into culture media at concentrations of 5 g/L lignin (0.5%). Such concentrations resulted in two-fold increase in laccase activity in cultures, as compared to the culture media containing only 0.4% glucose. The latter may be due to the synthesis of inducible laccase isozymes, which in turn is believed to be because of increased secondary metabolism. Sharma and associates [41] have also observed an increase in laccase yield in a *Ganoderma* sp., rckk-02, when cultured in the presence of increasing concentrations of lignin. Other compounds shown to increase laccase production in cultures include veratryl alcohol, syringic acid, and 2,5-xylidine [41–43]. We used peptone as a nitrogen source which is known to enhance laccase synthesis [38]. Due to the complex composition of peptones, they provide a wide range of benefits to the cells and cell performance. Copper ions (Cu²⁺) are part of the active site of laccases; accordingly, CuSO₄ is most frequently supplemented with growth media to enhance laccase production in fungi. Cu²⁺ ions are thus crucial for the synthesis of catalytically active laccase protein. Further, Cu²⁺ ions also induce the expression of certain genes including laccase gene [44]. The promoter region of the genes encoding for laccase contains various recognition sites specific for xenobiotics and heavy metals [3, 45]. It has been demonstrated that the *Pleurotus* laccase genes *poxc* and *poxalb* are transcriptionally induced by Cu²⁺ ions [45, 46]. Relatively high concentrations of Cu²⁺ have also been shown to suppress laccase production which could be due to activation of defense mechanisms [42]. In our experiments, the optimal concentration of Cu²⁺ required to express highest levels of laccases was 30 μM. This is consistent with other studies [45, 46]. In addition to Cu²⁺, distinct organic inducers with structural similarities or relationships with lignin are often used to induce the expression of laccase [47].

Many fungal laccase isozymes have been purified and their molecular properties have been summarized [3]. The laccase enzyme purified from *G. lucidum*-CDBT1 had a molecular mass of 43 kDa and exhibited *Kₘ*, *Vₘₐₓ*, and *Kₘ* values of 0.110 mM, 36 μmol/min/mg, and 246 min⁻¹, respectively. The *Kₘ* values for laccases reported previously ranged from 0.1 to 3.7 mM with ABTS as substrate [23]. The *Vₘₐₓ* and *Vₘₐₓ* values of recombinant laccase heterologously expressed in *Pichia pastoris* using ABTS substrate were found to be 0.521 mM and 19.65 μmol/min/mg, respectively [3, 23, 48]. *Kₘ* reported in this work is lower than most other reported values suggesting this enzyme has higher affinity towards nonphenolic substrate ABTS [47]. This is as expected as these enzymes originate from different organisms and their physicochemical characteristics have been shown to be different. The other physicochemical characteristics, for example, pH and temperature optima, of the laccase isozyme isolated from *G. lucidum*-CDBT1 were similar but not necessarily identical to those previously reported [25, 47]. The isolated enzyme was thermally stable (70°C for up to 1 h) and exhibited broad pH stability (pH 3.0–7.0).

Fungal species express a number of different laccase isozymes with molecular masses ranging from 24 to 80 kDa [40], and this variation could be attributed to the different ecological origins of the species or different culture conditions under which various isozymes were expressed and/or secreted. In particular, presence of the inducers in the media seems to play an important role in which laccase isozymes were secreted into the medium. For example, *P. pulmonarius* produced three laccase isozymes, two of which (Icl1 and Icc2) were constitutive and the 3rd is inducible (Icc3); Icc3 was detected only when the fungus was cultured in the presence of inducers [48]. This also appears to be true among various strains of *G. lucidum*, Table 2. In this study, native-PAGE analysis suggested the presence of three laccase isozymes in *G. lucidum*-CDBT1. As judged by native-PAGE two of the isozymes were minor components whereas the 3rd was a major inducible isozyme. The major isozyme has been successfully purified in this study. This enzyme has characteristics similar to the enzyme isolated by Ko and associates [25] although the molecular mass is different. Molecular weight of the laccase isozyme-3 (Figure 4), reported herein (43 kDa as judged by SDS-PAGE), is in agreement with the
Table 2: A comparison of the physicochemical characteristics of laccases isolated from several strains of *Ganoderma lucidum*.

| *G. lucidum* strain | Mol. mass (kDa) | pH optimum | *Kₘ* with ABTS as substrate (mM) | Country of origin | Reference |
|---------------------|-----------------|------------|----------------------------------|-------------------|-----------|
| Unknown             | 68              | 3.0        | 0.114                            | China             | [23]      |
| CBS229.93           | 62.5            | 5.0        | 0.107                            | Netherlands       | [24]      |
| Gala3               | 65–68           | 3.5        | 0.037                            | South Korea       | [25]      |
| IBL-05              | 38.3            | 5.0        | 0.047                            | India             | [26]      |
| MDU-7               | 24–66           | 5.0        | 0.026–0.029                      | India             | [27]      |
| CDBTI               | 43              | 5.0        | 0.110                            | Nepal             | This work |

Figure 5: pH and temperature optimum and pH and thermal stability of purified laccase. The experimental conditions were as described in Materials and Methods. Experiments described in (a) and (c) were performed at room temperature (~25°C). The experimental results depicted in (a) and (c) (pH optimum and pH stability) were performed using buffers of various pH specified, whereas the temperature optimum and thermal stability was performed at pH 5 (0.1 M sodium acetate, pH 5.0).
value reported by Murugesan and associates [49] based on SDS- and native-PAGE of crude extracts stained for enzyme activity. This study [49] did not undertake physicochemical characterization of the enzyme. The minor isoforms that we have observed in native-PAGE gels are being isolated using enrichment processes in an ongoing study. Ko and associates [25] have, in fact, reported the presence of three laccase isozymes in *G. lucidum* whose molecular masses ranged between 40 and 68 kDa [42]. The number of laccase isozymes secreted by other fungal species also varies. For example, *T. multicolor* is reported to secrete five isozymes [50] whereas *P. ostreatus* secretes eight different laccase isozymes in response to Cu$^{2+}$ ions [51]. The number of laccase isozymes expressed/produced by different strains of the same species was also different [23–27, 37]. In this regard, we have summarized some physicochemical characteristics of several laccases isolated from a number of different strains of *G. lucidum*, Table 2. Accordingly, it appears that the laccase isozyme 3 is isolated and purified in this study, although having similar enzyme characteristics, its molecular mass is different. Further molecular characterization, for example, sequence determination and its comparison to other known sequences, is required to call this a novel enzyme.

In conclusion, we have successfully cultured and optimized culture conditions, induced expression, and purified and characterized a laccase in *G. lucidum*-CDBT1 isolated from its native habitat in Nepal. The isozyme characterized from *G. lucidum*-CDBT1 appears to be different from those reported among various strains of *G. lucidum*. Identification of minor isoforms of laccases in *G. lucidum*-CDBT1 and various laccases present in other organisms screened in this study, that is, *G. japonicum* and *L. edodes*, and the use of these enzymes to depolymerize lignin to expose cellulose and hemicellulose fibers from lignocellulosic biomass for efficient hydrolysis and fermentation of sugars are being studied in our laboratory.

**Competing Interests**

The authors hereby declare that they have no commercial or other competing and/or conflicting interests with regard to the publication of this research paper.

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